PLANT CYTOGENETICS

Second Edition

Ram J. Singh



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Ram J. Singh, Ph.D.

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Cover figure courtesy of Professor Juan Orellana. The figure is a metaphase-I cell of hexaploid wheat cv. Chinese Spring: chromosomes of A genome are green, chromosomes of B genome are brown, and chromosomes of D genome are red.

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Dedication

To my wife and boys

Preface

Gregor Johann Mendel, a monk in the Augustinian monastery at Brün, Austria (now Brno, The Czech Republic), conceived and delivered the science of genetics to the scientific world in 1865. Unfortunately, the invaluable treasure was buried in libraries until 1900, when three scientists, Carl Correns (Germany), Hugo de Vries (The Netherlands), and Erich von Tschermak (Austria), independently, unearthed Mendel's laws. Three years later, Walter Sutton and Theodor Boveri (1903) proposed the chromosomal theory of inheritance, creating the science of cytogenetics. The pioneering cytogenetic investigations in maize, *Datura*, and *Drosophila* during the early 20th century showed that an individual has a fixed number of chromosomes. The genes are located on the chromosomes and are inherited precisely by mitosis and meiosis. Alteration in the genetic material induces mutations. Since its inception, cytogenetics has exploded in many disciplines, involving organisms ranging from viruses to mammals. The dawn of the 21st century entered into sequencing of the entire genome of the human and many crops.

Plant cytogenetics is progressing at an extremely rapid pace since the publication of the first edition of *Plant Cytogenetics* by CRC Press in 1993. The time is ripe to update this book (second edition) in an encyclopedic fashion that includes two new chapters on the mode of reproduction in plants and transgenic crops. The introduction flows directly into the stream of classical and modern cytological techniques after a brief exposure to Mendelism and major chronological discoveries in the science of genetics. The handling of plant chromosomes assembled in Chapter 2 has undergone monumental progress. Precise identification and nomenclature of plant chromosomes from aceto-carmine and Feulgen staining techniques have progressed to Giemsa banding, fluorescence in situ hybridization, and sorting and karyotyping of chromosomes by flow cytometry. Several protocols for determining nuclear DNA content by flow cytometry have been formulated and perfected. Cell division is a complex and mysterious phenomenon, and it has not been completely understood. Chapter 3 (cell division - mitosis and meiosis) and Chapter 4 (genetic control of meiosis) present mainly classical descriptions of mitosis and meiosis in the higher plants. The mode of reproduction (sexual, asexual, and sex chromosomes) in the higher plants is described in Chapter 5. Chapter 6 includes chromosome nomenclature based on kinetochore position on chromosomes, karyotyping by aceto-carmine, and Giemsa C- and N-banding methods. Chromosome numbers of some economically important plants are presented in one of the appendices (6-I.). Chapter 7 discusses chromosomal aberrations (structural and numerical chromosome changes). Utilization of primary, secondary, and tertiary gene pools for crop improvement depends upon the comprehension of genomic relationships between the cultigen and allied species and genera. Genomic relationship established based on cytogenetics and verified by biochemical and molecular methods is presented in Chapter 8. The cause of morphological aberrations in transformants from cell and tissue culture is described in Chapter 9. The dazzling progress achieved since 1986 in producing genetically modified organisms (GMO) by private and public institutions is briefly elaborated in Chapter 10. I have provided an adequate literature citation; it covers publications from ancient to the present. The effectiveness of the second edition was improved by including an extensive glossary. The appendices include recipes for rescuing immature embryos and karyotyping by flow cytometry and the current linkage map of barley.

I am optimistic that this book will be accepted by plant cytogeneticists and will encourage students to receive training in the exciting field of plant cytogenetics.

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Ram J. Singh

The Author

Ram J. Singh, Ph.D., is an Agronomist-Plant Cytogeneticist in the Department of Crop Sciences at the University of Illinois at Urbana-Champaign. Dr. Singh received his Ph.D. in Plant Cytogenetics under the guidance of the late Professor Takumi Tsuchiya from Colorado State University, Fort Collins, Colorado. Dr. Singh isolated monotelotrisomics and acrotrisomics in barley, identified them by Giemsa C- and N-banding techniques, and determined chromosome arm-linkage group relationships.

Dr. Singh conceived, planned, and conducted an excellent quality pioneering research related to many cytogenetic problems in barley, rice, rye, wheat, and soybean, and published in highly reputable national and international prestigious journals, including *American Journal of Botany, Caryologia, Chromosoma, Crop Science, Genetics, Genome, International Journal of Plant Sciences, Journal of Heredity, Plant Breeding, and Theoretical and Applied Genetics.* In addition, he summarized his research results by writing several book chapters. Dr. Singh has presented research findings as an invited speaker at national and international meetings. Recently, he agreed to edit a series entitled "Genetic Resources, Chromosome Engineering, and Crop Improvement."

Dr. Singh produced fertile plants with 2n = 40, 41, or 42 chromosomes, for the first time, from an intersubgeneric cross of the soybean and *Glycine tomentella* (2n = 78). The screening of derived lines shows that the genes for resistance to the soybean cyst nematode (SCN) have been transferred from *G. tomentella* to the soybean. He constructed a soybean chromosome map, for the first time, based on pachytene chromosome analysis, and this innovative research laid the foundation for creating a global soybean map. By using fluorescent *in situ* hybridization, he confirmed the tetraploid origin of the soybean and species of the genus *Glycine*.

Dr. Singh is a member of the Crop Science Society of America and the American Society of Agronomy. He has written *Plant Cytogenetics*, 1st ed., nine chapters in books, 70 refereed papers, 45 abstracts, and 33 articles in newsletters. He received an Academic Professional Award for Excellence: Innovative and Creativity.

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CHAPTER 1

Introduction

The foundation of the science of genetics was laid in 1865 (published in 1866) by Gregor Mendel, when he reported principles of segregation and an independent assortment based on his careful selection of seven contrasting traits and controlled artificial hybridization experiments with the garden pea (*Pisum sativum* L.). Unfortunately, the importance of Mendel's classic paper was not widely known or clearly appreciated until 1900, when three botanists, Carl Correns, Hugo deVries, and Erich vonTschermak, rediscovered Mendel's laws of inheritance as a result of their own investigations (Peters, 1959).

Mendel formulated two basic principles of heredity from his experiments with the garden pea: (1) the law of segregation and (2) the law of independent assortment. He selected pure-line parent varieties differing from each other by clear-cut contrasting traits, which helped him to classify the F_1 (first filial) generation and to score segregating progenies accurately in F_2 (second filial) and later generations. Although Mendel was unaware of the gene–chromosome relationship, his method of experimentation, keen observations, and statistical approach to analysis of data led him to discover and develop basic principles of heredity.

Mendel's first law relates to members of a single gene pair. In diploid species, factors (genes) occur in pairs, one member contributed from each of the parents. In hybrid individuals, in which the two members of a gene pair differ, there is no blending, but rather each of the two differing gene forms is transmitted unchanged to progeny. In the formation of sex cells (gametes), later shown to occur as a result of meiosis (followed by gametophyte generations in higher plants), members of factor pairs typically behave as if they were discrete particles that separate (segregate) clearly and are delivered in equal numbers to gametes. He postulated that when contrasting members of factor pairs are delivered in equal numbers to gametes and then are combined randomly at fertilization in zygotes in an F_2 generation, the progeny segregate (or may be separated into) the two grand parental types in a ratio of three-fourths dominant trait expression: one-fourth recessive trait expression. Mendel's knowledge of probability, his use of contrasting traits, and a clear testing of the genetic constitutions of parents, F_1 and F_2 generations enabled him to formulate the *law of segregation*.

Mendel also made crosses in reciprocal directions in which parents differ by two unrelated traits (two pairs of genes \rightarrow dihybrid). The four phenotypic categories occur in nearly equal frequencies in test crosses. That is, from the dihybrid parent, round (*R*) and wrinkled (*r*) seed traits are transmitted in equal frequencies, as are yellow (*Y*) and green (*y*) seed traits (demonstrating the law of segregation for both the *Rr* and *Yy* gene pairs). In addition, *RrYy* individuals produce four gametic genotypes in equal proportions, and in F₂, segregate in a phenotypic ratio of 9 (round, yellow; *R*-*Y*-):3 (wrinkled, yellow; *rrY*-):3 (round, green; *R*-*yy*):1 (wrinkled, green; *rryy*). This proves that during meiosis, copies of *R* and *r* are assorted (distributed) to the meiotic products (and eventually to gametes) independently of copies of *Y* and *y*. This is the law of *independent assortment*.

Mendel's laws are not universal when nonallelic genes are linked [usually by virtue of being located on the same chromosomes but less than 50 centiMorgans (cM is a map unit) apart]; segregation still occurs, but assortment is not independent. The frequencies of gametic genotypes can be estimated by observing phenotypes of zygotes either from F_2 or testcross populations. The closer the distance between the genes, the stronger the linkage. The absence of crossover products (also known as recombinants) suggests complete linkage. The strength of linkage expresses itself primarily in the production of gametes, and that depends upon the distance between two genes on the same chromosome. Discovery and exploitation of linkage and crossing over led to detailed chromosome mapping in numerous organisms, an activity that continues to the present day. In many species, linkage groups have now been assigned to each of the individual chromosomes of the genomes, especially through the use of chromosomal aberrations.

Shortly after the rediscovery of Mendel's laws of inheritance, Sutton and Boveri (1903), independently, reported that the hereditary factors (now known as genes) are located in the chromosomes (Sutton-Boveri Hypothesis; Chromosomal Theory of Inheritance), and cytogenetics was born (see Peters, 1959). Cytogenetics flourished once it was established that chromosomes are the vehicles of genes. Classical cytogenetic studies in plants and animals quickly led to numerous extensions of genetic information and interpretation. Cytogenetic studies have made landmark contributions to knowledge in biology. The combined discipline has been especially powerful in explaining hereditary phenomena that would have been extremely difficult to decipher by alternative approaches. In spite of a proliferation of powerful new disciplines such as molecular biology, and an ever-increasing level of complexity and performance of new apparatus, there remains a need for well-trained cytogeneticists to address questions that are most readily and economically resolved by cytogenetic approaches.

Cytogenetics is a hybrid science that combines cytology (the study of chromosomes and other cell components) and genetics (the study of inheritance). The science includes handling chromosomes (chromosome staining techniques), function and movement of chromosomes (cell division; mitosis, meiosis), numbers and structure of chromosomes (karyotype analysis), and numerous modifications of structure and behavior as they relate to recombination, transmission, and expression of genes. Succeeding chapters will summarize and document much of the cytogenetic information in higher plants that has been reported in recent decades. It is hoped that this information and literature citations will prove useful to practicing cytologists and cytogeneticists, as well as to members of other disciplines.

Year	Discoverer	Discovery
		1. Pre-Mendelian Period
300 B.C.	Euclid	Discovery of optical properties of curved surfaces
~1285	Degli	Invention of spectacles
1590	Janssen (Zachary and Francis, brothers)	Discovery of operational compound microscope
1610	Galileo	Construction of first microscope by combining lenses in a tube of lead
1625	Faber	Application of microscope for anatomical studies
1651	Harvey	Development of concept that life originates from eggs and semen; development of theory of epigenesis
1661	Malpighi	Discovery of capillaries in the lungs of animals
1665	Hooke	Discovery of cell by examining the texture of cork by magnifying lenses
1672	de Graafe	Discovery of follicles in the human ovary; incorrectly identified them as eggs
1677	Leeuwenhoek	Discovery of sperm of some animals but mistook them for "wild animalcules"
1679	Swammerdam	Development of preformation theory based on studies of the development of insects; a simple enlargement from a minute but preformed animal to adult

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Year	Discoverer	Discovery	
1682	Grew	The first explicit mention of the stamens as the male organs of the flower	
1694	Camerarius	Description of sexual reproduction in plants	
1707	Linnaeus	Father of plant taxonomy	
1761	Kölreuter	Production of hybrids in <i>Nicotiana, Dianthus, Mathiola</i> , and <i>Hyoscyamus</i> and equal hereditary contribution of both parents	
1809	Lamarck	Development of the theory of acquired inheritance known "Lamarckism" or "Lamarckianism"	
1824	Amici	Discovery of pollen tube	
1828	Brown	Discovery of cell nucleus in the flowering plant Tradescantia	
1835	Mohl	Creator of modern plant cytology (cell division and importance of protoplasm)	
1838	Schleiden	Founder of cell theory (all plant tissues are composed of cells)	
1839	Schwann	Establishment of the cell theory in definite form; father of modern cytology	
1841	Remak	The phenomenon of amitosis or direct division	
1842	Nägeli	Publication of the first drawings of chromosomes	
1845	von Siebold	Protozoa are animals consisting of a single cell	
1848	1848 Hofmeister Illustration of tetrad formation, identification of two guart the opposite poles of the embryo sac but failure to between the synergids and the egg		
1858	Darwin and Wallace	Theory of evolution through natural selection	
1859	Darwin	The origin of species	
		2. Post-Mendelian Period	
1865	Mendel	Discovery of the laws of heredity (segregation and independent assortment)	
1866	Darwin	Theory of pangenesis (mechanism of heredity)	
1871	Miescher	Isolation of nucleic acid and nucleoprotein	
1872	Hertwig	Fertilization is the union of sperm and egg nuclei (general statement)	
1879	Flemming	Discovery of mitosis	
1879	Strasburger	Description of megasporogenesis in plants	
1879	Schleicher	Indirect division is called karyokinesis	
1880	Hanstein	Coined the term protoplast; followed the sequence of early cell divisions in the development of the embryos	
1881	Focke	Xenia; the pollen produced a visible influence on the heredital characters of those parts of the ovule that surround the embr	
1882	Flemming	Discovery of tiny threads inside salamander, later known as chromosomes; mitosis and chromatin	
1884	Strasburger	Discovery of actual process of syngamy	
1890	Waldeyer	Fundamental fact in mitosis is the formation of the nuclear filaments named chromosomes	
1892	Hertwig	Protoplasm (proto, first and plasma formation) theory	
1896	Wilson	Publication of <i>The Cell in Development and Inheritance</i> (1st ed.)	
1898	Nawaschin	Discovery of double fertilization in angiosperm	
1900	deVries, Tschermak, Correns	Rediscovery of Mendel's laws (independently)	
1900	Juel	Parthenogenesis	
1901	McClung	Sex determination is related to the chromosomes	
1901	deVries	The mutation theory	
1902–1903	Sutton, Boveri	Chromosome theory of inheritance (independently)	
1905	Johannsen	Coined the terms gene, genotype, and phenotype	
1905	Wilson	Coined the term X-chromosomes	
1905–1908 1906	Bateson and Punnett Montgomery	Discovery of the first linkage Distinction made between heterochromosomes (X and Y) and	

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Year	Discoverer	Discovery
1907	Bateson	Coined the terms genetics (1906), allelomorph (= allele), F_1 and
		F_2 (1902), homozygote and heterozygote (1902), epistasis (1907)
1909	Jenssens	Chiasma type theory of crossing over
1910	Morgan and colleagues	Sex linkage and genes are located on chromosomes
1912	Morgan	Linkage map of Drosophila
1912	Baur	Sex-linked gene in Silene alba (= Melandrium album)
1913	Sturtevant	Linear arrangement of genes on Drosophila chromosomes
1913–1916	Bridges	Nondisjunction as a proof of the chromosome theory of inheritance
1913, 1917	Carothers	Correlation of Mendelian segregation and the distribution of heteromorphic homologues in meiosis
1917,1919, 1920	Bridges	Recognition of deficiencies, duplications, and translocations by genetic tests
1921	Blakeslee	Discovery of trisomics in Datura stramonium
1921	Sturtevant	Recognition of an inversion by genetic test
1922	L.V. Morgan	Attached X-chromosomes in Drosophila
1924	Feulgen and Rossenbeck	Specific stain for DNA (Feulgen stain)
1925	Morgan	Publication of The Theory of the Gene
1926	Kihara and Ono	Introduction of terms auto- and allopolyploidy
1927	Longley	Discovery of B-chromosomes in maize
1927	Muller	X-ray causes gene mutation
1928	Heitz	Distinction between euchromatin and heterochromatin
1928	Griffith	Transformation (a harmless strain of bacteria into a virulent one)
1928	Gowen	Discovery of a gene that affects meiosis
1929	Darlington	Coined the term chiasma terminalization
1930	Kihara	Establishment of genome analysis
1931	Creighton and McClintock	Cytological proof of crossing over in maize
1931	Stern	Cytological proof of crossing over in Drosophila
1931	Belling	Classical model of crossing over
1932	Darlington	Publication of Recent Advances in Cytology (1st ed.)
1933	Heitz and Bauer	Discovery of the giant chromosomes in dipteran insects
1933	Painter	Discovery of salivary gland (polytene) chromosomes in Drosophila melanogaster
1934	McClintock	Recognition of nucleolus organizer in maize
1936	Caspersson	Development of ultraviolet photomicrography for the study of nucleic acid within nucleus
1936	Stern	Discovery of mitotic crossing over
1936	Sturtevant	Preferential segregation in Drosophila
1936	Sturtevant and Beadle	Relation of inversions to crossing over and disjunction
1937	Blakeslee and Avery	Chromosome doubling by colchicine
1937	Sax	Beginning of x-ray cytology and cytogenetics
1939	Thom and Steinberg	Induction of mutations by nitrous acid
1941	Beadle and Tatum	Development of one gene-one enzyme concept
1944	Avery, MacLeod, McCarty	DNA is the hereditary material, not protein
1946	Lederberg and Tatum	Genetic exchange in bacteria
1947	Auerbach and Robson	Induction of mutations by nitrogen and sulfur mustard
1949	Delbruck and Bailey	Genetic exchange in bacteriophage
1949	Barr and Bertram	Discovery of small stainable body in nondividing nuclei of female and absent in males-sex chromatin or Barr body
1950	Levan and Tjio	Use of 8-hydroxyquinoline for pretreatment of chromosomes
1950	McClintock	Discovery of activator-dissociation system in maize
1951	Horowitz and Leupold	One gene-one enzyme hypothesis
1952	Chargaff	Discovery of A-T and G-C ratios in DNA
1952	Franklin	X-ray diffraction data of DNA
1953	Watson, Crick	Announcement of the double-helix structure of DNA
1955	Benzer	Fine structure of a genetic region in bacteriophage

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Year	Discoverer	Discovery
1956	Tjio and Levan	Determination of accurate chromosome number $2n = 46$ of humans
1960	Barski	Somatic cell fusion
1962	Ris and Plaut	Discovery of chloroplast chromosomes
1963	Nass and Nass	Discovery of mitochondrial chromosomes
1963	Marmur	DNA/DNA hybridization in vitro
1964	Britten and Waring	Discovery of repetitious DNA in eukaryotes
1965	Du Praw	Discovery of whole mount electron microscopy
1968	Arber, Smith and Nathan	Restriction and modification enzymes
1968–1970	Caspersson and colleagues	Development of fluorescent and Giemsa staining for eukaryotic chromosomes
1969	Jonathan and colleagues	Isolation of the first gene
1970	Pardue and Gall	DNA/DNA and RNA/RNA hybridization in situ
1972	Khorana and colleagues	Synthesis of artificial gene
1972	Boyer's research group (Berg and colleagues)	Production of recombinant DNA
1974	Thomas Jr. and Wilson	Discovery of the widespread occurrence of the hair-pin-like structures (palindromes) resulting from inverted repetitious DNA, located at intervals along the chromatids of eukaryotic chromosomes
	3. G	enetic Engineering Period
1973	Cohen, Boyer and colleagues	Insertion of a gene; DNA cloning
1976	Genentech	Establishment of the first genetic engineering company
1977–1978	Sanger and Coulson; Maxam and Gilbert	Rapid sequencing of DNA
1983	Gusella and colleagues	Location of Huntington's disease on human chromosome 4
1984	Jeffreys	Development of genetic fingerprinting
1986	Initiation of the Human Genome Project	
1987	Production of the first comprehensive human genetic map	
1988 1990	Creation of the National Center for Human Genome Research The launch of the Human Genome Project; the first gene therapy on a 4-year-old girl with an immune disorder known as ADA deficiency, but procedure did not work; Mary-Clair King demonstrated that a gene on chromosome 17 causes an inherited form of breast cancer and increases the risk of ovarian cancer	
1992		an genetic map by international collaboration
1995	The Institute for Genomic Research	
1995	Publication of the first complete DNA sequence of the genome of a free-living organism — the bacterium <i>Haemophilus influenzae</i>	
1995	Monsanto Company	Development of Roundup Ready soybeans
1996	An international collaboration involving 600 scientists in Europe, North America, and Japan	The first complete sequence of the genome of a eukaryote (the yeast <i>Saccharomyces cerevisiae</i>)
1996	Transgenic maize, potatoes, and cotton expressing BT toxins grown in the United States on a large scale	
1998	The first genome of a multicellular organism — the 97 megabase DNA sequence of the round worm	
2000	Celera Genomics	The first draft of the sequence of 90% of the human genome
2000	The Institute of Genomic Research	Complete Arabidopsis genome map
2001	Syngenta and Myriad Genetics	Complete Rice Genome map
2002	Syngenta	Complete Rice Genome — Japonica rice
2002	Beijing Genomic Institute	Complete Rice Genome — Indica rice

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Sources: De Robertis et al., 1954; Elliott, 1958; Lander and Weinberg, 2000; Maheshwari, 1950; Peters, 1959; Schultz-Schaeffer, 1980; Swanson, Merz, and Young, 1981; Singh, 1993; and many scientific journals.

CHAPTER 2

The Handling of Plant Chromosomes

I. INTRODUCTION

Cytologists have devised cytological techniques from time to time to obtain precise information on chromosome numbers and chromosome structures and to examine the mechanism of cell division in plant species. These properties are studied by cytological techniques. The basic principles for handling the mitotic and meiotic chromosomes of all plant species are similar and consist of collection of specimens, fixation, and staining. However, cytological procedures are modified depending upon crop species, objective of the experiments, and, above all, personal preference of the cytologists (Darlington and La Cour, 1969; Sharma and Sharma, 1965).

II. SQUASH TECHNIQUE — MITOTIC AND MEIOTIC CHROMOSOMES

A. Collection of Roots

Actively growing roots can be collected from germinating seeds in a Petri dish or from seedlings growing in pots or in the field. Generally, roots are collected in the laboratory after germinating the seeds in a Petri dish.

Place the seeds on a moistened filter paper in a Petri dish. For the analysis of cereal chromosomes (e.g., wheat, oat, barley, and rye), keep the Petri dish with seeds in a dark cold room or refrigerator (0 to 4° C) for 3 to 5 days. The cold treatment facilitates uniform and rapid seed germination. Remove the Petri dish with seeds from the refrigerator and leave at room temperature (20 to 25° C) for germination. For soybean chromosome analysis, germinating seeds are kept at 30° C for 4 h before collecting the roots. This is not required when roots are collected from a sand bench or vermiculite in the greenhouse.

Collect roots that are 1 to 2 cm long. It should be kept in mind that roots should touch the filter paper, otherwise the mitotic index will be very low. When it is desired to collect roots from pots, care should be taken not to break the actively growing roots. Transfer roots to vials containing cold water after washing soil from the roots. Keep the vials in ice-cold water in an ice chest.

B. Pretreatment of Roots

Pretreatment of roots is an essential step for the study of somatic chromosomes. It performs several purposes: stops the formation of spindles, increases the number of metaphase cells by arresting the chromosomes at the metaphase plate, contracts the chromosome length with distinct constrictions, and increases the viscosity of the cytoplasm. Numerous pretreatment agents, described below, have been developed.

1. Ice-Cold Water

Pretreatment with ice-cold water is effective and is widely used for cereal chromosomes. This particular pretreatment is preferred over other pretreatments when chromosomes of a large number of plants must be studied.

Keep roots in numbered vials two thirds filled with cold water. Transfer vials with roots into a container that will allow cold water to cover the top of the vials, and further cover the vials with a thick layer of ice (Tsuchiya, 1971a). Keep the container in a refrigerator for 12 to 24 h, depending on the materials and experimental objectives. The recommended pretreatment lengths for barley and wheat chromosomes are 16 to 18 and 24 h, respectively. Longer pretreatment will shorten the chromosome length considerably.

2. 8-Hydroxyquinoline

Tjio and Levan (1950) were the first to recognize the usefulness of 8-hydroxyquinoline for chromosome analysis. Treat roots with 0.002 M (0.5 g/L) aqueous solution [dissolve in double distilled water (ddH₂O) at room temperature (RT)] of 8-hydroxyquinoline for 3 to 5 h at 16 to 18°C. Warmer temperatures often cause sticky chromosomes. Pretreatment with 8-hydroxyquinoline has been effective for plants with small-size chromosomes. It makes primary (kinetochore) and secondary constrictions (nucleolus organizer region) very clear.

3. Colchicine

Because colchicine in higher concentrations induces polyploidy, a low concentration (0.1 to 0.5% for 1 to 2 h, at RT) is recommended for pretreatment. A treatment time of 1 1/2 h gives the best results for soybean chromosomes. This treatment schedule arrests chromosomes at the metaphase stage, giving a high proportion of cells with metaphase chromosomes, thus making chromosome count efficient. Roots should be washed thoroughly after colchicine pretreatment. This pretreatment facilitates better penetration of fixative at the subsequent stages of chromosome preparation. Because colchicine is a toxic chemical (carcinogenic and lethal poison), necessary precautions should be taken when it is used.

4. α-Bromonaphthalene

The effect of α -bromonaphthalene is almost the same as that of colchicine. It is sparingly water soluble. A saturated aqueous solution is used in pretreatment for 2 to 4 h at RT. This pretreatment is effective for wheat and barley chromosomes.

5. Paradichlorobenzene

Pretreatment with paradichlorobenzene (PDB) is effective for plants with small-size chromosomes. Like α -bromonaphthalene, it has a low solubility in water. Weigh 3 g of paradichlorobenzene, and add to it 200 mL distilled water. Incubate overnight at 60°C, then cool. Some crystals may remain undissolved. Shake thoroughly before using. Pretreat roots for 2 to 2 1/2 h at 15 to 20°C or at RT (Palmer and Heer, 1973).

C. Fixation

The science of chromosome study depends upon good fixative. The function of a fixative is to fix, or stop the cells at a desired stage of cell division without causing distortion, swelling, or shrinkage of the chromosomes. The most widely used fixatives described by Sharma and Sharma (1965) are listed below.

1. Carnoy's Solution 1

- 1 part glacial acetic acid
- 3 parts ethanol (95 to 100%)

Note: This fixative is prepared fresh each time and is used for the fixation of roots and anthers. For anthers, a trace (1 g/100 mL fixative) of ferric chloride (FeCl₂·6H₂O) is added to the fixative as a mordant, if aceto-carmine is used as a stain. The material should be kept in the fixative at least 24 h.

2. Carnoy's Solution 2

- 1 part glacial acetic acid
- · 3 parts chloroform
- 6 parts ethanol (95 to 100%)

A modification of 1:3:4 has also been used for wheat chromosomes.

3. Propionic Acid Alcohol Solution

- 1 part propionic acid
- 3 parts ethanol (95 to 100%)
- 1 g/100 mL fixative (for meiotic materials only) ferric chloride

This fixative is good for plants with small chromosomes. It provides clear cytoplasm and optimal staining for chromosomes (Swaminathan, Magoon, and Mehra, 1954). After 24 h of fixation, roots are transferred to staining solution. Inflorescence and flower buds are first washed with two changes of 70% ethanol. Materials to be stained for observations of meiosis should be stored in 70% ethanol under refrigeration (4 to 5°C). Materials can be stored for a long time at 20°C.

D. Staining of Chromosomes

Chromosome number and morphology and structure can be visualized through a microscope after optimal staining of the chromosomes. A good quality staining agent stains specifically the chromosomes, differentiates euchromatin and heterochromatin, and provides clear cytoplasm with stained nucleoli (but not with Feulgen stain). The following staining methods have been used extensively for staining plant chromosomes.

1. Aceto-Carmine Staining

a. Ingredients

For 1% stain:

- 1 g carmine powder (use a lot of carmine certified by the Biological Stain Commission)
- 45 mL glacial acetic acid
- 55 mL ddH₂O water

b. Preparation of Stain

- Heat to boil 100 mL 45% acetic acid under a fume hood.
- Add carmine powder to the boiling 45% acetic acid.

- Boil for 5 to 10 min, with occasional stirring, until color becomes dark red.
- Cool and filter into a colored bottle, and store in a refrigerator.

Note: For preparing aceto-orcein, carmine is replaced by orcein in the same procedure.

c. Staining of Root

- Aceto-carmine stain has been effective for somatic chromosomes of barley. Roots are transferred directly into the stain after pretreatment.
- The stain acts like a fixative and is routinely used for the analysis of F₁ and F₂ aneuploid populations of barley. Cells are fragile when slides are prepared immediately (Tsuchiya, 1971a).

2. Feulgen Staining

a. Ingredients

- 1 g basic fuchsin (use a stain commission type especially certified for use in Feulgen technique)
- 200 mL ddH₂O
- 30 mL 1 N HCl
- 3 g potassium metabisulfite ($K_2S_2O_5$)
- 0.5 g activated charcoal

b. Preparation of Stain

Method 1:

Dissolve 1 g basic fuchsin gradually in 200 mL of boiling ddH_2O and shake thoroughly.

Cool to 50°C, and filter.

Add to filtrate 30 mL 1 *N* HCl and then 3 g potassium metabisulfite (should not be from old stock). Close mouth of the container with a stopper, seal with parafilm, wrap container with aluminum foil, and store in a dark chamber at RT for 24 h.

If solution shows faint straw color, add 0.5 g activated charcoal powder, shake thoroughly, and keep overnight in a refrigerator (4°C).

Filter and store the stain in a colored bottle in a refrigerator.

Method 2: (J. Greilhuber, personal communication, 2002)

Dissolve 1g basic fuchsin in 30 mL 1 N HCl at RT (3 h, stirring).

Add 170 mL dH₂O + 4.45 g K₂S₂O₅ (3 h or overnight, stirring), tighten vessel.

Add 0.5 g activated charcoal (15 min, stirring), filter by suction. The reagent is colorless. Avoid evaporation of SO_2 during all steps.

Note: After decoloration, the solution has lost the red tint but may be brown, like Cognac. These compounds are removed by the charcoal treatment. The use of aged $K_2S_2O_5$ can result in precipitation of white crystal needles (leucofuchsin), which are insoluble in SO_2 water but soluble in dH_2O . This can cause serious problems with washing out residual reagent from the sample. The SO_2 water washes (0.5 g $K_2S_2O_5$ in 100 mL $dH_2O \pm 5$ mL 1 *N* HCl) should always be done for 30 to 45 min for removing nonbound reagent from the stained tissue. Plasmatic background can so be avoided.

Ready-to-use Feulgen stain is also available: Sigma, Schiff reagent, Fuchsin-sulfite reagent (lot # 51H5014).

c. Staining of Roots

Wash fixed root tips once with ddH₂O to remove the fixative. If formaldehyde has been used in the fixative, wash roots in running tap water thoroughly for 1 h or wash 3 times for 15 min in ddH₂O.

- Hydrolyze roots in 1 *N* HCl (60°C) for 6 to 10 min, depending upon the specimen. Instead of 1 *N* HCl at 60°C, 5 *N* HCl at 20°C for 60 min may be used. The latter is recommended for quantitative Feulgen staining (J. Greilhuber, personal communication, 2002).
- After hydrolysis, rinse root tips in ddH_2O , remove excess water, and transfer root tips to Feulgen stain. Adequate staining can be achieved after 1 to 2 h at RT.
- Cut stained region of root, place on a clean glass slide, add a drop of 1% aceto-carmine or 1% propiono-carmine, place a cover glass, and prepare slide by squash method.

Tuleen (1971) modified the above procedures for barley chromosomes. After pretreatment, transfer roots to 1 *N* HCl for approximately 7 min at 55°C. Transfer hydrolyzed roots immediately to Feulgen stain. Keep roots in stain at RT for 2 h. If it is desired to keep materials for a longer period, store in a refrigerator. Squash in 1% aceto-carmine.

Palmer and Heer (1973) developed a cytological procedure by using Feulgen stain for soybean chromosomes after comparing several techniques. Their modified version is as follows:

- Germinate seeds until roots are 7 to 10 cm long. Collect root tips after first 3 h of the 30°C period. Seed germinated in a sandbox or bench in a greenhouse yields a large number of secondary roots after 5 to 7 days of germination. It is advised to germinate seeds of soybean or large-seeded legumes in a sand bench or in vermiculite in the greenhouse.
- Excise 1 cm of root tips, slit last 1/3 with a razor blade. Pretreat tips in covered vials in saturated solution PDB 15°C for 2 h. Pretreatment of roots with 0.1% colchicine for 1 1/2 h at RT proved to be better than PDB treatment. Pretreatment with 8-hydroxyquinoline at 16 to 18°C for 3 to 5 h has been found to be efficient to arrest a large number of metaphase cells.
- Wash root tips with ddH_2O , fix in freshly prepared 3:1 (95% ethanol:glacial acetic acid) for at least 24 h in covered vials.
- Wash root tips in ddH₂O, drain the water, and hydrolyze in 1 N HCl 10 to 12 min at 60° C.
- Wash root tips once in ddH_2O , place in Feulgen stain in covered vials for 1 1/2 to 3 h at RT.

Wash root tips in ice-cold ddH₂O.

- Place root tips in pectinase in spot plates for 1 to 2 h, at 40°C. After treatment in pectinase, root tips may be stored in 70% ethanol in covered vials at 4°C in a refrigerator, or slides may be made immediately.
- *Note:* This step may be omitted. After appropriate staining, wash roots once with chilled ddH_2O and store in cold ddH_2O in a refrigerator. Chromosome staining is not distorted as long as roots are in chilled ddH_2O .
- Place a root tip on a clean slide; remove root cap with a razor blade. Place less than 1 mm of root tip (dark purple region only) in a drop of propiono-carmine or aceto-carmine stain. Place a cover glass and prepare slide by squash method.

d. Softening of Roots

Pectinase has been found to be effective to soften root tips and has been used in various combinations and concentrations depending upon the materials:

- 500 mg pectinase, 500 mg cellulase, 10 mL ddH₂O, 6 drops 1 *N* HCl; before use, wait 24 h after solution is made. The solution can be stored in a refrigerator (2°C) for 1 to 2 months: soften roots for 1 to 1 1/2 h at RT (Kimber et al., 1975).
- 2% cellulase (Onozuka-Rl0), 2% pectinase (Fluka): Soften roots for 5 to 8 h at RT (Hadlaczky and Kalmán, 1975).
- 2% cellulase, 2% pectinase, adjust pH 4.5 with 0.2 *N* HCl: soften roots for 1 to 4 h at RT (Stack and Comings, 1979).

3. Alcoholic-Hydrochloric Acid-Carmine

Snow (1963) developed this stain from the study of mitotic and meiotic chromosomes of several plant species.

a. Ingredients

- 4 g (certified) carmine powder
- 15 mL dH₂0
- 1 mL concentrated HCl
- 95 mL (85%) ethanol

b. Preparation of Stain

- 1. Add 4 g certified carmine to 15 mL dH_20 in a small beaker.
- 2. Add 1 mL concentrated HCl. Mix well and boil gently for about 10 min with frequent stirring.
- 3. Cool, add 95 mL 85% ethanol, and filter.

c. Staining of Roots

- 1. For staining, wash the fixed materials with two to three changes of 70% ethanol, allowing at least 1 h for each change.
- 2. The specimen can be stored in the last change of ethanol in a refrigerator.
- 3. Place the drained specimen (roots or anthers) in stain for at least 24 h. Sometimes, several days or weeks are needed to stain compact tissues (e.g., heads of *Compositae*).
- 4. Pour used stain back into stock bottle. It can then be reused.
- 5. Rinse the specimen with dH_20 or with 70% ethanol.
- 6. Maceration of tissues should be done in 45% acetic acid at 60°C, and slides should be prepared by squash method.

Note: Aceto-carmine stain is not required, and materials should not be hydrolyzed in 1 N HCl, because HCl bleaches the stain.

4. Lacto-Propionic-Orcein

Dyer (1963) used this particular stain for a large number of crops. This stain is useful for plants with small or numerous chromosomes. Chromosomes are intensely stained, and cytoplasm remains clear.

a. Ingredients

- 2 g orcein
- 50 mL lactic acid
- 50 mL propionic acid

b. Preparation of Stain

- 1. Add 2 g natural orcein to 100 mL of a mixture of equal parts of lactic acid and propionic acid at RT.
- 2. Filter and dilute the stock solution to 45% with dH_2O .

c. Staining of Roots

Stain specimens in lacto-propionic-orcein stain for 2 min, and prepare slide by squash method.

5. Carbol Fuchsin Staining

This technique was modified from Darlington and La Cour (1969).

a. Ingredients - 1

- Solution A
 - 3 g basic fuchsin in 100 mL 70% ethanol
 - 10 mL 3% basic fuchsin
 - Add 90 mL of 5% phenol in ddH₂O

Note: Solution A can be stored in a refrigerator for a long time.

- Solution B
 - 55 mL Solution A
 - 6 mL glacial acetic acid
 - 3 mL formalin

Note: Solution B can be stored for only 2 weeks in a refrigerator.

b. Staining Solution (100 mL)

- 20 mL Solution B
- 80 mL 45% acetic acid
- 1.8 g sorbitol

c. Ingredients - 2

This recipe has been formulated to reduce the concentration of formalin. The higher concentration of formalin is toxic.

- Solution A
 - 20 mL 3% basic fuchsin in 70% ethanol
 - 80 mL of 5% phenol in ddH_2O

Note: Solution A can be stored in a refrigerator for a long time.

- Solution B
 - 55 mL Solution A
 - 6 mL glacial acetic acid
 - 3 mL formalin

Note: Solution B can be stored for only 2 weeks in a refrigerator.

d. Staining Solution (100 mL)

- 10 mL Solution B
- 90 mL 45% acetic acid
- 1.8 g sorbitol

Caution! Handle carbol fuchsin stain under a fume hood, store stain for at least 2 weeks at RT before use (aging of stain), and store working stain in a refrigerator.

e. Staining of Roots

- 1. Stain roots for 14 to 16 h in a refrigerator.
- Wash roots in cold ddH₂O (store water bottle in a refrigerator) at least 3 times in order to remove carbol fuchsin stain.
- 3. Store roots in cold ddH₂O in a refrigerator.
- 4. Squash a root tip under a coverslip in a drop of 45% acetic acid.

Note: Roots can be stored for a long time in a refrigerator (Singh, Kollipara, and Hymowitz, 1998b; Xu et al., 2000a).

6. Giemsa Staining

a. Giemsa C-Banding Technique

The Giemsa C-banding technique has facilitated the identification of individual chromosomes in many plant species by its characteristic C- (constitutive heterochromatin) banding patterns. Several minor modifications in the C-banding procedure have been made to obtain the maximum number of C-bands in the chromosomes. Procedures of seed germination and pretreatments described earlier can be followed. Fix roots in 3:1 (95% ethanol:glacial acetic acid) for at least 24 h. Soften cereal roots in 45% acetic acid or in 0.5% aceto-carmine. Pectinase and cellulase can also be used to soften the roots. Prepare the slide by the squash method. Remove cover glass by dry-ice method (Conger and Fairchild, 1953).

i. Dehydration — The majority of researchers placed slides in 95 to 100% ethanol for 1 h to obtain C-bands in *Secale* chromosomes (Thomas and Kaltsikes, 1974; Singh and Lelley, 1975; Singh and Röbbelen, 1975; Kimber et al., 1975; Bennett, Gustafson, and Smith, 1977; Seal and Bennett, 1981) and *Hordeum vulgare* chromosomes (Linde-Laursen, 1975). However, Stack and Comings (1979) did not dehydrate rye chromosomes and obtained satisfactory C-banding patterns. In contrast, Limin and Dvořák (1976) and Fujigaki and Tsuchiya (1985) placed rye chromosomes slides in ethanol for 2.25 h and 2 h, respectively, and observed C-banding patterns similar to those observed after 1 h dehydration. Ward (1980) treated maize chromosomes slides in 75% (10 min), 95% (1.5 h), and 100% ethanol (20 min). On the other hand, Mastenbroek and de Wet (1983) rinsed maize and *Tripsacum* chromosome slides once in 90% ethanol and twice in 100% ethanol.

After dehydration, keep air-dried slides at RT overnight (Singh and Röbbelen, 1975). Linde-Laursen (1975) kept barley chromosome slides in a desiccator over silica gel for 1 to 2 weeks at 18°C. Air-dried slides can be processed through BSG (barium hydroxide/saline/Giemsa) method.

ii. Denaturation — Prepare fresh saturated solution of barium hydroxide [5 g Ba(OH)₂·8 H₂O + 100 mL dH₂O]. Solution is filtered in a Coplin jar. Keep Coplin jar in a water bath (50 to 55°C) or at RT. An alternative is to use hot dH₂O and cool to 50 to 55°C or 20°C. Filtering is not necessary. Replace Ba(OH)₂ solution by cold water and rinse slides (J. Greilhuber, personal communication, 2002).

Darvey and Gustafson (1975) prepared barium hydroxide by adding deionized distilled water at 80°C to remove CO_2 from the crystal of barium hydroxide. Shake the solution a few seconds and pour off the supernatant. Deionized water was then added to the crystals of barium hydroxide remaining in the bottom of the flask.

Treat slides in barium hydroxide for 5 to 15 min at 50 to 55°C depending upon the materials. Wash slides thoroughly in three or four changes of dH_2O for a total of 10 min, and air dry the slides or transfer the slides directly into 2 × SSC. Merker (1973) skipped the denaturation step and still produced satisfactory C-bands in rye chromosomes.

iii. Renaturation — Incubate slides in $2 \times SSC$ (0.3 M NaCl + 0.03 M Na₃C₆H5O₇·2 H₂O), pH 7 to 7.6 at 60 to 65°C in a water bath or oven for 1 h (Merker, 1973; Singh and Röbbelen, 1975; Weimarck, 1975; Thomas and Kaltsikes, 1974; Verma and Rees, 1974; Kimber et al., 1975; Hadlaczky and Kalmán, 1975; Fujigaki and Tsuchiya, 1985). Incubation periods and temperatures are variable and can be determined by experimentation. After saline sodium citrate (2 × SSC) treatment, wash slides in three changes of dH₂O for a total of 10 min, and air dry the slides.

iv. Staining — Stain slides for 1 to 2 min with Giemsa stain, 3 mL stock solution + 60 mL Sörensen phosphate buffer (0.2 M), pH 6.9 (30 mL KH_2PO_4 + 30 mL Na_2 HPO_4 ·2 H_2O), freshly mixed. Monitor staining regularly. The Leishman stain also produces results similar to those with Giemsa

stain (Darvey and Gustafson, 1975; Thomas and Kaltsikes, 1974; Ward, 1980; Chow and Larter, 1981). After optimal staining, place slides quickly in dH₂O, air dry, store in xylene overnight, air dry again, and mount cover glass in Euparal or Canada Balsam or Permount.

v. Preparation of Giemsa Stock Solution from Powder — The following describes the preparation of Giemsa stock solution from powder (Kimber et al., 1975):

- 1 g Giemsa powder
- 66 mL glycerine
- 66 mL methanol
- Dissolve Giemsa powder in the glycerine at 60°C for 1 h with constant stirring. Add methanol, and continue stirring at 60°C for 1 day (24 h). Filter and keep in a refrigerator. It can be kept for 1 or 2 months.

Noda and Kasha (1978) did not use BSG method for banding barley chromosomes. They developed the following protocol:

- 1. Collect barley roots, pretreat, and fix as described earlier.
- 2. Wash roots in tap water.
- 3. Hydrolyze in 1 N HCl at 60° C for 7 min.
- 4. Treat roots in 1% pectinase at 37°C for 13 min.
- 5. Transfer roots in tap water, and prepare slides by squash technique in a drop of 30% acetic acid.
- 6. After dehydration and maturation of slides, place slides in 1 *N* HCl at 60°C for 5 min, rinse slides in tap water, and air dry at RT for $\frac{1}{2}$ day.
- 7. Immerse slides in 0.07 N NaOH for 35 sec.
- Stain in 60 × diluted Giemsa solution in 0.067 M Sörensen phosphate buffer (pH 6.9) for about 30 min. After appropriate staining, rinse slides in tap water, air dry slides at RT, and mount cover glass in Canada balsam.

Limin and Dvořák (1976), working with rye, demonstrated that the combination of barium hydroxide and saline sodium citrate (SSC) treatment is necessary for obtaining clearly differentiated dark-staining bands.

The C-banding procedure of Giraldez, Cermeño, and Orellana (1979) for rye chromosomes is as follows:

- 1. Immerse slides in 0.2 N HCl at 60°C for 3 min.
- 2. Wash slides in tap water and treat in saturated solution of barium hydroxide [Ba(OH)₂] at RT for 10 min.
- 3. Wash in tap water, treat slides in 2 × SSC at 60°C for 1 h, and stain in 3% Giemsa in phosphate buffer pH 7.0.
- 4. Wash in tap water, air dry, immerse in xylene for 5 min, and mount in DPX.

Carvalho and Saraiva (1993) modified the C-banding technique for maize chromosomes:

- 1. Treat 2-week-old slides with freshly prepared 5% Ba(OH)₂·8 H₂0 solution at 56°C with continuous agitation for 10 to 15 sec.
- 2. Wash slides in 70% ethanol (two changes) and then in 100% ethanol, and transfer slides to methanol:acetic acid (8:1).
- 3. Dry slides on a hot plate for a few minutes and stain with 3% Giemsa (MERCK) in phosphate buffer, pH 6.8, for 8 to 10 min, then wash slides twice in dH₂O and air dry.

b. Giemsa N-Banding Technique

This technique was originally developed to stain nucleolus organizing regions of mammalian and plant chromosomes (Matsui and Sasaki, 1973; Funaki, Matsui, and Sasaki, 1975):

- 1. Incubate slides at $96 \pm 1^{\circ}$ C for 15 min in 1 N NaH₂PO₄ (pH 4.2 ± 0.2) and adjust pH with 1 N NaOH.
- 2. Rinse thoroughly in distilled water and stain in Giemsa (dilute 1:25 in 1/15 M phosphate buffer, pH 7.0) for 20 min.
- 3. Rinse slide in tap water and air dry.

Gerlach (1977) modified the Giemsa N-banding technique of Funaki, Matsui, and Sasaki (1975) for the staining of wheat chromosomes:

- 1. Incubate air-dried slides in 1 M NaH₂PO₄ (pH 4.15) for 3 min at 94 \pm 1°C.
- 2. Rinse slides in distilled water.
- 3. Stain for 30 min with a solution of 10% Gurr's Giemsa R66 in 1/15 M Sörensen's phosphate buffer (pH 6.8).
- 4. Rinse slides in tap water and mount in immersion oil.

The Giemsa N-banding technique has been used with slight modifications following Gerlach (1977) to identify chromosomes of barley (Islam, 1980; Singh and Tsuchiya, 1982a,b), wheat (Jewell, 1981; Endo and Gill, 1984) and lentil (Mehra, Butler, and Beckman, 1986).

Singh and Tsuchiya (1982b) suggested that the Giemsa banding technique should be considered as a qualitative tool to identify individual chromosomes, while conventional staining methods should be used as a quantitative approach to establish the standard karyotype. The combination of acetocarmine and Giemsa staining suggested by Nakata, Yasumuro, and Sasaki (1977), used by Singh and Tsuchiya (1982a,b) for barley and Endo and Gill (1984) for wheat chromosomes, demonstrated that karyotype analysis could be conducted with greater precision than was previously possible:

- 1. Barley roots can be collected and fixed according to the procedures described earlier (Singh and Tsuchiya, 1982b).
- 2. Transfer fixed roots to aceto-carmine (0.3%) for about 2 to 3 h.
- 3. Prepare slides by squash method.
- 4. Photograph the cells with well-spread chromosomes by phase contrast.
- 5. Remove cover glass by dry-ice method.
- 6. Place slides in 96% ethanol for 2 to 4 h. Endo and Gill (1984) treated wheat chromosome slides with hot (55 to 60°C) 45% acetic acid for 10 to 15 min.
- 7. Air dry slides overnight at RT. Linde-Laursen (1975) placed slides in a desiccator over silica gel for 2 to 4 weeks.
- 8. Incubate slides in 1 M NaH₂Po₄, pH 4.15, for 1.5 to 2 min (5 min for barley) at $94 \pm 1^{\circ}$ C.
- 9. Rinse slides briefly in dH_2O and then air dry.
- 10. Stain for 20 min to 24 h in 1% Giemsa (Sigma No. G04507) in 1/15 M Sörensen's phosphate buffer.
- 11. After optimal staining of the chromosomes, which were previously photographed from acetocarmine preparation, rinse slides in dH₂O, air dry, keep in xylene overnight, air dry again, and mount in Permount.

c. HCI-KOH-Giemsa (HKG) Technique

Shang, Jackson, and Nguyen (1988a,b) claimed that the HKG technique produces well-separated and sharply banded chromosomes, including centromeric bands of wheat chromosomes, and that the results are more highly reproducible than C- or N-banding techniques:

- 1. Pretreat roots for 2.5 h with an aqueous solution of α -bromonaphthalene (0.01 mL stock solution in 10 mL water; stock solution: 1 mL α -bromonaphthalene in 100 mL absolute ethanol).
- 2. Wash roots twice in distilled water and hydrolyze with 5 N HCl for 20 min at RT.
- 3. Wash roots twice in distilled water and store material in 45% acetic acid.
- 4. Clean slides in 95% ethanol and wipe dry.
- 5. Prepare slide by taking 1 mm meristematic part of the root in 45% acetic acid.

- 6. Remove cover glass by liquid nitrogen. The cover glass can be removed by placing prepared slides in liquid nitrogen, by dry ice, or by placing in an -80°C freezer. The purpose is not to lose the specimen or to distort the chromosome spread.
- 7. Air dry the slide and store for 3 to 7 days.
- 8. Treat air-dried slide with 1 *N* HCl at 60°C for 6 min and wash slides four times in distilled water for a total of 10 min at RT.
- 9. Air dry slide for half a day, dip slide into fresh 0.07 N KOH for 20 to 25 sec followed by dipping into 1/15 M Sörensen's phosphate buffer (pH 6.8) for 5 to 10 sec with shaking.
- 10. Stain in 3% Gurr's improved Giemsa stain solution (3 mL stain in 100 mL 1/15 M Sörensen's phosphate buffer, pH 6.8) for 1 to 2 h or until proper staining is obtained. The slides can be kept in the stain for 1 to 2 days without overstaining.
- 11. Rinse slide in distilled water, air dry, and mount in a synthetic resin (Preserveaslide, Mathesoin, Coleman and Bell Co., Norwood, OH).

d. Modified HKG-Banding Technique

The following explains the modified HKG-banding technique (Carvalho and Saraiva, 1993):

- 1. Hydrolyze 1- to 5-day-old slides in 1 N HCl at 60°C for 4 to 6 min.
- 2. Wash slides four times in dH_2O for a total of 10 min.
- 3. Immerse slides briefly in 0.9% NaCl, plunge 10 times in 70% ethanol, and dry slides on a hot plate (surface temperature 50°C) for a few seconds.
- 4. Immerse slides in 0.06 N KOH solution for 8 to 12 sec with continuous agitation at RT.
- 5. Wash slides in 70% ethanol (two changes) and 100% ethanol and transfer to methanol:acetic acid (8:1).
- 6. Air dry slides on a hot plate for few minutes and stain according to the procedure described for the modified C-banding procedure.

e. Giemsa G-Banding Technique

The G-banding technique was originally developed to identify human chromosomes. Kakeda et al. (1990) perfected the G-banding method for maize chromosomes:

- 1. Germinate maize seeds in small pots filled with moist vermiculite for 2 days at 32°C under continuous light.
- Excise three root tips about 1 cm long from each seed and pretreat with a 0.05% colchicine solution or 0.05% colchicine solution containing either 10 ppm actinomycine D or ethidium bromide for 2 h at 25°C.
- Dip root tips in the Ohnuki's hypotonic solution (55 mM KCl, 55 mM NaNO₃, 55 mM CH₃COONa, 10:5:2) for 30 min to 1 h at 25°C.
- 4. Fix root tips with methanol-acetic acid (3:1) for 1 to 4 days in a freezer (-20°C) or for at least 2 h at 25°C to actinomycine-D pretreated ones.
- 5. Remove meristem cells with a tweezer in a drop of fresh fixative or macerate enzymatically. *Note:* For enzyme maceration; wash fixed root tips for about 10 min and macerate in an enzyme mixture [2% Cellulase RS (Yakult Honsa Co., Ltd, Tokyo) and 2% MacerozymeR -200 (Yakult Honsa), pH 4.2] for 20 to 60 min at 37°C in a 1.5 mL Eppendorf tube. Rinse root tips with dH₂O two or three times. Pick up macerated root tip with the help of a Pasteur pipette, and place it on a glass slide.
- 6. Cut the actively growing tip into small pieces with a sharp-pointed tweezer with the addition of fresh fixative.
- 7. Observe the slide under phase contrast microscope, select a slide with well-spread chromosomes, and air dry for about 2 days in an incubator at 37°C.
- 8. Stain samples, prepared by the actinomycine-D, directly in 10% Wright solution diluted with 1/15 M phosphate buffer (pH 6.8) for 10 min at 25°C, wash, and air dry slides.
- 9. For slides prepared by enzyme maceration, fix sample again in a 2% glutaraldehyde solution diluted with phosphate buffer for 10 min at 25°C and wash.

- Immerse postfixed slides in 2% trypsin (MERCK, Art. 8367) dissolved in PBS (pH 7.2) for 10 min at 25°C or in 0.02% SDS dissolved Tris-HCl buffer (20 mM, pH 8.0) for 2 to 25 min at 25°C.
- 11. Wash slides briefly, and air dry.
- 12. Stain slides in 5% Wright solution in 1/30 M phosphate buffer (pH 6.8) for 5 min.

III. SMEAR TECHNIQUE FOR PLANT CHROMOSOMES

The removal of the cover glass is a prerequisite for Giemsa C- and N-banding techniques for plant chromosomes. To avoid this step, attempts are being made to prepare plant chromosome slides by smear technique, as used for the mammalian chromosomes. By smear technique, chromosome slides have been prepared from the roots (Kurata and Omura, 1978; Rayburn and Gold, 1982; Pijnacker and Ferwerda, 1984; Geber and Schweizer, 1988) and protoplasts (Mouras, Salessas, and Lutz, 1978; Malmberg and Griesbach, 1980; Murata, 1983; Hadlaczky et al., 1983) of several plant species.

A. Chromosome Preparation from the Root

- 1. Germination of seeds and collection and pretreatment of roots should be conducted as described earlier.
- Cut only the meristematic region (1 mm) of the roots, and treat roots with hypotonic 0.075 M KCl for 15 min (potato Pijnacker and Ferwerda, 1984), 20 min (rice Kurata and Omura, 1978) at RT. Geber and Schweizer (1988) did not treat roots with hypotonic solution.
- Fix roots in absolute ethanol/glacial acetic acid (3:1) or methanol/glacial acetic acid (3:1) for a minimum of 1 h (Geber and Schweizer, 1988) or 2 h or up to several months in a deep freezer (Pijnacker and Ferwerda, 1984).
- 4. Rinse fixed roots in 0.1 M citric acid-sodium citrate buffer, pH 4.4 to 4.8.
- Enzyme treatment is quite variable: 10% pectinase (Sigma P-5146) + 1.5% cellulase (Onozuka R-10) in citrate buffer, at 37°C for 30 min (Pijnacker and Ferwerda, 1984); 6% cellulase (Onozuka R-10) + 6% pectinase (Sigma), adjust pH 4.0 with HCl, treat roots for about 60 min at 35°C (Kurata and Omura, 1978).

A mixture of pectinase, 20 to 40% (v/v), (Sigma, from *Aspergillus niger*, P-5146, obtained as glycerol-containing stock solution) + 2 to 4% (w/v) cellulase (Calbiochem 21947 or Onozuka R-10) in 0.01 M citric acid-sodium citrate buffer was used for softening root tissues of *Sinapsis alba* (Geber and Schweizer, 1988). According to Kurata and Omura (1978), wash the roots in distilled water for 5 to 10 min at about 20°C to remove the enzyme. Place the root meristem on a clean slide with a drop of fresh fixative (three parts methanol:one part acetic acid), and break the root meristem into fine pieces with a needle. Add a few drops of the fixative, and flame dry the slide.

Pijnacker and Ferwerda (1984) prepared slides of potato (*Solanum tuberosum*) by the following smear method:

- 1. Transfer one root meristem to a clean slide.
- 2. Remove the excess buffer, add a drop of 60% acetic acid, heat slide (without boiling) over an alcohol flame, and leave for 2 to 5 min.
- 3. Suspend cells in a drop of acetic acid with the help of a fine needle, leave for 1 min, and heat slightly again with tilting of the slide.
- 4. Add Carnoy's solution to suspension when slide is cooled (some seconds), add three more drops of Carnoy's solution on the top of the suspension, air dry the slides, store overnight or longer, and stain as needed.

Geber and Schweizer (1988) used the following procedure:

1. Spin the cells at about 4000 rpm in small conical centrifuge tube (10 mL volume) at each step to change the solution.

- 2. Remove the supernatant carefully with a Pasteur pipette, and resuspend the pellet in approximately 5 mL liquid.
- 3. Transfer fixed roots to buffer, collect only meristem tissues, and rinse twice in buffer.
- 4. Soften the tissue in enzyme solution at 37°C for 1 to 2 h, depending upon plant material.
- 5. Wash twice in buffer, and finally, suspend the pellet in a drop of buffer to prevent the sticking together of the protoplasts.
- 6. Add an excess of fixative (freshly prepared) and change it twice, suspend the pellet in a small amount of fixative, and drop this suspension onto ice-cold tilted slides and air dry.
- 7. Age air-dried slides overnight or longer before further processing.

Note: In addition to *Sinapsis alba*, the above procedure produced an excellent chromosome spread of *Vicia faba*, *Pisum sativum*, *Crepis capillaris*, *Calla palustris*, and *Spirodela polyrrhiza*.

B. Chromosome Preparation from Cell Suspension and Callus

Murata (1983) developed an air dry method, described below, to study chromosomes from suspension and callus cultures. The procedure has been divided into three steps: (1) pretreatment to accumulate metaphase cells, (2) cell wall digestion and protoplast isolation, and (3) application of air-drying technique.

1. Suspension Culture of Celery (Apium graveolens)

- 1. Add 1 mL of 0.5% colchicine solution to 9 mL of cell suspension 2 to 3 days after subculture in a 100×15 mm Petri dish, and place on a gyratory shaker (50 rpm) for 2 h.
- Transfer 2 mL of cell suspension to 2 mL of enzyme solution in 100 × 15 mL Petri dish [enzyme solution: 2% cellulysin (Calbiochem), 1% macerase (Calbiochem), and 0.6 M sorbitol (pH 5.5 to 5.6)]. Substitution of 1% pectolyase Y-23 (Kikkoman) for macerase produced comparatively faster protoplast isolation.
- 3. Seal the Petri dish with parafilm, and place on the gyratory shaker (50 rpm) for 3 to 4 h at RT (25°C).
- 4. Filter the cells and enzyme mixture through 60 μ m nylon mesh into a 15 mL centrifuge tube, and centrifuge (65 × g) for 3 min.
- 5. Rinse twice with 0.6 M sorbitol, suspend in 5 mL hypotonic solution (0.075 M KCl), and allow to stand for 7 min at RT (25°C).
- 6. Remove the supernatant following centrifugation $(65 \times g)$ for 5 min, gradually add fresh fixative 3 (95% ethanol):1 (glacial acetic acid) up to 5 mL, and allow to stand for 1 h.
- 7. Resuspend in fresh fixative following centrifugation; repeat twice, and make the final volume of the fixed cells 0.5 to 1.0 mL.
- 8. Put five to six drops of the fixed cells, by using a Pasteur pipette, onto a wet cold slide, and air or flame dry.
- 9. Stain 3 to 4 min with 4% Giemsa (Gurr's R66, Bio/Medical Special) diluted with 1/15 M phosphate buffer (pH 6.8), rinse in phosphate buffer and distilled water, and air dry.
- 10. Mount slide in DePeX mounting medium (Bio/Medical Special).

2. Callus Culture of Brassica carinata

- 1. Put 10 to 20 mg calluses 5 to 7 days after subculture into a 15 mL centrifuge tube with 5 mL of Murashige and Skoog (1962) (MS) liquid medium containing 0.5% colchicine, and allow to stand for 5 h.
- 2. Discard the liquid medium following centrifugation (100 × g) for 5 min, and suspend in the fresh Carnoy's fixative (3:1) for 1 h.
- 3. Rinse twice with dH₂O, and add 5 mL of enzyme solution as in Step 2 of celery.
- 4. Place the centrifuge tube, sealed with a cap, horizontally, on the gyratory shaker (100 rpm) for 2 h.
- 5. Filter the cell suspension through 60 μ m nylon mesh into another centrifuge tube.
- 6. The subsequent procedures are as described in Steps 6 to 10 of celery.

Pijnacker, Walch, and Ferwerda (1986) studied the chromosomes of leaf explants of potato by following the smear technique:

- 1. Fix leaf explants (with developing callus) directly in 3:1 (ethanol 100%:glacial acetic acid) at 4°C for about 24 h or pretreat in a saturated α -bromonaphthalene for 3 h at RT before fixation.
- 2. Rinse leaf pieces in distilled water, incubate in 15% (v/v) pectinase (Sigma 5146) + 1.5% (w/v) cellulase R10 (Yakult) in citrate buffer pH 4.8 for 45 min at 37°C, rinse, and then keep in distilled water for a minimum of 2 h.
- 3. Transfer one leaf piece to a clean slide, and add a drop of 60% acetic acid.
- 4. Make leaf pieces into a fine suspension.
- 5. Surround this suspension with cold fixative (3:1), then add about three drops of fixative on top of the suspension, and air dry the slide.
- 6. Stain chromosomes by the fluorescent and Giemsa techniques.

C. Chromosome Preparation from Flower

Murata and Motoyoshi (1995) developed a cytological procedure for *Arabidopsis thaliana* floral tissues of young buds:

- 1. Dissect out sepals from young buds (1.5 to 2 mm in length) under a dissecting microscope.
- 2. Transfer buds without sepals into a 1.5 mL microtube with 1 mL dH₂O, and keep at 0°C in iced water for 24 h to allow accumulation of metaphase cells.
- 3. Fix buds in 3:1 (99% methanol:glacial acetic acid), and store at -20° C.
- 4. For slide preparation, rinse buds well with dH₂O.
- 5. Digest buds with an enzyme solution containing 2% (w/v) cellulase Onozuka R10 (Kinki Yakult) and 20% (v/v) pectinase (Sigma); incubate for 1.5 h at 30°C.
- 6. Suspend tissues with micropipette, rinse suspended cells with dH₂O, and fix again.
- 7. After three changes of fresh fixative, drop fixed cells onto wet, cold slides, and flame dry.
- 8. Stain slides with 4% (v/v) Giemsa solution diluted with 1/15 M phosphate buffer, pH 6.8 for 20 to 30 min, and examine with a microscope.

D. Chromosome Preparation from Shoot

Ma et al. (1996) developed a cytological procedure to study mitotic metaphase chromosomes from shoots of roses:

- 1. Collect actively growing terminal shoot (2 to 4 mm) in the morning during the springtime burst of growth. Place five to 10 shoot tips in one 5 mL tube, and keep on ice water (0°C) for transport to the laboratory.
- Remove the young outside leaves, cut terminal 2 to 4 mm portion of the shoot apex, and place immediately in a pretreatment solution (0.1% colchicine + 0.001 M 8-hydroxyquinoline) for 4 h in the dark at ≈25°C.
- 3. Fix shoot tips in a freshly prepared mixture of 2 acetone:1 acetic acid (v/v) + 2% (w/v) polyvinylpyrrolidone (MW 40,000). Fix tissues for 24 h at ambient pressure.
- 4. Soak shoot tips in dH₂O for 1 to 24 h to elute the fixative and soften the tissues.
- 5. Hydrolyze shoot tips in 1 *N* HCl for 20 min at $\approx 25^{\circ}$ C. Remove 1 *N* HCl, wash two times in dH₂O, and soak for 10 min in dH₂O.
- 6. Incubate shoot tips in 0.075 M KCl for 30 to 60 min. (Alternative: 0.01 M sodium citrate, pH 4.6, for 15 to 30 min.)
- Digest the shoot tips in 5% cellulase R10 (Yakult Honsha Pharmaceutical, Tokyo) + 1% pectolyase Y23 (Seishin, Tokyo) + 0.01 M sodium citrate at pH 4.6. Use 2 to 4 μL of enzyme mix per shoot tip. Digest shoot tip in enzyme for 3 to 4 h at 37°C.
- 8. Prepare slides by one-slide maceration or by suspension and spreading of protoplasts.
- 9. One-slide maceration:

- a. Place one shoot tip on a microscope slide, and blot excess water.
- b. Add two to three drops of 3:1 or 1 ethanol:1 acetic acid fixative.
- c. Disperse tissues gently with forcep tips, examine briefly at ×160, until most of the cells are settled onto the surface of the slide.
- d. Wash cytoplasmic debris with additional drops of fixative. Remove excess fixative by filter paper, air dry, stain, and cover with a coverslip.
- 10. Suspension and spreading of protoplast:
 - Add about 480 μL dH₂O to microcentrifuge tube containing the shoot tips and enzyme mix. Vortex tube vigorously for 30 to 60 sec to break up the shoot tips. Discard undissociated pieces.
 - b. Centrifuge at \approx 700 g_n fixed suspension protoplast for 5 min at 2 to 4°C, and discard the supernatant.
 - c. Resuspend the pellet in \approx 500 µL dH₂O. Transfer supernatant cell suspension to a new micro-centrifuge tube.
 - d. Centrifuge the suspension at ~700 g_n for 5 min at 2 to 4°C, discard the supernatant, and resuspend pellet in ~500 μ L of 3 ethanol:1 acetic acid.
 - e. Centrifuge the suspension at \approx 700 g_n for 5 min at 2 to 4°C, discard the supernatant, and resuspend the pellet in 15 to 20 µL freshly made 3 ethanol:1 acetic acid per shoot tip.
 - f. Apply 5 to 10 μ L of the protoplast suspension to a scrupulously clean microscope slide and allow it to air dry (\approx 30 to 50% relative humidity).
 - g. For FISH, allow slides to air dry for 2 to 3 days at low humidity to improve cell retention.

IV. POLLEN STAINING

A. Pollen Fertility

Pollen viability or fertility can be ascertained after staining the mature pollen with aceto-carmine (1%) or I_2 -KI (potassium iodide). Collect flowers during anthesis. Dust pollen grains on a clean slide and add a drop of aceto-carmine, place a cover glass, and heat the slide over a low flame but be sure not to boil. Heating dissolves the starch grains and facilitates the staining of sperm nuclei and the vegetative nucleus. Staining pollen grains with I_2 -KI does not require heating, and fertile pollen grains turn black, while sterile pollen grains are colorless.

Pollen grains with two well-developed sperm nuclei and one vegetative nucleus are considered functional pollen grains (Figure 2.1). Degenerated pollen grains are those containing two deformed

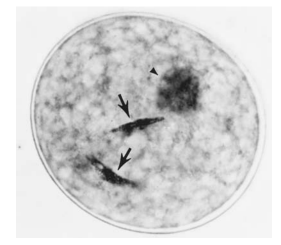


Figure 2.1 A fertile pollen grain with two sperm nuclei (arrows) and a vegetative nucleus (arrow head) in *Secale cereale* (2*n* = 14). (From Singh, R.J., unpublished results.)

sperm nuclei, two or one underdeveloped nuclei, or empty pollen grains that are shrunken and without cytoplasm and nucleus.

Sometimes, pollen fertility can be estimated by the pollen germination test. Dust fresh pollen grains onto the surface of 1 mL of germination medium in depression wells, cover to reduce evaporation, and observe after 1 to 3 h. Germination medium: 12.5, 20, or 25% sucrose in distilled water + 0.01% borate (Coleman and Goff, 1985).

B. Chromosome Count in Pollen

To determine the chromosome count in pollen, the following procedure is suggested (Kindiger and Beckett, 1985):

Stain

- *Solution A*: Dissolve 2 g hematoxylin in 100 mL 50% propionic acid. Allow the solution to age for about 1 week. The solution can be kept indefinitely in a stoppered brown bottle at RT.
- *Solution B*: Dissolve 0.5 g ferric ammonium sulfate [Fe NH₄ (SO₄)] in 100 mL 50% propionic acid. The solution can be kept indefinitely in a stoppered brown bottle at RT.
- *Note:* Mix equal volumes of Solutions A and B. The mixture, which turns dark brown, is ready for use immediately. The stain remains good for about 2 weeks.

The following protocol was developed to count chromosomes from maize pollen grains:

- 1. Collect fresh maize tassels (when about 3 cm of the tassel becomes visible above the leaf whorl) in a waxed short bag or other moisture-retaining container.
- 2. Fix tassel in a 14:1 mixture of 70% ethanol:formaldehyde, and store in a refrigerator. This fixative produces better results than 3:1 (95% ethanol:glacial acetic acid).
- 3. Pretreatment is unnecessary for the fresh material. However, anthers can be pretreated with α -bromonaphthalene with one drop of DMSO for 30 to 40 min, fix anthers in glacial acetic acid.
- 4. Dissect out anthers and place in a drop of 45% acetic acid on a clean slide, add chloral hydrate crystal, remove debris, add a drop of stain, macerate and stain for a min, apply coverslip, heat (but not to boil), and remove excess stain by filter paper by applying gentle pressure. Heat helps to darken the chromosomes and nuclei and clears the cytoplasm.
- 5. To make the slide permanent, seal the edges of the coverslip with Permount and allow to air dry. Stain begins to fade after 3 months.

C. Differential Staining of Pollen

A stain was developed by Alexander (1969) for examining aborted to nonaborted pollen grains. The pollen staining depends upon the concentration of stain, thickness of pollen walls, and pH of stain.

1. Ingredients

• 10 mL	95% ethanol
• 10 mg	Malachite green (1 mL of 1% solution in 95% ethanol)
• 50 mL	dH ₂ 0
• 25 mL	Glycerol
• 5 g	Phenol
• 5 g	Chloral hydrate
• 50 mg	Acid fuchsin (5 mL of 1% solution in dH_20)
• 5 mg	Orange G (0.5 mL of 1% solution in dH_20)
• 1 to 4 mI	Glacial acetic acid

1 to 4 mL Glacial acetic acid

Note: The amount of glacial acetic acid depends upon the thickness of the pollen walls: 1 mL (thin-walled pollen); 3 mL (thick-walled and spiny pollen); or 4 mL (nondehiscent anthers).

2. Staining of Pollen with Thin Wall

- 1. Thin-walled pollen occurs in *Phaseolus, Sorghum, Oryza, Triticum, Hordeum, Zea*, and *Lycopersicon*.
- 2. Stain pollen in one drop of stain, cover with a coverslip, heat over a small flame, and examine microscopically.
- 3. If differentiation is not satisfactory, keep slide in an oven at 50°C for 24 h.
- 4. Mounted pollen can be stored for a week without fading of stain.

3. Staining of Pollen with Thick and Spiny Walls, or Both

a. Nonsticky Pollen

- 1. Acidify about 100 mL of the stain with 3 mL glacial acetic acid.
- 2. Add a small quantity of pollen into a small vial, and pour enough stain to cover pollen.
- 3. Keep in a 50°C oven for 24 to 48 h.
- 4. Examine after 24 h for differentiation.
- 5. Add one drop of 45% acetic acid into the specimen vial, and mix if aborted and nonaborted pollen are not well differentiated.

b. Sticky and Oily Pollen

- 1. Acidify about 100 mL stain with 3 mL glacial acetic acid.
- 2. Fix mature but nondehisced anther for 24 h in a fixative (3 ethanol:2 chloroform:1 glacial acetic acid). This fixative removes sticky materials.
- 3. Transfer through an ethanol-water series to water.
- 4. Remove the excess water by placing anthers between filter papers.
- 5. Cover one or two anthers with one to two drops of stain on a slide, split the anther wall with a needle to release pollen, and remove debris.
- 6. Place a coverslip over the stain, and store at 50°C oven for 24 to 48 h.
- 7. After 24 h, add one to two drops of stain along the sides of the coverslip to replace the amount of stain lost during evaporation.
- 8. Examine the slide after 24 h. If the green color dominates, increase the acidity of the stain mixture. This stain is suitable for *Hibiscus* and *Cucurbita*.

c. Staining of Pollen Inside Nondehiscent Anther

- 1. Acidify 100 mL stain with 4 mL glacial acetic acid.
- 2. Collect anthers immediately after anthesis.
- 3. Fix for 24 h in a fixative:
 - a. 60 mL Methanol
 - b. 30 mL Chloroform
 - c. 20 mL dH₂0
 - d. 1 g Picric acid
 - e. 1 g Hgcl₂
- 4. Transfer through 70, 50, and 30% ethanol, 30 min each, change gradually to hydrate the anthers, and finally, rinse with dH_20 .
- 5. Remove excess water between two filter papers.
- 6. Stain anthers, and keep at 50° C oven for 24 to 48 h.
- 7. Remove excess stain with blotting paper and examine. If overstained, remount in 25% glycerol containing 4% chloral hydrate, and seal.

V. POLLEN-STIGMA INCOMPATIBILITY

A. Staining Pistils by Aniline Blue

To ascertain compatibility in wide crosses, pollen germination, pollen tube growth, and fertilization are determined by fluorescence microscopy. An intersubgeneric hybrid between soybean and a wild perennial species, *Glycine clandestina*, of the subgenus *Glycine*, is cited as an example:

- 1. Fix fertilized gynoecia in 1:1:18 [1 formaldehyde (HCHO approx. 37%):1 propionic acid:18 ethanol (70%)] 24 h postpollination.
- 2. After 24 h fixation, wash gynoecia once with ddH_2O , and transfer to 1 N NaOH at RT (25°C) for 24 h to soften the tissues.
- Wash softened gynoecia twice, 2 min each, with ddH₂O, and stain for 24 h in 0.1% water-soluble aniline blue solution prepared in 0.1 N K₃PO₄ (Kho and Baër, 1968).
- 4. Squash gynoecia gently in a drop of 80% glycerine under a cover glass on a clean slide, and observe by a fluorescence microscope. In a compatible cross of *G. clandestina* \times *G. max*, pollen tubes had already reached the ovules 24 h postpollination (Figure 2.2).

B. Cleared-Pistil Technique

Young, Sherwood, and Bashaw (1979) developed a protocol known as cleared-pistil and thicksectioning technique for detecting apomixis in grasses:

- 1. Collect inflorescences from field- or greenhouse-grown plants at the time of stigma exertion.
- Fix in formalin + glacial acetic acid + alcohol [known as FAA; 95% ethanol:water:40% formalin:glacial acetic acid; 40:14:3:3 (v/v)].

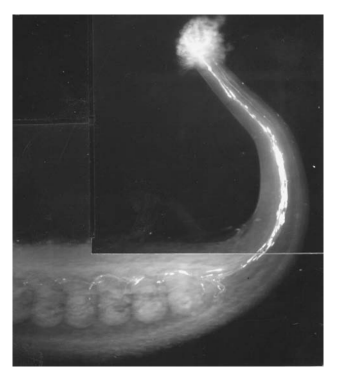


Figure 2.2 Pollen tube growth in a gynoecium following intersubgeneric hybridization in *Glycine clandestina* (2n = 40) × soybean (*G. max*) (2n = 40), 24 h postpollination. Pollen tubes are already in contact with ovules. (From Singh, R.J. and Hymowitz, T., *Plant Breed*, 98, 171–173, 1987. With permission.)

- 3. Excise fixed pistils, and place in 50% ethanol in a 15 mm \times 60 mm screw-cap vial.
- 4. Dehydration and clearing step:
 - a. 70% ethanol \rightarrow 85% ethanol \rightarrow 100 ethanol (three changes)
 - b. 1 (ethanol):1 (methyl salicylate) → 1 (ethanol):3 (methyl salicylate) → 100% methyl salicylate (two changes)

Note: Place one excise pistil in a separate vial with 1 mL liquid for 30 min. Later, wrap 10 to 20 pistils from an inflorescence in a moist envelope from $12 \text{ mm} \times 12 \text{ mm}$ square of single-thickness Kimwipes tissue, place in a vial, and clear using 2 mL solution for 2 h each step.

- 5. Change solution with a Pasteur pipette, store cleared pistils in methyl salicylate in vials, and examine with an interference contrast microscope.
- 6. Mount cleared pistils in methyl salicylate under an unsealed coverslip on a microscope slide.
- 7. Record components of embryo sac and photograph with 35 mm Kodak Technical Pan Film, ASA 100.
- 8. After examination by interference contrast microscopy, transfer each pistil to a vial with 50% ethanol for embedding in plastic.
- 9. The infiltration steps are as follows:
 - a. 50% ethanol → 75% ethanol → 100% ethanol (three changes) → 100% propylene oxide (two changes) → 50% Spurr's low-viscosity embedding medium → 100% Spurr's (four changes). Steps through 50% Spurr's are 30 min each and after 50% Spurr's are 1 h each. At each Spurr's medium step, agitate the vials briefly on a vortex mixture and evacuate.
 - b. Spurr's low-viscosity embedding medium (Spurr, 1969):
 - i. Epoxy resin (ERL 4206: vinyl, cyclohexene dioxide; MW 140.18): 10 g
 - ii. Flexibilizer (DER-736: curing agent (low viscosity), an epoxide MW 175-205, av. 380: 6 g
 - iii. NSA (nonenyl succinic anhydride): 26 g
 - iv. S-1 or DMAE (dimethylaminoethanol, curing agent): 0.2g
 - c. The cure schedule is 16 h at 70°C.
- 10. Remove styles after the final changes, align ovaries in a size 00, square-tipped BEEM capsules, and cure the resin for 16 h at 70°C.
- For sectioning, coat untrimmed blocks on the top and bottom edges with a mixture of Elmer's contact cement and toluene (1:2 v/v), dry overnight at RT, and section at 1.8 μm with a glass knife on a Sorvall MT 2-B ultramicrotome.
- 12. Collect ribbons, six to 10 sections long, on a teflon-coated slide, then transfer to a large drop of freshly boiled dH_20 on a slide coated with 0.5% gelatin. Expand ribbons, and dry on a hot plate at 65°C.
- 13. Heat slide carefully for 20 sec over an alcohol flame to secure the sections.
- 14. Stain sections by floating the slide with 1% toluidine blue O in 0.05 M borate buffer at pH 9 for 4 to 8 min at 65°C.
- 15. Rinse slide briefly with each of dH_20 , 95% ethanol, 100% ethanol, and xylene.
- 16. Affix a coverslip with Permount.

c. Modified Cleared-Pistil Technique

Following find the steps for the modified cleared-pistil technique (Crane and Carman, 1987; C. B., do Valle, personal communication, 2002). *Note:* Pistils should remain in each of the following solutions for at least 30 min:

- 1. Fix inflorescence in 6:3:1 (v/v/v/) 95% ethanol:chloroform:glacial acetic acid for 24 h.
- 2. Store in 75% ethanol at -20° C.
- 3. Transfer pistils in 95% ethanol $\rightarrow 2:1 (v/v) 95\%$ ethanol:benzyl benzoate $\rightarrow 1:2 (v/v) 95\%$ ethanol:benzyl benzoate $\rightarrow 2:1 (v/v)$ benzyl benzoate:dibutyle phthalate (tetratogen) clearing solution.
- 4. Caution! Avoid contact with skin.

- 5. Remove solution carefully from pistil-containing vials by a fine bent Pasteur pipette. Some solution may remain with the pistil, but remove as much as possible without sucking the pistil into the pipette.
- 6. Place cleared pistils on a microscope slide in a sagital optical section between two coverslips.
- 7. Fill spaces with the clearing solution.
- 8. A third coverslip is then placed on top, resting on the two other coverslips. Avoid formation of air bubbles.

Note: Time in clearing solution may be species specific. *Brachiaria* requires a longer time than *Panicum*.

VI. FLUORESCENCE IN SITU HYBRIDIZATION (FISH)

Fluorescence *in situ* hybridization is a powerful technique for detecting specific nucleic acid sequences and localizing highly repetitive DNA sequences in the specific regions of the chromosomes. Since the publication of Gall and Pardue (1969), numerous modifications and refinements of techniques have facilitated the diagnosis and identification of chromosomal aberrations, particularly for human and animal chromosomes (Viegas-Pequignot et al., 1989; Lichter et al., 1990).

The application of *in situ* hybridization technique in plants has lagged behind compared to its use in mammalian cytogenetics. The main handicap of utilizing this technique for plant chromosomes is obtaining high-frequency mitotic metaphase cells without cell wall and cytoplasmic debris. These obstacles hinder hybridization of low-copy-number sequences to the chromosomes (Gustafson, Butler, and McIntyre, 1990). *In situ* hybridization technique has been used to identify chromosomes in several plant species, particularly in wheat and its allied genera (Bedbrook et al., 1980; Hutchinson and Miller, 1982; Teoh, Hutchinson, and Miller, 1983; Rayburn and Gill, 1985a,b; Schubert and Wobus, 1985; Shen, Wang, and Wu, 1987; Visser et al., 1988; Huang, Hahlbrock, and Somssich, 1988; Bergey et al., 1989; Le, Armstrong, and Miki, 1989; Maluszynska and Schweizer, 1989; Mouras et al., 1989; Schwarzacher et al., 1989; Skorupska et al., 1989; Gustafson, Butler, and McIntyre, 1990; Mukai, Endo, and Gill, 1990; Schwarzacher and Heslop-Harrison, 1991; Dhaliwal, Martin, and Gill, 1991; Mukai and Gill, 1991; Friebe et al., 1991a; Lapitan, Canal, and Tanksley, 1991; Wang, Lapitan, and Tsuchiya, 1991; Leitch et al., 1996; Griffor et al., 1991; Leitch et al., 1996; Singh, Kim, and Hymowitz, 2001; Hayasaki, Morikawa, and Leggett, 2001).

Singh, Kim, and Hymowitz (2001) developed a procedure to detect repetitive DNA sequences on somatic metaphase chromosomes for soybean (2n = 40) root tips using fluorescence *in situ* hybridization after consulting the above published results for the mammalian as well as plant chromosomes.

A. Chromosome Preparation

Following are steps for chromosome preparation (Ambros, Matzke, and Matzke, 1986; Geber and Schweizer, 1988; Murata, 1983; Griffor et al., 1991; Singh, Kim, and Hymowitz, 2001).

1. Collection, Pretreatment, and Fixation of Root

- 1. Germinate soybean seeds in a sandbox or in vermiculite in the greenhouse (for cereals, seeds can be germinated on a moist filter paper in a Petri dish). The sandbox should be kept moist.
- 2. After about 7 days, collect actively growing root tips (3 to 5 mm) from secondary roots.
- Pretreat root tips in 0.1% colchicine for 2 h at RT. Often, metaphase chromosomes are found to be clumped. Sometimes, 0.05% 8-hydroxyquinoline at 16°C for 4 to 5 h yields a higher mitotic index. For cereal chromosomes, pretreatment with ice-cold water 18 h (barley) to 24 h (wheat) is recommended.

4. Fix root tips in a freshly prepared fixative of methanol, acetic acid, and chloroform (3:1:1) for 2 to 3 h at -20°C or in 3 (95% ethanol):1 (propionic acid) or 3 (95% ethanol):1 (glacial acetic acid). However, roots can be stored for 1 to 2 weeks at -20°C.

2. Preparation of Buffer

- 1. Preparation of CA-SC buffer:
 - a. 0.01 M citric-sodium citrate buffer pH 4.5 to 4.8 (CA-SC)
 - b. Prepare from 10 × stock solution consisting of three parts 0.1 M tri-sodium citrate and two parts 0.1 M citric acid. Check pH it should be in the range of 4.5 to 4.8.
- 2. Preparation of SSC buffer:
 - a. $20 \times SSC$
 - i. 701.2 g NaCl
 - ii. 52.8 g Na citrate
 - iii. 4 L ddH₂O, mix, filter, sterilize, and adjust pH 7.4 (stock)
 - b. $4 \times SSC + 0.2\%$ Tween-20 (designated as $4 \times SSC$)
 - i. 400 mL 20 × SSC
 - ii. 1596 mL ddH20
 - iii. 4 mL Tween-20
 - c. $2 \times SSC$
 - i. $100 \text{ mL } 20 \times \text{SSC}$
 - ii. 900 mL ddH_20

3. Enzyme Solution

The enzyme solution should be prepared fresh before use:

- 0.02 g cellulase (2%) cellulose, Onozuka R-10 (from Yakult, Tokyo, Japan)
- 0.01 g pectinase (1%) pectinase, Sigma (cat. # P-2401)
- Dissolve in 1.0 mL 0.01 M CA-SC buffer in an Eppendorf tube

4. Preparation of Slides

In between washes, cells are spun down at 2500 rpm for 5 min in an Eppendorf tube. Remove the supernatant carefully with a Pasteur pipette:

- 1. Wash root tips three times for 5 min each in 0.01 M CA-SC buffer in a spot plate at RT.
- 2. Place a Petri cover on some crushed ice, and dissect only 1 to 2 mm in length the meristematic region of the root tips (cream colored at the very tip).
- 3. Transfer the tips into an Eppendorf tube with 0.01 M CA-SC buffer on ice.
- 4. Spin down in a microfuge for a few seconds. Withdraw the supernatant.
- 5. Add 1.0 mL of enzyme solution, seal with parafilm, and incubate in a 37°C water bath for 1 h and 30 min.
- 6. Aspirate the tips in the enzyme solution by drawing them into a Pasteur pipette several times. The tips should break apart, and the solution should appear cloudy.
- 7. Again, incubate in a 37°C water bath for 10 to 15 min.
- 8. Spin down as stated above.
- 9. Add about a milliliter of 0.01 M CA-SC buffer, and let it sit on ice for 35 to 45 min.
- 10. Spin (3000 rpm, 3 min), resuspend the pellet in 75 mM KCl (150 mM KCl has also produced good results), and let it sit for 10 min at RT.
- 11. Spin (5000 rpm, 3 min) and suspend in fresh fixative (three parts 100% ethanol:one part glacial acetic acid) and let it stand 17 to 20 min at RT.
- 12. Spin (5000 rpm, 3 min), and suspend pellet in approximately 20 to 25 drops of fixative from a Pasteur pipette.

- 13. Place one drop of this suspension per slide using a Pasteur pipette. *Note*: Use slides that are ice-cold from -70°C freezer. Drop from about 3 cm above on the flat slide and blow gently on slide to aid in spreading.
- 14. Air dry slides. Once dried, they can be stored in a slide box sealed in a bag at -70° C for several weeks.

Note: Slides can be prepared by the squash method. Cut the root tip and squash on a clean slide under a clean coverslip with a drop of 45% acetic acid.

B. Fluorescence in Situ Hybridization Procedure

The following protocol provided by F. Ahmad, describes the fluorescence *in situ* hybridization procedure (Ahmad et al., 1999). Check slides for good metaphase cells by mounting in 0.2 μ g/mL Sigma DAPI in 1 × PBS, pH 7.0, or in 0.4 μ g/mL Sigma propidium iodide in 1 × PBS, pH 7.0. Mark the position of cells from the scale on the microscope stage. Place the slides in a -80°C freezer for 5 min and remove the cover glass with a razor blade. Destain the slides by immersing in 45% acetic acid for 1 to 2 sec, rinse in water, and air dry. *Scan slides for prometaphase and metaphase spreads using a phase-contrast lens. Remove the coverslip after dipping the slides in liquid nitrogen for a few seconds. Coverslips can also be removed by dry ice. Prepare eight slides.*

1. Prehybridization Method

- 1. Treat slides in a staining jar in 45% acetic acid for 10 min at RT, and air dry for 1 to 2 h or overnight at RT.
- 2. Turn on water bath at 70°C, prepare humid chamber.
- 3. Take 24 μL (from 1 mg/mL stock) RNase stock solution + 776 μL 2 × SSC (800 mL). Add 100 μL RNase to each slide, and cover with a colored plastic coverslip (24 × 30 mm). Incubate at 37°C for 1 h in a humid chamber, and drain slides. (*Caution! Slide should not get dry.*)
- 4. Treat slides in 70% formamide for 2 min at 70°C, and drain slides.
- Dehydrate slides at -20°C in 70% (5 min), 80% (5 min), 95% (5 min), and 100% (30 sec) ethanol. Air dry at RT for 1 to 2 h.

2. Hybridization

- 1. Probe (50 µL)
 - a. 200 ng rDNA; [internal transcribed spacer (ITS) region of nuclear ribosomal DNA probe (ITS1,5.8S, ITS2; approximately 700 nucleotides)] (5 μ L) + 19 μ L dH₂O sterilized H₂O.
 - i. Denature on PCR at 95°C for 5 min or in boiling water.
 - ii. Keep on ice 5 min (immediately).
 - iii. Add 5 μ L 10× dNTP + 20 μ L 2.5 × random primer solution.
 - iv. Mix (tapping by bottom).
 - v. Add 10 µL Klenow from the freezer.
 - vi. Mix (tapping by bottom).
 - vii. Incubate at 37°C for 3 h (Bio Prime[™] DNA Labeling System, Life Technologies, Inc., Gaithersburg, MD, cat. No.: 18094–011; lot no. EHDOO1)
- 2. Preparation of hybridization solution
 - a. Each slide needs 50 μL of solution.
 - i. 200 µL 50% formamide
 - ii. 80 μL 50% dextron sulfate
 - iii. 53 μL TE buffer
 - iv. 40 μL 20 \times SSC
 - v. 17 µL salmon sperm (ss) DNA
 - vi. 10 µL probe
 - b. Mix hybridization solution in 1.5 mL tube, give quick spin, and transfer into three PCR tubes (two contain 150 μ L, and the third contains 100 μ L).

- c. Denature at 80°C for 10 min (during this break, prewarm the humidity chamber), keep on ice for 10 min, give quick spin, mix, and keep on ice.
- d. Apply 50 µL denatured hybridization solution on each slide, cover, and keep at 80°C for exactly 8 min, move humid chamber with slides quickly to 37°C for overnight for hybridization [at this time, prepare 4 × SSC + Tween-20 (will be known as 4 × SSC) and 2 × SSC]
- 3. Posthybridization (next day)
 - a. Turn on water bath (40°C), place two Coplin jars (*designated as jar*) with 2 × SSC, one with 35 mL 2 × SSC, one tube containing 35 mL formamide, and one bottle with 400 mL 2 × SSC.
 - b. Remove coverslip, and drain slides.
 - c. Wash slides in 2 × SSC at 40°C for 5 min twice (discard used 2 × SSC, and fill with fresh 2 × SSC) (during second wash, prepare 5% BSA: dissolve 0.24 g albumin bovine in 4.8 mL 4 × SSC), and then drain slides.
 - d. Treat slides in 50% formamide for 10 min at 40°C, then drain slides.
 - e. Wash in $2 \times SSC$ for 5 min twice at 40°C, wash third time in $2 \times SSC$ at RT, and drain slides.
 - f. Wash slides in $4 \times SSC$ at RT for 5 min (*during this time, prepare humidity chamber*), and then drain slides.
 - g. Add 200 μL 5% BSA, cover, treat for 5 min at RT (during this time, prepare 4 μL [fluorescein-Avidin, DCS (FITC) + 796 μL 5% BSA = 800 μL; 10 μg/mL], and drain slides.
 - h. Add 100 μ L FITC to each slide, cover, incubate for 1 h at 37°C (*during this time, place three jars with 4 × SSC and one bottle with 4 × SSC in water bath at 40°C*), and drain slides.
 - i. Wash slide three times at 40°C in 4 × SSC for 5 min each [during this period, prepare 1600 μL in 2 mL tube (80 μL goat serum + 1520 μL 4 × SSC)], and drain slides.
 - j. Add 200 μ L goat serum to each slide, incubate for 5 min at RT in a humid chamber [*during this period, prepare 800 \muL: 8 \muL biotinylated anti-avidin (10 \mug/mL) + 792 \muL 4 × SSC], then drain slides.*
 - k. Add 100 µL biotinylated anti-avidin, cover, incubate at 37°C for 1 h, and drain slides.
 - 1. Wash slide three times at 40° C for 5 min, then drain slides.
 - m. Repeat Step g.
 - n. Repeat Step h.
 - Repeat Step i [during second wash, prepare propidium iodide (PI): 16 μL PI (1 mg/mL) + 784 μL 2 × SSC].
 - p. Add 100 µL PI, cover, incubate in dark for 20 min at RT in humid chamber, and drain slides.
 - q. Rinse slides briefly in $2 \times SSC$, drain, and wipe the back of the slides.
 - r. Add 50 μ L (two drops) Vectashield mounting medium for fluorescence (Vector H-1000), cover with 22 × 32 mm cover glass, and store in the dark overnight.

3. Observation and Photography

Use a Zeiss Axioskop or an Olympus fluorescence microscope. For the Zeiss Axioskop:

- 1. Observe slides by using D (dark)-field.
- 2. Turn on the UV power supply.
- 3. Use 63× oil lens.
- 4. There are two levels of UV: 50% for general observation and 100% for photography using 100× oil lens.
- 5. Use neutral filter 0.1 m (-90% of light) to 0.3 m (60% light).
- 6. Use 400 ASA slide Fuji film.

C. Genomic in Situ Hybridization (GISH)

The genomic *in situ* hybridization technique is a powerful tool used to distinguish the genome of one parent from the other by preferential labeling of the chromosome of one parent. It has proved to be a boon to cytogeneticists in identifying precisely the inserted region in the recipient parent from the alien species and in examining the evolutionary relationship of crops.

The protocol for GISH is essentially the same as for the FISH, except for the blocking genomic DNA. The GISH protocol is defined to eliminate most of the cross-hybridization between total genomic DNA from the two species. Optimum results are obtained when blocking DNA exceed the concentration of probe DNA by 100-fold. Blocking genomic DNA fragments of 100 to 200 bp length are obtained by autoclaving the total genomic DNA from each parental species.

For GISH mixture, the yield of labeled probe can be increased by use of carrier DNA. The role of block and carrier DNA are combined in addition to unlabeled total genomic blocking DNA at 35 times the concentration of probe DNA immediately after stopping the NT reaction with EDTA. The BIONICKTM Labeling System is used for biotin-labeled DNA probes by nick translation [BIONICKTM Labeling System: Life Technologies, Gaithersburg, MD (cat. no. 18247–015; lot no. KKF712)].

D. Multicolor Genomic in Situ Hybridization (McGISH)

Multicolor GISH (Fukui, Ohmido, and Khush, 1994) is an excellent cytogenetic tool for simultaneous discrimination of each genome and identification of diploid progenitors of allopolyploids, simultaneous mapping of different DNA sequences, physical ordering of multiple probes in a single chromosome, genome allocation of the gene of interest, detection of chromosomal aberrations, and examination of chromosome organization in interphase nuclei (Mukai, 1996). Shishido et al. (1998) developed multicolor genomic *in situ* hybridization to identify somatic hybrids between *Oryza sativa* cv. "Kitaake" (AA, 2n = 24) and *O. punctata* (BBCC, 2n = 48), and the progeny rescued from embryo culture. The procedure provided by K. Fukui, is as follows:

- Label total DNA of diploid rice species, O. sativa (AA), O. punctata (BB), and O. officinalis (CC) with biotin-16-dUTP (Roche Diagnostic, R. Hoffman-LaRoche, Ltd., Basel Switzerland) or digoxigenin-11-dUTP (Roche Diagnostic, R. Hoffman-LaRoche, Ltd., Basel Switzerland) by the standard random primed labeling protocol.
- 2. Denature hybridization mixture (100 ng labeled probe/slide + equal parts of 50% formamide and $2 \times SSC$) for 10 min at 90°C, and cool immediately on ice (0°C).
- 3. Denature chromosome spreads in 50% formamide/2×SSC for 6 min at 70°C with the hybridization mixture (Fukui, Ohmido, and Khush, 1994), and then hybridize for 3 to 4 days at 37°C.
- 4. Wash twice in 2 × SSC, once in 50% formamide/2 × SSC, and once in 4 × SSC, each for 10 min at 40°C.
- Apply a drop of Fluorescin-avidin (1% Vector Laboratories, CA; vector@vectorlabs.com; 800-227-6666) + 1% bovine serum albumin (BSA) in BT buffer (0.1 *M* sodium hydrogen carbonate, 0.05% Tween-20, pH 8.3) on chromosome spread.
- 6. Incubate at 37°C for 1 h.
- 7. Wash slides three times (10 min each) at 37°C.
- 8. Amplify again (repeat Step 5).
- Apply a drop on chromosome spread anti-digoxigenin-rhodamine in 5% goat biotinylated antiavidin and anti-digoxigenin-rhodamine in 5% goat serum in BT buffer, and then incubate at 37°C for 1 h.
- 10. Wash with BT buffer three times at 37°C (5 min each).
- 11. Apply a drop of 1% fluorescin-avidin and anti-sheep-Texas Red (1% Vector) in 1% BSA in BT buffer to chromosome spread.
- 12. Incubate at 37°C for 1 h.
- 13. Wash twice with BT buffer and once with $2 \times SSC$ at $40^{\circ}C$ (10 min each).
- 14. Counterstain chromosomes with 1 μ g/mL 4',6-diamidino-2 phenylindole (DAPI) in an antifadant solution (Vector Shield, Vector).

Note: Block chromosome spread with 5% bovine serum or goat serum albumin in BT buffer at 37°C for 5 min before each immunocytochemical step.

E. Primed in Situ (PRINS) DNA Labeling

The PRINS DNA labeling technique, first described by Koch et al. (1989), is an alternative to FISH for the detection of nucleic acid. This techniques involves labeling chromosomes by annealing an oligonucleotide DNA primer to the denatured DNA of chromosomes spread on slide glass and extending it enzymatically *in situ* with incorporation of labeled nucleotides. The PRINS technique has been used extensively in human cytogenetics for mapping of repetitive and low-copy sequence, for chromosome identification, for detection of aneuploidy in sperm cells, for the analysis of the human chromosome complement of somatic hybrids, for specifying chromosome rearrangements by combination of PRINS labeling of chromosome-specific alphoid sequences, and for chromosome painting (Pellestor et al., 1996; Gosden, 1997).

The reliable and reproducible detection of single-copy sequence below 10 kb in plants with large genome has been difficult with FISH. Menke, Fuchs, and Schubert (1998) developed a procedure of PRINS for detection of repetitive and low-copy sequences on plant chromosomes and protocol is provided by I. Schubert. For chromosome preparation, follow the protocol used for FISH:

- 1. Prior to PRINS, wash slides three times in $2 \times SSC$ for 5 min at RT.
- 2. Treat slides with RNase (50 μ g/mL in 2 × SSC) for 40 min at 37°C. Subsequently a fill-in reaction is carried out to reduce background signals caused by nicks within the chromosomal DNA, which may induce polymerase activity at sites of free 3' OH-ends.
- To reduce background signals, wash slides first in 2 × SSC and equilibrate in *Taq*-polymerase buffer (10 mM Tris-HCl, pH 8.4, 50 mM KCl, 1.5 mM MgCl₂) for 5 min.
- 4. Reaction mixture (20 μ L):
 - a. $1 \times PCR$ buffer (Boehringer)
 - b. 1.5 mM MgCl₂
 - c. 100 μM of each dATP, dCTP, dGTP
 - d. 100 µM of 2',3'-dideoxy (dd)TTP
 - e. 2 U of Taq-DNA polymerase (Boehringer)
- 5. Drop the reaction mixture on a slide, cover with a coverslip, and seal with Fixogum rubber cement.
- 6. Heat slides at 93°C for 90 sec, followed by 72°C for 20 min in a wet chamber (Zytotherm, Schutron). Before labeling, wash chromosomes and equilibrate again.
- 7. Labeling mixture (25 μ L):
 - a. $1 \times PCR$ buffer
 - b. 1.5 mM MgCl₂
 - c. 100 μ M of each dATP, dCTP, dGTP, 75 μ M dTTP
 - d. 25 μ M of digoxygenin-11-dUTP or fluorescein-12-dUTP (Boehringer)
 - e. 4 µM of each of the corresponding oligonucleotide primers and 2.5 U of Taq-DNA polymerase
- 8. Seal the coverslip, and denature chromosomes for 2 to 3 min at 93°C. Anneal primers at 55 to 60°C for 10 min, and extend at 72°C for 40 min.
- 9. Wash slides twice in $4 \times SSC$, 0.1% Tween-20 for 5 min at $42^{\circ}C$ in order to stop the reaction.
- 10. Counterstain the preparation labeled with FITC-dUTP with propidium iodide/4',6-diamidino-2phenylindole dihydrochloride (PI/DAPI, 1 μg/mL each in antifade), and examine immediately. Detect Dig-dUTP with anti-Dig-FITC-Fab fragments from sheep (2 μg/mL, Boehringer).
- 11. Incubate samples in blocking solution ($4 \times SSC$, 0.1% Tween-20, 3% BSA) at 37°C for 30 min.
- 12. Subsequently, apply Fab-fragments in detection buffer (4 \times SSC, 0.1% Tween-20, 1% BSA) for 50 min at 37°C.
- 13. Remove unspecifically bound conjugates by washing slides in wash buffer three times for 5 min at 42°C.
- 14. After counterstaining, described above, examine slides using a Zeiss epifluorescence microscope.

F. Fluorescence in Situ Hybridization on Extended DNA Fibers: Fiber-FISH

This protocol has been provided by S.A. Jackson, R.M. Stupar, and J. Jiang. The Fiber-FISH technique is a powerful cytological tool used to analyze large repetitive regions and increase the

resolution of FISH analyses to a few kb in the higher eukaryotic genomes (Stupar et al., 2001). It can be used to gauge the distances between adjacent clones up to \sim 500 kb and to measure repetitive loci up to \sim 1.7 mb. Combined with metaphase and interphase nuclei analysis, this tool allows molecular cytogeneticists to map loci to specific chromosomes and to determine the distance between loci from a few kb up to several mb.

1. Protocol 1: Fiber-FISH on Extended Nuclear DNA Fibers

a. Isolation of Plant Nuclei

Isolation of plant nuclei is the same as for the preparation of high-molecular-weight DNA embedded in agarose plugs. However, we suspend in 50% glycerol to store at -20° C. Almost all of the published protocols call for 20 µm or 30 µm filtrations, but this can be omitted with most of the plant materials in order to obtain the maximum number of nuclei. If there is too much debris in the suspension or on the slides, the last (20 µm) filtration can be added:

- 1. Freeze 2 to 5 g fresh leaf material in liquid nitrogen, and grind to a fine powder with a precooled (-20°C) mortar and pestle.
- Transfer powder to a 50 mL centrifuge tube, add 20 mL chilled nuclei isolation buffer (NIB), and mix *gently* (make sure to break up the clumps) on ice for 5 min (in an ice bucket on a shaker). [NIB: 10 mM Tris-HCl pH 9.5, 10 mM EDTA, 100 mM KCl, 0.5 M sucrose, 4.0 mM spermidine, 1.0 mM spermine, 0.1% mercapto-ethanol. Prepare a large stock and store it in a refrigerator (4°C). Mercapto-ethanol should not be included in the stock. It can be added just before use.]
- 3. Filter through nylon mesh: 148 and 48 μm sequentially, into cold (on ice) 50 mL centrifuge tubes using a cooled funnel. (Nylon filters were obtained from Tetko Inc., Lancaster, NY, U.S.A.)
- 4. Add 1 mL NIB containing 10% (v/v) Triton X-100 (premixed), and *gently* mix the filtrate. The final concentration of Triton X-100 should be 0.5%. It removes any chloroplast contamination. Centrifuge at 2000 × g for 10 min at 4°C. Decant the supernatant. If pellet is very small, skip the further cleaning steps and move directly to Step 7. Otherwise, resuspend the large pellet in 20 mL NIB (with mercapto-ethanol added).
- 5. Filter through nylon mesh: 48 and 30 μm sequentially (optional), into cold (on ice) 50 mL centrifuge tubes.

Note: A 22 µm filter step will lose a lot of nuclei. However, this makes the nuclei cleaner.

- Add 1 mL NIB containing 10% (v/v) Triton X-100 (premixed), and *gently* mix the filtrate, as in Step 4. Centrifuge at 2000 × g for 10 min at 4°C.
- 7. Decant the supernatant, and resuspend the pellet in 200 μ L to 5 mL of 1:1 NIB:100% glycerol (neither mercapto-ethanol nor Triton X-100 added), depending on the amount of nuclei harvested (concentration ~5 × 10⁶ nuclei/mL, can be checked by staining with DAPI and examining under a microscope). Store at -20°C.

b. Extension of DNA Fibers

Extending the fibers is a critical step in Fiber-FISH. There are several methods of extending the fibers. The dragging method with a coverslip seems to give the most uniform results. When dragging, it is imperative that it be done slowly and smoothly. Poly-L-lysine slides obtained from Sigma can be used. These slides are treated so as to promote the adhesion of one or both ends of the DNA molecule. Silinated slides can also be used but seem to generate too much background signal. The calibration of the method should be checked occasionally by using BACs or cosmids of a known length as probes:

1. Identify the nuclei portion in the nuclei stock. The nuclei tend to settle near the bottom of the tube, and the settling process can take a day or longer. The pellet may appear white, and the nuclei often sit right above this bottom film. The color of the nuclei is variable across species and samples, but normally, clean nuclei has a gray/white coloration. Any layers above the nuclei tend to contain

debris. Some people like to mix the nuclei stock prior to slide preparation by gently inverting the Eppendorf tube several times. This is not desirable, as it mixes the debris with the nuclei.

- 2. With a cut P20 pipette tip, pipette 1 to 10 μ L nuclei suspension (1 to 5 μ L/slide depending on the suspension concentration) into ~100 μ L NIB (minus mercapto-ethanol and Triton) in an Eppendorf tube to dilute the glycerol. Gently mix the nuclei with the buffer, and centrifuge at 3000 to 3600 rpm for 5 min. Carefully remove the supernatant with a pipette, leaving only the nuclei pellet.
- Resuspend the nuclei in PBS (the final volume is 2 μL per slide) [PBS: 10 mM sodium phosphate, pH 7.0, 140 mM NaCl].
- Pipette 2 μL suspension across one end of a clean poly-L-lysine slide (Sigma, Poly-Prep, cat. # P0425) and air dry for 5 to 10 min. The nuclei should dry to the point where it appears "sticky," neither wet nor dry.
- Pipette 8 μL STE lysis buffer on top of the nuclei, and incubate at RT for 4 min [STE: 0.5% SDS, 5 mM EDTA, 100 mM Tris, pH 7.0].
- 6. Slowly drag the solution down the slide with the edge of a clean coverslip held just above the surface of the slide. Do not touch the coverslip to the slide surface, as this will drag the nuclei completely off the slide. Air dry for 10 min.
- 7. Fix in fresh 3:1 100% ethanol:glacial acetic acid for 2 min.
- 8. Bake at 60°C for 30 min.
- 9. Slides can be used immediately, and they can be stored in a box for several weeks. But, it is suggested that they be used immediately.

c. Probe Application

- 1. Apply a 10 μ L probe to the slide, then cover with a 22 × 22 mm coverslip, and seal with rubber cement.
- 2. After the cement is dried, place the slide in an 80°C oven for 3 min in direct contact with a heated surface, then for 2 min in a wet chamber prewarmed in the 80°C oven.
- 3. Transfer wet chamber, with the slides, immediately to 37°C overnight. It is recommended to incubate at 37°C for longer periods, up to 3 or 4 days, especially for difficult probes.

d. Probe Detection

1.

Three-layer detection gives a much stronger signal than does the single layer of antibodies. All antibody layers are composed of the antibodies diluted in the appropriate buffers at the concentration specified below:

- Apply 100 μ L antibody to each slide and gently place a 22 × 40 coverslip upon the antibody solution to promote an even spreading. All antibody layers are incubated in a 37°C wet chamber for a minimum of 30 min. Often, the first layer is incubated for up to 45 to 60 min.
- *Notes*: The blocking step using 4 M buffer seems to help reduce some of the background noise. Dry bovine milk from Sigma works the best in the 4 M buffer other substitutes (i.e., Carnation dry milk) tend to reduce the amount of signal. The 4 M and TNB buffers can be prepared at 5× and stored at -20°C. The wash solutions, 4 T and TNT, can be prepared at 20× and 10×, respectively, and stored at RT.

One-Color Detection Protocol		Time, min (total)
i.	Wash in $2 \times SSC$	5
ii.	Wash in $2 \times SSC \ 42^{\circ}C$	10
iii.	Wash in $2 \times SSC$	5
iv.	Wash in 1×4 T	5
v.	Incubate at 37°C in 4 M	30
vi.	Wash in 1×4 T	2
vii.	Incubate FITC-avidin (1 µL antibody stock/100 µL TNB buffe	er) 30
viii.	Wash three times in $1 \times TNT$	5 (15)
ix.	Incubate Biotin anti-avidin (0.5 µL/100 µL TNB buffer)	30

	х.	Wash three times in $1 \times TNT$	5 (15)
	xi.	Incubate FITC-avidin (1 µL/100 µL TNB buffer)	30
	xii.	Wash three times in $1 \times TNT$	5 (15)
	xiii.	Wash two times in $1 \times PBS$	5 (10)
	xiv.	Add 10 µL Prolong (Molecular Probes) or Vectashield (Vector Labs	s), cover with a 22 \times
		30 mm coverslip, and squash	
	Note	es: All 30 min incubation periods are at 37°C.	
	i.	Antibodies:	
		A. FITC-avidin, 1 µL per 100 µL buffer	
		B. Biotin anti-avidin, 0.5 µL per 100 µL buffer	
		C. Mouse anti-dig, 1 µL per 100 µL buffer	
		D. Dig anti-mouse, 1 µL per 100 µL buffer	
		E. Rodamine anti-dig, 1 to 2 μ L per 100 μ L buffer	
	ii.	Solutions:	
		A. 4M: 3 to 5% nonfat dry milk [Sigma, cat. # M7409] in $4 \times$ SSC	l ,
		B. 4T : 4 × SSC, 0.05% Tween-20	
		C. TNB: 0.1 M Tris-HCl pH 7.5, 0.15 M NaCl, 0.5% blocking	reagent (Boehringer
		Mannheim)	
		D. TNT: 0.1 M Tris-HCl, 0.15 M NaCl, 0.05% Tween-20, pH 7.5	
		E. PBS : 0.13 M NaCl, 0.007 M Na ₂ HPO ₄ , 0.003 M NaH ₂ PO ₄	
2.	Two-Co	olor Detection Protocol Time,	, min (total)
	i.	Wash in $2 \times SSC$	5
	ii.	Wash in $2 \times SSC \ 42^{\circ}C$	10
	iii.	Wash in $2 \times SSC$	5
		Wash in 1×4 T	5
	v.	Incubate at 37°C in 4 M	30
		Wash in 1×4 T	2
	vii.	Incubate FITC anti-dig (1 μ L antibody stock/100 μ L TNB buffer) + T	exas Red streptavidin
		(1 µL/100 µL), 37°C for 30 min	
		Wash three times in $1 \times TNT$	5 (15)
	ix.	Incubate FITC anti-sheep (1 μ L/100 μ L TNB buffer) + Biotinylated a	anti-avidin (1 µL/100
		μL), 37°C for 30 min	
		Wash three times in $1 \times TNT$	5 (15)
		Incubate Texas Red streptavidin (1 μ L/100 μ L TNB buffer), 37°C f	
		Wash three times in $1 \times TNT$	5 (15)
		Wash two times in $1 \times PBS$	5 (10)
		Add 10 μ L Vectashield (Vector Labs), cover with a 22 \times 30 mm co	verslip, and squash.
	Note		
	i.	Preparation of antibody stocks:	
		A. FITC anti-sheep comes ready to use and is stable at 4° C.	
2	DII	B. FITC anti-dig needs 1 mL water resuspension.	
3.		lysine slide preparation	
	i. .:	Boil slides in $5 M$ HCl for 2 to 3 h.	
	ii.	Rinse thoroughly with dH_2O , then air dry.	000 0:
	iii.	Incubate overnight in filtered 10^{-6} g/mL poly-D-lysine (MW = 350,)	000, Sigma).
4	iv.	Rinse thoroughly.	
4.	i.	lanation 30 min. 1:1 HCl:methanol	
	ii.	Overnight 18 M sulfuric acid	
	iii.	Eight to 10 washes in ddH_2O	
	iv.	10 min boiling ddH_2O	
	1 V.	10 mm boming dull20	

- v. 1 h 10% 3-aminopropyltriethoxysilane in 95% ethanol
- vi. Rinse several times in ddH₂O
- vii. Wash in 100% ethanol

viii. 80 to100°C overnight before use

Note: The Poly-Prep slide from Sigma (cat. #P0425) is ready to use, is available, and can be used instead of the above procedure.

2. Protocol 2: Fiber-FISH Using BAC and Circular Molecules as Targets

a. Prior to Preparing Slides

- 1. Label appropriate probes (biotin and dig).
- 2. Miniprep BAC DNA (Solution 1, 2, 3 method followed by IPA precipitation; use 20 μL water for resuspension).
- 3. Silanize 22×22 coverslips by dipping in Sigmacote for 10 min, then air drying.

b. Slide Preparation

- 1. Prepare wet chamber at 37°C, and turn slide warmer up to 60°C.
- 2. Dilute BAC DNA (with cut P20 pipette tips) to appropriate level (dilute 1 μ L BAC into 9 μ L water). Add all 10 μ L of diluted BAC to Poly-Prep slide (Sigma #P0425).
- 3. Add 15 μ L of FISH lysis buffer to BAC drop. Allow drop to spread. Let this sit at RT for ~5 min. Add water to the slide if it dries.
 - a. FISH lysis buffer:
 - i. 2% Sarkosyl
 - ii. 0.25% SDS
 - iii. 50 mM Tris (pH 7.4)
 - iv. 50 mM EDTA (pH 8.0)
- 4. Gently place ("drop") a silanized coverslip directly over the liquid (use a tweezer to avoid air bubbles).
- 5. Transfer slides to slide warmer. Allow slides to "bake" for 15 min. At this point, one should see the liquid begin to recede.
- 6. Place slides in 3:1 (ethanol: glacial acetic acid), wait 1 min. Gently shake slide to promote removal of the cover slip. Once cover slip falls off, transfer slides to new container of 3:1 and incubate for 1 min 30 sec. Transfer slides back to slide warmer for an additional 15 min.
- 7. Add probe, denature, and detect as in nuclear fiber-FISH.

3. Protocol 3: Staining Fibers (Yo-Yo Staining)

- 1. Prepare slides for fiber analysis (as in BAC or nuclear fiber-FISH protocols).
- 2. Dilute Yo-Yo (molecular probes) in PBS following manufacturer's direction.
- 3. Add 100 μ L of the Yo-Yo dilution to the slide, and add a coverslip. Store in a dark place at RT for 10 to 20 min.
- 4. Wash three times in PBS (5 min each).
- 5. Short dry. Add anti-fade (Vectashield).
- 6. View slides.

4. Source of Chemicals

See Appendix 2-I.

VII. TOTAL DNA EXTRACTION — PLANT GENOMIC DNA

For more information on plant DNA preparation, see Dellaporta, Woods, and Hicks (1983).

A. Protocol 1

1. Solutions

- 1. Miniprep salts buffer, pH 8.0 per 1 L:
 - a. 29.02 g 500 mM NaCl
 - b. 12.1 g 100 mM Tris base
 - c. 18.6 g 50 mM EDTA disodium
 - d. Adjust volume to 1000 mL final vol, pH 8.0 with HCl, autoclave
- 2. Complete buffer (per sample), and make just before use:
 - a. 50 mL Miniprep salts buffer
 - b. 10 mM *o*-phenanthroline (mol wt 198; is prepared by dissolving 90 mg *o*-phenanthroline in 500 μL ethenol)
 - c. 0.5 g SDS (heat gently at 37°C to dissolve)
- 3. Immediately before use, add the following:
 - a. 35 μ L β -mercaptoethanol
 - b. 100 μ L of 10 mg/mL ethidium bromide stock solution (remember, this amount is needed for each sample, so multiply each amount by the number of samples extracted)
 - c. 5 M potassium acetate (mol wt 98) 98 g to 200 mL final volume in distilled water and no pH adjustment
- 4. Wash buffer per 500 mL
 - a. 1.51 g 25 mM Tris base
 - b. 1.86 g 10 mM EDTA disodium
 - c. to 500 mL final volume, pH 8.0 with HCl, autoclave

2. Procedure

- 1. Take ice and plastic bags, and collect fresh 2 g leaves from the growth chamber or greenhouse-grown seedlings.
- 2. Grind young leaves or seedlings into fine powder using a mortar and pestle in liquid nitrogen.
- Add 25 mL of complete buffer to mortar, and continue grinding until homogenized. Pour into sterile GSA bottle. Wash pestle with additional 25 mL aliquot of complete buffer. Pour into bottle, and cap.
- 4. Heat bottle at 65°C for 10 min, swirling occasionally.
- 5. Add 17 mL 5 M potassium acetate to bottle. Swirl to mix. Place in ice for 15 min.
- 6. Centrifuge at 9000 rpm for 10 min in JA14 rotor.
- Filter supernatant through mira cloth into a clean (sterile) GSA bottle containing 33 mL cold isopropanol. Place in -20°C freezer for 20 min (should see DNA spool out).
- 8. Spool out DNA, if possible, using a glass stirring rod. Dissolve DNA in 9 mL of wash buffer *(in an autoclaved plastic tube or sterile 30 mL COREX tube)*. If DNA will not spool, spin for 10 min at 9000 rpm, decant supernatant, dry pellets under airflow in hood, and redissolve in 9 mL wash buffer on ice. Be sure pellets are redissolved well before proceeding.
- 9. Add 500 μ L ethidium bromide (10 mg/mL stock).
- 10. Add 10 g CsCl, cover tightly with parafilm, and invert to mix. CsCl must go into solution. Incubate 1 h at 4°C.
- 11. If there is any debris, centrifuge at 8000 rpm for 15 min in the JA17 rotor. Rinse the rotor after using with CsCl.
- 12. If there is floating debris, filter supernatant through mira cloth, and collect in 15 mL graduated conical tubes. Protect from light. Adjust refractive index to 1.3870–1.3885 using the refractometer. Add wash buffer if the index is too high, and add solid CsCl if it is too low. (It is usually high).
- 13. Fill and balance 13.5 mL polyallomer ultracentrifuge tube (fit the type 65 or 75 Ti rotors) as follows: Pipette the solution into each tube using an appropriate syringe with an 18 gauge needle. (Do not use Pasteur pipette, because the tip breaks off too easily.) Balance the tube using the syringe needle to add or remove small amounts of the solution. If you do not have enough sample, fill and balance the tubes with a blank CsCl–TNE solution of the proper density. Leave a small

air bubble just below the neck of the tube and have the tubes as usual, check the seal, and recheck the balance.

- 14. Load the tubes properly in the Type 75 Ti rotor with the red tube spacer on top.
- 15. Spin at 20°C and 50,000 rpm for 40 h, or use 60,000 rpm for 22 h, but do not exceed 60,000 rpm. Use deceleration rate 9.
- 16. Remove tubes from rotor. Clean the rotor and parts after the run by rinsing out well with tap and distilled water, then drain. Cesium chloride is corrosive.
- 17. Visualize band with hand UV light if necessary. Collect band with 5 mL syringe and 18 gauge needle. Transfer to sterile 15 mL COREX tube.
- 18. Remove the ethidium bromide from the DNA by extracting with butanol saturated with CsCl. (Butanol saturated with CsCl is prepared by adding a few grams of solid CsCl to 100% butanol. As the butanol is used, more can be added to the CsCl.) Add two volumes of the butanol to the DNA, and mix well by vortexing. Spin at 1000 rpm for 1 min in the clinical centrifuge or the JA 17 rotor. Remove the upper butanol layer, and discard into the organic waste. Repeat the extraction until all of the ethidium bromide purple color has been removed. This will take about three to four extractions.
- 19. Can dialyse the DNA against $1 \times \text{TNE}$ to remove the CsCl, but it is faster to ethanol precipitate: a. Add two volumes of distilled water. (This will dilute the CsCl so that it does not precipitate out.)
 - b. Add six volumes of ethanol (i.e., six volumes of the original DNA volume). *Note*: Na acetate is not needed, as CsCl salt concentration is already high.
 - c. Incubate 1 h at -70°C, unless the DNA spools out immediately.
 - d. Centrifuge 15 min at 8000 rpm or in microfuge.
 - e. Add 80% ethanol to remove residual salts from the tube, spin decant, dry briefly in air, and redissolve DNA in 500 μ L of 1 × TE (10 mM Tris, 1 mM EDTA) in a microtube (*to be sure that DNA is in the Na salt form and there is not a lot of Cs that may inhibit enzymes*).
 - f. Reprecipitate the DNA by adding 1/10th volume of Na acetate and two volumes of ethanol; mix, freeze, spin, wash, and dry.
- 20. Dry pellet and resuspend in 500 μ L of 1 × TNE or more, if needed. Be sure all of the DNA is resuspended evenly, this may take 10 to 15 min on ice to resolve, with gently inverting. Scan a 1/100 dilution. Adjust to desired concentration with 1 × TNE, rescan, and store at -20°C.

B. Protocol 2

1. Solutions

- Solution 1: 15% sucrose, 50 mM Tris-HCl (pH 8.0), 50 mM Na₂EDTA, 250 mM NaCl
- Solution 2: 20 mM Tris-HCl (pH 8.0), 10 mM Na₂EDTA
- TE: 20% SDS, 7.5 M NH₄OAc, isopropanol, ethanol

2. Procedure

- 1. Grind the leaf tissue (50 to 100 mg) in a 1.5 mL tube with a "pellet pestle" or a glass rod in liquid nitrogen.
- 2. Add 1 mL of Solution 1 to the ground sample, mix, and keep on ice for a few minutes.
- 3. Spin for 5 min at 5 K, and discard the supernatant (nuclei in the pellet).
- 4. Resuspend the pellet in 0.3 mL of Solution 2, by gently tapping and add 20 μ L of 20% SDS, invert to mix.
- 5. Incubate the sample at 70°C for 15 min. (Remove parafilm, because it will melt at 70°C.)
- 6. Add 170 μ L of 7.5 M NH₄OAc and mix thoroughly. Cover with parafilm inside out, and keep on ice for 30 min.
- 7. Sediment the NH_4OAc -SDS pellet at 15 K for 5 min.
- 8. Transfer supernatant to a new tube, add 0.6 vol (300 μ L) of isopropanol, incubate on ice for 15 min, mix well, and whitish cotton material will precipitate. Keep on ice.
- 9. Pellet the DNA at 15 K for 5 min.
- 10. Wash the pellet with cold 80% ethanol, dry, and resuspend in 50 μ L of TE.

Note: Clean up: resuspend in 100 μ L TE + 50 μ L 7.5 M ammonium acetate ~30 min on ice, transfer supernatant to a new tube + 300 μ L ethanol, and wash with 80% ethanol.

VIII. DETERMINATION OF NUCLEAR DNA CONTENT OF PLANTS BY FLOW CYTOMETRY

A. Introduction

Determination of an accurate amount of nuclear DNA content is extremely important to understand the hereditary constituent of an organism. Flow cytometry, originally developed for medical studies, is an easy, rapid, accurate, and convenient tool for estimating plant genome size, ploidy level, assessing DNA content, and analyzing the cell cycle (Doležel, Binarová, and Lucretti, 1989; Doležel, 1991; Awoleye et al., 1994; Bharathan, Lambert, and Galbraith, 1994; Baranyi and Greilhuber, 1995, 1996; Greilhuber, 1998; Winkelmann, Sangwan, and Schwenkel, 1998). Voluminous information on the nuclear DNA contents in plants is being published by Bennett and his colleagues (Bennett and Leitch, 1995, 1997, 2001; Bennett, Leitch, and Hanson, 1998; Bennett et al., 2000; Hanson et al., 2001).

In addition to determining the nuclear DNA content, flow cytometry in higher plants is used for studying plant protoplast (protoplast size, cell wall synthesis, chlorophyll content, alkaloid content, RNA content, protein content, protoplast–microbe interaction, sorting of protoplast fusion products) and chromosomes (chromosome size, centromeric index, sorting of large quantities of chromosomes of single type for gene isolation and mapping) (Doležel, Macas, and Lucretti, 1999; www.ueb.cas.cz/olomoucl). Several protocols have been developed, and are described below, to estimate nuclear DNA contents in the higher plants.

B. Protocol 1

For more information on Protocol 1, see Arumuganathan and Earle (1991).

1. Stock Solutions

- 1. MgSO₄ buffer:
 - a. Dissolve in ddH₂O
 - b. 0.246 g (10 mM) MgSO₄·7 H₂O
 - c. 0.370 g (50 mM) KCl
 - d. 0.120 g (5 mM) Hepes
 - e. Adjust volume to 100 mL with ddH₂O, and adjust pH to 8.0.
- 2. Triton X-100 stock (10% w/v): 1 g Triton X-100 in 10 mL ddH_2O
- 3. Propidium iodide (PI) stock (5 mg/mL): PI (Calbiochem, 537059): 5.0 mg/mL ddH₂O; cover with aluminum foil and store in refrigerator
- 4. RNase (DNase free) (Roche Molecular Biochemicals, cat. # 1 119 915)
- 5. Alsever's solution:
 - a. Dissolve in ddH_2O
 - b. 0.055 g citric acid
 - c. 2.05 g glucose
 - d. 0.42 g sodium chloride
 - e. 0.8 g sodium citrate
 - f. Adjust volume to 100 mL with ddH_2O , adjust pH to 6.1, and autoclave.
- 6. Chicken red blood cells (CRBC):
 - a. Fresh chicken blood diluted with Alsever's solution to a concentration of ~10⁷ CRBC/mL (see "Estimation of Nuclear DNA Content," item 1). Store in refrigerator (can be used up to 4 weeks).

2. Final Solutions

The following solutions should be freshly prepared using the stock solutions, be protected from light with aluminum foil, and be kept on ice.

- 1. Solution A (25 mL, enough for 15 samples):
 - a. 24 mL MgSO₄ buffer (ice-cold)
 - b. 25 mg dithiothreitol (Sigma, D-0632)
 - c. 500 µL PI stock
 - d. 625 µL Triton X-100 stock
- 2. Solution B (7.5 mL, enough for 15 samples):
 - a. 7.5 mL Solution A
 - b. 17.5 µL RNase (DNase free)

3. Procedure

Note: The entire procedure should be done on ice or in a cold room.

a. Preparation of Suspension of Nuclei from Plant Tissues

- 1. Excise young leaves (without midrib) from healthy plants.
- 2. Place pieces of leaves (~20 mg from test plant and ~20 mg from standard plant) (see "Estimation of Nuclear DNA Content," item 2) in plastic Petri dishes (35 mm × 10 mm) on ice.
- 3. Add 1 mL of Solution A, and slice the tissues into thin pieces (≤0.5 mm pieces) using a sharp scalpel.
- 4. Filter the homogenate through 30 μ m nylon mesh into a microcentrifuge tube.
- 5. Centrifuge at high speed (15,000 rpm) for 15 to 20 sec, and discard the supernatant.
- 6. Resuspend pellet in 200 to 400 μ L of Solution B.
- 7. Incubate for 15 min at 37°C.
- 8. Run the sample on a flow cytometer.

b. Preparation of Suspension of Nuclei from Plant Protoplast

- 1. Isolate protoplasts from callus (see "Estimation of Nuclear DNA Content," item 3), cell suspensions, or tissues by standard methods using wall-degrading enzymes.
- 2. Add 1 mL of Solution A to the pellet of protoplasts (~105), and incubate on ice for 10 min.
- 3. Filter the lysate through 33 µm nylon mesh into a microcentrifuge tube.
- 4. Follow Steps 5 to 8 described above for plant tissues.

c. Flow Cytometric Analysis of Suspensions of Nuclei

- For estimation of the DNA content of nuclei, measure the relative fluorescence of nuclei by using an EPICS PROFILE flow cytometer (see "Estimation of Nuclear DNA Content," item 4) (Coulter Electronics, Hialeah, Florida, U.S.A.) with an argon ion laser operating at a wavelength of 488 nm. Filters included a 457 to 502 nm laser blocking filter and a 610 nm long-pass absorbance filter in front of the photomultiplier tube. The instrument is aligned, and analog digital converter linearity is checked according to the procedure recommended by Coulter Electronics.
- 2. Objects are analyzed for logarithmic (LFL) and linear integral fluorescence (FL) and forwardangle light scatter (FS), a parameter related to particle size. Instrument protocol is set to generate three histograms:
 - a. LFL versus FS (64×64 channel)
 - b. Frequency versus LFL (256 channel, 3-decade log scale)
 - c. Frequency versus FL (256 channel)

d. Analysis Procedure

- 1. Aspirate about 50 μL of stained nuclear suspension into EPICS PROFILE flow cytometer.
- 2. Adjust the high voltage of the photomultiplier tube (PMT) so that the signals corresponding to the populations of intact nuclei ($G_0 + G_1$ nuclei, $G_2 + M$ nuclei, CRBC nuclei or internal standard and nuclei of the highest ploidy) fall within the scale of the logarithmic fluorescence intensity (see "Estimation of Nuclear DNA Content," item 5). CRBC may be problematic as a standard for plant samples (R. Obermayer, personal communication, 2002).
- 3. Set a gating region around the zone containing signals from the intact nuclei on the bivariate histogram to eliminate extraneous signals from the conditioned histogram of fluorescence.
- 4. Adjust the gain of the fluorescence amplifier (and the high voltage of the PMT, if necessary) so that the modes of the peaks corresponding to the CRBC nuclei or internal standard and the $G_0 + G_1$ nuclei of the test sample fall at desired positions within the scale of the histogram of the linear fluorescence intensity (see "Estimation of Nuclear DNA Content," item 6).
- 5. Set analysis regions about peaks to calculate frequency of nuclei, mean relative fluorescence, standard deviation, and coefficient of variation (CV) for each peak.
- 6. Repeat the above steps for replicate measurements from samples.

e. Estimation of Nuclear DNA Content

Compare the mean position of the peaks due to the plant nuclei with the mean peak position of the internal standard nuclei.

Nuclear DNA amount = [mean position of unknown plant nuclear peak/mean position of known nuclear peak] × DNA content of known standard

Note: Suspensions of plant nuclei with nuclear DNA content that has previously been determined can also be used as internal standards.

- 1. CRBC are used as an internal standard. Fresh chicken blood is collected into heparnized tubes from 2- to 3-weeks-old chicken. Usually, 1:100 dilution in Alsever's solution gives ~10⁷ CRBC/mL. Human leukocytes (HLN) [(6.5 ± 0.21) pg/2C] or red blood cells from rainbow trout (TRBC) [(5.05 ± 0.18) pg/2C] can also be used as internal standards; the values of these standards were determined by comparison with CRBC in more than 20 experiments. Suspensions of nuclei from plants with nuclear DNA content that has previously been determined can also be used as internal standards.
- 2. Avoid the use of more than 50 mg tissue/mL Solution A for chopping. Larger amounts of tissue not only increase the time spent for chopping and dilute the stain, but also increase the amount of debris and decrease the frequency of intact nuclei. Wash tissues excised from greenhouse or field-grown plants to remove debris and chemical residues that may fluoresce.
- 3. It is sometimes easier to obtain clean preparations of nuclei from callus.
- 4. Preference depends upon the availability of flow cytometer. (With commercially available flow cytometers, two basic optical parameters of particles can be analyzed: light scatter and fluorescence emission.)
- 5. The logarithmic histogram can accommodate signals varying in amplitude by at least a factor of 1000. This is useful for detecting nuclear peaks corresponding to very large or very small genomes.
- 6. Peaks for the internal standard are kept at the highest possible position between Channel 32 and 224 of a 256-channel histogram. This is usually done by keeping amplifier gain at 5 and adjusting the PMT voltage. A gain setting of less than 5 on the PROFILE causes nonlinear gain amplification. A simple statistical procedure using confidence intervals is used to compare the mean DNA content of the strains.

C. Protocol 2

For more information on Protocol 2, see Obermayer (2000).

1. Flow Cytometry after DAPI Staining

a. Chemicals

- Citric acid 1-hydrate (C₆H₈O₇·H₂0) MERCK
- Triton X-100 (C₃₄H₆₂O₁₁) MERCK
- Anhydrous disodium hydrogen phosphate (Na₂HPO₄) MERCK
- Diamidino-2-phenylindole (DAPI)
- Sulfurhodamine 101 (SR)

b. Stock Solutions

- DAPI: 0.1 mg/mL dH₂O, store at -20°C
- SR: 0.3 mg/mL dH₂O, store at 4°C
- Triton X-100: 5% Triton (dissolve in dH_2O by incubation at 37°C and store at 4°C)

c. Isolation Buffer

- 0.1 M citric acid 1-hydrate
- 0.5% Triton X-100 from 5% stock solution in dH₂O (For example, 60 mL isolation buffer contains 1.26 g citric acid 1-hydrate + 6 mL (5%) Triton stock + 54 mL dH₂O)
- Store isolation buffer at 4°C for 3 weeks (maximum).

d. Staining Solution

- 1. Dissolve 0.4 M Na₂HPO₄ at 37° C (water bath) in dH₂O.
- Cool at RT, and mix with DAPI stock in proportion 1:20 (5 μg/mL DAPI) and with SR stock in proportion 1:100 (3 μg/mL SR). (Thus, 100 mL staining solution contains 5.679 g Na₂HPO₄, 5 mL DAPI stock, 1 mL SR stock, and 94 mL dH₂O).
- 3. The staining solution can be stored at RT for about 3 months.

e. Sample Preparation

- 1. Pour 0.3 mL isolation buffer into a small Petri dish (5 cm diameter), add 10 mg young leaves, and cut into thin pieces with a sharp razor blade (use each side only twice). Slope the Petri dish, and rinse down with 0.3 mL isolation buffer.
- 2. Filter 0.4 mL of the suspension through a 48 μ m nylon tissue into a test tube (55 × 12 mm).
- 3. Add 2 mL staining solution (proportion 1 + 5), and mix carefully.
- 4. Measure the sample at the flow cytometer immediately or within a few hours.

2. Flow Cytometry after Propidium Iodide (PI) Staining

a. Chemicals

- Citric acid 1-hydrate (C₆H₈O₇·H₂0) MERCK
- Triton X-100 (C₃₄H₆₂O₁₁) MERCK
- Anhydrous disodium hydrogen phosphate (Na₂HPO₄) MERCK
- Ribonuclease A (RNase): This is important for good digestion SIGMA Ribonuclease (Ribonuclease I; EC 3.1.27.5) Type I-AS; from Bovine Pancreas (9001–99–4) EC No. 232 646–6.

- Trisodium citrate dihydrate (C₆H₅Na₃O₇·2 H₂O) MERCK
- Anhydrous sodium sulfate (Na₂SO₄) MERCK
- Propidium iodide (PI) SIGMA (95 to 98% purity); P-4170, 25535-16-4

b. Stock Solutions

- 1. PI: Dissolve PI 1 mg/mL in dH₂O, and store at 4°C. Normally, prepare 120 mL, and store in 10 fractions.
- 2. Triton X-100: Dissolve 5% Triton in dH_2O by incubation at 37°C, and store at 4°C (e.g., 1.5 mL Triton + 28.5 mL dH_2O).
- RNase stock solution: Dissolve 3 mg RNase/mL dH₂O, heat to 80°C (to destroy DNase), and store at -20°C in small portions in 2 mL Eppendorf tubes. The RNase stock solution is added after the isolation of the nuclei in a proportion of 1:20, which results in a final concentration of 0.15 mg/mL RNase (e.g., 0.060 g RNase for 20 mL stock solution).
- 4. 10× stock solution: Sodium citrate (100 mM) and sodium sulfate (250 mM): 5.882 g sodium citrate + 7.102 g sodium sulfate for 200 mL 10x stock, and store at 4°C.

c. Isolation Buffer

- 0.1 M citric acid 1-hydrate, 0.5% Triton X-100 from 5% stock solution in dH₂O. (The 60 mL isolation buffer contains 1.26 g citric acid 1-hydrate and 6 mL 5% Triton stock solution in 54 mL dH₂O.)
- 2. The isolation buffer can be stored at 4°C for 3 weeks.

d. Staining Solution

- 1. Dissolve 0.4 M Na_2HPO_4 in dH_2O .
- Mix with 10x stock of sodium citrate (100 mM) and sodium sulfate (250 mM) in a proportion of 1:10. Finally, add PI stock in a concentration of 0.06 mL stock/mL staining solution. (100 mL staining solution contains 5.679 g Na₂HPO₄ + 6 mL PI-stock + 10 mL 10x stock of sodium citrate and sodium sulfate + 84 mL dH₂O).
- 3. The staining solution can be stored at RT for about 3 months in a dark bottle.
- 4. When a staining solution has a concentration of 60 μ g/mL, a proportion of one part (0.4 mL) nuclei suspension + five parts (2 mL) staining solution results in a final PI concentration of 50 μ g/mL.

e. Sample Preparation

- 1. Transfer 0.55 mL isolation buffer into a small Petri dish, add about 15 mg young leaves, and cut with a sharp razor blade. Slope the Petri dish, and rinse down with 0.55 mL isolation buffer.
- 2. Filter 0.95 mL of the suspension through a nylon tissue into a test tube (55×12 mm).
- 3. Add 0.05 mL RNase stock solution.
- 4. Digest at 37°C for 30 min in the water bath. (If necessary, sample can be stored overnight at 4°C.)
- 5. Divide the suspension of each sample into two test tubes, each containing 0.4 mL.
- 6. Add 2 mL staining solution to each test tube (proportion 1 + 5), and mix carefully.
- 7. Incubate for at least 20 min at RT.
- 8. Measure the sample with the flow cytometer after 20 min or within a few hours.

3. Materials for Flow Cytometry

See Appendix 2-II

D. Protocol 3

This protocol was developed by Doležel, Binarová, and Lucretti (1989; www.ueb.cas.cz/olomoucl). The main advantages of flow cytometric analysis are as follows:

- · Rapidity, precision, and convenience
- No need for dividing cells
- Nondestructive (requires small amount of tissue)
- Analysis of large population of cells (detection of mixoploidy)

1. Procedure

- 1. Chop 20 mg (higher quantities are usually needed for callus and cultured cells) leaves with a sharp scalpel in 1 mL of ice-cold LB01 buffer in a Petri dish.
- 2. It is preferable to include a DNA fluorochrome [DAPI (2 μg/mL) or PI (50 μg/mL + 50 μg/mL RNase)] in the buffer prior to chopping. *Note*: A modified version is as follows: isolate protoplast, and resuspend in ice-cold LB01 to a concentration of 10⁵ to 10⁶/mL. The concentration of Triton X-100 in the LB01 buffer should be increased to 0.5% (v/v). This improves the release of the nuclei from the protoplast. The protoplast
- should be 90 to 100% viable.
- 3. Filter the suspension through 42 μ m nylon mesh.
- 4. Store on ice prior to analysis (a few min to 1 h).
- 5. Analyze the relative DNA content by flow cytometry.

2. Fluorescent Staining of Nuclear DNA

Flow cytometry can quantify precisely the nuclear DNA content of cells by a suitable fluorescent dye. Nuclear DNA content can be estimated using several stains that bind specifically and stoichiometrically to DNA. Some of them quantitatively intercalate between pairs of double-stranded nucleic acids, others bind selectively to DNA either to AT- or GC-rich regions.

The selection of fluorescent stains is important. The fluorescent molecules absorb and emit light at a characteristic wavelength. Thus, fluorescent probes can be selectively excited and detected even in a complex mixture of fluorescent molecular species. The emitted light has always less energy, because some of the absorbed energy is lost as heat. This phenomenon is called "stokes shift," which facilitates discrimination between the exciting light and the emitted fluorescence.

3. Cell Cycle in Higher Plants

The cell cycle in higher plants (eukaryotes) is divided into cellular (growth phase and division phase — cytokinesis) and nuclear events. Nuclear events include mitosis (M) and interphase periods. Interphase is divided into a synthetic (S) phase, and two gaps, G_1 and G_2 . Nonproliferating cells may leave the cell cycle in G_1 or G_2 (Figure 2.3).

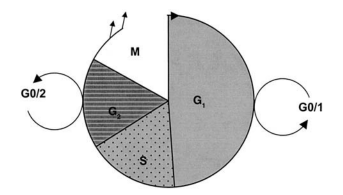


Figure 2.3 The cell cycle in plants. The interphase includes G_1 , S, and G_2 phases. (M = mitosis). (Redrawn from www.ueb.cas.cz/olomouc1.)

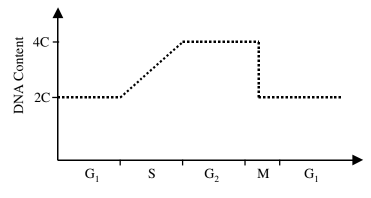


Figure 2.4 Changes of nuclear DNA content during the cell cycle. The G₁ phase constitutes 2 C DNA content and G₂ 4 C DNA content. (Redrawn from www.ueb.cas.cz/olomouc1.)

4. Nuclear DNA Content and the Cell Cycle

A diploid cell has a 2 C nuclear DNA content during G_1 phase (i.e., two copies of nuclear genome). Nuclear DNA content doubles from 2 C to the 4 C level during the S phase of the cell cycle. The S phase is followed by the G_2 phase (second period of cell growth), during which the DNA content is maintained at the 4 C level. The DNA content returns suddenly to the 2 C level, as the G_2 phase moves into mitosis, which generates two identical daughter nuclei (Figure 2.4).

5. Determination of Nuclear Genome Size

 $2 \text{ C DNA (pg)} = \text{Sample } G_1 \text{ Peak Mean} \times \text{Standard } 2 \text{ C DNA Content (pg)/Standard } G_1 \text{ Peak Mean}$

A comparison of relative positions of G_1 peaks corresponding to the sample nuclei and the nuclei isolated from a plant with known DNA content, respectively, provide precise estimation of the "unknown" DNA content.

6. Resolution of DNA Content Histograms

The resolution of histograms of nuclear DNA content is usually estimated by the coefficient of variation of the peaks corresponding to cells in the G_1 or G_2 phase of the cell cycles.

 $CV = Standard Deviation/(Mean \times 100)$

The CV reflects the variation in nuclei isolation, staining, and measurement. The smaller the CV, the more accurate the analysis of DNA content.

7. Instrument Calibration

The precision of flow cytometric measurement can be controlled using instrument standards (e.g., fluorescently labeled microspheres, nuclei isolated from chicken or fish red blood cells). Adjustment of the flow cytometer to give a maximum signal amplitude and minimal CV using an instrument standard defines the optimal operating conditions.

8. Preparation of Fixed CRBC Nuclei for Instrument Alignment

1. Mix 1 mL fresh chicken blood with 3 mL CRBC Buffer 1 (below). Centrifuge at 50 g for 5 min.

- 2. Discard the supernatant, add 3 mL CRBC Buffer 1, and gently mix. Centrifuge at 50 g for 5 min.
- 3. Discard the supernatant, resuspend the pellet in 2 mL CRBC Buffer 2 (below), and vortex briefly.
- 4. Immediately add 2 mL CRBC Buffer 3 (below), and mix briefly.
- 5. Centrifuge at 250 g for 5 min. Discard the supernatant. Add 2 mL CRBC Buffer 3, and mix gently.
- 6. Centrifuge at 120 g for 5 min, and discard the supernatant. Transfer the pellet to a clean tube, add 2 mL CRBC Buffer 3, and mix gently.
- 7. Centrifuge at 90 g for 5 min, and discard the supernatant.
- 8. Resuspend the pellet, add 2 mL ice-cold fresh fixative (3 ethanol:1 glacial acetic acid), and vortex briefly.
- 9. Leave overnight at 4°C, do not shake.
- 10. Gently remove the fixative, and resuspend the pelleted nuclei.
- 11. Add 6 mL of ice-cold 70% ethanol, vortex briefly, and syringe through a 30 G needle, three times.
- 12. Filter the nuclear suspension through a 42 μ m nylon filter to remove large clumps.
- 13. Store in aliquots of 2 mL at -20°C. If the concentration of the nuclei is too high, dilute it using ice-cold 70% ethanol.

Note: Isolated nuclei can be stored for up to several years without any sign of deterioration. Fixed nuclei are not suitable as a standard for estimation of nuclear DNA content in absolute units (genome size).

9. CRBC Buffers

- 1. CRBC Buffer 1:
 - a. 1.637 g 140 mM NaCl
 - b. 588.2 mg 10 mM sodium citrate
 - c. 24.23 mg 1 mM Tris
 - d. Adjust volume to 200 mL, and adjust pH to 7.1.
- 2. CRBC Buffer 2:

 - a. 0.41 g 140 mM NaCl b. 2.5 mL 5% (v/v) Triton X-100
 - c. Adjust volume to 50 mL.
- 3. CRBC Buffer 3:
 - a. 54.77 g 320 mM sucrose
 - b. 1.85 g
 - 15 mM MgSO₄· 7 H₂O 15 mM β-mercaptoethanol c. 530 μL
 - d. 60.57 mg 1 mM Tris
 - e. Adjust volume to 500 mL and pH to 7.1.

10. Determination of Ploidy in Plants by Flow Cytometry

Flow cytometry helps to determine the ploidy level in plants without counting chromosomes. Nuclear DNA content of G_1 reflects the ploidy level in cells, and that can be used to distinguish polyploid plants from the diploids.

Ploidy	DNA Content (G ₁ Phase)
2 <i>n</i> = <i>x</i>	1 C
2n = 2x	2 C
2n = 4x	4 C

11. Identification of Interspecific Hybrids

Flow cytometery may help to identify interspecific F_1 hybrids, where parents differ by nuclear DNA content. If applicable, flow cytometry permits screening of a large number of hybrids without counting chromosomes.

12. Identification of Aneuploid Lines

Flow cytometry can be used to identify an uploid lines based on quantity of nuclear DNA content. It depends on the precision of the measure (characterized by coefficient of variation of G_1 peaks) and on the difference in DNA content between diploid and an uploid plants. Basically, there are two approaches for the detection of an uploid plants using nuclear DNA flow cytometry. In both cases, internal standards should be used.

- 1. The use of euploid plant of the same species as an internal standard.
- 2. The use of a different species as an internal standard.

$$D = \frac{(G_1 \text{ Peak Ratio } 2) - (G_1 \text{ Peak Ratio } 1)}{(G_1 \text{ Peak Ratio } 1)} \times 100(\%)$$

3. where D = the relative difference in DNA content between euploid and aneuploid plants.

E. Protocol 4

This protocol has been provided by A.L. Rayburn.

1. Chemicals

- Hexylene glycol (2-methyl-2, 4-pentanediol; Aldrich cat. # 11,210-0)
- MgCl₂
- PEG 6000
- Propidium iodide (1 mg/mL stock)
- RNase
- · Sodium chloride
- Sodium citrate (trisodium salt)
- Triton X-100

2. Extraction Buffers

• dH ₂ O	425 mL
 Hexylene glycol 	65 mL
• 1 M Tris, pH 8.0	5 mL
• 1 M MgCl ₂	5 mL

3. Stain Solution A (10 mL)

	PEG 6000 (3% w/v)	0.2α
		0.3 g
٠	ddH ₂ O	4.0 mL
•	8 mM citrate buffer	4.4 mL
•	PI (100 μg/mL	1.0 mL
•	RNase-180 unit/mL	0.5 mL
•	Triton X-100 (0.1%)	0.1 mL

4. Stain Solution B (10 mL)

• PEG 6000 (3% w/v)	0.3 g
• ddH ₂ O	4.2 mL
• 0.8 mL NaCl	4.7 mL
• PI (100 μg/mL)	1.0 mL
• Triton X-100 (0.1%)	0.1 mL

5. Isolation and Staining of Nuclei

- 1. Germinate maize seeds in a 1:1 mixture of perlite and vermiculite. Expose plants to a day length of 14 h. The irradiance is approximately 100 μ M photons m⁻²S⁻¹. These conditions result in tissues that are most conducive to nuclear isolation.
- 2. Harvest seedlings after 2 weeks, remove 2.5 cm section of the seedling starting at the first mesocotyl, and remove coleoptile.
- 3. Slice seedling section into disks using a clean razor blade of approximately 3 mm in length, and place into an extraction buffer. *Note*: Keep the sample at 4°C at this and all subsequent steps. Remember to turn on the water bath and the centrifuge. Also, prepare the stain and salt solutions and place under the box on the bench so they are not exposed to light.
- 4. Place sample beakers in a beaker that is a little larger and full of ice. Turn the homogenizer (Biospec products, Bartlesville, Oklahoma, U.S.A.) on and grind the sample for 25 sec at 4500 rpm. Wash the homogenizer with ddH₂O on high power for several seconds, then dry it using Kimwipes, and repeat this after every sample.
- 5. Filter homogenate through 250 μ m mesh into the second beaker, and then filter the sample through 53 μ m into the third beaker. Place the sample in the labeled test tube.
- 6. Centrifuge the filtrate in a 15 mL Corex tube at 500 g, 4°C, for 15 min.
- 7. Remove the supernatant, resuspend the pellet in 300 µL of PI stain Solution A, and vortex for 10 sec.
- 8. Place in the water bath at 37°C for 20 min.
- 9. Add 300 µL stain Solution B to each microfuge tube.
- 10. Place on ice, in the dark, in the refrigerator, for at least 1 h.

Note: This method yields a large number of nuclei (maize: 250,000 to 30,000 nuclei per isolation).

F. Validity of Genome Size Measurements by Flow Cytometry

Despite numerous and basically correct assertions that flow cytometry is a fast and precise method for nuclear DNA content determination, published results obtained with this method have not always been reproducible (Greilhuber and Ebert, 1994; Baranyi and Greilhuber, 1996; Greilhuber and Obermayer, 1997; Greilhuber, 1998; Doležel et al., 1998; Temsch and Greilhuber, 2000; Vilhar et al., 2001; J. Greilhuber, personal communication, 2002). Greilhuber and Obermayer (1997) reinvestigated, applying flow cytometry with DAPI and ethidium bromide as DNA stains and Feulgen densitometry, the genome size in 19 of the 20 soybean cultivars examined by Graham, Nickell, and Rayburn (1994), for which intercultivar variation up to 11% had been claimed. No reproducible genome size differences were found among these cultivars with either technique, and a correlation of genome size with maturity group as reported by Graham, Nickell, and Rayburn (1994) was not confirmed. These discrepancies were attributed to the omission of internal standardization by Graham, Nickell, and Rayburn (1994). Temsch and Greilhuber (2000) re-evaluated genome size variation in Arachis hypogaea reported by Singh et al. (1996) and failed to reproduce their results. Bennett et al. (2000) examined nuclear DNA contents of Allium cepa cultivars, an out breeder with telomeric heterochromatic, from diverse geographical and environmental origins, and found no large difference in C-value (nuclear DNA content) between different cultivars.

Discrepancies in the genome size estimation from various laboratories may be attributed to technical variation and lack of reliable methods for genome size estimation. The two most widely used methods are Feulgen densitometry and flow cytometry, for determining DNA content. Flow cytometry is more convenient and rapid and has become an extremely more popular method than Feulgen densitometry. The simplicity of flow cytometry may be deceiving and may also, like densitometry, lead to the generation of flawed results as have been clearly demonstrated for the soybean, ground nut, pea, and *Allium*. It has been shown that fluorochromes that bind preferentially at AT- or GC-rich regions of DNA are not suitable for genome size estimation in plants. Thus,

preference of DAPI over PI is not a wise choice. The proper concentration of fluorochrome is critical (Doležel, 1991).

Second, plants such as *Helianthus annuus* may contain so-called "fluorescence inhibitors," i.e., still unidentified secondary metabolites that probably bind to chromatin and reduce the fluorescence yield (Price, Hodnett, and Johnston, 2000). This stoichiometric error in flow cytometry is comparable to the "self-tanning error" in Feulgen densitometry (Greilhuber, 1986, 1988), i.e., the diminishing of the Feulgen reaction by tannins and related compounds that are contained in many plant tissues.

In order to test the reproducibility of flow-cytometric data and promote better standardization procedures, Doležel et al. (1998) conducted collaborative experiments among four laboratories and compared interlaboratory results on plant genome size. Each laboratory used a different buffer or procedure for nuclei isolation. Two laboratories used arc-lamp-based instruments, and two used laser-based instruments. The results obtained after nuclei staining with PI (a DNA intercalator) agreed well with those obtained using Feulgen densitometry. The data recorded after nuclei staining with DAPI did not agree with those obtained using Feulgen densitometry. Small, but statistically significant, differences were observed between data obtained with individual instruments; differences between the same type of instruments were negligible, while larger differences were observed between lamp- and laser-based instruments.

Also of relevance for the evaluation of the reproducibility of flow-cytometric results is the study by Vilhar et al. (2001), who examined the reliability of the nuclear DNA content measurements in two laboratories. Nuclear DNA content obtained by image cytometry was comparable to photometric cytometry and flow cytometry. Image cytometry exhibited little variation among repeated experiments within each laboratory or among different operators using the same instruments. Image cytometry produced accurate and reproducible results and may be used as an alternative to photometric cytometry in plant nuclear DNA measurements. They proposed two standards for quality control of nuclear DNA content measurements by image cytometry: (1) the coefficient of variation of the peak should be lower than 6% and, (2) the 4 C/2 C ratio should be between 1.9 to 2.1.

IX. KARYOTYPING AND SORTING OF PLANT CHROMOSOMES BY FLOW CYTOMETRY

The sorting of individual chromosomes by flow cytometry is a valuable cytological tool in physical gene mapping, identification of chromosomes, isolation of molecular markers, and construction of chromosome-specific DNA libraries (Doležel, Macas, and Lucretti, 1999). However, this technique has lagged behind in plants because of the symmetric chromosomes of some plant species (Doležel, Lucretti, and Schubert, 1994). However, a range of tissues and culture types should permit the selection of a proper system for synchronization of cell division and collection of a high yield of metaphase chromosomes from the root tips of each plant species (Doležel, Lucretti, and Schubert, 1994). The karyotyping of symmetrical chromosomes is feasible by flow cytometer (Pich et al., 1995). The mitotic index (MI) is estimated as: (number of cells in mitosis/total number of cells) $\times 100(\%)$.

A. Protocol 1

1. Accumulation of Mitotic Metaphase

This protocol is highly efficient for the large-seeded legumes (*Vicia faba, Pisum sativum*) and cereals (*Hordeum vulgare, Secale cereale, Triticum aestivum*). The four prerequisites are as follows: (1) accumulation of cells at metaphase; (2) preparation of chromosome suspensions; (3) flow

analysis and sorting of chromosomes; and (4) processing of sorted chromosomes, cell-cycle synchronization, and metaphase chromosome accumulation (Doležel, Macas, and Lucretti, 1999).

Note: Perform all incubations at 25 ± 0.5 °C in darkness and all solutions are aerated. Keep aeration stones and tubing clean to avoid bacteria and fungi contamination.

a. Large-Seeded Legumes (Faba Bean, Garden Pea)

- 1. Materials for flow sorting:
 - a. Seeds
 - b. Perlite or vermiculite
 - c. $1 \times Hoagland's nutrient$
 - d. Hydroxyurea (HU)
 - e. Amiprophos-methyl (AMP)
 - f. Aquarium bubbler with tubing and aeration stones
 - g. Four-liter plastic tray (25 cm long \times 15 cm wide \times 11 cm high)
 - h. 750 mL plastic tray (14 cm long \times 8 cm wide \times 10 cm high) including an open-mesh basket to hold germinated seeds
 - i. Biological incubator (heating/cooling) with internal temperature adjusted to $25 \pm 0.5^{\circ}$ C
- 2. Germination of seeds:
 - a. Imbibe 30 to 35 seeds (needed to prepare one sample) for 24 h in dH₂O with aeration.
 - b. Wet an inert substrate with $1 \times$ Hoagland's nutrient solution, and put it into a 4 L plastic tray.
 - c. Wash seeds in dH_2O , spread them over the surface of the wet substrate, and cover them with a 1 cm layer of wet substrate.
 - d. Cover the tray with aluminum foil, and germinate seeds at 25 ± 0.5°C in a biological incubator in the dark. The optimum root length is ~4 cm, and it takes about 2 to 3 days.
 - e. Remove seedlings from the substrate, and wash in dH_2O .
- 3. Accumulation of metaphase cells in root tips:
 - a. Select ~30 seedlings with primary roots of similar length.
 - b. Thread seedling roots through the holes of an open-mesh basket placed in a 750 mL plastic tray filled with dH_2O .
 - c. Transfer the basket with seedlings to a second plastic tray containing HU treatment solution.
 - d. Incubate seedlings of faba bean with main roots about 2 cm long for 18.5 h or 18 h (garden pea) at 25 ± 0.5 °C.
 - e. Wash roots in several changes with dH_2O .
 - f. Immerse in HU-free 1 × Hoagland's nutrient solution for 4.5 h at 25 ± 0.5 °C.
 - g. Transfer the basket with seedlings to a tray filled with APM solution, and incubate for 2 h at $25 \pm 0.5^{\circ}$ C.

b. Cereals

- 1. Additional materials:
 - a. Cereal seeds
 - b. Glass Petri dishes 18 cm in diameter
 - c. Paper towels cut to 18 cm diameter
 - d. Filter papers cut to 18 cm diameter
- 2. Germination of seeds:
 - a. Place several layers of paper towel into an 18 cm Petri dish, and top them with a single sheet of filter paper.
 - b. Moisten the paper layers with dH_2O .
 - c. Spread the seeds on the filter paper. Approximately 50 seedlings are required to prepare one sample (1 mL chromosome suspension).
 - d. Cover the Petri dish, and germinate the seeds at $25 \pm 0.5^{\circ}$ C in a biological incubator in the dark. (The optimum root length is 2 to 3 cm and is achieved in 2 to 3 days).

- e. Select ~50 seedlings with roots of similar length, and process through HU and AMP treatment as described in "Accumulation of Metaphase Cells in Root Tips" for large-seeded legumes (Steps a to g). For HU treatment, incubate seedlings for 18 h. For HU-free Hoagland's nutrient solution, incubate for 6.5 h.
- f. Transfer the basket with seedlings to a plastic tray filled with an ice water bath (1 to 2°C).
- g. Place the container in a refrigerator, and leave overnight.

2. Analysis of the Degree of the Metaphase Synchrony

a. Materials

- · Root tips synchronized at metaphase
- 3:1 (v/v) ethanol/glacial acetic acid, freshly prepared
- 70 and 95% (v/v) ethanol
- 5 N HCl
- Schiff's reagent SIGMA (Lot # 79H5076)
- 45% (v/v) acetic acid
- Fructose syrup
- Xylene
- DePeX (Serva)
- Microscope slides
- 18 × 18 mm coverslips
- Coplin jars
- Microscope

b. Fixation and Staining

- 1. Collect 1 cm root tips in dH_2O ; five roots may be enough.
- 2. Fix root tips in 3:1 ethanol/glacial acetic acid overnight at 4°C.
- 3. Remove fixative with washes (three times) in 70% ethanol. Fixed root tips may be stored in 70% ethanol at 4°C for up to 1 year.
- 4. Wash root tips in several changes of dH_2O .
- 5. Hydrolyze root tips in 5 N HCl at RT for 25 min.
- 6. Wash in dH₂O, and incubate in Schiff's reagent for 1 h at RT.
- 7. Wash root tips in dH_2O , and soften for ~1 min in 45% acetic acid at RT.
- 8. To make slide for immediate count, cut purple dark-stained meristem tip, and squash it in a drop of fructose syrup between a microscope slide and a 18 × 18 mm coverslip. Prepare at least five different slides.
- 9. Analyze about 1000 cells from each slide, and determine the proportion of cells in metaphase.
- 10. To make a permanent slide, squash the purple dark-stained meristem tip in a drop of 45% acetic acid, and immediately place the slide on a block of dry ice or dip in the liquid nitrogen, remove the coverslip, dehydrate in two changes of 96% ethanol in a Coplin jar and air dry overnight, dip slide in xylene and mount on a drop of DePeX or Permount, analyze about 1000 cells from each slide, and determine the proportion of cells in metaphase.

B. Protocol 2

1. Suspensions of Plant Chromosomes

a. Materials

- Root tips synchronized in metaphase
- · Formaldehyde fixative
- Tris buffer

- LB01 lysis buffer
- 0.1 mg/mL DAPI stock solution
- 5°C water bath
- 5 mL polystyrene tubes (Falcon 352008; Becton Dickinson)
- Mechanical homogenizer (Polytron PT 1300D with a PT-DA 1305/2E probe; Kinematica)
- + 50 μm (pore size) nylon mesh in 4 \times 4 cm squares
- 0.5 mL tubes for polymerase chain reaction (PCR)
- Microscope slide
- Fluorescence microscope with $10 \times$ to $20 \times$ objectives and DAPI filter set

b. Preparation of Chromosome Suspension

- 1. Cut root tips (1 cm) immediately after the APM treatment, and rinse in dH_2O .
- 2. Fix in formaldehyde fixative at 5°C for 30 min (faba bean, garden pea, rye) or 20 min (barley).
- 3. Wash root tips three times (5 min each) in 25 mL Tris buffer.
- 4. Excise root meristem (1.2 to 2.0 mm) of 30 roots with a sharp scalpel in a glass Petri dish, and transfer them to 5 mL polystyrene tube containing 1 mL LB01 lysis buffer.
- 5. Isolate chromosomes by homogenizing at 9500 rpm using a Polytron PT 1300D for 15 sec (faba bean and garden pea) or 10 sec (barley and rye).
- Pass the suspension of released chromosomes and nuclei into a 5 mL polystyrene tube through 50 μm pore-size nylon filter to remove large tissues and fragments.
- 7. Store the suspension on ice. It is recommended that the chromosomes be analyzed on the same day, although they can be stored overnight.
- 8. Transfer 50 μ L chromosome suspension in a 0.5 mL PCR tube.
- 9. Add 1 μ L of 0.1 mg/mL DAPI stock solution, place a small drop (~10 μ L) of DAPI-stained suspension on a microscope slide.
- 10. Observe the suspension under low magnification $(10 \times \text{ or } 20 \times)$ using a fluorescence microscope. Do not cover with a coverslip.

Note: The suspension should contain intact nuclei and chromosomes. The concentration of chromosomes in the sample should be $\geq 5 \times 10^{5}$ /mL. If the chromosomes are damaged (broken or appear as long extended fibers), the formaldehyde fixation is too weak and should be prolonged. If the chromosomes are aggregated or the cells remain intact, the fixation is too strong and should be shortened.

2. Alignment of Flow Cytometer for Chromosome Analysis and Sorting

The alignment of the flow cytometer is crucial to achieve the highest purity in the sorted chromosomes fraction. This protocol describes suitable setups and fine tuning of the instrument for chromosome analysis and sorting, e.g., FACS Vantage. (For operation and alignment of the flow cytometer, read the manufacturer's instructions.)

a. Additional Materials

- Calibration beads (Polysciences): BB beads (univariate analysis) or YG beads (bivariate analysis)
- 530 ± 30 nm and 585 ± 42 nm band-pass filters (bivariate analysis)

b. Alignment of Flow Cytometer

- 1. Switch on the laser(s):
 - a. Univariate analysis Operate the argon ion laser in multi-UV mode (351.1 to 363.8 nm) with 300 mW output power.
 - Bivariate analysis Operate the first argon laser in multi-UV mode (351.1 to 363.8 nm) with 300 mW output power, and the second argon ion laser at 457.9 nm with 300 mW output power.
- 2. Allow the laser(s) to stabilize for 30 min. Peak the laser optics for maximum light output.

- 3. Empty the waste container, and fill the sheath container with sterile sheath fluid SF50.
- 4. Adjust sheath fluid pressure (10 psi for FACS Vantage instrument), and leave the fluid running to fill all plastic lines and filters in the instrument.
- 5. Install a nozzle (70 μ m orifice), and check for the air bubbles.
- 6. Install appropriate optical filters for alignment:
 - a. Univariate analysis Use a 424 ± 44 nm band-pass filter in front of the DAPI detector.
 - b. Bivariate analysis Use a 530 ± 30 nm band-pass filter in front of the DAPI detector and a 585 ± 42 nm band-pass filter in front of the mithramycin detector. Use a half mirror to split the fluorescence from the first and second lasers.
- 7. Trigger on forward scatter (FS), and select a threshold level.
- 8. Run fluorescent beads at a flow rate of 200 particles/sec, using BB beads for univariate analysis and YG beads for bivariate analysis.
- 9. Display the data on a dot plot of FS versus DAPI, and on one-parameter histograms of FS and DAPI fluorescence.
- 10. Align the instrument to achieve maximum signal intensity and minimum coefficient of variation of FS and DAPI signals.
- 11. For bivariate analysis, use a one-parameter histogram of mithramycin fluorescence, and align the second laser to achieve maximum intensity and the lowest coefficient of variation of the mithramycin signal. Change only settings specific for the second laser; do not adjust the controls. Adjust dual-laser delay and dead-time parameters as needed.

c. Adjust Sorting Device

- 1. Switch on the sorting device, and warm up the deflection plates for 30 min.
- 2. Run calibration beads at a flow rate of 200 particles/sec. It is important that the adjustment of the sorting device be done with the sample running.
- 3. Switch on the test sort mode.
- 4. Adjust the drop drive frequency and drop drive amplitude to break the stream at a suitable distance from the laser intercept point (check for satellite drops).
- 5. Adjust the drop drive phase to obtain single side streams.
- 6. Adjust the position of side streams so that they enter the collection tubes.
- 7. Switch off the test sort mode.
- 8. Calculate drop delay, and perform its optimization:
 - a. Define a sorting region for single beads (avoiding doublets and clumps).
 - b. Select a one-droplet sort envelope (number of deflected droplets) and sort mode, giving the highest purity and count precision (counter mode in FACS Vantage instrument).
 - c. Sort 20 beads onto a microscope slide.
 - d. Check the number of beads using a fluorescence microscope (do not cover the drop with a coverslip).
 - e. If the number of sorted beads is not correct, change the drop delay by a factor of 0.25, and repeat the steps above until the drop delay setting results in the highest number of sorted beads. To determine optimal drop delay, it may be convenient to sort 20 beads (at 0.25 step settings) on the same slide.

C. Protocol 3

1. Univariate Flow Karyotyping and Sorting of Plant Chromosomes

This protocol describes the analysis and sorting of plant chromosomes stained with DAPI. The flow cytometer must be equipped with a UV light source to excite this dye.

a. Materials

- Chromosome suspensions
- 0.1 mg/mL DAPI stock solution

- LB01 lysis buffer
- Collection liquid
- Sheath fluid SF50 for flow cytometric analysis; 40 mM KCl/10 mM NaCl (sterilize by autoclaving)
- Computer with spreadsheet or other software for theoretical flow karyotypes (www.ueb.cas.cz/olomoucl)
- 20 μ m pore size nylon mesh in 4 × 4 cm squares
- Flow cytometer and sorter (for example, Becton Dickinson FACS Vantage) with a UV argon laser (Coherent Innova 305) and a 424 ± 44 nm band-pass filter
- · Microscope slides
- · Fluorescence microscope with DAPI filter set

b. Preparation of Theoretical Flow Karyotypes

- 1. Prepare theoretical flow karyotypes using a spreadsheet or dedicated computer software.
- 2. Predict the assignment of chromosomes to chromosome peaks on a flow karyotype.
- 3. Determine the resolution (coefficient of variation) of chromosome peaks needed to discriminate individual chromosome types.

Note: Theoretical flow karyotypes can be modeled based on relative length or DNA content of individual chromosomes and are useful in planning experiments with chromosome analysis. The model predicts the complexity of the analysis and limitations of univariate flow karyotyping. It may be used to predict positions of peaks representing specific chromosomes on a flow karyotype and to study the effect of resolution (coefficient of variation) of chromosome peaks on discrimination of individual chromosome types.

c. Performing Flow Cytometry

- 1. Stain a chromosome suspension (~1 mL/sample) by adding 0.1 mg/mL DAPI stock solution to a final concentration of 2 μ g/mL. Analysis can be conducted immediately after addition of DAPI without incubation. If necessary, the stained suspension can be kept on ice.
- 2. Filter the suspension through a 20 μ m nylon mesh.
- 3. Make sure that the flow cytometer is properly aligned for univariate analysis and that a 424 ± 44 nm band-pass filter is placed in front of the DAPI fluorescence detector.
- 4. Run a dummy sample (LB01 lysis buffer containing 2 μg/mL DAPI) to equilibrate the sample line. This ensures stable peak positions during analysis and sorting.
- 5. Introduce the sample, and let it stabilize at an appropriate flow rate (200 particles/sec). If possible, do not change the flow rate during the analysis. Significant changes in the flow rate during the analysis may result in peak shifts.
- 6. Set a gating region on a dot plot of forward scatter (FS) and DAPI peak/pulse height to exclude debris, nuclei, and large clumps.
- 7. Adjust photomultiplier voltage and amplification gains so that chromosome peaks are evenly distributed on a histogram of DAPI signal pulse area/integral.
- 8. Collect 20,000 to 50,000 chromosomes, and save the results on a computer disk.

d. Sorting of Chromosomes

- 1. Make sure that the sorting device is properly adjusted.
- 2. Run the sample, and display the signals on a dot plot of DAPI signals pulse width versus area/integral.
- 3. Adjust the DAPI pulse width amplifier gain and width offset as needed to achieve optimal resolution of the width signal.
- 4. Check for the stability of the break-off point and the side streams.
- 5. Define the sorting region for the largest chromosome on the dot plot of DAPI pulse width versus DAPI pulse area/integral.

- 6. Select a one-droplet sort envelope (number of deflected droplets) and sort mode giving the highest purity and count precision (counter mode in FACS Vantage instrument).
- 7. Sort an exact number of chromosomes (e.g., 50) onto a microscope slide.
- 8. Check the number of chromosomes using a fluorescence microscope (do not cover the drop with a coverslip).
- 9. If the number is not correct, repeat adjustment of the sorting device using fluorescent beads.
- 10. Define a sorting region for the chromosome to be sorted on the dot plot of DAPI pulse width versus DAPI pulse area/integral.
- 11. Select sort mode and sort envelope according to required purity, number of chromosomes to be sorted, and desired volume for the sorted fraction. (Consult the manufacturer's instructions for an explanation of sort modes and sort envelopes.)
- Sort the required number of chromosomes into a polystyrene tube containing the appropriate amount of collection liquid.
 Note: The amount and composition of collection liquid depends on the number of sorted chromosomes and on their subsequent use. For PCR, use a small volume (20 to 60 μL) of sterile, dH₂O
- in a 0.5 mL PCR tube. 13. Microcentrifuge the tube for 5 to 10 sec at RT.
- 14. Sort chromosomes onto a microscope slide for determination of purity.

2. Bivariate Flow Karyotyping and Chromosome Sorting

This protocol provides information for chromosome isolation and bivariate analysis after dual staining with DAPI and mithramycin with preferentially bound AT-rich and GC-rich regions of DNA, respectively.

a. Additional Materials

- 100 mM MgSO₄ solution (filter through a 0.22 μ m filter; store at 4°C)
- 1 mg/mL mithramycin stock solution
- A 424 \pm 44 nm band-pass filter, and a 490 nm long-pass filter

b. Performing Flow Cytometry

- 1. To a chromosome suspension (~1 mL/sample), add 100 mM $MgSO_4$ solution to a final concentration of 10 mM.
- 2. Stain chromosomes by adding 0.1 mg/mL DAPI stock solution to a final concentration of 1.5 μg/mL and 1 mg/mL mithramycin stock solution to a final concentration of 20 μg/mL.
- 3. Allow to equilibrate for 30 min on ice.
- 4. Make sure that the flow cytometer is properly aligned for bivariate analysis. Use a half mirror to split the DAPI fluorescence through a 424 ± 44 nm band-pass filter and the mithramycin fluorescence through a 490 nm long-pass filter. *Note*: Because of the optical design of the dual-laser FACS Vantage instrument (which employs

spatially separated beam geometry), a half mirror is used to reflect all light from the second laser at 90 degrees toward the mithramycin detector. The DAPI fluorescence (excited by the first laser) is not reflected, and enters the DAPI detector directly.

- 5. Run a dummy sample (LB01 lysis buffer containing 1.5 μg/mL DAPI and 20 μg/mL mithramycin) to equilibrate the sample line. This ensures stable peak positions during analysis and sorting.
- 6. Filter the sample through a 20 μm nylon mesh.
- 7. Run the sample, and let it stabilize at the appropriate flow rate (200 particles/sec).
- 8. Set a gating region on a dot plot of FS versus DAPI peak/pulse height. Gating is used to exclude small debris and large clumps and nuclei. Use this gate to display other parameters (DAPI pulse area/integral and mithramycin pulse area/integral).
- 9. Adjust photomultiplier voltage and amplification gains so that chromosome peaks are evenly distributed on histograms of DAPI pulse area/integral and mithramycin pulse area/integral.

- 10. Display the data on a dot plot of DAPI pulse area/integral versus mithramycin pulse area/integral.
- 11. Collect 20,000 to 50,000 chromosomes, and save the results on a computer disk.

c. Chromosome Sorting

- 1. Make sure that the sorting device is properly adjusted.
- 2. Run the sample, and display the signals on a dot plot of DAPI pulse area/integral versus mithramycin pulse area/integral.
- 3. Check for stability of break-off point and the side streams.
- 4. Define the sorting region for the largest chromosome on the dot plot of DAPI pulse area/integral versus mithramycin pulse area/integral.
- 5. Proceed with cell sorting as described (Steps 6 to 14 in the section entitled "Sorting of Chromosomes" in "Univariate Flow Karyotyping and Sorting of Plant Chromosomes"), but define a sorting region on the dot plot of DAPI pulse area/integral versus mithramycin pulse area/integral.

D. Protocol 4

1. Physical Mapping of DNA Sequences Using PCR

This protocol describes the use of PCR on flow-sorted chromosomes for physical mapping of DNA sequences to individual chromosomes or their regions. Experiments involving PCR require extremely careful technique to prevent contamination.

a. Materials

- PCR premix
- Loading buffer
- 1.5% (w/v) agarose gel
- $1 \times TAE$ electrophoresis buffer
- DNA molecular weight markers
- 0.5 µg/mL ethidium bromide solution
- 0.5 mL PCR tubes
- Thermal cycler
- Horizontal gel electrophoresis apparatus and power supply
- · UV transilluminator and gel documentation system

b. Sorting Chromosomes

- 1. Prepare 0.5 mL PCR tubes containing 19 μL sterile $dH_2O.$ The final volume after sorting will be ~20 $\mu L.$
- 2. Sort 500 chromosomes into each tube.
- 3. Freeze the tube, and store at -20°C for up to 6 months. It is important to freeze the tubes, even if the reaction is to be performed on the same day of sorting.

c. Perform PCR

- 1. Thaw a chromosome fraction, and add 30 µL PCR premix. Vortex and microcentrigauge briefly.
- 2. Place tube in a thermal cycler, and perform PCR amplification using the following cycles:

nin	94°C (denaturation)
nin	94°C (denaturation)
nin	58°C (annealing)
nin	72°C (extension)
min	72°C (extension)
lefinitely	4°C (hold)
	nin nin nin nin min lefinitely

The annealing temperature must be optimized for a given primer pair and template.

d. Analyze PCR Products

- 1. Take equal amounts of PCR products (5 to $10 \,\mu$ L) from each tube, and add 1 to $2 \,\mu$ L loading buffer.
- Load samples onto a 1.5% agarose gel bathed in 1 × TAE electrophoresis buffer. Also, load DNA molecular weight markers.
- 3. Run electrophoresis at a constant voltage of 4 to 5 V/cm until the bromphenol blue reaches a point 3 cm from the edge of the gel.
- 4. Stain the gel with 0.5 μ g/mL ethidium bromide solution.
- 5. Photograph the gel, and analyze the presence of products in individual lanes.

2. Two-Step Sorting

This procedure is developed to sort chromosomes when their frequency in the original suspension is too low. This is frequently the case for large chromosomes. During the first sort, the sample is enriched for the required chromosome. During the second sort, the chromosomes are sorted with a high purity. In some cases, it may be practical to enrich the sample for more than one chromosome. Individual chromosomes are sorted during the second sort.

a. Additional Materials

- 1.5 mL polystyrene cup (Delta Lab, cat. # 900022)
- 1.5 mL polystyrene PCR tube

b. Enrich Sample for Desired Chromosome(s)

- 1. Make sure that the sorting device is properly adjusted.
- 2. Run the sample, and display the signals on a suitable distribution.
- 3. Perform trial sorting onto a microscope slide, and check the number of chromosomes using a dot plot of DAPI pulse width versus DAPI area/integral to define the sorting region for univariate analysis, or using DAPI area/integral versus mithramycin area/integral for bivariate analysis.
- 4. Select the sort mode and sort envelope that allow for the highest recovery (enrich mode and three deflected droplets in FACS Vantage instrument).
- 5. On a suitable distribution, define a sorting region for the chromosomes to be sorted.
- 6. Sort ≥100,000 chromosomes into 400 µL LB01 lysis buffer in a 1.5 mL polystyrene cup.

Note: The actual number of chromosomes that should be sorted depends on the number of chromosomes that will be sorted during the second sort. It is recommended to sort at least five times more chromosomes than the final number required.

c. Perform Second Sort

- 1. Add fluorescent dye(s) to reach recommended final concentrations.
- 2. Run the sample, and define a sorting region for the chromosome to be sorted.
- 3. Select sort mode, and sort envelope according to required purity, number of chromosomes to be sorted, and desired volume for the sorted fraction. (Consult manufacturer's instructions for explanation of sort modes and sort envelopes.)
- 4. Sort the required number of chromosomes into a 1.5 mL polystyrene PCR tube containing the appropriate amount of collection liquid. [For PCR, use a small volume (20 to 60 μ L) of sterile, dH₂O in a 0.5 mL PCR tube.]
- 5. Microcentrifuge the tube for 5 to 10 sec at RT.
- 6. Sort chromosomes onto a microscope slide for determination of purity.

3. Reagents and Solutions

See Appendix 2-III.

E. Remarks

- 1. It may take a week from the preparation of chromosome suspensions to their analysis and chromosome sorting by flow cytometer.
- 2. Although the protocols described above are not complicated, they require careful planning and expertise in several disciplines.
- 3. A large number of chromosomes can be sorted by the flow cytometer.
- Chromosomes isolated from the above procedures have been suitable for scanning electron microscopy, fluorescence *in situ* hybridization, PRINS, and immunolocalization of chromosome proteins. (See Doležel, Macas, and Lucretti, 1999.)

X. PRODUCTION OF WIDE HYBRIDS THROUGH IN VITRO TECHNIQUE

The media used for immature embryo or seed rescue differ from crop to crop. However, plants have been obtained from explants of cotyledon, hypocotyl, stem, leaf, shoot apex, root, young inflorescences, flower petals, patioles, ovular tissues, or embryo-derived calluses established either through embryogenesis or organogensis. Modification in media composition depends entirely on the source of explants and crops. The rescue of plants from immature seed harvested from 19 to 21 days postpollination is described here, as it is extremely difficult to obtain plants from wide crosses in the genus *Glycine*. The protocol described has been derived from the modification of several media formulations (Appendix 2-IV).

The following is the protocol for disinfection and excision of mature and immature embryos and seed:

- 1. Surface sterile seeds or pods with 5% sodium hypochlorite or 20% commercial bleach for 15 min followed by washing three to four times with sterile distilled water in hood.
- 2. Dissect the pods (soybean), and culture the immature seeds on initial medium. All media are prepared fresh by adding the compounds in order:
 - a. Adjust pH 5.8 using 0.1 or 1.0 N KOH prior to autoclaving.
 - b. Autoclave at 1.46 kg/cm² for 20 min at 121°C.
 - c. Add filter sterilized (Millipore HA 0.45 $\mu m)$ Glutamine (160 mL H₂O + 7.305 g Glutamine at 18°C and IAA).
 - d. Dispense under aseptic conditions.
- 3. All cultures are incubated at $25 \pm 1^{\circ}$ C, with a 16 h photoperiod under cool white fluorescent tubes (ca45 μ EM⁻²S⁻¹) in a Percival model LVL incubator.
- 4. Subculture every 2 weeks for the first 6 weeks, followed by 4-week intervals (total 10 weeks). Observe regularly for seed germination and also for callus formation:
 - a. Transfer germinating seedling to Step 5.
 - b. Remove embryo with cotyledonary tissue from nongerminating seeds, and place on the same medium for 2 weeks.
- 5. Place germinating seed on germination medium, and subculture at 2-week intervals until shoot is completely elongated. Transfer shoot to rooting medium if roots have not developed.
- 6. Transfer germinated seedlings to sterile vermiculite and peat moss (3:1) in a closed chamber (GA 7 vessels, Magenta Company) with enough moisture to maintain humidity.
- 7. Acclimatize seedlings to greenhouse condition by slowly displacing the lid of Magenta box.
- 8. Transfer plantlets to 15 cm clay pots in the greenhouse. The potting soil consists of 1:1:1 mixture of clay, peat, and sand.
- 9. Fertilize plants twice a week with a half-strength Hoagland's solution.

CHAPTER 3

Cell Division

I. INTRODUCTION

Cell division is a continuous process that occurs in all living organisms. It has been divided into two categories: mitosis and meiosis. Both forms of nuclear division occur in eukaryotes, and these processes comprise the cell cycle: G_1 (growth) \rightarrow S (synthesis of DNA) \rightarrow G_2 (growth) \rightarrow M (mitosis or meiosis) \rightarrow C (cytokinesis) (Smith and Kindfield, 1999). Mitosis occurs in somatic tissues, where each chromosome is divided identically into halves, both qualitatively and quantitatively. In contrast, meiosis takes place in germ cells, with the consequence that nuclei with haploid chromosome numbers are produced. Both types of cell division play important roles in the development and hereditary continuity of an organism.

II. MITOSIS

A. Process of Mitosis

The synonym of mitosis is karyokinesis, i.e., nuclear division. It is also known as equational division, because the exact longitudinal division of each chromosome into identical chromatids and their precise distribution into daughter nuclei leads to the formation of two cells identical to the original cell from which they were derived.

The process of mitotic cell division has been divided into six stages:

- 1. Interphase
- 2. Prophase
- 3. Metaphase
- 4. Anaphase
- 5. Telophase
- 6. Cytokinesis

1. Interphase

Two more terms, resting stage and metabolic stage, have been used to identify interphase cells. However, interphase cells should not be described as being in a "resting stage," because their nuclei are active as they prepare for cell division. The DNA replication and transcription occur during interphase (Manuelidis, 1990). Thus, "metabolic stage" is a more appropriate term for the interphase cells. The interphase nucleus contains one or more prominent nucleoli and numerous chromocenters,

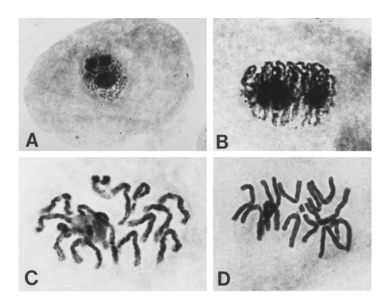


Figure 3.1 Mitosis in barley (2n = 14) without pretreatment and with aceto-carmine staining. (A) Interphase; (B) early prophase; (C) late prophase; and (D) metaphase. (From Singh, R.J., unpublished results.)

depending upon the heterochromatic nature of the chromosomes. Chromosomes cannot be traced individually, and they are lightly stained (Figure 3.1A).

2. Prophase

All the chromosomes are beginning to become distinct, and they are uniformly distributed in the nucleus. In early prophase, chromonemata become less uniform. The chromosomes are more or less spirally coiled and seem to be longitudinally double (Figure 3.1B). The two longitudinal halves of a chromosome are known as chromatids. As the prophase stage advances to mid- and late prophase, chromosomes become thicker, straighter, and smoother (Figure 3.1C). The two chromatids of a chromosome become clearly visible. The nucleoli begin to disappear in late prophase. It was formerly believed that chromosomes in a prophase nucleus are arranged in a haphazard fashion throughout the nucleus, but this is not true. Several studies have shown that in interphase and prophase, nuclei, kinetochores (centromeres), or primary constrictions are oriented toward one pole, while telomeres face opposite the kinetochores and are attached to the nuclear membrane. This orientation suggests that chromosomes maintain substantially the previous telophasic position (Figure 3.1B,C).

3. Metaphase

During metaphase, kinetochores move to the equatorial plate. Nucleoli and nuclear membrane disappear. Chromosomes are shrunk to the minimum length. Kinetochores are attached to spindle fibers, while chromosome arms may float on either side in the nucleus (Figure 3.1D). Karyotype analysis of a species is generally studied at metaphase, after pretreatment of specimen cells.

4. Anaphase

As the anaphase stage ensues, kinetochores become functionally double, and the two sister chromatids of each chromosome separate (Figure 3.2A). It appears that the kinetochores (spindle fiber attachment) regions of the two sister chromatids are being pulled to opposite poles by the spindle

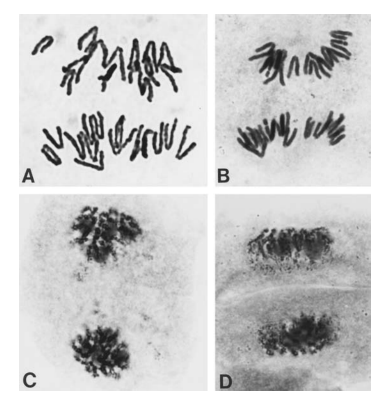


Figure 3.2 Mitosis in barley (2n = 14) without pretreatment and with aceto-carmine staining. (A) Anaphase;
 (B) late anaphase, chromosomes have already reached their respective poles; (C) telophase; and
 (D) cytokinesis. (From Singh, R.J., unpublished results.)

fibers. Chromatids are more slender and densely stained. Each daughter chromosome moves to a polar region (Figure 3.2B). By the end of anaphase, spindle fibers disappear, and the compact groups of chromosomes at the two poles are of identical genetic constitution.

Based on studies in frog eggs and egg extracts by Karsenti and Vernos (2001), mitotic spindle is a molecular machine capable of distributing the chromosomes to the daughter cells with stunning precision. The spindle is composed of microtubules that facilitate movement of chromosomes precisely during cell division. Microtubules in the spindle are arranged in two antiparallel arrays, with their plus ends at the equator, and their minus ends to the poles. Three kinds of microtubules are present in the spindle:

- 1. Astral microtubules originate from the poles and radiate out into cytoplasm.
- 2. Some microtubules connect the poles to specific sites (kinetochores) on the chromosomes.
- 3. A variable number of microtubules originate from the spindle poles and overlap in an antiparallel way at the spindle equator or interact with chromosome arms.

Most plants do not have morphologically distinct microtubules (Marc, 1997). However, plants contain functionally similar microtubule organizing centers. The key element of microtubules in plants is γ -tubulin, which is omnipresent in eukaryotes, including plants. The initiation of microtubules during cell division is initiated by γ -tubulin. Microtubule-organizing centers are dispersed in the cell. However, microtubules can assemble and self-organize into a bipolar spindle without a centriolar centrosome, but their regulation is not yet established.

Forer and Wilson (1994) presented four models for chromosome movement during postmetaphase of mitosis:

- 1. The microtubules and filaments are attached to the kinetochore.
- 2. At the onset of anaphase, motor molecules fixed in a spindle filament matrix push pole ward on the kinetochore microtubules.
- 3. Chromosome arms are pushed poleward in anaphase by forces independent of the forces that push the kinetochore fiber poleward; the forces on the chromosome arms also may arise from motor molecules associated with the spindle matrix.
- 4. The addition of tubulin subunit to kinetochore microtubules occurs at the kinetochore. The removal of tubulin subunits from kinetochore microtubules occurs at the pole and at the kinetochore. Polymerization and depolymerization are regulated at least in part by "compression" and "stretching" forces on the kinetochore microtubules. The compression of kinetochore microtubules arises when motor molecules push the kinetochore fiber into the kinetochore and chromosomes into the kinetochore fiber. The stretching force arises by motor molecules pulling kinetochore fiber out of the kinetochore.

5. Telophase

Chromosomes contract and form a dense chromatid boll. Chromosomes of the two daughter nuclei reorganize at the telophase. Nucleoli, nuclear membranes, and chromocenters reappear, and the chromosomes lose their stainability (Figure 3.2C).

6. Cytokinesis

The division of cytoplasm and its organelles between daughter cells is called cytokinesis, which begins during the late telophase stage at or near the equatorial (metaphase) plate (McIntosh and Koonce, 1989; Figure 3.2D). Cytokinesis in plants differs from that in animals. In plants, it takes place by the formation of a cell plate, but in animals, cytokinesis begins by furrowing.

B. Duration of Mitosis

The duration of the mitotic cycle varies with plant species, and ploidy levels appear to have no effect (Table 3.1). In general, interphase takes the longest time, and prophase is shorter than the remainder of the cycle. The results have shown that the greater the DNA content per cell, the longer the mitotic cycle (Van't Hof and Sparrow, 1963; Evans and Rees, 1971).

C. Chromosome Orientation at Interphase and Prophase

After mitotic anaphase, chromosomes remain localized during interphase, and eventually reappear in the same position during the next prophase (Rable model). Singh and Röbbelen (1975) demonstrated after Giemsa staining of chromosomes of several species of *Secale* that in an interphase cell, the chromocenter (telomeres), mostly lay at one side of nucleus, while opposite to it generally lay a region of dense filamentous structures — the kinetochore. The latter site could be visualized as being a spindle pole position retained from the previous division. Chromosome arms extending to the region of chromocenters represent their telomeric ends (Figure 3.3A). Therefore, chromosomes may have a relatively fixed position in the nucleus (Comings, 1980). Chromosomes are more closely associated during interphase and become more loosely oriented as mitosis progresses. The chromocenters are gradually distributed into the net-like structure of the condensing chromosomes. At early prophase, fused chromocenters can be clearly seen, and such association is still visualized during mid-prophase (Figure 3.3B). Thus, the orientation of chromosomes in interphase nuclei is loosened during later stages of the mitotic cycle, and this may be due to normal chromosome movement. Differences in squashing of material may also contribute to this effect.

Species	2 <i>n</i>	Ploidy Level	Mitotic Cycle	Authority
1. Haplopappus gracilis	4	2 <i>x</i>	10.50	Sparvoli, Gay, and Kaufmann, 1966
2. Crepis capillaris	6	2 <i>x</i>	10.75	Van't Hof, 1965
3. Trillium erectum	10	2 <i>x</i>	29.00	Van't Hof and Sparrow, 1963
4. Tradescantia paludosa	12	2 <i>x</i>	20.00	Wimber, 1960
5. Vicia faba	12	2 <i>x</i>	13.00	Van't Hof and Sparrow, 1963
6. Impatiens balsamina	14	2 <i>x</i>	8.80	Van't Hof, 1965
7. Lathyrus angulatus	14	2 <i>x</i>	12.25	Evans and Rees, 1971
8. Lathyrus articularis	14	2 <i>x</i>	14.25	Evans and Rees, 1971
9. Lathyrus hirsutus	14	2 <i>x</i>	18.00	Evans and Rees, 1971
10. Avena strigosa	14	2 <i>x</i>	9.80	Yang and Dodson, 1970
11. Secale cereale	14	2 <i>x</i>	12.75	Ayonoadu and Rees, 1968
12. Allium cepa	16	2 <i>x</i>	17.40	Van't Hof, 1965
13. Allium fistulosum	16	2 <i>x</i>	18.80	Van't Hof, 1965
14. Hyacinthus orientalis	16	2 <i>x</i>	24.00	Evans and Rees, 1971
15. Zea mays	20	2 <i>x</i>	10.50	Evans and Rees, 1971
16. Melandrium album	22	2 <i>x</i>	15.50	Choudhuri, 1969
17. Lycopersicon esculentum	24	2 <i>x</i>	10.60	Van't Hof, 1965
18. Tulipa kaufmanniana	24	2 <i>x</i>	23.0	Van't Hof and Sparrow, 1963
19. Avena strigosa	28	4 <i>x</i>	9.90	Yang and Dodson, 1970
20. Pisum sativum	28	4 <i>x</i>	12.00	Van't Hof, Wilson, and Colon, 1960
21. Triticum durum	28	4 <i>x</i>	14.00	Avanzi and Deri, 1969
22. Allium tuberosum	32	4 <i>x</i>	20.60	Van't Hof, 1965
23. Helianthus annuus	34	2 <i>x</i>	9.00	Van't Hof and Sparrow, 1963
24. Triticum aestivum	42	6 <i>x</i>	10.50	Bennett, 1971

Table 3.1 Mitotic Cycle Time (h) in Several Plant Species

D. Somatic Association

A nonrandom arrangement of chromosomes during mitosis has been reported even by earlier cytologists (Vanderlyn, 1948). Somatic association of homologous chromosomes has been observed in a large number of plants (Avivi and Feldman, 1980). It has been speculated that in plants, homologous chromosomes are closely associated during interphase (Kitani, 1963; Feldman, Mello-Sampayo, and Sears, 1966; Wagenaar, 1969). However, due to lack of suitable materials and techniques, all the somatic association studies in plants, reviewed by Avivi and Feldman (1980) are from observations or measurements of distances between homologous chromosomes at the metaphase stage, except for the observations of Singh, Röbbelen, and Okamoto (1976). Singh, Röbbelen, and Okamoto (1976) demonstrated that homologous chromosomes are more closely

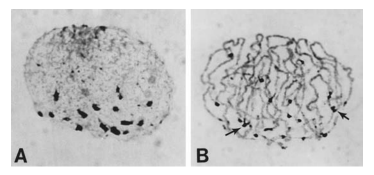


Figure 3.3 Mitotic nuclei of rye (2n = 14) after Giemsa staining. (A) Interphase, with chromocenters (telomeric bands) on the lower portion of nucleus, and spindle pole (kinetochores) on the upper side of the nucleus. (B) An early prophase nucleus with fused telomeres (arrows). (From Singh, R.J. and Röbbelen, G., *Z. Pflanzenzüchtg.*, 75, 270–285, 1975. With permission.)

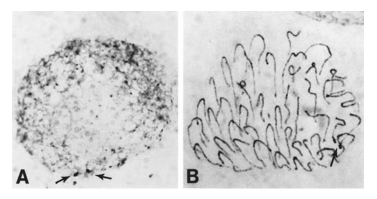


Figure 3.4 Mitotic nuclei of wheat-rye diteloaddition line (2n = 42 wheat + 2 telo rye) after Giemsa staining.
(A) Interphase with two chromocenters (arrows) representing telomeres of the rye telocentric chromosomes. (B) an early prophase with dark telomeric bands (arrow) showing somatic association. (From Singh, R.J., Röbbelen, G., and Okamoto, M., *Chromosoma (Berlin)*, 56, 265–273, 1976. With permission.)

associated during interphase of somatic cells than at metaphase. Recently, nonrandom chromatin arrangements in intact mitotic interphase nuclei of barley by confocal fluorescent microscope were recorded by Noguchi and Fukui (1995). Centromere were clustered at one site, and subtelomeric regions dispensed or were close to the nuclear membrane on the opposite site (polarity). FISH and EM studies also supported polarity results. In *Chrysanthemum multicore*, they recorded that a homologous satellite chromosome (SAT) pair fused with each other during telophase. The SAT fusion in a pair of homologous chromosomes occurs after the nuclear fusion.

Somatic association of homologous chromosomes of rye (*Secale cereale*) was studied by the Giemsa banding technique at interphase in wheat–rye addition lines. Telomeres of the rye chromosomes, appearing as chromocenters, showed close somatic association in disomic addition lines (Figure 3.4A,B), but they were distributed at random in double monosomic additions. This demonstrates directly that somatic association of homologs at interphase is even closer in nondividing nuclei than in metaphase cells. Feldman and Avivi (1973) suggested that the positions occupied by chromosomes, when examined at metaphase, were, in fact, interphase positions. However, it is unlikely that such measurements in metaphase determine interphase positions. Studies in other crop plants at metaphase also showed higher mean distances between homologous chromosomes than those reported for interphase nuclei (Table 3.2). The differences reflect dislocation of chromosomes from interphase to metaphase by chromosome contraction and spindle-mediated movement, reducing the association of homologous chromosomes.

	Mean	Distance ^a	
Crops	Homologous	Nonhomologous	Authors
	;	Somatic metaphase	
Hexaploid wheat	0.285	0.445	Feldman, Mello-Sampayo, and Sears, 1966
Hexaploid wheat	0.341	0.368	Darvey and Driscoll, 1972
Diploid oat	0.374	0.388	Sadasivaiah, Watkins, and Rajhathy, 1969
Hexaploid oat	0.332	0.440	Thomas, 1973
Barley	0.321	0.485	Fedak and Helgason, 1970b
	:	Somatic interphase	
Wheat-rye addition lines	0.250	0.387	Singh, Röbbelen, and Okamoto, 1976

Table 3.2 Comparison of Mean Distances Between Homologous Chromosomes Observed in Wheat, Oat, and Barley

^a Theoretical random distribution, 0.452.

In contrast, evidence of somatic association of homologous chromosomes has been rejected by several reports (Darvey and Driscoll, 1972; Dvořák and Knott, 1973; Therman and Sarto, 1977; Heslop-Harrison and Bennett, 1983).

Precise spatial position of chromosomes during prometaphase of human fibroplasts and He la cells was recorded by FISH (Nagele et al., 1995). Chromosomes were arranged in a wheel-shaped (rosette) pattern, and homologues were consistently positioned on opposite sites of the rosette, but the mechanism is unknown. Chromosome 16 was usually positioned adjacent to those of X chromosome. This observation does not support somatic association.

Studies of various plant species have elucidated that somatic association is a prerequisite for meiotic pairing (Brown and Stack, 1968; Stack and Brown, 1969; Chauhan and Abel, 1968; Loidl, 1990). Palmer (1971) observed no evidence for a nonrandom association of homologous chromosome in the premeiotic or ameiotic mitoses of homozygous ameiotic plants or in the premeiotic mitosis of normal sibs, and supported the classical view that homologous chromosomes do not synapse until zygonema. Furthermore, the formation (during the zygonema stage) and function of the synaptonemal complex do not necessarily favor somatic association (Moens, 1973).

Despite numerous examples that favor somatic association of homologous chromosomes (Avivi and Feldman, 1980), it is still a highly controversial issue. Therefore, a definite conclusion that the phenomenon of somatic chromosome association is common in all organisms cannot be drawn (Loidl, 1990).

III. MEIOSIS

Reduction of the somatic chromosome number to the haploid number occurs in the germ cells of plants and animals. Meiosis, known as reduction division, consists of two nuclear divisions. The first division is disjunctional. This division includes the initiation and maintenance of homologous chromosome association in prophase and metaphase and subsequent movement of homologous kinetochores of each bivalent to opposite poles during anaphase-I, thus, reducing the chromosome number by half in each daughter nucleus. Mitosis occurs in Cycle 2, producing four haploid cells from each of the diploid cells (meiocytes) that undergo meiosis. Meiotic stages have been divided into two cycles. The first cycle consists of stages responsible for producing daughter nuclei with the haploid chromosome number, and after the second cycle, four haploid nuclei will have been produced.

A. Process of Meiosis

- 1. Cycle 1
- a. Prophase-I

i. Leptonema — The chromonemata become more distinct from one another at leptonema and appear as long and slender threads. The chromosomes may be oriented with kinetochores toward the same side of the nucleus, forming a so-called "bouquet." Leptotene cells have large nucleoli and distinct nuclear membranes (Figure 3.5A).

ii. Zygonema — The mechanism of synapsis of homologous chromosomes is a complex phenomenon and is described by light microscope, electron microscope (synaptonemal complex-SC), and molecular experiments. The light microscope view is presented here.

Synapsis starts initially from one or more regions of homologous chromosomes and gradually extends at several secondary sites zipper-like, until it is complete (Burnham et al., 1972). Chromosomes begin to shorten and thicken (Figure 3.5B). A physical multiple interstitial interhomologue

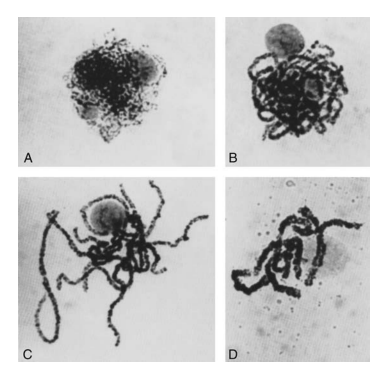


Figure 3.5 Meiosis in barley (2*n* = 14) after propiono-carmine staining. (A) Leptonema; (B) Zygonema; (C) Pachynema; and (D) an early diplonema. (From Singh, R.J., unpublished results.)

connection is a prerequisite for homologous chromosome pairing and recombination (Kleckner, 1996). The site of chromosome pairing initiation is still debated. The intimate association of homologous chromosomes in the synaptonemal complex, alignment of homologous chromosomes, and recombination have been critically reviewed by Santos (1999). Recombination should be required, for synapsis was concluded.

Centric heterochromatin contains multiple pairing elements. These elements act, initiating the proper alignment of achiasmate chromosomes early in meiosis (Karpen, Lee, and Le, 1996). This suggests that heterochromatin in eukaryote is one of the extremely mysterious components. It is distinguished from euchromatin by being sparsely populated with genes, inhibiting the function of euchromatic gene (position effect variegation), replicating late in the S phase, and being rich in tandemly repeated satellite sequences (Karpen, Lee, and Le, 1996).

However, in trisomics and in triploids, three homologous chromosomes associate in a trivalent synapsed arrangement.

iii. Pachynema — The synapsis of homologous chromosomes is completed. Chromosome regions unpaired in pachynema will usually remain unpaired. Chromosomes are noticeably thicker and shorter than those in the leptonema, and paired entities visibly represent the haploid number of that species. Each pair of chromosomes is termed a bivalent. The term bivalent is used for the paired homologous chromosomes. Some scientists still use an outdated term, tetrad. Tetrad should be used only for the four meiotic products such as the tetrad after meiosis in anthers. The unpaired chromosome is a univalent, and three paired homologous chromosomes are trivalent, etc.

Nucleoli are clearly visible, and certain chromosomes may be attached to them. These chromosomes are known as nucleolous organizer chromosomes or satellite (SAT) chromosome (Figure 3.5C).

iv. Diplonema — Chromatids composing bivalents begin to separate, or repel, at one or more points along their length (Figure 3.5D). Each point of contact is known as a chiasma (plural:

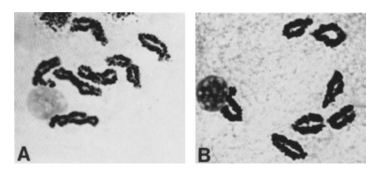


Figure 3.6 Meiosis in barley (2*n* = 14) after propiono-carmine staining. (A) Late diplonema, showing seven bivalents. (B) Diakinesis, a bivalent associated with nucleolus. (From Singh, R.J., unpublished results.)

chiasmata). At each point of contact, two chromatids have exchanged portions by crossing over. The number of chiasmata varies from organism to organism. In general, longer chromosomes have more chiasmata than shorter chromosomes.

Chiasmata may be interstitial or terminal. Interstitial chiasmata may be found anywhere along the length of a chromosome arm, but terminal chiasmata are located at a chromosome tip (telomere). In most cases, a terminal chiasma occupies an interstitial position earlier, but as cell division progresses, its position moves to the tip of the chromosome arm in which it occurred (Figure 3.6A). This phenomenon has been termed terminalization.

Torrezan and Pagliarini (1995) examined the influence of heterochromatin on chiasma localization and terminalization in maize. Chiasma frequency was higher during the diplotene, with a predominance of interstitial chiasmata located in euchromatin. During diakinesis, reduction in chiasma frequency was recorded in euchromatin, with a consequent increase in the heterochromatin. The chiasmata localized in euchromatin appear to move to the region of the heterochromatin blocks. It appears that heterochromatin serve as a barrier against terminalization, because terminalization beyond the bands was not recorded in any heteromorphic bivalent.

v. Diakinesis — Chromosomes continue to shorten and thicken. In a squash preparation, compact and thick chromosomes lie well spaced in the nucleus, often in a row near the nuclear membrane. This is a favorable stage in which to count the chromosomes. The number of chiasmata is reduced due to terminalization. The nucleolus begins to decrease in size (Figure 3.6B).

b. Metaphase-I

As soon as cells reach the metaphase stage, the nuclear membrane and nucleoli disappear, and spindle fibers appear. Bivalents are arranged at the equatorial plate, with their kinetochores facing the two poles of the cell. Chromosomes reach their maximum contractions (Figure 3.7A).

c. Anaphase-I

Homologous chromosomes begin to separate toward the opposite poles of the spindle (Figure 3.7B). The dissociation of chromosomes initiates on schedule, leaving unpaired chromosomes behind. Chromosome movement to their respective poles is coupled with the shortening of spindle micro-tubules, or kinetochore motors chew microtubules as they drag chromosomes to the poles (Zhang and Nicklas, 1996). This generates two groups of dyads with the haploid chromosome number (Figure 3.7C). The meiotic anaphase-I chromosomes are much shorter and thicker than the chromosomes of mitotic anaphase.

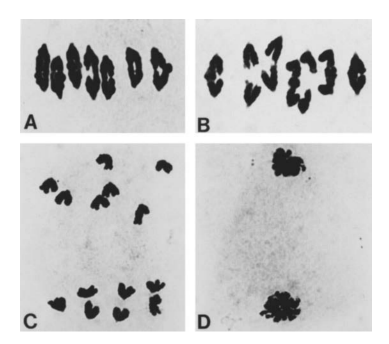


Figure 3.7 Meiosis in barley (2*n* = 14) after propiono-carmine staining. (A) Metaphase-I showing seven bivalents. (B) An early anaphase-I, where chromosomes are beginning to disjoin. (C) Anaphase-I, showing 7–7 chromosome migration. (D) Telophase-I. (From Singh, R.J., unpublished results.)

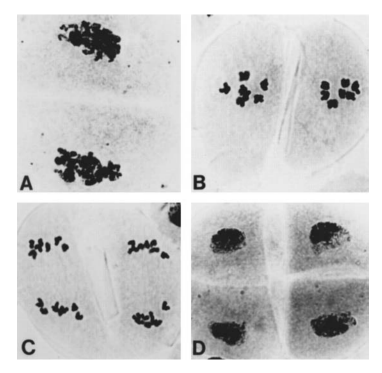


Figure 3.8 Meiosis in barley (2*n* = 14) after propiono-carmine staining. (A) Prophase-II; (B) Metaphase-II; (C) Anaphase-II; and (D) a quartet or tetrad cell. (From Singh, R.J., unpublished results.)

d. Telophase-I

At the end of anaphase-I, chromosomes reach their respective poles, and polar groups of chromosomes become compact (Figure 3.7D). Nuclear membranes and nucleoli start to develop, and eventually, two daughter nuclei with haploid chromosomes are generated. The chromatids are widely separated from each other and show no relational coiling. As a result of crossing over, each chromatid derived from a particular bivalent may be different genetically from either of the parental homologs that entered the bivalent.

e. Interkinesis

Interkinesis is the time gap between Division 1 and Division 2. Generally, it is short or may not occur at all. At the end of telophase-I, cytokinesis does not invariably follow Division 1. In many vascular plants, two daughter nuclei lie in a common cytoplasm and undergo second division.

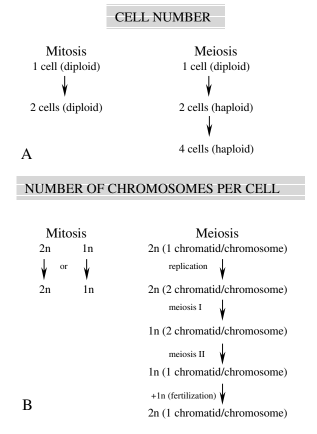
2. Cycle 2

In the second division of meiosis, prophase-II (Figure 3.8A), metaphase-II (Figure 3.8B), anaphase-II (Figure 3.8C), and telophase-II, resemble similar stages of mitotic divisions. This second division yields a quartet (tetrad) of uninucleate cells (Figure 3.8D) called microspores (in male) produced from pollen mother cells (PMC), and megaspores (in female) produced from megaspore mother cells (MMC). Tetrad analysis is a powerful tool for scientists in determining gamete development, cell division, chromosome dynamics, and recombination (Copenhaver, Keith, and Preuss, 2000).

The difference between mitosis and meiosis has been shown diagrammatically in Figure 3.9. Mitosis occurs in all growing tissues: 2n cells produce only 2n cells, or 1n cells produce only 1ncells during mitosis; while meiosis is confined to reproductive tissue: 2n tissues produce 1n cell (Figure 3.9A), and after fertilization with 1n spore, produces 2n cells. Chromosomes usually do not pair in mitotic prophase: 1-chromatid chromosomes replicate and produce 2-chromatid chromosomes, and that separates at anaphase; while in meiosis, synapsis and exchange of 2-chromatid homologue in early prophase leads to the reduction of chromosome number (Figure 3.9B). During mitosis, kinetochores lie on the metaphase plate, but in meiosis, kinetochores lie on either side of the metaphase plate. Kinetochores divide in mitotic anaphase, while kinetochores do not divide during meiotic anaphase-I. Mitosis concludes with the production of two identical diploid (2n) daughter cells, in absence of mutation, while the end of meiosis results in production of four haploid (n) daughter nuclei — two identical and two altered (resultant of synapsis, crossing over, recombination, and independent (Figure 3.9B) assortment of nonhomologous chromosomes) (Figure 3.9A). When egg and sperm unite, a zygote 2n is produced with the same amount of genetic materials as its parents, maintaining the continuity of chromosome number, but variation in genotypic constitution and phenotypic appearance evolves by meiosis.

Although most people use mitosis in reference to Division 2 of meiosis, Division 2 differs from mitosis three ways: (1) mitosis is always preceded by an interphase in which DNA synthesis occurs — meiotic Division 2 is not; (2) the chromatids within meiotic Division 2 chromosomes are not necessarily identical (due to crossing over), but in mitosis, they are; and (3) chromatids within a chromosome of mitotic prophase and metaphase are tightly wound around each other — in meiotic Division 2, the chromatids appear as if they are repelling each other in late prophase and metaphase. [This could be a carry over from anaphase-I (Figure 3.7C), when the chromatids appear to be repelling each other (e.g., four chromatid arms for a metacentric chromosome in anaphase-I)]. Division 2 could be described as superficially appearing as a mitotic division.

The following characteristic features of meiosis suggest that cell division is under genetic control (Baker et al., 1976):



- Figure 3.9 The product of mitosis and meiosis. (A) Cell number. (B) Number of chromosomes per cell. (From Smith, M.U. and Kindfield, A.C.H., *Am. Biol. Teach.*, 61, 366–371, 1999. With permission.)
 - 1. The replication of bulk DNA (premeiotic S-phase)
 - 2. Synapsis of homologous chromosomes, chiasma formation, and crossing over
 - 3. Segregation of homologous kinetochores at anaphase-I, followed by segregation of sister kinetochores to opposite poles at anaphase-II
 - 4. Formation of nuclei with the haploid chromosome number

B. Duration of Meiosis

Bennett (1971) reviewed published results on meiotic duration. The results showed that in diploid species, the duration is positively correlated with the DNA content per nucleus and with the mitotic cycle time (Table 3.3). Meiotic duration is also influenced by chromosomal organization, DNA structure, and the developmental pattern of the plant species.

C. Gametogenesis

The end products of meiosis in higher plants are four microspores in the male and four megaspores in the female, containing the haploid (n) chromosome number. Two mitotic divisions occur in microspores. The first division produces a tube nucleus and a generative nucleus. The second division occurs only in the generative nucleus, producing two sperm cells (Figure 3.10A).

Megaspores are produced in the ovules from a megaspore mother cell. Of the four megaspores produced by meiosis, only one normally survives. Three mitotic divisions occur during megagametogenesis. In the first division, a megaspore nucleus divides to give rise to the primary

Species	2 <i>n</i>	Meiotic Cycle	DNA per Cell (in picograms)
1. Antirrhinum majus	16	24.0	5.5
2. Haplopapus gracilis	4	36.0	5.5
3. Secale cereale	14	51.2	28.7
4. Allium cepa	16	96.0	54.0
5. Tradescantia paludosa	12	126.0	59.0
6. Tulbaghia violacea	12	130.0	58.5
7. Lilium henryi	24	170.0	100.0
8. Lilium longiflorum	24	192.0	106.0
9. Trillium erectum	10	274.0	120.0

^a Results from several authors, and data taken at different temperatures, however, these data provide convincing evidence.

Source: From Bennett, M.D., Proc. Roy. Soc. London Ser. B, 178, 277-299, 1971. With permission.

micropylar and primary chalazal nuclei. The second mitosis produces two nuclei each at the micropylar and at the chalazal region of the embryo sac. The third mitosis results in four nuclei at each of the opposite poles of an embryo sac. One nucleus from each pole, known as polar nuclei, moves to the middle of the embryo sac, and the two nuclei fuse to give rise to a secondary nucleus, or polar fusion nucleus, with 2n chromosome constitution. At the micropylar end of the embryo sac, of the three nuclei remaining, one nucleus differentiates into an egg cell, and the remaining two become synergids (member of the egg apparatus). The three nuclei at the chalazal end of the embryo sac are called antipodal nuclei (Figure 3.10B). Cell wall formation around the antipodals, synergids, and egg occurs. At maturity, the embryo sac consists of the following cells: antipodals (three), synergids (two), egg (one), and the large central cell that contains the polar nuclei. There are many variations to embryo sac development, and they are detailed in Maheshwari (1950). The present text covers a typical gametogenesis of many plants, particularly the grasses so important in plant breeding.

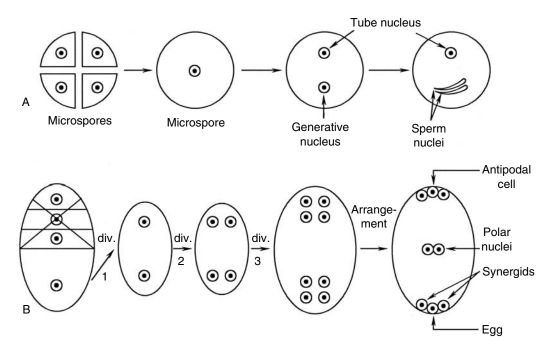


Figure 3.10 Gametogenesis in higher plants. (A) Formation of pollen grains. (B) Formation of embryo sac.

During pollination and fertilization in higher plants, pollen grains fall onto the stigma. The tube nucleus of a pollen grain directs growth of the pollen tube down the style, its travel through the micropyle, and its entrance into the nucellus. The two sperms are released into the embryo sac. One sperm nucleus unites with the egg nucleus to give rise to a 2n zygote (embryo). The other sperm fuses with the secondary nucleus to form an endosperm nucleus with triploid chromosome number. From this double fertilization, a seed develops. Higashiyama et al. (2001) demonstrated by laser cell ablation in flowering plants that two synergid cells adjacent to the egg cell attract pollen tubes. Once fertilization is completed, the embryo sac no longer attracts the pollen tube, and this cessation of attraction might be involved in blocking polyspermy.

CHAPTER 4

Genetic Control of Meiosis

I. INTRODUCTION

Meiosis is a complex process that occurs in all sexually reproducing organisms. It is a unique cell division that produces haploid gametes from diploid parental cells. During sexual reproduction, the union of two haploid spores restores the diploid chromosome complement of an organism. Thus, meiosis helps maintain chromosome numbers at a constant level from generation to generation, and ensures the operation of Mendel's laws of heredity. The steps of gametogenesis are premeiotic, meiotic [pairing of homologous chromosomes, assembly of the synaptonemal complex (SC), formation of chiasma (formation of stable connections between homologs formed at the site of crossovers), and recombination and creation of haploid meiotic products], and postmeiotic mitosis (gametophytogenesis — the equational division). The meiotic events are divided into a series of substages based on changes in chromosome morphology (Table 4.1).

The entire process of meiosis is under the control of a large number of genes (Darlington, 1929, 1932; Prakken, 1943; Rees, 1961; Katayama, 1964; Riley and Law, 1965; Baker et al., 1976; Maguire, 1978; Golubovskaya, 1979, 1989; Gottschalk and Kaul, 1980a,b; Koduru and Rao, 1981; Kaul and Murthy, 1985; Kaul, 1988; Roeder, 1997; Zickler and Kleckner, 1999; Pagliarini, 2000; Villeneuve and Hillers, 2001). Grishaeva and Bogdanov (2000) estimated more than 80 genes specifically controlling meiosis and meiotic recombination in *Drosophila melanogaster*.

The precise sequence of meiosis is sometimes disturbed by mutations. Meiotic mutants drastically change the normal behavior of chromosomes beginning with the initiation of premeiotic DNA synthesis (premeiotic mutants), during prophase-I (synaptic mutants), anaphase-I to telophase-II (disjunction mutants) (meiotic mutants), and after the completion of the second division of meiosis (postmeiotic mutants) (Figure 4.1). Meiotic mutants are rather common in the plant kingdom and are distributed in a large number of crop species covering a wide range of crop families. In the literature, meiotic and postmeiotic mutants predominate, while premeiotic and disjunction mutants are relatively rare.

Meiotic mutants have been identified chiefly on the basis of cytological observations, genetic evidence, and pollen or ovule abortion. Sometimes, they exhibit changes in the general growth habits of plants. Meiotic mutants occur by spontaneous origin in natural populations, may be induced by mutagenesis, or may result from interspecific hybridization (Tables 4.2, 4.3, and 4.4).

II. SYNAPTIC MUTANTS

It has been established by classical cytology that pairing of homologous chromosomes is initiated during zygonema by a longitudinal zipping up. It has been demonstrated in many plants that pairing sites are numerous (multiple sites) and fairly uniformly distributed along chromosomes. This

Meiotic Prophase	Chromosome Morphology	Bouquet Formation	DSB Repair	Cytological Signs
Leptotene	Axial elements begin to develop	Telomeres begin to cluster	DSBs appear	Early nodules
Zygotene	Chromosome synapsis initiates	Telomeres tightly clustered	DSBs disappear	Early nodules
Pachytene	Chromosomes fully synapsed	Telomeres disperse	DHJ	Late nodules
Diplotene	SC disassembled		Mature recombinants	Chiasma
Diakinesis	Further chromosome contraction	—	—	Chiasma

Table 4.1 Major Events in Meiotic Prophase

Note: DSB = Double-strand break; DHJ = double Holliday junction; SC = synaptonemal complex. *Source:* Adapted from Roeder G.S., *Genes and Develop.*, 11, 2600–2621, 1997.

phenomenon is not universal. In nematode (*Caenorhabditis elegans*), each chromsome contains a single site-homologue recognition region (HRR), and in every case, it is located at one end of the chromosome. This region promotes homologous chromosome pairing (Roeder, 1997). By mid-pachynema, pairing is completed. At the end of metaphase-I, chromosomes are highly contracted, the repulsion of homologues is maximum, and chromosome association is maintained only by chiasmata (Maguire, 1978). Meiotic events are precisely checked at each checkpoint (station) to ensure that one event does not occur until the preceding event has been completed. Two checkpoints operate in meiosis. The recombination checkpoint guards the cells so they do not exit pachynema until recombination intermediates have been resolved. The metaphase checkpoint impedes cells from exiting metaphase-I until all chromosome pairs have been properly oriented on the metaphase plate (Roeder, 1997). Asynaptic and desynaptic (asyndetic) mutations ignore the checkpoints (Baker et al., 1976).

Rhoades (1956) identified an ameiotic mutant (*am1*) in maize, in which meiosis did not occur, and pollen mother cells degenerated. The trait is a monogenic recessive, and homozygous plants carrying the *am1* gene are completely male sterile and partially female sterile. Palmer (1971) studied *am1* mutants cytologically and recorded normal premeiotic mitoses.

A. Distribution of Synaptic Mutants

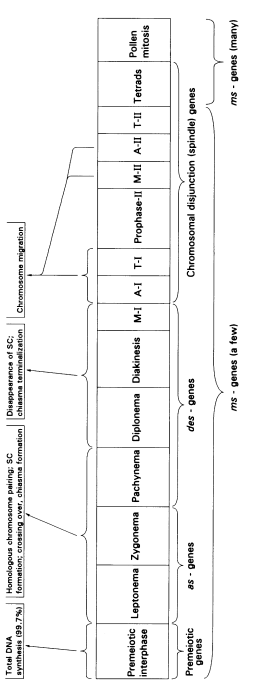
Since the discovery of the first synaptic mutants in maize by Beadle and McClintock (1928), the occurrence of such mutations has been recorded in a large number of plant species. In 1964, Katayama reported synaptic mutants in 20 families, consisting of 50 genera and about 70 species of higher plants. Koduru and Rao (1981) estimated synaptic mutants to have been observed in 126 species belonging to 93 genera. The estimates of Koduru and Rao (1981) are subject to some reduction. For example, they considered *Triticum vulgare* and *T. aestivum* to represent different species. Similarly, they considered red clover and *Trifolium pratense* to be distinct species. Therefore, they counted four species instead of two.

The family Gramineae comprises the largest number of species with identified synaptic mutants, followed by Leguminosae, Liliaceae, Solanaceae, and Malvaceae. The majority of synaptic mutants reported in higher plants are in diploid species.

Although desynaptic mutants have been reported in a large number of plant species, their origin, inheritance, gene symbol designation, and meiotic chromosome behavior have been examined in only a few plant species (Table 4.2).

B. Origin of Synaptic Mutants

Synaptic mutants of spontaneous origin have been isolated from natural populations. Similar mutations may also be induced by mutagenesis or derived from species hybrids. Generally, these mutants are





Species	2n	Gene Symbol	Origin	Genetics	Description	Authority
				Asyna	Asynaptic Mutants	
Brassica campestris	20	<i>as</i> , <i>as</i> 3	<u>n</u>	3:1	Chromosome pairing was not apparent during pachynema, diakinesis, and MI	Stringam, 1970
Datura stramonium	24	þq	S	3:1	A complete lack of homologous chromosome pairing (241) during first meiosis	Bergner, Cartledge, and Blakeslee, 1934
Hordeum vulgare	14	as	-	3:1	Absence of pachytene chromosome pairing; at MI, 14 univalent were obtained	Sethi, Gill, and Ghai, 1970
Oryza sativa	24	I	-	3:1	Absence of chromosome pairing and 24I at diakinesis and MI	Kitada and Omura, 1984
Pisum sativum	14	as	-	3:1	Chromosome pairing lacked completely at pachynema and MI	Gottschalk and Klein, 1976
Secale cereale	14	sy-1	S	3:1	Absence of meiotic chromosome pairing	Sosnikhina et al., 1992
Solanum commersonii	24	sy2	_	3:1	Absence of chromosome pairing at pachynema, diplonema, diakinesis, and MI	Johnston et al., 1986
Sorghum vulgare	20	as4	S	3:1	Characterized by lack of normal chromosome association in prophase	Stephens and Schertz, 1965
Triticum durum	28	Ι	-	3:1	Majority of the sporocytes showed 28I at MI; lacked SC (La Cour and Wells, 1970)	Martini and Bozzini, 1966
Zea mays	20	as	S	3:1	Twenty univalent were most frequent, and 18I + 1II were less frequent	Beadle, 1930
Zea mays	20	as	S	3:1	Bivalent ranged 0–9.95 at MI	Miller, 1963

Mutants
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Table 4.2

				Desyr	Desynaptic Mutants	
Allium ascalonicum	ω	I	თ	¢.	At pachynema, chromosome pairing is complete; at MI, bivalent ranged from 0-4	Darlington and Hague, 1955
Avena strigosa	14	ds	S	ю. 1:	Chromosomes were synapsed in earlier stages of meiosis; bivalent ranged from 0–7	Dyck and Rajhathy, 1965
Collinsia tinctoria	14	Q ^{et}	_	3:1	Seven bivalents were observed in 26.87% cells, while 141 in 12.8% sporocytes	Mehra and Rai, 1972
Glycine max	40	st5	S	ю:1	Almost complete pachytene chromosome pairing; very little pairing at MI	Palmer and Kaul, 1983
Hordeum vulgare	14	des1–des 15	S,	ю. 1:	Pachytene chromosome pairing is normal; chromosomes undergo desynapsis during diplonema	Ramage, 1985
Lathyrus odoratus	14	I	_	3:1	Frequencies of bivalent varied from plant to plant, and bivalents ranged from 0–7	Khawaja and Ellis, 1987
Lycopersicon esculentum	24	as1–as5	S	э: 1	Variable chromosome pairing at pachynema, diakinesis, and MI	Soost, 1951
Pennisetum glaucum	14	I	_	3:1	Chromosome pairing at pachynema is normal; bivalents at MI ranged from 0–7	Subba Rao et al.,1982
Pisum sativum	14	sp	_	3:1	Mean bivalents ranged from 0–6; the most frequent (31.5–57.6%) was 14I	Gottschalk and Baquar, 1971
Secale cereale	14	sy7, sy10	ა	3:1	Variable chromosome pairing at pachynema	Fedotova et al., 1994
Triticum aestivum	42	ds	S	ю. 1:	Chromosome synapsis occurred in normal fashion but fell apart after pachynema	Li, Pao, and Li, 1945
Avena sativa	42	syn	_	3:1 1:0	Homologous chromosomes were paired in early meiosis but disassociated prematurely at late prophase-I	Rines and Johnson, 1988
a = induced by mutagenesis; S = spontaneous origin.	sis; S = s	spontaneous origin.				

GENETIC CONTROL OF MEIOSIS

Types of Mutagen	Crops	2 <i>n</i>	Authors
X-rays	Capsicum annuum	24	Morgan, 1963
-	Collinsia tinctoria	14	Mehra and Rai, 1970
	Nicotiana sylvestris	24	Goodspeed and Avery, 1939
	Oryza sativa	24	Katayama, 1961
	Pisum sativum	14	Gottschalk and Klein, 1976
	Zea mays	20	Morgan, 1956
Gamma rays	Allium cepa	16	Konvička and Gottschalk, 1974
	Brassica oleracea	18	Gottschalk and Kaul, 1980b
	Hordeum vulgare	14	Sethi, Gill, and Ghai, 1970
	Vicia faba	12	Sjödin, 1970
Fast neutrons	Triticum durum	28	Martini and Bozzini, 1966
	Vicia faba	12	Sjödin, 1970
Colchicine	Capsicum annuum	24	Panda, Aneil Kumar, and Raja Rao, 1987
	Lathyrus odoratus	14	Khawaja and Ellis, 1987
	Lathyrus pratensis	14	Khawaja and Ellis, 1987
Diethyl sulfate	Hordeum vulgare	14	Prasad and Tripathi, 1986
Ethyleneimine	Brassica campestris	20	Stringam, 1970
-	Oryza sativa	24	Singh and Ikehashi, 1981
Ethylenemethane sulfonate	Nigella sativa	12	Datta and Biswas, 1985
5	Vicia faba	12	Sjödin, 1970
N-nitroso-N-methylurea	Oryza sativa	24	Kitada et al., 1983; Kitada and Omura, 1984
	Zea mays	20	Golubovskaya, 1979

Table 4.3 Synaptic Mutants Induced by Mutagens

Table 4.4 Synaptic Mutants of Hybrid Origin

		Gene		
Hybrids	2 <i>n</i>	Symbol	Genetics	Authors
Avena abyssinica \times A. barbata	28	ds2	3:1	Thomas and Rajhathy, 1966
Gossypium hirsutum × G. barbadense	52	as	15:1	Beasley and Brown, 1942
Nicotiana rustica × N. tabacum	42	as	3:1	Swaminathan and Murty, 1959
Sorghum durra $ imes$ S. candatum	<i>lurra × S. candatum</i> 20 <i>as1</i> 3:1 Krishn 1957		Krishnaswamy and Meenakshi 1957	
Triticum monococcum \times T. aegilopoides	14	_	15:1	Smith, 1936

identified in segregating generations, in which they are distinguished from normal plants because of pollen or ovule abortion. Meiotic mutants mostly exhibit monogenic recessive inheritance (Table 4.2; Koduru and Rao, 1981). However, the reported exception is a digenic recessive inheritance in cotton (Beasley and Brown, 1942; Weaver, 1971). Katayama (1964) and Koduru and Rao (1981) cited Hollingshead (1930a,b), in which she reported a dominant gene for asynapsis in *Crepis*. Katayama (1964) cited, "Hollingshead, L. 1930. Univ. Calif. Publ. Agr. Sci. 6: 107–134." The title of this paper is "A cytological study of haploid *Crepis capillaris* plants." I could not conclude from the paper that a dominant gene was controlling pairing. Koduru and Rao (1981) cited "Hollingshead, L. (1930): Cytological investigations of hybrids and hybrid derivatives of *Crepis capillaris* and *C. tectorum*, Univ. Calif. Publ. Agr. Sci. 6, 55–94." However, I was unable to locate a claim of monofactorial dominant inheritance. These two citations appear to present misinformation.

Synaptic mutants of spontaneous origin predominate in the literature. In barley, of the 15 desynaptic genes identified, 13 were of spontaneous origin, and two were induced by irradiation with x-rays (Table 4.5). Furthermore, synaptic mutants have been induced by various mutagens, with or without intention.

Singh and Ikehashi (1981) isolated two desynaptic mutants of rice by ethyleneimine treatment. They identified 93 sterile and partial sterile segregating M_2 families among 1924 M_2 lines.

Desynaptic Genes	Origin	Chromosome Behavior (Range Bivalents)	Female Fertility (%)	Chromosomes	Inheritance
des1	X-rays	7 O II to 5 rod II + 4 I	45	1	3:1
des2	X-rays	7 \odot II to 2 rod II + 10 I	1	3	3:1
des3	S	7 ○ II to 4 ○ II + 2 rod II + 2 I	1–33	_	3:1
des4	S	7 O II to 3 rod II + 8 I	18	1	3:1
des5	S	7 \odot II to 14 I	7	1	3:1
des6	S	7 \odot II to 14 I	16	1	3:1
des7	S	7 \odot II to 14 I	33	2	3:1
des8	S	7 \odot II to 14 I	22	—	3:1
des9	S	7 O II to 5 rod II + 4 I	90	_	3:1
des10	S	7 ○ II to 3 ○ II + 2 rod II + 4 I	60–80	—	3:1
des11	S	6 〇 II + 1 rod II to 2 rod II + 10 I	40	—	3:1
des12	S	7 O II to 1 rod II + 12 I	35	_	3:1
des13	S	7 ○ II to 4 ○ II + 2 rod II + 2 I	20–30	—	3:1
des14	S	7 \odot II to 1 rod II + 12 I	35	—	3:1
des15	S	5 〇 II + 2 rod II to 1 rod II + 12 I	25	_	

 Table 4.5
 Desynaptic Genes in Barley

Sources: des1-8 from Hernandez-Soriano, J.M., Ramage, R.T., and Eslick, R.F., Barley Genet. Newsl., 3, 124–131, 1973; des 9–14 from Hernandez-Soriano, J.M. and Ramage, R.T., Barley Genet. Newsl., 4, 137–142, 1974; and des 15 from Hernandez-Soriano, J.M. and Ramage, R.T., Barley Genet. Newsl., 5, 113, 1975.

Cytological observations revealed chromosomal interchanges, triploids, desynapsis, and genic male sterility (Table 4.6). However, the main objective of the research was to induce monogenic male sterility.

C. Cytological Behavior of Synaptic Mutants

The term asynapsis was originally proposed by Randolph (1928) to describe the absence of normal chromosome pairing during first meiotic division. The meiotic mutant was first described by Beadle and McClintock (1928) in maize. They reported that the majority of the sporocytes at metaphase-I showed 20 univalents (I) and rarely 10 bivalents (II). Mutant plants showed variable pollen and ovule abortion. Beadle (1930) assigned the gene symbol *as* to this particular asynaptic mutant.

Table 4.6	Cytological Observation and Pollen Fertility of Sterile and Partially Sterile Lines from Two Ethyleneimine Treatments
-	Ethylonoiming Treatment

	Ethyleneimine Treatment			
Cytological Observation	1 h, 0.4%	3 h, 0.2%	Total	Pollen Fertility Range (%)
Reciprocal translocations	39	22	61	1.6-80.0
Triploids	1	1	2	0.9-2.1
Desynaptic plants	1	1	2	2.1-22.1
Meiosis not studied	4	7	11	33.0-89.0
Normal meiosis	4	9	13	80-100
Normal meiosis	2	2	4	0.0-4.0
Total	51	42	93	

Source: From Singh, R.J. and Ikehashi, H., Crop Sci., 21, 286–289, 1981. With permission.

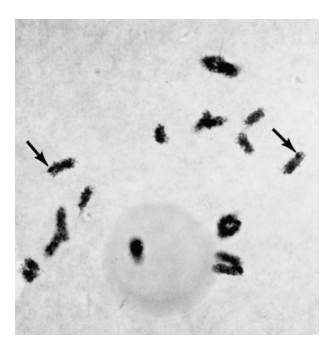


Figure 4.2 A microsporocyte at diakinesis from a desynaptic maize plant showing eight bivalents [One is nearly separated to univalents and two pairs of widely separated univalents. In one pair of univalents, there is an equationally separated distal knob (arrows), which indicates that a crossover has occurred between the knob and the kinetochore, so that in each univalent, there is a knob-carrying and knobless chromatid.] and four univalents. (From Maguire, M.P., Paredes, A.M., and Riess, R.W., *Genome*, 34, 879–887, 1991. With permission.)

Asynaptic mutants do not exhibit normal pairing of homologous chromosomes at pachynema (failure to synapse in the first place), while failure to maintain association after the first synapsis is known as desynapsis. In asynapsis, most or all of the chromosomes remain univalents at diakinesis and metaphase-I (Table 4.2). This behavior indicates that the asynaptic mutant of maize described by Beadle (1930) is probably asynaptic rather than desynaptic. Miller (1963) examined *as* mutants cytologically and observed zero to nearly complete synapsis of homologous chromosomes at early pachynema and at metaphase-I. The range of bivalents was 0 to 9.95. In the desynaptic (*dy*) mutant of maize, bivalents are mostly loosely associated (Figure 4.2). The maize asynaptic (*as*) mutant also induces polyploid meiocytes, elongated and curved spindles, misdivision of univalents, and monod centromere and partial or complete failure of cytokinesis after the meiotic divisions (Miller, 1963).

Asynaptic mutants have been reported in several plants (Table 4.2). In *Brassica campestris* (2n = 20), Stringam (1970) isolated three meiotic mutants. Mutants *as* and *as3* displayed no chromosome pairing at pachynema, and no bivalents were recorded at metaphase-I. The *as2* mutant showed a variable number of bivalents (up to four loosely paired) at diakinesis. Kitada and Omura (1984) identified one asynaptic (MM-19) and two desynaptic (MM-4, MM-16) mutants from *N*-methyl-*N*-nitrosourea-treated rice seeds. In the MM-19 mutant, the homologous chromosomes lacked complete synapsis at pachynema, resulting in 24I at metaphase-I. In contrast, mutants MM-4 and MM-16 showed variable chromosome pairing from zygonema to pachynema and displayed univalents and bivalents at metaphase-I.

Chromosome disjunction at anaphase-I to telophase-I is highly irregular in asynaptic mutants. The second division is essentially normal, but cells inherit chromosomal abnormalities resulting from the first meiotic division. Asynapsis produces chromosomally unbalanced male and female spores, resulting in a high level of pollen and ovule abortion. Gottschalk (1987)

found in x-ray-induced pea desynaptic mutants that *ds* genes influence microsporogenesis more strongly than megasporogenesis.

It has been demonstrated by electron microscopy (EM) that in asynaptic mutants, formation of the synaptonemal complex (SC) is blocked (La Cour and Wells, 1970; Golubovskaya and Mashnenkov, 1976). La Cour and Wells (1970) examined two synaptic mutants of *Triticum durum* (2n = 4x = 28) by light and electron microscopes. The light microscope showed suppression of zygotene and pachytene pairing, and EM revealed the absence of SC. When SC is eliminated in the bivalents, the two lateral elements (cores) are set free (Figure 4.3). Recently, Maguire, Paredes, and Riess (1991) observed normal crossing over followed by failure of chiasma maintenance in a desynaptic mutant of maize. They examined normal and desynaptic stocks by EM and found statistically significant wider dimension of the SC central region and less twisting of the synapsed configuration

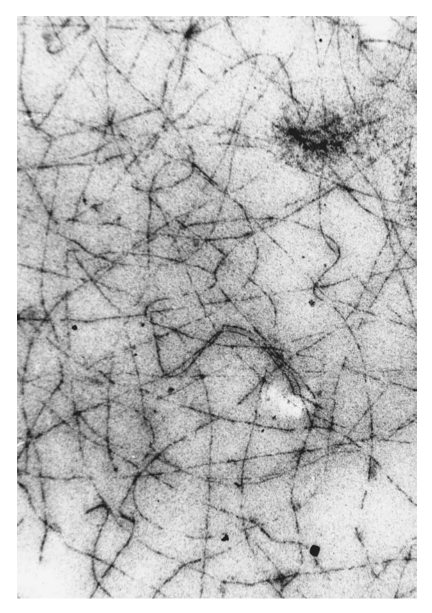


Figure 4.3 Asynaptic (*as*) nucleus at mid to late pachynema. Note the widespread lack of homologue pairing and short triple association. (Courtesy of M.P. Maguire.)

at pachynema in desynaptic mutants compared to normal. Chromosomes undergo desynapsis after pachynema to diakinesis, and by metaphase-I, the desynapsis is completed. The SC is apparently rapidly disintegrated following pachynema (Maguire, personal communication, 1992). A reduction of chiasma frequency or complete failure of chiasma formation occurs at diplonema to metaphase-I, resulting in various frequencies of univalents and bivalents. The action of desynaptic genes differs among sporocytes as well as among plants.

Because pachytene chromosome analysis is not feasible for a large number of plant species, the action of desynaptic genes is often ascertained on the basis of studies of diakinesis and metaphase-I. Chromosome associations at these later stages of meiosis appear to correlate well with the amount of pachytene chromosome synapsis. The degree of desynapsis is reflected by the number of bivalents at metaphase-I and the frequency of chiasmata per cell. Chiasmata are normally not randomly distributed among cells, chromosomes, and bivalents. They also vary between genotypes and between and within cells (Rees, 1961; Jones, 1967, 1974). Chiasmata in desynaptic plants are mostly terminal at metaphase-I and are rarely interstitial (Li, Pao, and Li, 1945).

Prakken (1943) classified desynaptic mutants depending upon their expressivity: weakly desynaptic (several univalents), intermediate desynaptic (many univalents), and completely desynaptic (exclusively univalents and rarely any bivalents).

Bivalents move to the equatorial plate at metaphase-I, while univalents tend to be distributed at random in the cytoplasm. The number of univalents varies within different microsporocytes in the same plant. This suggests that within a chromosome complement of a species, there may be differences among the different chromosomes concerning their requirements for the initiation of pairing (Rees, 1958; Swaminathan and Murty, 1959; Koduru and Rao, 1981).

Disjunction of bivalents at anaphase-I is usually normal. Univalents sometimes move to the poles at random without dividing, while in other cases, they divide equationally (Soost, 1951; Miller, 1963). Univalents that fail to move to either pole remain as laggards at the equatorial plate. At telephase-I, those chromosomes that reach the poles organize dyad nuclei, while laggards often form micronuclei. The second meiotic division is essentially normal, and irregularities are restricted to the first meiotic division. As a consequence of meiotic irregularities in asynaptic and desynaptic mutants, chromosomally unbalanced male and female microspores and megaspores are generated, resulting in reduced pollen and ovule viability.

D. Synaptic Mutants and Recombinations

Homologous chromosome pairing, synaptonemal complex formation, crossing over, chiasma formation, and gene recombination in normal plants occur during zygonema and pachynema (Henderson, 1970; Palmer, 1971; Gillies, 1975). One expects to observe a reduction in recombination frequencies in synaptic mutants because of the variable level of chromosome pairing. However, conflicting views prevail in the literature. Enns and Larter (1962) recorded reduction in recombination percentage (40–43% to 14–16%) between marked loci on Chromosome 2 in homozygous desynaptic plants of barley. The reduction in recombination was positively associated with reduced chiasma formation. Nel (1979) demonstrated in maize that recombination was reduced between the marked loci on Chromosome 3 in the asynaptic heterozygotes, and recombination between the same loci was reduced to a greater extent in asynaptic homozygotes. This, however, contradicts his earlier report (Nel, 1973).

Instances have also been reported in which recombination values between two genes are not affected by the synaptic mutants. In maize, the crossing-over value was similar in the *sh1-wx* region in asynaptic and nonasynaptic plants (Beadle, 1933; Dempsey, 1959). In two asynaptic mutants (as_1, as_4) of tomato, Soost (1951) recorded apparently no significant difference in percent crossing overs between the d_1 (dwarf growth habit) and *Wo* (wooly) marker genes in *as* and *As* plants.

A higher recombination between certain marked loci in asynaptic (*as*) plants was reported in maize (Miller, 1963) and tomato (Moens, 1969). Miller (1963) proposed that higher than normal

rates of recombination occurring in *as* plants are due to the fact that the genetic markers tested were in distal segments of the short arm or in segments near the centromere, where chromosomes pair more often than in intercalary regions. The increased recombination in synaptic mutants may be due in some cases to compensation for the loss of recombination in some other parts of the genome (Miller, 1963; Sinha and Mohapatra, 1969; Omara and Hayward, 1978).

E. Factors Influencing Pairing in Synaptic Mutants

The degree of chromosome pairing in synaptic mutants may be influenced by temperature, humidity, and chemicals (Prakken, 1943; Ahloowalia, 1969; Koduru and Rao, 1981). Furthermore, it also varies from plant to plant, day to day, year to year, and between specimens collected at different times during the same day (Prakken, 1943; Soost, 1951). Moreover, the degree of expression of each synaptic gene is variable (Table 4.2). Goodspeed and Avery (1939) reported with regard to an asynaptic mutant of *Nicotiana sylvestris* that high temperature and low humidity greatly increased asynapsis, and high temperature and high humidity decreased asynapsis. Ahloowalia (1969) recorded in a desynaptic mutant of rye grass (2n = 14) that at a lower temperature (11° C), the mean bivalent/cell was 6.71 ± 0.040, but at 28°C, desynapsis (5.39 ± 0.068 /cell) was observed. In contrast, Li, Pao, and Li (1945) observed a greater extent of pairing at a higher temperature and a decreased pairing at a lower temperature in desynaptic mutants of wheat.

Magnard et al. (2001) examined a temperature-sensitive, *tardy asynchronous meiosis (tam)*, mutant in *Arabidopsis* isolated from an EMS-mutagenized M₂ population. Although *tam* allele is temperature sensitive, *tam* plants are fertile, even at the restrictive temperatures. The *tam* mutant is a recessive trait, nonallelic to *mei1*, and regulates delayed and asynchronous Meiosis I and II. The range of aberrant tetrads at 22°C in *tam* plants was 5 to 90%, and it reached to 100% at 27°C, and 12 to 24 h exposure at 27°C induced total aberrant tetrads. It is unique that the percentage of this aberration decreased when the mutant was returned to 22°C for 24 h. Chromosome abnormalities during meiosis in *tam* mutant included condensed chromosome and chromosome lagging or scattering that resulted in the formation of aberrant tetrads with more than four spores and sometimes more than one nucleus in a cell. The delayed and asynchronous meiosis produced mostly dyads in the *tam* mutant. In contrast, wild-type (WT) plants were not temperature sensitive, as meiosis was completely synchronous.

Experiments have shown that an increase in ion content of potassium (Law, 1963) and phosphate (Bennett and Rees, 1970; Fedak, 1973) increases the number of bivalents in desynaptic plants. Ahloowalia (1969) recorded an increased bivalent frequency in a desynaptic mutant of rye grass treated with 5-ethyl, 5-phenylbarbituric acid and 5,5-diethylbarbituric acid. It was hypothesized that hydrogen bonds keep chromosomes paired after their initial pairing. The desynaptic mutants may be defective in a thermosensitive compound controlling hydrogen bonds in the condensation of chromosomes. These studies suggest that desynaptic mutants may be deficient in certain ions required for normal synapsis, and that when these chemicals are added, chromosome pairing is enhanced. This aspect needs to be investigated further.

III. GENETIC CONTROL OF RECOMBINATION

The universal rule of meiosis is to segregate parental and recombinant chromosomes into gametes precisely in equal (50%) frequencies. The foundation of Mendelism is laid on this principle. Non-Mendelian segregation is recorded during meiosis in the yeast *Saccharomyces carevisiae*, which is regulated by a *ntd 80* gene (Allers and Lichten, 2001a). Two types of meiotic recombinations in yeast were discovered: reciprocal crossing over and nonreciprocal crossing over (gene conversion). Both mechanisms are highly correlated, and their relationships are altered in mutants such as *zip1*, *msh 4*, *msh 5*, and *mer 3*, where frequencies of gene conversion are normal, but reciprocal crossing over is reduced about twofold (Storlazzi et al., 1996; Allers and Lichten, 2001a).

IV. GENES RESPONSIBLE FOR CHROMOSOME DISJUNCTION

Clark (1940) reported a meiotic divergent spindle mutation (dv) induced in maize pollen treated with UV light. The dv gene disrupts the structure and function of spindles. The mutation is inherited as a simple Mendelian recessive, showing partial male and female sterility. The dv mutant exhibits normal chromosome behavior until metaphase-I (bivalents assume a normal equatorial orientation). The spindles do not converge at each pole as is observed for normal maize. The migration of chromosomes at anaphase-I is impaired, because chromosomes disjoin irregularly in several groups. As a result, single telophase-I nuclei are not formed. The second meiotic division is synchronous in all the nuclei of both daughter cells; instead of normal tetrads of microspores, one to several small nuclei (polyads) are formed. Golubovskaya and Sitnikova (1980) induced three mutants (ms28, ms43, mei025) in maize by N-nitroso-N-methylurea. These mutations are similar to dv, being inherited as single recessive genes causing complete pollen sterility.

A meiotic mutant that shows precocious centromere division (pc) in *Lycopersicon esculentum* was described by Clayberg (1959). Chromosome pairing is normal until metaphase-I. The precocity first appears at anaphase-I in some bivalents, which often lag and undergo premature centromere division. The centromeres of those chromosomes not lagging in the first division divide, in most cases, by prophase-II. All the chromosomes were regularly oriented at the metaphase-II plate. The precociously divided chromosomes move to the poles at random without further division. Many chromosomes lag in the second division and frequently form restitution nuclei. Irregular chromosome segregation results in gametes of unbalanced chromosome numbers, as revealed by the appearance of trisomic progeny. The mutation segregates as a single recessive gene (pc) and shows 0 to 10% pollen fertility.

Beadle (1932) identified a recessive, partially fertile mutant of maize (*va*). The homozygous plant (*va/va*) exhibits normal prophase-I but cytokinesis is often absent at telophase-I, resulting in gametes with diploid (2n = 20) and tetraploid (2n = 40) chromosome constitutions. A failure in cytokinesis may occur at the first or at the second meiotic division. In *Datura stramonium*, Satina and Blakeslee (1935) observed a mutant that displayed a completely normal first meiotic division, but the second division did not occur, and instead of tetrads, dyads were formed with doubled (2n = 24) chromosome numbers. Ploidy increased with each subsequent generation. The mutant was designated by the gene symbol *dy*.

Golubovskaya and Mashnenkov (1975) isolated a recessive mutant, absent from the first meiotic division (*afd*), by treating dry maize seeds with 0.012% solution of *N*-nitroso-*N*-methylurea. Meiotic prophase-I was absent in the mutant. Homologous chromosomes failed to pair and remained as 20I at metaphase-I. Chromosomes resembled C-mitotic chromosomes. At anaphase-I, 20 chromatids migrated to each pole. Because the kinetochores divided in the first division, chromatid movement to the poles at anaphase-II was random, leading to complete male and female sterility.

A failure in chromosome migration may result in the formation of multiploid microsporocytes, i.e., mixoploidy (incomplete formation of cell walls following nuclear division). An example was reported by Smith (1942) in barley. The mutant segregated as a recessive gene, designated mu. The number of bivalents in the mu mutant varied in different metaphase plates from fewer than seven to more than 100, in multiples of seven. Cytokinesis was suppressed in some premeiotic divisions, and in other groups, migration of chromosomes took place in the formation of the prometaphase plate. Pollen grains were of various sizes and devoid of starch. Sharma and Reinbergs (1972) also isolated a mutant, ms-au, in barley with comparable effects. Takahashi, Tsuchiya, and Moriya (1964) reported a dwarf plant, governed by a recessive gene and an intensifier gene. The dwarf plant carried cells with chromosome numbers ranging from 2n = 28 to 210, together with cells carrying a normal chromosome complement of 2n = 14.

A triploid inducer gene (*tri*) in barley was reported by Ahokas (1977) and studied by Finch and Bennett (1979). The mutant apparently suppresses at random the second meiotic division in about half of the megaspore mother cells. This results in the formation of embryo sacs with diploid nuclei. The triploid inducer gene does not appear to affect microsporogenesis. Rhoades and Dempsey (1966) found a mutant for elongate chromosomes (*el*) in the open pollinated maize variety Hays Golden. The mutant segregated as a monogenic recessive. The despiralization of the chromosomes occurred at both meiotic anaphases. The most significant effect of the *el* gene was the production of unreduced eggs with variable chromosome composition and varying frequencies. Ears borne on *el/el* plants had plump and shriveled seeds as well as aborted ovules. Chromosome counts revealed that plump seeds were diploid (2n = 20). A considerable frequency of shriveled seeds did not germinate, but chromosome numbers ranging from 2n = 25 to 33. It is interesting to note that all the offspring (961) were normal diploids (2n = 20) in crosses between normal female plants and *el/el* male plants.

The *el* gene generates a series of polyploid plants. Rhoades and Dempsey (1966) obtained vigorous plants up to the 5x (2n = 50) level, but beyond this level, the plants were dwarfed and female sterile. It was determined that the unreduced eggs originated by omission of the second meiotic division.

Beadle (1929) identified a meiotic mutant in maize that showed apparently normal first and second meiotic divisions of microsporogenesis, but microspore quartet cells underwent a series (at least four) of meiosis-like divisions in which the chromosomes were distributed to the two poles at random and without splitting. Cells with only one chromosome in the spindle were seen, and fragmentation of chromosomes was rather common. Plants were completely pollen sterile and partially female sterile. Beadle (1931) named the mutant polymitotic and assigned the gene symbol *po*. The polymitotic trait inherited as a simple Mendelian recessive and was not allelic to the *as* (asynaptic) gene.

V. OTHER MEIOTIC MUTANTS

Genes for "long" and "short" chromosomes have been induced by x-ray in barley. The gene for long chromosome causes a high frequency of elongated rod bivalents at metaphase-I (Burnham, 1946). On the other hand, Moh and Nilan (1954) found a recessive mutant for short chromosomes among the progenies derived from barley seed that had been subjected to atom bomb irradiation. In this mutant, chromosomes paired at early meiotic prophase, and there was a considerable shortening of the chromosomes during prophase, especially at diakinesis; the bivalents usually underwent precocious terminalization of all chiasmata, resulting in 14 univalents at metaphase-I.

A recessive gene for chromosome stickiness (*st*) has been recorded in *Zea mays* (Beadle, 1937) and *Collinsia tinctoria* (Mehra and Rai, 1970). Chromosomes were clumped at prophase-I and frequently were associated. Thus, chiasma formation was impaired, and chromosomes did not orient at the equatorial plate. Chromosomal fragmentation was common at anaphase-I, and plants homozygous for the sticky gene were male sterile and female sterile. Stout and Phillips (1973) did not record any difference in histone composition between normal plants and *st/st* mutant maize plants.

Meiotic mutants in maize have been extensively studied. They are summarized by Carlson (1988) as follows:

- 1. Absence of meiosis: Ameiotic (am1, am2)
- 2. Absence or disruption of synapsis: Asynaptic (*as*), desynaptic (*dy*, *dys1*, *dys2*), and absence of first division (*afd*)
- 3. Change in structural organization of chromosome: Elongate (el) and sticky (st)
- 4. Improper meiotic segregation or defective meiotic spindle: Divergent spindle (dv, ms28, ms43, mei025)
- 5. Failure of cytokinesis or irregularities of cell shape: Variable sterile (ms8, ms9)
- 6. Extra divisions following meiosis: Polymitotic (po)
- 7. Mutants with several effects on meiosis: Plural abnormalities of meiosis (pam2, ms17)

VI. ROLE OF HETEROCHROMATIN IN CHROMOSOME PAIRING

Heterochromatin plays an important role in chromosome pairing and chromosome alignment. The heterochromatin of centromeric regions, supernumerary chromosomes (B-chromosomes or accessory chromosomes), nucleolus organizer regions, and knobs (e.g., Knob 10 in maize) consists of highly repetitive DNA. Distal euchromatin of chromosomes is constituted mainly of unique sequences interspersed with moderately repeated sequences that have regulatory and functional roles in transcription, replication, synapsis, and recombination (Gillies, 1975; Loidl, 1987).

Several reports show that heterochromatin (highest degree of DNA synthesis-S phase) increases chiasma frequency (Rees and Evans, 1966), increases the frequency of nonhomologous association at diplonema (Church, 1974), increases crossing over (Nel, 1973), and lengthens meiotic prophase. Thus, heterochromatin prolongs the time during which crossing over takes place (Rhoades and Dempsey, 1972). Supernumerary chromosomes suppress homoeologous chromosome pairing in *Triticum aestivum* (Sears, 1976) and in *Lolium* (Evans and Macefield, 1973). On the other hand, Romero and Lacadena (1982) observed a promoter effect of rye B-chromosomes on homoeologous chromosome pairing in hexaploid wheat. This suggests that the mechanism of heterochromatin-mediated homologous chromosome pairing is undetermined (Roeder, 1997).

VII. DIPLOID-LIKE MEIOSIS IN ALLOPOLYPLOIDS

Diploid-like meiosis is under genetic control and is a common occurrence in allohexaploids of several crop plants of the family Gramineae, e.g., *Triticum aestivum* (Okamoto, 1957; Sears and Okamoto, 1958; Riley and Chapman, 1958a), *Avena sativa* (Rajhathy and Thomas, 1972; Jauhar, 1977), *Festuca arundinacea* (Jauhar, 1975), and *Hordeum parodii* (Subrahmanyam, 1978).

A. The 5-B System in Wheat (2n = 6x = 42)

Meiotic chromosome pairing under genetic control in the hexaploid wheat cultivar Chinese Spring was first demonstrated by Okamoto (1957). He studied meiotic chromosome pairing in the F_1 between plants that were monosomic for a telocentric chromosome 5BL (2n = 40 + 1 telo 5BL) and AADD (2n = 28; derived from amphidiploid *T. aegilopoides* × Ae. squarrosa) plants. Two kinds of F₁ plants were expected: 2n = 34 chromosomes without the 5BL chromosome, and 2n =35, with the 5BL chromosome. Chromosome pairing data in 34-chromosome plants revealed a low frequency of univalents and an increased number of bivalents and multivalents compared to 35-chromosome plants (Table 4.7). Based on these results, Okamoto suggested that chromosome 5BL carries a gene or genes for asynapsis. Subsequently, Riley and Chapman (1958a) reported similar results. Wall, Riley, and Gale (1971) assigned the gene symbol Ph (homoeologous pairing suppressor). A single Ph gene was postulated to control meiotic pairing of homoeologous chromosomes. Mello-Sampayo (1972) suggested two loci, acting additively, and Dover (1973) reported two linked loci for Ph. It has been demonstrated that the three genomes (A, B, and D) are genetically closely related, and Ph controls only homologous chromosome pairing. When Ph is removed or its activity is suppressed, not only do homoeologous chromosomes pair, but they also pair with the chromosomes of related species and genera, making alien gene transfer possible (Sears, 1975, 1976).

Since the discovery of Ph, several chromosome pairing suppressor and promoter genes in Chinese Spring wheat have been identified (Sears, 1976). In addition to Ph, there are three more minor pairing suppressors located on chromosomes 3AS, 3DS (Mello-Sampayo and Canas, 1973), and 4D (Driscoll, 1973); the chromosome 4D gene is as effective as the gene on chromosome 3AS, while the gene of chromosome 3DS is only half as effective as the Ph gene.

Besides suppressors, there are several promoters of chromosome pairing located on chromosome arms 5DL, 5AL, 5BS (Feldman, 1966; Riley et al., 1966b; Feldman and Mello-Sampayo, 1967;

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Table 4.7

					Average	Average Number of			
	Number		Bival	Bivalents					
	of PMCs	Univalents	Closed	Open	Trivalents	Quadrivalents	5-valents	6-valents	7-valents
34-chromosome plant	200	8.24	5.07	3.445	0.735	0.685	0.19	0.22	0.02
35-chromosome plant	200	23.82	1.025	4.115	0.320	0.005	0	0	0

Source: From Okamoto, M., Wheat Inf. Serv., 5, 6, 1957. With permission.

Riley and Chapman, 1967), 5DS (Feldman, 1968), and 5AS (Dvořák, 1976). Furthermore, genes on 2AS, 3BL, and 3DL may also be considered promoters for pairing (Sears, 1977). It has been observed in Chinese Spring wheat that asynapsis occurs at a low temperature (15°C or below), when chromosome 5D is absent (nulli-5D tetra-5B), and a lesser reduction is shown at 20°C. But, pairing is more or less normal at 25°C (Riley, 1966).

Diploid-like meiosis under genetic control has been suggested in allopolyploid species of several genera: *Chrysanthemum* (Watanabe, 1981), *Glycine* (Singh and Hymowitz, 1985a), *Gossypium* (Kimber, 1961), *Pennisetum* (Jauhar, 1981), and *Solanum* (Dvořák, 1983a).

B. Origin of Ph

Three theories have been postulated regarding the origin of the Ph gene:

- 1. It was already present in the diploid B-genome species (Okamoto and Inomata, 1974; Waines, 1976). Ekingen, Attia, and Röbbelen (1977) suggested that the homologous pairing suppressor of chromosome 3D of hexaploid wheat already existed in *Ae. squarrosa*. However, this possibility needs to be investigated further.
- 2. The *Ph* gene was transferred to chromosome 5B from an accessory chromosome. It has been shown that B-chromosomes appear to have a suppressive effect on chromosome pairing similar to that of the 5B chromosome. Furthermore, the accessory chromosomes of rye have a slight suppressive effect on pairing in hybrids with *T. aestivum*, whether 5B is present or not (Dover and Riley, 1972).
- 3. The *Ph* gene arose as a single mutation following formation of the AABB amphidiploid (Riley and Chapman, 1958a). Riley, Chapman, and Miller (1973) suggested "an activity like that of the *Ph* allele must have occurred in the first hybrids and allotetraploids from which polyploid wheat was evolved." According to Sears (1976), "an origin of *Ph* by mutation in the newly formed AABB tetraploid seemed a more likely possibility."

VIII. HAPLOIDY

Haploids have been isolated in diploid as well as polyploid species of higher plants as spontaneous occurrence in natural populations (Kimber and Riley, 1963), by induction in anthers, pollen, and ovule culture (Sunderland, 1974; Maheshwari, Tyagi, and Malhotra, 1980; Morrison and Evans, 1988) and after interspecific crosses (Kasha, 1974; Rowe, 1974; Choo, Reinbergs, and Kasha, 1985; Gupta and Gupta, 1973). The genetics and mechanisms of barley haploid (2n = x = 7) and potato dihaploid (2n = 2x = 24) production have been extensively studied and well documented (barley — Symko, 1969; Kasha and Kao, 1970; potato — Hougas and Peloquin, 1957; Hougas, Peloquin, and Ross, 1958).

A. Mechanism of Chromosome Elimination

The gradual and selective elimination of *Hordeum bulbosum* (wild species) chromosomes in diploid (2n = 14) interspecific hybrids of *H. vulgare* (cultigen) and *H. bulbosum* (alien) is attributed to chromosome fragmentation, micronuclei formation and degradation of chromatin (Subrahmanyam and Kasha, 1973; Thomas, 1988), lagging chromosomes and bridges (Lange, 1971; Bennett, Finch, and Barclay, 1976), noncongressed chromosomes at metaphase, or the failure of chromosome migration to anaphase poles (Bennett, Finch, and Barclay, 1976). These chromosomal abnormalities have been postulated to relate to *vulgare* (V) and *bulbosum* (B) genome ratios (Subrahmanyam and Kasha, 1973), asynchronous cell cycle phases and mitotic rhythms (Lange, 1971; Kasha, 1974), differential amphiplasty (Bennett, Finch, and Barclay, 1976; Lange and Jochemsen, 1976), inactivation of alien (B) DNA by nuclease (Davies, 1974), and formation of multipolar spindles and asynchrony in nucleoprotein synthesis (Bennett, Finch, and Barclay, 1976).

It has been reported by Subrahmanyam and Kasha (1973) and Bennett, Finch, and Barclay (1976) that normal double fertilization occurs in V × B crosses. Both reports recorded selective elimination of *bulbosum* chromosomes in embryos on the third day. Subrahmanyam and Kasha (1973) found 93.69% haploid nuclei (2n = 7) 11 days after pollination (DAP). In contrast, Bennett, Finch, and Barclay (1976) recorded 93.6 to 100% haploid nuclei five DAP; the discrepancy in results was attributed to different genetic backgrounds (Bennett, Finch, and Barclay, 1976).

The mechanism of dihaploid (2n = 24) isolation from *Solanum tuberosum* (2n = 4x = 48; female) and *S. phureja* (2n = 2x = 24; pollinator) is different from that observed in *vulgare–bul-bosum* crosses. Wagenheim, Peloquin, and Hougas (1960) observed that developing dihaploid embryos were associated with hexaploid endosperms, suggesting that both sperm nuclei fused to form a restitution nucleus and then combined with the secondary nucleus. Montelongo-Escobedo and Rowe (1969) recorded restitution sperm nuclei (single male gametes) in over 30% of the pollen tubes of the superior pollinators, while an inferior pollinator had two sperm nuclei in 97% of the pollen tubes. They suggested that the production of pollen grains containing a single male gamete was under genetic control.

B. Genetics of Chromosome Elimination

The entire process of the elimination of *bulbosum* chromosomes in interspecific hybrids of *H. vulgare* and *H. bulbosum* is under genetic control, and the stability of chromosomes in hybrids is influenced by the balance of parental genomes. Furthermore, haploids (n = 7) are produced in high frequency (98.2%) from diploid (2n = 2x = 14) VV by BB crosses (Table 4.8). Chromosome elimination does not occur in true hybrids with the genome ratio of 1V:2B (Davies, 1974). Subrahmanyam and Kasha (1973) concluded that based on development and chromosomal abnormalities, endosperm stability could be placed in the following descending order: 1V:4B > 1V:2B > 1V:1B > 2V:1B. The more stable embryos (VBB) were much larger and better developed at the time of embryo culture than the other embryos, regardless of the endosperm constitution (Kasha, 1974).

Ho and Kasha (1975) demonstrated from primary and telotrisomic analysis that Chromosomes 2 (both arms) and 3 (long arm) most likely carry a gene or genes, located on *H. vulgare* chromosomes, for the elimination of *H. bulbosum* chromosomes (Table 4.9). They crossed seven primary trisomic stocks (2n = 2x + 1 = 15) of barley with pollen from autotetraploid (2n = 4x = 28) *H. bulbosum*. Because primary trisomics generate 8- and 7-chromosome gametes, two kinds of hybrid progenies are expected, namely: (1) 21-chromosome triploid hybrids and (2) 22-chromosome hypertriploid hybrids. The 21-chromosome triploids should be similar to the relatively stable hybrids produced from the cross *H. vulgare* (2x) × *H. bulbosum* (4x).

Cross	Number	Gen	otype and	d Chromo	some Nur	mber		ed Genomic stitution
Combination	of	V	VV	VB	VBB	VVBB		
(female × male)	Plants	7	14	14	21	28	Embryo	Endosperm
$VV\timesBB$	1544	1517	_	26	1	_	1V:1B	2V:1B
$BB\timesVV$	35	35	_	_	_	—	1V:1B	1V:2B
$VV\timesBBBB$	87	_	_	_	87	—	1V:2B	2V:2B
$BBBB\timesVV$	6	_	_	_	6	—	1V:2B	1V:4B
$VVVV\timesBB$	4	_	4	_	—	_	2V:1B	4V:1B
$VVVV\timesBBBB$	79	_	76	_	_	3	2V:2B	4V:2B
$BBBB\timesVVVV$	34	—	34	—	—	—	2V:2B	2V:4B

Table 4.8 Types and Frequencies of Progeny Obtained from Interspecific Crosses between *Hordeum* vulgare (V) and *H. bulbosum* (B)

Source: Modified from Kasha, K.J., in *Haploids in Higher Plants: Advances and Potential*, Ainsworth Press, Canada, 1974, 67–87.

	Number of	Total	Frequency and	(%) of Progeny w	vith 2 <i>n</i> =	Expected	
Trisomic Type	Florets Pollinated	Progeny Obtained	21 Chromosomes	22 Chromosomes	Others ^a	Trisomic Transmission	χ² Value
1	2537	81	69(85.2)	11(13.6)	1(1.2)	15.0	0.06
2	2843	57	55(96.5)	1(1.8)	1(1.8)	22.5	10.73 ^b
3	1015	75	72(96.0)	2(2.7)	1(1.3)	20.3	11.44 ^b
4	975	39	28(71.8)	10(25.6)	1(2.6)	29.6	0.19
5	1651	60	52(86.7)	8(13.3)	0(0.0)	20.0	1.31
6	2283	99	80(80.0)	18(18.2)	1(1.0)	16.1	0.17
7	1924	95	79(83.2)	14(14.7)	2(2.1)	20.0	0.60
Total	13,228	506	435(86.0)	64(12.6)	7(1.4)		

Table 4.9Chromosome Number Segregation in Progenies of Crosses between Seven Primary
Trisomics (2n = 2x + 1 = 15) of Barley and Autotetraploid (2n = 4x = 28) Hordeum bulbosum

^a Aneuploids.

^b Significant at 1% level.

Source: From Ho, K.M. and Kasha, K.J., Genetics, 81, 263-275, 1975. With permission.

The frequency of 22-chromosome hybrids should be similar to the usual transmission frequency of the extra chromosome through female gametes in trisomic plants. However, if a specific chromosome carries a factor or factors controlling chromosome elimination, the 22-chromosome plant is not expected to survive, or its frequency will be much lower in crosses with a plant trisomic for that chromosome (Table 4.9). This is because the controlling genes on this chromosome will be in the ratio of 1V:1B, and the *bulbosum* chromosomes will be eliminated. Therefore, it is expected to find 8-chromosome (2n = x + 1) haploid plants of *H. vulgare* if they are able to survive following chromosome elimination.

C. Haploid-Initiator Gene in Barley

A partially dominant haploid-initiator gene (*hap*) was identified by Hagberg and Hagberg (1980) in barley. The haploid-initiator gene controls the abortion or the survival of abnormal embryos and endosperms. Plants homozygous for the *hap* gene produce progeny that include from 10 to 14% haploids. It is clearly shown in barley that male sperm nuclei reach the synergid cells about 1 h after pollination. One of the two male sperm nuclei reaches the two polar nuclei and forms a triploid endosperm. In plants of *hap/hap* and *hap/+* genotypes, the egg cell is not always reached by the other sperm nucleus. Thus, in a *hap/hap* plant, evidently about half of the eggs stay unfertilized, and some of these develop into haploid embryos. The frequency of haploid occurrence is highly influenced by the genotype and also by the environment. Using the *hap* system with marker genes, the breeders do not need to use embryo culture technique. However, they have to make a large number of crosses — a greater number than is needed using the *H. bulbosum* technique (Hagberg and Hagberg, 1987).

IX. MALE STERILITY

Male sterility is common in higher plants. It occurs spontaneously in natural populations and may be induced in mutation experiments. Kaul (1988) estimates that in about 175 species, male sterility (*ms* genes) has arisen spontaneously, and in about 35 species, male sterile genes have been induced by mutagenesis.

The *ms* genes cause complete breakdown of microsporogenesis, but macrosporogenesis typically remains uninfluenced. By contrast, prophase-I and prophase-II meiotic mutants affect microand macrosporogenesis, resulting in chromosomally unbalanced spores, causing male and female sterility. On the other hand, male and female spores are functionally normal in plants with selfincompatibility systems, but self-fertility is genotypically controlled. Compared to normal fertile plants, male sterile plants possess smaller anthers, shrunken and nondehiscent.

A. Classification of Male Sterility

In general, male sterility is divided into three different types: (1) genetic, (2) cytoplasmic, and (3) cytoplasmic–genetic. Genetic male sterility produces nonfunctional androeciums or absence of pollen in plants in several ways. One of these is termed structural and includes structural malformation of stamens into pistils, nondehiscence of anthers because of the absence of anther pores, or the absence of stamens. In a second type, termed functional, viable pollen is formed, but normal anther dehiscence is prevented by barriers such as faulty, or lack of, exine formation. A third basis, termed sporogenous, is the result of a scarcity of pollen, due to the abortion of microsporogenous cells in premeiotic, meiotic, and postmeiotic mutants (Figure 4.1).

Genetic male sterility exhibits Mendelian inheritance. The majority of *ms* genes represent monogenic recessive genes. In some cases, male sterility is controlled by a single dominant gene. Occasionally, it is due to several recessive genes (Jain, 1959; Gottschalk and Kaul, 1974; Driscoll, 1986; Kaul, 1988). The soybean contains 9 (*ms1* to *ms9*) genetic male sterile (female sterile and male fertile) lines. All are recessive in nature (Palmer, 2000).

Cytoplasmic male sterility is inherited maternally, and regardless of the nuclear constitution, plants carrying sterile cytoplasm are male sterile (Edwardson, 1970). Cytoplasmic-genetic male sterility involves the interaction of cytoplasm and nuclear factors. The progenies of cytoplasmic-genetic male sterile plants can be fertile when certain genetic stocks carrying nuclear fertility-restorer genes are used as pollinators (Allard, 1960).

Male sterility that is not genetically inherited is also produced by chemicals (male gametocides) or by changing the atmospheric temperature, light intensity, soil conditions, and growing conditions (field versus greenhouse).

B. Mechanism of Male Sterility

Most male sterility in plants has a sporogenous basis. On the other hand, functional and structural *ms* genes are relatively less prevalent (Kaul, 1988). The *ms* genes act with remarkable precision at definite meiotic stages (premeiotic, meiotic, and postmeiotic, as shown in Figure 4.1) and impair the normal development of sporogenous tissues, tapetal cells, pollen mother cells, and microspores.

A relatively smaller number of premeiotic *ms* genes act on sporogenous cells by causing degeneration of anthers (anthers remain rudimentary in the flowers), archesporial tissues, and pollen mother cells. Some genes are effective during meiotic divisions. They disturb the normal sequence of meiosis or affect a particular stage of meiosis, causing abnormal chromosome behavior and microspore formation (Kaul, 1988).

Many *ms* genes act during postmeiosis, particularly immediately after the tetrad stage. Their action typically blocks normal pollen formation, resulting in empty, nonfunctional pollen grains. At the same time, the tapetal cell layer begins to disorganize, and much of the tapetal cytoplasm is replaced by vacuoles. The tapetum plays an important role in microsporogenesis, especially by transporting nutrients to developing microspores. A deformed tapetum provides an insufficient supply of nutrients to microspores, thus starving them (Cooper, 1952; Rick and Butler, 1956; Filion and Christie, 1966; Mian et al., 1974; Chauhan and Kinoshita, 1979; Albertsen and Phillips, 1981; Dundas, Saxena, and Byth, 1982; Graybosch and Palmer, 1988; Kaul, 1988).

CHAPTER 5

Mode of Reproduction in Plants

I. INTRODUCTION

The mode of reproduction in plants may be sexual (amphimixis), asexual (apomixis or agamospermy), or by specialized vegetative structures (vegetative reproduction). Sexual reproduction involves pollination, germination of pollen grains on the stigma, growth of pollen tubes down the style, the entrance of pollen tube to the ovule through the micropylar opening, the discharge of two sperm nuclei into the embryo sac and fusion of a sperm nucleus with the egg nucleus, producing an embryo, and conjugation of a second sperm nucleus with the two polar nuclei (secondary nucleus) forming a triploid endosperm. Thus, seeds develop as the result of double fertilization, and the majority of plants reproduce sexually (Figure 5.1). The disruption of the sexual process may lead to apomixis.

Apomictic plants of some species produce seed directly from chromosomally unreduced megaspore mother cells or from somatic cells of the nucellus or ovule, without fertilization. In addition, several plants propagate by specialized vegetative structures, like bulbs, corms, cuttings, runners, rhizomes, tubers, and by grafting.

This chapter reviews the salient cytogenetic information on the mode of reproduction in plants (Stebbins, 1950; Maheshwari, 1950; Nygren, 1954; Fryxell, 1957; Bashaw, 1980; Grant, 1981; Nogler, 1984; Hanna, 1991, 1995; Asker and Jerling, 1992; Chapman and Peat, 1992; den Nijs and van Dijk, 1993; Ramachandran and Raghavan, 1992; Naumova, 1993; Koltunow, Bicknell, and Chaudhury, 1995; Jefferson and Bicknell, 1996; Ramulu et al., 1999). The genetic basis of apomixis, its prevalence in polyploids, and exploitation to enhance yield of major crops such as wheat, rice, and maize (apomixis-revolution) are discussed.

II. SEXUAL REPRODUCTION

Sexual reproduction is a complex biological activity that facilitates genetic diversity and speciation. In flowering plants, the reproductive organs are in the flower. Meiosis and fertilization are two essential processes in the sexual cycle of higher plants. Sexual reproduction consists of two generations — sporophytic and gametophytic. The sporophytic generation begins when an egg nucleus unites with a sperm nucleus, producing an embryo, and the second sperm nucleus fuses with the polar nuclei (secondary nucleus), producing triploid endosperm, and continues with the development of seed, seedling, mature plant, and flowers. Sporophytic or somatic tissues contain a diploid (2n) chromosome number. The flower contains spore-forming organs called anthers and ovaries through meiosis. Anthers and the ovaries produce haploid (n) microspores and

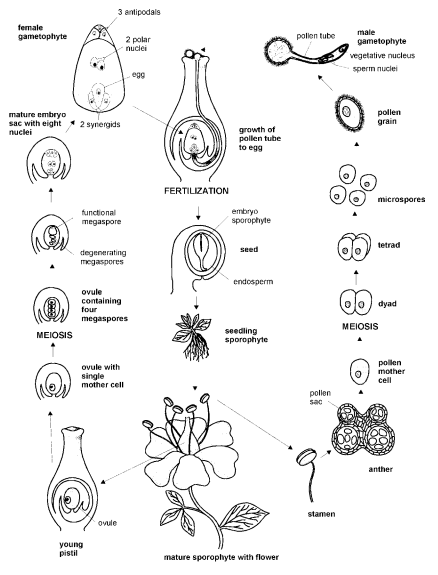


Figure 5.1 The sexual life cycle (amphimixis) in plants. (Redrawn from Nelson, G.E., Robinson, G.G., and Boolootian, R.A., *Fundamental Concepts of Biology*, 2nd ed., John Wiley, New York, 1970, 189.)

megaspores, respectively. Thus, alternation of sporophytic and gametophytic generations is a rule in sexual reproduction (Figure 5.1).

III. ASEXUAL REPRODUCTION

Asexual reproduction (uniparental) in plants may occur without sexual fusion of the spores (apomixis) or by vegetative organs (specialized vegetative structures). Apomixis is common in forage grasses, citrus, mango, blackberries, guayule, and ornamental shrubs. Vegetative propagation occurs by stem modification (bulbs, corms, runners, rhizomes, and tubers), root modification (tuber roots), and flowers, by inflorescence modification (vivipary). Furthermore, generation of plants by grafting and cell and tissue cultures is also included in vegetative propagation.

A. Apomixis

The term "apomixis" was coined by Winkler (1908) and replaced the misused term "apogamy." Apomixis is a type of asexual reproduction in which seed (sporophyte) is produced from the female gametophyte without fertilization. The offspring of apomictic plants are exact genetic replicas of the maternal plant. Apomixis is synonymous with asexual seed formation or agamospermy (Asker and Jerling, 1992). Solntseva (1976) defined apomixis in angiosperms as a method of seed production in which the embryo develops from the cells of the gametophyte with various disruptions of the sporogenesis and sexual processes.

Apomixis is widely distributed in flowering plants (Grant, 1981), and the literature is inundated with reviews, research articles, and proceedings of conferences and symposia. Darlington (1939a, 1958) considered apomixis as a reproductive mechanism for escaping sterility but an evolutionary blind alley; an escape guided in one or several steps by natural selection; but it is an escape that leads to extinction. Stebbins (1950) proposed in *Variation and Evolution in Plants* that there is no evidence that apomicts have ever been able to evolve a new genus or even a subgenus. In this sense, all agamic complexes are closed systems, resulting in an evolutionary blind alley. Their proposal can be accepted under a single dominant gene hypothesis and at the same ploidy level. The predominance of apomixis in polyploid apomictic populations strongly supports this model (Grimanelli et al., 1998b).

However, the current knowledge of apomixis does not support Darlington's blind alley concept or the Stebbins' speciation statement. On the contrary, de Wet and Harlan (1970) contended, based on experimental evidence from *Capillipedium*, *Dichanthium*, and *Bothriochloa*, and a compilospecies *B. intermedia*, that sexual and asexual polyploid species are in an active stage of evolution. Evolutionary mechanisms like hybridization, recombination, polyploidy, clonal selection, and somatic mutation are constantly generating new sexual and apomictic forms or species.

1. The Compilospecies Concept

Harlan and de Wet (1963) stated: "The term compilospecies is taken from the Latin compilo: to snatch together and carry off, to plunder, or to rob. A compilospecies is genetically aggressive, plundering related species of their heredities, and in some cases it may completely assimilate a species, causing it to become extinct." The compilospecies concept is based on the behavior of the *Bothriochloa intermedia* and cytogenetic structure of *Bothriochloa–Dichanthium–Capillipedium* apomictic complex (Figure 5.2). In this complex, sexual and asexual reproduction are independent and genetically active. Their habitats are always contiguous, producing a number of intertaxonomic gene flow interactions among species and even genera.

The major agamospecies are tetraploid, and all are connected directly or indirectly with the compilospecies *B. intermedia* (Figure 5.2). The tetraploid cytotype predominates and cytogenetically behaves like segmental alloploids. Preferential bivalent formation insures production of genetically and physiologically balanced and, hence, functional male and female sexual spores, even among the most apomictic biotypes. It is likely that chromosome pairing in alloploids may be under genetic control. The extensive introgression generates high degrees of heterozygosity and polymorphism. The entire complex (18 species) has an extremely dynamic cytogenetic system, ploidy level 2*x* to 6*x*, and is in a progressive state of evolution (de Wet and Harlan, 1970).

2. Sources of Apomixis

Apomixis is widespread throughout the plant kingdom and is most common in members of the families Poaceae, Compositae, and Rosaceae (Nygren, 1954; Asker and Jerling, 1992; Ramulu et al., 1999). The majority of wild polyploid perennial relatives of cultivated species harbor the apomixis, and only a small proportion of cultivated crops in natural populations are apomictic.

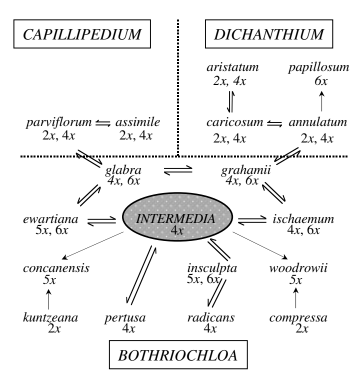


Figure 5.2 Three major agamospecies (*Capillipedium, Dichanthium, Bothriochloa*) are directly or indirectly connected with the compilospecies *intermedia*. The gene flow is continuous introgression (double line with half arrow) and hybridization without further introgression (solid line). (Redrawn from de Wet, J.M.J. and Harlan, J.R., *Evolution*, 24, 270–277, 1970.)

Apomixis can be induced by chemicals and irradiation and is isolated by wide hybridization. The occurrence of apomixis is rare in the diploid species, and more than 90% of the apomictics are usually polyploids (Grant, 1981; Asker and Jerling, 1992).

Alien species are the most reliable source for isolating apomixis. Wild relatives of wheat [e.g., *Elymus scabrus* (Hair, 1956) and *Elymus rectisetus* (Liu, Wang, and Carmen, 1994)]; maize [e.g., *Tripsacum dactlyloides* (Farquharson, 1955; de Wet, 1979; Dewald and Kindiger, 1994; Leblanc et al., 1995a,b)]; and pearl millet [e.g., polyploid *Pennisetum* species (Dujardin and Hanna, 1984; Chapman and Busri, 1994)] carry gene(s) for apomixis. Furthermore, many tropical grass genera such as *Panicum*, *Paspalum, Cenchrus, Eragrostis, Poa, Elymus, Hordeum, Brachiaria, Setaria, Saccharum, Sorghum*, etc., are a rich source of apomixis (Chapman, 1992). A number of apomictic species have been identified in the *Bothriochloa–Dichanthium–Capillipedium* complex (de Wet and Harlan, 1970). Some species of *Malus, Rubus*, and wild species of strawberry (*Fragaria*) are apomictic (Nogler, 1984). Apomixis predominates in cross-fertilizing plants but has not developed in self-fertilizing plants.

Certain types of apomixis can be induced by mutagenesis in some crops. Ramulu et al. (1998) developed various strategies to induce and isolate mutants for apomixis in *Arabidopsis* and *Petunia* by treatment with ethylemethanesulfonate (EMS) and by transposon mutagensis. Ohad et al. (1996) screened ~50,000 EMS treated M_1 plants of *Arabidopsis thaliana* and identified a total of 12 lines from M_2 and M_3 families that displayed elongated siliques in the absence of fertilization-independent endosperm (*fie*). The *fie* allele (female gametophyte) is the source of signals that activate sporophytic fruit and seed coat development, not transmitted by the female gametophyte. The inheritance of the *fie* allele by the female gametophyte results in embryo abortion, even when the pollen carries the wild-type *FIE* allele (essential for female reproductive development). Ohad et al. (1996) concluded that *fie* gene has the genetic potential of inducing autonomous endosperm formation in *Arabidopsis thaliana*, as observed in certain apomictic plants.

A similar approach by Chaudhury et al. (1997) in *Arabidopsis thaliana* was pursued. They isolated six putative fertilization-independent seed mutants (*fis*) by treating the stamenless pistillata (*pi*) mutant with EMS. The *pi* mutant is characterized by short siliques without seed. The *fis* mutant in the *pi* background bears long siliques with developing seeds, though anthers are pollenless. Three mutant alleles (fis_1 , fis_2 , fis_3) are independent and are associated with three different chromosomes. They suggested that fis_3 and *fie* (described by Ohad et al., 1996) may be allelic. The gametophytic nature of *fis* mutants indicates that their time of action is downstream from the point at which apomixis operates in plants. In normal sexual reproduction, *FIS* genes are likely to play a key regulatory role in the development of seeds after normal pollination and fertilization.

Facultative apomixis has been induced in *Pennisetum glaucum* (Hanna and Powell, 1973; Arthur, Ozias-Akins, and Hanna, 1993) and *Sorghum bicolor* (Hanna, Schertz, and Bashaw, 1970). The expression of apomixis in these experiments was variable and low. These examples indicate that apomixis in diploid plants can be induced by mutagenesis.

Apomixis can be produced through hybridization of sexual and apomictic plants and may be transferred to the cultigen through cytogenetic or molecular methods (Hanna and Bashaw, 1987). The introgression of gene(s) for apomixis from hexaploid (2n = 6x = 54) perennial East African grass species *P. squamulatum* to synthetic autotetraploid pearl millet has been extensively attempted (Dujardin and Hanna, 1983, 1984, 1987; Hanna et al., 1993; Chapman and Busri, 1994). Hanna et al. (1993) isolated seven apomictic BC₄ plants. The expression of the apomictic trait showed that most highly apomictic plants had 2n = 27 or 29 chromosomes. No BC₄ plants were obligately apomictic, although one plant produced 89% maternal type, and six of the seven BC₄ plants produced a few offspring that formed only aposporous embryo sacs. This study failed to transfer the apomictic trait from a noncultivated wild perennial species to the cultigen. Lubbers et al. (1994) isolated molecular markers that may facilitate the identification of gene(s) for apomixis. It may be possible to transfer these genes to pearl millet through nonconventional methods, such as somatic hybridization and genetic transformation.

No gene(s) for apomixis have been found in the major economically important crops (wheat, barley, rice, maize, and soybean). Polyembryony is common in several cultivars of Citrus (Koltunow, Hidaka, and Robinson, 1996). Apomixis is being utilized to produce disease-free scion and uniform root stocks in Citrus.

3. Distribution of Apomixis

Apomixis is distributed throughout the plant kingdom from algae to angiosperms (Asker and Jerling, 1992; Koltunow, Bicknell, and Chaudhury, 1995). It has been estimated that apomixis occurs in about 300 genera belonging to 80 families (Khokhlov, 1976). Apomixis has an uneven taxonomic distribution occurring mostly in polyploid species of Poaceae, Rosaceae, and Asteraceae, and is often associated with fertile hybrids that otherwise would have been sterile (den Nijs and van Dijk, 1993). In the Citrus group, however, apomixis is not related to polyploidy, but it is related to hybridization.

4. Types of Apomixis

Two main types of apomixis are known: gametophytic apomixis (gametophytic agamospermy) and adventitious embryony (nucellar) (Figure 5.3). The two forms of apomixis are unrelated (Asker and Jerling, 1992). In gametophytic apomixis, an egg cell develops parthenogenetically from an unreduced embryo sac, producing only maternal-type offspring. In contrast, adventitious embryony does not have a gametophytic stage but only a sporophytic stage, and it is connected with parthenogenesis. Embryos develop from somatic cells of the nucellus or integument. Thus, alternation of generation is eliminated (Grant, 1981).

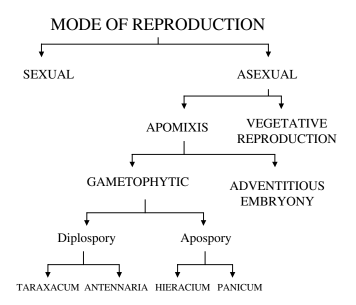


Figure 5.3 Classification of the mode of reproduction in plants.

a. Gametophytic Apomixis

Gametophytic apomixis has been divided into diplospory and apospory (Figure 5.3). Both forms maintain the alternation of generations, but the gametophytic and sporophytic stages contain the same level of ploidy (Nogler, 1984). Meiosis and fertilization are circumvented in both forms, and an unreduced egg cell develops asexually by parthenogenesis. Although the predominant mode of reproduction in gametophytic apomixis is obligate, it may sometimes be combined with sexuality. Such a type is known as facultative apomixes, and frequency is influenced by several factors.

i. Diplospory — Diplospory (generative apospory) occurs in plants where meiosis is absent or development of a linear tetrad of megaspores does not occur. It is subdivided into two forms named after the genera in which they were first discovered (Figure 5.3):

- *Taraxacum*: In this type, the megaspore mother cell enters the meiotic prophase-I, but chromosomes remain as univalents resulting in restitution nuclei with full somatic chromosome complement. The second division is normal producing dyads with unreduced chromosome number. Usually the chalazal dyad cell gives rise to an eight-nucleate embryo sac after three mitoses (Figure 5.4). There are four divisions of nuclei and cells instead of the five observed in normal embryogenesis (Figure 5.4). The *Taraxacum* type of apomixis, in which embryo and endosperm develop without fertilization, is known as mitotic diplospory or autonomous apomixis and is found among genera of the Compositae (*Taraxacum, Erigeron, Chondrilla*), rarely in *Arabis holboellii* and *Agropyron scabrum*, and in certain *Paspalum* species (Nogler, 1984; Asker and Jerling, 1992).
- *Antennaria:* The megaspore mother cell does not undergo meiosis but directly undergoes three mitotic divisions, producing an eight-nucleate unreduced (diploid) embryo sac (Figure 5.4). The *Antennaria* type is also called mitotic diplospory and has a wide taxonomic distribution (Nogler, 1984).
- Other types of diplospory: Deviation from the Taraxacum and Antennaria types of diplospory has been observed.
 - *Eragrostis*: It is similar to the *Antennaria* type, but with the difference that the embryo sac formed has four nuclei, because the third mitosis is omitted. Thus, cells and nuclei have two divisions (Figure 5.4).
 - *Ixeris*: An asyndetic meiotic chromosome pairing produces a restitution nucleus, as observed in the *Taraxacum* type, which then undergoes a second meiotic division without cytokinesis.

Type of cell development of the spore		Sporoç	genesis	Mother	Game	etophytoge	nesis		visions and
Type of development	of the		ision	cell of embryo sac		Division		Mature embryo sac	of div uclei cells
		1	2		1	2	3		No. of n the
Normal				٦					5
Taraxacum	۲			٢					4
Antennaria									3
Eragrostis									2

Figure 5.4 Megagametogenesis in amphimixis and various apomictics.

This results in two unreduced instead of four reduced nuclei. The two mitoses produce an eight-nucleate embryo sac. The *Ixeris* type of diplospory is found in *Ixeris dentata*.

Allium: Premeiotic chromosome doubling by endomitosis or endoreduplication in the female produces unreduced nuclei. Meiosis is normal and yields a tetrad with unreduced nuclei of parental genotypes. Two subsequent mitoses in the chalazal dyad result in an eight-nucleate embryo sac. It is observed in *Allium nutans* and *A. odorum*.

ii. Apospory — Apospory, earlier termed "somatic apospory" (Nogler, 1984), is characterized by the development of unreduced embryo sacs directly from the somatic (vegetative) cells located in the center of the nucellus. There may be multiple embryo sacs in an ovule, but only one of them matures into an aposporous embryo sac (Nogler, 1984; Asker and Jerling, 1992). Aposporous apomixis is of two main types: the bipolar type *Hieracium* and the monopolar *Panicum* type (Figure 5.3).

Hieracium: The initial cell produces an eight-nucleate bipolar embryo sac by mitotic divisions along with the development of the Polygonum type observed in the related sexual taxa. Thus, an ovule may contain two or more eight-nucleate embryo sacs, one is reduced which may degenerate, and others are unreduced (Khokhlov, 1976). Apospory is widely distributed in several families, e.g., in the genera of *Hypericum*, *Poa*, *Ranunculus*, *Crepis*, *Hieracium*, *Hierochloe*, and *Beta* (Nogler, 1984).
Panicum: The unreduced being four-nucleate monopolar embryo sac is produced after the second mitosis, and the third mitosis is absent. This type of apomixis was discovered in *Panicum maximum* (Warmke, 1954) and is found in grasses belonging to Panicoideae and Andropogoneae (Nogler, 1984; Asker and Jerling, 1992). It is widely distributed in the genera of the *Bothriochloa–Dichanthium–Capillipedium* complex, *Cenchrus*, *Chloris*, *Digitaria*, *Eriochloa*, *Heteropogon*, *Hyparrhenia*, *Panicum*, *Paspalum*, *Pennisetum*, *Sorghum*, *Themeda*, and *Urochloa* (Nogler, 1984). An exceptional case is recorded within the Panicoideae. The unreduced aposporous sacs are usually four-nucleate, and the reduced ones are eight-nucleate and bipolar.

b. Adventitious Embryony

The terms nucellar or adventive embryony are synonymous with adventitious embryony (Asker and Jerling, 1992). Adventitious embryony is an asexual reproduction in which adventitious embryos develop from one or more of the nucellar and integumentary cells of the ovule.

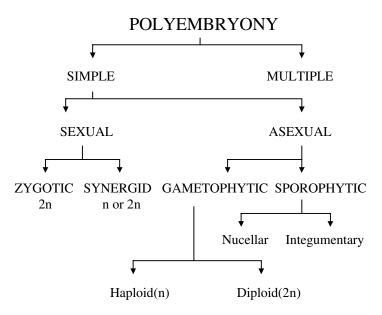


Figure 5.5 Classification of polyembryony in plants.

Adventitious embryony differs from diplospory and apospory in that it lacks embryo sacs and produces viable seeds. Thus, these embryos do not contain polar nuclei or endosperm. Double fertilization is apparently absent and, even if present, it is influenced by environmental conditions (Naumova, 1993). Meiosis is usually normal with reduced chromosome numbers in embryo sacs.

Adventitious embryos can be induced in plants experimentally through tissue culture. The production of somatic embryos (embryoids, i.e., embryo-like structures) *in vitro* is a prerequisite for regeneration of plants; however, adventitious embryos and embryoids are not always produced in a similar way (Naumova, 1993).

Adventitious embryony is widespread in economically important crops belonging to the genera *Citrus*, *Euphorbia*, *Mangifera*, *Malus*, *Ribes*, *Beta* and several genera of grasses. It has been identified in more than 250 species of 121 genera belonging to 57 families of flowering plants (Naumova, 1993).

Polyembryony: Polyembryony, the occurrence of more than one embryo in a seed, may be simple or multiple (Figure 5.5). Simple polyembryony is divided into sexual and asexual. Polyembryony from sexual reproduction may be zygotic–zygotic polyembryony, develop from suspensor–suspensor polyembryony or synergid cells — synergid polyembryony (Lakshmanan and Ambegaokar, 1984). Polyembryony is controlled by several genes. It may be autonomous where no induction is necessary, or induced when pollination is required. Asexual polyembryony may arise from gametophyte or sporophyte. The gametophyte polyembryony can be haploid or diploid, and the sporophytic form is from nucellar and integumentary cells (Figure 5.5).

5. Endosperm Development in Apomixis

a. Autonomous Endosperm Development

Plants with autonomous apomixis diploid parthenogenesis do not require pollination, and neither the egg nucleus nor the polar nuclei is fertilized. The central cell (polar nuclei) develops parthenogenetically, producing endosperm without fertilization, resulting in autonomous endosperm development (Asker and Jerling, 1992). Autonomous endosperm usually contains variable chromosomes due to the fusion of polar nuclei and endomitosis. Autonomous endosperm development is rather common in most apomictic Compositae (Nogler, 1984) and Asteraceae (Asker and Jerling, 1992).

b. Pseudogamous Endosperm Development

Fertilization of the polar nuclei is required for endosperm development. According to Asker and Jerling (1992), pseudogamy includes cases where embryo formation takes place independently of pollination, i.e., embryos develop before pollination, which is necessary for the production of endosperm and germinable seed. Pseudogamous apomicts often exhibit poor seed set. Seed development depends on a precise ratio of maternal (m) and paternal (p) genomes in the endosperm (Haig and Westoby, 1991). This ratio is called endosperm balance ratio (EBN) and is usually 2 m:1 p. The failure in crosses between autotetraploid and diploid-related species is attributed to the departure from EBN.

A parthenogenic embryo may develop in association with a viable endosperm. Thus, the development of an embryo into a seed is a preadaptation for apomixis. Morgan, Ozias-Akins, and Hanna (1998) isolated a tetraploid, partially male-fertile, aposprous apomictic line 169–46 from *Pennisetum glaucum* $(2n = 4 \ x = 28) \times P$. *squamulatum* (2n = 6x = 54). This interspecific F₁ hybrid (2n = 42; expected 41) was crossed with a colchicine-induced amphidiploid (2n = 6x = 42) between pearl millet and Napier grass (*P. purpureum*, 2n = 4x = 28). This double-cross apomictic hybrid was crossed as a male to a tetraploid pearl millet three times. In each generation, Morgan, Ozias-Akins, and Hanna (1998) selected apomictic and partially fertile plants and eventually identified a tetraploid apomictic pearl millet line 169–46. The central-cell nuclei (secondary nucleus) of 169–46 are unreduced (2n = 4x = 28), and on fusion with 14 chromosome gametes from pearl millet, produce an endosperm with 4 m:1 p genomic ratio. The maternal and paternal genomes are similar in 4x pearl millet, and thus, the m:p ploidy ratio in the endosperm should equal the m:p endosperm balance ratio (2:1). Therefore, 4 m:1 p endosperm imbalance causes poor seed set in 169–46. By BC₃, autotetraploid pearl millet exhibits meiotic irregularities but has fairly good seed set. Apomictic diploid pearl millet has not yet been identified (Hanna, W.W., personal communication, 2002).

6. Identification of Apomixis

Apomixis can be identified by morphological, cytological, genetic, biochemical, and molecular methods:

- 1. Uniform progeny from a single plant is an indicator, i.e., matromorphic progeny.
- 2. Maternal type progeny appears in the F_1 ; thus, no genetic variation is expected in the F_2 .
- 3. In crosses between a recessive genotype as a female parent and dominant genotype as a male, exclusively recessive genotype is obtained.
- 4. A high degree of polyploidy is usually associated with apomixis and is constant from parent to the progeny.
- 5. An unusually high degree of fertility in aneuploids, triploids, wide crosses, or other plants expected to be sterile is an indication of apomixis.
- 6. Multiple embryos, seedlings, stigmas, and ovules suggest an indication of apomixis (Hanna, 1991).
- 7. Cytohistological examinations are useful tools in identifying and verifying various types of apomixis.
- Both biochemical and molecular techniques are currently being used to screen apomixis at the seedling stage in *Pennisetum* species (Lubbers et al., 1994) and *Poa pratensis* (Mazzucato et al., 1995). Mazzucato, den Nijs, and Falcinelli (1996) developed an auxin test to determine frequency of parthenogenesis.

7. Classification of Plants Based on Natural Crossing

- 1. *Dioecious plants*: Staminate and pistillate flowers are produced on separate plants of a species; entirely cross pollinated.
- 2. Apomictic plants: Union of spores is not involved; cross pollination is required.
- 3. *Self-incompatible plants*: Both male and female spores are produced at the same time and are functional but fail to fertilize and produce seeds; entirely cross pollinated.

- 4. Dichogamous plants: Plants include protandry and protogyny ensuring cross pollination.
- 5. Cleistogamous plants: Flowers remain closed and are completely self-fertilized.

8. Regulation of Apomixis

Despite numerous publications, the inheritance of apomixis is not clearly demonstrated and the genetics of apomixis are not clearly understood. Although apomixis is genetically controlled, it is also influenced by environmental factors like temperatures and light regimes changes (Gerstel, Hammond, and Kidd, 1955; Bashaw, 1980; Hanna and Bashaw, 1987; Hanna, 1991, 1995; Asker and Jerling, 1992; Ramachandran and Raghavan, 1992; den Nijs and van Dijk, 1993; Lutts, Ndikumana, and Louant, 1994). Hanna et al. (1973) observed that in guineagrass, sexuality was controlled by a dosage effect of two or more dominant alleles present at different loci or at a single locus. It has been demonstrated that apomixis is controlled by qualitative traits, mainly by recessive and dominant genes, and in some species where apomixis is facultative, polygenes may be involved. An intermediate to recessive inheritance with quantitative dosage effects for apomixis has been suggested in polyploid species of Rubus (Chirsten, 1952; Berger, 1953), but a dominant gene for apomixis in apple was reported (Sax, 1959). Based on inheritance of apospory in buffelgrass, Sherwood, Berg, and Young (1994) postulated a two-locus model for tetrasomic transmission in which the dominant allele A of one locus is required for apospory but is hypostatic to the dominant allele B of the second locus, which confers sexuality. Bicknell, Borst, and Koltunow (2000) recorded monogenic dominant inheritance in *Hieracium piloselloides* (2n = 3x = 27) and *H. aurantiacum* (2n = 3x + 4 = 31). They could not recover diploid apomictic plants from these crosses due to selection against the survival of diploid zygotes. The dominant genes in both species are closely linked or possibly allelic. This shows that apomixis has a genetic basis, but how it is regulated is still unresolved. Genetic analysis of Paspalum simplex showed apomixis to be under the control of a single dominant gene (Pupilli et al. 2001). den Nijs and van Dijk (1993) concluded that different genetic mechanisms for apomixis may exist even within the same species. More critical studies are needed to clearly understand these mechanisms.

Grimanelli et al. (1998a) supported a one-dominant-gene hypothesis for the failure of meiosis in *Tripsacum*. However, the apomictic process may be controlled by a cluster of linked genes inherited in a tetrasomic fashion. van Dijk et al. (1999) did not find a single locus control for apomixis in *Taraxacum* and suggested that several genes were involved in the control of apomixis. In a subsequent study, Grimanelli et al. (1998b) proposed non-Mendelian transmission of apomixis in maize–*Tripsacum* hybrids. A gene (or genes) controlling apomixis in *Tripsacum* was linked with a segregation distortion-type system, promoting the elimination of the apomixis allele when transmitted through haploid gametes. This protects the diploid level from being invaded by apomixis.

Occasional occurrence of unreduced egg cells or parthenogenesis in sexual plants may not be related to apomixis. Three sets of genes, acting at different reproductive stages of plants, may be responsible for apomixis: (1) failure of reduction in number of chromosomes, (2) failure of fertilization, and (3) development of nonreduced unfertilized egg cells.

The mode of reproduction in facultative apomictic plants is often influenced by environmental factors such as day length, light intensity, temperature shocks, plant age, nutrition supply, exposure to certain growth hormones, or growing environments (greenhouse versus field). Knox and Heslop-Harrison (1963) reported that in *Dicanthium aristatum*, an 8-h light treatment produced inflores-cence with 79% aposporous embryo sacs, but a 16-h photoperiod produced only 47%. Chapman (1992) recorded 60% apomictic embryos in the same species with a photoperiod excess of 14 h, while more than 90% were found with less than 14 h. Contradictory results were observed in *Poa ampla*, where a 20-h daylight regime produced more sexuality (Williamson, 1981). Sexuality was promoted in *Poa pratensis* when plants were grown and flowered in early spring in the glasshouse, but it was not recorded in plants grown in the field (Grazi, Umaerus, and Åkerberg, 1961). Temperature and seasonal variations have been found to have significant influence on the expression

of apomixis in *Malus* (Schmidt, 1977). Similar results were reported in *Cenchrus ciliaris* (syn. *Pennisetum ciliare*), where frequency of sexual pistils differed between sampling dates in one field, indicating that environment may influence development of embryo sac type (Sherwood, Young, and Bashaw, 1980).

In *Brachiaria decumbens*, reproductive behavior appeared similar between greenhouse- and field-grown plants, but in *B. brizantha*, a much lower level of sexuality was observed in field-grown plants. Sexual reproduction was more frequent in adverse climatic conditions in *Paspalum cromoy- orrhizon* (Quarin, 1986).

Occasional isolation of apomixis in plants is due to pollinations: condition of pollination, choice of pollinators, condition of pollen applied (fresh versus old), or time of pollination (early versus delayed). Martínez, Espinoza, and Quarin (1994) observed that an early pollination by 2x and 4x pollen in an apomictic tetraploid (2n = 4x = 40) *Paspalum notatum* sometimes prevented parthenogenetic development of the unreduced egg cell in an aposporous embryo sac allowing fertilization. Wide hybridization often induces apomictic seed development. It may be due to slow pollen tube growth, or sperm nuclei may fail to fertilize the egg. In sexual plants, wide hybridization, pollination with irradiated pollen, or chemically treated pollen promote haploid formation (Asker and Jerling, 1992).

To conclude, apomixis is under complex genetic control; however, eight genetic models for the inheritance of apomixis can be proposed from the published results:

Model I: One disomic locus, apospory recessive (aa).

Model II: One disomic locus, apospory dominant (AA, Aa).

- Model III: Two disomic loci, additive gene action, requires at least two dominant alleles for sexuality (*Aabb*, *Aabb*, *aaBb*, *aabb* genotypes apomictic)
- Model IV: Two disomic loci, dominant allele A is required for apospory, dominant allele B confers sexuality and is epistatic to dominant gene A (Aabb, Aabb \rightarrow apomictic).

Model V: One tetrasomic locus — apospory recessive (aaaa) to sexuality.

- Model VI: One tetrasomic locus, dosage effect (Aaaa, AAAa, AAAA); sexual genotype aaaa.
- Model VII: One tetrasomic locus, diplospory dominant (Aaaa, Aaaa, AAAa, AAAA).

Model VIII: One disomic dominant but controlled by a cluster of linked loci.

9. Apomixis in Crop Improvement

Apomixis helps fix heterosis in a desired heterozygous gene combinations. Extreme heterozygosity is a characteristic of plants reproducing by apomixis. It fixes new heterozygous genotypes with valuable agronomic traits and resistance to pests and pathogens (Asker and Jerling, 1992; Nassar, 1994). Although apomixis is a potentially powerful tool in plant breeding, its application in crop improvement is limited to turf and forage grasses (Hanna, 1991; Voigt and Tischler, 1994).

Obligate apomixis may suppress sexuality completely. Thus, a significant reduction in cost of hybrid seed production could be avoided. Because outcrossing within a population in the commercial field with obligate apomicts is not a problem, it can be used for other vegetatively propagated crops. Purity can be maintained by controlling mechanical mixture during harvest. Farmers will not be required to buy hybrid seed every year, because they could use seeds indefinitely without the risk of contamination from recombination. Apomixis would be disadvantageous to seed industries, because it would result in a loss of control of commercial hybrids and reduction in seed sales. In obligate apomixis, progeny testing for stability and field isolation are not needed in commercial seed production. It can be effective in a breeding program, provided that cross-compatible sexual and partial sexual plants are available to allow generation of new gene combinations.

Introgression of the apomixis from *Tripsacum dactyloides* (2n = 2x = 36; 2n = 4x = 72) to maize (*Zea mays*) was extensively attempted by a number of research groups (Harlan and de Wet, 1977; Grimanelli et al., 1998a,b; Hoisington et al., 1999; Blakey, Goldman, and Dewald, 2001). Two pathways were used: maize $2n = 20 \times Tripsacum 2n = 2x = 36 \rightarrow 2n = 28 \rightarrow 2n = 20$ pathway (the most common way) and maize $2n = 20 \times Tripsacum 2n = 4x = 72 \rightarrow 2n = 46 \rightarrow 2n = 56 \rightarrow 2n = 56$

 $2n = 38 \rightarrow 2n = 20$ pathway (Harlan and de Wet, 1977). Harlan and de Wet (1977) isolated highly tripsacoid maize lines with 2n = 20 chromosomes. Dominant resistance to six maize diseases were found in BC₈ populations, but no apomictic line was recovered.

As of March 1999, 10 patents related to apomixis, including for maize, pearl millet, and rice, have been issued (Bicknell, 1999). Currently, 20 patents related to apomixis have been granted (http://gb.espacenet.com). Despite enormous research and financial resources from public, private, and international institutes like CIMMYT, apomictic maize has not been commercialized.

Facultative apomixis has been identified in some plant groups for utilization in breeding programs. The best source of apomixis is the distantly related species of the cultigens. The main emphasis is on isolation, identification, and incorporation of obligate or facultative apomixis in several major crops such as *Citrus*, berries, apple, fodder grasses, rubber plants (*Parthenium*), pearl millet, maize, wheat, barley, rice, *Sorghum*, sugar beet, *Brassica*, pea, soybean, alfalfa, cotton, potato, tobacco (Asker and Jerling, 1992). In rice, wild relatives failed to induce apomixis (Rutger, 1992); however, Zhou et al. (1993) discovered a rice line SAR-1 that showed a high degree of pollen sterility, but seed set under isolation was 55.33%. Cytoembryological examination revealed division of eggs without fertilization, or adventitious embryo developed from cells of ovary wall and produced seed. Endosperm developed normally to provide nutrients for embryo development. The apomixis trait is heritable and is being incorporated for rice varietal improvement.

Several factors should be examined carefully before apomixis can be successfully used to produce new cultivars. These are facultative behavior, number of genes involved, modifiers, environmental factors, ploidy levels, and seed sterility. Although apomixis provides a unique opportunity to develop and maintain superior genotypes, its use is restricted because of lack of basic understanding of the mechanism of apomixis (Asker and Jerling, 1992; Hanna, 1995). Studies on synteny among grass species have revealed homoeologous regions in chromosomes. For example, a chromosome segment conditioning apomixis in *Paspalum simplex* is homoeologous to the telomeric region of the long arm of rice chromosome 12 (Pupilli et al., 2001). Ortiz et al. (2001) discovered that *Paspalum notatum* (2n = 20) linkage groups 1, 3, 4, 5, 6, 8, 10 have synteny regions with maps of maize and rice. Apomictic gene(s) could be isolated and transferred to desired sexual diploid crops through transformation, provided such a system is available.

B. Vegetative Reproduction

Vegetative reproduction is an asexual form of reproduction in higher plants by which new identical individuals are generated from a single parent without sexual reproduction. The offspring of asexual propagation is known as a clone. The principal method of asexual reproduction in higher plants is by vegetative propagation. It is characterized by mitosis which occurs in the shoot and root apex, cambium, intercalary zones, callus tissues, and adventitious buds.

Vegetative propagation facilitates easy, rapid multiplication of economically valuable heterozygous plants without alteration and does not induce genetic diversity. The perpetuation of seedless orange, banana, and grapes are excellent examples.

1. Vegetative Reproduction by Specialized Vegetative Organs

Many plants multiply through specialized vegetative structures by modifying stems. Common examples are as follows:

- Bulbs Bulbs are a short basal, underground stem surrounded with thick, fleshy leaves, common in the onion, daffodil, hyacinth (family Liliaceae). Bulbils or bulblets are miniature bulbs used in propagation.
- Corms Corms are short, upright, hard or fleshy bulb-like stems usually covered with papery, thin, dry leaves and do not contain fleshy leaves. It is common in gladiolus, crocus, and water chestnut. Cormels are miniature corms. Fleshy buds develop between the old and new corms.

- 3. *Runners (Stolons)* Runners are a horizontal above-ground stem that usually produce plants by rooting at nodes. Strawberry reproduces vegetatively via stolons.
- 4. *Rhizomes* A rhizome is a horizontal, prostrate, or underground stem that contains nodes and internodes of various length and readily produces adventitious roots. Species with rhizomes are easily propagated by cutting the rhizomes into small pieces that contain a vegetative bud. Examples of species with underground rhizomes are Johnsongrass, brown grass, and hops.
- 5. *Tubers* A tuber is fleshy portion of a rhizome, underground storage stem. Potato is the best example of a species with tubers. A small piece of potato with a bud (known as an eye) is planted to produce more potatoes. This ensures uniformity in the next generation, as each eye generates a replica of its parent.

Vegetative propagation is also common by tuberous roots like sweet potato. Tuberous roots of some species may contain shoot buds at the "stem end" as part of their structure. Other examples include dahlia and begonia. The primary tap root develops into an enlarged tuberous root that can be propagated by dividing into several portions, each with a bud (Janick et al., 1981).

2. Vegetative Reproduction by Adventitious Roots and Shoots

Reproduction of an entire plant from a buried branch or stem is called layering or layerage. Once new roots and shoots emerge, plantlets are separated from the mother plants. Cutting is one of the most important methods of vegetative reproduction. Small pieces of stems (cuttings) are used by horticulturists and nurserymen for multiplying and reproducing ornamental crops.

3. Vegetative Reproduction by Grafting

Reproduction of an entire plant by union of a small actively growing shoot (scion) grafted onto root-stock that is resistant to pathogens and pests is an invaluable tool in plant propagation. Grafting is quite common for a large number of domestic fruit crops to produce disease-free crops.

4. Vegetative Reproduction by Tissue Culture

Plant propagation through cell and tissue cultures is termed micropropagation. It involves regeneration of plants aseptically from cells (including protoplasts) and tissues (immature embryos, leaves, roots, and stems) in artificial cultures. A single protoplast can regenerate a whole plant. Thus, a large number of plantlets can be generated via embryogenesis and organogenesis from a small piece of the stock plant. Micropropagation can result in the isolation of disease-free plants and has proven efficient for orchid propagation, where the natural propagation rate is slow.

5. Advantages of Vegetative Reproduction

Vegetative reproduction (propagation) has numerous advantages, such as efficient commercial crop production (potato and sugarcane), exploitation of heterosis (hybrid vigor is not lost), avoidance of dormancy and juvenile period (grafting on older root stocks allows new wood of seedling to produce fruit sooner than if it remains on its own root stock), maintenance of sterile or lethal genotypes, facilitation of physiological and genetic studies, and increased plants of unique genotypes in breeding programs.

Despite many advantages, vegetative reproduction has a serious problem. All vegetatively propagated plants from the same source are genetically uniform. This means that genetic vulnerability exists. If the genotype is susceptible to a pest or pathogen, or if a new pest or a pathogen develops that can infect the genotype, then all plants of the clone will be susceptible. If a disease strikes the members of a clone, production of disease-free seed or shoot is extremely difficult.

IV. THE CHROMOSOMAL BASIS OF SEX DETERMINATION IN PLANTS

The majority of flowering plant species are hermaphroditic, having flowers with stamens and carpels. In a relatively small number of plant species, sex is determined by the presence or absence of a pair of sex chromosomes (chromosomal mechanism of sex determination). The species may be heterogametic (produce two kinds of gametes and offspring with two sexes) or homogametic (one kind of gamete and offspring). The sex chromosomes (heteromorphic sex pair = heterochromosomes; X and Y) are distinct from all other chromosomes (autosomes). In certain plants, sex determination is under genetic control, where discrete sex-chromosome systems are not discovered and usually influenced by autosomal genes (Burnham, 1962). Heteromorphic sex chromosomes are clearly defined in only a few species (Table 5.1).

A. System of Sex Determination

1. Male Heterogametic (Female XX; Male XY = X-Y System)

The female is homogametic, and all eggs carry an X chromosome. The male is heterogametic and 50% male spores carry an X chromosome, and 50% will contain a Y chromosome. Random fertilization ensures a sex ratio of 1 female:1 male in every generation. The X-Y system is common in *Cannabis sativum* (hemp), *Humulus lupulus* (hop), *Rumex angiocarpus, Silene latifolia* (syn. *Melandrium album*), and *M. rubrum* (Table 5.1). The X-Y system is also prevalent in mammals, including humans, *Drosophila*, and many other species.

2. Male Heterogametic (Female XX; Male XO = X-O System)

The male produces two types of gametes: one with the X chromosome, and the other without a sex chromosome (O). The female produces gametes only with the X chromosome. The X-O system is found only in *Vallisneria spiralis* (2n = 20), and *Dioscorea sinuata* (yam). The X-O system is common in grasshoppers, crickets, and roaches.

3. Male Heterogametic (but with One Extra Chromosome)

This system is proposed in *Phoradendron flavescens* (American mistletoe) and *Phoradendron villosum* [more evidence is needed (Burnham, 1962)].

Species	2 <i>n</i> Female	2 <i>n</i> Male
Cannabis sativum	18 + XX	18 + XY
Humulus lupulus	18 + XX	18 + XY
Humulus lupulus var. cordifolius	$16 + X_1 X_1 X_2 X_2$	$16 + X_1Y_1X_2Y_2$
Humulus japonicus	14 + XX	$14 + XY_1Y_2$
Rumex angiocarpus	12 + XX	12 + XY
Rumex tenuifolius	24 + (XX) XX	24 + (XX) XY
Rumex acetosellas. str.	36 + (XXXX) XX	36 (XXXX) XY
Rumex graminifolius	48 (XXXXXX) XX	48 (XXXXXX) XY
Rumex hastatulus	6 + XX	$6 + XY_1Y_2$
Rumex acetosa	12 + XX	$12 + XY_1Y_2$
Rumex paucifolius	24 + (XX) XX	24 + (XX) XY
Melandrium albumª	22 + XX	22 + XY
Melandrium rubrum	22 + XX	22 + XY

 Table 5.1
 Established Heteromorphic Sex Chromosomes in Plants

a Silene latifolia.

Source: From Westergaard, M., Adv. Genet., 9, 217-281, 1958. With permission.

4. Female Heterogametic [Female XY (ZW); Male XX (ZZ) (Z-W System)]

The female is heterogametic, and the egg determines the sex. The male is homogametic and produces only one kind of gamete. The Z-W system is found in 6x (2n = 42) strawberry (*Fragaria elatior*). This system is absent in all animals, but is found in birds and some insects, including butterflies and moths.

5. Compound Chromosomes (Interchanges among X, Y, and Autosomes)

These chromosomes are found in certain races of common hop and in garden sorrel (*Rumex acetosa*), where a chain of three chromosomes is observed in the heterogametic sex, arranged at meiotic metaphase-I as $Y_1 X Y_2$, with the X oriented to pass to one pole and $Y_1 Y_2$ to migrate to the other pole. In certain cases, there is a chain of five chromosomes in the male. This may be due to interchanges between autosomes and sex chromosomes (Burnham, 1962).

Several dioecious species of plants are without sex chromosomes, but sex expression is under genetic control. These are *Spinacia oleracia, Ribes alpium, Vitis vinifera, Carica papaya, Asparagus officinalis,* and *Bryonia dioica.* The sex-determining gene in the spinach is on Chromosome 1.

B. Determination of Heterogametic Sex in Plants

1. Cytological Identification

A classical example of a heteromorphic pair of sex chromosomes (XX female; XY male) that has been widely studied in plants is white campion, *Silene latifolia* (syn. *Melandrium album*). Cytological investigation establishes that the Y chromosome is larger than the X, and both are larger than the autosomes. Westergaard (1958) divided X and Y chromosomes into four hypothetical segments. Segment IV is homologous in X and Y chromosomes, and these ends pair during meiosis. The other segments of X and Y are nonhomologous (Figure 5.6). In the Y chromosome, Segment I carries the gene (s) that suppresses the development of female sex organs, Segment II initiates anther development, and Segment III controls the last stages in anther development. The sex expression gene in the Y chromosome is present on the segment that is nonhomologous with the X chromosome and is never separated by crossing over (Westergaard 1958).

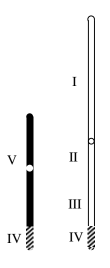


Figure 5.6 Sex chromosomes (X Y) of Silene latifolia (syn. Melandrium album). Segments I, II, and III are different in Y. (Segment IV is the homologous region, and Segment V is the differential region of X.) (Redrawn from Westergaard, M., Adv. Genet., 9, 217–281, 1958.)

Doležel and Göhde (1995) analyzed nuclear DNA content of *S. latifolia* by high-resolution flow cytometry. They found lower DNA content in female than male plants due to sex chromosome heteromorphism. Kejnovský et al. (2001) identified, for the first time, male-specific genes, *MROS*, expressed in *S. latifolia*. The *MROS* genes were located on chromosomes using the flow-sorted X chromosomes and autosomes as a template for PCR with internal primers. Their results indicate that at least two copies of the *MROS3* gene are located in tandem on the X chromosome with additional copy(ies) on autosomes, while *MROS1*, *MROS2*, and *MROS4* are exclusively located on autosomes. They conclude that *MROS3* is a low-copy gene family that is connected with the normal pollen development, present in dioecious and other dicots. For example, *MROS3* homologues are also discovered in a nonrelated *Arabidopsis thaliana* genome. This suggests an ancient origin of the *MROS3* gene. It is possible that the X chromosome of *S. latifolia* has some regions homologous to Chromosomes III and IV of *A. thaliana*.

Another dioecious crop, hemp (*Cannabis sativum*), was extensively investigated and has a sexual system similar to *S. latifolia*. Female plants contain two X chromosomes, whereas male plants have one X and one Y chromosome. The Y chromosome is much larger than the X chromosome and autosomes. By using flow cytometry, Sakamoto et al. (1998) reported a genome size of 1638 Mbp for diploid female (2n = 18 + XX), and 1683 Mbp for diploid male (2n = 18 + XY) plants. Karyotype analysis revealed that the X chromosome was submetacentric and the Y chromosome subtelocentric. The Y chromosome has the longest long arm, with a satellite in the terminal of its short arm (Figure 5.7).

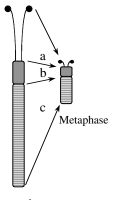
2. Sex Linkage Inheritance

In *Lychnis alba*, broad-leaved female (normal) \times narrow-leaved male produce F_1 plants in which all female plants are broad-leaved, but part of the males are broad-leaved and part are narrow-leaved (Burnham, 1962). The sex-determining gene is on the X chromosome, and no allele is present on the Y chromosome.

3. Crosses between Dioecious and Monoecious Bryonia Species

The segregation in crosses between dioecious and monoecious Bryonia species is shown below:

- 1. Bryonia dioica (dioecious) female \times Bryonia alba (monoecious) male \rightarrow all female
- 2. Bryonia alba (monoecious) female \times Bryonia dioica (dioecious) male \rightarrow 1 female:1 male



Prometaphase

Figure 5.7 Specific condensation of the long arm and satellite of the Y chromosome of *Cannabis sativum.* (a) NOR region; (b) short arm; and (c) long arm. (Redrawn from Sakamoto et al., *Cytologia*, 63, 459–464, 1998.)

- 3. Bryonia dioica (dioecious) female \times Bryonia macrostylis (monoecious) male \rightarrow all female
- 4. Bryonia macrostylis (monoecious) female \times Bryonia dioica (dioecious) male \rightarrow 1 female:1 male
- 5. Bryonia dioica (dioecious) female \times Bryonia multiflora (dioecious) male \rightarrow all monoecious
- 6. Bryonia multiflora (dioecious) female \times Bryonia dioica (dioecious) male \rightarrow 1 female:1 male
- 7. Bryonia macrostylis (monoecious) female \times Bryonia alba (monoecious) male \rightarrow all hermaphrodite
- 8. Bryonia alba (monoecious) male \times Bryonia macrostylis (monoecious) female \rightarrow 1 female:1 male

a. Conclusion

- 1. Dioecious × monoecious species produce 1 female:1 male. This suggests the male is the heterogamete of the dioecious species.
- 2. Crosses between dioecious \times dioecious species may give monoecious progeny.
- 3. Crosses between monoecious × monoecious species may give dioecious species.

4. Competition Tests in Pollination

In *Silene*, pollination by excess pollen produces excess female progeny, but sparse pollination gives 1 female:1 male (male heterogametic). By contrast, in *Cannabis sativum* (hemp), sparse pollination increases the proportion of males.

5. Self-Pollination of Plants that are Normally Unisexual

In Asparagus (2n = 20), only a few seeds are obtained from a male plant, and they segregate into 3 males:1 female. No sex chromosome has been identified cytologically, but sex-determining genes must be on the chromosomes. The XY male $[\otimes] \rightarrow 1XX$ female:2XY males:1YY male; XX female \times YY male \rightarrow all XY and male. This suggests that a dominant gene for maleness is on the Y chromosome.

6. Crosses between Diploids and Autotetraploids

When the male is heterogametic, chromosome doubling in dioecious species (XX and XY) will produce autotetraploids with genotypes XXXX and XXYY. When the female is homogametic, crosses between XXXX female \times YY male \rightarrow 1XXX female:1XXY male; XX female \times XXYY male \rightarrow 1 female:5 males. When the female is heterogametic, crosses between XXYY \times XX \rightarrow 4XXY:1XYY:1XXX (5 females:1 male). In *Spinacia oleracea* (spinach) and also in *Silene latifolia*, male plants are heterogametic.

7. Sex Ratio among Progeny of Primary Trisomics

In spinach, a diploid female \times primary trisomic (Triplo 1) male cross-produces an F₂ population that segregates in a trisomic ratio of 2 females:1 male. This suggests that a gene for sex determination is dominant and is located on Chromosome 1.

C. Sex Expression

Table 5.2 compares sex expression in *Silene latifolia* and *Drosophila melanogaster*. Investigations on sex expression in diploid, triploid, and tetraploid with different dosages of X and Y chromosomes show that a plant is male when one or more Y chromosomes is present, and a plant is female when Y is absent. The expression of female traits is possible when the ratio of X:Y reaches 1:4. Plants with XX chromosomes are females.

Sex expression on the same plant or on different plants is controlled by qualitative or quantitative genes confined to sex chromosomes (X or Y or both) that, through interaction

Chromosome		
Constitution	Silene	Drosophila
1. 2A + XX	Female	Female
2. 2A + XXX	Female	Female
3. 3A + X	—	Male
4. 3A + XX	Female	Hermaphrodite
5. 3A + XXX	Female	Female
6. 4A + XX	Female	Male
7. 4A + XXX	Female	Hermaphrodite
8. 4A + XXXX	Female	Female
9. 4A + XXXXX	Female	—
10. 2A + XY	Male	Male
11. 2A + XYY	Male	Male
12. 2A + XXY	Male	Female
13. 2A + XXYY	—	Female
14. 3A + XY	Male	—
15. 3A + XXY	Male	Hermaphrodite
16. 3A + XXXY	Male	Female
17. 4A + XY	Male	—
18. 4A + XXY	Male	—
19. 4A + XXYY	Male	—
20. 4A + XXXY	Male	—
21. 4A + XXXYY	Male	—
22. 4A + XXXXY	Male \rightarrow Hermaphrodite	—
23. 4A + XXXXYY	Male	_

Table 5.2 Sex Expression (The Ratio of Sex Chromosomes and Autosomes) in *Silene latifolia* (syn. *Melandrium*) and *Drosophila*

Source: From Westergaard, M., Adv. Genet., 9, 217–281, 1958. With permission.

Table 5.3	Terminologies	for Sex	Expression	in Plants
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Type of Plants	Expression of Sex
Hermaphrodite	Perfect flower (bisexual; a flower with androecium and gynoecium)
Monoecious	Separate male and female flowers on the same plant (not synonymous with imperfect and should never be applied to individual flower)
Dioecious	Separate male and female flowers on different plants
Andromonoecious	Perfect and male flowers on the same plant
Gynomonoecious	Perfect and female flowers on the same plant
Trimonoecious	Perfect, female, male flowers on the same plant (polygamous)
Androdioecious	Perfect and female flowers on different plants
Gynodioecious	Perfect and female flowers on different plants

Source: From Westergaard, M., Adv. Genet., 9, 217-281, 1958. With permission.

with sex genes on the autosomes, determine which sex will be expressed (Westergaard, 1958). Terminologies have been assigned for sex expression in plants based on presence or absence of sex organs (Table 5.3)

Based on the system of sex determination in diploid and polyploid plants, two kinds of mechanisms are evident: the active Y chromosome plays a decisive role as an enhancer for maleness and a suppressor for gynoecism in determining the sex, and the sex-deciding genes are in the X chromosome, and the sex depends upon X chromosome/autosome ratios, with the Y chromosome being inactive (Westergaard 1958).

D. The Evolution of Dioecism

It has been demonstrated that in higher plants, hermaphroditism condition is the original form of sexuality. Dioecism originated from bisexual or monoecious species through mutation and natural selection. The unisexual condition developed as a result of a trigger mechanism that suppresses the potentialities of the opposite sex in males and females. The forward evolution is from bisexual to unisexual, and the reverse evolution is from dioecism to bisexuality.

CHAPTER 6

Karyotype Analysis

I. INTRODUCTION

An individual displays its characteristic 2n chromosome number — half maternal and half paternal. A deviation in either direction (+ or -) results in chromosomal imbalance. Cytological techniques, described in Chapter 2, determine the chromosome constitution of an organism and facilitate recognition of the individual chromosomes. Three terms, namely, karyotype, karyogram, and idiogram, are often referred to in the identification of chromosomes. Karyotype is the number, size, and morphology of a chromosome set of a cell in an individual or species (Battaglia, 1994). Karyogram is the physical measurement of the chromosomes from a photomicrograph, where chromosomes are arranged in descending order (longest to shortest). An idiogram represents a diagrammatic sketch (interpretive drawing) of the karyogram (Figure 6.1). The classification of chromosomes is based on physical characteristics, such as size of chromosomes, features of telomere, position of kinetochore, secondary constriction, size and position of heterochromatic knobs, and relative length of chromosomes (Figure 6.1).

Karyotype analysis is usually based on somatic mitotic metaphase chromosome measurement. This can be estimated with two assumptions: the exact length (image parameter) and chromosomes within a complement exhibit an even condensation (condensation pattern) (Fukui and Kakeda, 1994). Metaphase chromosomes (maximum condensation) are obtained after pretreatment (Figure 6.2). However, differential condensation among chromosome arms may be the cause of variation in relative length values.

Occasionally, conventional staining techniques do not distinguish chromosomes of similar morphological features. In this case, Giemsa C- and N-banding techniques have helped identify these chromosomes (Singh and Tsuchiya, 1981b; Schlegel, Melz, and Nestrowicz, 1987; Gill, Friebe, and Endo, 1991). Furthermore, the recent adaptation of the *in situ* hybridization technique for plant chromosomes, particularly for cereals, has also facilitated the process of distinguishing morphologically similar chromosomes.

II. NOMENCLATURE OF CHROMOSOMES

The kinetochore (centromere) position is a useful landmark for the morphological identification and nomenclature of chromosomes (Battaglia, 1955; Levan, Fredga, and Sandberg, 1964; Matérn and Simak, 1968; Naranjo, Poggio, and Brandham, 1983). Battaglia (1955) disagreed with using the term kinetochore for the centromeric region of the chromosome body. A monocentric chromosome contains one centromere and nomenclatured chromosomes based on the position of the centromere in the chromosomes:

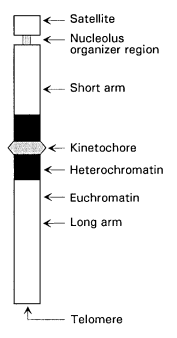
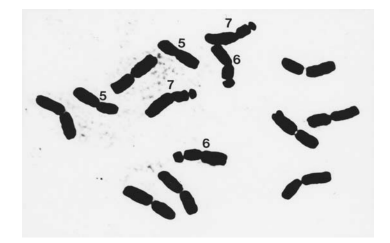
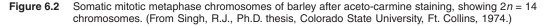
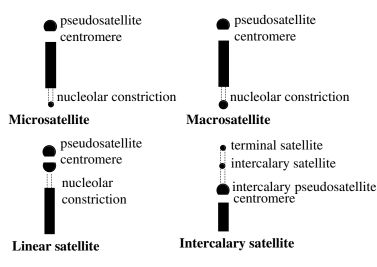


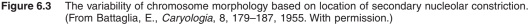
Figure 6.1 An idiogram of a metaphase chromosome.

- 1. *Median centromere* (isobrachial chromosomes): Centromere is situated in the middle of the chromosome, resulting in an arm ratio of 1:1.
- 2. *Submedian centromere* (heterobrachial chromosome): Centromere is located near the middle of the chromosome, resulting in an arm ratio of more than 1:1 but less than 1:3 (from 1:1 to 1:2.9).
- 3. *Subterminal centromere* (hyperbrachial chromosome): Centromere is near one extremity of the chromosome, resulting in a ratio of 1:3 or more.
- 4. *Terminal centromere* (monobrachial chromosome): Centromere is situated at one extremity of the chromosome, resulting in an arm ratio of 0:1.









Battaglia (1955) considered satellite as a part of the chromosome distal to a nucleolar constriction, and it is universally accepted that for each satellite, there is one nucleolus. It is an established fact that a satellite is a spheroidal body and diameter that is the same or smaller than the diameter of the chromosomes, situated at one extremity, and connected to the chromosome body by a thin thread (Figure 6.3). The position of a satellite may be terminal (a satellite between its nucleolar constriction and one extremity) or intercalary (a satellite between two nuclear constrictions). Satellite chromosome is designated based on size and location (Figure 6.3):

- 1. *Microsatellite*: A spheroidal satellite of small size, i.e., having a diameter equal to or less than one half the chromosomal diameter.
- 2. *Macrosatellite*: A spheroidal satellite of large size, i.e., having a diameter greater than one half the chromosomal diameter.
- 3. Linear satellite: A satellite having the shape of a long chromosomal segment.

Levan, Fredga, and Sandberg (1964) discussed thoroughly the nomenclature of chromosomes, and only salient points of their paper will be taken into consideration in this chapter. The relative lengths of the long arm (l) and short arm (s) are shown by the arm ratio (r = l/s). Based on arm ratio, Levan, Fredga, and Sandberg (1964) grouped chromosomes in six categories (Table 6.1).

Table 6.1 Nomenclatu	re of Chromosomes		
Centromere Position	Arm Ratio (I/s)	Chromoso	ome Designations
Median <i>sensu stricto</i>	1.0	М	(Metacentric) ^a
Median region	1.7	m	(Metacentric) ^a
Submedian	3.0	Sm	(Submetacentric)
Subterminal	3.0	St	(Subtelocentric)
Terminal	7.0	t	(Acrocentric)
Terminal sensu stricto	~	Т	(Telocentric)

^a Levan, Fredga, and Sandberg (1964) recommended terms to discard. Because these terms are often used in chromosome nomenclature, I suggest they not be abandoned. *Source:* Based on Levan, A., Fredga, K.F., and Sandberg, A.A., *Hereditas*, 52, 201–220, 1964. Naranjo, Poggio, and Brandham (1983) proposed a modified version of the Levan, Fredga, and Sandberg (1964) nomenclature. They divided chromosomes into eight equal units. Like Levan, Fredga, and Sandberg (1964), they felt difficulties in assigning chromosomes to a particular type when the arm ratio was exactly 5:3, 6:2, or 7:2, which were three boundary ratios separating the four intermediate groups m (median region), sm (submedian), st (subterminal), and t (terminal). They suggested that the chromosomes with these arm ratios must be classified as m-sm, sm-t, or st-t. In a random sample of chromosomes, only a few will fit into these categories, and most of the chromosomes will fall in one of the six types identified in Table 6.1.

III. KARYOTYPE ANALYSIS BY MITOTIC METAPHASE CHROMOSOMES

Karyotype analysis has played an important role in the identification and designation of chromosomes in many plant species. Barley (*Hordeum vulgare* L.) is cited here as an example. Barley is a basic diploid, contains 2n = 2x = 14 chromosomes, and the chromosome-linkage group relationship has been established based on cytogenetics and molecular studies.

In barley, Chromosome 5 is the smallest chromosome, and Chromosomes 6 and 7 are the nucleolus organizer chromosomes that are morphologically distinct. Based on conventional staining techniques, Chromosomes 1 to 4 are difficult to distinguish (Figure 6.2). Chromosome designations of Tjio and Hagberg (1951) were accepted by barley cytogeneticists until Tuleen (1973) and Künzel (1976) questioned the identity of Chromosomes 1, 2, and 3. Both authors observed, based on the results of multiple translocation analysis, that Chromosome 2 is the longest chromosome in the barley complement. The application of the Giemsa C-banding (Figure 6.4) and N-banding techniques helped identify all the barley chromosomes, while it was not possible by conventional staining techniques (Linde-Laursen, 1975; Singh and Tsuchiya, 1981b, 1982a, 1982b; Kakeda, Fukui, and Yamagata, 1991). Furthermore, the combination of aceto-carmine and Giemsa staining technique applied to the same cell (Nakata, Yasumuro, and Sasaki, 1977) helped to construct karyotype analysis better than either technique alone.

The salient features of the seven barley chromosomes based on conventional and Giemsa Cand N-banding techniques (Singh and Tsuchiya, 1982a,b), and homoeologous (in parenthesis) groups (Costa et al., 2001) with wheat are described below.

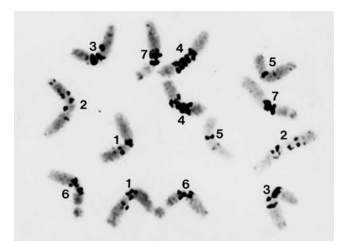


Figure 6.4 Somatic mitotic metaphase chromosomes of barley after Giemsa C-banding. (From Singh, R.J. and Tsuchiya, T., *Z. Pflanzenzüchtg.*, 86, 336–340, 1981b. With permission.)

A. Chromosome 1 (7H)

This is the third longest chromosome and is metacentric (Figure 6.5; Table 6.2). In four cells (Cells 1, 3, 7, 9) out of 10 cells measured, the long and short arms of Chromosome 1 were equal in length. In three cases (Cells 4, 5, 10), the long arm was longer than the short arm, and in three other instances (Cells 2, 6, 8), the short arm was longer (Table 6.2). Because both arms were almost equal in measurement, their designation in karyogram and idiogram depended on the morphological effects of telotrisomic plants, gene-chromosome arm relationships, and also on the Giemsa N-banding pattern (Singh and Tsuchiya, 1982a,b).

B. Chromosome 2 (2H)

This is the longest chromosome among the five nonsatellited chromosomes of the barley complement. It carries its kinetochore at the median (l/s = 1.26) region (Tables 6.2 and 6.3; Figure 6.5). Similar results were reported by Tuleen (1973) and Künzel (1976).

C. Chromosome 3 (3H)

Tjio and Hagberg (1951) identified Chromosome 3 as a median (arm ratio = 1.09) chromosome. If Chromosomes 1 and 3 of Tjio and Hagberg (1951) are switched, their results will agree with the results presented in Table 6.2. Chromosome 3 showed a dark centromeric band. The band on the short arm appeared as a large block at metaphase. The long arm had a dark interstitial band (close to the kinetochore) and a faint dot on each chromatid in the middle of the long arm (Figure 6.5).

D. Chromosome 4 (4H)

This chromosome contains its kinetochore at the median region (l/s = 1.21) and was correctly identified in all of the studies. Conventional staining techniques do not distinguish Chromosome 4 from Chromosomes 1, 2, and 3. However, based on Giemsa C- and N-banding techniques, Chromosome 4 was easily distinguished from the rest of the chromosomes, because it is the most heavily banded in the barley complement; about 48% of the chromosome is heterochromatic. Sometimes it is difficult to locate the centromere position in condensed Giemsa-banded metaphase chromosomes (Figure 6.4). However, the appearance of a diamond-shaped centromere position and the use of aceto-carmine stained Giesma N-banding technique facilitated the precise localization of the kinetochore (Figure 6.5).

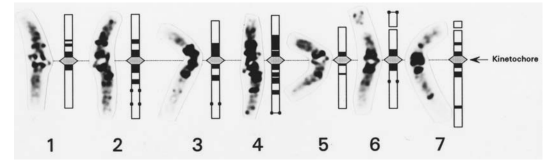


Figure 6.5 Karyogram and idiogram of Giemsa N-banded chromosomes of barley. (Redrawn from Singh, R.J. and Tsuchiya, T., J. Hered., 73, 227–229, 1982b.)

						Chror	nosome	Chromosome no. and relative arm length (%)	relative a	rm lengt	h (%)					
		1		2		3		4		5		6		~		
Cell Number	–	s	–	s	–	s	-	s	_	s	-	s	-	s	6 Sat	7 Sat
-	7.69	7.69	9.23	6.92	9.23	6.15	7.69	6.92	7.69	5.38	7.69	4.61	8.46	4.61	2.30	1.53
2	7.69	8.09	8.90	7.29	9.72	6.47	8.90	5.67	7.29	4.85	7.29	4.04	9.72	4.04	2.42	1.62
e	6.98	6.98	9.56	8.08	8.46	6.61	8.08	6.99	7.35	5.15	6.61	4.42	9.56	5.14	2.20	1.47
4	8.43	7.23	8.43	7.23	9.04	6.62	7.83	6.63	7.23	5.42	7.23	4.21	9.64	4.82	2.41	1.80
5	8.41	7.96	8.85	7.52	8.85	6.19	7.08	6.19	7.08	5.31	7.08	4.42	10.62	4.42	1.77	1.77
6	7.14	7.93	9.52	7.14	9.52	6.74	7.93	5.95	6.74	4.76	7.14	4.37	9.92	5.16	2.38	1.58
7	7.41	7.41	9.05	6.58	9.47	6.58	7.41	6.58	7.41	4.94	8.23	4.94	9.05	4.94	2.47	1.65
8	6.36	7.07	9.19	7.06	9.89	7.06	7.77	7.36	7.77	5.65	7.42	4.24	9.54	4.59	2.82	2.12
6	7.64	7.64	8.92	7.32	9.55	7.01	7.64	6.37	7.64	5.09	7.01	4.45	8.92	4.77	2.22	1.27
10	8.62	7.66	8.62	6.71	8.62	5.75	7.66	6.70	7.66	5.75	6.70	3.83	10.92	4.78	1.91	1.91
Mean	7.64	7.57	9.03	7.19	9.24	6.52	7.80	6.44	7.39	5.23	7.24	4.36	9.64	4.73	2.29	1.67
95% confidence limit	±0.51	±0.27	±0.26	±0.29	±0.34	±0.28	±0.34	±0.29	±0.23	±0.23	±0.34	±0.21	±0.53	±0.24	±0.21	±0.17
Relative length ^b	65.89	65.28	77.88	62.06	79.68	56.24	67.29	55.53	63.72	45.14	62.46	37.57	83.12	40.78	19.75	14.42
95% confidence limit	±4.39	±2.34	±2.23	±2.56	±2.95	±2.48	±2.93	±2.56	±1.97	±2.04	±2.90	±1.85	±4.58	±2.07	±1.81	±1.48
Arm ratio (I/s)	1.01		1.26		1.42		1.21		1.41		1.66 ^a		2.04ª			
l = long arm; s = short arm.	t arm.															

Table 6.2 Relative Chromosome Arm Length (%), Mean Arm Ratios of Hordeum vulgare cv. Shin Ebisu (SE 16)

^a Arm ratios do not include satellite. ^b Based on 100 units for both arms of Chromosome 6.

Source: From Singh, R.J. and Tsuchiya, T., Theor. Appl. Genet., 64, 13-24, 1982a. With permission.

			Arm			Arm			Arm			Arm			Arm			Arm			Arm
			ratio	,	.:	ratio	ы.		ratio	4	,	ratio	5.		ratio	6.		ratio	7		ratio
Authors	–	s	l/s	–	s	s/I	–	s	l/s	–	s	l/s	–	s	l/s	–	s	l/s	–	s	l/s
Tijo and Hagberg 78.4 (1951) 63.7	78.4 63.7	Ω Ω	8.5 1.34 7 8.6 1.09	71.7	61.1	61.1 1.17	63.7 78.4	58.6 58.5	1.09 1.34	67.2	67.2 51.8 1.30 60.7 44.3 1.37 62.1 37.9 1.64 78.5 32.2	1.30	60.7	44.3	1.37	62.1	37.9	1.64	78.5	32.2	2.44
Tuleen (1973)	64.5	ဖ	1.05	75.0	60.5	1.24	74.5	56.8	1.31												
Künzel (1976)	66.5	64.2	1.04	73.9	63.0	1.17	72.4	58.5	1.24	68.2	58.0	1.18	63.6	44.4	44.4 1.43	62.1	36.7	1.69	79.8	37.7	2.12
Singh and Tsuchiya (1982a)	65.9	65.3	1.01	77.9	62.1	1.25	79.7	56.2	1.42	67.3	55.5		63.7	45.1	1.41	62.5	37.6	1.66	83.1	40.8	2.04
Source: From Singh, R.J. and Tsuchiya, T., Theor.	jh, R.J.	and Ts	uchiya,	T., The		Appl. Genet., 64, 13-24, 1982a. With permission.	., 64, 1	3-24, 1	982a. M	Vith per	mission										

Table 6.3 Comparison of Relative Chromosome Arm Length and Arm Ratios in Barley, Observed by Several Authors

KARYOTYPE ANALYSIS

E. Chromosome 5 (1H)

This chromosome is the shortest among the five nonsatellited chromosomes of barley and has an arm ratio (1.42) similar to Chromosome 3 (Tables 6.2 and 6.3). It has a centromeric band and an intercalary band on the long arm and a band on the short arm that is darker than those on long arm (Figure 6.5).

F. Chromosome 6 (6H)

This chromosome has a larger satellite than Chromosome 7 and has an arm ratio of 1.66 (without the satellite). A similar observation was also recorded by other workers (Table 6.3). Chromosome 6 showed a dark centromeric band in both arms, a faint intercalary band on the long arm, and a faint dot on each chromatid on the telomere of the satellite (Figure 6.5).

G. Chromosome 7 (5H)

This chromosome has the longest long arm in the barley karyotype and carries a submedian kinetochore (Tables 6.2 and 6.3). It showed an equally dense centromeric band at the distal portion of the long arm, and a faint intercalary band was also observed in the short arm (Figure 6.5).

The literature on karyotype studies is voluminous. It has been shown here that a combination of several techniques facilitates the construction of a karyogram of a crop species better than one technique alone.

IV. KARYOTYPE ANALYSIS BY PACHYTENE CHROMOSOMES

Sometimes, pachytene chromosomes are used when somatic chromosomes do not show distinguishing landmarks. The classical examples for conducting karyotype analysis based on pachytene chromosomes are in maize (McClintock, 1929b), tomato (Barton, 1950), *Brassica* (Röbbelen, 1960), and rice (Khush et al., 1984). Singh and Hymowitz (1988) constructed a karyogram (idiogram) for the soybean (an economically important oil seed crop) pachytene chromosomes for the first time. The soybean contains a high chromosome number (2n = 40), small and similar (symmetrical) chromosome size (1.42–2.84 µm), and lacks morphological landmarks. Individual somatic chromosomes of the soybean have not been clearly distinguishable, however, only a pair of satellite chromosomes is occasionally visible (Singh, Kim, and Hymowitz, 2001).

No reliable techniques are known that consistently produce a high degree of success with squash preparations of soybean meiocytes (Palmer and Kilen, 1987). Singh and Hymowitz (1988) found that it was not possible to trace all 20 pachytene chromosome bivalents in a single cell, but the isolation of 1–3 bivalents, observations on euchromatin and heterochromatin differentiation, and chromosome measurements (Figure 6.6) facilitated the identification and construction of a pachytene chromosome map (an idiogram) of the soybean for the first time (Figure 6.7). The heterochromatin is distributed proximal to and on either side of the centromeres on the long and short arms, and 6 of the 20 short arms are totally heterochromatic. This latter feature makes soybean pachytene chromosomes unique.

V. KARYOTYPE ANALYSIS BY FLOW CYTOMETRY

Flow cytogenetics may be defined as the use of flow cytometry to sort and analyze individual chromosomes and physical mapping of genes of economic importance using FISH technology.

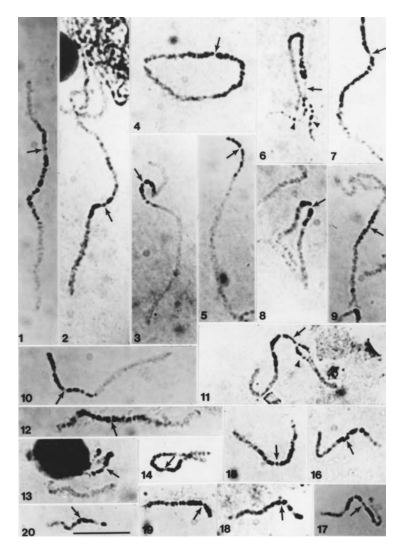


Figure 6.6 The pachytene chromosome complement of *Glycine max* × *G. soja* F₁ hybrid. Each figure shows a different chromosome. For example: (1) Chromosome 1 to (20) Chromosome 20. Arrows indicate centromere location. (From Singh, R.J. and Hymowitz, T., *Theor. Appl. Genet.*, 76, 705–711, 1988. With permission.)

However, this technique has not been universally successful in plants because of lack of quality chromosomes and an inability to resolve a single chromosome on a flow karyotype.

Lucretti et al. (1993) invented a procedure to sort only metacentric chromosome of *Vicia faba* by flow cytometry. They located rDNA locus by FISH. Preparation of high-quality chromosome suspensions is a prerequisite for successful chromosome sorting and karyptyping. They listed several factors for poor quality chromosomes in suspensions, such as the splitting of metaphase chromosomes into chromatids, chromosome breakage, chromosome clumping, presence of interphase nuclei, and presence of cellular and chromosomal debris.

Flow cytometry may be a valuable tool if we can distinguish aneuploid from diploid plants based on relative surplus or deficit of DNA content. With this view in consideration, Samoylova, Meister, and Miséra (1996) examined a relative surplus of DNA content in *Arabidopsis* primary and telotrisomics in interphase nuclei measured by flow cytometer to distinguish diploid (wildtype) from trisomic plants. They measured differences in nuclear fluorescence intensity between

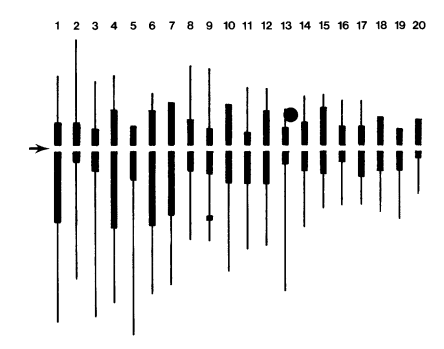


Figure 6.7 Proposed idiogram, based on Figure 6.6, of the pachytene chromosomes of the soybean. Arrow indicates centromere location. (From Singh, R.J. and Hymowitz, T., *Theor. Appl. Genet.*, 76, 705–711, 1988. With permission.)

diploid and trisomics. The relative surplus of genomic DNA recorded by primary and telotisomics was attributed to the extra chromosome (Figure 6.8). However, flow karyotype contradicts cytological observation. Cytologically, Chromosomes 5 and 3 are larger than Chromosomes 4 and 2 (the smallest), but relative DNA content (%) was found lesser for Chromosomes 5 and 3 than 4 and 2 (Figure 6.8). At this stage, we may conclude that flow karyotyping for plants needs to be perfected.

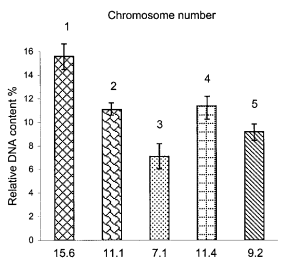


Figure 6.8 The flow karyotype of *Arabidopsis thaliana* interphase chromosomes. (From Samoylova, T.I., Meister, A., and Miséra, S., *Plant J.*, 10, 949–954, 1996. With permission.)

VI. KARYOTYPE ANALYSIS BY IMAGE ANALYSIS

This procedure, chromosome image analyzing system (CHIAS), was provided by K. Fukui. The beginning of chromosome image analysis goes back to the early 1980s, when computer systems were at the cradle stage in handling huge digital data of images. There were only a few expensive image analyzing systems available, and imaging techniques suitable for plant chromosomes analysis were under development. Some trial studies were carried out using human chromosomes, especially in the field of semiautomatic identification of human chromosomes (Casperson, Lomakka, and Moller, 1971; Castleman and Melnyk, 1976; Lundsteen et al., 1980).

In 1985, the first comprehensive chromosome image analyzing system (CHIAS) with software fulfilling the basic requirements of cytologists and cytogeneticists was developed (Fukui, 1986). Then, further development of imaging methods, such as quantifying chromosome morphology and its band patterns in barley (Fukui and Kakeda, 1990), quantifying uneven condensation patterns appearing at the prometaphase chromosomes in rice (Fukui and Iijima, 1991), simulating human vision for identifying and quantifying chromosome band patterns in *Crepis* (Fukui and Kamisugi, 1995), etc., followed. An imaging method for quantification of pachytene chromomeres was soon released.

No personal computers with enough imaging capability allow image analysis in every cytology and cytogenetics laboratory. The basic points that should be in mind with the application of imaging methods are as follows:

- 1. Importance of the quality of chromosome images No imaging method can create new information that is not originally included in the chromosome images.
- The information of the images is basically reduced by each application of image manipulation Imaging methods present the essence of the chromosome image of the original image as visible and, thus, perceptible.
- 3. Imaging methods can present the image information by numerical data.

The standard and basic procedures for image analyses and the manuals (Kato et al., 1997) can be obtained by written form or via the Internet (http://mail.bio.eng.osaka-u.ac.jp/cell/).

VII. CHROMOSOME NUMBER OF ECONOMICALLY IMPORTANT PLANTS

The comprehension of chromosome number of plants is important for the systematist, cytogeneticist, evolutionist, plant breeder, and molecular geneticist. The chromosome number of plants is compiled in *Chromosome Atlas of Flowering Plants* (Darlington and Wylie, 1955), and in several journals, such as *Taxon, Rhodora, American Journal of Botany, International Organization of Plant Biosystematists Newsletter (ISPOB), Annals of the Missouri Botanical Garden, New Zealand Journal of Botany*, and *Systematic Botany*. Chromosome numbers of some plants are listed in Appendix 6.I.

CHAPTER 7A

Chromosomal Aberrations — Structural Chromosome Changes

I. DEFICIENCIES

A. Introduction

The loss of a segment from a normal chromosome is known as deficiency (Df). The term deletion is often used as a synonym of deficiency. Deficiencies indicate any chromosomal loss, and the term deletions should be confined to a deficiency involving an internal region of a chromosome (McClintock, 1931). Thus, deficiency may be intercalary (Figure 7.1A) or terminal (Figure 7.1B). The deficiency method, which is also known as the pseudodominant method, has been effectively utilized for locating genes in the chromosomes of maize (McClintock, 1931, 1941a; Chao et al., 1996) and tomato (Khush and Rick, 1967b, 1968a).

B. Origin and Identification of Deficiencies

Induced deficiencies may be generated by x-raying pollen carrying normal (wild-type) alleles and applying the pollen to the stigma of female flowers. The female parent carries recessive alleles at loci in the genome that will hopefully be represented in segments of induced deficiency. A majority of the plants show dominant phenotype, but occasionally, a few plants with the homozygous recessive phenotype appear in the progenies. Cytological examination of these plants with recessive phenotype at pachynema often reveals normal association along the entire length of each chromosome, except in one region of one chromosome. This observation is a strong indication that the recessive allele may be carried in hemizygous condition. A loop is generally observed if the deficiency is long enough and is located in an interstitial region. A terminal deficiency results in an unpaired end region.

X-ray-induced deficiencies occur at nonrandom positions in chromosomes. Two reports provided conflicting results in tomato. Gottschalk (1951) found 73.2% centromeric breaks, 18.4% breaks in heterochromatin, and 8.4% in the euchromatin. Khush and Rick (1968a) recorded the highest frequency (60%) of breaks in the heterochromatin. The frequency of breaks in kinetochores was intermediate (20%), and only 15% of breaks occurred in euchromatin.

Fast neutron also induces breaks in chromosomes in a nonrandom fashion (Khush and Rick, 1968a). However, fast neutron was found to be much more efficient than x-rays in inducing breaks in euchromatin in tomato. Both types of radiation produced breaks preferentially in heterochromatin. A terminal deficiency with one break in heterochromatin was the most frequent; a terminal deficiency in euchromatin was not observed. Terminal deficiencies were observed for only those arms

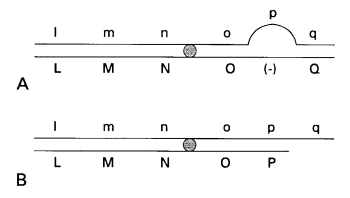


Figure 7.1 Types of deficiencies. (A) Intercalary; (B) terminal.

with losses that were tolerated by gametophytes or sporophytes. Broken ends in a kinetochore or euchromatin did not heal without reuniting with other broken ends. The most frequent type of interstitial deficiency obtained in tomato was the type that resulted from both breaks in the euchromatin. All such breaks were obtained from fast neutron treatment of pollen, and none were induced by x-ray treatment.

In maize, the *r*-X1 deficiency, a small intercalary deficiency located in the long arm of Chromosome 10 that includes the *R1* locus, induces terminal deficiencies, monosomics and trisomics, and nondisjunction of chromosomes in the early embryos (Weber, 1983; Lin, Marquette, and Sallee, 1990; Weber and Chao, 1994).

Several Aegilops species such as Ae. caudata (2n = 14; CC), Ae. cylindrica (2n = 28; CCDD), Ae. geniculata (2n = 28; UUMM), Ae. longissima (2n = 14; S^S), Ae. sharonensis (2n = 14, S^{sh}S^{sh}), Ae. speltoides (2n = 14; SS), and Ae. triuncialis (2n = 28, UUCC) contain a gametocidal gene (Gc) that induces chromosome aberrations in common Chinese Spring wheat (2n = 42, AABBDD) (Endo, 1988, 1990; Endo and Mukai, 1988; Kota and Dvořák, 1988; Ogihara, Hasegawa, and Tsujimoto, 1994; Endo and Gill, 1996; Nasuda, Friebe, and Gill, 1998). The Gc gene in Aegilops species was identified during the production of alloplasmic chromosome alien addition lines between crosses of common wheat and Aegilops species (Endo, 1990). Taxonomic nomenclature of Aegilops species by van Slageren is used as suggested by B.S. Gill (personal communication, 2002; Table 7.75).

The following procedures (Figure 7.2) generated 436 deficiencies in Chinese Spring wheat by using gametocidal gene (Endo, 1990; Endo and Gill, 1996):

- 1. Back cross the monosomic alien addition and translocation lines as female to euploid Chinese Spring wheat as a male.
- 2. Examine the chromosome constitutions of the progeny by C-banding technique.
- 3. Select plants with deficiency or deficiencies and without alien chromosome.
- 4. Self-pollinate deficiency heterozygote plants. Screen the self-progeny cytologically for deficiency homozygous plants with the least degree of aberrations in the other chromosomes. For example, translocations and aneuploidy.
- 5. In case the deficiency homozygote is not found among 10 or more offsprings from the structural heterozygotes, cross the deficiency heterozygote as female with an appropriate nullisomic-tetrasomic, or an appropriate ditelosomic of Chinese Spring to create the deficiency hemizygotes in the F₁ progeny.
- 6. Screen F₂ progeny for deficiency homozygote.

Endo (1988) observed deficiencies and translocations in almost half the progeny of a monosomic addition line of common wheat that carried a chromosome of *Ae. cylindrica*. Chromosome breaks induced by Gc gene occur in various regions of the wheat chromosomes and also in the *Ae*.

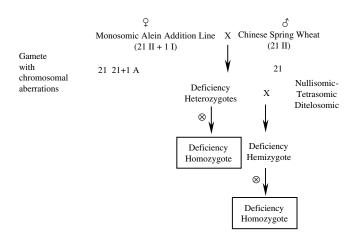


Figure 7.2 Production of the deficiency stocks in common wheat. The "A" represents the *Aegilops* chromosome causing chromosomal aberrations.

cylindrica chromosomes (Endo and Gill, 1996; Tsujimoto et al., 2001). The frequency of chromosome structural changes was far less in the self-progeny of disomic addition plants and in the F_1 monosomic alien addition line progeny derived from reciprocal crosses with common wheat. Nasuda, Friebe, and Gill (1998) observed that *Gc* gene-induced chromosomal breakage probably occurs prior to first pollen mitosis and during first and second pollen mitosis (Figure 7.3).

Kota and Dvořák (1988) found deficiencies, translocations, ring chromosomes, dicentric chromosomes, and paracentric inversions during the production of a substitution of chromosome $6B^{s}$ from *Ae. speltoides* for chromosome 6B of Chinese Spring wheat; 49 of the 138 plants contained chromosome aberrations. Chromosome rearrangements were recorded in wheat and *Ae. speltoides* chromosomes. The B-genome chromosomes showed the highest frequency of structural changes, followed by A-genome and D-genome chromosomes. The chromosome aberrations were nonrandom in the B-genome. Chromosomes 1B and 5B were the most frequently involved. Chromosome aberrations are induced prior to syngamy and are seldom transmitted through the gametophyte if the parent is used as a male. The other assumption is that the factor responsible for genomic instability is active only in the maternal germ line, different from that recorded by Endo (1988).

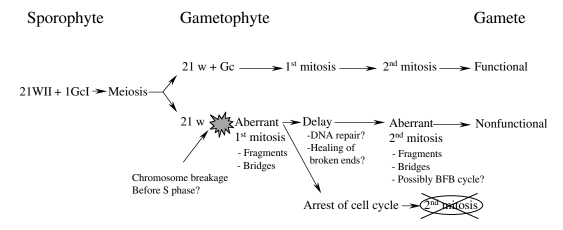


Figure 7.3 The mode of action of *Gc* genes in Chinese Spring wheat background, causing semisterility. (Redrawn from Nasuda, S., Friebe, B., and Gill, B.S., *Genetics*, 149, 1115–1124, 1998.)

Homoeologous	Genome \rightarrow	4	A	I	В	I	D	
Group	Arms →	S	L	S	L	S	L	Total
1		5	6	22	18	5	8	64
2		9	6	13	11	6	12	57
3		4	8	10	12	9	3	46
4		4	13	9	14	5	15	60
5		11	23	9	18	4	12	77
6		5	8	11	15	7	11	57
7		13	25	6	16	6	9	75
Total		140 ((32%)	184 ((42%)	112 ((26%)	436

Table 7.1 Frequency of Deficiencies in Chromosome Arms in Wheat Cultivar Chinese Spring

Source: From Endo, T.R. and Gill, B.S., J. Hered., 87, 295-307, 1996. With permission.

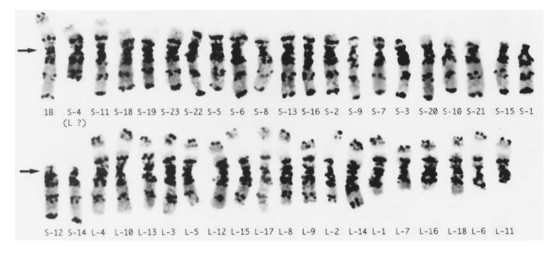


Figure 7.4 A series of deficiencies in Chromosome 1B (normal Chromosome 1B, extreme left) detected by Giemsa C-banding technique. The horizontal line (arrows) represents the kinetochore. (From Endo, T.R. and Gill, B.S., *J. Hered.*, 87, 295–307, 1996. With permission.)

The number of deficiencies predominates (42%) for B-genome chromosomes (Table 7.1). A series of deficiencies in Chromosome 1B is shown in Figure 7.4. Breakage occurs generally adjacent to the heterochromatic regions. The B-genome chromosomes are more vulnerable to breakage, because these chromosomes are more heterochromatic than A and D genome chromosomes. Deficiencies are being identified by Giemsa C- and N-banding and by *in situ* hybridization techniques or by GISH in wheat–rye and wheat–barley addition lines. The rate of deficiency was highest (42%) in chromosomes of B-genome followed by A-genome (32%) and D-genome(26%). The distribution of breakpoints correlated well with the relative size of the genomes (Table 7.1).

In another study, Tsujimoto et al. (2001) induced a total of 128 chromosomal aberrations (terminal deficiencies, 110; translocations, 7; dicentrics, 8; insertions, 1; highly chimeric, 2) in 1B chromosome of common wheat by Gc gene. These lines were produced by crossing a monosomic alien chromosome addition line of Chinese Spring carrying Chromosome 2C of *Ae. cylindrica* (abbreviated as CS + 2C) with nullisomic 1B-tetrasomic 1D.

Barley and rye are diploid (2n = 14) and unable to tolerate deficiencies. Production of deficiencies in wheat–barley and wheat–rye addition lines by *Gc* factor will enhance the gene mapping in barley and rye. By using the *Gc* gene from *Ae. cylindrica*, Shi and Endo (1999) produced chromosome aberrations for barley Chromosomes 2H, 3H, 4H, 5H, and 7H (Table 7.2). They identified a total of 31 deficiencies, 26 translocations, and two isochromosomes. Serizawa et al. (2001) produced seven deficiencies and 15 translocations in barley Chromosome 7H. The breakpoints of

	Selfed	progeny	Back-cros	k-cross progeny		
Barley Chromosome	Number of Plants Examined	Number of Plants with Aberrations (%)	Number of Plants Examined	Number of Plants with Aberrations (%)		
2H	15	1 (6.7)	22	3 (13.6)		
3H	35	2 (5.7)	60	9 (15.0)		
4H	48	16 (33.3)	20	3 (15.0)		
5H	7	1 (14.3)	18	3 (16.7)		
6H	43	8 (18.6)	51	7 (16.7)		
7H	28	6 (21.4)	156	26 (16.7)		
Total	176	34 (19.3)	327	51 (15.6)		

Table 7.2 Frequency of Plants with Aberrant Barley Chromosomes

Source: From Shi, F. and Endo, T.R., Genes Genet. Syst., 74, 49-54, 1999. With permission.

the deficiencies and translocations by N-banding, FISH, and GISH suggest that they are localized, in general, to the distal and proximal regions of barley chromosomes.

Friebe, Kynast, and Gill (2000) produced 56 deficiencies for rye chromosomes by Gc gene. They crossed seven disomic alien addition lines (DAALs) of CS wheat-"Imperial" rye (2n = 44;21W II + 1R II) as a female with DAALs of CS wheat — Ae. cylindrica (2n = 44; 21W II + 2C II) as male. The double monosomic alien addition line (2n = 44; 21W II + 1R I + 1C I) is expected to produce four types of gametes: 21W, 21W + R, 21W + 2C, and 21W + R + C. Because univalents in wheat are expected to be eliminated in about 3/4 of the gametes, the expected frequencies are in the proportion of 9/16, 3/16, 3/16, and 1/16, respectively. It has been established that a chromosome containing the Gc gene induces chromosome structural changes in gametes without a Gcchromosome. The target gametes for producing the deficiencies in rye chromosome, in this case, are 3/16, which is considerably low. To enhance the screening of higher frequencies of deficiencies, they back-crossed the double MAALs (2n = 21 W II + R I + 2C I) with the corresponding wheat-rye DAALs, and screened the BC_1 progenies cytologically to identify disomic plants for a given rye chromosome and monosomic for the gametocidal chromosome (2n = 45, 21 II + R II + 2C I). All gametes of these plants have the target rye chromosome, and 3/4 of the gametes are expected to be without 2C (21 W + R) and, thus, are not subjected to chromosome aberrations. These plants were back-crossed with the corresponding wheat-rye addition line or selfed. The derived lines were screened by C-banding for chromosomal structural changes in the rye chromosomes. By using this procedure, 33 deficiencies, 22 wheat-rye dicentrics, and seven wheat-rye and rye-rye translocations were recovered.

C. Meiotic Chromosome Pairing in Wheat Deficiency Line

The degree of meiotic chromosome association at metaphase-I between homologous chromosome arms of wheat is drastically influenced by a deficiency in one arm. Curtis, Lukaszewaski, and Chrzastek (1991) examined meiotic metaphase-I pairing in 4AL arm containing relative length of deficiency from 6 to 50%. The pairing frequency continued to decline as the deficiency length increased (Table 7.3). Pairing frequency in 4AL deficient for 34% reduced to 3% and was absent (0%) when deficiency reached 36%. The lack of pairing between the deficient arm and its normal homologue is probably related to an inability of homologues of unequal length to initiate pairing, and this may result in no recovery of recombinant chromosomes (Curtis, Lukaszewaski, and Chrzastek 1991). Hohmann et al. (1995) recorded a pairing reduction of 60% in the deficiency of the most distal 1% of chromosome arm 7AL.

Deficiencies in wheat, rye, and barley chromosomes induced by Gc gene mechanism are mostly terminal. Progeny of deficiency stocks of wheat breed true, suggesting that broken chromosomal

		Pairing Fi	requency (%)	of Deficient A	Arm with
	Relative Length	Complete H	omologue	Complete ⁻	Telosome
Chromosome	of Deficiency	S	L	S	L
Df4A06L	6	90.0	32.0	_	_
Df4A08L	8	84.0	64.0	85.2	29.8
Df4A11L	11	89.5	50.0	89.0	16.4
Df4A17L	17	70.6	11.8	76.1	5.4
Df4A23L	23	82.7	6.9	70.2	3.8
Df4A34L	34	83.0	3.0	68.6	0.0
Df4A36L	36	87.0	0.0	77.8	0.0
Df4A39L	39	_	_	41.7	0.0
Df4A50L	50	84.0	0.0	—	_

Table 7.3 Pairing Frequencies at Metaphase-I of Deficient Chromosomes 4AL with Complete Homologous and Telosome Chromosomes

S, short arm; L, long arm.

Source: From Curtis, C.A., Lukaszewaski, A.J., and Chrzastek, M., *Genome*, 34, 553–560, 1991. With permission.

ends heal after breakage by the synthesis of telomeric sequences or undergo fusions to produce dicentric or translocated chromosomes. Dicentric chromosomes undergoing breakage-fusion-bridge (BFB) cycles in the first few divisions of sporophyte are particularly healed before germ line differentiation, and ends are totally healed in the ensuing gametophytic stage (Friebe et al., 2001).

D. Transmission of Deficiencies

Deficiencies in maize were transmitted through eggs but not through pollen (McClintock, 1938a; Rhoades and Dempsey, 1973). The morphologically and cytologically detectable deficiencies were rarely transmitted to the next generation in tomato. Small cytologically undetectable deficiencies were transmitted through male and female normally. In the case of a deficiency of the entire heterochromatic 2S arm, only one individual of 1416 plants examined showed transmission of the deficiency. No tomato euchromatic deficiencies were known to be transmitted, unless they involved heterochromatin (Khush and Rick, 1967b). Even small deficiencies are lethal during gametophyte development in *Vicia faba* (Schubert and Reiger, 1990).

Of a total of 338 deficiencies, homozygous lines of common wheat (289 single, 39 double, seven triple, and three quadruple deficiency lines) are maintained as they were more or less fertile (B.S. Gill, personal communication, 2002). About 67% of the wheat deficiencies are homozygous, and they transmit normally (Endo and Gill, 1996). Some deficiency homozygotes are sterile or could not be obtained. These lines have to be maintained as heterozygotes, as they behave just like a monosomic chromosome, and the progeny must be screened cytologically for homozygotes. For example, all homozygous plants for 2AS deficiencies were highly male and female sterile, because all deficiencies were larger than that of 2AS-5. All 4BS deficiency homozygotes were totally male sterile. These stocks are being maintained, producing new disomic stocks for the 2AS or 2BS deficiency chromosomes and monosomic for the 2A or 4B short-arm telocentric chromosomes. They recovered deficiency homozygotes at a high rate in the progeny of these stocks. Most of the barley deficiencies in wheat are heterozygous and are transmitted to offspring like monosomic addition in wheat. Barley deficiencies and translocations in wheat–barley addition lines are differentiated by GISH (Shi and Endo, 1999).

E. Genetic Studies

Deficiencies have been utilized effectively to locate marker genes by the pseudodominant technique in maize (McClintock, 1941a), tomato (Rick and Khush, 1961; Khush and Rick, 1967b, 1968a)

and wheat (Gill et al., 1996; Sukta et al., 1999; Tsujimoto et al., 2001). Khush and Rick (1968a) located 35 marker genes belonging to 18 arms of the 12 tomato chromosomes by the induced-deficiency technique. In addition to gene location, induced deficiencies helped better the understanding of tomato genomes in ways briefly listed below:

- 1. Association of a linkage map with its respective chromosome: Rick and Khush (1961) associated the unlocated *a-hl* linkage group with Chromosome 11 by induced deficiencies.
- 2. *Identification of unmarked chromosome*: By induced deficiencies, Khush and Rick (1966) associated the marker *alb* with Chromosome 12.
- 3. *Location of a marker with a particular arm*: Numerous induced deficiencies were used to assign markers to particular arms of the tomato chromosomes.
- 4. Location of markers to a specific region of a chromosome: Several markers, such as *sy*, *ru*, *bls*, *sf*, *ra*, *var*, *ag*, and *t*^y, were located to a specific segment of a tomato chromosome by induced deficiencies.
- 5. Assignment of unlocated markers to respective chromosomes: Induced deficiencies helped to assign unlocated markers *clau*, *alb*, *lut*, and *fd* to their respective chromosomes.
- 6. *Orientation of the linkage maps*: Proper arm assignments of linkage maps of Chromosomes 2, 4, 6, 8, and 11 were made.
- 7. *Location of centromere positions*: Induced deficiencies facilitated the precise locations of the centromeres of nine chromosomes and the approximation of the other three chromosomes of tomato.
- 8. *Production of aneuploids*: Induced deficiencies were utilized to generate tertiary trisomics, secondary trisomics, and telotrisomics.

Deficiency stocks are an excellent tool for physical mapping of molecular and qualitative trait loci in wheat (Gill et al., 1996; Sutka et al., 1999; Tsujimoto et al., 2001). Gill et al. (1996) developed physical maps by locating 80 DNA and two morphological markers using 65 deficiency lines for homoeologous Group 5 chromosomes. The maps were constructed for Chromosomes 5B in wheat and 5D in *Aegilops tauschii*. They recorded suppression of recombination in the centromeric regions and found it was prominent in the gene-rich regions.

F. Use of Deficiencies in Hybrid Maize Breeding

Patterson (1973) proposed a system for producing hybrid maize by utilizing nuclear male sterility genes (*ms*) in conjunction with chromosome deficiencies in the background of a normal (nonsterile) cytoplasm. Male sterile stocks are normally maintained by *ms/ms* (female) \times *ms/*+ (male) crosses. The progeny are expected to segregate 1/2 male sterile (*ms/ms*):1/2 male fertile (*ms/*+) (Figure 7.5A). On the other hand, all plants from this test cross are expected to be sterile if pollen grains carrying the + allele are nonfunctional in fertilization. Detasseling could be greatly reduced or eliminated in female rows in hybrid production if this result could be approximated. As shown in Figure 7.5B, this differential transmission of *ms* locus alleles might be achieved by a male fertile allele carried immediately proximal to a terminal deficiency that is female transmissible, but not male transmitted. Differential transmission could equally well be achieved by a male fertile allele carried immediately adjacent to an internal deficiency showing the same transmission characteristics. In maize, however, few simple deficiencies showing the required transmission characteristics have been recognized and saved.

Patterson (1973) demonstrated that the required level of differential transmission of *ms* alleles could be accomplished by utilizing female-transmissible duplicate-deficient (Dp-Df) chromosome complements derived by Adjacent 1 disjunction from some heterozygous reciprocal translocations. In suitable Dp-Df complements, the deficient segment is closely linked in the coupling phase to the + allele of an *ms* gene in *ms/ms* (female) by Dp-Df +/*ms* or Dp-Df + *ms/ms* (male) crosses. As shown in Figure 7.6, there are two options with the same chromosome structures. In Figure 7.6 (upper) is shown the wild-type allele (+) of a Chromosome 6 male sterile gene located proximal

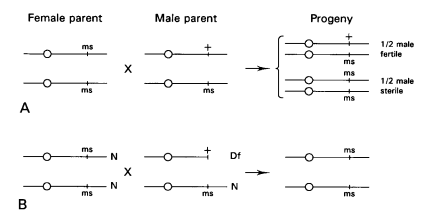


Figure 7.5 Use of chromosome deficiencies to produce male sterile progeny in maize. (A) Expected segregation in male sterile (female) × male fertile (male); (B) a proposed method to produce male sterile progeny in maize. (Redrawn from Patterson, E.B., *Proc. 7th Meeting Maize and Sorghum Sect.*, 1973.)

to the interchange point in the 6⁹ chromosome; it is thus located just proximal to a terminal deficiency for the tip of the long arm of Chromosome 6. In Figure 7.6 (lower) is shown the alternative option of using the wild-type allele of a Chromosome 9 male sterile gene located distal to the interchange point on the 6⁹ chromosome; in this position, the locus is part of a Chromosome 9 segment that becomes triplicated in the derived Dp-Df plant. Suitably marked Dp-Df plants will have received the recessive *ms* allele from the male parent. The interchange point on the 6⁹ chromosome 6. A locus on either side of that interchange point is thus linked genetically and physically to the same terminal deficiency. Because the 6⁹ chromosome cannot be male transmitted in the absence of the 9⁶ chromosome, a wild-type allele of an *ms* gene carried on it cannot be transmitted to progeny, except if it is transferred to a normally arranged homologue as a result of crossing over between the locus and the adjacent interchange point.

In commercial use, Dp-Df stocks would be derived in inbred lines destined to be used as female parents, or components of female parents, in production of hybrid maize. In initial crosses,

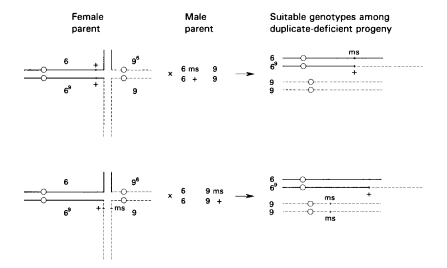


Figure 7.6 The derivation of suitably marked duplicate-deficient progeny from a reciprocal translocation in maize. (Redrawn from Patterson, E.B., *Proc. 7th Meeting Maize and Sorghum Sect.*, 1973.)

suitably marked Dp-Df plants like those shown in Figure 7.6, are self-pollinated to propagate plants of the parental Dp-Df constitution and are crossed as male parents to male sterile plants of the same line. The latter cross confirms the constitution of the Dp-Df male parent and, at the same time, produces seeds that will yield virtually all male sterile plants for use in female rows of foundation or production fields. In routine foundation field practice, male fertile Dp-Df plant produced by self or sib open pollination would serve as pollinators in male rows. Female rows would consist of male sterile plants; seed produced on these plants is suitable, in turn, for planting additional male sterile rows in foundation or production fields. Male rows in production fields would consist of standard inbred line, because such lines normally carry wild-type alleles of recessive nuclear male sterile genes and thus function as natural restorers of male fertility in hybrid seed sold to farmers.

Four different Dp-Df complements were identified (Patterson, 1973) that when used in suitable combinations with three different male sterile genes in test crosses each gave progenies in which only one or two plants per 1000 were male fertile. Commercial procedures require that Dp-Df plants be used as pollinators in foundation fields. Thus far, Dp-Df complements showing strong linkage with suitable *ms* loci have not met maize industry requirements for agronomic performance with respect to pollen and seed production.

G. Ring Chromosomes

Ring chromosomes have been found in several plant species, including Zea mays, Nicotiana tabacum, Antirrhinum majus, Petunia hybrida, and Hordeum vulgare, as well as in Drosophila and man. Cytological behavior of ring chromosomes is unique. McClintock (1931, 1938b, 1941a, 1941b) studied thoroughly the mitotic and meiotic behaviors of ring chromosomes in maize and developed the breakage-fusion-bridge (BFB) cycle hypothesis to explain changes in chromosome sizes (duplications and deficiencies). She also followed these chromosome aberrations in genetic studies (Figure 7.7). The breakage-fusion-bridge cycle phenomenon has been confirmed by Morgan (1933) and Braver and Blount (1950) in Drosophila, by Stino (1940) in tobacco, Schwartz (1953a, 1953b) and Fabergé (1958) in maize, Michaelis (1959) in Antirrhinum majus, Tsunewaki (1959) in wheat-Agropyron hybrids, Frost, Lesley, and Locke (1959) in Matthiola incana, Maizonnier and Cornu (1979) in Petunia hybrida, and Singh and Tsuchiya (1981e) in barley. Most of the ring chromosomes studied were in the disomic condition, in which a ring chromosome compensated for a deficiency in the standard homologues.

Ring chromosomes may occur spontaneously due to meiotic irregularities, or they may be induced by x-ray or gamma ray treatment. Only ring chromosomes are produced from the ring chromosomes. According to McClintock (1941a), ring chromosomes originate from a univalent chromosome by nonhomologous chromosome synapsis of two arms followed by crossing over. In barley, it is believed that the ring chromosomes obtained in the progenies of Triplo 7 (semi-erect) may have originated in a similar fashion (Singh and Tsuchiya, 1981e).

Various sizes of ring chromosomes have been observed in disomic wheat-rye addition lines by BFB cycle (Lukaszewaski, 1995). However, this study could not discover a mechanism like transposable element in wheat as was discovered in maize.

Ring chromosomes are not stable during cell division and often are eliminated during cell divisions. Their numbers and sizes are changed in successive cell cycles depending upon the position of the breakage in dicentric double-sized ring chromosomes (Figure 7.7). During mitosis, small ring chromosomes exhibit the following features:

- 1. Reduced frequency with which double-sized or interlocking rings originate
- 2. Frequent loss of ring chromosomes from the nuclei
- 3. Less frequent changes in size of the ring chromosomes
- 4. Increased number of rings

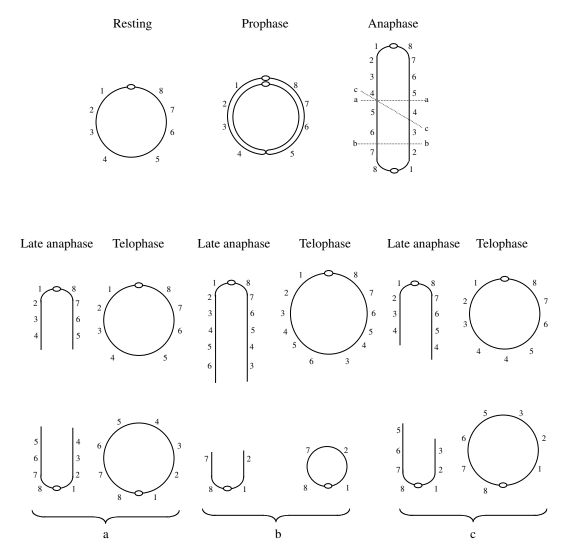


Figure 7.7 Origin of ring chromosomes of various sizes by breakage-fusion-bridge cycle. Breakage points (a — a; b — b; c — c), shown at anaphase-I in a dicentric bridge produce ring chromosomes of various sizes. (Redrawn from McClintock, B., *Genetics*, 26, 542–571, 1941a.)

McClintock (1938b) utilized a ring chromosome to locate a marker gene in a particular region of Chromosome 5. The ring chromosome carried the locus Bm1 (allele bm1 shows brown midrib when homozygous or hemizygous). The rod chromosomes were of three types: both copies of chromosomes lacked the Bm1 locus, one copy was deficient for the locus, while the other carried bm1, or both carried bm1. The plants expressed variegation for Bm1 and bm1 expression through the frequent loss of the ring chromosomes from somatic tissues.

II. DUPLICATIONS

A. Introduction

An extra piece of chromosome segment attached to the same homologous chromosome or transposed to one of the nonhomologous members of the genome is termed a duplication. Based on

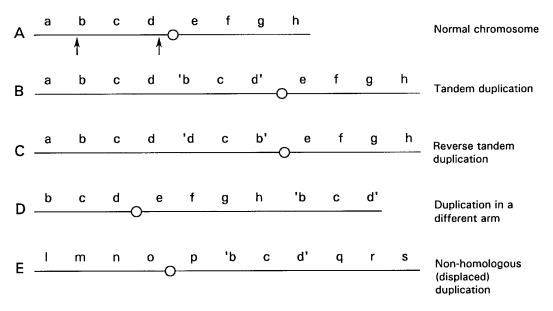


Figure 7.8 Types of duplications are shown in parts A to E. Arrows show a segment of chromosome duplicated in several fashions.

transposition of the segment, Burnham (1962) classified duplications shown in Figure 7.8 as follows: (1) tandem (Figure 7.8B); (2) reverse tandem (Figure 7.8C); (3) duplication in a different arm (Figure 7.8D); and (4) displaced nonhomologous (Figure 7.8E). Duplications are more frequent and less lethal to organisms than are deficiencies. Combined duplications and deficiencies, abbreviated as (Dp-Df) for duplicate and deficient, produce changes in the amount of genetic material in spores, pollen, gametes, seeds, and plants.

B. Origin of Duplications and Deficiencies

- 1. Duplications occur in nature and are also produced experimentally. In several diploid crops, duplicate genes, determined by the F₂ segregation (15:1 or 9:7), have been recorded. Furthermore, the formation of occasional rod- or ring-shaped bivalents at meiosis in haploid suggests the presence of duplications (Burnham, 1962).
- 2. Generally, ionizing types of radiation break chromosomes, producing duplications and deficiencies.
- Viable Dp-Dfs are isolated in the progenies of chromosomal interchanges in maize (Gopinath and Burnham, 1956; Phillips, Burnham, and Patterson, 1971; Patterson, 1978; Carlson, 1983), barley (Hagberg, 1962), and cotton (Menzel and Brown, 1952, 1978; Menzel and Dougherty, 1987; Contolini and Menzel, 1987).
- 4. Dp-Dfs are produced from breakage of dicentric bridges at meiotic anaphase-I and anaphase-II, and spore mitotic anaphase (Figure 7.9) in the progenies of inversion heterozygotes (McClintock, 1938a; Rhoades and Dempsey, 1953). The observed pseudo-alleles near the end of the short arm of Chromosome 9 in maize suggested to McClintock (1941c) the presence of a series of similar genes that arose through duplications. A similar situation can originate with regard to duplications from breakage-fusion-bridge cycles.

C. Identification of Duplications and Deficiencies

In favorable species, Dp-Df plants may be identified cytologically at pachynema by an unpaired segment that is present in single copy and by pairing patterns displayed by a segment present in triplication. In maize, Dp-Df plants may be recognized by pollen phenotypes, because they produce equal numbers of usually distinguishable normal and Dp-Df pollen grains.

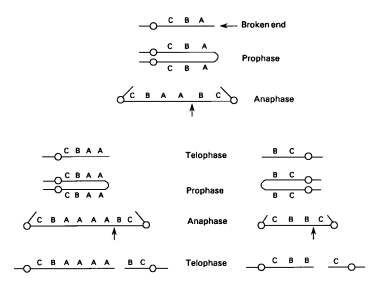


Figure 7.9 Production of Dp-Dfs from breakage-fusion-bridge cycle. Breakage points are shown by arrows. (Redrawn from McClintock, B., *Genetics*, 26, 234–282, 1941c.)

D. Transmission of Duplications and Deficiencies

The frequency of Dp-Df individuals in progenies depends on frequencies of Dp-Df megaspores and viability of Dp-Df megaspores, megagametophytes, gametes, and zygotes. In maize, Dp-Df kernels borne on Dp-Df plants often comprise less than one-third of the total kernels (Patterson, 1978). Dp-Df complements are transmitted almost exclusively through the female in maize. The frequencies are variable, depending especially upon the lengths of the deficient and duplicated segments. There was no transmission of Dp-Df chromosomes through the pollen in material studied by Rhoades and Dempsey (1953). However, Patterson (personal communication, 1992) has evidence that at least three Dp-Df complements derived from reciprocal translocations in maize may be male — transmitted even under conditions of pollen competition. The usual lack of pollen transmission may be attributed to pollen inviability or the inability of Dp-Df pollen to compete with chromosomally balanced pollen in effecting fertilization. By utilizing B-A translocations, Carlson (1986) and Carlson and Roseman (1991) produced heritable segmental (proximal and distal) duplications in maize. In cotton, Dp-Dfs obtained from adjacent-1 disjunction of heterozygous translocation tetravalents were usually ovule viable and, occasionally, pollen viable. Adjacent-2 Dp-Dfs were rarely recovered (Menzel and Dougherty, 1987).

E. Use of Duplications and Deficiencies in Genetic Studies

Duplications can modify genetic ratios because of inviable gametes or alter rates of transmission, and they may change phenotypes due to dosage effects of particular alleles. The Dp-Df stocks in maize are useful in localizing and sequencing genes in chromosomes. Distortion in genetic segregation ratios of linked genes may be used to determine the positions of translocation points with respect to gene loci and the sequence of gene loci with respect to each other (Patterson, 1978). In the simplest application, translocation heterozygotes homozygous for a counterpart dominant allele are used as females and are pollinated by plants homozygous for (or carrying) a recessive mutant allele. In this initial extraction of a female-transmissible Dp-Df complement arising from adjacent-1 disjunction, Dp-Df plants in maize may often be identified by pollen phenotype (Figure 7.10). If the gene is located in the Df segment, a recessive allele will be in a hemizygous condition and will be expressed. If the gene is not located in the Df segment, Dp-Df plants carrying the recessive

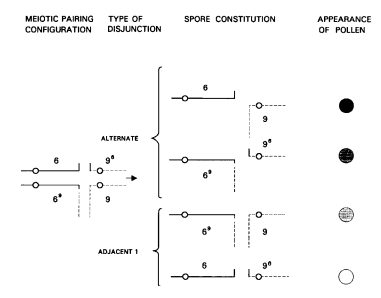


Figure 7.10 Chromosome configurations and spore constitutions after meiosis in a maize plant heterozygous for a reciprocal translocation. Appearance of pollen grains with Dp-Df chromosome constitutions reflects an unbalance of spores. (Redrawn from Patterson, E.B., *Maize Breeding and Genetics*, John Wiley & Sons, New York, 1978, 693–670.)

allele are crossed as males to a tester stock homozygous for the mutant allele. Recombination frequency of the mutant locus with the interchange point is measured, and this recombination frequency can be compared with the independently or simultaneously measured recombination frequencies of various mapped genes in the same chromosome with the same interchange point in Dp-Df plants. If the mutant locus is in fact linked to the deficiency, this procedure will furnish an approximate positioning of the new mutant locus in the chromosome map. Further elaborations of Dp-Df mapping techniques were also discussed (Patterson, 1978).

Modern techniques that sequence RFLPs with respect to each other and to marker genes have found various unbalanced chromosome complements useful in these studies. Particularly useful have been Dp-Df complements derived from heterozygous reciprocal translocations and inversions, simple monosomics, tertiary trisomics, and heterozygous deficiencies generated from B–A translocations (Carlson and Curtis, 1986).

III. INTERCHANGES

A. Introduction

Interchanges are known as segmental chromosomal interchanges, reciprocal translocations, or simply translocations (Burnham, 1956; Ramage, 1971). Translocations are the result of the reciprocal exchange of terminal segments of nonhomologous chromosomes (Figure 7.11).

Translocations were first observed by Gates (1908) in *Oenothera rubrinervis*, where more than two chromosomes were attached to form a ring. Subsequently, Cleland (1922) suggested in studies based on *Oenothera* species that chromosomes in a ring are associated in a specific and constant order. Belling and Blakeslee (1926) put forth an hypothesis of segmental interchange from chromosome pairing studies in *Oenothera* and suggested that the formation of a circle is due to reciprocal translocation. However, McClintock (1930) was the first to provide cytological evidence of interchanges between two nonhomologous chromosomes in maize.

Normal chromosomes

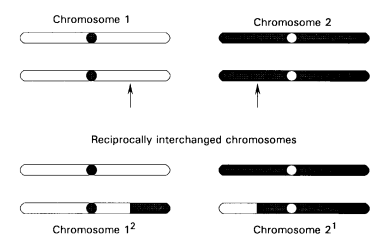


Figure 7.11 Origin of interchanged (reciprocal translocation) chromosomes (arrows show the breakage points).

B. Identification of Interchanges

An interchange heterozygote may be identified cytologically, by its effects on partial pollen and seed sterility, or by genetic tests. A reciprocal translocation is obtained when each member of two nonhomologous chromosomes is broken, as shown in Figure 7.11, and the two terminal segments exchange reciprocally, producing the interchanged chromosomes designated as chromosomes 1^2 and 2^1 . Thus, an organism carrying chromosomes 1, $1,^2 2,^1 2$ is known as an interchanged heterozygote or a translocation heterozygote. Interchanged heterozygote chromosomes form a cross-shaped configuration at pachynema of meiosis (Figures 7.12A, 7.13, 7.14). A group of four chromosomes, known as a quadrivalent or tetravalent, coorient at diakinesis and metaphase-I in such a fashion that several types of configurations are observed, depending upon the positions of kinetochores (Figure 7.12B).

In coorientation configuration, the chromosomes are distributed in equal numbers to the opposite poles. In alternate (zigzag) and Adjacent 1 (open) segregations, homologous kinetochores move to opposite poles (Figure 7.12B). In constrast, in Adjacent 2 (open) segregation, homologous kinetechores move the same pole (Figure 7.12B).

The region between an interchange point and a kinetochore is termed an interstitial region. If there is no crossing over in an interstitial region, alternate chromosome migration results in viable male and female gametes: 1 + 2 standard normal or $1^2 + 2^1$ translocated, but balanced, chromosomes. Meiotic products from adjacent disjunctions are typically inviable due to duplications and deficiencies. However, crossing over in the interstitial segments will transfer blocks of genes between the interchanged and normal chromosomes, altering the types of products from alternate and Adjacent 1 disjunctions, but all of the products will be abortive from Adjacent 2 disjunction (Figure 7.13; Table 7.4).

When an interchange heterozygote orients on the meiotic metaphase plate, an approximate ratio of 1 alternate: 1 adjacent arrangement of chromosomes is frequently observed because of the random behavior of two pairs of cooriented kinetochores (Endrizzi, 1974; Chochran, 1983; Rickards, 1983). In alternate orientations, the two standard chromosomes or the two interchanged chromosomes may orient to a given pole. The two configurations may be distinguished by chromosome morphological traits (Endrizzi, 1974; Lacadena and Candela, 1977).

Chromosome configurations in interchange heterozygotes can be identified in cotton because of differences in chromosome lengths of A and D genomes (Endrizzi, 1974).

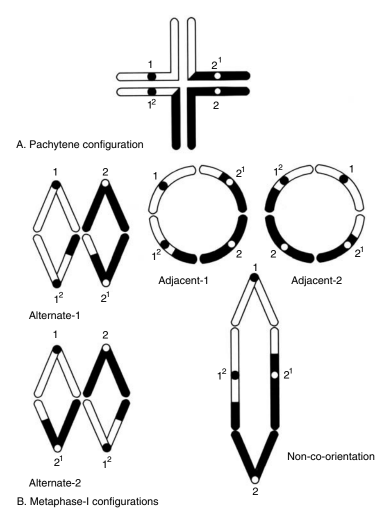


Figure 7.12 Chromosome configurations in a reciprocal translocation heterozygote. (A) A cross-shaped configuration at pachynema. (B) Possible chromosome configurations at metaphase-I.

The deficiency in the Adjacent 2 class in a T4–5 interchange is most likely due to the knob or may be attributed to nonterminalized chiasmata that may have caused a higher frequency of coorientation of homologous kinetochores to opposite poles (Table 7.5).

In noncooriented arrangement, normal chromosomes carrying nonhomologous kinetochores are on opposite sides of a quadrivalent located equidistant from the equatorial plate. The two noncooriented interchanged chromosomes are stretched in the middle and are not attached to the poles (Figure 7.12B). At anaphase-I, one cooriented and two noncooriented chromosomes move to the one pole, while the other cooriented normal chromosome moves to the opposite pole, producing 3–1 chromosome segregation. This type of segregation generates tertiary trisomics and monosomics in the progenies. Furthermore, chromosome segregation of 2:2 results in unbalanced gametes (deficiency–duplication).

In diploid plant species, heterozygous interchanges generally exhibit approximately 50% pollen sterility. In the *Glycine max* \times *G. soja* cross, studied by Singh and Hymowitz (1988), pollen fertility ranged from 49.2 to 53.3% (Table 7.6). This result suggests that alternate (zigzag) and adjacent (open) configurations are in nearly equal proportions. Other plant species with 50% sterility in translocation heterozygotes have been listed by Burnham (1962). Interchanged heterozygotes usu-

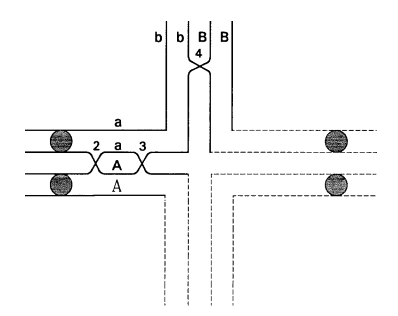


Figure 7.13 A reciprocal translocation heterozygote at pachynema showing crossing overs between kinetochore and marker gene *a* (2), between marker gene *a* and breakage point (interstitial region) (3), and between breakage point and marker gene *b* (distal region) (4).

Table 7.4	Nature of Crossing Over, and Types of Gamete Formations in Three Types of Chromosome
	Coorientation

Nature of Crossing Over	Alternates 1 and 2	Types of Orientation Adjacent 1	Adjacent 2
1. No crossing over	Original combinations are viable	All gametes are nonviable	All gametes are nonviable
2. Crossing over between centromere and marker gene <i>a</i>	Original combinations are viable	Only cytological crossover products are viable	All gametes are nonviable
3. Crossing over within interstitial segment; between marker gene <i>a</i> and breakage point	Original combinations are viable	Only cytological and genetical crossover products are viable	All gametes are nonviable
4. Crossing over in the distal region; between breakage point and marker gene <i>b</i>	All combinations are viable	All combinations are nonviable	All gametes are nonviable

Table 7.5	Frequency of Chromosome Configurations in the
	Interchange Heterozygotes of Gossypium hirsutum

Translocation	Types of Chromosome Orientation								
Stocks	Adj. 1ª	Adj. 2ª	Alt. 1 ^b	Alt. 2 ^b					
T4–5	48	16	45	37					
T10–19	68	31	60	30					
AG184	20	6	20	13					

^a Adj. = adjacent.

^b Alt. = alternate.

Source: From Endrizzi, J.E., Genetics, 77, 55–60, 1974. With permission.

Hybrids	2 <i>n</i>	Ш	IV	Total PMC	Pollen Fertility (%)
Bonus ^a × PI 81762 ^b	40	18.3	0.9	38	49.2
PI 81762 \times Bonus	40	18.2	0.9	25	52.0
Essex ^a × PI 81762	40	18.2	0.9	48	53.3
PI 81762 \times Essex	40	18.4	0.8	25	51.2

Table 7.6Meiotic Chromosome Configuration at Diakinesis and Pollen Fertility
(%) in *Glycine max* × *G. soja* and in Its Reciprocal Crosses

^a Glycine max.

^b G. soja.

Source: From Singh R.J. and Hymowitz, T., *Theor. Appl. Genet.*, 76, 705–711, 1988. With permission.

ally do not show 50% pollen and seed sterility in polyploids, even though parents sometimes differ by two interchanges. Tetraploid cotton (Brown, 1980), hexaploid wheat (Baker and McIntosh, 1966), and hexaploid oat (Singh and Kolb, 1991) have been extensively studied. In hexaploid oat, pollen fertility in intercultivar F_1 hybrids ranged from 91.6 to 99.7%, and in parents, it ranged from 94.1 to 95.5% (Table 7.7). These results suggest that male and female spores containing Dp-Df in polyploids are as competitive as normal spores.

C. Robertsonian Translocations

The kinetochore is an important part in the chromosome and is responsible for a remarkable and accurate movement of meiotic prometaphase homologous chromosomes to metaphase plate. It participates in spindle checkpoint control and moves chromosomes poleward at anaphase (Yu and Dawe, 2000). Meiotically unaligned (unpaired-univalents; nonhomologous) univalents remain laggard at anaphase-I and telophase-I and often misdivide at the kinetochore. Centric misdivision followed by the fusion of broken arms from different chromosomes produces Robertsonian translocation (Robertson, 1916). Robertsonian translocation is found in plants, animals, and humans and plays a key role in the karyotype evolution (see Zhang et al., 2001). Two different fused kinetochores can be distinguished by FISH.

			%						
	Frequen	cy of Chr	omosome	Associati	onsª at D	iakinesis	Total		
F₁ Hybrids	2111	1IV + 19II	1111 + 1911 + 21	1IV + 18II + 2I	2IV + 17II	1VI + 18II	Pollen Mother Cells	1IV + 19II	Pollen Fertility
Andrew/Gopher	10	22	_	_	_	_	32	68.8	99.7
Andrew/Hazel	3	40	1	1	7	_	52	76.9	92.0
Andrew/PA12422	4	16	1	_	2	_	23	69.6	_
Andrew/Otee	15	5	_	_	_	_	20	25.0	99.7
Hazel/Gopher	72	69	1	_	_	_	142	48.6	_
PA12422/Gopher	88	25	_	_	_	_	113	22.1	_
Hazel/Otee	87	40	1	_	_	_	128	31.3	_
Otee/PA12422	74	14	_	_	_	_	88	15.9	_
Hazel/Ogle	25	_	_	_	_	_	25	0.0	94.7
Hazel/PA12422	56	46	_	_	_	11	113	40.7	91.6

Table 7.7 Meiotic Chromosome Configurations at Diakinesis in Intercultivar Hybrids of Hexaploid Oat

^a I = univalent; II = bivalent; III = trivalent; IV = quadrivalent; VI = hexavalent.

Source: From Singh, R.J. and Kolb, F.L., Crop Sci., 31, 726-729, 1991. With permission.

D. Interchanges in Genetic Studies

Interchange stocks have been used to identify linkage groups, to associate new mutants to specific chromosomes, and to construct new karyotypes. This technique may be superior to the gene-marker stock method, because an interchange, in contrast to gene markers, usually does not affect the expression of other traits. Semisterility is often more clearly expressed than genetic markers, and two chromosomes may be tested for linkage instead of only one (Lamm and Miravalle, 1959).

The interchange-gene-linkage analysis is based on the association of contrasting traits with partial sterility. An interchange behaves as a dominant marker for partial sterility located simultaneously in the two interchanged chromosomes at the points where the original breakage and exchange occurred. For example, translocated chromosomes are designated *T*, and their normal chromosome homologues are symbolized as *N*. An individual heterozygous for a translocation (*T/N*) and for a gene pair (*Aa*) produces four kinds of gametes *AT*, *AN*, *aT*, and *aN*. In an F_2 generation, plants that are *T/T* (translocation homozygote) or *N/N* (normal homozygote) genotypes are fertile (F), while plants with *T/N* (translocation heterozygote) genotype are partial sterile (PS). Thus, four classes in progenies are expected in F_2 :

- A PS (T/N) = Normal, partial sterile, translocation heterozygote (six)
- A F (T/T N/N) = Normal, fertile, translocation homozygote or normal homozygote (six)
- a PS (T/N) = Mutant, partial sterile, translocation heterozygote (two)
- a F (*T/T*, *N/N*) = Mutant, fertile, translocation homozygote or normal homozygote (two)

Based on the above information, Tuleen (1971) used translocation tester stocks of barley carrying unequal interchanged chromosome pieces that could be readily identified at somatic metaphase. By using qualitative seedling mutants, four phenotypic classes can be recognized in F_2 by identifying *N*/*N* and *T*/*T* individuals. A linkage can be detected by studying the recessive fraction of F_2 population; plants with *T*/*Naa* genotype are generated from the union of a nonrecombinant gamete (*aN*) and a recombinant gamete (*aT*), while *aaT*/*T* and *aa N*/*N* plants are generated from the union of two recombinant or two nonrecombinant gametes, respectively. In F_2 , a deviation from 1 (*N*/*N*):2 (*T*/*N*):1(*T*/*T*) in recessive homozygotes can be used to determine the linkage. The mutants glossy seedling and virido-xantha are located on chromosome 4 of barley, because both mutants showed linkage with T4–5e and independent segregation with T1–5a. Furthermore, both mutants are close to a breakage point, because no recombinant (*T*/*N*) genotype was recovered (Table 7.8).

This technique is efficient and quite useful, because only a small sample of recessive F_2 seedlings, classified for the translocation by root tip squashes, is needed to detect linkage. However, a limitation of this technique is that it is time-consuming, because classification for translocation by the root tip method is tedious, applicable mainly to plants with large identifiable chromosomes, and requires expertise in cytological technique.

Interchanges have been useful for testing the independence of linkage groups established genetically. This was demonstrated by translocation analysis in barley. Kramer, Veyl, and Hanson (1954) observed that chromosome 1 of barley carried two linkage groups (III and VII). They used

		. <u>.</u>		3				
		T4–5e						
Mutant	NN	NT	TT	X ²	NN	NT	TT	X ²
Glossy seedling	7	0	0	14.0ª	1	5	1	0.0
Virido-xantha	10	0	0	20.0ª	2	4	1	0.3

Table 7.8 Interchange-Gene Linkages for T4–5e and T1–5a, F₂ Recessive Seedlings in Barley Classified Cytologically for the Interchanges

^a Significant at the 1% level.

Source: From Tuleen, N.A., Barley Genetics II. Proc. 2nd. Int. Barley Genet. Symp., 1971, 208–212. With permission.

eight translocation lines involving six chromosomes (a-b, b-d, c-d, a-e, c-e, c-b, b-f, e-f) and crossed with marker genes representing seven linkage groups of barley. Three interchange testers showed that linkage groups III and VII are not independent. With this information, Haus (1958) studied gene a_{c2} (white seedling) of linkage group III and y_c (virescent seedling) of linkage group VII of barley. The measured recombination value between these two genes was 28.14 ± 1.59% from combined F_2 and F_3 data, confirming that these two genes, a_{c2} and y_c belong to the same linkage group rather than to two different linkage groups.

E. Principles of Producing Interchange Testers

Complete interchange tester sets have been established and are being utilized in cytogenetic studies in several plant species, such as barley (Burnham, White, and Livers, 1954), garden pea (Lamm and Miravalle, 1959), maize (Burnham, 1954), tomato (Gill et al., 1980), rye (Sybenga and Wolters, 1972; Sybenga et al., 1985; Sybenga, 1996), cotton (Ray and Endrizzi, 1982), and pearl millet (Minocha et al., 1982).

Chromosomal interchanges occur spontaneously and may also be induced by irradiation (x-ray, gamma ray, fast neutron, thermal neutron, ultraviolet light) and chemical mutagens (ethyleneimine, ethyl methane sulfonate). Translocations have also been reported in the progenies of maize carrying the sticky gene (*st*) (Beadle, 1937) as well as activator and dissociation controlling elements (McClintock, 1950). They have also been found in aged seeds of barley and wheat (Gunthardt et al., 1953).

The chromosomes involved in different translocations may be identified by studying the F_1 s generated by intercrossing them with translocations involving known chromosomes, then observing the F_1 s at diakinesis or metaphase-I. The three possibilities are as follows:

- The occurrence of two quadrivalents (IV) suggests that both translocation testers involve different chromosomes.
- The presence of a hexavalent (VI) in F₁ sporocytes indicates that one of the chromosomes involved in the two translocation stocks is common.
- When both translocation stocks involve the same chromosomes, F₁ hybrids may show bivalents only.

F. Identification of Interchanged Chromosomes

Unidentified interchanged stocks can be identified cytologically by studying chromosome configurations at pachynema (Figure 7.14), diakinesis, and metaphase-I in an F_1 hybrid involving known translocation testers and aneuploid stocks. For example, reciprocal translocations in various wheat cultivars have been recorded in intervarietal F_1 hybrids, and the chromosomes involved in interchanges are identified using aneuploids of wheat cv. Chinese Spring. Because Chinese Spring wheat is considered the most primitive cultivar with standard chromosomes (Sears, 1953b), it is hybridized with other wheat cultivars. Chromosome pairing for chromosomal structural changes at meiotic metaphase-I is analyzed in F_1 plants. The observation of 1IV + 19II suggests that the parental cultivars differ from Chinese Spring by one reciprocal translocation. Similarly, the chromosome association of 2IV + 17II suggests that two independent interchanges are involved. In cases where two interchanges are observed, the frequency of cells with 1IV is higher than cells with 2IVs. This suggests that interchanged segments are different (Vega and Lacadena, 1982). Furthermore, the higher the frequency of quadrivalents, the larger the chromosome segments involved in translocation. Similar results were observed in intervarietal F_1 hybrids of oat (Singh and Kolb, 1991).

Singh and Kolb (1991) observed three types of multivalent associations in all intercultivar F_1 hybrid combinations among six parental lines (Andrew, Gopher, Hazel, Ogle, Otee, and a breeding line PA 12422) of oat (2n = 6x = 42): 1IV + 19II (one interchange), 2 IV + 17II (two independent interchanges), and 1VI + 18II (progressive or successive interchanges involving three chromosomes). The telomeres of chromosomes were designated arbitrarily as follows: 1.2, 3.4, 5.6, 7.8, 9.10 ... 41.42.

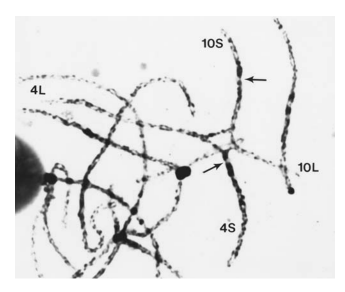


Figure 7.14 A cell at pachynema showing a cross-shaped configuration in an heterozygous reciprocal translocation in maize. Chromosomes involved in the interchange are numbered, and kinetochores are shown by an arrow. (Courtesy of D.F. Weber.)

Andrew and Gopher are differentiated by a single interchange, because 68.8% of the sporocytes showed a 1IV + 19II chromosome configuration (Table 7.7). Suppose a reciprocal translocation occurred between the chromosomes of Andrew designated 5.6 and 7.8, the resultant chromosomes designated 5.7 and 6.8 would correspond to the chromosome structures of Gopher; however, Andrew and Gopher are not related.

Based on results presented in Table 7.7, it is evident that Gopher and Otee have a similar chromosome arrangement, i.e., 1.2, 3.4, 5.7, 6.8, 9.10 ... 41.42. Also, Hazel and Ogle cultivars carry a similar chromosome constitution. Because Hazel differs from Gopher and Otee by a single interchange and from Andrew by two independent interchanges, a reciprocal translocation occurred between chromosomes 1.2 and 3.4 that resulted in 1.3, 2.4. 5.7, 6.8, 9.10 ... 41.42 chromosome ends for Hazel and Ogle. It should be mentioned that the frequency of sporocytes with the 1IV + 19II configuration in the Andrew/Hazel hybrid was higher (76.9%) compared with the sporocytes (13.4%) with 2IV + 17II (Table 7.7). This suggests that the chromosome segments involved in the second interchange may be small. The recorded chain quadrivalent configurations in the Andrew/Hazel and Hazel/Gopher F_1 hybrids also suggest that the second interchange (1.3, 2.4) is a small segment.

The F₁ hybrids between Hazel/PA 12422 showed 1VI + 18II, together with 1IV + 19II and 21II configurations. If it is assumed that a reciprocal translocation occurred between chromosomes 3.4 and 9.10 of Otee, then PA 12422 has 3.9 and 4.10 interchanged chromosomes, giving 1.2, 3.9, 4.10, 5.7, 6.8 ... 41.42 chromosome ends in PA 12422. Thus, the chromosome arrangement in a hexavalent configuration observed in Hazel/PA 12422 cross can be 1.2-2.4-4.10-10-9-9.3-3.1. Furthermore, when any four of the six chromosomes (e.g., 1.2-2.4-4.10-10.9) associate in a quadrivalent configuration, the configuration should be an open chain, because the two ends of the quadrivalent will be nonhomologous with each other.

The low frequency of 1IV + 19II configuration in PA 12422/Gopher (22.1%) and Otee/PA 12422 (15.9%) hybrids suggests that the interchanged segments (3.9 and 4.10) of PA 12422 are short. The small homologous segments are not large enough to form chiasmata with homologous counterparts, or by the time the chromosomes reach diakinesis and metaphase-I, the chiasmata are already terminalized, producing an open ring, a chain quadrivalent, or two bivalents.

Once a cultivar is recognized for a reciprocal translocation, attempts are made to identify the interchanged chromosome with the help of monosomics or monotelodisomics. Generally,

Unidentified Translocation Stocks	Translocation Lines	Primary trisomics	Telotrisomics	Monosomics	Mono	telodisomics
Tra. stock A	VI + II	V + II	V ^t + II	+	IV ^t + II	Associated
Tra. stock B	2IV + II	IV + III + II	$IV + III^t + II$	IV + I + II	$1IV + 1^{t}II$	Independent

Table 7.9 Cytological Identification of Unidentified Interchange Stocks with Various Known Aneuploid Stocks

aneuploids are used as a female parent and cultivars are crossed as a pollen parent. The F_1 population will segregate for disomic and monosomic or monotelosomic plants. In a critical combination, if a cultivar differs by one reciprocal translocation, the majority of sporocytes will show 1III + II (in monosomic) or $1IV^{het}$ + II (in monotelosomic) chromosome association. This suggests that the monosomic chromosome is involved in an interchange. However, in a noncritical combination, a chromosome configuration of 1IV + II + 1I (in monosomic) and $1IV + 1^tII$ (in monotelodisomic) will be recorded (Table 7.9). It should be indicated that the frequency of trivalents or quadrivalents depends upon the size of chromosome segments involved in a translocation; the higher the frequency of quadrivalents, the larger the segments interchanged. Furthermore, the occurrence of chains or open quadrivalent configurations suggests that one of the chromosome segments involved in an interchange is shorter than the others.

G. B–A Interchanges

B–A interchanges or translocations are useful for studying the breakage-fusion-bridge cycle (Zheng, Roseman, and Carlson, 1999) and genetic analysis (Beckett, 1978; Carlson, 1986) in maize. These interchanges have been obtained by translocation between a member of the standard, "A" chromosome, set and a type of supernumerary chromosome, "B" chromosome, found in certain strains of maize. After reciprocal exchanges between A and B chromosomes, A^B and B^A chromosome types are produced (Figure 7.15A).

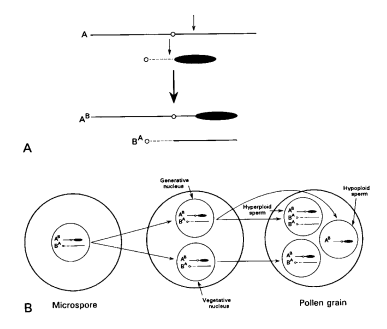


Figure 7.15 B–A translocations. (A) Production of a B–A (A,^B B^A) translocation in maize (arrows show breakage points). (B) Development of hypoploid and hyperploid sperms due to nondisjunction of the B^A chromosome during second mitosis of a microspore. (Redrawn from Beckett, J.B., *J. Hered.*, 69, 27–36, 1978.)

Dominant and recessive traits can be located on A chromosomes by a pseudodominance technique, because B-chromosomes are believed to be largely genetically inert. Roman and Ullstrup (1951) were the first to locate a gene (*hm*) for reaction to *Helminthosporium carbonum* using B-A interchanges.

For locating a recessive gene (r), homozygous recessive (rr) plants are used as female and are pollinated with a B-A interchange carrying a dominant gene (R) for aleurone and plant color on the B^{A} chromosome. As expected, each egg and polar nucleus will have r genotype, and the female contribution to the endosperm will be of rr genotype. In contrast, hyper- and hypoloid sperm nuclei will be generated in B-A interchanges because of nondisjunction of a B^A chromosome at the second mitotic division in the microspore (Figure 7.15B). When an egg (r) is fertilized by a hyperploid (RR)sperm, the constitution of embryo tissue will be colored, while aleurone will be colorless, because it lacks a B^A chromosome and, thus, an R allele. By contrast, the embryo will be colorless when an egg is fertilized by a hypoploid sperm, and aleurone cells of the endosperm will be colored after the polar nuclei are fertilized by a hyperploid sperm carrying an R allele on each of two B^A chromosomes. In maize, endosperm and embryo color traits may be scored on unshelled ears, and if a gene is located on a B^A chromosome, the kernels will segregate into colored and colorless endosperm types, reflecting the frequencies of transmission and functioning of hyper- and hypoploid sperms. In the example given here, noncorrespondence between aleurone and embryo color is evidence of nondisjunction. On the other hand, if the dominant allele of a gene is not located on a particular BA chromosome, hypoploidy for that chromosome will not uncover the relevant recessive phenotype.

A dominant gene can also be located on a chromosome by a further test. A dominant gene in hemizygous condition will be expressed in F_1 plants. If selfed or test crossed to recessive plants, all progeny will be dominant. To test whether the dominant gene is located on an A^B chromosome proximal to the interchange point, the hypoploid is selfed or back crossed to recessive plants. A locus just proximal to the interchange point in A^B may yield all, or nearly all, dominant progeny. Positions on an A^B farther from the interchange point will indicate by recombination values whether there is linkage and its strength.

If a gene is not located on either interchanged chromosome of a B–A translocation, a 3:1 or a 1:1 ratio will be observed in F_2 or test cross progenies of hypoploid plants.

B–A interchanges may be maintained in the heterozygous condition by crossing B–A translocation heterozygotes with pollen from plants carrying normal karyotype. About 1/4 of the plants in progenies are expected to be heterozygous for the translocation. These plants can be identified by pollen sterility or by chromosome analysis. A modern maintenance procedure consists of developing stocks of homozygous B–A translocations. These are maintained by selfing or sibbing. For use in research, these homozygous stocks may be used directly. Alternatively, they may be crossed as male parents to vigorous standard strains. All nonhypoploid progeny will be heterozygous or hyperploid, suitable in either case for locating genes (Beckett, 1978).

H. Role of Interchanges in Evolution of Crops

Interchanges in nature have played a major role in the evolution and speciation of several crop species. Earlier investigations on interchanges in *Oenothera* led to the discovery of several genetic systems. Among cultivated crop species, the role of reciprocal translocations in the speciation of *Secale* species is a classical example, and numerous cytogenetic investigations have been conducted to determine the progenitor of cultivated rye, *Secale cereale*.

1. Oenothera Species

Hugo de Vries (1901) formulated a mutation theory based on the peculiar behavior of *Oenothera lamarkiana* and developed the *Theory of Intracellular Pangenesis* to explain the mutants. However, it was proved later that the majority of deVries' mutants were not the result of gene mutation but

were due to chromosomal aberrations: trisomics, triploids, tetraploids, and interchanges. Furthermore, all the deVries' lines studied bred true and behaved like pure lines, because they were permanent heterozygotes known as *Renner Complex* (Cleland, 1962). This peculiar behavior is attributed to several mechanisms, including gametophytic and zygotic lethals, microspore and megaspore competition (Renner effect), and self-incompatibility. Burnham (1962) and Cleland (1962) discuss cytogenetics of *Oenothera* at great length. Thus, only salient points will be discussed here.

a. Gametophytic and Zygotic Lethals

Gametophytic and zygotic lethals are found in most of the *Oenothera*. Gametophytic lethals are megaspore lethal or pollen lethal. In such cases, sperm do not fertilize eggs of the same genotype due to the slow growth of pollen tubes or the slow development of embryo sacs. Therefore, only gametes of differing genotypes are compatible, producing only heterozygous plants. In *O. lama-rkiana*, one gamete complex is called *velans*, and the other is called *gaudens*. Homozygous *velans* or *gaudens* are lethal in the zygotes and do not survive; thus, only heterozygous *velans/gaudens* survives. This particular mechanism is known as a balanced lethal (Cleland, 1962).

b. Microspore and Megaspore Competition

Megaspore competition probably occurs in all the *Oenothera*. The pattern of female transmission is due to genetic compatibility or incompatibility regarding embryo sac development. The distortion in genetic ratio depends upon problems related to interacting effects of megaspore and pollen tube growth. In *O. hookeri*, the embryo sac develops regularly from the micropylar spore. In contrast, the chalazal cell often develops in *O. muricata* to form an embryo sac. Virtually all the eggs carry "rigens" spores, and thus, "curvans" spores rarely reach the egg. Therefore, "rigens" finds itself in the mycropylar spore, and this particular cell develops into an embryo sac (Cleland, 1962).

c. Chromosome Designation in Oenothera

Chromosomes in *Oenothera* were designated based on their meiotic associations. Attempts have not been made to conduct karyotype analysis, because somatic chromosomes are small and morphologically similar. Considering the "hookeri" complex primitive and having standard and original chromosome end arrangement, the chromosomes of other *Oenothera* species were designated. It has been observed that reciprocal translocation was not limited to the 14 ends of the chromosome complement, but every one of the 14 ends has been found associated with every other end among the chromosome arrangements. Thus, there are 91 possible associations of 14 ends by twos, and they have been observed among the complexes where segmental arrangements have been fully determined.

O. hookeri:	1.2	3.4	5.6	7.8	9.10	11.12	13.14	
[Standard]	1.2	3.4	5.6	7.8	9.10	11.12	13.14	
O. hookeri:	1.2	3.4	5.6	7.10	9.8	11.12	13.14	
								= 🔾 4 + 5II
[Johansen arrangement]	1.2	3.4	5.6	7.10	9.8	11.12	13.14	

This arrangement is quite common among the races of *O. hookeri* found in California and adjacent areas.

O. lamarkiana:	1.2	3.4	5.6	7.8	9.10	11.12	13.14	hookeri
	1.2	3.4	7.6	5.8	9.10	11.12	13.14	velans
	1.2	3.4	5.6	7.8	9.10	11.12	13.14	hookeri
	1.2	3.9	5.6	7.11	14.8	4.12	13.10	gaudens

Thus, there have been 15 possible types of arrangement of chromosomes into circle pairs. Chromosomes in all the *Oenothera* are equal in size, with median centromeres leading to the regular separation of adjacent chromosomes to opposite poles. Although repeated interchanges occurred in the genomes of *Oenothera*, most of the chromosomes still carry their unaltered chromosome morphology. Interchanges were of equal length, and breaks occurred near centromeres. In contrast, chromosome morphologies in *Rhoeo* differ significantly from each other, resulting in uneven spacing of centromeres in a circle at metaphase-I. This arrangement leads to a high degree of irregularity at anaphase-I, when chromosomes move to their respective poles.

2. Secale Species

The genus *Secale* L. consists of a cultigen, rye (*S. cereale*), and four major wild species, namely, *S. vavilovii*, *S. africanum*, *S. montanum*, and *S. silvestre*. All contain 2n = 14 chromosomes. Wild and cultivated species are separated from each other by an effective reproductive isolation, and reciprocal translocations have played an important role in the evolution of cultivated rye.

Of the four wild species of *S. cereale*, *S. silvestre* is an unique species. It differs from others morphologically and cytogenetically (Khush, 1962, 1963) and shows the least Giemsa C-banding pattern (Singh and Röbbelen, 1975; Bennett, Gustafson, and Smith, 1977). Furthermore, the cross-ability rate with other species is extremely low. Khush (1962) obtained 0–0.8% seed set in *S. cereale* \times *S. Silvestre*, but Singh (1977) did not record mature seed set from any combinations involving *S. silvestre* (Figure 7.16). In these crosses, seed developed for about 15 days but collapsed later.

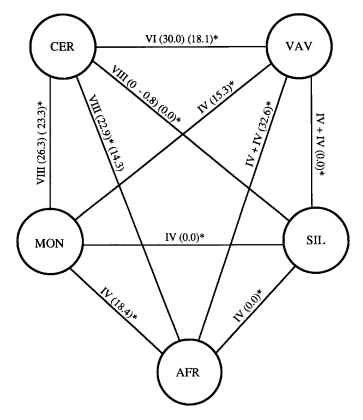


Figure 7.16 Cytogenetic relationships among five major Secale species. Abbreviations: AFR, africanum; CER, cereale; MON, montanum; SIL, silvestre; VAV, vavilovii. Crossability rates (%) are in parentheses (From Khush, G.S., Evolution, 16, 484–496, 1962. With permission.) and in parentheses with an * (From Singh, R.J., Cereal Res. Commun., 5, 67–75, 1977. With permission.).

Plants could have been obtained through embryo culture. It is interesting to note that *S. silvestre* and *S. vavilovii* did not produce hybrid seed, although both are annual and self-pollinating. These results support the suggestion of Khush (1963) that *S. silvestre* should be placed in a separate section, *Silvestria*.

The nature of the barriers among *Secale* species is partly geographical, and the isolated habitats (Stutz, 1972) rarely allow natural hybridization (Kranz, 1963, 1976). On the other hand, translocations have introduced limitations of genetic exchange by chromosomal sterility.

Secale montanum is assumed to be the oldest within the four major species, S. silvestre, S. africanum, S. vavilovii, and S. cereale (Riley, 1955; Khush and Stebbins, 1961; Khush, 1963). Among the different attempts to describe the translocation system within Secale (Riley, 1955; Khush, 1962; Kranz, 1963), the first consistent evidence on the relationships between these species based on Giemsa C-banding patterns (chromosomal differentiation) was presented by Singh and Röbbelen (1977).

Interspecific hybrids of S. montanum with S. vavilovii and with S. africanum, respectively, showed a ring of four chromosomes. But in the two cases, different chromosomes are involved. Evolution apparently occurred separately from these two species through one different translocation each. The same translocation that distinguishes S. montanum from S. vavilovii was found in S. cereale but differed for the second interchange, resulting in IIV + 3II configuration in F_1 of S. cereale and S. vavilovii. Additional translocations occurred to the extent that S. cereale is now separated from S. vavilovii by two interchanges and from S. africanum and S. montanum and S. silvestre by three interchanges (Figure 7.16). As expected, the F_1 involving S. africanum and S. vavilovii showed two quadrivalents (Figure 7.17A). In S. cereale \times S. africanum, but more frequently in crosses with S. montanum, the configuration of 1VIII + 3II was observed (Figure 7.17B), but this association was reduced in certain nuclei to 1VI + 4II (Singh and Röbbelen, 1977). Obviously, one of the three translocations carried only a rather small segment. Riley (1955) also found a chain of eight chromosomes in S. cereale \times S. delmaticum hybrids, though with even smaller frequency (1/70). Secale delmaticum and S. montanum are similar in gross chromosome structure (Riley, 1955). Stutz (1972), on the other hand, described 7II in hybrids of S. cereale \times S. vavilovii, supposing complete structural similarity between these species. For these differences, the only plausible explanation is that the material of Stutz was of different origin, and that the easy pitfall of misclassification of species was not effectively avoided (Khush, 1963).

Secale vavilovii and S. africanum showed 2IV + 3II, but the 1IV + 5II configuration was found in combinations of S. montanum and S. vavilovii (Figure 7.16; Khush, 1962). Secale montanum and S. africanum also gave 1IV + 5II. Thus, three species differ from each other with respect to the translocated chromosomes. It is evident that the same chromosomes cannot be involved to distinguish S. cereale from S. vavilovii and S. montanum, as Heemert and Sybenga (1972) postulated.

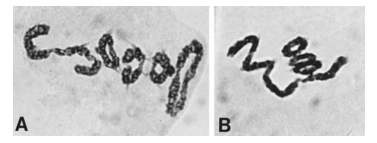


Figure 7.17 Meiotic metaphase-I configurations in *Secale* interspecific F₁ hybrids. (A) *S. africanum* × *S. vavilovii* (2 IV + 3 II); (B) *S. cereale* × *S. montanum* (1VIII + 3 II). (From Singh, R.J. unpublished results.)

In none of the above interspecific hybrids studied was the nucleolus organizer chromosome (SAT-chromosome) included in translocations. This confirms previous findings (Jain, 1960; Kranz, 1963; Heemert and Sybenga, 1972) and suggests that the SAT-chromosome is less subject to structural changes, possibly due to functional restrictions. However, the SAT-chromosome is vulnerable to structural changes when incorporated into a wheat nucleus, producing translocations and substitution lines (Zeller, 1973).

IV. INVERSIONS

A. Introduction

A change in the linear sequence of the genes in a chromosome that results in the reverse order of genes in a chromosome segment is called an inversion. Inversions are widespread in plants, insects, and mammals, and are found in nature or induced by radiation and chemical mutagens. Chromosomal breakage followed by healing produces new chromosomes. Inversions are responsible for speciation (Stebbins, 1950). In plants, McClintock (1931) was the first to show cytologically in maize that when a long inversion is heterozygous, a large loop is seen at pachynema, and bridge(s) and acentric fragment(s) may occur at meiotic anaphase-I and anaphase-II.

B. Types of Inversions

Based on the positions of the two breaks of an inversion in relation to the kinetochore of a chromosome, inversions are of two types: (1) paracentric inversion and (2) pericentric inversion.

1. Paracentric Inversion

In a paracentric inversion, both breaks occur in the same arm, so the inverted region does not include a kinetochore. In a heterozygous inversion, a loop is observed at pachynema, and chromatin bridges and acentric fragments may be found at anaphase-I and anaphase-II. A dicentric bridge and acentric fragment at anaphase-I may arise due to an error in the normal process of crossing over (C.O.). However, if the bridge and fragment are moderately constant, then this is most likely due to a paracentric inversion (Sjödin, 1971).

a. Cytological Behavior

Different configurations of dicentric bridges and fragments at anaphases-I and -II are expected depending upon the types of crossing over within the loop (Figure 7.18; Table 7.10). A single crossing over within the loop will produce a BF (bridge fragment); 2-strand double crossing over in the loop will yield two normal (*in*) and two inversion (*IN*) chromatids, and B and F will not be observed; a 3-strand double C.O. will produce B and F together with one *in* and one *IN* chromatids. A 4-strand double will give rise to two B and two F. This will result in all the unbalanced chromatids (Table 7.10). It gets more complicated when simultaneous crossing over occurs at position 6, and there is a single C.O. in the loop. This will produce LF (loop and fragment) and BF together with *in* and *IN* chromatids. Thus, 50% of the chromatids are expected to be genetically balanced (Table 7.10). Furthermore, the situation gets even more complicated following a triple C.O. (one at 6 and a double C.O. in the loop). Crossing over at 6 and 1 and 2 (3-strand) will produce only *in* and *IN* chromatids. Crossing over at 3 (2-strand) and 4 (4-strand) and simultaneously at 6 will generate only LLFF chromatids, and likewise, triple C.O. at 6, 1, and 5 will produce BBFF chromatids. In those cases, all the produced gametes are unbalanced (McClinock, 1938a; Burnham, 1962; Ekberg, 1974) (Table 7.10).

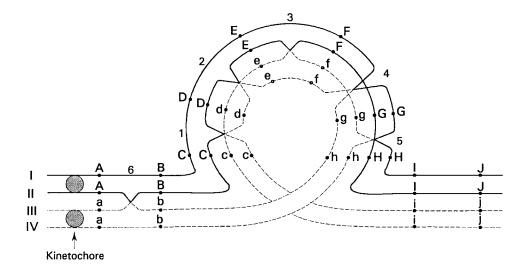


Figure 7.18 A paracentric inversion heterozygote at pachynema. The numbers represent the positions of crossing overs. Chromatids are designated by numbers I, II, III, and IV. Chromatids I and II contain dominant gene loci, and their corresponding recessive alleles are on Chromatids III and IV. Crossover products are listed in Table 7.10.

	Chromatid	Constitution			Balanced
Crossing-Over (C.O.) in Inversion	Normal (in)	Inversion (IN)	Anaphase-I	Anaphase-II	Chromatids (%)
		I — C.O. in inv	version loop at:		
1,2 = 2-Strand double	2	2	in	No bridge	100
1,3 = 3-Strand double	1	1	BF	No bridge	50
3,5 = 3-Strand double	1	1	BF	No bridge	50
3,4 = 4-Strand double			2B, 2F	No bridge	0
	II — C.O. a	t 6 and single	C.O. in inversi	on loop at:	
3 = 2 Strand	1	1	LF	Bridge in one cell	50
1 = 3 Strand	1	1	BF	No bridge	50
5 = 3 Strand	1	1	BF	No bridge	50
4 = 4 Strand	1	1	LF	Bridge in one cell	50
	III — C.O. a	t 6 and double	e C.O. in invers	ion loop at:	
1,2	2	2	in	No bridge	100
1,3	1	1	BF	No bridge	50
1,4	1	1	LF	Bridge in one cell	50
1,5			2B2F	No bridge	0
3,4			2L2F	Bridge in both cells	0

Table 7.10 Chromosome Constitution in a Paracentric Inversion

Abbreviations: in = normal; IN = inversion; B = dicentric bridge; F = acentric fragment; L = dicentric loop; LF = loop and fragment.

Anaphase configurations, where Chromosome 4 in maize contained an inverted segment (paracentric), were examined thoroughly by McClintock (1938a). The frequencies (%) of various configurations were as follows: 1B + 1F = 41.4; 2B + 2F = 3.1; and F = 2.6. Anaphase-II configurations depended upon the preceding anaphase-I configurations: no B + 1F = 42.7%; no B + 2F = 1.3%; 1B + 1F = 3.0%. Anaphase-II configurations with 1B + 1F in each daughter nucleus can be readily

	N	BF	LF	BBFF	LLFF	Number PMCs Studied
NI501 AI	41	34	22	3	1	959
All	38	35	22	3	2	333
CI501 AI	42	36	20	2	1	1327
All	35	34	29	1	1	315
XI501 AI	76	13	10	0.3	0.3	689
All	78	14	8	0	0	100

Table 7.11 Normal and Aberrant Sporocytes (%) at Anaphase-I and Anaphase-II in Three Heterozygous Paracentric Inversions in Barley

Abbreviations: N = normal sporocytes; BF = bridge fragment; LF = loop fragment. *Source:* From Ekberg, I., *Hereditas*, 76, 1–30, 1974. With permission.

recognized, and this configuration is due to a triple crossing over. The observed frequency suggests that this rarely occurs. This conclusion can be confirmed from the observation of Russell and Burnham (1950). They recorded no bridges at anaphase-II in a paracentric inversion heterozygote for Chromosome 4 of maize. They observed the higher frequencies of cells with B and F (average 11.4%) at the second meiotic division.

Ekberg (1974) examined different configurations at anaphase-I and anaphase-II in three paracentric inversion heterozygotes of barley (Table 7.11). Two inversion stocks (NI501, CI501) showed similar configurations, while XI 501 differed significantly, showing 76% normal sporocytes. The frequencies of BBFF and LLFF were low (0 to 3%). A crossover within the inversion accompanied by a proximal crossover was the most common event.

In a *Vicia faba* paracentric inversion, the frequencies (%) of anaphase-I (anaphase-II), studied by Sjödin (1971) were as follows: in = 27 (21); BF = 40(24); LF = 30(52); BBFF = 2(0.2); and LLFF = 0.4(2). Again, the frequencies of BBFF and LLFF were the lowest.

Das (1955) examined four x-ray-induced paracentric inversions in barley. The frequencies of 1B + 1F at anaphase-I ranged from 49.99% (D59–14) to 60.61% (D58–15). In contrast, only a few cells (approximately 2% of the sporocytes) of hybrids involving barley cultivar OAC 21 with B and F were detected (Powell and Nilan, 1968). This suggests that the long sections of chromosomes around kinetochores are not involved in crossing over. Thus, this particular paracentric inversion cannot be detected by B and F. Therefore, the occurrence of localized chiasmata in barley may be the reason for the rarity of reported inversions.

Detection of inversions is influenced by temperature. Swanson (1940) observed in *Tradescantia* F_1 hybrids that warmer temperatures increased chiasma formation, particularly in the interstitial regions, with a correlated increase in bridge frequency (Table 7.12). Detection of inversions is also influenced by temperature in barley. Powell and Nilan (1963) found optimal C.O. in an inversion heterozygote of barley between 15 and 21°C. Thus, under field conditions when temperatures are adverse, crossing over in short inversions in varietal hybrids may be too low for detection.

The breakage of dicentric bridges occurs at various positions in different sporocytes at telophase-I and telophase-II. Fragment chromatids remain in the cytoplasm and can be distributed at random to either pole at telophase-I (dyad) or telophase-II (quartet). Fragments not included in nuclei may form micronuclei.

Table 7.12 Effect of Temperature on Production of Bridges in Harvard Hybrid Tradescantia

Temperature (°C)	Interstitial Chiasmata per Cell	Terminal Chiasmata per Cell	Univalent (%)	Bridges (%)
12–15	1.6	6.07	78.5	17.5
19–22	2.88	7.7	54.0	26.1
27	3.8	7.62	53.0	47.8

Source: From Swanson, C.P., Genetics, 25, 438-465, 1940. With permission.

Paracentric inversions have been studied in several plant species, including maize (McClintock, 1931; Morgan, 1950; Russell and Burnham, 1950; Rhoades and Dempsey, 1953), barley (Smith, 1941; Das, 1955; Holm, 1960; Powell and Nilan, 1963, 1968; Kreft, 1969; Ekberg, 1974; Yu and Hockett, 1979), *Tradescantia* (Swanson, 1940), *Vicia faba* (Sjödin, 1971), and *Lilium formosanum* (Brown and Zohary, 1955). The widespread occurrence of inversions in *Tradescantia* suggests that inversions have played a major role in speciation. Paracentric inversion is rather common in *Drosophila*, because crossover products, dicentric bridges, and fragments, are selectively eliminated into polar bodies during oogenesis by preferential segregation (Sturtevant and Beadle, 1936).

b. Fertility

The characteristic feature of chromosomal interchanges and inversions is that these chromosomal aberrations produce varying degrees of pollen and ovule abortion. Paracentric inversion causes pollen and ovule abortion, because single crossovers, certain double crossovers in the inversion loop, and triple crossovers (rare) generate bridges and fragments at anaphase-I and anaphase-II, producing spores containing chromatids with duplications and deficiencies. These spores generally abort, and the degree of pollen abortion can be predicted from the frequencies of anaphase-I and anaphase-II cells containing bridges and fragments. If crossing over does not occur in the inversion loop, all the spores would be functional, because two spores would contain normal chromatids, and two spores would have inverted chromatids.

Morgan (1950) observed 28.2% pollen abortion and 4% ovule abortion in an In4a paracentric inversion of maize. The normal sibs showed 2.9% aborted pollen. Russell and Burnham (1950) studied a paracentric inversion involving Chromosome 2 (In2a). They did not find ovule abortion, but pollen abortion ranged from 12.4 to 21.6% with an average of 16.5%. Pollen abortion in In3a of maize ranged from 11.5 to 27.6% (Rhoades and Dempsey, 1953). They stated that the observed percentage of abortion was always less than that indicated by the cytological observations.

In a paracentric inversion of *Vicia faba*, Sjödin (1971) observed an average of 37% (range 28 to 59%) pollen abortion, which agreed with the predicted (38%) value. In barley, Ekberg (1974) studied pollen and seed fertility of three paracentric inversion stocks. The degree of pollen fertility and seed set did not differ significantly (Table 7.13).

According to McClintock (1938a), if a fragment chromosome is included in a tube nucleus and if the fragment compensates a deficiency portion, the tube nucleus would contain a complete genomic complement. Thus, it could be expected to function in the growth of a pollen tube with a deficient Chromosome 4 into an embryo sac. The resulting zygote would be heterozygous for a broken Chromosome 4 with a terminal deficiency of the long arm. Such plants were recovered.

c. Linkage Studies

Paracentric inversions have been used in locating genes in specific segments of chromosomes in maize (Morgan, 1950; Russell and Burnham, 1950; Rhoades and Dempsey, 1953), barley (Ekberg, 1974), and *Drosophila* (Burnham, 1962).

Table 7.13	Pollen Fertility, Seed Set, and Segregation Ratios for Three
	Heterozygous Paracentric Inversion Stocks in Barley

Inversion Stock	Pollen Fertility (%)	Seed Set (%)	Segregation Ratio (F:PS)	χ^2 for 1:1 Segregation
NI501	78.2 ± 2.3	78.3 ± 1.1	101:91	0.52
XI501	83.6 ± 2.8	83.3 ± 0.9	233:209	1.30
CI501	74.6 ± 4.6	78.2 ± 1.1	47:33	2.4

Abbreviations: F = fertile; PS = partial sterile.

Source: From Ekberg, I., Hereditas, 76, 1-30, 1974. With permission.

Parental genotype gametes containing *in* and *IN* chromosomes are mostly viable in paracentric inversions. Therefore, inversions are often known as "crossover suppressors." Generally, an inversion homozygote (*IN IN*) carrying a dominant allele, for example *X*, is pollinated by a genetic stock that contains a recessive allele *x* and a normal karyotype (*in*). In inversions and translocations, a breakage point is considered a dominant locus that behaves as a single factor. F_1 (inversion heterozygote) plants are either selfed to produce an F_2 or are back-crossed to the recessive parent. Reciprocal crosses can also be made.

There are two possibilities. A marker gene may be located outside an inversion loop or inside the inversion. In independent assortment, four types of gametes are expected from an F_1 (*X IN*, *X in*, *x IN*, *x in*) in an equal proportion of 1:1:1:1. Partial sterility is used as a marker in the linkage test to identify members of the plant population with the inverted chromosome. Recombinants in BC₁ are scored to determine the association between a marker gene and a break point. If no, or a few, recombinants (*X in*, *x IN*) are found, the break point is located close to the *x* locus. On the other hand, if a higher frequency of recombinants is found, the locus *x* is located farther away.

The other possibility is that the marker locus x is located within the inversion loop. A single crossover within the inversion is expected to result in transmission of only parental-type gametes. Recombinants are produced following 2-strand and 3-strand double crossovers and in equal proportions. Furthermore, simultaneous crossing over at the proximal Region 6 (Figure 7.18) and double crossover or triple crossover within the inversion will not alter the proportions of recombinants.

Rhoades and Dempsey (1953) investigated breakage points of the inversion In3a in relation to genes Rg, lg, A, and et. All genes lie in the long arm of Chromosome 3 in the order given, with Rg nearest to the kinetochore. The standard recombination values between these loci are Rg-Lg = 16%, lg-A = 34%, A-et = 12%, and Rg is 15 map units from the kinetochore. F₁ plants heterozygous for the inversion and for these genetic markers were test crossed to structurally normal plants homozygous for the respective recessive alleles. Table 7.14 shows 16.3% recombination between Rg and Lg in the inversion heterozygote and 15.4% recombination in normal plants. The standard value is 16%. Reduction in recombination (0.6%) occurred between Lg and A in the IN/in fraction, while the recombination value in the *in/in* fraction was 24.9%. Based on the reduction of genetic recombination in the Lg-A region, Rhoades and Dempsey (1953) suggested that these loci are located in the inverted segment, while Rg is in the proximal region.

Dobzhansky and Rhoades (1938) wrote whether an inversion causes the genes in a chromosome to be inherited as a unit depends directly upon certain types of crossover.

2. Pericentric Inversion

In a pericentric inversion, the two breaks occur in opposite arms of a chromosome (Figure 7.19). Dicentric bridges and acentric fragments at anaphase-I and anaphase-II are not observed in pericentric inversions.

Colla	silution		
	IN/in	in/in	Recombinations
rg Lg A (0)	612	181	
<i>Rg lg a</i> (0)	470	162	Rg - Lg = IN/in = 16.3%
<i>rg lg a</i> (1)	108	32	= in/in = 15.4%
<i>Rg Lg A</i> (1)	111	35	Lg - A = IN/in = 0.6%
rg Lg a (2)	0	59	= in/in = 24.9%
Rg lg A (2)	3	60	
<i>rg lg A</i> (1.2)	2	4	
<i>Rg Lg a</i> (1.2)	3	13	

 Table 7.14 Test Cross Data from a Plant of $rg^{(1)} Lg^{(2)} A/Rg Ig a$

 Constitution

Source: From Rhoades, M.M. and Dempsey, E., *Am. J. Bot.*, 40, 405–424, 1953. With permission.

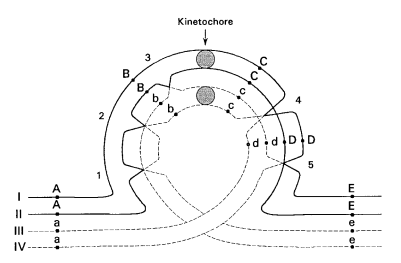


Figure 7.19 A pericentric inversion heterozygote at pachynema. (See Figure 7.18 for explanations.) Cross-over products are listed in Table 7.15.

Pericentric inversions have been found in maize (Morgan, 1950; Zohary, 1955), *Vicia faba* (Sjödin, 1971), *Allium thunbergii* (Watanabe and Noda, 1974), *Scilla scilloides* (Noda, 1974), and in many grasshopper species (White and Morley, 1955). Pericentric inversions are rare in *Drosophila*, because single crossovers between normal and inverted segments in heterozygotes produce Dp and Df chromatids, which are lethal in zygotes or the embryos.

a. Cytological Behavior

Any single crossover at any point and 3-strand double crossovers in pericentric inversion heterozygotes produce chromatids consisting of one normal (*in*), one inversion (*IN*), and two Dp and two Df constitution (Figure 7.19). A 2-strand double crossover generates two normal and two inverted chromatids. However, a 4-strand double crossover (at 1,5) results in all four chromatids with duplications and deficiencies (Table 7.15). Pericentric inversions in Orthoptera are usually characterized by nonhomologous pairing between normal and inverted chromosomes, which prevents crossing over in the inverted region. Thus, pericentric inversions are also known as "*crossover suppressors*," because recombinant strands are not recovered (White and Morley, 1955; Cabrero and Camacho, 1982).

Pericentric inversions may shift the kinetochore position in a chromosome and result in a change of arm ratio. This feature facilitates the identification of inverted chromosomes by karyotype analysis at mitotic and meiotic stages. At metaphase-I of meiosis, inverted chromosomes may contribute to asymmetrical bivalents.

Morgan (1950) identified two pericentric inversions in maize at pachynema. In *Vicia faba*, Sjödin (1971) identified two pericentric inversions at mitotic metaphase. In both cases, Chromosome 1 was involved in the inversion.

Crossing Over in		Chromatid Constitution after C.O.			
Inversion	Position of C.O.	Normal (in)	Inversion (IN)	Dp + Df	
1. Any single C.O. 2. Double C.O.	Any point	1	1	Dp + Df (2)	
2 Strand	1, 2	2	2	_	
3 Strand	1,3 or 1,4	1	1	Dp + Df (2)	
4 Strand	1,5	0	0	Dp + Df (4)	

Table 7.15 Chromosome Constitutions in a Pericentric Inversion after Crossing Over

b. Fertility

Heterozygous pericentric inversions cause pollen and ovule abortion, because spores are produced that contain chromosomes with deficiencies and duplications. In *Vicia faba*, Sjödin (1971) observed about 50% pollen abortion in two heterozygous pericentric inversions. Pollen abortion in heterozygous pericentric Inversion 2b in maize ranged from 18.3 to 25.6% (19.1% average), and ovule abortion was 20.1%. Normal plants showed 2.7 to 4.1% pollen abortion. In pericentric Inversion 5a in maize, the average ovule abortion was 12.5%, and pollen abortion was 28.3%. The differences between ovule and pollen abortion are related to the higher frequencies of crossing over in male than that in female flowers (Morgan, 1950). In the natural population, four pericentric inversion heterozygotes of *Scilla scilloides* (2n = 18) showed an average of 46.4% pollen abortion (Noda, 1974).

CHAPTER 7B

Chromosomal Aberrations — Numerical Chromosome Changes

I. EUPLOIDY

A. Introduction

An individual carrying chromosome numbers other than true monoploid or diploid numbers is called heteroploid (Sharp, 1934). Heteroploidy is divided into euploidy and aneuploidy. In the euploid, an individual carries an exact multiple of the basic chromosome number, while in the aneuploid, the chromosome number is some number other than an exact multiple of the basic set.

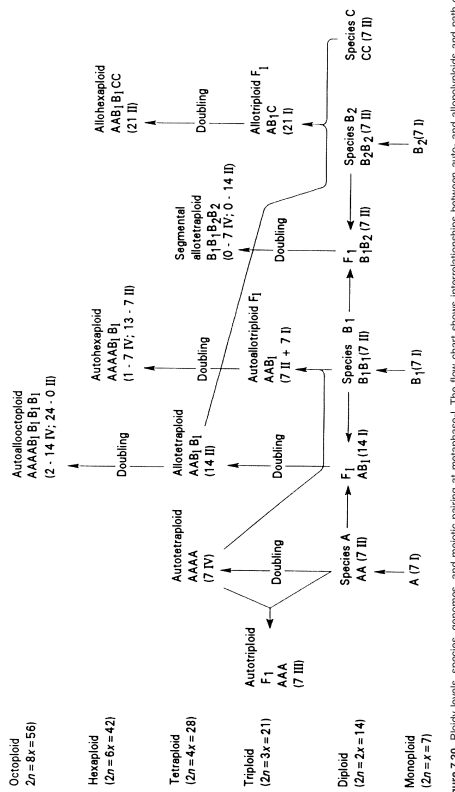
In polyploids, x is the basic (monoploid) chromosome number, n is the gametic chromosome number of chromosomes, and 2n is the zygotic or somatic chromosome number. For example, the genomic formula of *Triticum aestivum* is 2n = 6x = 42 and *Hordeum vulgare* is 2n = 2x = 14. In both cases, the basic chromosome number x is seven. The basic set of chromosomes in a diploid is called a genome.

B. Classification of Euploidy

Euploidy is divided into auto- and allopolyploidy. In the autopolyploids, the genomes are alike, because one basic genome is multiplied (x, monoploid; 2x, diploid; 3x, triploid; 4x, tetraploid; 5x, pentaploid; 6x, hexaploid; 7x, heptaploid; 8x, octoploid). In allopolyploids, two or more genomes derived from different genomically unlike, distinct species are present (Figure 7.21).

Stebbins (1950) recognized two additional euploid categories, namely, segmental allopolyploids and autoallopolyploid combinations. Segmental allopolyploids carry genomes intermediate in degree of similarity and generally exhibit preferential pairing. For example, a segmental allotetraploid with genomes $B_1 B_1 B_2 B_2$ usually forms bivalents and occasionally quadrivalents. Autoallopolyploidy is confined to hexaploidy and higher levels of polyploidy (Stebbins, 1950). The term amphiploidy or amphidiploidy denotes polyploids derived after hybridization between two or more genomically dissimilar species separated by chromosomal sterility (Figure 7.20).

Polyploidy has played a major role in the speciation of higher plants. In angiosperms, 30 to 35% (Stebbins, 1950, 1971), 47% (Grant, 1971), and approximately 70% (Masterson, 1994) of the species are of polyploid origin. The species in which the basic chromosome number is x = 10, or higher, evolved by polyploidization.





1. Autopolyploidy

a. Autotriploids (3x, AAA)

An individual possessing three basic sets of homologous chromosomes is known as autotriploid.

i. Sources of Autotriploids — Autotriploids arise from the progenies of diploid (2x) species by spontaneous origin at low frequencies. They originate from the sexual fusion of an unreduced (2n) egg and haploid (n) male sperm. Harlan and deWet (1975) recorded polyploid individuals in 68 genera that originated from unreduced gametes. Triploids are identified and distinguished from diploids in the natural population because of their more vigorous vegetative growth, profuse tillering, gigas phenotypes and high ovule abortion.

Spontaneously induced triploids have been isolated and identified in numerous sexually and vegetatively propagated crops such as *canna* (Belling, 1921), *Datura* (Belling and Blakeslee, 1922), maize (McClintock, 1929c), tomato (Lesley, 1928; Rick, 1945), rice (Ramanujam, 1937; Rao and Reddi, 1971; Khush et al., 1984), barley (Sandfaer, 1975), pearl millet (Rao et al., 1988; Dujardin and Hanna, 1988), sorghum (Schertz and Stephens, 1965), and others (Burnham, 1962; Balog, 1979, 1984).

Rick (1945) found in tomato that triploids appeared spontaneously at the rate of 0.08% (45/55,000 plants). Sandfaer (1975) observed 0.05% (179/381,563 seeds) triploids of spontaneous origin in barley. Seeds of 39 barley cultivars were examined to determine the frequency of auto-triploids. Barley seeds were divided into three groups: Group A included fully developed seeds that constituted 95.8%, group B contained shriveled seed (2.0%), and Group C had a light "empty" (2.2%). Seeds of Group A were assumed to be diploid (2n = 2x = 14), and chromosomes were not determined. Germination of Group B seeds was 73.9%, and the frequency of triploid plants (2n = 3x = 21) was 2.86%. The germination rate of Group C seeds was only 2.4%, but its members consisted of 10.29% triploid plants (Table 7.16). The frequency of triploid plants may have been influenced by the genotypes of the cultivars, because of the 39 cultivars examined, triploids were recorded (range: 1 to 25 plants) in 22 cultivars.

Autotriploids are produced experimentally from tetraploid (4x) by diploid (2x) crosses. Håkansson and Ellerström (1950) suggested that seed development was better in rye when a plant with the higher chromosome number was used as a female parent rather than the reverse. In the $2x \times 4x$ cross of rye, 1275 spikelets were pollinated and 36 well-developed kernels were obtained. Seventeen seeds germinated; four seedlings were triploid (2n = 3x = 21), and two plants were diploid (2n = 14). In $4x \times 2x$ hybridization, 783 pollinated spikelets yielded 20 well-developed seeds; six seeds germinated, giving four triploids and two tetraploids. Diploid and tetraploid plants may have originated after selfing. Similar results were reported by Tsuchiya (1960b) in barley. In a $4x \times 2x$ cross, 980 seeds were obtained from 2721 pollinations, but only 31 seeds germinated (1.13%). By contrast, triploid plants were not found in a $2x \times 4x$ cross (Table 7.17). In rice, the success rate of finding triploids depended upon the crossing methods and the frequency of triploids ranged from 0.0% to 1.09% (Morinaga and Kuriyama, 1959).

In *Triticum monococcum* (2n = 2x = 14), Kuspira et al. (1986) obtained a total of 1750 paperthin, shriveled seeds from 7700 pollinated florets in a $4x \times 2x$ cross (23.0% seed set). In a $2x \times 4x$ cross, 9200 pollinations resulted in 260 seeds (3% seed set). Not all seeds (2010) germinated. However, 15-day-old embryos from a $4x \times 2x$ cross that were excised aseptically produced triploid plants. This result indicates that endosperm collapse 15 days postfertilization caused the death of embryos.

Twin seedlings in some cases produce triploid. Müntzing (1939) cytologically examined a total of 2201 twin plants from 16 species. Triploid plants were the most frequent (77) followed by haploid (11), and tetraploid plants were rare (2).

Cultivars
39 Barley
33
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of
Summary
16
Table 7.16

	(%)				
	Triploid (%)	Ι	2.86	10.29	
	Aneuploid	I	7	۲	
Number of Plants	iploid (2 <i>n</i> = 14) Triploid (2 <i>n</i> = 21)	I	158	21	
Number	Diploid $(2n = 14)$	1	5364	182	
Germination	(%)	I	73.9	2.4	
Number of	Seeds	365,645	7480	8438	
1000 Kernel	Weight (g)	41.2	13.7	3.5	
Seed Size	Groups	٨	В	o	
	of Seeds		381,563		
Number of Number	Spikes		18,719		

Source: From Sandfaer, J., Hereditas, 80, 149–153, 1975. With permission.

arley
Diploid B
and
Tetraploid
between
Crosses
Reciprocal
from
Success
Crossing
Table 7.17

	Numk	Number of	Fertility		Number of Seeds	
Cross	Florets	Seeds	(%)	Sown	Germinated (%) Success (%)	Success (%)
$4x \times 2x$	2721	980	36.01	980	31 (3.16)	1.13
$2x \times 4x$	77	58	75.32	58	0 (0.0)	0.0
Source: From T	suchiya, T., Sei	ken Zihô, 11, 2	Source: From Tsuchiya, T., Seiken Zihô, 11, 29–37, 1960b. With permission	Vith permissio		

7 Chromosome
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Table 7.

Table 7.18 Frequencies of Trivalents at	encies of	f Trivaleı		taphase-	-l in Autc	triploid€	s in Five	Species	with $x = 7$ (Metaphase-I in Autotriploids in Five Species with $x = 7$ Chromosomes				
		Fre	Frequencies	ies of Trivalents at Metaphase-I	ents at l	Metapha :	se-l			Total Number	Mean	Mean Number/Cell	/Cell	
Species	0	-	2	e	4	2	9	7	III 2 %	of Cells	-	=	≡	Author
Hordeum spontaneum	-	5	4	27	45	53	44	13	6.77	192	2.33	2.34	4.66	Tsuchiya, 1952
Lolium perenne	0	0	0	ო	8	32	53	68	41.46	164	0.79	0.79	5.34	Myers, 1944
Pennisetum glaucum	0	0	0	N	0	7	o	20	52.63	38	0.82	0.82	6.18	Pantulu, 1968
Secale cereale	-	0	4	4	ო	Ŋ	ო	0	00.0	20	3.25	3.25	3.75	Lamm, 1944
Triticum monococcum	2	ω	37	89	123	126	87	29	5.80	500	2.65	2.60	4.38	Kuspira et al., 1986

It has been demonstrated that normal seed development depends on a correct ploidy ratio (2:3) between embryo and endosperm that occurs normally in a $2x \times 2x$ cross (Watkins, 1932; Esen and Soost, 1973). Departure from this quantitative ratio such as 3:4 ($2x \times 4x$) or 3:5 ($4x \times 2x$) between embryo and endosperm genome sets disturbs physiological and genetic balance, leading to the collapse of endosperm and then death of the embryos. In $2x \times 4x$ crosses of citrus, about 92 to 99% of the seeds containing triploid embryos and tetraploid endosperm aborted at different stages of embryogenesis (Esen and Soost, 1973). Lin (1984) dismissed the hypothesis of embryo and endosperm relations based on his studies of diploid and tetraploid maize that involved a gametophyte mutant (*ig*), and suggested, "development of maize endosperm evidently is affected by the parental source of its set of chromosomes." Because he used a gametophyte mutant stock (*ig*) that generates chromosome abnormalities, however, the embryo and endosperm ratio (2:3) theory should perhaps be investigated further.

A large number of autotriploids have been isolated from the progeny of different lines of homozygous recessive male-sterile (Ms1 ms1 ms1) soybean (Xu et al., 2000b). Autotriploids derived from ms1 ms1 soybean always carry the recessive ms1 gene in homozygous (ms1 ms1 ms1) conditions and have poor seed set. Chen and Palmer (1985) isolated 138 plants with greater than 40 chromosomes (2n = 44 to 2n = 71), but not 41 chromosomes, from 32 male-fertile (Ms1 ms1 ms1) triploid plants.

Autotriploids can be induced by mutagens. Pantulu (1968) obtained triploids (2n = 3x = 21) from gamma-ray-irradiated pearl millet seed. Singh and Ikehashi (1981) isolated 93 totally sterile to semisterile plants in an M₂ population of rice treated with a chemical mutagen, ethyleneimine. Two plants were found to be triploid (2n = 3x = 36). Morphologically triploid rice plants were dark green, tillered profusely, and showed extremely low pollen (0.9 to 2.1%) and ovule fertility. The triploids probably originated by the fertilization of unreduced female gametes (2n) (caused by mutagen) by *n* male gametes. Colchicine induces formation of tetraploids as well as triploids (Myers, 1944; Jauhar, 1970).

ii. Cytological Behavior in Autotriploids — In autotriploids, only two of three homologous chromosomes associate at any point during pachynema. Pairing patterns determine the frequencies and types of trivalent association at metaphase-I (Darlington and Mather, 1932; Benavente and Orellana, 1984). Various combinations of univalents, bivalents, and trivalents are possible at diakinesis and metaphase-I in autotriploids (Table 7.18). Although all five species shown are 2n = 3x = 21 and have intermediate chromosome size, they exhibit highly variable trivalent frequencies. The mean numbers of trivalents per cell ranged from 3.75 (Secale cereale) to 6.18 (Pennisetum glaucum). The highest frequencies of sporocytes with 5III + 2II + 2I was observed in barley, rye, and T. monococcum. In autotriploid maize (2n = 3x = 30), 9III + 1II + 1I chromosome association was the most frequent (McClintock, 1929c).

Trivalent association is expected to be more frequent with long chromosomes than with shorter chromosomes (Darlington, 1929; Darlington and Mather, 1932; Burnham, 1962). However, it has been suggested that aside from environmental influences, trivalent frequencies are genetically determined (Kuspira et al., 1986).

Based on random chromosome pairing, crossing over, and chiasma formation during pachynema, there are four possible trivalent configurations at diakinesis: (1) chain (V-shaped), (2) ring-rod (frying-pan), (3) triple arc (birdcage), and (4) Y-shaped (Figure 7.21A). The V-shaped trivalent can arise by distal chiasmata involving one chromosome on both arms. A frying-pan (ring-rod) requires three crossover and three chiasmata (one interstitial and two distal). The triple arc configuration is rarely observed and arises by four reciprocal exchanges (two interstitial and two distal) in both arms. A Y-shaped trivalent needs a minimum of two reciprocal exchanges (one interstitial and one distal) involving only the same arm of all three homologues (Figure 7.21A).

Each group of three homologous chromosomes at diakinesis and metaphase-I can coorient in three different ways: (1) linear, (2) convergent, and (3) indifferent. The convergent pattern, partic-

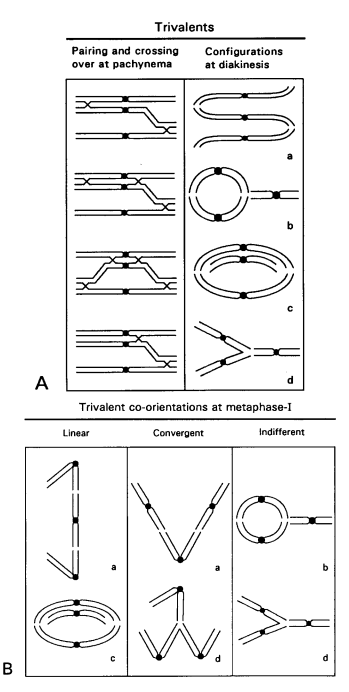


Figure 7.21 (A) Possible trivalent configurations at diakinesis in an autotriploid, based on crossing over and pachytene chromosome pairing. (B) Types of trivalent coorientations at metaphase-I in an autotetraploid. (Redrawn from Kuspira, J.R. et al., *Can. J. Genet. Cytol.*, 28, 867–887, 1986.)

ularly V-shaped trivalent coorientation at metaphase-I [Figure 7.21B(a)], is the most frequent. Convergent Type d [Figure 7.21B(d)] is an hypothetical configuration and is generally not observed. Linear types rod-shaped [Figure 7.21B(a)] and triple-arc [Figure 7.21B(c)] and indifferent, i.e., ring-rod [frying-pan, Figure 7.21B(b)] and Y-shaped [Figure 7.21B(d)] orientations are relatively rare (Table 7.19). It has been elucidated that orientation of the kinetochores within trivalents at

		Trivalent Coor	ientations at	t Metaphase-I		
Species	2 <i>n</i>	Convergent (%)	Linear (%)	Indifferent (%)	Total	Authors
Hordeum spontaneum	21	806 (90.2)	76 (8.5)	12 (1.3)	894	Tsuchiya, 1952
Triticum monococcum	21	561 (93.5)	0 (0.0)	39 (6.5)	600	Kuspira et al., 1986

Table 7.19 Frequencies of Trivalent Coorientations at Metaphase-I in Autotriploids

metaphase-I toward the poles in meiocytes is random (Satina and Blakeslee, 1937; Balog, 1979; Kuspira et al., 1986).

Disjunction of chromosomes at anaphase-I in trivalents is mainly determined by the orientation and coorientation of their kinetochores at metaphase-I. In convergent coorientations, two chromosomes regularly go to one pole and the middle one to the opposite pole. Chromosomes of indifferent orientation often migrate in a 1:1 fashion. Occasionally, a 2:1 segregation is found. Linear alignments usually segregate in a 1:1 fashion, and the middle chromosome behaves like a lagging univalent. These observations indicate that chromosome migration at anaphase-I is nonrandom (Darlington, 1929; Burnham, 1962; Balog, 1979).

In autotriploids (2n = 3x = 21) of barley and *T. monococcum*, the highest frequency of sporocytes at anaphase-I showed ten to 11 chromosome disjunction followed by nine to 12, eight to 13, and seven to 14 (Tsuchiya, 1952; Kuspira et al., 1986). Lagging chromosomes in barley and *Lolium perenne* ranged from zero to six, and in *T. monococcum*, ranged from zero to three (Table 7.20). A contrasting difference among three autotriploids was the frequencies of sporocytes without laggards. In *T. monococcum*, 89.2% of the sporocytes did not carry laggards, but the frequencies were much lower in barley and ryegrass (Table 7.20). Some of the laggards may be included in dyads. Tsuchiya (1952) found 50% telophase-I cells without laggards, while anaphase-I cells showed 29.92%. Lagging chromosomes that do not reach either pole are eventually eliminated.

iii. Fertility and Breeding Behavior in Autotriploids — Ovule fertility in autotriploids ranges from high sterility to complete fertility. Complete sterility has been found to be useful for breeding seedless fruits, particularly seedless watermelon (Kihara, 1958), tuber-bearing plants, vegetatively propagated ornamentals, and fruit trees.

Although pollen fertility was high in triploids of *Datura* (Satina and Blakeslee, 1937), barley (Tsuchiya, 1952), *T. monococcum* (Kuspira et al., 1986), and pearl millet (Dujardin and Hanna, 1988), seed set after selfing was poor. In triploid *Datura*, while Satina and Blakeslee (1937) recorded pollen fertility between 50 and 60%, only 15% or less of pollen germinated, and of the latter, nearly two-thirds burst within or near the stigma, leaving about 5% functional pollen. These results suggest that microgametophytes with other than haploid chromosome numbers are unviable, because pollen grains with the aneuploid chromosome numbers (n + 1, n + 2, or higher) are genetically and physiologically unbalanced and cannot compete in fertilization with *n* chromosome spores. Pollen transmission of an n + 1 gamete is rare.

		Freq	uency o	f Cells w	vith Lago	jards		Total Number of	
Species	0	1	2	3	4	5	6	Cells	Authors
Lolium perenne Hordeum spontaneum	37.0 29.92	26.0 26.26	17.0 21.26	9.0 15.35	9.0 2.36	1.0 3.15	1.0 0.79	1103 127	Myers, 1944 Tsuchiya, 1952
Triticum monococcum	89.20	7.2	1.20	2.4	0.0	0.0	0.0	250	Kuspira et al., 1986

Table 7.20 Frequencies (%) of Laggard at Anaphase-I in Three Autotriploid (2n = 3x = 21) Species

On the female side, gametes with n, n + 1, n + 2, n + 3, and rarely n + 4 and n + 5, are functional. The maximum limit of tolerance of extra chromosomes is usually three in true diploids. Therefore, gametes and zygotes containing more than three extra chromosomes in addition to their normal chromosome complement will typically abort. However, there are a few exceptions where autotriploids set normal seed. These cases indicate that male and female gametes can sometimes tolerate unbalanced chromosome numbers resulting in a high frequency of viable gametes, and viable progeny that are produced may include diploid as well as most of the possible aneuploids. In autotriploids of spinach, Janick and Stevenson (1955) observed normal seed set. Progenies of $3x \times 2x$, and the reciprocal, yielded plants ranging from diploid (2n = 2x = 12) to triploid (2n = 3x = 18) (Tabushi, 1958).

b. Hypertriploids

i. Sources of Hypertriploids — Occasionally, trisomics generate a low frequency of triploids (3x) and near triploids, such as hypotriploids (2n = 3x - 1) and hypertriploids (2n = 3x + 1), in their progenies (Tsuchiya, 1960a, 1967; Singh and Tsuchiya, 1975a; Xu et al., 2000b). Hypertriploid (2n = 3x + 1 = 22) plants in barley were relatively short as compared to diploid, but there was a marked increase in leaf size and stomatal length in hypertriploids compared to diploid sibs (Singh and Tsuchiya, 1975a). Hypertriploid soybean plant was more vigorous in vegetative growth with robust main branches and large, dark-green leathery leaves, and matured much later than its disomic sib, a typical trait of an autotriploid plant (Xu et al., 2000a). A spontaneous hypertriploid (2n = 3x + 1 = 61) plant in the soybean was identified in the progeny of a cross between T31 (glabrous, p2 p2) and an unidentified primary trisomic (2n = 41) (Xu et al., 2000a). It is assumed that hypertriploid plants probably originated by the fusion of an unreduced egg (n = 40) from T31 and an n + 1 (n = 21) sperm from the primary trisomic line. The frequency of hypertriploid in the soybean is extremely low. All 146 F₁ plants involving T31 (female) crosses with 14 primary trisomics contained either 2n = 40 or 2n = 41 chromosomes.

ii. Cytological Behavior in Hypertriploids — In hypertriploids (2n = 3x + 1), one chromosome is present four times, while others occur in three copies. Hypertriploid plants have been found in several plant species. Singh and Tsuchiya (1975a) isolated four hypertriploid barley plants (2n = 3x = 22) in the progenies of Triplo 4 (Robust). Thus, it was assumed that Chromosome 4 was present four times. As was expected, chromosome configurations at metaphase-I were various combinations of quadrivalents, trivalents, bivalents, and univalents. The maximum chromosome association was 1IV + 6III (Figure 7.22A). Quadrivalents were either N-shaped or with the fourth chromosome associated with tandem-V trivalent (Figure 7.22A,B). An association of 6III + 1II + 2I in some cells indicates that a quadrivalent dissociated to form 1III + 1I and a trivalent 1II + 1I. The minimal chromosome association was 1IV + 1III + 5II + 5I. The frequency of univalents ranged from zero to five. An average frequency of chromosome association per cell in all hypertriploids was 0.35IV + 4.91III + 2.06II + 1.73I. Meiotic chromosome pairing in hypertriploid soybean was similar to the barley hypertriploid. Theoretically, the chromosome configurations at metaphase-I in soybean hypertriploid are mainly the following three types:

- 1. 1IV + 19III, 1IV + 18III + 1II + 1I ... 1IV + 19II + 19I.
- 2. 19III + 2II, 18III + 3II + 1I, 17III + 4II + 2I \dots 1III + 20II + 18I.
- 3. 20III + 1I, 19III + 1II + 2I, $18III + 2II + 3I \dots 1III + 19II + 20I$.

Meiotic chromosome pairing in soybean hypertriploid showed the above three types of chromosome association in 90% of the studied PMCs, which is similar to the barley hypertriploids. In soybean hypertriploid, the second type of chromosome configuration was observed in 63% of the PMCs, suggesting that the four homologous chromosomes primarily formed two

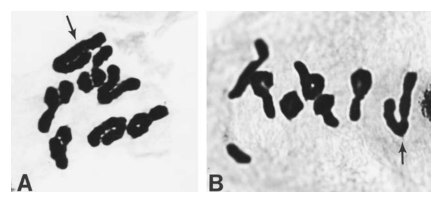


Figure 7.22 Meiotic metaphase-I configurations in an hypertriploid (2n = 3x + 1 = 22) in barley. (A) 1IV (arrow) + 6III; (B) 1IV (arrow) + 5III + 1II + 1I (From Singh, R.J. and Tsuchiya, T., *Caryologia* 28, 89–98, 1975a. With permission.)

bivalents (Xu et al., 2000a). In barley hypertriploid, the frequency of the first type was marginally higher than that of the second type. These differences are probably due to the difference in chromosome size between soybean and barley. The soybean chromosomes are about ten times smaller than those in barley.

The genetic male sterile soybean autotriploids showed chromosome association of enneavalent (association of three trivalents), hexavalent, pentavalent, and quadrivalent at diakinesis and metaphase-I (Chen and Palmer, 1985). Nonhomologous chromosome association was proposed. However, Xu et al. (2000a) recorded only quadrivalent, and higher multivalents were not found in hypertriploid. Precocious chromosome separation in some bivalents and trivalents was common in some PMCs. This may be due to an early terminalization of chiasma in small soybean chromosomes.

The most frequent trivalent type in hypertriploids studied in barley was ring-rod (frying-pan) followed by tandem-V and tandem chain. The frequencies of triple arc and Y-shaped trivalents were low and similar to the autotriploids. It is interesting to note that the tandem-V trivalent was found in the highest frequency in autotriploids of barley (Tsuchiya, 1952). It was not possible to examine the shape of trivalents in the soybean hypertriploid because of small chromosomes. Chromosome disjunction at anaphase-I was fairly normal in 42.5% cells, without lagging chromosomes. The number of lagging chromosomes ranged from zero to four. Chromosome disjunction at anaphase-I in soybean hypertriploid was mostly 30 to 31 resulting meiotic products with higher chromosome numbers.

iii. Fertility and Breeding Behavior in Hypertriploids — Hypertriploid plants in the soybean showed 63% fertile pollen, while diploids had 98%. The first flush of flowers generally abort, and that was also observed for the hypertriploid. The hypertriploid produced 98 selfed seeds, 16 seeds in hypertriploid (female) × diploid (male), and no seed in the reciprocal cross.

The self seeds of the soybean hypertriploid germinated poorly, and only 43 (44%) of 98 seeds could germinate. Most of the shrunken and irregular-shaped seeds were not viable, which may be attributed to poor endosperm development. The chromosome number ranged from 2n = 50 to 2n = 69, and plants with 2n = 40 or 41 were not recovered. This suggests that soybean male and female spores tolerate higher chromosome numbers than those recorded in autotriploid and hyper-triploid of barley. One more back cross is needed to isolate primary trisomics. This study clearly demonstrates that soybean is not a true diploid but is of tetraploid origin.

c. Autotetraploids (4x, AAAA)

i. Sources of Autotetraploids — Autotetraploids (4x) occasionally appear spontaneously in natural diploid (2x) populations by nondisjunction in somatic tissue (meristematic chromosome doubling)

or in reproductive tissues by formation of unreduced gametes (de Wet, 1980). Autotetraploids of spontaneous origin are also found among the progenies of twin seedlings and male steriles or in response to genes causing abnormal meiosis. A classic example of spontaneous origin of an autotetraploid is *Oenothera gigas*, which contained 2n = 28 chromosomes and that was originally considered to be a genic mutant (Lutz, 1907). Furthermore, autotetraploids are successfully induced from temperature shocks, cell and tissue cultures, irradiations, and chemicals (colchicine, certain growth hormones, e.g., naphthalene-acetic acid) (Burnham, 1962). Chromosome doubling occurs in zygotes from fusion of an unreduced egg (2x) and an unreduced sperm (2x), and in somatic tissues, from nondivision, where chromosomes split, but nuclear division fails, resulting in tetraploid tissues.

Colchicine: Of all the above means of obtaining tetraploids (polyploids), colchicine has been found to be the most effective chemical to induce polyploidy in a large number of plant and animal species (Blakeslee and Avery, 1937; Blakeslee, 1939; Eigsti and Dustin, 1955; Burnham, 1962). Colchicine is an alkaloid and is a highly poisonous (carcinogenic) chemical that should not be absorbed by the skin (Blakeslee and Avery, 1937). Colchicine is extracted from the seeds and bulbs of the wild meadow saffron or autumn-flowering *Crocus (Colchicum autumnale* L.) (Eigsti and Dustin, 1955).

Colchicine acts by inhibiting spindle formation and preventing anaphase. Chromosomes stay at the equatorial plate but split longitudinally. Divided chromosomes remain in a single restitution nucleus, doubling the chromosome number. This process will be continued as long as the drug is present. Colchicine is also used as a pretreatment agent for counting somatic chromosomes. Colchicine appears to affect only actively dividing cells. Therefore, colchicine is applied to growing organs of plants, to rapidly germinating seeds, to growing shoots and buds, and axillary nodes. Colchicine is water soluble.

Seed treatment: Colchicine concentrations and treatment durations depend upon the crop species. Blakeslee and Avery (1937) treated rapidly germinating *Datura* seeds with colchicine concentrations ranging from 0.003125% to 1.6% for 10 days. Concentrations up to 0.1% were not effective, but tetraploids were obtained in the higher concentrations (0.2 to 1.6%).

Seed treatment with 0.25% aqueous colchicine was found to be the most effective in producing autotetraploids in chickpea (*Cicer arietinum* L., 2n = 16), but seedling treatment failed (Pundir, Rao, and van der Maesen, 1983). Colchicine-treated seeds and seedlings should be washed thoroughly in running tap water before planting.

Seedling treatment: Actively growing shoots of young seedlings may be immersed in aqueous colchicine solution, or colchicine may be applied to shoots by absorbent cotton or swab. Immersion of roots should be avoided. Sears (1941) obtained amphidiploid sectors from sterile intergeneric hybrids of *Triticinae* by use of colchicine. The crowns and bases of plants were wrapped with absorbent cotton, and the cotton was soaked with 0.5% aqueous solution of colchicine. The plants were transferred to a high-humidity chamber to maintain humidity. Treated plants can be covered with clear plastic bags. Cotton was kept wet for 2 to 5 days by applying colchicine twice a day. Cotton was removed, and plants were transferred to the greenhouse for further growth. He obtained tetraploid sectors in 39 of the 60 surviving plants, representing 17 different hybrids. Stebbins (1949) isolated autotetraploids from 20 different species of the family Poaceae (Gramineae) by treating shoots with 0.1 to 0.2% aqueous solution of colchicine to *Glycine* species hybrids failed to produce amphiploids, but shoot treatment was successful. Jauhar (1970) obtained autotetraploid pearl millet by treating 10-day-old shoots and 2-week-old seedlings with 0.2% aqueous solution of colchicine for 11 to 12 h.

Schank and Knowles (1961) recorded that the most successful colchicine treatment to induce tetraploidy in safflower was a 0.1% aqueous solution applied to a cotton swab wedged between cotyledons of 3-day-old seedlings. The shoot treatment was less successful.

For potato, Kessel and Rowe (1975) attempted ten different colchicine treatments to induce autotetraploids. The success rate was low (Table 7.21). Axillary treatment produced the best results,

Treatment	Methods	Chromosome Doubling (%)
1. Smearing	Lanoline paste containing 1% colchicine was smeared on the eyes of the tuber, and the tubers were allowed to sprout	1.9
2. Dropping I	Absorbent cotton with 1% colchicine was placed on the eyes of the tubers, and cotton was kept moist for 5 days; untreated eyes and sprouts should be removed	1.2
3. Dropping II	Cotton was placed in the bottom of a Petri dish, and tubers with marked eyes were placed on the cotton; cotton was kept moist with 1% colchicine for 5 days	0.0
4. Soaking	The tubers were soaked in a Petri dish containing 1% colchicine for 5 days	0.0
5. Seedling	A drop of lanoline paste containing 1% colchicine was placed on a growing point of approximately 7.5 cm tall seedling	0.7
6. Subaxillary	Potato was grafted onto tomato root stock; after scions were 25 to 30 cm tall, the axillary bud was removed; after 24 h, cotton was wrapped in leaf axils, and cotton was soaked with 1% colchicine; the treated plant was placed in a plastic bag; after 24 h, the bag and cotton were removed	2.8
7. Agar	Seeds were placed in a Petri dish on agar jelly containing 1% colchicine; germinated seeds were washed, and seedlings were transplanted to pots	0.3
8. Colchicine solution	Seeds were germinated on filter paper in a Petri dish soaked in either 0.5 or 0.1% colchicines; duration of treatment was not indicated; seedlings were washed and transplanted to small pots	0.6 (0.5 colchicine); 1.5 (0.1 colchicine)
9. Postgermination 10. Soaking	Germinated seeds were soaked for 3 to 7 days in 1% colchicine Dry seeds were soaked for 3 to 7 days in colchicine containing: 0.5%; 0.25%	0.9 0.4; 0.0

Source: From Kessel, R. and Rowe, P.R., Euphytica, 24, 65–75, 1975. With permission.

though only 2.8% plants were chromosomally doubled. However, the duration of colchicine treatment was 24 h. The longer treatment period (4 to 6 days) could have increased the frequency of tetraploids. Any of these treatments can be used to induce polyploidy in crop plants.

Thiebaut et al. (1979) recommended treating barley haploid seedlings at the three-leaf stage using a solution containing 0.1% colchicine, 2% DMSO, 0.3 mL/L (10 drops) of Twin 20, and 10 mg/L GA3 under 25 to 32°C. Seedlings were treated for 5 h.

Colchicine (0.1 to 0.2%) can be incorporated in liquid or solid media before autoclaving. Actively growing young shoots or axillary buds are exposed to colchicine for 7 to 14 days, then are transferred to colchicine-free medium. Chromosomally doubled shoots are slow growing with dark-green, thick leaves. Lyrene and Perry (1982) preferred colchicine treatment for blueberry in liquid medium on a rotating wheel for 24 h with 0.2% colchicine.

Nitrous Oxide: It has been demonstrated that nitrous oxide is a chromosome doubling agent. Compared to colchicine, it induces a relatively higher frequency of chromosome-doubled plants, causes lesser lethality, and the gas is relatively harmless. Chromosome doubling by nitrous oxide treatment is effective for plants still in tissue culture but is unsuitable for large plants (Hansen et al., 1988).

Taylor et al. (1976) excised heads of diploid red clover containing 2 cm stems 24 h after crossing and placed them in vials containing a 2% aqueous sucrose solution. Vials with heads were placed in a gas-tight chamber, and nitrous oxide was maintained to 6 bars atmospheric pressure (approximately 90 psi). After 24 h, the heads were removed to a dark incubator at 20°C for a seed maturation period of 3 weeks. A total of 226 plants was treated with nitrous oxide, and 160 plants (71%) were identified as putative tetraploid (2n = 4x = 28) based on pollen size. Of 136 plants examined cytologically, 119 plants were tetraploids, three plants were diploid, two plants died, and 12 plants were aneutetraploid (2n = 26, 27, or 29). No plants with chromosomal chimaerism were found. Hansen et al. (1988) compared the effect of nitrous oxide and colchicine on chromosome doubling of anther culture-derived young seedlings of wheat still in culture. Two colchicine treatments (0.005 and 0.01% for 24 h) were compared with two nitrous oxide treatments (24 h and 48 h at 6 atm.). Both nitrous oxide treatments were as effective as 0.01% colchicine; however, colchicine treatment killed a significant proportion of the treated plants, while nitrous oxide treatment was non-toxic. The low concentration of colchicine (0.005%) showed lower chromosome doubling efficiency.

A mixture of 0.1% colchicine and 2.0 or 4.0% DMSO (dimethyl sulfoxide) proved to be more efficient in producing doubled haploids in barley than that observed with colchicine alone or by nitrous oxide treament (Subrahmanyam and Kasha, 1975).

ii. Morphological Characteristics of Autotetraploids — Generally, autotetraploids are slow in growth and exhibit dark-green, large leaves. It is a common conception that autotetraploids usually produce *gigas* phenotype, plants larger than their diploid ancestors (Ramanujam and Parthasarathy, 1953; Kuspira, Bhambhani, and Shimada, 1985). The *gigas* phenotype has been expressed in autotetraploid rye (Müntzing, 1951), pearl millet (Jauhar, 1970; Hanna, Powell, and Burton, 1976), *Sorghum* (Schertz, 1962), safflower (Schank and Knowles, 1961), chickpea (Pundir, Rao, and van der Maesen, 1983), and *Triticum monococcum* (Kuspira, Bhambhani, and Shimada, 1985) and is attributed to the increased sizes of cells, stomata, leaves, pollen, and seed. In self-pollinating species, autotetraploids are either smaller or similar in height to their diploid counterparts.

In *T. monococcum* autotetraploids (2n = 4x = 28), plant height was reduced by 15% compared to diploid sibs. Furthermore, the average reduction of number of tillers per plant was 37.5% (Kuspira, Bhambhani, and Shimada, 1985). Autotetraploids are usually a few days later in flowering than the counterpart diploids.

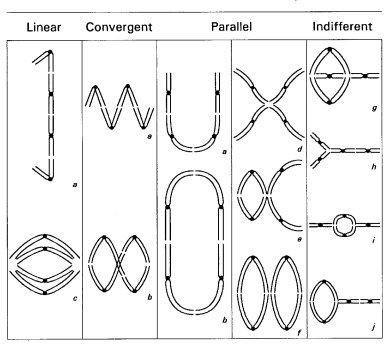
iii. Cytological Behavior in Autotetraploids — In autotetraploids, each chromosome is present four times. Thus, chromosome associations such as quadrivalents, trivalents, bivalents, and univalents are expected based on random association of four homologous chromosomes. Chromosome pairing is usually studied at diakinesis or metaphase-I of meiosis, because chromosomes at these stages are in condensed forms. However, the majority of chiasmata are usually terminalized.

Kuspira, Bhambhani, and Shimada (1985) diagrammed ten possible quadrivalent configurations for each group of the four homologous chromosomes (Figure 7.23). However, they observed only chains (convergent-a; parallel-a) and rings (convergent-b; parallel-b) of four chromosomes (Figure 7.23). Of the 272 quadrivalents examined, chain configurations comprised 70, and ring quadrivalents numbered 202. The open-ring quadrivalent was the highest (197/272). The average chromosome association per cell was 0.62I + 9.86II + 0.2III + 1.74IV.

Table 7.22 compares the frequencies of chromosome configurations observed in autotetraploids of nine species containing basic chromosome number x = 7. The frequencies of bivalents and quadrivalents are different among species and even vary within a species. For example, Morrison and Rajhathy (1960a) recorded a higher frequency of quadrivalents (5.1 per cell) than those recorded (1.74 per cell) by Kuspira, Bhambhani, and Shimada (1985) in *T. monococcum*; these differences may be contributed by genetic factors.

Chromosome migration at anaphase-I in autotetraploids is determined by the coorientation of their kinetochores at metaphase-I. In *T. monococcum* autotetraploids, Kuspira, Bhambhani, and Shimada (1985) recorded 14–14, 16–13 + 1F (false univalent), 14-12 + 2F in 67%, 6.5%, and 1.3% of sporocytes, respectively. Moreover, 15–13, 16–12, and 15–12 + 1F chromosome migrations were observed in 20.0%, 3.5%, and 1.7% of sporocytes, respectively. Based on anaphase-I chromosome movement, 70.90% balanced gametes (n = 14) are expected.

iv. Fertility and Breeding Behavior in Autotetraploids — Autotetraploids are partially pollen and ovule abortive because of unbalanced spore constitution. The synthesized chickpea autotetraploid



Quadrivalent co-orientations at metaphase-I

Figure 7.23 Expected ten types of quadrivalent coorientations at metaphase-I in an autotetraploid. Generally, quadrivalent coorientations convergent a and b and parallel a and b are observed. (Redrawn from Kuspira, J.R., Bhambhani, R.N., and Shimada, T. *Can. J. Genet. Cytol.*, 27, 51–63, 1985.)

Species	Univalents	Bivalents	Trivalents	Quadrivalents	Source
Arrhenatherum elatius	0.0	4.3	0.0	4.8	Morrison and Rajhathy, 1960a
Avena strigosa	0.1	5.0	0.1	4.4	Morrison and Rajhathy, 1960a
Hordeum bulbosum	0.1	5.6	0.2	4.0	Morrison and Rajhathy, 1960a
Hordeum vulgare	0.3	5.7	0.1	3.9	Morrison and Rajhathy, 1960a
Pennisetum glaucum	2.64	8.97	0.38	1.49	Hanna, Powell, and Burton, 1976
Petunia hybrida	0.1	4.6	0.1	4.6	Morrison and Rajhathy, 1960b
Pisum sativum	0.3	5.21	0.16	4.20	Mercy-Kutty and Kumar, 1983
Secale cereale	0.5	6.1	0.2	3.7	Morrison and Rajhathy, 1960a
Triticum monococcum	0.2	3.5	0.1	5.1	Morrison and Rajhathy, 1960a
Triticum monococcum	0.62	9.86	0.23	1.74	Kuspira, Bhambhani, and Shimada, 1985

Table 7.22 Mean Number of Chromosome Configurations Observed per Microsporocyte at Metaphase-I in Tetraploids of Six Species with 2n = 4x = 28

showed 84.4% (range 79.2 to 90.2%) pollen fertility compared to 96.0% in diploids (Pundir, Rao, and van der Maesen, 1983).

Selection for higher fertility is accompanied by a higher frequency of bivalents in autotetraploid maize (Gilles and Randolph, 1951), pearl millet (Gill et al., 1969; Jauhar, 1970), Coix lacryma-jobi (Venkateswarlu and Rao, 1976), and safflower (Schank and Knowles, 1961). Gilles and Randolph (1951) observed fewer quadrivalents (7.46 per cell) and more bivalents in autotetraploids maize at the end of a 10-year period than were present at the beginning of the period (8.47 per cell). They suggested that the original seed stocks had a gene or genes in heterozygous condition for influencing quadrivalent formation. In safflower, Schank and Knowles (1961) examined tetraploids from C_1 to C_3 generations and found a decrease in frequency of quadrivalents and an increase in pollen fertility and seed set. Gill et al. (1969) studied autotetraploid pearl millet for five generations (C_0 to C_5) and observed a reduction in the frequency of multivalents and an increase in the frequency of bivalents. Seed fertility improved through six generations but was still lower in tetraploids compared to diploids. Similarly, Jauhar (1970) observed a shift from multivalent to a bivalent type pairing in autotetraploid pearl millet from the C_0 to C_6 generations. Some of the quadrivalents formed in the C_5 and C_6 generations appeared to be loosely associated. He suggested that structural divergence coupled with genetic factors were the cause of sterility in raw autotetraploids.

Selection for high fertility in autotetraploids is under genetic control (Roseweir and Rees, 1962; Doggett, 1964; Bender and Gaul, 1966; Narasinga Rao and Pantulu, 1982). Selection for high fertility in autotetraploid rye was influenced by increasing the quadrivalent frequency and reducing trivalent frequency, and the bivalent association was under genetic control. However, the high frequency of quadrivalents and high fertility found in rye may not occur in all other tetraploids (Roseweir and Rees, 1962). Doggett (1964) observed variations in seed set (5 to 80%) among individual heads of *Sorghum*, but it was not possible to isolate lines of higher fertility after eight generations; fertility, however, was suggested to be under genetic control.

Morrison and Rajhathy (1960b) contradicted the above findings. They examined ten autotetraploids representing different families and found no evidence of an increase in the numbers of bivalents. Approximately two-thirds of homologous chromosomes formed quadrivalents, and no evidence of genetic control over quadrivalent frequencies was found. Plants with small chromosomes contained as many quadrivalents as plants with large chromosomes. They concluded there is a fairly uniform number of quadrivalents for all species, and that either presentday estimates of chiasmata are out of line or else their frequency has no effect on the number of quadrivalents formed.

Mastenbroek, de Wet, and Lu (1982) studied chromosome associations in early and advanced generations of tetraploid maize (2n = 4x = 40). The increased yield in tetraploid maize selected for 22 years was not due to a change in chromosome associations during diakinesis. It is interesting to note that frequency of quadrivalent and bivalent formation remained constant from generation to generation, and increase in yield was associated with selection against production and functioning of cytologically unbalanced gametes. Selection for fertility was a heritable unit.

Bender and Gaul (1966) suggested that diploidization of autotetraploids could be achieved artificially by an extensive genome reconstruction by means of induced chromosome mutations and gene mutations. Based on these assumptions, Friedt (1978) selected highly fertile lines derived from the hybridization of five autotetraploid barley varieties previously irradiated up to seven times, but the results were negative.

The above results suggest that autotetraploids do not breed true but throw a low frequency of aneuploids in their progenies because of occasional 3:1 disjunction during anaphase-I (Müntzing, 1951; Doyle, 1986).

v. Genetic Segregation in Autotetraploids — The genetics of autotetraploids is quite complex if pursued to any depth. In autotetraploids, every chromosome and gene locus is present in four

copies. Quadrivalent formation and tetrasomic inheritance are the criteria to identify autotetraploids. Multivalents are also formed in reciprocal translocations, but the genetic ratios distinguish autotetraploids from reciprocal translocations.

The number of possible genotypes in a diploid with two alleles is AA, Aa, and aa. On the other hand, the number of genotypes in 4x level is five:

AAAA or A ⁴	Quadruplex
AAAa or A ³ a	Triplex
AAaa or A ² a ²	Duplex
Aaaa or A a ³	Simplex
aaaa or a4	Nulliplex

Methods of calculating theoretical genetic ratios for genotypes triplex (AAAa), duplex (AAaa), and simplex (Aaaa) in random chromosome, random chromatid, and maximum equational segregations have been described in detail by Allard (1960) and Burnham (1962) and are summarized in Table 7.23.

- Only certain ploidy level is beneficial. For example, 4x apple is not desirable, but 3x apples are good, and some of the best varieties of the apple are 3x. Red clover and turnip perform the best at 4x, but fodder beets are generally unfavorable at this level.
- Sterility associated with ploidy may be desirable. Certain floral plants have longer-lasting blooms
 if sterile. Autotriploids may be beneficial if propagated vegetatively. Seedless banana and watermelon are triploid and commercially successful.
- Among cereals, 4*x* rye is successful but may be of outbreeding nature. Fertility in 4*x* is low because of cytological abnormalities that result in genetic unbalance and cross pollination with 2*x* plants. This produces 3*x* seeds that abort prematurely.
- An incompatibility system that successfully maintains cross fertilization at 2x may break down and allow self-fertilization at 4x.

2. Allopolyploidy

a. Origin of Allopolyploidy

An allopolyploid individual is derived by interspecific hybridization from two or more genomically distinct and distantly related diploid species followed by chromosome doubling of sterile F_1 hybrids. Interspecific and intergeneric (wide hybrid) F_1 hybrids are usually sterile, because genomes are highly divergent, and chromosomes lack affinity or form only a small number of loosely synapsed bivalents. This type of synapsis is known as heterogenetic association. In contrast, homogenetic association occurs between chromosomes of the same genome. However, normal chromosome pairing and seed fertility are restored by doubling the chromosomes to a condition known as amphiploidy or amphidiploidy (Stebbins, 1950; Allard, 1960). The majority of allopolyploids are either tetraploids or hexaploids.

Allopolyploidization in nature has played the most important role in the evolution of the crop species wheat, oat, cotton, tobacco, brassicas, and sugar cane. The origin, cytology, and breeding behavior of a few allopolyploids have been described in a chapter on genome analysis.

The classical examples of artificially synthesized allopolyploids are *Raphanobrassica* and Triticale. Karpechenko (1927) produced an intergeneric hybrid between radish (*Raphanus sativus*, 2n = 2x = 18; genome RR) and cabbage (*Brassica oleracea* 2n = 2x = 18; genome BB). Chromosomes of the sterile F₁ hybrid showed absolutely no homologies. Fertility was restored by doubling the chromosomes through the formation and union of unreduced male and female gametes.

		Gametic Types		Phen	Phenotypic Ratios (F ₂)	(F ₂)	Pheno	Phenotypic Ratios (BC ₁)	BC ₁)
Genotype Segregation Chromosome	Iromosome	Chromatid	Maximum Equational Segregation	Chromosome	Chromatid	Maximum Equational Segregation	Chromosome	Chromatid	Maximum Equational Segregation
AAAa 1	1AA + 1Aa	IAA + 1Aa 15AA + 12Aa + 1aa	+ 1aa 13AA + 10Aa + laa	allA	783A:1a	575A:1a	-AIIA-	27A-:1a	23A-:1a
AAaa 1A	4A + 4Aa + 1aa	AA + 4Aa + 3AA + 8Aa + 3aa 1aa	2AA + 5Aa + 2aa	35A:1a	20.8A:1a	77A:4a	5A-:1a	3.7A-:1a	7A-:2a
Aaaa 1	1Aa + 1aa	1Aa + 1aa 1AA + 12Aa + 15aa 1AA + 10Aa + 13aa	1AA + 10Aa + 13aa	3A:1a	2.5A:1a	407A:169a	1A-:1a	0.87A-:1a	11A-:13a

Table 7.23 Expected Gametic Types and E₂ and BC₁ Phenotypic Ratios in Autotetraploids for Random Chromosome, Random Chromatid and Maximum Equational

b. Characteristics of Allopolyploidy

Plant geneticists are optimistic to create a new crop through allopolyploidization. The success of new crops depends upon the following factors:

- It is extremely difficult to predict the nature of interaction between two genomes. The interaction for any trait may be intermediate or heterotic or lethal.
- Interaction between nuclear and cytoplasmic genes may produce cytoplasmic genetic male sterility.
- Economically beneficial traits can be transferred to cultigen from the alien species by hybridization followed by chromosome doubling and backcrossing.

c. Man-Made Crop

Triticale, the first man-made cereal of great potential economic value, are amphiploids of *Triticum* and *Secale*. Concerted cytogenetic and breeding efforts have been made for the past several decades to improve the yield and qualities of triticales, particularly hexaploid (2n = 6x = 42) and octoploid (2n = 8x = 56) cytotypes. Tetraploid (2n = 4x = 28) and decaploid (2n = 10x = 70) triticales also have been studied. Triticales may be primary (original "raw" amphiploids) or secondary (advanced lines derived from crosses between triticales and hexaploid wheats) (Gupta and Priyadarshan, 1985).

Hexaploid triticales (AABBRR) are amphiploids produced from tetraploid (AABB) wheat and diploid (RR) rye. Due to intensive breeding efforts, several secondary hexaploid triticale lines and cultivars have been released. However, cytological instability, reduced seed set, and shriveled seed are a few of the major constraints for the success of triticales. The cytological instability was attributed to tetraploid cytoplasm, allogamous rye, the differences between duration of meiosis, DNA content between rye and wheat, and interaction between the genes of wheat and rye chromosomes.

Octoploid triticales (AABBDDRR) are derived from doubling the chromosomes of F_1 hybrids of hexaploid wheat (AABBDD) and diploid rye (RR). Weimarck (1973) found a lower degree of aneuploidy, more stable meiosis, and higher fertility than with primary triticale. Fertility was not correlated with cytological disturbances. Octoploid triticales contain good winter hardiness, high protein content, good baking quality, and have early flowering and seed maturity and large kernel size. However, seed sterility and early sprouting are undesirable traits that have hampered the popularity of octoploid triticale.

Tetraploid triticales are produced in three ways: (1) by hybridizing *T. monococcum* (AA) with diploid rye (RR) and doubling the chromosomes of the sterile F_1 (AR) to obtain fertile amphiploids (AARR); (2) by isolating tetraploid triticale lines (AARR) after crossing autoallohexaploid wheat (AAAABB) with diploid rye (RR), followed by the elimination of B genome chromosomes; and (3) by hybridizing hexaploid triticale (AABBRR) with diploid rye (RR), and after selfing, to select AARR, BBRR, and (AB) (AB) RR lines. Tetraploid triticales do not have commercial value at present but can be used to improve hexaploid triticale and diploid rye (Krolow, 1973).

Decaploid triticale (AABBDDRRRR) is obtained by crossing hexaploid wheat and tetraploid rye, followed by chromosome doubling. Decaploid triticale is found to be useless because of poor vigor and seed fertility.

II. ANEUPLOIDY: TRISOMICS

A. Primary Trisomics

1. Introduction

An organism containing a normal chromosome complement and one extra chromosome is known as a trisomic. The extra chromosome may be primary, secondary, tertiary, telocentric, acrocentric,

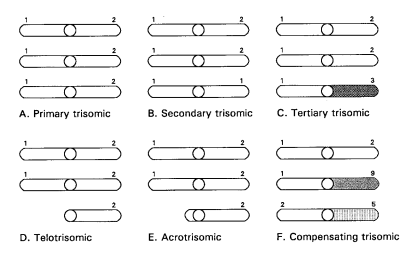


Figure 7.24 Various types of trisomics.

or compensating (Figure 7.24). An individual with a normal chromosome complement plus an extra complete chromosome (2n = 2x + 1) is designated a primary trisomic, and the individual is called Triplo. Blakeslee (1921) coined the term "trisome" for a plant with the addition of one member of the basic chromosome complement, and Bridges (1921) was the first to apply trisomic technique to associate a gene *ey* with the fourth chromosome of *Drosophila melanogaster*. Primary trisomics, since then, have been used extensively for the determination of gene-chromosome-linkage group relationships in several plant species (Khush, 1973; Singh, 1993).

2. Sources of Primary Trisomics

Generally, triploids are considered to be one of the best and most dependable sources for establishing primary trisomic series (Table 7.24). However, primary trisomics have been isolated occasionally from the progenies of normal diploids (spontaneous origin), asynaptic and desynaptic plants, mutagen treated progenies, interchange heterozygotes, and in the progenies of related and unrelated trisomics. The requirement to all sources of primary trisomics is the nondisjunction of chromosomes during meiosis. This results in aneuploid (n + 1, n + 2, n + 3,...) gametes, and when these gametes are fertilized by a male sperm with n chromosome number, aneuploid plants (2n + 1, 2n + 2, 2n + 3,...) are originated.

a. Autotriploids

A complete set of primary trisomics and other aneuploids have been isolated in a majority of diploid species from the progenies of autotriploids (3x) and autotriploid (3x) by diploid (2x) crosses (Tables 7.24 and 7.25). Triploid plants occasionally occur in natural populations by spontaneous origin. They are frequently contrasted from diploids by their taller height, more vigorous growth, and profuse tillering habits. They may show higher sterility. Autotriploid plants are also produced by crossing tetraploid and diploid plants. Generally, autotriploid plants are pollinated by the diploid of the same variety in order to isolate a complete trisomic series in a uniform genetic background (Table 7.25).

Various combinations of chromosome configurations, trivalents, bivalents, and univalents, are observed at diakinesis and metaphase-I in triploids. Chromosomes move at random during anaphase-I, and gametes with chromosome numbers ranging from *n* to 2n are expected at the end of meiosis. Thus, it is possible to obtain plants with 2x to 3x chromosome numbers in the progeny of an autotriploid plant. However, plants with 2x (diploid), 2x + 1 (primary trisomic), 2x + 1 + 1

	Species	2n	Authors
1.	Antirrhinum majus	16	Rudorf-Lauritzen, 1958; Sampson, Hunter, and Bradley, 1961
2.	Arabidopsis thaliana	10	Sears and Lee-Chen, 1970
3.	Avena barbata	14	Nishiyama, 1981
4.	Beta vulgaris	18	Levan, 1942; Kaltsikes and Evans, 1967; Romagosa et al., 1986
5.	Clarkia unguiculata	18	Vasek, 1956
6.	Collinsia heterophylla	14	Garber, 1964
7.	Corchorus olitoris	14	lyer, 1968
8.	Crepis capillaris	6	Babcock and Navashin, 1930
9.	Datura stramonium	24	Blakeslee and Avery, 1938
10.	Fragaria bracteata	14	Yarnell, 1931
11.	Helianthus annuus	34	Jan, Chandler, and Wagner, 1988
12.	Hordeum chilense	14	Cabrera, Remirez, and Martin, 1999
13.	Hordeum spontaneum	14	Tsuchiya, 1960a
14.	Hordeum vulgare	14	Kerber, 1954; Tsuchiya, 1967
15.	Humulus lupulus	18	Haunold, 1970
16.	Lolium perenne	14	Myers, 1944; Meijer and Ahloowalia, 1981
17.	Lotus pedunculatus	10	Chen and Grant, 1968a,b
18.	Lycopersicon esculentum	24	Lesley, 1928, 1932; Rick and Barton, 1954
19.	Lycopersicum peruvianum	24	Sree Ramulu et al., 1977
20.	Medicago sativa	16	Kasha and McLennen, 1967
21.	Nicotiana sylvestris	24	Goodspeed and Avery, 1939
22	Psathyrostachys juncea	14	Wei et al., 1995
23.	Oenothera blandina	14	Catcheside, 1954
24.	Oryza sativa	24	Hu, 1968; Iwata, Omura, and Nakagahra, 1970; Iwata and Omura, 1984; Khush et al., 1984; Cheng et al., 2001
25.	Oryza glaberrima	24	Ishiki, 1991
26.	Pennisetum glaucum	14	Gill, Virmani, and Minocha, 1970
27.	Petunia axillaris	14	Reddi and Padmaja, 1982
28.	Petunia hybrida	14	Smith, de Jong, and Oud, 1975
29.	Secale cereale	14	Kamanoi and Jenkins, 1962; Zeller, Kimber, and Gill, 1977; Melz, Schlegel, and Sybenga, 1988; Fujigaki and Tsuchiya, 1990
30.	Solanum chacoense	24	Lee and Rowe, 1975
31.	Solanum tuberosum	24	Wagenvoort, 1995
32.	Sorghum bicolor	20	Schertz, 1966, 1974
33.	Spinacia oleracea	12	Tabushi, 1958; Janick, Mahoney, and Pfahler, 1959
34.	Triticum monococcum	14	Kuspira et al., 1986
35.	Verbena tenuisecta	10	Arora and Khoshoo, 1969
36.	Zea mays	20	McClintock, 1929c; McClintock and Hill, 1931

(double trisomic), and 2x + 1 + 1 + 1 (triple trisomic) chromosome numbers are frequently obtained in varying frequencies (Table 7.25). These numbers indicate that the maximum number of extra chromosomes tolerated by gametes in most diploid species is one, or two, or rarely three. This limitation may be due to the fact that male and female gametes, or zygotes and embryos, with higher than three extra chromosomes abort in the progenies of triploids of the diploid species, because the duplication of extra genetic material causes genetic and physiological imbalance.

The true nature of a species, whether it is a diploid or a polyploid, can be ascertained by looking at the frequency of aneuploids in progeny from autotriploids. For example, the basic chromosome number of rice is disputed by several rice scientists. Nandi (1936) proposed, based on karyotype analysis, that rice (*Oryza sativa* L.) is a balanced secondary allotetraploid with x = 5 as a basic chromosome number, where A and B chromosomes duplicated after hybridization between putative ancestral species. However, it was evident from the progenies of autotriploids that the maximum number of extra chromosomes tolerated by rice gametes was five, and plants with 2n + 1 and 2n + 2 chromosomes were in the highest frequency (Table 7.26). The narrow

					Frequer	ICIES OF	Plants V	vith Chr	omosor.	Frequencies of Plants with Chromosome Numbers	ers				
													Total Numbers		
Species	u	2x	2x+1	2x+2	2x+3	2 <i>x</i> +4	2x+5	2 <i>x</i> +6	2 <i>x</i> +7	2x+8	3х	Other	Plant	% 2x+1	Authority
Datura stramonium	12	58	138	79	10								285	48.4	Satina, Blakeslee, and Averv. 1938
Hordeum spontaneum	7	29	59	22	9	-					-	œ	126	46.8	Tsuchiya, 1960a
Lolium perenne	7	30	52	27	9	4							119	43.7	Myers, 1944
Lotus pedunculatus	2	126	73	2								-	202	36.1	Chen and Grant, 1968a
Lycopersicon esculentum	12	303	342	131	7						N	14	799	42.8	Rick and Barton, 1954
Nicotiana sylvestris	12	-	17	20	ω	ო	ო		-	-		ო	57	29.8	Goodspeed and Avery, 1939
Oryza sativa	12	0	20	25	14	80	ო						72	27.8	Khush et al., 1984
Oryza glaberrima	12	9	1	б	5	-							32	34.4	Ishiki, 1991
Pennisetum glaucum	~	59	20								-	11	91	22.0	Gill, Virmani, and Minocha, 1970
Secale cereale	7	12	23	12	9	ო						N	58	39.7	Kamanoi and Jenkins, 1962
Spinacia oleracea	9	30	39	14	8	9	15	16					128	30.5	Tabushi, 1958
	9	06	118	23	ი	0	41	43					327	36.1	Janick, Mahoney, and Pfahler, 1959
Triticum monococcum	7	74	59	N									135	43.7	Kuspira et al., 1986

176

Total

(%)

30

29

28

2n 27

26

25

24

Reference Ramanujam, 1937

Katayama, 1963 Hu, 1968

2x + 1

50 21 87 72

18.0 28.5 30.2 40.2 27.8

m o - o o

- o - o

<u>+ ~ ~ ~ ~ </u>

0 - 4 6 4

20 4 2 0 9 25 2 0 9

ь 6 20 а 6

Watanabe et al., 1969 Khush et al., 1984

	From	Autotriploid Pr	ogeny	From Double	
Type of Trisomics	1953 (%)	1954 (%)	Total (%)	and Triple Trisomics (%)	Grand Total (%)
Bush	5 (20.8)	3 (8.6)	8 (13.6)	1 (5.6)	9 (11.7)
Slender	3 (12.5)	8 (22.9)	11 (18.6)	3 (16.7)	14 (18.2)
Pale	6 (25.0)	10 (28.6)	16 (27.2)	3 (16.7)	19 (24.7)
Robust	2 (8.3)	2 (5.7)	4 (6.8)	3 (16.7)	7 (9.1)
Pseudonormal	1 (4.2)	3 (8.6)	4 (6.8)	3 (16.7)	7 (9.1)
Purple	3 (12.5)	6 (17.2)	9 (15.3)	3 (16.7)	12 (15.6)
Semierect	4 (16.6)	3 (8.6)	7 (11.8)	2 (11.1)	9 (11.7)

Table 7.27 Frequency of Seven Types of Primary Trisomics in the Progenies of Autotriploids and Double or Triple Trisomic Plants of Barley

Source: From Tsuchiya, T., Jpn. J. Bot., 17, 177-213, 1960a. With permission.

tolerance limit of the extra chromosomes in gametes shows that rice is truly diploid with x = 12 basic chromosome number. Polyploid species tolerate much higher numbers of extra chromosomes (Ising, 1969; Khush, 1973).

The frequency of occurrence of each trisomic type in progenies of autotriploids varies with types of trisomics and year of planting — above all, it is influenced by the genetic background of an autotriploid. Tsuchiya (1960a) reported in barley that Pale (Triplo 3) trisomic appeared most frequently in each of two years (Table 7.27) followed by Slender (Triplo 2), Purple (Triplo 6), Bush (Triplo 1), and Semierect (Triplo 7). The frequencies of Robust (Triplo 4) and Pseudonormal (Triplo 5) were lowest (6.8%) among 59 primary trisomics (Table 7.27). Similar results have been reported in Datura stramonium (Blakeslee and Avery, 1938), Lycopersicon esculentum (Lesley, 1928, 1932; Rick and Barton, 1954), Nicotiana sylvestris (Goodspeed and Avery, 1939), Zea mays (McClintock and Hill, 1931), Oryza sativa (Hu, 1968), and Beta vulgaris (Romagosa et al., 1986). Chen and Grant (1968a), working with the trisomics of *Lotus pedunculatus*, were unable to isolate Triplo 2. They suggested two possibilities: the first explanation was that Triplo 2 may have been grouped with other types. This outcome is possible, because somatic chromosomes of Lotus pedunculatus are small and do not allow karyotype analysis. Furthermore, pachytene chromosome analysis was not conducted, and genetic tests were not performed. A second possibility is that Triplo 2 may be unviable or less viable than other Triplos. A similar situation was reported in Oenothera (de Vries and Boedijn, 1923) and Arabidopsis thaliana (Steinitz-Sears, 1963).

Isolation of primary trisomics from autotriploids is preferred, because the frequency of aneuploids is extremely high in the progenies of autotriploids compared to such other sources as synaptic mutants, translocation heterozygotes, induced mutations, and normal diploids. Autotriploids generally produce 30 to 80% simple primary trisomic plants, while other sources yield 1 to 3%.

It has been observed that the initial phase of seed development is normal in autotriploids, but the endosperm shrivels after a week, resulting in the death of the embryo. The failure in endosperm development is caused by an extremely unbalanced chromosome number, and this is the most likely explanation for the occurrence of plants with 2x + 1, 2x + 2, 2x + 3 chromosome numbers in the progenies of autotriploids such as barley, rice, maize, tomato, and others (Table 7.25). Rick and Notani (1961) reported that the primitive variety of tomato Red Cherry tolerates a significantly higher frequency of aneuploids than does a large fruited cultivated variety San Marzano. It was also reported that extra chromosomes caused far less anatomical modification in Red Cherry than in a large fruited tomato. The triploids and aueuploids of Red Cherry also had higher fertility. Similar observations were reported in barley, where trisomics of wild barley (*Hordeum spontaneum*) were vigorous and fertile compared to the trisomics of cultivated barley (Tsuchiya, 1960a, 1967). In the cultivated varieties, the gene balance may be upset, as they represent a short time selection of individuals which are superior in one or a few special traits. It has also been suggested that primitive varieties tolerate significantly higher chromosome numbers than cultivated varieties.

b. Synaptic Mutants (Asynaptic and Desynaptic)

A few trisomic plants have been isolated from the progenies of mutant plants. Synaptic mutants show a high frequency of univalents at diakinesis and metaphase-I of meiosis. Their occurrence is due to the disturbance in normal bivalent pairing governed by a homozygous recessive gene. The random movement of chromosomes generates (n + 1) gametes, and when these gametes are fertilized by a haploid male, gamete primary trisomics are produced. Koller (1938) isolated a primary trisomic in asynaptic progenies of *Pisum sativum*. Katayama (1963) obtained 9.2% (21/227) plants with 2n = 25 chromosomes from the progeny of asynaptic rice plants. However, no attempts were made to identify trisomics morphologically, cytologically, or genetically. Palmer (1974) and Palmer and Heer (1976) obtained a few trisomics among progeny of homozygous recessive asynaptic soybean, *Glycine max*. Recently, Xu et al. (2000c) identified 20 possible primary trisomics of the soybean from the aneuploid lines obtained from the progenies of asynaptic and desynaptic and male sterile lines and male sterile induced autotriploids.

Dyck and Rajhathy (1965) isolated six of the seven possible primary trisomics from the progeny of desynaptic *Avena strigosa*. Rajhathy (1975) further reported all the expected seven primary trisomics from the progeny of desynaptic plants. Several back crosses were made to eliminate the desynaptic gene.

Five morphologically distinct primary trisomics of *Vicia faba* were isolated in the progenies of asynaptic mutant and were identified cytologically by translocation tester sets (Barceló and Martin, 1990).

Primary trisomics obtained from the progenies of synaptic mutants are not suitable for use in cytogenetic analysis, because they include sterile plants that cause the release of unrelated trisomics in their progeny (Rick and Barton, 1954).

c. Mutagen Treatment

Primary trisomics have been isolated sporadically from the progenies of plants treated by mutagens. Soriano (1957) isolated trisomics in *Collinsia heterophylla* by colchicine treatment. However, trisomics did not express contrasting morphological traits (Dhillon and Garber, 1960; Garber, 1964). Their independence was not tested cytologically and genetically. Therefore, it is likely that not all the possible (seven) trisomics were induced. Parthasarathy (1938) accidentally obtained a trisomic plant from the progenies of x-irradiated rice seeds. Martin (1978) isolated four primary trisomics and two double trisomics from the progeny of an x-ray induced mutant in *Vicia faba*. Simeone, Blanco, and Giorgi (1985) produced 14 possible primary trisomics in *Durum* wheat from the progenies of mutagen (x-ray, thermal, or fast neutrons) treated seeds; only transmission of the extra chromosome was reported. Thakare, Joshua, and Rao (1974) isolated five morphologically distinct primary trisomics by irradiating seeds of *Corchorus olitorius* cv. JR063 (2n = 14) with gamma rays and thermal neutrons. These primary trisomics were produced from chromosomal interchanges. One trisomic was secondary for Chromosome 5, and another was tertiary for Chromosome 2.

Chemical irradiation treatment disturbs cell division, leading to nondisjunction of chromosomes during anaphase-I. This treatment may produce female gametes with n + 1 chromosome constitution and may give rise to a trisomic plant after fertilization with an n chromosome sperm. However, it is possible that trisomics isolated from mutagen treatment are not always true primary trisomic types but may, in some cases, include tertiary trisomics.

d. Normal Diploids

Earlier reports on the occurrence of primary trisomics were based mainly on their occasional isolation from the progenies of normal diploids: *Datura* (Blakeslee, 1924); *Matthiola* (Frost and Mann, 1924); *Crepis* (Babcock and Navashin, 1930); *Secale* (Takagi, 1935); *Hordeum* (Smith, 1941); *Lycopersicon* (Rick, 1945); and *Nicotiana* (Goodspeed and Avery, 1939). In diploid (2n =

24) progenies of marigold (*Tagetes erecta*), Lin and Chen (1981) isolated trisomic plants. The occurrence of primary trisomics from diploid progenies is possible if one bivalent fails to move to the metaphase plate and is included in one telophase nucleus. Trisomic plants are obtained when n + 1 gametes produced in this way are fertilized by normal sperm. Belling and Blakeslee (1924) demonstrated cytologically eight cases of 11–13 chromosome separation in 1137 PMC of normal disomic *Datura* (0.4% n + 1 pollen grains).

e. Other Sources

Another good source of primary trisomics is among the progenies of double trisomics or multiple trisomics. Sometimes, they are an excellent source for obtaining missing trisomics. Goodspeed and Avery (1941) found the missing Trisomic 12 in the progeny of a multiple trisomic, while the other 11 primary trisomics of *Nicotiana sylvestris* were isolated from other sources. Tsuchiya (1960a) obtained all seven primary trisomics from double and triple trisomics in barley. The frequency varied with the trisomic types (Table 7.27). The double and triple trisomics are frequently obtained as siblings of primary trisomics in the progenies of autotriploids. They are relatively vigorous and more or less seed fertile.

Sometimes unrelated primary trisomics are obtained from the progenies of primary trisomics. They have been recorded in *Lycopersicon esculentum* (Lesley, 1928), *Datura stramonium* (Blakeslee and Avery, 1938), *Nicotiana sylvestris* (Goodspeed and Avery, 1939) and *Lotus pedunculatus* (Chen and Grant, 1968b). No unrelated primary trisomics were isolated in barley (Tsuchiya, 1960a, 1967). Avery, Satina, and Rietsema (1959) suggested from their studies of *Datura* that the presence of an extra chromosome encourages in some way nondisjunction of other chromosome sets. However, such events are rare, and one should not depend on this source to isolate primary trisomics.

Occasionally, progenies of interchange heterozygotes produce primary trisomics due to a 3–1 chromosome disjunction at anaphase-I from a noncooriented quadrivalent. The frequency of primary trisomics ranged from 0.94 to 5.65% (Table 7.28).

The progenies of secondary, tertiary, and telotrisomics throw primary trisomics with a low frequency, and the trisomic types are those to which secondary, tertiary, or telotrisomics belong. This outcome happens when secondary, tertiary, or telocentric chromosomes disjoin from a trivalent and move to one pole, while two normal chromosomes move to the opposite pole. Gametes with a complete normal extra chromosome are generated, and after fertilization with normal haploid sperm, primary trisomics result. Singh and Tsuchiya (1977) isolated as high as 1.5% plants with 2n = 15 chromosomes in the progenies of seven monotelotrisomics of barley. In general, aneupoid plants of hybrid origin have a heterotic effect, which may give them a better tolerance against numerical imbalance.

3. Identification of Primary Trisomics

In general, each of the primary trisomics in a diploid species differs from its normal diploid sibs and also from each other in several distinctive traits. The differences are morphological, anatomical,

of Barle	У		
Interchange	Number of Plants	Number of Trisomics	Percentage Trisomics
a + b	1008	26	2.58
b + c	1708	16	0.94
b + d	1380	53	3.84
b + g	1290	59	4.57
e + f	1629	92	5.65

Table 7.28 Frequency of Trisomics from Interchanged Heterozygotes of Barley

Source: From Ramage, R.T. and Day, A.D., Agron. J., 52, 590–591, 1960. With permission.

cytological, physiological, or genetic. Each chromosome carries distinctive genes that affect qualitative as well as quantitative traits responsible for plant growth, vigor, and development. The modifications in expression are such that each trisomic type is distinct and easily distinguishable.

a. Morphological Identification

The identification of a complete primary trisomic series established in *Datura*, maize, barley, diploid oat, pearl millet, rice, *Arabidopsis*, tomato, and several other crop species is based on morphological features such as growth habit, plant height, degree of branching and tillering, leaf size, leaf shape, color and texture of leaf surface, internode length, days to flower, seed fertility, and many other visible morphological traits.

The main diagnostic feature of a plant with 2n = 2x + 1 chromosome constitution is that plant shows slower growth habit than its diploid sibs. Primary trisomic plants can be distinguished fairly easily from normal diploids at seedling, maximum tillering, flowering, or maturity stages. The nomenclature of a trisomic type is based on a most distinctive morphological feature. The 12 trisomic types of *Datura*, a classical example, were named according to the shape and size of *capsules*. The extra chromosome that modified the phenotypic appearance of the plant also contained a factor responsible for the change in capsule size and shape: Globe — shortening and widening the capsule; Cocklebur — narrow capsule; Echinus — long spines; Glossy — shiny surface; microcarpic — downy surface; and so on (Figure 7.25).

The seven primary trisomics of barley were classified into seven separate types based on distinct, easily noticeable morphological features of different trisomic plants: Bush — bushy growth habit; Slender — slender appearance; Pale — pale color leaves; Robust — vigorous growth habit; Pseudonormal — similar to diploid; Purple — dark purple color in leaf sheaths; and Semierect — semierect growth habit. Five (Bush, Slender, Pale, Pseudonormal, Semierect) of the seven primary trisomics are readily distinguishable from each other as well as from diploids at an early seedling stage, because they exhibit distinguishing seedling traits (Tsuchiya, 1960a, 1967). The remaining two types, Robust and Purple, are easier to identify in the time span of tillering to heading.

The primary trisomics of pearl millet were significantly shorter in plant height, later to flower, and narrower in leaf width than their diploid sibs. They were designated Tiny, Dark green, Lax, Slender, Spindle, Broad, and Pseudonormal. Tiny was weak in growth and vigor, while Pseudonormal was similar to diploid sibs (Gill, Virmani, and Minocha, 1970).

Sears and Lee-Chen (1970) identified four of the five primary trisomics of *Arabidopsis thaliana* based on their easily identifiable morphological appearance of leaves: Concave (C), Round (R), Yellow (Y), and Narrow (N). The fifth trisomic type, Fragilis (F), was identified as early maturing with protruding stigmas. It is self-sterile, because pollen is shed before the stigma becomes receptive.

Primary trisomics have been named in maize (Rhoades and McClintock, 1935), tomato (Rick and Barton, 1954), rice (Khush et al., 1984), and soybean (Xu et al., 2000c), based upon the length of extra chromosome at pachynema. For example, in rice, 12 possible primary trisomics are expected. Thus, Triplo 1 carries the longest and Triplo 12 the smallest extra chromosome. The same nomenclature is employed in maize, tomato, and soybean.

i. Effect of the Length of Extra Chromosome on Plant Morphology — The degree of growth, vigor, and development of a trisomic plant depends roughly upon the length of extra chromosome present in the trisomic condition. The trisomics in which the extra chromosome is long are distinct morphologically at earlier stages of plant growth than those primary trisomics carrying a short extra chromosome. This relationship has been observed in maize (McClintock, 1929c), tomato (Rick and Barton, 1954), barley (Tsuchiya, 1960a, 1967), pearl millet (Gill, Virmani, and Minocha, 1970), rice (Khush et al., 1984), and in many other species (Khush, 1973; Singh 1993). However, there are some exceptions. Triplo 7 and Triplo 8 of tomato are comparatively weak, while Triplo 3 and 6 (longer chromosomes) are relatively vigorous (Rick and Barton, 1954).

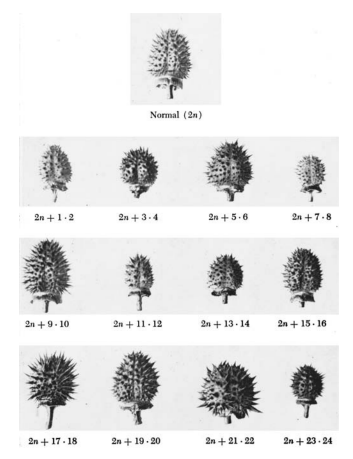


Figure 7.25 Capsules of diploid (2*n* = 24) and 12 possible primary trisomics (2*x* + 1) in *D. stramonium*. (From Avery, A.G., Satina, S., and Rietsema, J., *Blakeslee: The Genus Datura*, Ronald Press, New York, 1959. With permission.)

The soybean contains 2n = 40 chromosomes and always express diploid-like meiosis. Of the 20 primary trisomics, 17 did not express obvious differences in morphology from the their disomic sib. Triplo 1 showed the largest leaves, pods, and seeds among the 20 primary trisomics, and they were about one-third larger than those of the disomic sib. Gray-colored saddle seeds in Triplo 17 distinguished it from the disomic and other primary trisomics (Xu et al., 2000c).

Phenotypic changes in trisomic plant organs have been explained on the basis of Bridges' (1922) gene balance theory. This theory assumes that the genetic complement of a diploid permits an individual to develop and function as an integrated organism. However, when an individual chromosome is added to the normal complement of a diploid species, the gene balance is greatly disturbed. Such imbalance is reflected in physiological, morphological, and developmental deviations. Each chromosome affects the anatomy, physiology, and morphology of the plant in a distinctive way, reflecting the differential gene content of the different chromosomes. The gene balance theory was formulated by Bridges to explain the *Datura* trisomics. Sinnott, Houghtaling, and Blakeslee (1934) supported the theory of Bridges and found a good fit between means of all trisomics and disomics for most of the characters studied. However, Sampson, Hunter, and Bradley (1961) and Rajhathy (1975) pointed out that the gene balance theory does not consider additive and optimum gene action as well as epistatic and compensating genes located on the other chromosomes.

The imbalance an extra chromosome exerts over the balanced condition may best be shown by capsules of the Globe trisomic of *Datura*. Blakeslee and Belling (1924b) observed that the Globe

trisomic with two extra chromosomes (2n = 2x + 2) has a greater imbalance (2/24) compared with its respective primary trisomic (2n = 2x + 1). In tetrasomic condition, the depression of capsules is severe in expression. Similarly, the tetraploid Globes have their capsules relatively more depressed and their spines relatively stouter as we pass from 4x to 4x + 2 and 4x + 3 chromosome constitutions. A 4x + 4 Globe was also expected but presumably was not isolated because the imbalance from the extra chromosomes would be 4/48. This is the same imbalance found in 2x + 1 Globe. It is, therefore, evident that the tolerance limit of imbalance in *Datura* is only two extra chromosomes to its diploid chromosome complement. Tetrasomic (2n = 2x + 2 = 44) soybean also exerts substantial alteration in morphology compared to those recorded in primary trisomics. All the available tetrasomics in the soybean are viable, are slow in vegetative growth, have altered morphology, and are partial fertile (Singh and Xu, unpublished results).

ii. Effect of a Nucleolus Organizer Chromosome on Plant Morphology — It is interesting to note that Triplo 9 of rice, which is a principal nucleolus organizer chromosome, has stoutness in some plant characters such as thick, dark-green leaves and culms, along with a large panicle with bold grains. Triplo 10, a weak nucleolus organizer, is not distinguishable from diploids at early plant growth stages but has small grains (Khush et al., 1984). Triplo 6 and Triplo 7 of barley carry extra nucleolus organizer chromosomes and are vigorous (Tsuchiya, 1960a, 1967). In maize, Triplo 6 is not distinguishable from disomic sib on gross plant phenotype (Rhoades and McClintock, 1935). Similarly, in rye (Kamanoi and Jenkins, 1962) and also in pearl millet (Gill, Virmani, and Minocha, 1970), plants trisomic for a nucleolus organizer chromosome are pseudonormal. Rick and Barton (1954) observed in tomato that Triplo 2, in which the extra chromosome is the second longest as well as a nucleolus organizer chromosome, has straggly growth habit, a large terminal leaf segment, and nearly normal leaf color and flowers. The nucleolus organizer chromosome of moss verbena when in trisomic condition increases the leaf and flower size (Arora and Khoshoo, 1969). By contrast, Triplo 13 of the soybean showed shorter nodes, dark green leaves, and smaller pods and seeds than disomic (Xu et al., 2000c).

iii. Effect of Genetic Background — The main morphological distinguishing features are similar for each trisomic type established in several cultivated varieties of barley (Kerber, 1954; Ramage, 1955; Tsuchiya, 1967) and are almost like those of wild barley, Hordeum spontaneum (Tsuchiya, 1960a). Similar observations have been reported in rice, where trisomic sets have been established in indica and japonica rices. The extra chromosome exerts the same modification in morphological traits, though different trisomic names were given depending upon observations of the authors (Hu, 1968; Iwata, Omura, and Nakagahra, 1970; Iwata and Omura, 1984; Khush et al., 1984). From these results, it is suggested that trisomic plants have identical or close distinguishing characters regardless of the genetic background, provided detailed and close observations are made on many plant organs throughout the entire growing period, from early seedling to maturity. However, some primary trisomics established from other than autotriploid sources do not express clear diagnostic traits in Clarkia unguiculata (Vasek, 1956, 1963), Collinsia heterophylla (Dhillon and Garber, 1960; Garber, 1964), Solanum chacoense (Lee and Rowe, 1975), and Lycopersicum peruvianum (Sree Ramulu et al., 1977). Primary trisomics in these cases were not distinguishable morphologically from each other or from disomic sib but were vigorous and fertile. Furthermore, even multiple trisomics isolated from interspecific hybrids of Solanum did not differ morphologically from diploid sibs. This is probably due to the genetic variability present in the material (Lee, Kessel, and Rowe, 1972).

The primary trisomic series established in barley was partly similar morphologically but different in seed fertility (Tsuchiya, 1960a, 1967; Ramage, 1955, 1960). These differences may be attributed mainly to their sources of isolation. Tsuchiya (1960a, 1967) obtained his trisomics from the progenies of autotriploids, while Ramage's trisomic set was isolated from translocation heterozygotes. The high sterility in Ramage's trisomic series is attributed to chromosomal abnormalities, originated from x-ray-induced segmental interchanges. These may have introduced disadvantageous changes into the chromosomes and genes.

b. Cytological Identification

The true nature of a trisomic plant, identified morphologically, is confirmed by cytological observations. Identification of the extra chromosome in trisomic plants is based first on somatic chromosome count and is verified by the analysis of meiotic stages, such as pachynema, diakinesis, metaphase-I, anaphase-I, or telophase-I. At diakinesis or at metaphase-I, a trivalent plus bivalents (III + II) or an univalent plus a bivalents (I + II) chromosome configuration predominates, suggesting the presence of an extra chromosome.

Somatic metaphase chromosomes have been utilized to some extent to identify primary trisomics of barley (Tsuchiya, 1960a, 1967), moss verbena (Arora and Khoshoo, 1969), diploid oat (Rajhathy, 1975), rye (Zeller, Kimber, and Gill, 1977; Fujigaki and Tsuchiya, 1990), and Triticum monococcum (Friebe et al., 1990b). The smallest chromosome number 5 of the five nonsatellited chromosomes of the barley complement is carried by trisome Pseudonormal. The nucleolus organizer Chromosomes 6 and 7 were easily identified from karyotype analysis, as they exhibited distinctive morphology. Purple primary trisomic of barley contains three Chromosomes 6 with large satellite (Figure 7.26). Moreover, it has been difficult to identify Chromosomes 1 through 4 in barley by standard staining methods, because they are similar in length and also lack morphological landmarks. Hence, cytological identification of primary trisomics Bush, Slender, Pale, and Robust is difficult. However, in this situation, identification of the extra chromosome is possible by Giemsa C- and N-banding techniques (Singh and Tsuchiya, 1981b, 1982a,b). The application of the Giemsa C-banding method helped Zeller, Kimber, and Gill (1977) to identify six of the seven possible primary trisomics of Secale cereale. After Giemsa staining, chromosomes of S. cereale exhibit telomeric heterochromatin and intercalary euchromatin differentiation. Such differentiation is not feasible by aceto-carmine or Feulgen staining techniques used routinely. Three nucleolus organizer chromosomes in soybean Triplo 13 were frequently observed by Xu et al. (2000c) at mitotic metaphase by Feulgen staining, while other chromosomes were indistinguishable.

Pachytene chromosome analysis has played a major role in identifying individual chromosomes in trisomic condition in maize (Rhoades and McClintock, 1935), tomato (Rick and Barton, 1954; Rick et al., 1964), and rice (Khush et al., 1984). Centromere positions in tomato have been precisely located, because centromeric regions are flanked by heterochromatin. Such differentiation helped to identify individual chromosomes by their relative length (Barton, 1950). The presence of heterochromatic knobs at specific positions on certain chromosomes of maize facilitated an easy morphological identification (Rhoades and McClintock, 1935).



Figure 7.26 Mitotic metaphase chromosome of a primary trisomic (2*n* = 41) for Triplo 6 (Purple) of barley showing three (arrows) nucleolus organizer chromosomes. (From Tsuchiya, T., *Jpn. J. Bot.*, 17, 177–213, 1960a. With permission.)



Figure 7.27 A trivalent configuration of Triplo 9 in rice at pachynema associated with the nucleolus. Arrow shows extra Chromosome 9 partially paired nonhomologously. (From Khush, G.S. et al., *Genetics*, 107, 141–163, 1984. With permission.)

The three homologous chromosomes in primary trisomics compete to pair with one another, but only two of the three homologues are synapsed in a normal fashion at a given position to form a bivalent. The third attempts to associate with the paired homologues in a random manner and may form a loose trivalent configuration or pair itself (Figure 7.27). When an unpaired chromosome does not have the opportunity for two-by-two association as proposed by McClintock (1932), the unpaired chromosome frequently folds back, showing nonhomologous chromosome association. Nonhomologous chromosome association (long arm paired with short arm) has been frequently observed at pachynema in primary trisomics of maize (McClintock, 1932; McClintock and Hill, 1931; Rhoades, 1940; Rhoades and McClintock, 1935), tomato (Sen, 1952; Rick and Barton, 1954; Rick, Dempsey, and Khush, 1964), *Sorghum* (Poon and Wu, 1967; Venkateswarlu and Reddi, 1968), *Solanum* (Vogt and Rowe, 1968; Ramanna and Wagenvoort, 1976; Wagenvoort, 1995), rice (Khush et al., 1984), and soybean (Xu et al., 2000c). Such nonhomologous pairing could be misinterpreted as a secondary trisomic chromosome in the case of a chromosome with a median kinetochore. However, close observation of the several pachytene trivalents for the mode of chromosome pairing leaves no doubt for correct identification.

At pachynema, some chromosomes, particularly long ones, show a chance association with the nucleolus. The analysis of several well-spread cells of Triplo 9 and Triplo 10 of rice indicated that Chromosome 9 is the main nucleolus organizer (Figure 7.27), and Chromosome 10 is a weak nucleolus organizer. In barley, Chromosome 6 is the main nucleolus organizer, while Chromosome 7, with a smaller satellite than Chromosome 6, has weak nucleolus organizer ability.

In general, trivalent + bivalents (Figure 7.28A) or univalent + bivalents (Figure 7.28B) chromosome associations are observed at diakinesis or metaphase-I in primary trisomics established in several diploid plant species. The frequency varies from crop to crop, stage of meiosis, and also among primary trisomic type. In barley, the range of trivalent association (1III + 6II) at metaphase-I in seven primary trisomics was 63.1% (Pseudonormal) to 78.6% (Slender), averaging 75.9%. The trivalent configuration was higher at diakinesis, averaging 89.1% (Table 7.29). Sears and Lee-Chen (1970), working with *Arabidopsis* primary trisomics, also observed regular trivalent formation (75%) at metaphase-I. It is expected to record about 66% trivalent configuration in primary trisomics if one chiasmata is formed in each arm.

The types of trivalents at metaphase-I differ in their frequencies from those recorded at diakinesis. The frequency of the chain type increases at metaphase-I, while the frying-pan and other complicated types are significantly reduced (Table 7.29).

It has been generally considered that there is a good correlation between chromosome length and trivalent formation — the longer the chromosome, the higher the frequency of sporocytes with

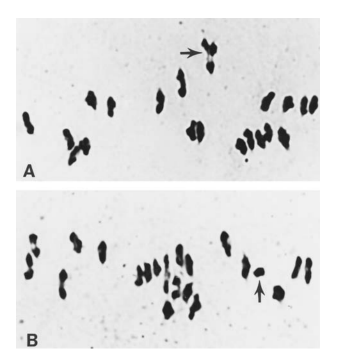


Figure 7.28 Meiotic chromosome configurations at metaphase-I in Triplo 20 of the soybean. (A) A cell with 1III (arrow) + 19II. (B) A cell with 20II + 1I (arrow). (From Xu, S.J. et al., *Crop Sci.*, 40, 1543–1551, 2000c. With permission.)

	С	hromosome	Associatio	ons		Types of	Trivalents	
Types of						Frying-		Triple-
Trisomics	Stage	1III + 6II	7ll + 1l	Others	Chain	Pan	Ү-Туре	Arc
Bush	DK	82.9	17.1	_	27.6	60.3	0	12.1
	MI	70.7	28.7	0.7	67.3	31.8	0.9	0
Slender	DK	89.3	10.7	_	53.1	43.2	1.1	2.7
	MI	78.6	20.0	1.4	69.2	29.8	0.4	0.6
Pale	DK	94.3	5.7	_	40.8	56.0	0	1.0
	MI	76.0	23.8	0.2	62.4	36.1	0.9	0.6
Robust	DK	92.9	7.1	_	46.2	48.7	0	5.1
	MI	77.7	21.9	0.4	64.9	33.0	1.1	1.1
Pseudonormal	DK	79.5	20.5	_	49.7	47.1	0.6	2.5
	MI	63.1	36.5	0.4	72.0	25.6	1.8	0.6
Purple	DK	89.0	11.0	_	41.6	53.9	0	4.5
	MI	77.7	22.3	0.1	60.2	37.8	0.5	0.1
Semierect	DK	93.9	6.1	_	20.3	70.7	0.8	8.1
	MI	71.7	28.2	0.2	47.1	46.7	5.7	0.4
Average	DK	89.1	10.9	_	40.6	54.1	0.4	4.9
-	MI	75.9	23.7	0.5	62.5	35.5	1.3	0.02

Table 7.29 Chromosome Associations and Types of Trivalents at Diakinesis (DK) and Metaphase-I (M	VII)
in Seven Trisomic Types of Barley (%)	

Source: From Tsuchiya, T., Jpn. J. Bot., 17, 177–213, 1960a. With permission.

trivalent configuration. It has been observed in maize (Einset, 1943), tomato (Rick and Barton, 1954), and also in *Lotus pedunculatus* (Chen and Grant, 1968a), that a long extra chromosome has a greater opportunity of forming chiasmata with its homologue than does a shorter one. This expectation can be well documented when we compare frequencies of trivalent association among barley, maize, and tomato; barley has the longest and tomato has the shortest chromosomes. The mean of sporocytes with trivalent configuration at diakinesis for primary trisomics of barley was 89.1% (Tsuchiya, 1960a) and for tomato 48.2% (Rick and Barton, 1954). A similar trend was not observed in primary trisomics of *Avena strigosa* (Rajhathy, 1975), where only 41.3% of the sporocytes showed trivalent configuration, though the karyotype of *Avena strigosa* is similar to the karyotype of barley. These differences may be attributed to the source of trisomics. *Avena strigosa* primary trisomics. Although *Avena strigosa* trisomics were grown several generations, it is likely that they may not yet be completely homozygous (Rajhathy, 1975).

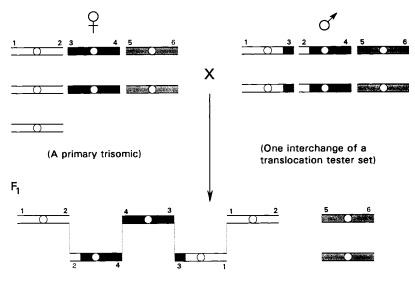
The disjunction of chromosomes in primary trisomics is nonrandom. Generally, two homologues (n + 1) disjoin from a trivalent and move to one pole, while the third homologue goes to the opposite pole in a majority of sporocytes. When an extra chromosome remains an univalent, it may either be included in one of the daughter nuclei, or it may lag behind and fail to be included in either telophase nucleus. At late telophase-I, the lagging chromosome divides equationally, and sister chromatids pass toward opposite poles. However, rarely do both reach these poles. When they reach the poles on time, they are included in the respective daughter nuclei. Otherwise, each may form a micronucleus. The lagging chromosome is sometimes located at the equatorial region or near one of the telophase nuclei. If present elsewhere in the cytoplasm, it is often eliminated. The second division is generally normal. If micronuclei are not lost permanently, they may be included in one of the daughter nuclei. The frequency of univalents may not provide an accurate picture for estimates on the transmission rate of an extra chromosome.

Goodspeed and Avery (1939), working with *Nicotiana sylvestris* primary trisomics, observed no apparent difference in trivalent frequencies among four different primaries. And, they did not find length differences in the extra chromosomes or significant differences in transmission frequencies of these four primaries. In contrast, Einset (1943) reported in maize that when a short chromosome is present in triplicate, fewer trivalents and more univalents are present at metaphase-I. Similar observations were recorded in triploid *Hyacinths* (Belling, 1925) and *Tulipa* (Newton and Darlington, 1929; Darlington and Mather, 1932), where shorter chromosomes form fewer chiasmata at metaphase-I than do longer ones.

c. Identification of Primary Trisomics with Translocation Testers

In most diploid species, simple primary trisomics have been distinguished among themselves and also from diploid sib by morphological appearances and to some extent by karyomorphology. The independence of primary trisomics may also be verified by translocation tester sets. Tsuchiya (1961) used Burnham's translocation tester sets which marked all seven chromosomes of barley (Burnham, 1956; Burnham and Hagberg, 1956) to test mainly the independence of trisomics for Chromosomes 1, 2, 3, and 4. These chromosomes are difficult to distinguish cytologically, because they are similar in karyotype.

Generally, trisomic plants are used as female parents and are pollinated by the translocation testers, because pollen with an extra chromosome does not compete successfully with normal pollen grains in fertilization. Chromosome pairing in F_1 trisomics is analyzed at diakinesis or at metaphase-I. If the extra chromosome of a certain trisomic type is partly homologous to one of the interchanged chromosomes in the testers, a chain of five chromosomes (pentavalent) plus bivalent (Figures 7.29 and 7.30), or the derivative types are obtained. On the other hand, if the extra chromosome is not partly homologous with either of the interchanged chromosomes, a quadrivalent plus a trivalent plus bivalents are observed.



[1 pentavalent (V) + bivalents]

Figure 7.29 Procedure to identify the extra chromosome in primary trisomics by translocation tester sets.



Figure 7.30 A meiotic metaphase-I chromosome configuration of 1V (arrow) + 4II in an F₁ hybrid between primary trisomic Bush and interchange suggests that the extra chromosome in Bush is for Chromosome 1. (From Singh, R.J., unpublished results.)

4. Transmission of the Extra Chromosome in Primary Trisomics

a. Female Transmission

Theoretically, about 50% primary trisomic plants are expected in the progenies of primary trisomics crossed as females. However, it is rarely observed. The transmission rate of the extra chromosome depends upon such factors as the meiotic behavior of the extra chromosome, seed size, genetic background, growing conditions, types of primary trisomics, and number of seeds studied.

It has been established in maize that trisomics for long chromosomes (2, 3, 5) transmit at higher frequencies of primary trisomics (an average of 48%) than the trisomics for the medium (6, 7, 8; average 37%) and short (9, 10; average 25%) chromosomes (Einset, 1943). A positive relationship was observed between transmission rate of an extra chromosome and the frequency of trivalents at metaphase-I. In maize, a higher frequency of trivalents was observed in trisomics for long chromosomes than for shorter ones (Einset, 1943). This observation indicates that the extra chromosome remains as an univalent in the majority of the sporocytes in trisomics for the shorter chromosomes. The univalent lags in meiotic divisions and eventually gets eliminated, yielding gametes without an extra chromosome.

,				
	Hordeum sp	oontaneumª	Hordeum vul	gare S.E. 16 ^b
Types of Trisomics	2x + 1 Selfed	$(2x+1) \times 2x$	2x + 1 Selfed	$(2x=1)\times 2x$
Bush	31.4	24.9	12.5	_
Slender	21.2	27.5	25.7	28.3
Pale	21.0	27.2	25.5	13.3
Robust	28.8	11.3	36.7	19.5
Pseudonormal	24.6	9.8	27.9	31.1
Purple	19.3	17.4	25.9	23.4
Semierect	23.4	24.5	33.3	21.9
Average	25.9	22.7	29.1	20.4

Table 7.30 Frequency of Primary Trisomics in Progenies of Two Primary Trisomic Sets of Barley (%)

^a Tsuchiya, T., Jpn. J. Bot., 17, 177–213, 1960a. With permission.

^b Tsuchiya, T., Can. J. Genet. Cytol., 9, 667–682, 1967. With permission.

The above relationship, however, was not observed in primary trisomics of *Datura*, barley, tomato, rice, diploid oat, or always even in maize. McClintock and Hill (1931) reported a transmission rate of Chromosome 10 of maize to be 33.06%, but Einset (1943) observed only 28%. Rhoades (1933a) obtained 31% female transmission in Trisome 5, but a much higher transmission rate (52%) was reported by Einset (1943). This discrepancy was explained as due to environmental effects or possibly to genic effects (Einset, 1943).

In *Datura*, the smallest trisome showed a transmission rate of 32.68%, while the third smallest transmitted only at a frequency of 2.99% (Blakeslee and Avery, 1938). In barley, Chromosome 5 is the shortest chromosome (Tjio and Hagberg, 1951) and is carried by trisomic Pseudonormal. It showed the lowest female transmission (9.8%) rate in *H. spontaneum* genetic background, but it had the highest transmission rate (31.1%) for the same trisome in *H. vulgare* (Table 7.30). No relationship was established between female transmission and length of the extra chromosome in the primary trisomics of the potato (Wagenvoort, 1995) and soybean (Xu et al., 2000c). An average (range) female transmission of the extra chromosome in 20 primary trisomics of the soybean was 41.6% [27% (Triplo 20) — 59% (Triplo 9)]. Primary trisomics of the soybean showed higher female transmission than those recorded for other diploid species (Xu et al., 2000c). This again suggests that soybean is of tetraploid origin. This concludes that there is no consistent relationship between the length of chromosome and female transmission rate of an extra chromosome in the primary trisomic progenies.

Generally, transmission of an extra chromosome is somewhat higher in selfed or hybrid progenies of primary trisomics (Goodspeed and Avery, 1939; Tsuchiya, 1960a, 1967; Rick and Barton, 1954; Liang, 1979; Lin and Coe, 1986; Curtis and Doyle, 1992). Tsuchiya (1960a, 1967) reported an average of 25.9% and 29.1% trisomic plants in selfed progenies of wild (*H. spontaneum*) and cultivated (*H. vulgae*) trisomic series, respectively. Sears and Lee-Chen (1970) reported a 25.8% transmission rate of the extra chromosome in selfed progenies of five primary trisomics of *Arabidopsis*. The range was 21 to 30%. Kaltsikes and Evans (1967) studied three trisomics of *Beta vulgaris* and reported an overall 22.07% transmission rate of the extra chromosome. Vogt and Rowe (1968) observed 20 to 24% female transmission in three trisomics of potato.

It has been observed that the occurrence of a higher frequency of trivalents in the sporocytes of primary trisomics for the longer chromosomes is due to the fact that long chromosomes have more physical opportunity to pair with homologues than do shorter chromosomes. It is therefore expected that higher female transmission rates for trisomics of the longer chromosomes will be found, because gametes with n and n + 1 chromosome numbers are expected to be formed almost in equal (50% each) numbers. However, such a relationship has been observed only in maize primary trisomics by Einset (1943). In spite of the equal frequencies of female gametes with n and n + 1 chromosome and result in increased gametic and zygotic lethality. Gametic competition among megaspores has also been suggested to occur,

Types of	Seed		Number of Seeds		Trisomics	Obtained
Trisomics	Size	Sown	Germinated	%	Number	%
Bush	Large	261	247	(94.63)	44	17.81
	Small	108	88	(81.48)	71	80.68
Slender	Large	128	108	(84.37)	17	15.74
	Small	46	20	(43.47)	18	90.00

Table 7.31 The Number and Percentage of	Trisomics in Two	vo Seed Groups of	Bush and Slender
Differing in Size		-	

Source: From Tsuchiya, T., Jpn. J. Bot., 17, 177-213, 1960a. With permission.

as has been shown for microspores. McClintock and Hill (1931) suggested that a basal megaspore with n + 1 chromosome constitution may not function to produce the embryo sac but may be replaced by a cell with the *n* chromosome number. However, such megaspore substitution was not reported in later studies of maize (Singleton and Mangelsdorf, 1940; Rhoades, 1942). Rédei (1965) reported some postreduction selection between the basal megaspores of different constitutions in *Arabidopsis*. The transmission of an extra chromosome has been observed (0 to 22%) through male gametes in primary trisomics. The low transmission of primary trisomics was accounted for by the preference in development of 2n cells over 2n + 1 cells (Sears and Lee-Chen, 1970).

Frequently, small seeds transmit a higher frequency of primary trisomic plants than large and plump seeds (Lesley, 1928; Einset, 1943; Tsuchiya, 1960a; Ramage and Day, 1960; Liang, 1979). Trisomic seeds are narrower, thinner, and later germinating, while large seeds are generally disomic. Tsuchiya (1960a) separated large and small seeds of two primary trisomics, Bush and Slender. He observed, in both cases, that small seeds showed poor germination but transmitted a considerably higher frequency of primary trisomic plants (Table 7.31). Ramage and Day (1960) suggested the use of a seed blower to enrich seed lots to contain a high frequency of trisomics, because trisomic seeds are lighter.

b. Pollen Transmission

The transmission of extra chromosomes through pollen in general is very low, because in diploid species, pollen with n + 1 chromosome constitution is unbalanced and generally unable to compete in fertilization with pollen carrying the balanced, n, chromosome number. The extra chromosome was not transmitted through pollen in $2x \times 2x + 1$ crosses of *H. spontaneum* primary trisomics (Tsuchiya, 1960a). Tetrasomic plants were obtained in the selfed progenies of Robust (0.5%), Pseudonormal (0.4%), and Semierect (5.9%) of cultivated barley, but no trisomic plants were recovered in $2x \times 2x + 1$ crosses (Tsuchiya, 1967). Male transmission of the extra chromosome has been observed in *D. stramonium* (Blakeslee and Avery, 1938), maize (McClintock and Hill, 1931), and tomato (Lesley, 1928, 1932). Pollen transmission of the extra chromosome was observed in only four primary trisomics of *Datura* (Blakeslee and Avery, 1938).

Lesley (1928) reported in tomato that male transmission of the extra chromosome was as frequent as through eggs in two trisomic types. Sears and Lee-Chen (1970) observed an average male transmission of 12% in *Arabidopsis thaliana*. The value ranged from 0.0% (F trisome) to 22.0% (N trisome). Buchholz and Blakeslee (1922) found in Glove trisomic of *Datura* that pollen tubes with the extra chromosome are slower growing than those of normal pollen. Tetrasomic plants have been isolated from the selfed progenies of primary trisomics of the soybean. All are viable, vigorous, and partially fertile, suggesting male transmission of the 21 chromosome gametes (Singh and Xu, unpublished data).

c. Progenies of Primary Trisomics

Generally, normal diploids and related primary trisomics are obtained in the progenies of primary trisomics. However, unrelated chromosomal variants are occasionally isolated, and they have been

reported in tomato (Lesley, 1928; Khush, 1973), *Datura* (Blakeslee and Avery, 1938), *Nicotiana* sylvestris (Goodspeed and Avery, 1939), barley (Tsuchiya, 1960a, 1967); and *Avena strigosa* (Rajhathy, 1975). The unrelated chromosome variants from the progenies of primary trisomics are telotrisomics, acrotrisomics, hypertriploids, triploids, hypotriploids, haploids, fragments of various sizes, ring chromosomes, secondary trisomics, tertiary trisomics, and various other chromosomal types. Einset (1943) studied the chromosomes of 1916 plants derived from $2x + 1 \times 2x$ crosses. There were 658 trisomics, five monosomics, one plant with 19 chromosomes + one fragment, one plant with 20 chromosomes + one fragment, three plants with 21 chromosomes + one fragment, two haploids, and three triploids.

Tetrasomic plants [an individual carrying two extra chromosomes in addition to its normal somatic chromosomes complement (2n = 2x + 2) is designated as tetrasomics] are expected in the progenies of primary trisomics if an extra chromosome is transmitted through male spore. Tetrasomics are rare and die prematurely in the progenies of barley, maize, rice, and tomato primaries. However, primary trisomics of the soybean transmit tetrasomics in low frequencies. The plants are viable, and compared to their counterpart, primary trisomics are slow in vegetative and reproductive growth and are partially fertile to completely fertile. Tetrasomics mostly breed true, and occasionally related trisomics are identified. Similarly, the progeny of tetrasomics × disomic is primary trisomics. Tetrasomics in the soybean are of unique cytogenetic stock, and that may reduce time devoted for cytological identification of tetrasomics and F_1 hybrids. Tetrasomic for Chromosome 13 with four NOR are, sometimes, observed at mitotic metaphase (Figure 7.31). The isolation of viable tetrasomics in the soybean suggests the tetraploid nature of the soybean genome.

5. Genetic Segregation in Primary Trisomics

Primary trisomics are helpful for locating a gene on a particular chromosome, verifying the independence of linkage groups, and also associating the genetic linkage groups with the individual chromosomes. The use of primary trisomics is often more efficient and dependable than other conventional and translocation procedures.

The principle of genetic segregation in primary trisomics has been described in detail by Burnham (1962), Hermsen (1970), and Khush (1973). When a primary trisomic is used to locate a gene on a particular chromosome, the genetic ratios are modified from 3:1 (F_2) or 1:1 (BC₁). The ratios encountered depend on the genotypes of the F_1 primary trisomic plants, whether duplex (AAa) or simplex (Aaa), on the type of chromosome segregation, and on the female transmission rate of the extra chromosome.

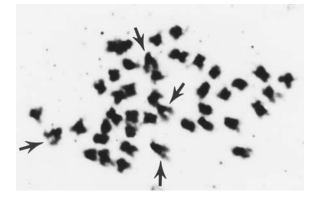


Figure 7.31 A mitotic metaphase cell of soybean tetrasomic for Chromosome 13 showing four SAT chromosomes (arrows). (From Singh, R.J., unpublished results.)

a. Association of a Recessive Gene

A mutant with the recessive genotype *aa* is crossed as a pollen parent onto a primary trisomic of the genotype *AAA*. In the F_1 , primary trisomic plants are identified morphologically and cytologically. Generally, one disomic plant is saved (control), and the remaining are discarded. The F_1 trisomic plants are allowed to self-pollinate or are test crossed (if sterile) to the homozygous recessive parent (BC₁). Two types of gametes are produced from the F_1 primary trisomics, *n* and *n* + 1. If *n* + 1 gametes are not male-transmitted with 50% frequency, total phenotypic ratios of 17A:1*a* and 5A:1*a* are expected in F_2 and BC₁, respectively. However, these ratios are never recorded, because usually only *n* gametes function in male-transmission, *n* + 1 gametes function in less than 50% frequency in female transmission.

On the basis of 50 and 33.3% female transmission of n + 1 gametes through females, assuming no male transmission of the extra chromosome, the expected frequencies of gametic types, genotypic frequencies, and phenotypic ratios derived from random chromosome, random chromatid, and maximum equational segregations in primary trisomic plants with AAa genotype are shown in Tables 7.32, 7.33, and 7.34, respectively. As evident from Table 7.34, the genetic ratios are modified if a gene is located in the extra chromosome in primary trisomic analysis, and results are quite different from disomic F_2 (3A:1a) and BC₁(1A:1a) ratios.

i. Random Chromosome Segregation — The expected frequencies of gametic types from a primary trisomic F_1 of duplex (AAa) genotypic constitution having 50% transmission of the extra chromosome through female are n + 1, 1AA + 2Aa, and n, 2A + 1a (Figure 7.32; Table 7.32). The expected genotypic frequencies in F_2 are as follows: 2x + 1 = 2AAA + 5AAa + 2Aaa and 2x = 4AA + 4Aa + 1aa (Table 7.33). Phenotypic frequencies will show as 9A-:0a::8A-:1a (Table 7.34). Similar modifications are expected in the BC₁: 3A-:0a::2A:1a. These ratios indicate that no recessive homozygous plants will be obtained in the trisomic fraction.

However, the expected 50% female transmission of n + 1 gametes is not observed. If we assume the female transmission of the extra chromosome is 33.3%, the frequency of an n gamete will become twice (4A + 2a), and the n + 1 gamete's (1AA + 2Aa) proportion will be unchanged (Table 7.32). A similar proportion of change will be recorded in the F₂ genotypic frequencies (2x + 1 = 2AAA + 5AAa + 2Aaa and 2x = 8AA + 8Aa + 2aa) (Table 7.33). The F₂ phenotypic ratio in trisomic (2x+1) fraction will be unaltered, but it will be changed in the disomic fraction (16A-:2a). Thus, the overall phenotypic ratio modifies from 17A-:1a to 12.5A-:1a (Figure 7.32). A similar change occurs in BC₁ (Table 7.34).

ii. Random Chromatid and Maximum Equational Segregation — Sometimes, homozygous recessive plants are obtained in the trisomic fraction of the F_2 and BC_1 populations. It is due to random chromatid crossing over or maximum equational segregation (Burnham, 1962). The expected gametic type frequencies in random chromatid segregation with 50% female transmission of n + 1 gametes are 6AA + 8Aa + 1aa, and n gametes are 10A + 5a, but with the 33.3% female transmission of n + 1 gametes, the frequency of n gametes becomes 20A + 10a (Table 7.32). The genotypic frequencies in an F_2 population having 50% female transmission of n + 1 gametes are expected as follows: 2x + 1 = 12AAA + 22AAa + 10Aaa + 1aaa and 2x = 20AA + 20Aa + 5aa. In F_2 , 8A (40A-:5a):1a phenotypic ratio is expected for the disomic fraction and 44A-:1a for the trisomic fraction. With 33.3% female transmission of the extra chromosome, the F_2 phenotypic ratio for the disomic fraction will be modified, and the total ratio is slightly smaller (11.27A-:1a) than that recorded in random chromosome assortment (12.5 A-:1a). A similar proportion of changes occurs in BC₁ (Tables 7.32, 7.33, and 7.34).

As in random chromatid segregation, the occurrence of recessive homozygotes in the trisomic fraction of the F_2 population can be explained based on maximum equational segregation (Burnham, 1962). Assume that only one chiasma is formed between a locus and a kinetochore in each of the three chromosomes of a trivalent (Figure 7.33). The gametic genotypes, n + 1 = 5AA + 6Aa + 1aa and n = 8A + 4a, are expected at the end of meiosis. There is not a real difference between the

	50% Female Transmission of <i>n</i> + 1 Gametes	iission of <i>n</i> + 1 es	33.3% Male Transmission of <i>n</i> + 1 Gametes	nission of <i>n</i> + 1 tes
Type of Segregation	<i>n</i> + 1	u	n + 1	u
Random chromosome	1AA + 2Aa	2A + 1a	1AA + 2Aa	4A + 2a
Random chromatid	6AA+ 8Aa+ 1aa	10A + 5 <i>a</i>	6AA+8Aa+1aa	20A + 10 <i>a</i>
Maximum equational	5AA+6Aa+1aa	8A + 4a	5AA+6Aa+1aa	16A + 8 <i>a</i>

Source: Burnham, C.R., Discussions in Cytogenetics, Burgess, Minneapolis, Minnesota, 1962. With permission.

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	50% Female Transmission of $n + 1$ Gametes	of n + 1 Gametes	33.3% Female Transmission of <i>n</i> + 1 Gametes	of n + 1 Gametes
Type of Segregation	2x + 1	2 <i>x</i>	2 <i>x</i> + 1	2 <i>x</i>
Random chromosome	2AAA + 5AAa + 2Aaa	4AA + 4Aa + 1aa	2AAA + 5AAa + 2Aaa	8AA + 8Aa + 2aa
Random chromatid	12AAA + 22AAa + 10Aaa + 1aaa	20AA + 20Aa + 5aa	12AAA + 22AAa + 10Aaa + 1aaa	40AA + 40Aa + 10aa
Maximum equational	10AAA + 17AAa + 8Aaa + 1aaa	16AA + 16Aa + 4aa	10AAA + 17AAa + 8Aaa + 1aaa	32AA + 32Aa + 8aa
		::		

Source: Burnham, C.R., Discussions in Cytogenetics, Burgess, Minneapolis, Minnesota, 1962. With permission.

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	20	% Fema	0% Female Transmission of n + 1 Gametes	sion of <i>n</i> .	+ 1 Gan	netes	33	.3% Fem	33.3% Female Transmission of n + 1 Gametes	sion of r	2 + 1 Gai	netes
	F2	~	Total	BC,	5	Total	Ľ	-5	Total	B	BC ₁	Total
Type of Segregation	2x + 1	2 <i>x</i>	Ratio A:a	2x + 1	2x	Ratio A:a	2x + 1	2 <i>x</i>	Ratio A:a	2 <i>n</i> + 1	2n	Ratio A:a
Random chromosome	0:6	8:1	17:1	3:0	2:1	5:1	0:6	16:2	12.5:1	3:0	4:2	3.5:1
Random chromatid	44:1	40:5	14:1	14:1	10:5	4:1	44:1	80:10	11.27:1	14:1	20:10	3.1:1
Maximum equational	35:1	32:4	13.4:1	11:1	8:4	3.8:1	35:1	64:8	11:1	11:1	16:8	3:1

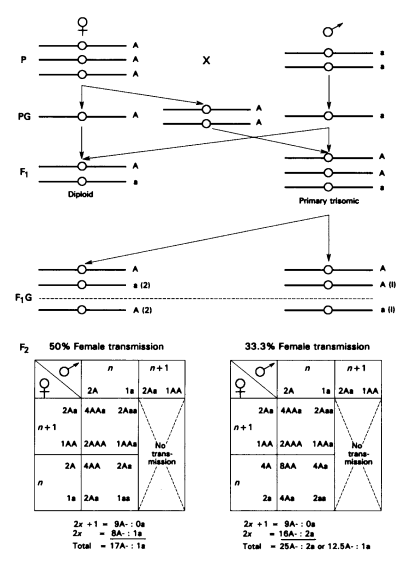


Figure 7.32 Random chromosome segregation showing association of a recessive marker (*a*) gene with a particular chromosome by primary trisomic method.

results expected from random chromatid segregation and maximum equational segregation (Tables 7.32, 7.33, and 7.34). In both cases, a low frequency of recessive homozygotes is expected with similar proportions in the trisomic fraction.

b. Association of a Dominant Gene

In this case, the genotype of the selected primary trisomic is *aaa*, and the genotype of the mutant is *AA*. The genotype of the F_1 primary trisomic is *Aaa* (simplex). The expected gametic types together with the F_2 genotypic, and phenotypic ratios based on 50% transmission of the extra chromosome in random chromosome assortment, random chromatid, and maximum equational segregations are provided in Table 7.35. It should be pointed out that separation of the trisomic and disomic fractions is advised when the genotype of the F_1 primary trisomic is in simplex (*Aaa*) condition.

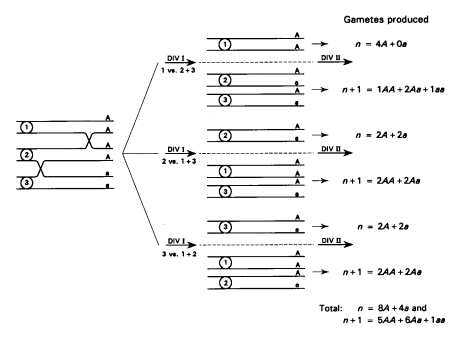


Figure 7.33 Maximum equational segregation in a primary trisomic heterozygote (Aaa).

c. Association of Codominant Gene

Primary trisomics have been used to associate isozyme markers to the specific chromosomes of tomato (Tanksley, 1983), rice (Delos Reyes, Khush, and Brar, 1998), and faba bean (Torres et al., 1995). These markers segregate in a codominant fashion (genotypic ratios) in the F_2 . In a critical combination (random chromosome), F_2 population segregates in a trisomic ratio, depending upon female transmission of the extra chromosome:

Genotypic Ratio	Female Transmission
5:7:1	30%
5:7.5:1	33.3%
6:11:1	50%

Molecular markers such as simple sequence repeats (SSR) can be associated with the chromosomes by primary trisomics. Cregan et al. (2001) associated SSR loci of molecular linkage group A_1 to Triplo 5, and F to Triplo 13 of the soybean. All SSR markers were segregated in a ratio of 6:11:1. The SSR loci not associated with Triplo 5 and Triplo 13 segregated in a disomic fashion of 1:2:1 (Table 7.36). This elucidates that primary trisomics form an excellent cytogenetic stock with which to associate molecular markers with the specific chromosomes by modification in the F_2 genetic ratio.

d. Association of Gene by Dosage Effect

The RFLP markers are assigned to specific arms of the chromosomes by gene dosage comparison of the autoradiographs in rice. By using secondary trisomics, telotrisomics, and primary trisomics of rice cv. IR36, Singh et al. (1996) crossed with a tropical japonica variety, MaHae. Both are highly polymorphic. An F_1 disomic has one allele each of IR36 and MaHae, and the intensities of

Iransmissio	Iransmission of <i>n</i> + 1 Gamete is	IS 50%)						
	Gametic Genotype	notype	F ₂ Ger	F ₂ Genotype	Phenotypic F ₂ Ratio	F ₂ Ratio	Phenotypic BC ₁ Ratio	BC, Ratio
Type of Segregation	n + 1	c	2x + 1	2 <i>X</i>	2x + 1	2х	2x + 1	2 <i>X</i>
Random chromosome	2Aa + 1aa	1A + 2a	2AAa + 5Aaa + 2aaa	1AA + 4Aa + 4aa	7:2	5:4	3:2	1:2
Random chromatid	1AA + 8Aa + 6aa	10 <i>A</i> + 20 <i>a</i>	1AAA + 10AAa + 22Aaa + 12aaa	10AA + 40Aa + 40aa	11:4	5:4	9:6	1: 2:
Maximum equational	1AA + 6Aa + 5aa	4 <i>A</i> + 8 <i>a</i>	1AAA + 10AAa + 17Aaa + 10aaa	4AA + 16Aa + 16aa	26:10	5:4	7:5	1.

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Source: Burnham, C.R., Discussions in Cytogenetics, Burgess, Minneapolis, Minnesota, 1962. With permission.

		• •		•
Triplos and SSR	Linkage	Segregation	Goodnes	ability for s of Fit to
Markers	Groups	Ratio	1:2:1	6:11:1
		Triplo 5		
Satt276	A1	27:26:3	0.001	0.05
Satt364	A1	26:27:3	0.001	0.10
Satt471	A1	25:27:2	0.001	0.14
Satt300	A1	26:27:3	0.001	0.10
Satt155	A1	25:28:3	0.001	0.18
Satt006	L	6:11:3	0.58	
		Triplo 13		
Satt569	F	24:34:5	0.0027	0.43
Satt193	F	24:34:5	0.0027	0.43
Satt030	F	24:34:5	0.0027	0.43
Satt343	F	24:34:5	0.0027	0.43
Satt657	F	20:38:5	0.0074	0.71
Satt022	Ν	7: 9:5	0.67	

Table 7.36 Segregation Ratios in F₂ Generation of the SSR Markers with Triplo 5 and Triplo 13 of the Soybean

Source: From Cregan, P.B. et al., *Crop Sci.*, 41, 1262–1267, 2001. With permission.

both autoradiographic bands in Southern blots were similar. An F_1 primary trisomic has two copies of the IR36 allele and one of the MaHae allele. The intensity of the IR36 band is expressed as twice that of the MaHae band. An F_1 secondary trisomic has three copies of the IR36 allele (located on the arm for which it is secondary) and one copy of MaHae allele. The intensity of the IR36 band for a marker present on that arm is three times that of MaHae band (Figure 7.34). If the marker in question is not located on the arm for which it is secondary trisomic, then the F_1 secondary trisomic shows similar intensity of IR36 and MaHae bands. The telotrisomic behaves the same way as the secondary trisomics, except that the intensity of the IR36 band is twice the intensity of the MaHae band. Based on these principles, RFLP markers were assigned to specific chromosome arms, and the positions of the centromeres were mapped between the nearest two markers located on opposite arms of a chromosome of rice.

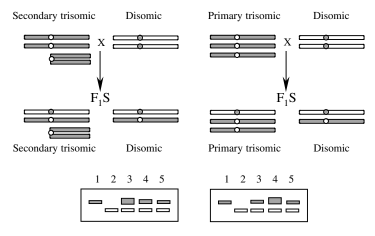


Figure 7.34 Dosage analysis in F₁ primary and secondary trisomics. Lanes 1–5 are IR 36, MaHae, F₁ secondary trisomic, F₁ primary trisomic, and F₁ disomic, respectively. (From Singh, K. et al., *Proc. Natl. Acad. Sci. USA*, 93: 6163–6168, 1996. With permission.)

e. Gene-Chromosome-Linkage Group Relationships

Primary trisomics have been used to associate a marker gene with a particular chromosome, to determine chromosome-linkage group relationships, and to test the independence of linkage groups (Burnham, 1962; Khush, 1973). Rick and Barton (1954) disproved the independence of the linkage groups X and XII and VI and VIII with the help of primary trisomic analysis in tomato. The following relationships between chromosomes-genes-linkage groups were established: 1-*y* (III); 2-*d* (I); 7-*wt* (X), *mc* (XII); 8-*l* (VI), *al*(VIII), *dl*; 9-*wd*; 10-*H* (VII). Moreover, Rick, Dempsey, and Khush (1964) analyzed 30 more marker genes with 11 of the 12 primary trisomics. Several linkage groups that were previously supposed to be independent were associated with the same chromosome, and none of the tested genes showed association with Triplo 12, the shortest chromosome. In rice, three markers (gh₁, nl₁, and gl₁) belonging to three different linkage groups (VI, IX, and XII) gave trisomic segregations with Triplo 5. Thus, these three linkage groups belong to Chromosome 5 (Khush et al., 1984). Similar results were obtained by Iwata and Omura (1976), who obtained trisomic ratios for *dl* (VI), nl₁, and gl₁ with their trisomic L. The gene d₂ (VII) and I-Bf (V) gave trisomic segregations with Triplo 9. Thus, linkage groups V and VII belong to the same chromosome.

Tsuchiya (1960a) studied ten marker genes belonging to four linkage testers with the seven primary trisomics of barley. The observed results indicated that Chromosome 6 carried no genetic linkage group previously established; two genetic linkage groups III and VII were located on Chromosome 1. Thus, the linkage groups were reduced to six. However, several new genes e_c (early), x_n (Xantha seedling), and o (orange lemma) were associated with Chromosome 6 by reciprocal translocations. Tsuchiya (1967) verified the association of gene o with Chromosome 6. A saturated linkage map of barley with molecular markers is shown in Appendix 7-I.

Lee-Chen and Steinitz-Sears (1967) studied linkage relationships in *Arabidopsis thaliana* by the primary trisomic method. Trisomic types NC and NS both gave a trisomic ratio with a marker gene *tz*. This demonstrates that NC and NS trisomics carry the same chromosome. A gene for narrow leaf (*an*) of linkage group-I did not show a trisomic ratio with any of the four tested primary trisomics, suggesting that the gene *an* may be located on the fifth trisomic type, which was not isolated.

By utilizing primary trisomics, Khush et al. (1984) identified all 12 possible linkage groups of rice. They selected 22 marker genes, one or two genes representing each linkage group, and crossed them with the trisomics. As soon as a marker was located on a specific chromosome, its tests with the remaining chromosomes were discontinued. They tested 120 of the possible 264 combinations involving 22 genes before markers for all of the chromosomes were identified.

Two dominant genes, Cl (clustered spikelets) and Ps1 (purple stigma-1) gave trisomic ratios with Triplo 6. The F₂ ratios in both cases were modified to a ratio of four normal:five mutant (instead of 1:3) in the diploid fraction. In the trisomic fraction of the F₂ of the Triplo $6 \times Ps1$, the ratio agreed with the expected two normal:seven mutant. However, the ratio of normal to mutant in the trisomic section of the F₂ population of Triplo $6 \times Cl$ was 7:2. This apparent reversal was due to the fact that $Cl \ cl \ cl$ individuals have normal phenotype (as determined from trisomic F₁ phenotypes) instead of mutant. Therefore, the expected genotypic ratio of 2 $Cl \ cl \ cl:5 \ Cl \ cl \ cl:2$ $cl \ cl \ cl$ was modified into seven normal to two mutant.

B. Secondary Trisomics

1. Introduction

Secondary trisomic plants carry an extra isochromosome (both arms homologous) in addition to their normal somatic chromosome complement. Secondary trisomics have been reported in *Datura stramonium* (Belling and Blakeslee, 1924; Blakeslee and Avery, 1938; Avery, Satina, and Rietsema, 1959), *Zea mays* (Rhoades, 1933b; Schneerman et al., 1998), *Lycopersicon esculentum*

(Sen, 1952; Moens, 1965; Khush and Rick, 1969; Rick and Gill, 1973), *Hordeum spontaneum* (Tsuchiya, 1960a), *Avena strigosa* (Rajhathy and Fedak, 1970), and *Oryza sativa* (Singh, Multani, and Kush, 1996). Secondary trisomics have been studied morphologically and cytologically in *Datura*, maize, tomato, and rice, and were utilized extensively in genetic and linkage studies in tomato (Khush and Rick, 1969) and rice (Singh, Multani, and Kush, 1996).

2. Sources of Secondary Trisomics

The secondary chromosome (isochromosome) might have arisen directly or progressively by way of an unstable telocentric fragment that would undergo a further misdivision or segregation without division (Darlington, 1939b).

In *Datura*, secondary trisomics have appeared spontaneously from unrelated primary trisomics occasionally, but rarely from unrelated secondaries or diploid parents. They most frequently appear spontaneously in the progenies of related primary trisomics. Rhoades (1933b) isolated a secondary trisomic plant for the short arm of Chromosome 5 from the offspring of an unrelated primary trisomic for Chromosome 6. In tomato, Sen (1952) identified two secondaries (2n + 8L.8L; 2n + 9L.9L) from the progenies of a mutation experiment, and Khush and Rick (1969) obtained nine of the possible 24 secondaries among the offspring of haplo-triplo disomics, double isocompensating trisomics, tertiary monosomics, and segmental deficiencies. Rick and Gill (1973) isolated three more new secondaries (2n + 5L.5L; 2n + 7L.7L; 2n + 11L + 11L) from the progenies of $2x + 1 \times 2x$ crosses. Singh, Multani, and Khush (1996) isolated secondary trisomics for both arms of chromosomes 1, 2, 6, and 7 and one arm of chromosomes 4, 5, 8, 9, and 12 from the progenies of a large population of each respective primary trisomics of rice (Table 7.37). The rarity of secondaries may be due to the poor viability of gametes and zygotes carrying an extra secondary chromosome.

3. Identification of Secondary Trisomics

a. Morphological Identification

In general, primary trisomics are morphologically, with a few exceptions, intermediate between those of their secondaries. In *Datura*, the seedlings of the 2n + 1.1 secondary showed narrow leaves due to

	-minary msoning	5		
	Total Plants	Secondary	/ Trisomics	Frequency
Trisomics	Grown	Short Arm	Long Arm	(%)
Triplo 1ª	_	1	1	_
Triplo 2	1632	2	1	0.18
Triplo 3ª	_	0	0	_
Triplo 4	1812	1	0	0.05
Triplo 5	1536	1	0	0.13
Triplo 6	2112	3	1	0.19
Triplo 7	3300	2	1	0.09
Triplo 8	1608	0	3	0.25
Triplo 9	2127	0	3	0.19
Triplo 10	1776	0	0	_
Triplo 11	600	1	1	3.33
Triplo 12	1632	2	0	0.12

Table 7.37 Frequency of Secondary Trisomics in the Progenies of Rice Primary Trisomics

^a Triplo 1 and Triplo 3 are highly sterile, and large populations could not be grown. 2n + 1S.1S and 2n + 1L.1L and .3L were selected from the progenies of primary trisomics.

Source: From Singh, K., Multani, D.S., and Khush, G.S., *Genetics*, 143, 517–529, 1996. With permission.

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factors in the .1 half, but the 2n + 2.2 secondary carried relatively broad leaves because of factors located on the half chromosome (.2). The primary trisomic 2n + 1.2 was intermediate in leaf width (Blakeslee and Avery, 1938). In tomato, secondary trisomics for the long arms (6L, 7L, 9L, 10L) showed much slower growth rates at all stages of plant growth than those recorded for the secondaries composed of short arms (3S, 7S, 9S). Secondary trisomics for the short arms were indistinguishable morphologically from disomics. The secondary 2n + 12L.2L was quite distinct at all stages of growth, but it showed only about half of the traits of its primary (Khush and Rick, 1969). In rice, most of the secondary trisomics resembled their counterpart primary trisomics for several morphological traits. In general, secondary trisomics. Some of the morphological features of primary trisomics were exaggerated in the secondary trisomics, while other secondary trisomics were indistinguishable from the normal diploid sibs. For example, secondary trisomic 2n + 7L.7L was weak and sterile, 2n + 7S.7S was partial fertile, and 2n + 4S.4S was morphologically like diploid sibs (Singh, Multani, and Khush, 1996).

b. Cytological Identification

Secondary trisomics are identified by the observation of a ring trivalent at diakinesis or metaphase-I (Figure 7.35A). At pachynema, based on euchromatin and heterochromatin differentiations, chromosome arm length and pairing pattern, Khush and Rick (1969) identified nine secondaries of tomato. Figure 7.35B shows a trivalent configuration at pachynema for a secondary trisomic. When a secondary chromosome remains as a univalent, it forms a "doughnut" (ringlet) or pair with the other homologous arms of the two normal chromosomes to form a trivalent. Ring trivalents in tomato secondaries for long arms are more frequent than those for the short arms (Table 7.38). In maize secondary 2n + 55.55, the highest frequency of sporocytes showed a ring trivalent (43.6%), followed by a ring univalent (32.3%) (Rhoades, 1933b). Similarly in *Datura*, the highest (44.9%) frequency was for the ring trivalent, followed by a V-shaped trivalent (22.0%) and a ring univalent (16.9%) (Belling and Blakeslee, 1924).

Secondary trisomics in rice were identified precisely at pachynema, and that facilitated pinpointing the position of kinetochores with certainty (Singh, Multani, and Khush, 1996). At diakinesis, various types of chromosome association were recorded in the secondary trisomics. Like tomato's secondaries, secondary trisomics in rice for the long arms showed higher frequencies of

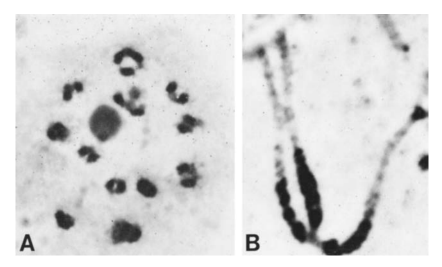


Figure 7.35 Chromosome configurations in secondary trisomics of tomato. (A) A ring trivalent configuration at diakinesis in secondary 2*n* + 10L.10L. (B) A trivalent configuration at pachynema in secondary 2*n* + 12L.12L; one chiasma can be seen in the 12 Ls. (From Khush, G.S. and Rick, C.M., *Heredity*, 24, 127–146, 1969. With permission.)

		osome Asso % Sporocytes	
Secondary Trisomics	11 + 1	12ll + 1l	11II + 3I
2n + 3S.3S	44	56	0
2n + 7S.7S	32	68	0
2 <i>n</i> + 9S.9S	40	56	4
2 <i>n</i> + 6L.6L	68	32	0
2 <i>n</i> + 8L.8L	58	42	0
2 <i>n</i> + 9L.9L	62	38	0
2 <i>n</i> + 10L.10L	60	40	0
2 <i>n</i> + 12L.12L	72	28	0

Table 7.38	Chromosome Association at Diakinesis in
	Secondary Trisomics of Tomato

Source: From Khush, G.S. and Rick, C.M., *Heredity*, 24, 129–146, 1969. With permission.

ring trivalent than those for the short arms. The frequency of ring trivalent ranged from 0.5% (2*n* + 4S.4S) to 25.6% (2*n* + 8L.8L) (Table 7.39).

4. Transmission of the Extra Chromosome in Secondary Trisomics

The transmission rate is correlated with the size of the extra secondary chromosome. The extra chromosomes with long arms cause greater imbalance than those for the short arms (Khush and Rick, 1969). Of the 14 secondaries studied in *Datura*, the female transmission rate of the secondary chromosome ranged from 2.4 (2n + 1.1) to 29.3% (2n + 19.19).

Secondary trisomics regularly throw related primaries. The frequency in *Datura* ranged from 0.06 (2n + 19.19) to 9.9% (2n + 9.9). Occasionally, unrelated primary trisomics appear in the offspring of secondaries. This suggests that the presence of one extra chromosome in some way stimulates nondisjunction in other chromosome pairs.

The female transmission rate of the extra secondary chromosome in tomato ranged from 0.0 (2n + 6L.6L) to 33.6% (2n + 7S.7S), and the frequencies of related primaries ranged from 0.0 (2n + 6L.6L)

		Chromoso	me Assoc	iation (% Sp	orocytes)	
Secondary	Total Cell			11 II + 1 III		
Trisomics	Observed	12 ll + 1l	Ring	Chain	Others	11II + 3I
2 <i>n</i> + 1S.1S	101	46.5	15.9	27.7	7.9	1.9
2 <i>n</i> + 1L.1L	165	29.7	24.8	23.0	18.2	4.2
2n + 2S.2S	147	74.8	6.8	6.8	11.6	0.0
2 <i>n</i> + 2L.2L	60	48.3	21.7	18.3	10.0	1.7
2 <i>n</i> + 4S.4S	215	81.4	0.5	6.5	11.6	0.0
2n + 5S.5S	127	65.4	10.2	14.2	7.1	3.1
2 <i>n</i> + 6S.6S	110	58.2	11.8	18.2	5.5	6.4
2 <i>n</i> + 6L.6L	142	50.7	19.7	15.5	11.3	2.8
2n + 7S.7S	156	57.7	15.4	17.9	9.0	0.0
2 <i>n</i> + 7L.7L	141	55.3	15.6	12.1	12.1	4.9
2 <i>n</i> + 8L.8L	125	60.0	25.6	11.2	3.2	0.0
2 <i>n</i> + 9L.9L	118	44.1	18.6	16.9	15.3	5.1
2 <i>n</i> + 11S.11S	146	65.1	11.6	12.3	10.3	0.7
2 <i>n</i> + 11L.11L	115	75.6	6.9	10.1	5.2	2.6
2 <i>n</i> + 12L.12L	93	55.9	15.1	14.0	12.9	2.1

Table 7.39 Chromosome Association at Diakinesis in Secondary Trisomics of Rice

Source: From Singh, K., Multani, D.S., and Khush, G.S., *Genetics*, 143, 517–529, 1996. With permission.

Secondary Trisomics	Total Plants	Disomics	Secondary Trisomics	Related Primary
2n + 5S.5S	181	69.1	28.2	2.7
2n + 6S.6S	41	63.4	19.5	17.1
2n + 8L.8L	116	55.2	24.1	20.7
2n + 11S.11S	103	68.0	25.2	6.8
2n + 11L.11L	124	69.3	23.4	7.3
2n + 12S.12S	123	54.5	40.6	4.9

Table 7.40 Transmission Rate (%) of the Extra Isochromosomes in the Selfed Progenies of Rice

Source: From Singh, K., Multani, D.S., and Khush, G.S., *Genetics*, 143, 517–529, 1996. With permission.

+ 7S.7S) to 14.9% (2n + 10L.10L). Like *Datura*, unrelated primaries were isolated in tomato, but the frequency was low (Khush and Rick, 1969).

The female transmission rates of the extra isochromosome in rice ranged from 19.5 (2n + 6S.6S) to 40.6% (2n + 12S.12S) (Table 7.40). Male transmission of isochromosome in rice was recorded for 2n + 4S.4S, as two plants in its progeny carried two extra 4S.4S isochromosomes (Singh, Multani, and Khush, 1996).

5. Genetic Segregation in Secondary Trisomics

Secondary trisomics can be used to locate genes in a particular half-chromosome in much the same way as with primary trisomics (Avery, Satina, and Rietsema, 1959). If a marker gene is located in the extra chromosome, a ratio of 3:1:: all (4):0 will be recorded for disomic and secondary trisomic fractions in random chromosome segregation. This suggests that no recessive homozygous plants will be obtained in the secondary trisomic fraction (Figure 7.36). In random chromatid segregation, the possible gametes with genotypes AAA (11), AAa (12), and Aaa (1) are expected in secondary trisomics. Thus, in F_2 , recessive homozygous plants will not be obtained in the secondary trisomic portion (all A-:0 a), but a very low frequency is expected in the primary trisomic portion (27.8 A-:1 a). The disomic portion is expected to segregate on a ratio of 4.76 A-:1 a (Figure 7.37). If a gene is not located on the secondary chromosome, a disomic ratio (3:1::3:1) in F_2 and in BC₁ (1:1::1:1) are observed for both diploid and secondary trisomic fractions.

Secondary trisomics were used for the first time by Khush and Rick (1969) to associate genes with a particular arm of a chromosome. They studied genetic segregation in the back crosses of five secondaries and located precisely the centromere positions for Chromosomes 8, 9, and 10, while they disproved the previously reported linkage map for Chromosome 10.

Secondary trisomics in rice have been used to locate marker genes on a particular arm of the rice chromosomes (Singh, Multani, and Khush, 1996). Secondaries for the short arm for Chromosomes 2, 4, 5, 6, and 12 and long and short arms for Chromosome 11 were used in locating genes on their respective arms (Table 7.41). For example, segregation of four genetic markers, z-1, v-4, la, and z-2, was analyzed in their crosses with 2n + 11L.11L as well as with 2n + 11S.11S. The segregation results show that z-1 is located on 11S, and the remaining three genes are located on 11L (Table 7.41).

C. Tertiary Trisomics

1. Introduction

A tertiary trisomic individual consists of an interchanged nonhomologous chromosome in addition to the normal somatic chromosome complement. Tertiary trisomics have been utilized effectively in

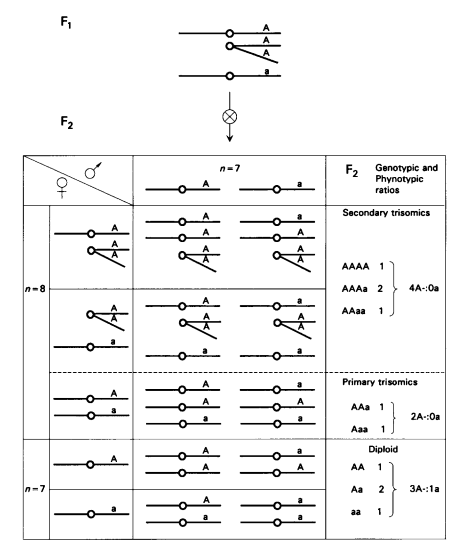
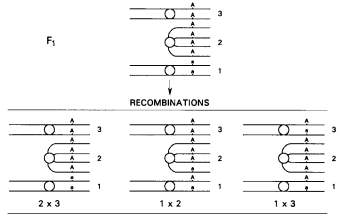


Figure 7.36 Association of a recessive gene (*a*) with a particular arm of a chromosome by use of secondary trisomics (random chromosome segregation).

tomato for determining centromere positions on the linkage maps and for associating a gene with a particular arm of a chromosome (Khush and Rick, 1967c). They have been used in barley for the construction of "balanced tertiary trisomics" (BTTs) for hybrid seed production (Ramage, 1965).

2. Sources of Tertiary Trisomics

In general, interchange heterozygotes that throw tertiary trisomics in their progenies have been described in maize (Burnham, 1930, 1934), pea (Sutton, 1939), *Oenothera lamarckiana* (Emerson, 1936), *Oenothera blandina* (Catcheside, 1954), *Datura stramonium* (Avery, Satina, and Rietsema, 1959), barley (Ramage, 1960), tomato (Khush and Rick, 1967c; Gill, 1978), *Phaseolus vulgaris* (Ashraf and Bassett, 1987), *Pennisetum glaucum* (Singh et al., 1982), *Secale cereale* (Janse, 1985, 1987), and lentil (Ladizinsky, Weeden, and Muehlbauer, 1990). Among the published results on tertiary trisomics, the investigation of Khush and Rick (1967c) in tomato is the most thorough. They reported seven tertiaries; two were isolated from tertiary monosomics and five from the interchanged heterozygotes.



Frequency of possible gametes with different genotypes:

			<i>n</i> =	- 8			n	= 7
Anonhoos	7 +	Secon	dary	7+	Prim	ary		
Anaphase separation	AAA	AAa	Aaa	AA	Aa	aa	A	a
2 x 3								
1.3 vs 2 1 vs 2.3 1.2 vs 3	0 4 0	0 0 4	0 0 0	0 0 0	4 0 0	0 0 0	0 0 2	0 2 0
1 x 2								
1.3 vs 2 1 vs 2.3 1.2 vs 3	0 2 1	0 2 2	0 0 1	2 0 0	2 0 0	0 0 0	0 1 2	0 1 0
1 x 3								
1.3 vs 2 1 vs 2.3 1.2 vs 3	0 2 2	0 2 2	0 0 0	1 0 0	2 0 0	1 0 0	0 1 1	0 1 1
Total								
n = 7 + Secondary n = 7 + Primary n = 7	11 	12 	1 	3	 8 	 1 	 7	 5

Figure 7.37 Association of a recessive gene (*a*) with a particular arm of a chromosome by use of secondary trisomics (random chromatid segregation).

In *Datura*, the tertiaries appeared spontaneously in a low frequency. In only six instances did tertiary trisomics occur spontaneously among approximately two million *Datura* plants grown to the adult stage. The rarity of occurrence of tertiaries and secondaries in *Datura* was attributed to the stable structure of the chromosomes (Avery, Satina, and Rietsema, 1959).

Tertiary trisomics originate from an interchange heterozygote in the following way. An interchange heterozygote occasionally forms a noncooriented quadrivalent configuration at diakinesis and metaphase-I, and a 3:1 random disjunction of chromosomes at anaphase-I will generate n + 1 gametes. Thus, eight possible types of 2x + 1 individuals are expected in the progeny of a selfed interchange heterozygote. Of the four tertiaries, two will be in homozygous background and the other two in translocation heterozygous background. For isolating tertiaries in tomatoes, Khush and Rick (1967c) hybridized interchanged heterozygotes with normal diploids. According to this procedure, the progeny should segregate in a proportion of two tertiaries (homozygous) to two primaries (interchange heterozygous) in the 2x + 1 fraction, and one normal to one interchange heterzygote in the 2x fraction (Figure 7.38). A plant is designated as a primary trisomic if an extra chromosome is normal; the trisomic may be a primary trisomic, a primary trisomic interchange heterozygote, or a primary trisomic interchange homozygote. On the other

			Disomics			Trisomics		
Cross	Total Plants (Number)	Normal (Number)	Recessive (Number)	χ² (3:1)	Normal (Number)	Recessive (Number)	Recessive (%)	Arm Location of the Gene
2n + 11S.11S/z-1	98	60	17	0.35	21	0	0.0	11S
2n + 11L.11L/z-1	91	51	16	0.04	19	5 2	0.8	11S
2n + 11S.11S/v-4	214	100	37	0.25	62	15	19.5	11L
2n + 11L.11L/v-4	207	111	28	1.75	68	0	0.0	11L
2n + 11S.11S/la	214	100	37	0.29	59	18	23.4	11L
2n + 11L.11L/ <i>la</i>	207	109	30	0.87	68	0	0.0	11L
2n + 11S.11S/z-2	200	119	29	2.31	42	10	19.4	11L
2n + 11L.11L/z-2	245	147	39	1.61	59	0	0.0	11L
Source: From Singh, K., Multani, D.S., and Khush, G.S., Genetics, 143, 517–529, 1996. V	K., Multani, D.S.	, and Khush, G	.S., Genetics, 1	43, 517–529	i, 1996. With p€	Vith permission.		

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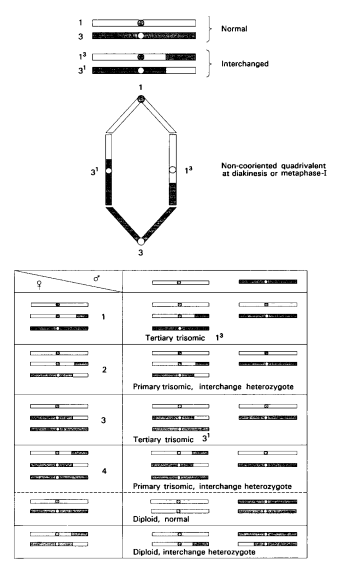


Figure 7.38 Expected types of plants from the progeny of an interchanged heterozygote.

hand, if the extra chromosome is a translocated chromosome, a plant is designated as a tertiary trisomic (Ramage, 1960).

3. Identification of Tertiary Trisomics

a. Morphological Identification

Tertiary chromosomes are composed of parts of two nonhomologous chromosomes. Thus, tertiaries inherit certain morphological features of the two related primaries. Like telotrisomics for the long arm, tertiary trisomics carrying a long arm generally exert a greater influence on plant morphology than when carrying a short arm.

Furthermore, if tertiary chromosomes contain two long arms, the longer of the two arms usually exerts a greater phenotypic effect, and such tertiaries frequently resemble the primary trisomics carrying the longer arm. In contrast, tertiaries for the short arms appear to contribute little to alter the phenotypic expression of the tertiary trisomics. In tomato, tertiary trisomic 2x + 5S.7L closely resembled Triplo 7, while the 2x + 7S.11L tertiary trisomic was similar to Triplo 11 in that the effect of the short arm was relatively minor. Tertiary trisomics for short arms such as 2x + 9S.12Sresembled neither Triplo 9 nor Triplo 12 (Khush and Rick, 1967c). This suggests that the short arms have little effect on phenotypes of tertiaries in tomato.

b. Cytological Identification

In tertiary trisomics, a maximum association of five chromosomes (pentavalent) or derived configurations, such as 1IV + I, 1III + 2I, 1III + II, 2II + 1I, 1II + 3I or 5I, are expected during diakinesis and metaphase-I. The derived configurations are attributed to the failure of chiasma formation or chiasma maintenance. Khush and Rick (1967c) analyzed chromosome associations at diakinesis in the seven tertiaries of tomato. The highest frequency of sporocytes showed 1III + 11II, and the next most frequent configurations were 1V + 10II and 12II + 1I. Sporocytes with such other configurations as 1IV + 10II + 1I, 1III + 10II + 2I, 11II + 3I, and 10II + 5I were rarely recorded. The formation of a pentavalent association depends upon the length of the tertiary chromosome. In tomato, tertiary chromosomes composed of short arms showed lower frequencies of the 1V + 10II configuration than those carrying long arms (Table 7.42).

The true nature of tertiary chromosomes can be established precisely by pachytene chromosome analysis. Such investigation has not been feasible in crops other than tomato, where breakage point and arm identity may be accurately determined (Khush and Rick, 1967c).

4. Transmission of the Extra Chromosome in Tertiary Trisomics

Tertiary trisomics usually throw, in addition to the parent type, small proportions of related primaries (Avery, Satina, and Rietsema, 1959; Khush and Rick, 1967c). In *Datura*, a very low frequency of related secondaries was also isolated, but these were not recorded in tomato (Khush and Rick, 1967c). However, Janse (1985) observed in a tertiary trisomic of rye that 58.1% of all microspores going through pollen mitosis carried seven chromosomes, and 41.9% microspores contained eight chromosomes. This suggests that up to the end of first pollen mitosis, aneuploid spores are not significantly fewer than euploid spores. Therefore, the failure of n + 1 gametes to be transmitted through pollen presumably results from the failure of n + 1 pollen to compete with n chromosome pollen. In tomato, tertiary chromosomes are transmitted through females, and frequencies ranged from 7.43 (2x + 2L.10L) to 40.78% (2x + 7S.11L). A high female transmission rate of tomato tertiaries was attributed to the genetically heterozygous background (Khush and Rick, 1967c). In common bean, the transmission rate of the extra tertiary chromosome after selfing ranged from 28 to 41% (Ashraf and Bassett, 1987), and in *Datura*, it ranged from 14.14 (2x + 1.18) to 32.05% (2x + 2.5).

Table 7.42 Frequencies of Various Chromosomal Associations Observed at Diakinesis in Tomato Tertiary Trisomics

	,							
Tertiary Trisomics	Number of Cells	10II + 1V	11 + 1	12ll + 1l	10II + 1IV + 1I	10 + 1 + 2	11II + 3I	10ll + 5l
2n + 1L.11L	50	15	20	13	1	1	0	0
2 <i>n</i> + 2L.10L	50	12	17	12	1	6	2	0
2 <i>n</i> + 4L.10L	50	22	20	6	1	1	0	0
2n + 5S.7L	50	13	20	15	0	2	0	0
2 <i>n</i> + 7S.11L	50	5	22	21	1	0	1	0
2n + 9L.12L	50	14	16	19	0	0	0	1
2 <i>n</i> + 9S.12S	50	11	17	17	2	2	0	1

Source: From Khush, G.S. and Rick, C.M., Can. J. Genet. Cytol., 9, 610-631, 1967c. With permission.

5. Genetic Segregation in Tertiary Trisomics

Genetic ratios in F_2 populations of tertiary trisomics are modified in a fashion similar to those recorded in secondary trisomics, telotrisomics, and acrotrisomics. Assuming duplex genetic constitution (*AAa*) of an F_1 , no male transmission of the tertiary chromosome, 50% female transmission, and a marker gene located on either arm of the tertiary chromosome, a trisomic ratio 3:1 (2*x*)::all 4:0 (2*x* + 1) or a total of 7:1 ratio is expected. Furthermore, a 5:1 ratio should be obtained when a female transmission rate of the tertiary chromosome reaches 33.3%. On the other hand, if a marker gene is not located on the tertiary chromosome, a disomic ratio should be recorded in the F_2 for 2*x* and 2*x* + 1 portions. Tertiary trisomics have been used in associating a gene with a particular arm of a chromosome in barley (Ramage, 1965) and tomato (Khush and Rick, 1967c).

Khush and Rick (1967c) utilized five tertiary trisomics of tomato in genetic and linkage studies. Genetic segregation of tertiary trisomic 4L.10L will be cited as an example. The markers *clau*, *ra*, *di*, *ful*, and *w*-4 are located on Chromosome 4. The genes *clau* and *ful* are on 4S, and *ra*, *di*, and *w*-4 are on 4L. The two markers *ag* and *tv* are on Chromosome 10. As expected, markers *clau* and *ful* showed a disomic ratio, and the remaining markers showed trisomic ratios. The occurrence of a low frequency of recessive homozygotes in the trisomic fraction can be explained by double reduction (Table 7.43).

6. Balanced Tertiary Trisomics

Ramage (1965) proposed a scheme utilizing balanced tertiary trisomics (BTT) for production of hybrid barley seed. Balanced tertiary trisomics are tertiary trisomics constituted in such a way that the dominant allele of a marker gene closely linked with the interchange breakpoint is carried on the tertiary chromosome. The recessive allele is carried on each of the two normal chromosomes. The dominant marker allele for a mature plant character, such as red plant color (R), may be carried on either the centromere portion or on the interchanged segment of the extra chromosome and should be linked with a male fertile gene (Ms). The two normal chromosomes should carry the corresponding recessive male sterile allele (ms). All balanced tertiary trisomics would be male fertile and red, and all diploids would be male sterile and green. All functioning pollen produced by the balanced tertiary trisomics would carry the male sterile (*ms*) and the green plant color alleles (r). The self-progeny of such a trisomic would be planted in an isolation block. Diploid plants would be green and male sterile. All seed set on them would produce male sterile diploids in the next generation. The diploid plants would be harvested separately, and seed produced on them would be used to plant the female rows in the hybrid seed production field. Balanced tertiary trisomics planted in isolation would be dominant and would be harvested separately. Seed produced on balanced tertiary trisomic plants would produce approximately 70% male sterile diploids and 30% balanced tertiary trisomics (Figure 7.39). This seed would be used to plant an isolation block the following year.

The prerequisites for utilizing the BTT system in hybrid seed production are that balanced tertiary trisomics produce abundant pollen and that during anthesis, sufficient wind and insects are available for pollen dissemination. Several other alternatives to distinguish tertiary trisomics from diploids have been suggested, such as resistance or susceptibility to phytocides, seed size or shape, and differential plant heights (Ramage, 1965).

D. Telotrisomics

1. Introduction

A telocentric chromosome consists of a kinetochore (centromere) and one complete arm of a normal chromosome. The plant with a normal chromosome complement plus an extra telocentric chromo-

					Progeny				
			2n			21	2 <i>n</i> + 1		
				%				%	
Gene	Chromosome	Total	Normal	Recessive	Recessive	Normal	Recessive	Recessive	χ² (3:1)
clau	4	496	263	95	26.2	94	44	31.8	3.58
ra	4	496	272	86	24.0	138	0	0.0	46.00
di	4	496	300	58	16.2	136	0	1.4	40.81
ful	4	239	107	33	23.5	73	26	26.2	1.17
W-4	4	239	107	33	23.5	66	0	0.0	33.00
ag	10	202	91	25	21.5	83	ი	3.4	21.80
tv	10	202	89	27	23.2	83	Ю	3.4	21.80
Source:	Source: From Khush, G.S. and Rick, C.M., Can. J. Genet. Cytol., 9, 610-631, 1967c. With permissior	3. and Rick, C.	M., Can. J. C	Genet. Cytol., 9,	610-631, 1967c	. With permis	sion.		

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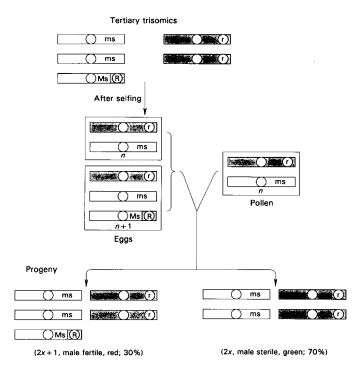


Figure 7.39 Utilization of balanced tertiary trisomics (BTT) to produce hybrid barley. (Redrawn from Ramage, R.T., *Crop Sci.*, 5, 177–178, 1965.)

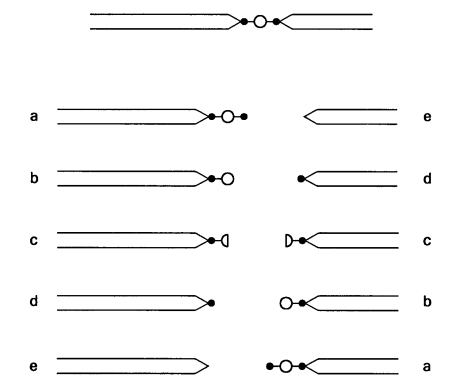


Figure 7.40 Origin of various types of telocentric chromosomes. (Redrawn from Steinitz-Sears, L.M., *Genetics*, 54, 241–248, 1966.)

some (2x + 1 telocentric; designated as telo) is designated as monotelotrisomic or telotrisomics (Kimber and Sears, 1969; Tsuchiya, 1972a). The telocentric chromosome contains a terminal kinetochore (Figure 7.40).

2. Sources of Telotrisomics

Monotelotrisomics originate spontaneously, often in the progenies of primary trisomics [*Zea* mays (Rhoades, 1936); *Datura stramonium* (Blakeslee and Avery, 1938); *Lotus pedunculatus* (Chen and Grant, 1968b); *Hordeum vulgare* (Tsuchiya, 1971b; Fedak, Tsuchiya, and Helgason, 1971; Singh and Tsuchiya, 1977; Siep, 1980; Furst and Tsuchiya, 1983; Shahla and Tsuchiya, 1990; Wang and Tsuchiya, 1990; Tsuchiya, 1991); *Avena strigosa* (Rajhathy, 1975); *Secale cereale* (Zeller, Kimber, and Gill, 1977; Sturm and Melz, 1982; Melz and Schlegel, 1985; Zeller, Cermeño, and Friebe, 1987); *Oryza sativa* (Singh, Multani, and Khush, 1996; Cheng et al., 2001)] and triploids [*H. vulgare* (Tsuchiya, 1971b; Fedak and Tsuchiya, 1975) and *Lotus pedunculatus* (Chen and Grant, 1968a)]. Of the 11 reported telotrisomics of barley, six (Triplo 2L, 2S, 3L, 4L, 5S, 6S) appeared in the progenies of their respective primary trisomics, and three (Triplo 1L, 1S, 5L) were isolated from triploids. Only, Triplo 3S and Triplo 7S originated in the progenies of plants carrying 2n = 13 + 1 acro $3L^{3S} + 1$ telo 3S, and 2n = 15 + 1 telo 7S, respectively (Table 7.44).

In rice, Singh, Multani, and Khush, (1996) isolated and identified telotrisomics for the short arm of Chromosomes 1, 8, 9, and 10 and for the long arm of Chromosomes 2, 3, and 5 from the progenies of their respective primary trisomics. Cheng et al. (2001) isolated all 24 telotrisomics of rice from approximately 180,000 plants derived from the trisomics and other aneuploids. *This report now claims the establishment of a complete set of telotrisomics in an economically important diploid crop*. They were able to complete the set by examining cytologically the variants morphologically distinct from the primaries and disomics. This study demonstrates that a complete set of telotrisomics can be established in any crop if a large population of primary trisomics is grown.

Khush and Rick (1968b) isolated six telotrisomics of the possible 24 of tomato from the progenies of terminal deficiency (Triplo 4L), tertiary monosomics (Triplo 3L, 8L, 7L) and compensating trisomics (Triplo 3S, 10S).

Proposed Designation of Telotrisomics	Source	Authority
Triplo 1L (7HL)ª	Autotriploid	Tsuchiya, 1971b, 1972a
Triplo 1S (7HS)	Triploid hybrid	Tsuchiya, 1971b
Triplo 2L (2HL)	Slender	Tsuchiya, 1971b
Triplo 2S (2HS)	Slender	Tsuchiya, 1971b
Triplo 3L (3HL)	Pale (F1 hybrid)	Tsuchiya, 1971b
Triplo 3S (3HS)	Novel compensating diploid ($2n = 13 + 1 \text{ acro } 3L^{3S} + 1 \text{ telo } 3S$)	Singh and Tsuchiya, 1981d
Triplo 4L (4HL)	Robust	Tsuchiya, 1971b
Triplo 5L (1HL)	Triploid hybrid	Tsuchiya, 1971b, 1972b
Triplo 5S (1HS)	Pseudonormal	Furst and Tsuchiya, 1983
Triplo 6S (6HS)	Purple (F_2)	Siep, 1980
Triplo 7S (5HS)	Semierect (F_1 hybrid) ($2n = 15 + 1$ telo 7S)	Shahla and Tsuchiya, 1983

Table 7.44 Sources of Eleven Telotrisomics of Barley

^a Homoeologous designation.

Source: Costa, J.M., Theor. Appl. Genet., 103, 415–424, 2001. With permission.

The low frequency of occurrence of telocentric chromosomes in the progenies of aneuploids of diploid crop plants, such as maize, barley, tomato, and many others, may be ascribed to the following reasons (Tsuchiya, 1972a).

The frequency of univalents in primary trisomics is low. The overall frequency of the sporocytes having one univalent was 23.7% (range of 20.0 to 36.5%) for the seven primary trisomics of Hordeum spontaneum and 22.5% (range of 15.3 to 29.3%) in H. vulgare (Tsuchiya, 1960a, 1967). In Beta vulgaris, the univalent frequency of three primary trisomics averaged 23.9%, ranging from 21.36 (Triplo 2) to 29.88% (Triplo 3) (Kaltsikes and Evans, 1967). Working with the primary trisomics of Arabidopsis thaliana, Sears and Lee-Chen (1970) found the 5II + 11 chromosome configuration to range from 10.3 (Triplo F) to 30.1% (Triplo R), with an average of 23.5%. In Nicotiana sylvestris, the average frequency of 12II + 3I was 4% (range of 0 to 8%) (Goodspeed and Avery, 1939). A much higher frequency of univalents was observed in tomato by Rick and Barton (1954), where 11 primaries averaged 45.6%. Values ranged from 26 (Triplo 3) to 70% (Triplo 12). These results indicate that the probability of the appearance of telocentric chromosomes in the progenies of primary trisomics will be low, because only a low frequency of univalents will lag at anaphase-I to telophase-I and will be subjected to misdivision — a prerequisite for the origin of telocentrics. In contrast, in wheat monosomics, the 20II + 1I configuration is sometimes observed in all the sporocytes (Person, 1956). This suggests that a univalent in wheat monosomics has a greater chance of producing telocentric chromosomes by misdivision than those recorded in primary trisomics. Misdivision of a kinetochore is a potent source of obtaining telocentric chromosomes in cotton (Brown, 1958) and wheat (Sears, 1952a,b; Steinitz-Sears, 1966).

3. Identification of Telotrisomics

a. Morphological Identification

In general, it has been observed in diploid species that a telotrisomic for the long arm of a chromosome has a similar effect on plant morphology as the corresponding primary trisomic. Goodspeed and Avery (1939) observed that a plant with 2n + 1f (fragment = telocentric) chromosome constitution was similar in plant morphology to the corresponding primary trisomics. Rhoades (1936) discovered a plant with 2n + 1 telo 5S in maize that did not resemble its primary trisomic or diploid sibs but was intermediate in appearance.

The effects of the short and long arms seem to depend upon the genetic material present in the extra telocentric chromosome. Khush and Rick (1968b) recorded in tomato that Triplo 4L, 7L, and 8L resembled their respective primary trisomics. Similarly, in barley Triplo 1L, 2L, 3L, 4L, and 5L were morphologically similar to their corresponding primaries (Singh and Tsuchiya, 1977). Triplo 2S did not resemble the primary trisomic, Slender, or diploid sibs. In contrast, morphological features of Triplo 1S, 3S, 5S, 6S, and 7S of barley (Singh and Tsuchiya, 1977, 1981b; Siep, 1980; Shahla and Tsuchiya, 1983) and Triplo 3S and 10S in tomato (Khush and Rick, 1968b) were similar to their diploid sibs.

The influence of long and short arm in telotrisomic condition can be precisely compared only in rice, because all 24 telotrisomics are described (Cheng et al., 2001). Triplo 4S, 5S, 7S, 9S, 10S, and 11S were morphologically indistinguishable from the disomic sibs. In this situation, chromosome count is required to establish the telotrisomic nature. Triplo 1S, 2S, 3S, 6S, 8S, and 12S expressed some characteristic morphological alterations. Telotrisomics for the long arm (Triplo 4L, 5L, 6L, 7L, 9L, 10L, 12L) resembled their corresponding primaries. All telotrisomics were fertile, even telotrisomics for Triplo 3, which was highly sterile. Telotrisomics for Chromosomes 1, 2, and 3 did not resemble diploid sibs or their respective primary trisomics. This study demonstrates that the short arm may contain less genetic material compared to the long arm, and thus, its influence is not that apparent on plant morphology than the long arm.

b. Cytological Identification

The extra telocentric chromosome is identified by somatic as well as meiotic chromosome counts. The precise arm identification is performed by karyotype analysis, utilizing somatic metaphase or meiotic pachytene chromosomes. The somatic karyotype analysis, however, does not provide enough evidence to distinguish between long and short arms, where both arms are of equal size and also lack euchromatin and heterochromatin differentiation. In the barley complement, Chromosomes 1, 2, 3, and 4 have almost median kinetochores and are indistinguishable by aceto-carmine or Feulgen staining techniques. Therefore, the initial identification of telotrisomics was based on morphological features and genetic studies (Fedak, Tsuchiya, and Helgason, 1971, 1972; Singh and Tsuchiya, 1977). Furthermore, translocation testers also did not elucidate the arm identity. Tsuchiya (1972a) analyzed meiotic pairing of three F_1 hybrids of Triplo 1L and translocation testers T1–4a, T1–6a, and T1–7a. He observed a 1V + 5II chromosome configuration at metaphase-I. This only demonstrated that the telocentric chromosome traced to Chromosome 1.

Based on genetic analysis, Fedak, Tsuchiya, and Helgason (1971) reported one telocentric chromosome to be the short arm of Chromosome 5, because it showed trisomic ratios with genes *trd* (third outer glume) and a_t (albino seedling); these genes are located on the short arm of linkage group 5 (Robertson, 1971). From karyotype analysis, Tsuchiya (1972b) demonstrated that previously identified telo 5S was a telo 5L (Figure 7.41). On this basis, he corrected the genetic linkage map of Chromosome 5.

By utilizing the Giemsa N-banding pattern, Singh and Tsuchiya (1982a,b) and Tsuchiya (1991) correctly identified and designated 11 telotrisomics (Telo 1L, 1S, 2L, 2S, 3L, 3S, 4L, 5L, 5S, 6S, 7S) of barley. Chromosome 3 contains its kinetochore at a median region, and its long arm is the second longest in the barley complement. Telo 3L showed a centromeric (fainter than telo 3S) and an intercalary band, and it also showed a faint dot on each chromatid at its distal region. Telo 3S showed a dark centromeric band only.

Similarly, the N-banding technique demonstrated that the previously identified telo 4S (Linde-Laursen, 1978; Singh and Tsuchiya, 1977, 1981a; Tsuchiya and Singh, 1982) was actually telo 4L. The morphology of correctly identified Triplo 4L is similar to Triplo 4. Giemsa N-banding technique revealed a deficiency in two telocentric chromosomes, telo 2S and 4L, of barley. Telo 2S showed a 50% distal deletion, while telo 4L had a 32% deficiency. The apparent loss of a distal portion of telo 2S and 4L did not affect morphology, transmission rate (Singh and Tsuchiya, 1977), or meiotic

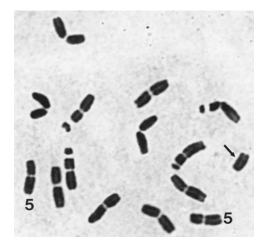


Figure 7.41 Mitotic metaphase chromosomes of Triplo 5L after aceto-carmine staining of barley. Compare long arm of Chromosome 5 (Chromosome 5 numbered) and Telo 5L (arrow). (From Singh, R.J., *Cytogenetics of Telotrisomics in Barley*, Ph.D. thesis, Colorado State University, 1974.)

behaviors (Singh and Tsuchiya, 1981a). In this case, it is likely that the broken end healed and started functioning as a normal telomere (Hang, 1981). If telotrisomic plants having deficiency in the telocentric chromosome were used in genetic-linkage analysis, a wrong conclusion could be drawn, because genes located in the deficient segment would show a disomic ratio. Telotrisomic analysis with Triplo 2S - *yst3* may be ascribed to a deficiency in telo 2S (Tsuchiya and Hang, 1979). These results indicate the importance of detailed karyotype study of telotrisomic plants.

Pachytene chromosome analysis has played a major role in maize and tomato cytogenetics. In both crops, long and short arms could be distinguished without doubt because of characteristic euchromatin and heterochromatin differentiation. Khush and Rick (1968b) were able to identify the available six monotelotrisomics of tomato at pachynema. Triplo 3S, 7L, and 10S possessed a truly terminal kinetochore, and telo 3L, 4L, and 8L had subterminal kinetochores.

Rice telotrisomics for 9S and 10S were easily identified, as they were associated with the nucleolus, while pachytene chromosome failed to identify other telotrisomics (Singh, Multani, and Khush, 1996). Cheng et al. (2001) identified telocentric nature by FISH using a rice centromeric-specific bacterial artificial chromosome (BAC) clone, 17p22, as a marker probe.

c. Meiotic Behavior of Telotrisomics

In telotrisomic plants, a telocentric chromosome may associate with its normal homologues, forming a trivalent (Figure 7.42A,B) or remaining an univalent. Singh and Tsuchiya (1981a) studied meiotic behavior of seven telocentric chromosomes in telotrisomic conditions. The frequencies of the different configurations are shown in Table 7.45. In the trivalent configurations, the telocentric chromosome was frequently tightly associated with an interstitial chiasma that persisted into metaphase-I (Figure. 7.42B). The average frequency of 1III + 6II chromosome configuration was 77% at diakinesis for the seven monotelotrisomics, ranging from 60.2 (Triplo 2L) to 89.1% (Triplo 5L). The remaining cells had 7II + 1I. As the meiotic stage proceeded from diakinesis to metaphase-I, the frequency of the 1III + 6II configuration. Rhoades (1940) observed that a telocentric chromosome in maize was associated at metaphase-I with its normal homologue in 59% of the sporocytes. Smith (1947) reported the 1III + 6II configuration in 65.9% sporocytes in *Triticum monococcum*, while the telocentric chromosome remained as an univalent in 34.1% cells.

The association of telocentric chromosome with its normal homologue depends upon the length of the extra chromosome. Khush and Rick (1968b) observed in tomato that the telocentric chromosomes for long arms (Triplo 3L, 4L, 7L, 8L) showed more trivalent formation than those for the short arms. Similar results were observed in the telotrisomics of rice (Singh, Multani, and Khush, 1996). In barley, the long and short arms do not have significantly different effects on the formation of trivalents, because the difference between the lengths of the long and short arms is not large enough to exert such an effect on trivalent formation. For example, Triplo 2L showed

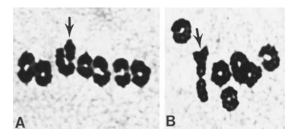


Figure 7.42 Chromosome configurations at metaphase-I in Triplo 3L of barley. (A) Tandom V-shaped trivalent.
 (B) Rod-shaped trivalent; interstitial chiasma (arrow) between telocentric and normal chromosome is evident. Arrows show a telocentric chromosome. (From Singh, R.J. and Tsuchiya, T., *Bot. Gaz.*, 142, 267–273, 1981a. With permission.)

	Diaki	Diakinesis	Metap	aphase-I		Trivalents	nts		Anap	hase-I ration	
		;		;	:		Ring-				Pollen
Triplo	1111 + 61I	711 + 117	1111 + 611	711 + 11	>	Rod	Rod	8-7	7-1-2	Others	Fertility
1L	80.0 ^a	20.0	74.8	25.2	67.4	7.9	24.7	82.0	12.0	6.0	86.5
1S	80.0	20.0	59.9	40.1	59.6	31.2	9.2	72.6	12.2	15.2	93.1
2L	60.2	39.8	54.6	45.4	64.0	13.8	22.2	80.0	18.0	2.0	83.7
2S	86.0	14.0	70.0	30.0	71.4	17.2	11.4	76.0	12.0	12.0	96.9
3L	77.8	22.2	71.0	29.0	57.2	15.9	26.9	59.4	20.3	20.3	75.4
4L	66.0	34.0	63.0	37.0	60.3	27.0	12.7	75.0	19.0	6.0	95.0
5L	89.1	10.9	71.0	29.0	44.0	31.2	24.8	56.0	20.0	24.0	81.1
Average	77.0 ^b	23.0	66.3	33.7	60.6	20.6	18.8	71.5	16.2	12.2	87.4
			:								

^a Observed χ^2 -value of trivalent association among telotrisomics at diakinesis (96.88) and metaphase-I (36.58) was significant at P < 0.001for 6 df. ^b Significant (χ^{2} -value 48.8, P < 0.001).

Source: From Singh, R.J. and Tsuchiya, T., Z. Pflanzenzüchtg., 78, 327–340, 1977; Singh, R.J. and Tsuchiya, T., Bot. Gaz., 142, 267–273, 1981a. With permission.

the lowest (60.2%) frequency of trivalent association at diakinesis, even though 2L is the thirdlongest long arm in the barley chromosome complement (Tjio and Hagberg, 1951). A great deal of difference was found in Triplo 2L studied at different times (Tsuchiya, 1971b). This result suggests that telo 2L may be sensitive to environmental conditions or that some other factors may be involved in chromosome association.

A complete chromosome in primary trisomic condition may have greater physical opportunity for synapsis with its homologues than a telocentric chromosome. This holds true for barley but not for tomato. The average metaphase-I trivalent configuration for the seven telotrisomics of barley was 66.3%, while the corresponding simple primary trisomics (Triplo 2, 3, 4, 5) showed trivalent in 77.9% sporocytes (Tsuchiya, 1967). This trend, however, was not recorded in tomato. Rick and Barton (1954) reported trivalent frequencies as: Triplo 3 = 70%; Triplo 4 = 56%; Triplo 7 = 36%; Triplo 8 = 24%; and Triplo 10 = 46%. When the respective telotrisomics are compared, it is evident that telo 7L (58.0%) and 8L (58.0%) showed a higher trivalent association than those recorded for their respective primaries (Khush and Rick, 1968b). This feature, therefore, differs from crop to crop, and a definite conclusion cannot be drawn.

Different types of trivalents are observed at diakinesis and metaphase-I (Table 7.45). These are tandem V-shaped (Figure 7.42A), rod-shaped (Figure 7.42B), and ring-and-rod-shaped. In barley telotrisomics, tandem V-shaped trivalents were predominant, with an average frequency of 60.6% of the sporocytes. Rod-shaped trivalents were next, with an average of 20.6%. Ring-and-rod trivalents were lowest in frequency with an average of 18.8% (Table 7.45). The last type has a more complicated chiasma pattern than the other two types of trivalents. Rhoades (1940) also observed tandem V-shaped trivalent in the majority of the sporocytes of a maize monotelotrisomic. The movement of these members of a trivalent is not random at anaphase-I. Among monotelotrisomics of barley, in an average of 71.5% of the sporocyte, the telocentric chromosome moved to a pole with one of its homologues, giving 8–7 chromosome disjunction (Table 7.45; Figure 7.43A). This has been observed in *Z. mays* (Rhoades, 1936), *T. monococcum* (Smith, 1947), and *L. esculentum* (Khush and Rick, 1968b).

Sometimes the telocentric chromosome does not move to either pole and remains as a laggard at the equatorial plate (Figure 7.43B). In barley, an average of 16.2% sporocytes showed 7–1–7 separation in the seven monotelotrisomics, ranging from 12 (Triplo 1L, 2S) to 20.3% (Triplo 3L). At late anaphase-I, the lagging telocentric chromosome started to divide, and as anaphase pro-

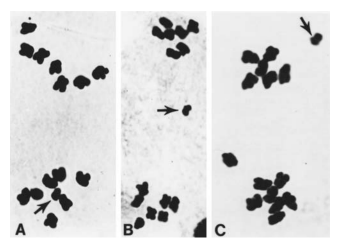


Figure 7.43 Separation of chromosomes at anaphase-I in Triplo 3L of barley. (A) 8–7 chromosome disjunction; arrow indicates telocentric chromosome. (B) 7–1–7 chromosome disjunction; telocentric chromosome arrow as a laggard. (C) 8–7 chromosome disjunction; telocentric chromosome (arrow) is on the upper pole with six chromosomes and normal chromosomes on the other pole. (From Singh, R.J. and Tsuchiya, T., *Bot. Gaz.*, 142, 267–273, 1981a. With permission.)

gressed, it divided, giving an 8–8 separation, and sometimes laggards were not included in the telophase nuclei. Occasionally, the telocentric moved to one pole, and the two normal homologues passed to the other pole (Figure 7.43C). This type of separation may be responsible for the occurrence of primary trisomics, with 15 complete chromosomes in the progenies of telotrisomics of barley (Singh and Tsuchiya, 1977).

At telophase-I, an average of 91.7% of the sporocytes for the seven telotrisomics showed normal separation of chromosomes, and at interkinesis, normal and abnormal cells with micronuclei were found. Meiotic abnormalities of the first division were reflected in the second division, which included laggards and micronuclei at the quartet stage (Singh and Tsuchiya, 1981a). Pollen fertility in seven telotrisomics of barley ranged from 75.4 (triplo 3L) to 96.9% (Triplo 2S), while the pollen fertility of normal diploid sibs was 97.8% (Table 7.45; Singh and Tsuchiya, 1977).

i. Meiotic Abnormalities — In the progenies of telotrisomics of barley, Singh and Tsuchiya (1981a) observed multiploid cells in Triplo 3L (Figure 7.44). A similar phenomenon was observed in Triplo 3 of *H. spontaneum* (Tsuchiya, 1960a) and Triplo 4 in *H. vulgare* (Tsuchiya, 1967).

Asynaptic cells were observed in Triplo 2L at a much higher frequency (4%) than in Triplo 2 (Tsuchiya, 1960a). Triplo 2S also showed asynaptic cells in 0.3% of the sporocytes. This indicates that both arms of Chromosome 2 may have an effect on the formation of asynaptic cells.

ii. Stability of Telocentric Chromosomes — The cytological stability of a telocentric chromosome depends upon the structure of its kinetochore and its behavior during cell division. Marks (1957) classified telocentric chromosomes into two categories: one group is stable, and the other is unstable. The instability of a telocentric chromosome is caused by nondisjunction and misdivision of univalents.

Rhoades (1940) indicated that telocentric chromosomes undergo structural changes in somatic cells. The loss and modification of telocentric chromosomes in somatic tissues suggests that a terminal kinetochore is unstable. Thus, the stability of a telocentric chromosome depends upon the kinetochore constitution. The kinetochore of most mitotic chromosomes is observed as a constriction, and in meiotic chromosomes, it appears as a simple, homogeneous, translucent body (Rhoades, 1940).

Based on her cytogenetic analysis of telocentric chromosomes in hexaploid wheat, Steinitz-Sears (1966) suggested that the relative instability of a telocentric chromosome may be attributed to the degree of completeness of its kinetochore (Figure 7.40). Generally, telocentric chromosomes of categories d and e are unstable. Darlington (1939b) explained the absence of telocentric chromosomes in plants by their instability. It appears that his generalization of unstable telocentric chromosomes was based on the study of the d and e categories (Figure 7.40).

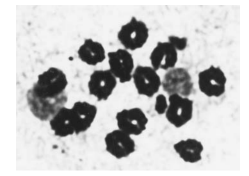


Figure 7.44 A multiploid sporocyte observed in Triplo 3L at diakinesis showing 14II + 2I. (From Singh, R.J., unpublished results.)

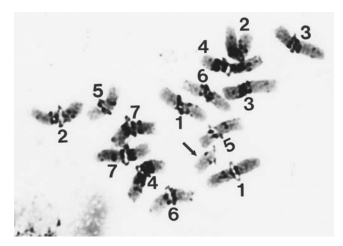


Figure 7.45 Giemsa N-banded mitotic mid-metaphase chromosomes of Triplo 5L of barley. Telo 5L contains half of the kinetochore (arrow). (From Singh, R.J. and Tsuchiya, T., *J. Hered.*, 73, 227–229, 1982b. With permission.)

Barley telotrisomics are fairly stable, except for Triplo 1L, which shows chimaerism (Tsuchiya, 1972a). Singh and Tsuchiya (1981a) speculated that barley telocentric chromosomes contain complete kinetochores, because they are highly stable. However, Giemsa N-banding technique revealed that they contained half of the kinetochore (Figure 7.45) and belonged to Category c of Steinitz-Sears (1966). The appearance of the diamond-shaped kinetochore in complete chromosomes and a half-diamond in telocentric chromosomes with the N-banding technique demonstrate that breakage occurred in the middle of the kinetochore. The stability of barley telocentrics (Singh and Tsuchiya, 1981b) and the lack of secondary trisomics in the progenies of monotelotrisomics (Singh and Tsuchiya, 1977) indicate that stability of barley telocentric chromosomes does not depend upon completeness of the kinetochore. As a matter of fact, there have been at least two cases of stable telocentric chromosomes in the natural population. Tsunewaki (1963) observed them in tetraploid wheat (2n = 28), and Strid (1968) recorded them in *Nigelle doerfleri* (2n = 12). Stable telocentric chromosomes were also produced in grasshoppers (*Myrmeleotettix maculatus*) following centric misdivision (Southern, 1969).

4. Transmission of the Extra Chromosome in Telotrisomics

The transmission of an extra telocentric chromosome depends upon the length of the telocentric chromosome, its meiotic behavior, size of seeds, genetic background, and also sometimes depends on environmental growing conditions. An average transmission of 34.6% (range; 30.5% — Triplo 4L to 38.4% — Triplo 1S) of the 7 telocentric chromosomes of barley through selfing was recorded (Table 7.46). The average female transmission of 11 telocentric chromosomes in barley was 30.7%, with less variation (range) among 11 types than in selfed progenies (Singh, 1993). These values are higher than those observed by Tsuchiya (1967) in primary trisomics. These results clearly demonstrate that transmission rates vary depending upon genetic and environmental conditions.

The female transmission rate of telocentric chromosomes in the progenies of telotrisomics of rice ranged from 28.6 (Triplo 2L) to 47.5% (Triplo 9S) (Table 7.47). It is evident that telocentrics for the short arm transmitted in greater frequencies than telocentrics for the long arm (Singh, Multani, and Khush, 1996).

The transmission of the barley telocentric chromosome has been observed to be a little higher, 34.6% (mean of seven telotrisomics), in a hybrid genetic background (Table 7.46). Similar results have been reported by Khush and Rick (1968b) in tomato telotrisomics.

		Chromsome	Numbers (2 <i>n</i>)		Number of	Plants
Triplo	14	14 + 1 telo	14 + 2 telo	15	Others	Studied
1L	67.4	31.4	_	1.2		86
1S	56.4	38.4	0.7	3.6	0.5	426
2L	60.9	37.5	0.3	1.3	0.1	1179
2S	59.8	37.6	1.1	1.0	0.6	797
3L	64.6	34.5	0.1	0.6	0.2	1935
4L	67.9	30.5	0.7	0.9	0.1	1100
5L	66.7	32.3	_	1.0	—	96
Average (total)	63.4	34.6	0.4	1.4	0.2	(5619)

Table 7.46 Female Transmission Rate (%) of the Extra Telocentric Chromosome in Selfed F₂ Population in Seven Telotrisomics of Barley

Source: From Singh, R.J. and Tsuchiya, T., Z. Pflanzenzüchtg. 78, 327–340, 1977. With permission.

The infrequent male transmission of the extra telocentric chromosome has been observed in maize (Rhoades, 1936), tomato (Khush and Rick, 1968b), barley (Singh and Tsuchiya, 1977; Siep, 1980), and rye (Melz and Schlegel, 1985). The range of male transmission of a barley telocentric chromosome was 0.0 (Triplo 2S, 5L) to 6.7% (Triplo 5S). Siep (1980) recorded 10.53% in Triplo 6S. By contrast, a high rate of male transmission of the extra telocentric chromosome 8S and 9S was recorded in the rice (Table 7.47). The occurrence of ditelotetrasomic plants in the progenies of telotrisomics of barley (Fedak and Helgason, 1970a; Singh and Tsuchiya, 1977; Tsuchiya and Wang, 1991), rice (Singh, Multani, and Khush, 1996), and also in *Datura* (Blakeslee and Avery, 1938), indicates that sometimes male gametes with an extra telocentric chromosome can take part in fertilization. It is generally believed that unbalanced gametes. If delayed pollination occurs, chances may be better for unbalanced gametes to take part in fertilization.

Related primary trisomics have been obtained in the progenies of tomato (Khush and Rick, 1968b), barley (Singh and Tsuchiya, 1977) and rice (Singh, Multani, and Khush, 1996). Rhoades (1936) obtained a secondary trisomic in the progenies of maize telotrisomics. In barley, secondary trisomic plants have not been isolated so far in the progenies of telotrisomics (Singh and Tsuchiya, 1977).

5. Genetic Segregation in Telotrisomics

Telotrisomics have been used in genetic linkage studies in several plant species, such as maize (Rhoades, 1936, 1940), diploid wheat (Moseman and Smith, 1954), tomato (Khush and Rick, 1968b), barley (Fedak, Tsuchiya, and Helgason, 1972; Tsuchiya, 1971b, 1972a,b; Tsuchiya and

Triplo	Total Plants	Disomics	Telotrisomics	Related Trisomics
1S	80	60.0	38.8	1.2
2L ^a	56	67.8	28.6	3.6
3L ^a	150	68.7	29.3	2.0
5L	136	66.2	30.9	2.9
8S ^b	324	48.1	40.1	0.0
9S⁵	80	32.5	47.5	0.0
10S	105	65.7	31.4	2.9

 Table 7.47
 Female Transmission Rate (%) of Extra Telocentric Chromosomes in Selfed or Back-Cross Progenies of Seven Telotrisomics of Rice

^a Back-cross progeny.

^b About 12 and 20% plants, respectively, contained a pair of extra telocentric chromosomes.

Source: From Singh, K., Multani, D.S., and Khush, G.S., *Genetics*, 143, 517–529, 1996. With permission.

Singh, 1982), rice (Singh, Multani, and Khush, 1996), hexaploid wheat (Sears, 1962, 1966a), cotton (Endrizzi and Kohel, 1966), and oat (McGinnis, Andrews, and McKenzie, 1963). These chromosomes are useful in associating a gene with a particular arm of a chromosome. With the use of multiple marker stocks, centromere position and gene sequence on a linkage map can be determined (Rhoades, 1936, 1940; Khush and Rick, 1968b; Reeves, Khush, and Rick, 1968).

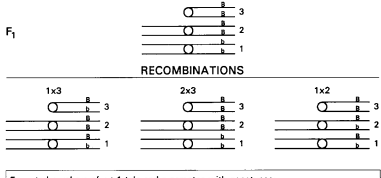
The principle of linkage studies with telotrisomics is different from that in simple primary trisomics. Generally, mutants are crossed as male parents, and telotrisomics are used as female parents. Telotrisomics are identified cytologically and morphologically in F_1 hybrids. Telotrisomics and one diploid sib (control) are saved and are allowed to self-pollinate. The chromosome number of each F_2 plant should be counted in order to separate disomics, telotrisomics, and other chromosomal types. Segregation ratios are calculated separately for the disomic and monotelotrisomic portions of the F₂ populations. Because the extra chromosome is derived from one arm of a standard chromosome, the genetic ratios are modified depending upon the type of segregation and transmission rate of the extra chromosome (Reeves, Khush, and Rick, 1968). If a gene is not located on a particular arm of a chromosome, a disomic ratio is obtained for the disomic and trisomic portions (3:1::3:1), and the result is known as a noncritical combination. If a gene is on the telocentric chromosome (critical combination), no recessive homozygotes will be obtained in the telotrisomic portion, although the diploid portion will show a disomic ratio (3:1::4:0), provided that the gene is close to the centromere. When both disomic and trisomic portions are combined, a 7:1 ratio is expected in random chromosome segregation, with a 50% female transmission rate of the telocentric chromosome. This ratio is further narrowed to 5:1 when the female transmission rate approaches 33.3%. Because the rate of pollen transmission of the extra telocentric chromosome is extremely low, male gametes with the extra chromosome have been considered nonfunctional.

Sometimes, recessive homozygotes are obtained in the telotrisomic fraction. This is due to random chromatid crossing over, and it happens when a gene is far from the centromere. It can be explained as follows. Let us assume Chromosome 1 has a recessive gene *b* that came from the male parent and that Chromosomes 2 (complete) and 3 (telocentric) came from the female parent with dominant genes *BB*; Chromosomes 1, 2, and 3 designate homologous chromosomes. There are three possibilities of recombinations: 1×3 ; 2×3 ; 1×2 (Figure 7.46). At the end of meiosis, gametes in a genotypic ratio of 11BB:12Bb:1bb (*n* + 1 telo) and 7BB:5bb (*n*) are expected to be generated. If the female transmission of the telocentric chromosome is 50%, a 119B::25bb (4.76*B*::1*bb*) ratio in disomic portion, 283B::5bb (56.6*B*::1*bb*) ratio in telotrisomic portion, and a total ratio of 402B::30bb (13.4*B*::1*bb*) are expected. However, with 33.3% female transmission of the extra chromosome, there will be no change in the ratio of disomic and trisomic sections, but segregation in the total population should be 521B::55bb, and the ratio should be 9.47B::1bb. That is, 9.55% of the F₂ population should be recessive homozygotes (*b*), while 6.95% recessive homozygotes are expected when the female transmission rate of the extra telocentric chromosome is 50%.

Thus, the genetic ratio is modified from 3:1 to 5:1 or 7:1 (random chromosome) and 13.4:1 or 9.47:1 (random chromatid) in telotrisomic analysis when a gene is located on the telocentric chromosome. This modification, however, depends upon the transmission rate of the telocentric chromosome through the female and the distance of a gene from the centromere.

Linkage studies with telotrisomics for both arms of a chromosome are desirable whenever these stocks are available. This provided a definite association of a gene with a particular arm of Chromosome 2 of barley (Tsuchiya and Singh, 1982). The four genes f (= lg), gs6, v, and gs5 were analyzed with Triplo 2L and 2S. The genes f (= lg) and gs6 showed disomic ratio with Triplo 2L but a trisomic ratio with Triplo 2S. Likewise, genes v and gs5 exhibited a trisomic ratio with Triplo 2L but a disomic ratio with Triplo 2S. These results suggest that genes v and gs5 are on the long arm, and f and gs6 are on the short arm of Chromosome 2 (Table 7.48).

The gene order can be reversed on the linkage map from telotrisomic analysis. In barley, the genes *gs5* and *e* were assigned to the short and long arm of Chromosome 2, respectively (Robertson,



Anaphase		n = 7+1 teio		n =	= 7
separation	BB	Bb	bb	В	b
1x3		_			
1.3 vs. 2	1	2	1	2	0
1 vs. 2.3	2	2	0	1	1
2x3					
1.3 vs. 2	0	4	0	2	0
1 vs. 2.3	4	0	0	0	2
1x2					
1.3 vs. 2	2	2	0	1	1
1 vs. 2.3	2	2	0	1	1
Total $n=7+1$ telo	11	12	1		
n=7				7	5

 F_2

Expected genotypic and phenotypic ratios:

Q O™	7B	5b	
11 BB	77 BBB	55 BBb	Trisomic Portion:
12 Bb	84 BBb	60 Bbb	Segregation ratio 283 : 5 or 56.6 : 1
1 bb	7 Bbb	5 bbb	
7 B	49 BB	35 Bb	Disomic Portion:
5 b	35 Bb	25 bb	Segregation ratio 119:25 or 4.76:1

Figure 7.46 Maximum equational chromatid segregation telotrisomic analysis. (Modified from Reeves, A.F., Khush, G.S., and Rick, C.M., *Can. J. Genet. Cytol.*, 10, 937–940, 1968.)

1971). From telotrisomic analysis, gene gs5 showed the trisomic ratio and gene e the disomic ratio with Triplo 2L. This indicates that the gene order should be reversed.

With the use of multiple marker stocks in telotrisomic analysis, the precise gene order on a linkage map can be obtained (Rhoades, 1940; Khush and Rick, 1968b; Tsuchiya and Singh, 1982). Khush and Rick (1968b) studied four genes (r, wf, rv, sf) with Triplo 3L of tomato. The genes r and wf showed disomic ratio, while rv and sf were trisomic; r and wf should be located on 3S, and rv and sf should be located on 3L (Table 7.49). The orientation of the linkage group, reading from the end of the short arm, was suggested as r-wf-sy-ru-centromere-rv-sf (Khush and Rick, 1968b). Tsuchiya and Singh (1982) used a multiple marker stock cu2-uz of barley for telotrisomic analysis. From conventional linkage analysis, gene cu2 was located farther from the centromere than gene uz. With Triplo 3L, both genes showed a trisomic ratio. However, more recessive homozygotes were obtained for gene cu2 than for uz in the diploid portion of an F₂ population, indicating cu2 is closer to the centromere than uz in the long arm of Chromosome 3.

	Marker		2x		8	2x + 1 telo			Total			χ²	
Triplo	Triplo Genes	Aª	ø	Total	A	a	Total	A	a	Total	3:1	5:1	7:1
2L	f(= lg)	62	9	68	30	11	41	92	17	109	5.13	0.09	
2L	gse	69	26	95	30	14	44	66	40	139	1.05		
2L	7	122	50	172	124	0	124	246	50	296	10.37	0.00	5.21
2L	gs2	55	14	69	30	0	30	85	14	66	6.23	0.46	0.24
2S	f(= lg)	95	16	111	71	0	71	166	16	182	25.51	8.12	2.29
2S	gse	50	ო	53	41	0	41	91	ო	94	23.84	12.29	7.44
2S	7	121	26	147	56	1	67	177	37	214	6.77	0.06	4.59
2S	gs2	56	80	64	19	£	24	75	13	88	4.91	0.23	0.41
^a A = norm	al: a = mutant	t.											

Table 7.48 F_2 Segregation Results of Genes f (= lg), gs6, v, gs5 with Triplo 2L and Triplo 2S of Barley

A = normal; a = mutant.

Source: From Tsuchiya, T. and Singh, R.J., Theor. Appl. Genet, 61, 201-208, 1982. With permission.

Genes Total A ^a a	2X		6	2x + 1 telo	
	% Recessive	A	a	% Recessive	χ² (3:1)
	28.7	39	13	25.0	0.00
<i>i</i> f 184 99 33	25.0	40	12	23.1	0.10
v 180 102 18	15.0	59	-	1.6	17.41
f 180 104 16	13.3	59	-	1.6	17.41

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Source: From Khush, G.S. and Rick, C.M., Cytologia, 33, 137–148, 1968b. With permission.

Sometimes, a smaller number of recessive homozygotes than the expected 25% for a disomic ratio are obtained in the F_2 population, even if the gene is not located on the telocentric chromosome. A wrong conclusion can be drawn if simply a chi-square value is calculated for a total population without a chromosome count. The results obtained in barley for genes *als*, *yst2*, *f2*, and *x*_s on Chromosome 3 are such examples. Chi-square values for these genes fit the trisomic ratio (Table 7.50). However, when chromosomes were studied in the F_2 , a high proportion of trisomic plants was found to consists of recessive homozygotes. Their frequencies were too high to consider them as a trisomic ratio, even if the possibilities of random chromatid crossing over were considered. From the results, it seems to be essential to count the chromosome number of every F_2 plant in telotrisomic analysis. There is an alternative to testing the trisomic or disomic ratios. The chromosome counts of recessive homozygotes only will provide fairly accurate results regarding the segregation ratios, disomic or trisomic.

Singh, Multani, and Khush (1996) studied the segregation of 26 genes with seven telotrisomics (Triplo 1S, 2L, 3L, 5L, 8S, 9S, 10S) of rice. For example, of the five genes of chromosome 1 tested with Triplo 1S, marker gene d-18 showed a trisomic ratio, suggesting that this gene is on the short arm of chromosome 1, while genes *chl*-6, *spl*-1, *z*8, and *gf2* segregated in a disomic fashion, suggesting that these genes are on the long arm of chromosome 1 (Table 7.51). Based on segregation of markers in telotrisomics and secondary trisomics centromere position on eight linkage groups and orientation of 10 linkage groups of rice were established (Singh, Multani, and Khush, 1996).

Telotrisomic analysis by the back-cross procedure is useless, because clear-cut segregation ratios are not obtained, regardless of parental genotypes or segregation of chromosomes. Because monotelotrisomics are fertile enough to generate F_2s , it is not advisable to conduct back crossing.

E. Acrotrisomics

1. Introduction

Trisomic plants carrying an extra acrocentric chromosome are designated acrotrisomics. In barley, six acrotrisomics (1L^{1S}, 3L^{3S}, 4L^{4S}, 5S^{5L}, 6S^{6L}, 7S^{7L}) have been studied and identified morphologically and cytologically, and have been used in locating genes physically in chromosomes (Tsuchiya, Shahla, and Hang, 1986).

2. Sources of Acrotrisomics

Acrotrisomics in barley have originated from the progeny of related primary trisomics ($4L^{4S} =$ Triplo 4), telotrisomics ($1L^{1S} =$ Triplo 1S), triploid hybrids ($3L^{3S}$), and unrelated telotrisomics ($5S^{5L} =$ Triplo 1S). In sugar beets, Romagosa et al. (1985) identified an acrotrisomic $9S^{9L}$ in the progeny of a primary trisomic, Triplo 9.

Acrocentric chromosomes in barley may have originated by a breakage in chromosome arm(s), preferably adjacent to heterochromatic regions, and followed by the healing of the broken ends or fusion with a second broken end. If the telomere concept of Muller (1940) is accepted, all the breakage should have at least two breaks with an intact telomere. However, according to McClintock (1941c), broken ends of meiotic chromosomes will heal when chromosomes enter into sporophytic tissues. In barley and also in sugar beets, acrotrisomics originated in the progenies of primary trisomics, telotrisomics, and triploids. It is reasonable to assume that a single breakage occurred at one site $(3L^{38}, 5S^{5L})$ or at two sites $(1L^{18}, 4L^{4S})$ in the extra chromosome, which went through a breakage-fusion-bridge cycle, and the broken end(s) healed as suggested by McClintock (1941c).

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HL)	
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Triplo	
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f 15 (
Results o	
Segregation	
50 F ₂	
Table 7.5	

Genes A^a a $cu2$ 140 18 uz 148 10 uz 148 10 wst 53 4 als 41 2 $gs2$ 49 9 zb 99 11 $cer-zn^{348}$ 61 12 yst 107 27 yst 38 21	a 800		İ		5		lotal			*	
140 148 53 41 49 61 61 38	18 10	Total	A	a	Total	٩	a	Total	3:1	5:1	7:1
148 53 49 99 61 38	10	158	54	0	54	194	18	212	30.81	10.20	3.11
53 41 49 61 107 38		158	54	0	54	202	10	212	46.52	21.79	11.74
41 49 61 38 38	4	57	44	0	44	97	4	101	23.84	14.95	6.73
49 99 61 38	2	43	38	0	38	79	0	81	18.62	11.75	7.45
99 61 38	6	58	45	0	45	94	6	103	14.52	4.66	1.84
61 107 38	11	110	43	0	43	142	11	153	25.88	9.90	3.94
107 38	12	73	52	÷	53	113	13	126	14.49	3.66	0.55
38	27	134	50	14	64	157	41	198	1.94		
	21	59	25	7	32	63	28	91	1.61		
61	œ	69	29	9	35	06	14	104	7.39	0.77	0.08
163	25	188	114	ო	117	277	28	305	40.71	12.27	3.07
46	8	54	29	4	33	75	12	87	5.82	0.52	0.14
65	6	74	42	ო	45	107	12	119	14.12	3.71	0.63
37	6	46	17	7	24	54	16	70	0.17		

^a A = normal; a = mutant.

Source: From Tsuchiya, T. and Singh, R.J., Theor. Appl. Genet., 61, 201–208, 1982. With permission.

Marker		2x		¢,	2x + 1 telo	_		Total		χ ² (2x)
Genes	A	ø	Total	۷	a	Total	A	a	Total	3:1
1-18	325	122	447	102	0	102	427	122	549	1.25
chl-6	262	105	367	62	32	94	324	137	461	2.25
9-1d:	91	29	120	42	=	53	133	40	173	0.04
8-	186	56	242	69	29	98	255	85	340	0.45
gf-2	116	54	170	24	6	33	140	63	203	4.12

CHROMOSOMAL	ABERRATIONS -	– NUMERICAL	. CHROMOSOME CHANGES
0111101110000111712			

3. Identification of Acrotrisomics

a. Morphological Identification

Morphologically, acrotrisomics 1L^{1S}, 3L^{3S}, 4L^{4S}, and 5S^{5L} were similar to their corresponding primary trisomics and telotrisomics for the long arm of chromosomes 1L, 3L, 4L, and 5L. Acrotrisome 1L^{1S} plants possess long, narrow leaves, many tillers, rather small spikes, long awns, and narrow seeds. Acrotrisomic 3L^{3S} plants show pale green color, revoluted leaves with tips extremely twisted, prominent hairs on the surfaces of leaf blades, compact spike, slightly shorter awns, and high pollen and ovule sterility. Acro 4L^{4S} plants carry short, thick culms, slightly shorter, dark-green revoluted leaves, and short and compact spikes. They resemble the Triplo 4 (Robust). Although acro 5S^{5L} is deficient of the 60% of distal portion of the long arm of Chromosome 5 and carries a complete short arm, morphologically, acro 5S^{5L} plants are similar to Triplo-5 and Triplo-5L. This suggests that the 40% proximal segment of the long arm has almost the same effect on plant morphology as the complete long arm.

b. Cytological Identification

Precise identification of acrotrisomic plants in barley has been possible based on combining the aceto-carmine and Giemsa N-banding techniques (Singh and Tsuchiya, 1982a,b; Tsuchiya et al., 1984; Shahla and Tsuchiya, 1986, 1987). Shahla and Tsuchiya (1987) reported that in acro $1L^{1S}$, long and short arms were 37.5% and 73.0% deficient, respectively. Acro $3L^{3S}$ carried an intact long arm (3L) without an apparent deficiency, while 3S had 77.8% deficiency (Figure 7.47A,B). The proximal band in the 3S was approximately 33%, and the breakage occurred in the proximal heterochromatic band (Tsuchiya et al., 1984). Acrocentric $4L^{4S}$ was found to be deficient for both arms, 4L = 31.7% and 4S = 59.3% (Tsuchiya et al., 1984). In acro $5S^{5L}$, the short arm was intact, while 60% of distal 5L was deficient (Shahla and Tsuchiya, 1986).

The only acrotrisomic, acro 9S^{9L}, reported in sugar beets carried a complete short arm, but 45% of the distal portion of 9L was missing (Romagosa et al., 1985).

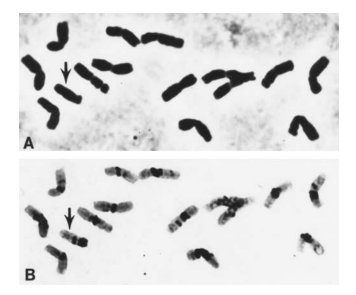


Figure 7.47 Aceto-carmine (A) and Giesma N-banded (B) acrotrisomics of a plant with 2n = 14 + 1 acro 3L.³⁵ Arrow shows an acro 3L³⁵. (From Singh, R.J., unpublished results.)

Acrotrisomic	Metapha Configu	• • •	Anapha	ase-I (%) C Migratio	hromosome on	
Types	1III + 6II	1I + 7II	8–7	7–1–7	7 + 1 – 1 + 7	Authors
1L ^{1S}	71.0	29.0	73.7	22.2	4.1	Shahla and Tsuchiya, 1987
3L ^{3S}	66.9	33.1	77.8	13.4	8.8	Singh, unpublished data
4L ^{4S}	65.2	34.8	81.0	14.7	3.5	Hang, 1981
5S⁵L	71.0	29.0	53.7	29.9	16.4	Shahla and Tsuchiya, 1986

Table 7.52 Meiotic Chromosome Behavior of Acrotrisomics of Barley

Although Giemsa C- and N-banding techniques have facilitated the precise identification of individual chromosomes of barley and also various aneuploid stocks, location of correct breakage points has not been possible (Singh and Tsuchiya, 1982a,b). Barley pachytene chromosomes have not been utilized to identify deficiencies and translocations, because they lack morphological landmarks such as euchromatin and heterochromatin differentiation (Singh and Tsuchiya, 1975b). Utilizing pachytene chromosomes, break points have been located precisely in the aneuploids of tomato (Khush and Rick, 1968a,b; Khush, 1973) and maize (Rhoades, 1955).

c. Meiotic Behavior of Acrocentric Chromosome

The extra acrocentric chromosome pairs with normal homologue at diakinesis and metaphase-I forms 1III + 6II in a majority of sporocytes (Table 7.52) or remains as a univalent, giving the 7II + 1I configuration (Figure 7.48A). The trivalents may be V-shaped, rod-shaped, ring-and-rod shaped (Figure 7.48B), or Y-shaped. As expected from metaphase-I configurations, 8–7 chromosome migration predominates in a large number of sporocytes (range of 53.7 to 81.0%). Sometimes, an acrocentric chromosome lags at the equatorial plate, while the remaining chromosomes have already reached their respective poles. Occasionally, acrocentric chromosomes divide longitudinally, giving 8–8 chromosome separation, though the frequency is low (Table 7.52). The lagging chromosomes will be eliminated before reaching the poles in the first division, and if entered into dyads, will be eliminated in the second meiotic division.

4. Transmission of the Extra Chromosome in Acrotrisomics

Transmission of acro $1L^{15}$, $3L^{38}$, $4L^{48}$, and $5S^{5L}$ in selfed populations (F₂) was 29.4%, 33.3%, 30.5%, and 37.6% respectively (Tsuchiya et al., 1984; Shahla and Tsuchiya, 1986). The transmission rate is lower in acrotrisomics × diploid crosses. For example, acro $1L^{18}$ showed 25.4% transmission (Shahla and Tsuchiya, 1987), acro $4L^{48}$ transmitted in 22.1% of plants (Hang, 1981),

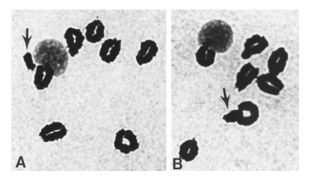


Figure 7.48 Chromosome configurations at diakinesis in plants with 2n = 14 + 1 acro $3L^{3S}$. (A) 7II + 1I (arrow shows an univalent). (B) 6II + 1III (ring- and rod-shaped trivalent, arrow). (From Singh, R.J., unpublished results.)

while 5S^{5L} was found in 35.8% of plants (Shahla and Tsuchiya, 1986). Acrocentric chromosomes may be transmitted through pollen in low frequencies, 1.3% (4L^{4S}) and 9% (5S^{5L}).

5. Genetic Segregation in Acrotrisomics

The theoretical segregation ratios in acrotrisomic analysis are the same as those observed for telotrisomic analysis (Reeves, Khush, and Rick, 1968). All of the available four acrotrisomics $(1L^{15}, 3L^{38}, 4L^{48}, 5L^{58})$ of barley have been utilized in the physical localization of genes (Tsuchiya et al., 1984; Shahla and Tsuchiya, 1987). Genetic segregation ratios in F₂ population between acro $3L^{38}$ and four genes (a_c , yst2, a_n , and x_s) are cited as an example. These genes are located on 3S. Two genes (a_c and yst2) showed a trisomic ratio, suggesting that these genes are located within the proximal segment consisting of 22.2% of the heterochromatic segment of the short arm of the acro $3L^{38}$ (Table 7.53).

Six morphological markers were analyzed with acro $5S^{5L}$ of barley by Shahla and Tsuchiya (1986). The segregation results are summarized in Table 7.54. Genes *fs*2, *g*, and *f3* segregated in a trisomic ratio with acro $5S^{5L}$, and no homozygous recessives were observed among acrotrisomic progeny. Genes *f7*, *trd*, and *int-a¹* segregated in a disomic ratio, suggesting that these genes are located on the missing region of the long arm of Chromosome 5.

Acrotrisomic linkage mapping is an excellent tool to locate genes precisely in a particular region of a chromosome. However, this method has a serious problem, because acrocentric chromosomes originate by breakage in chromosome arm(s) and healing of the broken ends. The mode of production of deficiencies is of two types, shown in Figure 7.49. A terminal deficiency needs only one break in the arm (Figure 7.49A), but an intercalary segmental deficiency needs at least two breaks. An acentric fragment gets eliminated, and a telomere attaches to the broken end of the acrocentric chromosome (Figure 7.49B). Precise physical location of genes may not be possible if genes are located in the attached telomeric segment, and, furthermore, if an intercalary deficiency (deletion) is small and cannot be detected by the Giemsa banding technique.

Examination of pachytene chromosomes should be attempted.

		2 <i>x</i>		2	x + 1 ac	ro		Total	
Genes	Aa	а	Total	Α	а	Total	Α	а	Total
Albino (a _c)	45	14	59	32	0	32	78	14	92
Yellow streak 2 (yst2)	84	12	96	34	0	34	121	12	133
Albino (a _n)	49	17	66	32	9	41	81	26	107
Xantha (x_s)	59	15	74	30	12	42	89	27	116

Table 7.53 Genetic Segregation Ratios in F_2 Population of Acrotrisomics $3L^{3S} \times$ Four Recessive Genetic Stocks

a + = normal; a = mutant.

Source: From Tsuchiya, T. et al., Theor. Appl. Genet., 68, 433-439, 1984. With permission.

Table 7.54 Genetic Segregation Ratios in F₂ Population of Acrotrisomics 5S^{5L} × Six Recessive Genetic Markers

		2 <i>x</i>		2	x + 1 ac	ro		Total	
Genes	Aa	а	Total	Α	а	Total	Α	а	Total
Fragile stem 2 (fs2)	66	13	79	63	0	63	129	13	142
Golden (g)	89	25	114	30	0	30	119	25	144
Chlorina 3 (f3)	77	16	93	53	0	53	130	16	146
Chlorina 7 (f7)	59	19	78	51	11	62	110	30	140
Third outer glume (trd)	65	26	91	42	16	58	107	42	149
Intermediate spike (int-a1)	68	23	91	43	15	58	111	38	149

^a A = normal; a = mutant.

Source: Shahla, A. and Tsuchiya, T., Can. J. Genet. Cytol., 28, 1026–1033, 1986. With permission.

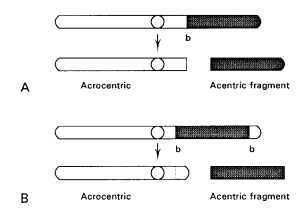


Figure 7.49 The mode of origin of an acrocentric chromosome. (A) Single break at Point b produces acrocentric chromosome with healed broken end and acentric fragment. (B) Two breaks at Points b, with the result of an acentric chromosome and the original telomere attached to form an acrocentric chromosome. (From Tsuchiya, T. et al., *Theor. Appl. Genet.*, 68, 433–439, 1984. With permission.)

F. Compensating Trisomics

1. Introduction

In a compensating trisomic, a missing normal chromosome is replaced either by two tertiary chromosomes or by a secondary and a tertiary chromosome. Blakeslee (1927) was the first who reported a compensating trisomic, "Nubbin," where a missing Chromosome 1.2 was compensated by tertiary Chromosomes 1.9 and 2.5, and the trisomic was designated by the chromosome formula 2n - 1.2 + 1.9 + 2.5. Compensating trisomics have been reported in *Datura stramonium* (Blakeslee, 1927; Avery, Satina, and Rietsema, 1959), *Triticum monococcum* (Smith, 1947), *Lycopersicon esculentum* (Khush and Rick, 1967a), *Pennisetum glaucum* (Saini and Minocha, 1981), and *Secale cereale* (de Vries and Sybenga, 1989).

2. Sources of Compensating Trisomics

The compensating trisomic type Nubbin in *Datura stramonium* was isolated among the progeny of a plant that was exposed to radium (Blakeslee, 1927). Furthermore, Avery, Satina, and Rietsema (1959) summarized the results of seven compensating trisomics in *D. stramonium*; all were produced experimentally. Khush and Rick (1967a) isolated three compensating trisomics in tomato in the progenies of tertiary or translocated chromosomes. Similarly, Saini and Minocha (1981) reported a compensating trisomic in pearl millet that was isolated from the progeny of a multiple interchange trisomic.

According to Avery, Satina, and Rietsema (1959), compensating trisomics can be isolated in several ways. One way is by crossing a secondary trisomic, such as 2n + 3.3 as a female with an interchange heterozygote of the formula 2n = 3.21 and 2n + 4.22 as a male. The chromosome end arrangement in F₁ would be 3.3-3.21-21.22-22.4-4.3. In the F₂ generation, they found 68 plants were normal diploid, two plants were 2n + 3.3, two plants were compensating trisomic, 2n - 3.4 + 3.3 + 4.22, and four plants were not identified.

De Vries and Sybenga (1989) isolated four telocentric-tertiary compensating trisomics in rye from four different reciprocal translocations and three different telocentrics.

3. Identification of Compensating Trisomics

a. Morphological Identification

Morphologically, depending upon the chromosome involved, compensating trisomic types exhibit morphological features of the corresponding trisomics. The tomato compensating trisomics 2n –

3S.3L + 3L.3L + .3S and 2n - 3S.3L + 3S.3S + 3L.3L expressed several morphological traits of Triplo 3 (Khush and Rick, 1967a).

b. Cytological Identification

The true identification of compensating trisomics is possible by observing the meiotic chromosome pairing pattern. Chromosome configurations during meiosis depend upon the types of compensating trisomics. Khush and Rick (1967a) identified three compensating trisomics of tomato at pachynema. In diisocompensating trisomics (2n - 3S.3L + 3S.3S + 3L.3L and 2n - 7S.7L + 7S.7S + 7L.7L), isochromosomes paired internally, leaving normal chromosomes (3S.3L; 7S.7L) as univalents. In the monoteloisocompensating trisomic, 2n - 3S.3L + 3L.3L+ .3S, a trivalent configuration was more frequent. In a ditertiary compensating trisomic, 2n- 1S.1L + 1S.2L + 3S.1L of pearl millet, Saini and Minocha (1981) recorded the maximum association of seven chromosomes (1VII + 4II) in 14.4% of sporocytes. The other chromosome configurations observed were 1V + 5II (31.8%), 1III + 6II (28.8%), 1IV + 1III + 4II (10.6%), 1IV + 5II + 1I (6.1%), and 7II + 1I (6.1%). At anaphase-I, an 8–7 chromosome migration predominated (91.7%).

4. Transmission of the Extra Chromosome in Compensating Trisomics

In seven compensating trisomics of *Datura stramonium*, one normal chromosome was replaced by two tertiary chromosomes. As expected, compensating trisomics threw diploids, parent types, related tertiaries and primaries, and also unrelated chromosomal types. A tomato compensating trisomic type (2n - 3S.3L + 3L.3L + .3S) transmitted diploids (56.3%), compensating trisomic-parental types (40.5%), telotrisomics (2.7%), and diisocompensating trisomics (0.5%) (Khush and Rick, 1967a).

5. Genetic Segregation in Compensating Trisomics

Compensating trisomics have been used to locate marker genes to their respective chromosomes in *Datura stramonium* (Avery, Satina, and Rietsema, 1959), *Triticum monococcum* (Smith, 1947), and *L. esculentum* (Khush and Rick, 1967a). Moreover, markers can be located in a particular arm of a chromosome, because certain compensating types throw secondary, tertiary, and telotrisomics in their progeny.

A disomic ratio (3:1::3:1) is observed if a gene is not located on the compensated chromosome. In case a marker gene is located on the compensated chromosome, a compensating trisomic ratio (0:1::1:0) is expected that is a completely different ratio from ratios observed in other trisomics (Khush, 1973).

Generally, compensating trisomics are made heterozygous for marker genes to be studied, and the F_1 heterozygous compensating trisomic is crossed onto the diploid recessive homozygote. It is expected that all the *n* chromosome male gametes will carry a recessive allele, with the exception of random chromatid crossing over, and all 2*x* progenies will be recessive homozygote. Utilizing this procedure, Avery, Satina, and Rietsema (1959) located a gene for albino-11 in Chromosome 11.12. The procedure is as follows: A compensating trisomic for chromosome 1.2 (2n - 1.2 + 1.9 + 2.5) was hybridized by a recessive homozygote mutant albino-11 (albino-11 grafted on green stock produced flowers and capsules). The F_1 compensating trisomic plants were crossed onto albino-11, and the progeny was segregated into 199 albino seedlings and 36 normal. The occurrence of 36 normal plants indicated either that the gene albino-11 is located away from the centromere and by double reduction, *n* gametes with normal alleles were produced, or that the compensating chromosomes were transmitted through male gametes at a low frequency.

A similar approach was used by Khush and Rick (1967a) to assign a marker gene ru to the short arm of Chromosome 3 of tomato. A compensating trisomic, 2n - 3S.3L(+) + 3L.3L(-) + .3S(+)

with normal + allele, was crossed by a recessive mutant gene ru [3S.3L(ru) + 3S.3L(ru)]. Two kinds of female gametes [n = 12, 3S.3L(+); n = 11 + 3L.3L + 3S(+)] and only one kind of male gamete [n = 11 + 3S.3L(ru)] are expected. In the F₁ diploid (2n = 24 + ru) and compensating trisomic [2n - 3S.3L(ru) + 3L.3L + .3S(+)] carrying ru gene in heterozygous state are expected; generally, diploids are discarded. The elimination of telo 3S(+) in somatic tissues resulted in chimaeric condition and expression of the ru character. This indicated that gene ru is located in the short arm of Chromosome 3.

Various types of an uploids in the progeny of a test cross between a compensating trisomic for Chromosome 3 and the diploid marker stock (ru) are expected:

- 1. Diploid: [3S.3L (*ru*) + 3S.3L (*ru*)].
- 2. Compensating trisomic (parental type): [3S.3L(ru) + 3L.3L(-) + 3S(+)].
- 3. Secondary trisomic: [3S.3L (*ru*) + 3S.3L (*ru*) + 3L.3L (–)].
- 4. Telotrisomic: [3S.3L (ru) + 3S.3L (ru) +.3S (-)].
- 5. Compensating trisomic (double isochromosome): [3S.3L (ru) + 3S.3S (+) + 3L.3L (-)].

G. Novel-Compensating Diploid

1. Introduction

In compensating diploids, one normal chromosome is replaced by two telocentric chromosomes representing both arms, or four telocentric chromosomes compensating for a pair of normal homologue. The compensation results in genetically balanced plants.

2. Sources of Novel-Compensating Diploid

The occurrence of compensating diploid plants is rare in diploid species. Such plants have been reported in *Nigella doerfleri* (Strid, 1968), *Hordeum vulgare* (Tsuchiya, 1973; Singh and Tsuchiya, 1981c, 1993), *Pennisetum glaucum* (Pantulu and Narasimha Rao, 1977; Manga et al., 1981), and *Secale cereale* (Zeller, Lintz, and Kunzmann, 1982; Melz and Winkel, 1986). Tsunewaki (1963) identified a tetraploid strain of Emmer wheat from Tibet with 2n = 26 + 4 telo (15 pairs). In *Nigella doerfleri*, Strid (1968) observed about half of the plants in a natural population with 2n = 14 (10 + 4 telocentrics) originating from the Island of Los, Kikladhes (Greece). The diploid chromosome constitution of *N. doerfleri* is 2n = 12.

Tsuchiya (1973) found a plant with 2n = 13 + 2telos (1L+1S) in the progeny of a cross between diploid (2n = 14) and 2n = 14 + 1telo1S of barley. Singh and Tsuchiya (1981c) isolated a plant with 2n = 13 + 1acro3L^S + 1telo3S in an F₂ population of a cross 2n = 14 + 1acro3L^S × yellow streaks (*yst2*). In 2n = 13 + 1telo1L + 1telo1S plants, 1telo1L and 1telo1S compensate for one normal Chromosome 1, and similarly, in plants with 2n = 13 + 1acro3L^S + 1telo3S, one acro3L^{3S} and one telo3S replace a normal Chromosome 3. Hang, Burton, and Satterfield (1998) isolated and identified barley compensating (partial) diploid (2n = 13 + 1acro6S^L) from the progenies of 2n = 14 + 1acro6S.^L

The origin of plants with $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ has been assumed as follows. This particular plant originated in the F₂ population of $2n = 14 + 1 \operatorname{acro3L^{3S}} \times yst2$. The misdivision of a normal Chromosome 3 may have occurred at anaphase-I in a sporocyte in one of the spikelets of an F₁ acrotrisomic $3L^{3S}$ plant, and by chance, seven chromosomes moved to one pole and $6 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ (telo3S is the product of misdivision) moved to the other pole. Thus, one gamete of this spikelet contained $6 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ is genetically balanced and may have survived, a seed with $6 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ is genetically balanced by fertilization with a normal seven-chromosome gamete. Compensating diploid plants in other plant species may have originated in a similar fashion.

3. Identification of Novel-Compensating Diploid

a. Morphological Identification

Morphologically, no visible differences are expected among compensating diploid plants. In barley, plants with the formulas 2n = 14, 13 + 1telo1L + 1telo1S, 12 + 2telo1L + 2telo1S (Tsuchiya, 1973) and 2n = 14,13 + 1acro3L^{3S} + 1telo3S, 2n = 12 + 2acro3L^{3S} + 2telo3S (Singh and Tsuchiya, 1981c; Singh and Tsuchiya, 1993) were indistinguishable morphologically. In this situation, chromosome count is required to separate these cytotypes.

b. Cytological Identification

True identity of novel-compensating diploids is based on chromosome count and karyotype analysis (Figure 7.50A,B). Banding techniques can be used where arms are similar.

4. Meiotic Behavior in Novel-Compensating Diploid

In general, meiosis is normal in compensating diploids. Meiotic chromosome behavior in plants with $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ and $2n = 12 + 2 \operatorname{acro3L^{3S}} + 2 \operatorname{telo3S}$ will be described in detail.

a. Plants with $2n = 13 + 1acro3L^{3S} + 1telo3S$

The general trend of chromosome association in plants with $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ was 3S-3S.3L-3L^{3S}, and this type of pairing behavior was readily recognized at late diplotene to diakinesis because of the differences in length of telo3S and acro3L^{3S} (Figure 7.51A,B,C). Of the

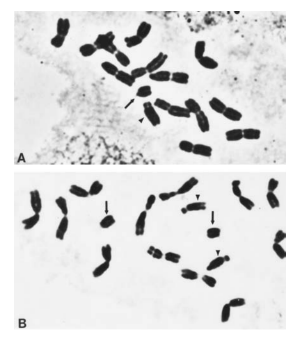


Figure 7.50 Somatic metaphase chromosomes of novel-compensating diploids of barley. (A) $2n = 13 + 1acro3L^{3S} + 1telo3S$, 22.2% of proximal portion of the short arm of Chromosome 3 is in trisomic condition; (B) $2n = 12 + 2acro3L^{3S} + 2telo3S$, 22.2% of proximal portion of the short arm of Chromosome 3 is in tetrasomic condition. Arrows show telo3S, and arrowheads show $acro3L^{3S}$. (From Singh, R.J. and Tsuchiya, T., *Genome*, 36, 343–349, 1993. With permission.)

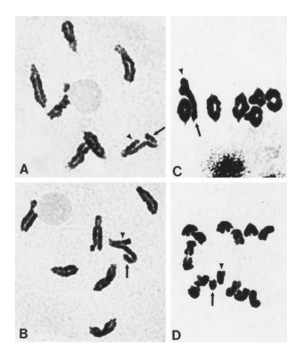


Figure 7.51 Meiotic chromosomes of plants with 2n = 13 + 1acro3L^{3S} + 1telo3S of barley. (A,B) Early diakinesis showing 1III + 6II chromosome configuration; (C) metaphase-I showing one rod-shaped trivalent + six bivalents; arrow indicates an interstitial chiasma between long arm of Chromosome 3 and acro3L^{3S}; (D) anaphase-I showing 7–8 (6 + 1acro3L^{3S} + 1telo3S) chromosome separation. Arrows show telo3S, and arrowheads show acro3L^{3S}. (From Singh, R.J. and Tsuchiya, T., *Genome*, 36, 343–349, 1993. With permission.)

312 metaphase-I sporocytes analyzed, 262 sporocytes (84%) showed 1III + 6II, and the remaining 50 sporocytes (16%) had 6II + 1 heteromorphic (het.) II + 1I configurations. A heteromorphic bivalent constituted a normal Chromosome 3 and an $acro3L^{3S}$ and often telo3L remained as univalent. The $acro3L^{3S}$ was more tightly associated with the long arm of Chromosome 3 by forming a interstitial chiasma than was telo3S (Figure 7.51C).

Chromosome migration at anaphase-I was influenced by the types of trivalent configuration. The highest frequency (78.2%) of sporocytes with V-shape trivalent resulted in the highest frequency (87.9%) of sporocytes with seven (normal chromosomes) to eight (six normal + 1acro3L³⁸ + 1telo3S) chromosome disjunction (Figure 7.51D). Moreover, low frequencies of sporocytes showed the following chromosome separation: seven normal + 1acro3L³⁸ – six normal + 1telo3S = (9.3%); seven normal + 1telo3S – seven normal + 1acro3L³⁸ = (2.8%). The spores generated from 7–8 chromosome disjunction are genetically balanced and function normally. However, spores with 7 + 1acro3L^{3S} chromosomes and 7 + 1telo3S are functional only through the female and produced plants with 2*n* = 14 + 1acro3L^{3S} and 2*n* = 14 + 1telo3S after fertilization with normal seven-chromosome gametes. Such gametes are expected from a rod-shaped trivalent, where one normal + 1telo3S or one normal + 1acro3L^{3S} move to one pole, leaving 1telo3S or 1acro3L^{3S} for the other pole. The spores with *n* = 6 + 1acro3L^{3S} and *n* = 6 + 1acro3L^{3S} + 1telo3S will be aborted because of deficiency. This indicates that male and female spores with *n* = 7 and *n* = 6 + 1acro3L^{3S} + 1telo3S will be functional in equal frequencies. The almost normal anaphase-I resulted in almost completely normal telophase-I (97.3% cells).

b. Plants with $2n = 12 + 2acro3L^{3S} + 2telo3S$

Chromosome association in plants with $2n = 12 + 2acro3L^{3S} + 2telo3S$ from diplotene to early diakinesis was exclusively 8II (Figure 7.52 A,B). Six bivalents were from the normal 12 chromosomes,

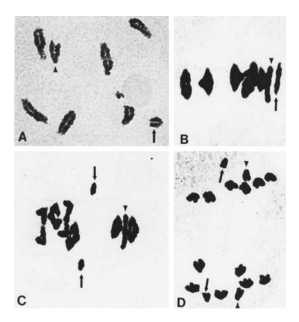


Figure 7.52 Meiotic chromosomes of plants with 2n = 12 + 2acro3L^{3S} + 2telo3S of barley. (A) Diakinesis with 8II (6II + 1IIacro3L^{3S} + 1IItelo3S); (B) metaphase-I showing 8II; (C) metaphase-I showing 7II + 2I; (D) anaphase-I showing 8–8 chromosome separation. Arrows show telo3S, and arrowheads show acro3L^{3S}. (From Singh, R.J. and Tsuchiya, T., *Genome*, 36, 343–349, 1993. With permission.)

and two bivalents were from 2acro3L³⁸ and 2telo3S. A bivalent from the telo3S was easily identified, because it showed pointed centromeric structure toward the spindle pole and was shorter in length than the bivalent derived from acro3L³⁸. Although 22.2% of the short arm of Chromosome 3 was in tetrasomic condition, a quadrivalent configuration was not recorded at diplotene or later meiotic stages.

The majority of the sporocytes (97.2%) at metaphase-I in plants with $2n = 12 + 2acro3L^{3S} + 2$ telo3S showed 8II (Figure 7.52B). Only six sporocytes of the 216 studied had 7II + 2I, in which two univalents were always telo3S (Figure 7.52C). At anaphase-I, 90% of the sporocytes showed 8(6 + $1acro3L^{3S} + 1telo3S$) – $8(6 + 1acro3L^{3S} + 1telo3S)$ chromosome separation (Figure 7.52D). Sometimes, bridges without fragments were recorded. At metaphase-II, all of the 200 dyads carried eight chromosomes. This indicates that meiosis is fairly normal in plants with $2n = 12 + 2acro3L^{3S} + 2telo3S$.

The normal meiosis in the above two novel-compensating diploids and also in other crops resulted in normal pollen and seed fertility. Fertility was not different from the diploid (Table 7.55).

Fedak (1976) studied meiosis and breeding behavior of an eight-paired barley produced by Wiebe, Ramage, and Eslick (1974). He observed 8II in 87.7% of the sporocytes. A low frequency (8.8%) of cells showed 1IV + 6II. This suggests that the material is still heterozygous for a small interchange involving a normal chromosome. Despite quadrivalent association, eight-paired barley bred true. However, seed fertility was only 65.7%.

5. Breeding Behavior of Novel-Compensating Diploid

Plants with various chromosome constitutions are expected in the progenies of compensating diploids. The results obtained in compensating diploid with $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ of barley will be cited here as an example. Somatic chromosome counts of 172 plants were obtained from the selfed progeny of a plant with $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$. Plants with chromosome constitutions $14,13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ (Figure 7.50A), and $12 + 2 \operatorname{acro3L^{3S}} + 2 \operatorname{telo3S}$ (Figure 7.50B) were obtained in a ratio of 1:2:1 (Table 7.56). Several other aneuploid types were also obtained with low frequencies (Table 7.56). A similar 1:2:1 ratio was observed by Tsuchiya (1973) and Strid (1968).

telo3S and 2 <i>n</i>		Spike	(Range)
3 + 1acro3L ^{3s} + 1	Seed Fertility	Plant	(Range)
ution of 2 <i>n</i> = 13			Average
omes Constitu		Number	of Plants
⁻ ertility (%) of Novel-Compensating Diploid with Chromosomes Constitution of $2n = 13 + 1$ acro3L ^{3s} + 1 telo3S and $2n + 2$ telo3S Compared with $2n = 14$ Siblings in Barley	llen Grains)		(Range) Degenerated of Plants
Fertility (%) of Novel-Compensating Diploid with Chro + 2telo3S Compared with $2n = 14$ Siblings in Barley	Pollen Fertility (% of Pollen Grains)		(Range)
of Novel-Com ompared with	Pollen Fe		Good
ed Fertility (%) 3L ^{3s} + 2telo3S C		Number	of Plants
Table 7.55 Pollen and Seed F = 12 + 2acro3L ^{3s} +		Chromosome	Constitution 2 <i>n</i>

14	5 2	98.2	(98.0–99.3)	1.8	ъ	96.61
13 + 1acro3L ^{3S} + 1telo3S	5 2	97.3	(95.1 - 98.8)	2.7	8	89.52
12 + 2acro3L ^{3S} + 2telo3S	5	98.3	(98.1–99.2)	1.2	Ð	90.48
Source: From Singh, R.J. and Tsuchiya, T., Genome, 36, 343-349, 1993. With permissio	Tsuchiya, T.,	Genome, 36,	343-349, 1993. Witl	h permission.		

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	Frequency (%)	Frequency (%) of Plants with Somatic Chromosome Numbers	Somatic Chromos	some Numbers	Total Number of	Chi s	Chi square
Parental Material 2 <i>n</i>	14	Type I	Type II	Other Types ^a	Plants Studied	1:2:1	÷
13 + 1acro3L ^{3S} + 1telo3S self	46(26.7)	80(46.6)	39(22.7)	7 ^a (4.1)	172	0.41	
13 + 1acro3L ^{3S} + 1telo3S \times 2x	48(52.2)	44(47.8)			92		0.17
Range (%)	(42.9–57.0)	(43.0–57.1)	I	I	Ι		
$2x \times 13+1$ acro3L ^{3S} +1 telo3S	56(44.1)	71(55.9)	I	I	127		1.77
Range (%)	(31.8 - 54.5)	(45.5 - 68.2)	Ι	I	Ι		
12 + 2acro3L ^{3S} + 2telo3S self	1	1	94(100.00)	4 L	94		
12 + 2acro3L ^{3S} + 2telo3S \times 2x	I	284	ő	I	287		
$2x \times 12 + 2acro3L^{3S} + 2telo3S$	သိ	107	I	I	110		

^b Triploid $(2n = 18 + 3 \arctan 3^{3s} + 3 \arctan 3^{3s})$. ^c Resulted from possible selfing.

Source: From Singh, R.J. and Tsuchiya, T., Genome, 36, 343-349, 1993. With permission.

(88.9–100) (82.1–100) (76.9–100)

(88.9–100) (84.0–100) (76.9–100)

As expected, a 1:1 ratio was observed for 2n = 14 and $2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ plants in diploid $(2n = 14) \times 2n = 13 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ and reciprocal crosses. However, the percent range of plants with $2n = 13 + \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ was higher compared to diploid in the progenies where plants with $2n = 13 + \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ were used as male parents. This suggests that male gametes with $n = 6 + 1 \operatorname{acro3L^{3S}} + 1 \operatorname{telo3S}$ are genetically balanced, and in this particular case, compete better than normal n = 7 chromosome gametes in fertilization, although 22.2% of the short arm was in a duplicate disomic condition in the gametes. The 22.2% segment, however, is heterochromatic.

Plants with $2n = 12 + 2 \operatorname{acro3} L^{38} + 2 \operatorname{telo3} S$ chromosomes were completely stable. Somatic chromosomes of 95 seedlings in the selfed progenies of plants with $2n = 12 + 2 \operatorname{acro3} L^{38} + 2 \operatorname{telo3} S$ were counted. Ninety-four seedlings were parental type, and one plant was triploid ($2n = 18 + 3 \operatorname{acro3} L^{38} + 3 \operatorname{telo3} S$). This triploid plant may have originated from the fertilizaton of an unreduced female gamete ($12 + 2 \operatorname{acro3} L^{38} + 2 \operatorname{telo3} S$) with a male gamete containing $n = 6 + 1 \operatorname{acro3} L^{38} + 1 \operatorname{telo3} S$. All the plants were $2n = 13 + 1 \operatorname{acro3} L^{38} + 1 \operatorname{telo3} S$ in crosses between plants with $2n = 12 + 2 \operatorname{acro3} L^{38} + 2 \operatorname{telo3} S$ and 2x. Similar results were reported by Manga et al. (1981) in pearl millet.

6. Usefulness of Novel-Compensating Diploid

The present case of an eight-paired barley is another example of establishing a barley strain with the basic chromosome number x = 8. Eight-paired barleys have been reported by Tsuchiya (1969, 1973) and Wiebe, Ramage, and Eslick (1974). Tsuchiya (1969) isolated an eight-paired barley from the progenies of Triplo 6, which had a paracentric inversion in the extra Chromosome 6, and showed 61% seed set, although meiosis was normal. The eight-paired barley reported by Wiebe, Ramage, and Eslick (1974) was derived from the progeny of tertiary trisomics and showed reduced vigor and partial seed set. However, another eight-paired barley (2n = 12 + 2telo1L + 2telo1S), as expected, was indistinguishable from the diploid, because it was genetically balanced (Tsuchiya, 1973). Recently, Hang and Tsuchiya (1992) produced nine-paired barley by crossing two eight-paired $(2n = 12 + 2\text{acro3L}^{3\text{S}} + 2\text{telo3S}; 2n = 12 + 2\text{telo1L} + 2\text{telo1S})$ barleys. The nine-paired barley was qualitatively and quantitatively similar to diploid barley.

Hagberg (1962) successfully produced a duplication for barley breeding by crossing two translocation lines involving the same chromosomes, using the principle established by Gopinath and Burnham (1956). The novel-compensating plant with $2n = 12 + 2 \operatorname{acro3L^{3S}} + 2 \operatorname{telo3S}$ is tetrasomic for the 22.2% proximal segment of the short arm in the acro3L^{3S} and may be useful in barley breeding programs. If a dominant gene (genes) for desired traits is located in the 22.2% segment, this particular region will carry four doses and will be superior to two doses carried by diploid barley. Furthermore, the four doses of a gene are assumed to be fixed in the population, because a quadrivalent was not observed at meiosis. This assumption was confirmed by growing the selfed progeny of plants with $2n = 12 + 2 \operatorname{acro3L^S} + 2 \operatorname{telo3S}$; all the plants were green. The gene *yst2* is in duplex condition (*Yst2*, *Yst2*, *yst2*); two telo3S carry the recessive allele and $2 \operatorname{acro3L^{3S}}$ carry the dominant allele. If a quadrivalent formation occurs, the selfed progeny should segregate in a 35:1 ratio for green and yellow streak. All the plants were green, confirming the absence of quadrivalent formation.

III. MONOSOMICS AND NULLISOMICS

A. Introduction

The term monosome (2n - 1), first coined by Blakeslee (1921), designates a primary monosome (monosomic) where one of the chromosomes is missing from the normal diploid complement. Tertiary monosomics possess a translocated chromosome, i.e., arms from two different chromosomes. The loss of a chromosome in a diploid species has a more drastic effect on plant morphology than when it occurs in a polyploid species. Monosomics are useful for locating genes to specific

chromosomes, in the assignment of linkage groups, in the study of the genetic control of chromosome pairing, and in the manipulation of chromosomes. Monosomics have been described in great detail by Burnham (1962), Khush (1973), and Weber (1983, 1991). Therefore, only classical examples will be discussed in this chapter.

An individual lacking one chromosome pair is called nullisome (nullisomic) and is symbolized as 2n - 2. For example, hexaploid wheat constitutes 2n = 6x = 42 chromosomes. If a pair of chromosome such as 1A is deleted, the plant will carry 2n = 6x = 40, and this particular plant is identified as a nulli-1A. A complete series of nullisomics in wheat cv. Chinese Spring has been established (Sears, 1944, 1954).

B. Sources of Monosomics and Nullisomics

Occasionally, monosomics originate spontaneously in the progenies of normal diploid and polyploid populations (Table 7.57). However, the frequency of occurrence of monosomic plants is relatively higher after treatment with various physical or chemical mutagens; in the progenies of haploids, polyploids, and aneuploids; in interspecific crosses; and in the progenies of plants with certain genetic systems (Khush, 1973; Edwards et al., 1980; Weber, 1983, 1991).

1. Spontaneous Origin

Monosomic (2n - 1) plants appear spontaneously, although rarely, in the progenies of diploids and polyploids. Blakeslee and Belling (1924a) observed sectoral chromosomal chimeras for monosomics (2n - 1), trisomics (2n + 1), and tetraploids (4x) in *Datura stramonium*. The deficiency was for one of the largest chromosomes, and offspring from the 2n - 1 branch failed to show individuals of the parental types. In addition, Bergner, Avery, and Blakeslee (1940) reported only one 2n - 1 plant among 20,879 plants of diploid *Datura stramonium*.

Monosomics can be isolated through screening of cultivated varieties. However, it requires considerable effort. Hacker and Riley (1963) identified six nullisomics, 40 monosomics, four trisomics, and three with telocentrics among 3453 plants in an hexaploid (2n = 6x = 42) oat cultivar Sun II (1.53% aneuploids). McGinnis (1962) reported 0.6% aneuploids in the sampled population of oat (Table 7.58). Riley and Kimber (1961) screened four wheat cultivars, and 0.69% of the population proved to be monosomic.

2. Chemical Treatments and X-Rays

Monosomics have been generated in populations after colchicine treatments and also from x-rayed pollen (Table 7.57). Bergner, Avery, and Blakeslee (1940) found seven monosomic (2n - 1 = 23) plants among 2135 grown plants after treating seeds of *Datura stramonium* with colchicine. They suggested that the occurrence of monosomics was 70 times as frequent as was observed in the control. Smith (1943) recorded a total of 26% off-type in colchicine-treated plants. One monosomic was in *Nicotiana langsdorffii*, and three were in F₁*Nicotiana langsdorffii* × *N. sanderae* colchicine-treated germinating seeds.

Bergner, Avery, and Blakeslee (1940) assumed the chromosome loss was due to anaphase-lagging or nondisjunction in cells recovering from the effect of colchicine treatment. However, according to Smith (1943), the loss of chromosomes was due to an abnormal scattering of chromosomes in colchicine-arrested cells, which was followed during recovery by multipolar spindle formation.

Irradiation of pollen often produces monosomics. The tube nucleus of a pollen grain mediates metabolism of the pollen grain, and the loss of one chromosome from the generative nucleus or one of the sperm nuclei, relatively inert metabolically, is compatible with survival and functioning of the pollen grain (Weber, 1983). Rick (1943) found two monosomic plants (2n = 13) in *Petunia hybrida* after x-raying pollen: one was a primary monosome, and the other was a tertiary monosome.

Crop Species	n	Methods	References
Datura stramonium	12	Spontaneous	Blakeslee and Belling, 1924a; Blakeslee and Avery, 1938
		Colchicine treatment	Bergner, Avery, and Blakeslee, 1940
Lycopersicon esculentum	12	X-ray of pollen	Rick and Khush, 1961; Khush and Rick 1966
Nicotiana longsdorfii	10	Colchicine treatment	Smith, 1943
Zea mays	10	$3x \times 2x$	McClintock, 1929a
		X-ray of pollen	Morgan, 1956; Baker and Morgan, 1966
		Genetic system	Weber, 1983
		Trisomics	Einset, 1943
		Knob and B-chromosome interaction	Rhoades, Dempsey, and Ghidoni, 196
Petunia hybrida	7	X-ray of pollen	Rick, 1943
Nicotiana tabacum	24	Interspecific hybrids, asynapsis	Clausen and Cameron, 1944
Nicotiana alata	9	Spontaneous	Avery, 1929
Gossypium	26	Spontaneous, trisomics,	Endrizzi, 1963; Brown and Endrizzi,
hirsutum		monosomics	1964; Endrizzi and Ramsay, 1979
		X-ray, neutron, gamma ray, intervarietal hybrids, interspecific hybrids, cytological combinations, desynapsis	Edwards et al., 1980
Avena byzantina	21	Haploids, aneuploids, autotriploids	Nishiyama, 1970
Avena sativa	21	Interspecific hybrids	Rajhathy and Dyck, 1964
		Spontaneous	McGinnis, 1962; Hacker and Riley, 1963; Ekingen, 1969
		X-ray	Costa-Rodrigues, 1954; Rajhathy and Dyck, 1964; Andrews and McGinnis, 1964; Chang and Sadanaga, 1964; Schulenburg, 1965
		Gamma ray	Singh and Wallace, 1967b
		Nitrous oxide	Dvořák and Harvey, 1973
Triticum aestivum	21	Haploids, nullisomics, trisomics	Sears, 1954
		Monosomics, translocations, x- ray pollen	Riley and Kimber, 1961
Clarkia amoena	7	Spontaneous	Snow, 1964
Glycine max	20	Desynapsis	Skorupska and Palmer, 1987
	20	Primary trisomics	Xu, Singh, and Hymowitz, 2000b
Brassica napus	19	Intergeneric and interspecific hybrids	Fan and Tai, 1985
Oryza sativa	12	Interracial hybrid	Seshu and Venkataswamy, 1958
		Gamma ray	Wang and Iwata, 1996

	Table 7.57	Sources	of	Monosomics
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Rick and Khush (1961) x-rayed pollen of nonmutant tomato plants and pollinated plants containing recessive markers. Among 2312 plants analyzed, nine plants monosomic (2n = 23) for Chromosome 11 were identified. Subsequently, Khush and Rick (1966) identified cytologically a primary monosome, mono-11, and 18 tertiary monosomics. All the interchanges occurred in the kinetochore, and deficiencies were tolerated for only 15 of the 24 arms of the complement.

Baker and Morgan (1966) x-irradiated maize pollen containing the dominant asynaptic allele (*As*), and pollinated a recessive asynaptic (*as*) female. Of the 5593 X_1 kernels expressing markers flanking the kinetochore, three plants were monosomic (2n = 19) for Chromosome 1. Two more plants were recovered, which had lost two of the three markers, and they probably were tertiary

		Numbers of Seedlings	%	
Methods	Oat Cultivars	Screened	Aneuploid	References
Haploidy	Garry	224,000	0.0	Rajhathy and Dyck, 1964
	Victory	224,000	0.0	Rajhathy and Dyck, 1964
	Rodney	224,000	0.0	Rajhathy and Dyck, 1964
Selection	Garry	4203	0.6	McGinnis, 1962
	Sun II	3453	1.5	Hacker and Riley, 1963
	Interspecific populations	2970	30.5	Rajhathy and Dyck, 1964
X-irradiation				
300–500r	Garry	200	30.8	Rajhathy and Dyck, 1964
600r	Garry	293	9.9	Andrews and McGinnis, 1964
600r	Rodney	160	23.1	Andrews and McGinnis, 1964
30kr	Borreck	220	13.6	Schulenburg, 1965
700r	Cherokee	233	9.8	Maneephong and Sadanaga, 1967
300r	Missouri 04047	279	7.2	Costa-Rodrigues,1954
Sonic vibration (20 min)	Garry	24	4.2	Andrews and McGinnis, 1964
Myleran 10 ³ M	Garry	113	0.9	Andrews and McGinnis, 1964
-	Rodney	90	0.0	Andrews and McGinnis, 1964
8-ethoxycaffeine 10 ³ M	Garry	72	1.4	Andrews and McGinnis, 1964
	Rodney	126	0.8	Andrews and McGinnis, 1964
Nitrous oxide	Garry	_	36.0	Dvořák and Harvey, 1973
Selection	Borreck	244	0.5	Schulenburg, 1965
Selection	Borreck	3500	2.4	Ekingen, 1969
	Zenshin	3000	2.5	Ekingen, 1969
EMS (3%)	Borreck	158	4.4	Schulenburg, 1965

Table 7.58 Aneu	ploids Resulting	from Physica	I and Chemical	Treatments of	f Hexaploid Oat

monosomics. This suggests that the monosomic in maize for Chromosome 1, the longest, is tolerated by the maize genome, but it is difficult to maintain.

Five monosomics (2n = 23) of rice were produced by treatment of pollen with gamma rays by Wang and Iwata (1996). One monosomic was tertiary, and the remaining four contained normal chromosomes. Mono-9 and mono-10 were identified cytologically.

By utilizing x-rays, gamma rays, and chemical mutagens, several attempts were made to induce monosomics in hexaploid oat (Table 7.58). The frequencies of monosomic plants ranged from 0.0 to 36.0%. Rajhathy and Thomas (1974) suggested that the production of aneuploids by radiation or chemical mutagens is not desired in hexaploid oat, because these mutagens also induce minor chromosome structural changes that cannot be readily detected in polyploid species, because polyploid species can tolerate considerable amounts of deficiency and duplication.

3. Haploids, Polyploids, and Aneuploids

Haploids have been an excellent source for isolating aneuploids, particularly monosomics, in hexaploid wheat (2n = 6x = 42) cv. Chinese Spring. Nineteen of the possible 21 monosomics were derived from the progenies of haploids (Sears, 1939, 1954). The remaining two monosomics were obtained from a nullisomic III (3B). Based on studies of meiosis in haploids published before 1939, Sears (1939) summarized four ways to obtain monosomics in wheat from the progenies of haploids.

Haploids, obtained from twin seedlings, also generated monosomics in hexaploid oat (Rajhathy and Dyck, 1964; Nishiyama et al., 1968; Nishiyama, 1970). Nishiyama (1970) isolated seven (1 to 7) of the 21 possible monosomics in *Avena byzantina* cv. Kanota. The remaining monosomics were found from the progenies of aneuploids (2n +) and autotriploids (3x).

In maize, McClintock (1929a) found an individual in an F_2 generation of $3x \times 2x$, where microsporocytes showed a 2n - 1 chromosome complement. The monosomic plant was small, with a poorly developed tassel, and in all cases, a 9II + 1I chromosome configuration was observed.

Monosomics have also been isolated from the progenies of trisomics (Einset, 1943; Brown and Endrizzi, 1964; Xu, Singh, and Hymowitz, 2000b). Einset (1943) obtained five monosomics and one monosomic and one fragment chromosome plants among 1916 plants in the crosses of $2x + 1 \times 2x$ maize plants. Xu, Singh, and Hymowitz (2000b) discovered mono-3 and mono-6 of the soybean in the back-cross progenies of Triplo 3 (BC₃) and Triplo 6 (BC₄), respectively. Triplo 3 (BC₂) was male and Triplo 6 was female in back cross to cv. Clark 63. Mono-3 probably originated from an n - 1 sperm and mono-6 from an n - 1 egg. The derivation of monosomics from the primary trisomics is generally associated with nondisjunction and lagging of chromosomes during meiosis, leading to male and female spores lacking one complete chromosome. Mostly, the missing chromosome in a monosomic plant is the same as the extra chromosome in the primary trisomic parent.

4. Asynaptic and Desynaptic Lines

Clausen and Cameron (1944) obtained monosomics in *Nicotiana* in asynaptic male sterile female \times normal male crosses. Aside from an occasional triploid, the offspring consisted of unbalanced diploids; relatively few were simple monosomics or trisomics, and most of them were double or triple monosomics or monosomic–trisomic combinations. Sears (1954) isolated 17 of the possible 21 monosomics in the progenies of partially asynaptic Nulli-III (3B) of hexaploid wheat cv. Chinese Spring. In *Gossypium hirsutum*, a monosomic M13 was obtained from an asynaptic strain (Brown and Endrizzi, 1964). Skorupska and Palmer (1987) identified a monosomic (2n = 39) in soybean from the progenies of a desynaptic mutant.

5. Intra- and Interspecific Hybrids

Clausen and Cameron (1944) isolated monosomics in *N. tabacum* x *N. sylvestris* or *N. tomentosa* crosses. However, a considerable amount of heterozygosity was found that confused the recognition and establishment of monosomic series.

A high frequency of monosomics in oat was obtained by Rajhathy and Dyck (1964) among the progenies of back-crossed pentaploid hybrids (Table 7.58). However, they questioned the usefulness of this source of monosomic lines because of the heterogeneity of the progeny. Chromosome rearrangements that could arise from such wide crosses could result in disturbances in meiotic behavior and lead to seed sterility in hybrids between the isolated monosomic lines and established varieties, when the latter are used in monosomic analysis. However, these lines could be back crossed to a desired cultivar to establish the aneuploid series in a uniform genetic background.

6. Genetic System

An unique genetic system, the *r*-X1 deficiency in maize, has been discovered for generating primary monosomics (Weber, 1991). The *r*-X1 deficiency includes the *R* locus on Chromosome 10. The dominant *R* locus is necessary for anthocyanin production in the aleurone of the endosperm of maize kernels. In induction of monosomics, if *R/r*-X1 plants (heterozygous for the deficiency) are test crossed as female parents by an *r/r* male parent, among the test cross progeny, about 55 to 66% of the kernels are colored (*Rr*), while the remainder are colorless (*r/r*-X1), deficiency-bearing kernels. The deficiency is only transmitted through the female and is not transmitted through pollen. The progeny of *Rr* are always diploid, but the progeny of *r/r*-X1 segregate in equal (10 to 18%) proportions of monosomics and trisomics; the remaining plants are diploid or occasionally multiply aneuploid. Thus, aneuploids are produced only from ovules bearing the *r*-X1 deficiency.

		Chromosome		_
Female (<i>R</i> / <i>r</i> -X1)	Male (<i>r/r</i>)	Number	Marker Gene	Frequency
Bm2	bm2	1	Brown midrib	0.03
Lg	lg	2	Liguleless	1.24
A	а	3	Anthocyaninless	0.21
Su	su	4	Sugary endosperm	0.29
Pr	pr	5	Red aleurone	0.06
Υ	У	6	Yellow endosperm	1.89
GI	gl	7	Glossy seedling	1.01
J	j	8	Japonica striping	3.46
Wx	WX	9	Waxy endosperm	0.49
G	g	10	Golden plant	1.57

Table 7.59 Mangelsdorf's Multiple Chromosome Markers in Maize

Source: From Weber, D.F., In Chromosome Engineering in Plants: Genetics, Breeding, Evolution. Pt. A., Gupta, P.K. and Tsuchiya, T., Eds., Elsevier, Amsterdam, 1991, 181–209. With permission.

Weber (1983) suggested that a gene is located on Chromosome 10 within the segment corresponding to the deficiency which is necessary for normal chromosomal disjunction at postmeiotic divisions during the megagametophyte (embryo sac) mitotic divisions. The loss occurs postmeiotically in the *r*-X1 deficiency-bearing embryo sacs. Thus, the full haploid chromosome complement is present in these embryo sacs, and nondisjunction occurs at one or more of the three mitotic divisions during embryo sac development, generating some aneuploid nuclei in these embryo sacs. Even though nullisomic nuclei are present in these embryo sacs, they do not abort, because other nuclei in the same embryo sac are haploid or disomic for the monosomic chromosome. When n - 1 egg in such embryo sacs is fertilized by haploid pollen, a monosomic is produced.

Monosomics generated utilizing the r-XI system are selected as follows. A male parent recessive for a sporophyte-expressed mutation is crossed onto an R/r-XI female homozygous for the dominant allele of this mutation. Progeny expressing the recessive phenotype are usually monosomic for the chromosome carrying this mutation; however, some plants with deficiencies including the marker mutation are also recovered.

Weber (1983) recovered primary monosomics for each of the ten maize chromosomes from a cross between two inbred lines. He crossed a r/r male that also carried a recessive marker for each of the ten chromosomes (Mangelsdorf's multiple chromosome tester) onto a R/r-X1 female that bore the corresponding dominant alleles (Table 7.59). In the F₁, r/r-X1 plants were selected. Five of the markers in Mangelsdorf's tester are expressed in the sporophyte (*bm2*, *lg*, *gl*, *j*, and *g* on Chromosomes 1, 2, 7, 8, and 10, respectively). Progeny of this cross expressing one of these markers are usually monosomic for the chromosome that carries the marker; however, partial chromosome losses are also occasionally observed.

The other five markers express in the endosperm of kernels. Thus, plants monosomic for the chromosomes bearing these markers cannot immediately be detected. Plants from the above cross that are of subnormal stature and have at least 50% aborted pollen are identified as putative monosomics. These are crossed with a stock that is *a*, *su*, *pr*, *y*, *w*, *x*, and *R*. If a plant is monosomic for Chromosomes 3, 4, 5, 6, or 9, the test cross will produce only kernels of the recessive phenotype for the chromosome bearing that gene. Diploid and all other monosomics will give a 1:1 ratio.

In hexaploid wheat monosomics, one expects to obtain an average of about 3% nullisomic plants from the selfed progeny (Sears, 1953b). This result is not always recorded experimentally, however, because the frequency of nullisomics depends upon the genetic background of monosomics, chromosome type, and growing conditions. The selfed progenies of monosomics in Chinese Spring transmitted nullisomics with the range of 0.9% (5B) to 10.3% (3B) (Sears, 1944). In *Avena byzantina* cv. Kanota, Morikawa (1985) found nullisomics in the progeny of only four monosomic lines, and the frequency was highly variable (mono-8, 42.1%; mono-19, 41.2%; mono-9, 2.7%, and mono-17, 2.8%). Seventeen monosomic lines did not transmit nullisomics.

Kramer and Reed (1988) pollinated mono-F, H, L, M, P, and R of *N. tabacum* – Purpurea by irradiated pollen of *N. glutinosa*. Two nulli haploid plants (2n = 23) were recovered only from a cross with mono-H. Subsequently, nullisomic (2n = 46) plants were obtained after chromosome doubling.

C. Identification of Monosomics and Nullisomics

1. Morphological Identification

Monosomic plants in diploids are generally more drastically modified morphologically than those in polyploids. Monosomics in *Petunia hybrida* (Rick, 1943), *Lycopersicon esculentum* (Rick and Khush, 1961; Khush and Rick, 1966), *Avena strigosa* (Andrews and McGinnis, 1964) *Zea mays* (McClintock, 1929a; Einset, 1943; Baker and Morgan, 1966), and *Oryza sativa* (Wang and Iwata, 1996) were weak and reduced in size.

Monosomics in maize generated by the r-XI deficiency procedure were relatively more vigorous than those monosomics in maize reported earlier. Weber (1983) emphasized that monosomic plants need "tender care." Monosomics in maize are smaller than their diploid sibs, and as in primary trisomics, chromosome imbalance affects the plant in a distinctive manner.

Maize monosomics mature slower than diploids, however, they were not retarded at any stage of their growth (Weber, 1983, 1991). In contrast, the growth state of monosomic tomato plants was extremely slow in the seedling stage; after 2 to 3 months, they showed vigorous growth, and some reached large size (Khush and Rick, 1966). Monosomics in rice were dwarf and completely sterile.

Two monosomics in the soybean expressed contrasting morphological features. Mono-3 was slow in vegetative growth (dwarf) with few branches but produced plumper pods and larger seeds than those in the disomic sibs. Mono-6 was indistinguishable from the disomic prior to flowering stage. Mono-6 set wrinkled pods and smaller seeds than those in the disomic plants (Xu, Singh, and Hymowitz, 2000b). A definite conclusion regarding the loss of a chromosome from the diploid complement can be drawn only after isolating all possible 20 monosomics in the soybean.

At the tetraploid level, the loss of one chromosome is not quite as severe as it is in diploids. In *Nicotian tabacum*, the 24 monosomics produced by Clausen and Cameron (1944) differ from one another and from normal sibs in a specific ensemble of quantitative and qualitative morphological features that can be classified accurately on the basis of their morphology; some monosomics were quite distinctive, while others required more careful examination. Similarly, monosomics in *Gossypium hirsutum* can be recognized by distinct morphological characteristics, including modifications of vegetative and reproductive structures, such as smaller or narrower leaves, smaller flowers or flower parts, and smaller, longer, or partially collapsed bolls (Brown and Endrizzi, 1964; Endrizzi and Brown, 1964).

Fan and Tai (1985) reported two monosomics in *Brassica napus*, and both were morphologically indistinguishable from normal disomics.

In hexaploid wheat and oat, monosomics differ little from normal sibs. Monosomics in wheat cv. Chinese Spring, grown under favorable conditions, were difficult or impossible to distinguish from normal, except for mono-IX (5A) (Sears, 1954).

Hacker and Riley (1965) compared the morphological features of monosomics and nullisomics with the euploids of oat cultivar Sun II and were able to identify 13 distinct monosomic lines; but only a few discernible differences were observed between monosomic plants. Ekingen (1969) observed no morphologically distinguishable features in monosomics of oat, while nullisomics were clearly distinguished from each other as well as from the corresponding monoor disomic plants.

Nullisomic plants in Chinese Spring wheat can be distinguished from normal sibs as well as from the corresponding monosomics by morphological features at seedling and maturity stages (Table 7.60). However, nulli-1B, 7A, 7B, and 7D are difficult to distinguish from disomics; they

Homoeologous Group	Morphological Features
1. (1A, 1B, 1D)	Three nullisomics of this group are reduced in plant height in varying degree; spikes are a little less dense than normal, with slightly stiffer glumes; they are both female and male sterile
2. (2A, 2B, 2D)	All three nullisomics are dwarfish with greatly reduced tillers; all are male fertile but female sterile; the spikes have thin, papery glumes and are completely awnless
3. (3A, 3B, 3D)	Nullisomics are identified at the seedling stage by their narrow, short, stiff leaves, and at maturity, nullisomics are dwarfed with narrow leaves and short spikes
4. (4A, 4B, 4D)	Nullisomics of all three have narrow leaves and slender culms; mature plants are dwarfed and male sterile
5. (5A, 5B, 5D)	Nullisomics have narrow leaves and slender culms, are late in maturity, spikes are reduced in size and have small glumes and seeds; they are female fertile and male sterile
6. (6A, 6B, 6D)	Nullisomics have narrow leaves, slender culms, and narrow, spreading outer glumes; they are straggly in appearance; all are female fertile
7. (7A, 7B, 7D)	Nullisomics differ little from normal at the seedling and maturity stages, are distinguishable only by a slight reduction in vigor and height and by certain spike characteristics; the seed fertilities of 7B and 7D are nearly normal but are greatly reduced in 7A by pistilloidy

Table 7.60 Morphological Features of Nullisomics of Triticum aestivum cv. Chinese Spring

Source: Adapted from Sears, E.R., Mo. Agric. Exp. Stn. Res. Bull., 572, 1-58, 1954.

are identified by intercrossing (Sears, 1954). Nullisomics from the homoeologous group 7 differed little from disomics at the seedling and adult plant stages and were distinguishable only by a slight reduction in vigor and height and by certain spike traits.

Four of the 21 possible nullisomics of hexaploid oats were reduced in vigor and plant height (Chang and Sadanaga, 1964; Ekingen, 1969; Morikawa, 1985). Morikawa (1985) described morphological characteristics of four nullisomics of *Avena byzantina* cv. Kanota. Nulli-8 showed strong desynapsis, complete sterility, and fatuoid character. Nulli-9 was extremely weak, often dying before flowering, produced many tillers, and was self-sterile. Nulli-17 was weak and grass-like, with comparatively large florets, and was self-sterile. Nulli-19 was shorter in height, with virescent appearance in some environments, and was sterile.

Nulli-H of *Nicotiana tabacum* carried smaller flowers than those of the disomic and mono-H (Kramer and Reed, 1988).

2. Cytological Identification

Through observations of pachytene chromosomes, monosomics were identified in tomato (Khush and Rick, 1966) and maize (Weber, 1983). Khush and Rick (1966) cytologically identified a primary monosome (mono-11) and 18 tertiary monosomes in tomato. In tertiary monosomes, all interchanges occurred in the kinetochore. Wang and Iwata (1996) identified mono-9 and mono-11 in rice based on mitotic metaphase chromosome karyotype.

Like trisomics, monosomics are identified based on karyotype analysis of mitotic metaphase chromosomes, pachytene chromosomes, and univalents and lagging chromosomes. However, karyotype analysis alone could not facilitate precise identification of monosomics. Olmo (1936) was not able to distinguish cytologically the monosomics of *Nicotiana tabacum*, despite differences between *N. sylvestris* and *N. tomentosa* chromosomes.

Hexaploid wheat (2n = 6x = 42) is an allopolyploid and contains chromosomes of A, B, and D genome diploid (2n = 2x = 14) species. Thus, it is necessary to know which chromosomes belong to A, which to B, and which to D. Morrison (1953) and Sears (1954) measured univalents at meiotic metaphase-I and telophase-II of the monosomics of *Triticum aestivum* cv. Chinese Spring. It was, however, not possible to distinguish all of the chromosomes based on karyotype measurement.

Homoeologous			
Group	Genome A	Genome B	Genome D
1	1A (XIV)	1B (I)	1D (XVII)
2	2A (XIII) [II] ^a	2B (II) [XIII]	2D (XX)
3	3A (XII)	3B (III)	3D (XVI)
4	4A (IV) (4B) ^b	4B (VIII) (4A)	4D (XV)
5	5A (IX)	5B (V)	5D (XVIII)
6	6A (VI)	6B (X)	6D (XIX)
7	7A (XI)	7B (VII)	7D (XXI)

Table 7.61 The Homoeologous Groups, Chromosomes Belonging to the A, B, and D Genomes, and the Original Numbers (Parentheses) Assigned to These Chromosomes in Hexaploid Wheat cv. Chinese Spring

^a Chapman and Riley (1966).

^b Dvořák (1983b).

Source: From Okamoto, M., Can. J. Genet. Cytol., 4, 31-37, 1962. With permission.

To assign chromosomes to the D genome, Sears (1958) and Okamoto (1962) crossed 2n - 1 plants with a tetraploid wheat (*Triticum dicoccum*, AABB) and examined the pattern of meiotic pairing in the F₁. When the pattern was 14II + 6I, the monosome belonged to the D genome, and when 13II + 8I, the monosome belonged to either the A or the B genome. By utilizing monotelosomics (2n - 1 telo), they distinguished the chromosomes belonging to the A-genome from those belonging to the B-genome. Okamoto (1962) crossed all monotelosomics, except for IV (4A), with synthetic tetraploid AADD (*T. aegilopoides* × *T.tauschii*). In the F₁, if a telocentric chromosome belonged to A-genome, chromosomes formed heteromorphic bivalent. Failure to form heteromorphic bivalent suggested that the telocentric belonged to the B-genome (Table 7.61). He assigned VIII to 4B and IV to 4A.

Subsequent reports (Chapman, Miller, and Riley, 1976; Dvořák, 1976) disputed the designation of 4A, because telosome 4A failed to pair with any *T. urartu* (A genome) chromosomes. Based on karyotype (Dvořák, 1983b), Giemsa C- and N-banding techniques (Chen and Gill, 1984), and *in situ* hybridization (Rayburn and Gill, 1985a), it was confirmed that 4A belonged to the B genome. Heterochromatic banding procedures helped to solve the controversy of 4A and also facilitated identification of all the 42 chromosomes of Chinese Spring (Gill and Kimber, 1974; Gerlach, 1977; Endo, 1986; Shang, Jackson, and Nguyen, 1988a; Naranjo et al., 1988; Dvořák, Resta, and Kota, 1990). However, at the Seventh International Wheat Genetics Symposium held at Cambridge, England, the following conclusion was reached: *"The proposal was not fully accepted and it was finally agreed to recommend that 4 A be designated 4 B and that 4 B designated 4?"* (Kimber and Tsunewaki, 1988).

Several attempts were made to identify monosomics of hexaploid oat by karyotype analysis of metaphase chromosomes (McGinnis and Taylor, 1961; McGinnis and Andrews, 1962; Hacker and Riley, 1965; McGinnis and Lin, 1966; Singh and Wallace, 1967a; Ekingen, 1969; Nishiyama, 1970; Hafiz and Thomas, 1978; Morikawa, 1985). It was not possible to identify all monosomic lines due to such factors as pretreatment, fixation, staining procedure, and chromosome preparation technique. Morikawa (1985) could only divide oat karyotype into four groups: (1) three pairs of satellite chromosomes (Sat); (2) four pairs of metacentric chromosomes (M); (3) seven pairs of submetacentric chromosomes (SM); and (4) seven pairs of subterminal chromosomes (ST). The A- and B-genome chromosomes of oat do not express a diagnostic pattern of C-bands (Fominaya, Vega, and Ferrer, 1988a,b). Thus, assignment of chromosomes to their respective genomes has not yet been established clearly.

Another technique often used is to intercross the same or different monosomic lines. The chromosome pairing at metaphase-I is examined. The failure of two monosomes to pair with each other (double monosomics) will indicate that the monosomic lines are different. Monosomics can also be identified by translocation tester stocks, if available (Endrizzi and Brown, 1964).

3. Meiosis in Monosomics and Nullisomics

In monosomic plants (2n - 1), generally, chromosome associations of bivalents (II) and one univalent (I) are observed at diakinesis and metaphase-I. Nonhomologous chromosome association was frequently recorded at pachynema in the univalent. The frequency of trivalent association is substantially lower in monosomics of diploids than in monosomics of tetraploid and hexaploid plants. In maize monosomics (2n - 1 = 19), Weber (1983) observed 9II + 1I in 99.3% of the cells and a trivalent configuration in 0.7% cells. He suggested that a trivalent configuration might be an artifact when a univalent chromosome is adjacent to or superimposed upon a bivalent. Figure 7.53 shows a sporocyte of mono-6 at diakinesis, where an univalent Chromosome 6 is attached with the nucleolus.

It is interesting to note that maize, being a diploid, tolerates a considerable degree of deficiencies as doubly and triply monosomic plants have been isolated and studied cytologically. Double and triple monosomics are identified by the simultaneous loss of marker genes on two and three nonhomologous chromosomes. Nonhomologous univalents rarely pair at diakinesis or metaphase-I (Figure 7.54A,B) and segregate independently at anaphase-I (Weber, 1991).

In four rice monosomics (2n = 23), 100% sporocytes contained 11II + 1I chromosome configuration (Wang and Iwata, 1996). Similarly, two monosomics of the soybean showed exclusively 19II + 1I (Xu, Singh, and Hymowitz, 2000b).

Nicotiana tabacum (2n = 4x = 48) and *Brassica napus* (2n = 38) are allotetraploids. Clausen and Cameron (1944) observed in monosomics of *N. tabacum*, in addition to usual configurations of 23II + 1I, two more types of associations: (1) the occurrence of a trivalent, 1III + 22II, and (2) desynaptic cells with 22II + 3I, 21II + 5I, etc. Both abnormalities were associated with mono-D and mono-S. And in other monosomes, trivalent configurations were rare.

Chromosomal affinities were also recorded in *B. napus*. Fan and Tai (1985) observed in mono-1, 18II + 1I in 85.3% cells and 1III + 17II in 14.7% cells. But in mono-2, the 1III + 17II configuration occurred in 61.9% of cells, a figure substantially higher than with mono-1.

The trivalent configuration was not recorded in monosomics of hexaploid (2n = 6x = 42) wheat, but desynaptic cells were recorded. Morrison (1953) studied meiotic pairing in eight of the 21 monosomes of wheat, and observed 20II + 1I in 97.6% of the sporocytes. Low frequencies of 19II

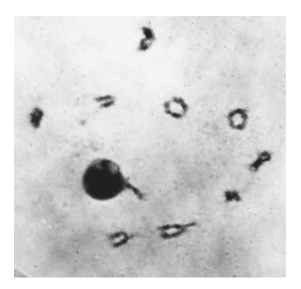


Figure 7.53 Mono-6 in maize at diakinesis, showing 9II + 1I configuration. Note the univalent Chromosome 6 associated with the nucleolus. (From Plewa, M.J. and Weber, D.F., *Genetics*, 81, 277–286, 1975. With permission.)

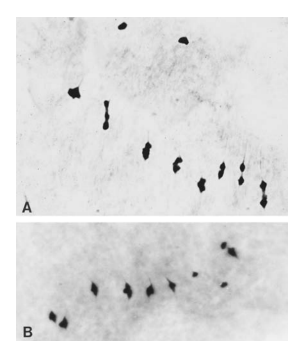


Figure 7.54 Double (A) and triple (B) monosomics in maize. (A) A metaphase-I cell with 8II + 21. (B) A metaphase-I cell with 7II + 3I. (Courtesy of D.F. Weber.)

+ 3I (2.1%) and 18II + 5I (0.3%) were also found. A definite conclusion cannot be drawn from these results, because not all monosomics were studied.

Person (1956) reported that the levels of meiotic irregularities were much higher in heterozygous wheat materials than when observed in stable, more or less homozygous varieties. The percentage of cells showing desynapsis was the highest (35.8%) in F_1 intervarietal hybrids, but the extent of desynaptic cells showed progressive reduction during successive back crosses: (BC₁ = 20.2%, BC₂ = 19.2%, BC₃ = 15.7%, BC₄ = 15.3%, BC₅ = 12.7%, BC₆ = 10.2%, and BC₇ = 7.6%). Meiotic irregularities sometimes produce a monosomic plant deficient for a chromosome other than that deficient in the monosomic parent. Person (1956) termed this phenomenon "univalent shift." Analogous results were reported in three monosomics of oat, where early generations had a higher incidence of partial asynapsis, and the recovery of homozygosity in later back-cross generations resulted in a more stable meiosis (Rajhathy and Thomas, 1974).

One expects to observe n - 1 bivalents at metaphase-I in nullisomics, but it is not always observed. Nulli-8 of oat showed 5II + 30I, while nulli-9, 17, and 19 exhibited 20II (Morikawa, 1985). Sears (1954) recorded desynapsis in nulli-2A and nulli-3B. Desynapsis occurred in nulli-5D at low temperatures (12 to 15°C), but pairing was normal at 20 and 28°C (Riley et al., 1966b). By contrast, in nulli-5, homoeologous chromosome pairing prevailed that resulted in multivalent association. It was demonstrated that a pairing control gene, *Ph* is located on 5BL, because there was no homoeologous synapsis in the presence of 5BL, but it occurs in the presence of 5BS (Riley and Chapman, 1967).

4. Behavior of Univalent Chromosomes at Meiosis

Behavior of univalents at meiosis has been studied thoroughly in monosomics of *N. tabacum* (Olmo, 1936) and *T. aestivum* cv. Chinese Spring (Sears, 1952a,b; Morrison, 1953; Person, 1956). During the disjunction of chromosomes at anaphase-I to telophase-I or anaphase-II to telophase-II, the univalent may be included in either daughter nucleus or may lag, divide, or misdivide. In maize,

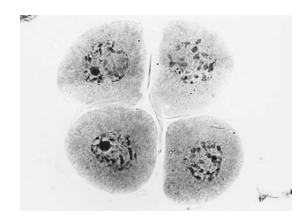


Figure 7.55 A quartet of microspores from a mono-6 plant. Nucleoli are present in the two haploid cells, each containing Chromosome 6, whereas nucleolar blebs are present in the two cells nullisomic for Chromosome 6, because they do not contain Chromosome 6, a NOR. (From Weber, D.F., Can. J. Genet. Cytol., 20, 97–100, 1978. With permission.)

the univalent passes to one pole at telophase-I in over half of the cells, because at prophase-II, 58.3% of the cell pairs contain nine chromosomes in one cell and ten in the other, 17.9% of the prophase-II cell pairs have 9 + 1 monod in each cell, indicating the univalent divided at telophase-I, and the rest have nine chromosomes, indicating that the monod chromosome was lost. (Weber, personal communication). This can be demonstrated by analyzing tetrads in mono-6. Of the four microspores, two left microspores contain one nucleolus (presence of Chromosome 6), and two right microspores are without nucleolus, indicating that Chromosome 6 was lost during meiosis. (Figure 7.55).

During the early phase of anaphase-I, bivalents disjoin normally, but the univalent remains more or less stationary at the equatorial plate and does not show any visible sign of longitudinal split; however, it undergoes division later, while chromosomes of the bivalents have already reached their respective poles. Thus, the daughter univalents lagging between the poles are eliminated from the macronuclei. In wheat, the univalent divided in 96% of sporocytes at first meiotic division (Sears, 1952a). Similarly, Darlington (1939b) found 98% misdivision of univalents in *Fritillaria kamtschatkensis*.

The misdivision of univalent mono-5A in four different strains of hexaploid wheat has been elaborated extensively in classic papers of Sears (1952a,b). The misdivision observed in wheat can be divided into three classes based on the number and kinds of arms going to one pole, as follows: (1) one normal chromatid passes to one pole, and two arms of the other chromatid either pass separately to the other pole or one or both remain acentric on the plate; (2) two identical arms move to one pole, and the other two arms either go to the other pole or one or both remain acentric on the plate; and (3) three arms go to one pole, and the fourth arm either moves to the other pole or remains on the plate. There is a fourth type, not observed in wheat, in which all four arms migrate to one pole.

The misdivision of mono-5A in Chinese Spring wheat was observed at telophase-I in 39.7% of cells, and at telophase-II, at least 29.3% of cells contained a laggard. The rate of misdivision of Chromosome 5A was influenced by the genetic background, because it ranged from 13.7% (Hope) to 39.7% (Chinese Spring) and also varied from chromosome to chromosome (Table 7.62). The misdivision of a univalent at telophase-I produced telocentrics and isochromosomes. At telophase-II, some of the isochromosomes misdivide again to yield additional telocentrics. The telocentrics and isochromosomes generated during microsporogenesis are lost or not transmitted through pollen. Thus, all of the telocentrics and isochromosomes recovered are produced during megasporogenesis. However, it has been suggested in *Fritillaria* (Darlington, 1940) and also in maize (Rhoades, 1940) that isochromosomes are produced through misdivision of telocentrics during pollen mitosis and can be transmitted.

		Misdivision at TI	
Chromosome	Variety	%	Total Number of Cells
5A	Chinese Spring	39.7	126
5A	Норе	13.7	95
5A	Thatcher	17.5	63
5A	Red Egyptian	23.6	191
2B	Red Egyptian	11.4	44
5B	Chinese Spring	26.7	15
7A	Hope	29.4	17

Table 7.62 Frequency of Misdivision of Various Chromosomes at Telophase-I in Several Wheat Cultivars

Source: From Sears, E.R., Chromosoma (Berlin), 4, 535-550, 1952a. With permission.

D. Fertility of Monosomics and Nullisomics

Pollen and seed fertility in monosomic lines depends upon the ploidy level, the chromosome involved, environmental conditions, and genetic background. Pollen fertility in mono-11 of tomato was 25%, and in tertiary monosomes, it ranged from 5 to 40% around a mode of 25%. As expected, the monosomics with higher pollen fertility generally set fruits with a higher number of seeds (Khush and Rick, 1966). Pollen fertility in maize monosomics ranged from 3.4 (mono-8) to 46.9% (mono-6). It is expected to observe 50% pollen fertility when the univalent chromosome in a monosomic plant is not lost during meiosis. However, observations on pollen fertility in maize monosomics revealed the following levels: mono-2 = 16.8%, mono-4 = 40.2%, mono-6 = 46.9%, mono-7 = 37.0%, mono-8 = 25.8%, mono-9 = 3.4%, and mono-10 = 43.9%. The univalent chromosome is frequently lost, producing a high frequency of microspores with nine chromosomes. (Weber, 1983). Pollen fertility (seed set) in soybean monosomics was mono-3, 8.8% (59 seeds) and mono-6, 20% (176 seeds). Disomic plant had 99% pollen fertility and produced 342 seeds (Xu, Singh, and Hymowitz, 2000b).

In *Nicotiana tabacum*, Clausen and Cameron (1944) recorded that most of the monosomics produced fewer seeds per capsule than disomics, with the exception of mono-A, which produced more seeds. Brown and Endrizzi (1964) found that seed and mote counts in *Gossypium* offered a reliable technique for recognition of monosomics, and that pollen fertility in monosomics proved to be inconclusive in distinguishing disomic and monosomic plants.

Seed set in monosomics of *Avena byzantina* cv. Kanota, studied by Morikawa (1985), ranged from 69.1 (mono-21) to 99.4% (mono-19). The disomic plants showed a seed set of 99.4%.

E. Breeding Behavior of Monosomics and Nullisomics

The monsomic condition is not transmitted through females or males in diploid species. Examples include *Datura stramonium* (Blakeslee and Belling, 1924a), *Oryza sativa* (Seshu and Venkataswamy, 1958), and *Zea mays* (Weber, 1983). In diploid species, the n - 1 spores abort and cannot compete with normal spores, because they are physiologically and genetically unbalanced. In cotton, male transmission of n - 1 gametes is rare (Endrizzi and Ramsay, 1979).

Unbalanced gametes are functional to a certain degree in polyploid species. The classical work of Sears (1953b, 1954) in hexaploid wheat cv. Chinese Spring demonstrates that after selfing monosomics, three types of offspring (disomic, monosomic, and nullisomic) occur in frequencies that vary. Some wheat plants lacking the chromosome concerned are obtained, and about 75% of the female gametes carry n = 20 chromosomes. The deviation from the expected 50% is due to the irregular behavior and resultant frequent loss of the unpaired chromosome during meiosis. Functioning male gametes predominantly contain n = 21 chromosomes, because pollen competition strongly favors pollen with the n = 21 chromosome number, and pollen of that constitution is involved in 90 to 99% of fertilization.

Female/Male	n (21-Chromosome Pollen) 0.96	n – 1 (20-Chromosome Pollen) 0.04
n (21-chromosome eggs) 0.25	2 <i>n</i> (42) plants 0.24	2 <i>n</i> – 1 (41) plants 0.01
<i>n</i> – 1 (20-chromosome eggs) 0.75	2 <i>n</i> – 1 (41) plants 0.72	2 <i>n</i> – 2 (40) plants 0.03

Table 7.63 Expected Transmission of the Monosomics in Triticum aestivum

Euploids (2n = 42) = 0.24; monosomics (2n = 41) = 0.73; nullisomics (2n = 40) = 0.03. *Source:* From Sears, E.R., *Am. Nat.*, 87, 245–252, 1953b. With permission.

The frequency of nullisomics in hexaploid wheat depends upon the degree of functioning of 20-chromosome pollen. A selfed progeny of monosomic plants should segregate, chromosomally, 24% disomics (2*n*), 73% monosomics (2*n* – 1), and 3% nullisomics (2*n* – 2) (Table 7.63). The frequencies of nullisomics in Chinese Spring wheat ranged from 0.9 (nulli-5B, nulli-6B) to 7.6% (nulli-3B) (Sears, 1954).

Tsunewaki and Heyne (1960) examined the transmission rate of 21 monosomics cytologically in five monosomic series established in five wheat cultivars. The frequencies of the monosomics ranged from 52.2 (mono-2B) to 89.2% (mono-3B).

Avena sativa is also an allohexaploid, but it differs from hexaploid wheats in degree of an euploid tolerance (Morikawa, 1985). Nullisomics were not found in the 17 monosomic lines, and monote-losomics (2n = 40 + 1 telo) with low frequencies were obtained in the progenies of four monosomics. The selfed progenies of monosomics threw monosomic plants at a high frequency ranging from 35.5 (mono-12) to 97.8% (mono-5).

The female transmission rate of monosomics in *N. tabacum* (2n = 4x = 48) ranged from 5.1 (mono-W) to 81.9% (mono-E). Monosomics that exhibited high ovular abortion showed a strongly depressed transmission rate (Clausen and Cameron, 1944).

The results discussed thus far suggest that the frequency of monosomics does not fall within the range of expectation. The actual frequency of monosomics is either too high or too low. It can be attributed to the elimination of the univalent. Greenleaf (1941) investigated cytologically mono-P of *N. tabacum* with the objective of determining the cause of aberrant transmission rates. Two main causes were established: (1) slow rate of development of the n - 1 embryo sacs and (2) a high frequency of aborted embryo sacs prior to the attainment of the eight-nucleate condition. According to Rajhathy and Thomas (1974), the genetic background of the monosomic lines and the particular chromosome involved have a significant effect on breeding behavior of monosomics.

Nullisomics are expected to be stable and to breed true. Sears (1954) examined selfed progenies of nine nullisomics of Chinese Spring wheat. With the exception of nulli-3B, the eight nullisomics were fairly stable (Table 7.64). Nulli-3 transmitted aberrant plants that helped in isolating monosomics. The stability of all nullisomics has not been further studied (B.S. Gill, B. Friebe, and P.P. Jauhar, personal communication).

F. Genetic Studies

The concept of using monosomics for associating genetic markers with a particular chromosome was first discovered by Bridges (1921) from the studies of "diminished" monosome (mono-4) in *Drosophila melanogaster*. Since then, monosomics (primary) have effectively been utilized in genetic and linkage studies in tomato (Khush and Rick, 1966), maize (Weber, 1983), tetraploid tobacco (Clausen and Cameron, 1944), tetraploid cotton (Endrizzi, 1963; Endrizzi and Ramsay, 1979; Endrizzi and Ray, 1991), and in hexaploid wheat (Unrau, 1950; Sears, 1953b; Khush, 1973).

As in simple primary trisomics, genes are tested against whole chromosomes when monosomics and nullisomics are used. By the monosomic method, not only are single genes assigned to a chromosome, but multigenes are also located.

Nullisome	Number of Offspring Grown	2 <i>n</i> = 40	2 <i>n</i> = 39	2 <i>n</i> = 41	% 2 <i>n</i> – 2
1B	68	65	1	2	95.6
7B	66	61	3	2	92.4
7A	15	10	3	2	66.7
ЗA	9	9	0	0	100.0
1A	12	11	0	1	91.7
1D	4	4	0	0	100.0
6D	2	2	0	0	100.0
7D	15	13	1	1	86.7

Table 7.64 Chromosome Constitution of Selfed Progenies of Nullisomic Plants in Triticum aestivum

Source: From Sears, E.R., Mo. Agric. Exp. Stn. Res. Bull., 572, 1-58, 1954. With permission.

1. Locating Recessive Genes

All monosomic stocks carrying a dominant allele, such as *DD*, are crossed as a female by disomic stocks with the recessive trait (*dd*) as a male. Monosomic plants produce two kinds of gametes: n(D), n - 1(-); and disomic male plants are expected to produce only one type, n(d), of gametes. In critical crosses, all F₁ monosomic offspring are recessive [*d*(-)], and disomic plants are dominant (*Dd*). This suggests that a recessive mutant is located in the missing chromosome. Thus, monosomics give results like sex-linked characters in animals. A monosomic is hemizygous for all genes on the univalent (pseudodominance).

In noncritical combinations, all F_1 offspring (monosomics, disomics) should show the dominant character, and the F_2 should segregate in a normal disomic 3 dominant: 1 recessive fashion. However, n - 1 gamete does not function in diploid species. Thus, this technique is applicable only for tetraploid and hexaploid crops.

Bridges (1921) crossed *Drosophila melanogaster* mono-4 (diminished) as a female with the recessive mutants bent and eyeless as male. All F_1 flies that were mono-4 were bent or eyeless. However, crosses of mono-4 with recessive markers from Chromosomes 1, 2, and 3 showed all F_1 flies with dominant character.

2. Locating Dominant Genes

All monosomic stocks carrying recessive characters (d-) are crossed by disomic stocks that carry the dominant allele (DD). All F₁ disomic and monosomic plants will show the dominant trait. Backcross or F₂ progeny will identify the critical cross. In F₂, all recessive homozyotes are expected to be nullisomics if a gene is located on the missing chromosome; otherwise, nullisomic plants should segregate for dominant and recessive characters.

Unrau (1950) associated seven of nine genes with a particular chromosome in wheat by monosomic and nullisomic methods. For example, red glumes in Federation-41 wheat are dominant, and in crosses with Chinese Spring, segregates in a 3(red):1 (white) ratio. Federation-41 wheat was crossed onto 17 monosomics of Chinese Spring wheat. In F₂, all except mono-1B, showed a 3:1 ratio (Table 7.65). The proportion of white glume was very low, and white glumed plants were nullisomic. This conclusively demonstrates that a gene for red glume color is associated with Chromosome 1B.

3. Locating Duplicate Genes

Duplicate genes can be located in the chromosomes by monosomic analysis in polyploid species. In *Nicotiana tabacum* (2n = 4x = 48), Clausen and Cameron (1944) assigned hairy filament to

F1 Plants	Red Glumes	White Glumes	% White Glumes ^a	Total
Mono-1B Total (excluding mono-1B) of 16 monosomics	528 10,975	38 3462	6.7 24.0	566 14,437

Table 7.65 Summary of the Segregation for Glume Color in F₂ from Crosses between 17 Monosomics of Chinese Spring and Federation 41, Red Glume Color

^a All white glumed plants were nullisomics.

Source: From Unrau, J., Sci. Agric., 30, 60-89, 1950. With permission.

Chromosomes A (subgenome *tomentosa*) and O (subgenome *sylvestris*) and yellow-burley to Chromosomes B (subgenome *tomentosa*) and O by means of monosomics.

Association of hairy filaments (*hf1*, *hf2*) to their respective chromosomes was accomplished by examining the segregation of smooth versus hairy filament in F_2 . A ratio of three normal:one hairy filament was observed in the mono-A fraction, and a 15 normal:1 hairy-filament ratio was observed in the disomic fraction. If a gene is not associated, it should segregate in a 15:1 ratio. Thus, hairy filament showed association with Chromosomes A and O, because a high proportion of hairy filament, and with the other monosomics, the segregation is in satisfactory agreement with a 15:1 ratio for duplicate genes (Table 7.66). Similarly, they associated another duplicate gene, yellow-burley (*yb1*, *yb2*), with mono-B [F_2 = green, 66: yellow-burley, (*yb1*), 23] and with mono-O (BC₁ = green, 19: yellow-burley (*yb2*), 17]. Thus, *yb1* was located in B chromosome of subgenome *tomentosa* and *yb2* in the O chromosome of subgenome *sylvestris*.

Thus, modifications in ratios in F_2 and BC_1 generations, produced from heterozygous F_1 monosomics, help to associate a gene or genes with particular chromosomes.

IV. ALIEN ADDITION LINES

A. Introduction

The transfer of agriculturally important traits from alien species to cultigens by interspecific and intergeneric crosses, known as wide hybridization, has been demonstrated in numerous crops (Goodman et al., 1987). Thus, wide hybridization is a valuable tool for creating genetic variability

Monosomic Types	Smooth Filament	Hairy Filament	% Hairy Filament
A	25	15	60.0
В	47	1	2.1
С	44	4	9.1
D	40	2	5.0
E	33	2	6.1
G	36	0	0.0
Н	38	2	5.3
М	39	5	12.8
Ν	41	1	2.4
0	30	15	50.0
Р	37	2	5.4
S	37	1	2.7
Totals (excluding A and O)	394	20	5.1
Expected 15:1	388	26	6.7

 Table 7.66 Segregation of Hairy Filament in F2 after Selfing F1 Monosomics Heterozygous for Hairy Filament in Nicotiana tabacum

Source: From Clausen, R.E. and Cameron, D.R., Genetics, 29, 447–477, 1944. With permission.

in plant breeding by broadening the germplasm base of the cultigens (Smith, 1971; Harlan, 1976; Hadley and Openshaw, 1980; Stalker, 1980; Zenkteler and Nitzsche, 1984; Tanksley and McCouch, 1997; Zamir, 2001). Exploitation of wild relatives of the crop plants is often hampered because of poor crossability, early embryo abortion, hybrid seed inviability, hybrid seedling lethality, and hybrid sterility due to low chromosome pairing. However, these barriers have been overcome by (1) the assemblage of diverse germplasm; (2) application of growth hormones to reduce embryo abortion; (3) improved culture conditions; (4) restoration of the seed fertility by doubling the chromosomes of sterile F_1 hybrids; and (5) utilization of bridge crosses where direct crosses are not possible. Introgression of useful genetic traits from wild relatives (donor parent) to cultigens (recipient parent) is achieved by producing alien addition, substitution, and translocation lines.

The objective of the following two sections is not to report a comprehensive review on alien addition and alien substitution lines. The literature on wheat and its allied species and genera are so voluminous that it is beyond reach to review it all here.

B. Production of Alien Addition Lines

Monosomic alien addition lines (MAALs) and disomic alien addition lines (DAALs) have been produced in polyploid and diploid species according to the procedure described by O'Mara (1940). Modifications in the technique depended on ploidy levels and the nature of the crops. In some species, all the possible alien addition lines (AALs) are available, while in others, a complete set is lacking (Table 7.67). The procedure involves the production of interspecific or intergeneric F_1 hybrids, induction of an amphiploid, generation of BC₁, isolation of monosomics in BC₂ and BC₃, and selection of disomic additions after selfing of the MAAL. The term "alien addition races" was originally proposed by Clausen (Gerstel, 1945).

1. Hexaploid Species

Hexaploid wheat (2n = 6x = 42, genome formula AABBDD) and diploid rye (2n = 2x = 14, RR) crosses result in F₁ polyhaploid (2n = 4x = 28, ABDR) sterile plants. Sometimes seeds are obtained because of unreduced gamete formation. Generally, octoploid (2n = 8x = 56, AABBDDRR) triticale is synthesized by doubling the chromosomes with colchicine. The synthesized octoploid triticale is fertile and is back crossed to wheat (BC₁), resulting in heptaploid (2n = 7x = 49, AABBDDR) plants. The heptaploid plants mostly exhibit 21II + 7I chromosome pairing at metaphase-I. The seven rye univalents are randomly excluded or included in the gametes. MAALs (2n = 42W + 1R — -7R) are isolated either after selfing the heptaploid or the heptaploid is again crossed by wheat (BC₂). DAALs are isolated in the selfed population of MAALs (Figure 7.56).

In general, the higher ploidy (female) × lower ploidy (male) crosses are more compatible than the reciprocals. However, there are a few exceptions (McFadden and Sears, 1944, 1946; Röbbelen and Smutkupt, 1968). Thomas (1968) isolated six of the possible seven MAALs of *Avena hirtula* (2n = 2x = 14, As) from crosses to *A. sativa* (2n = 6x = 42, AACCDD) by way of an *A. hirtula* (female) by *A. sativa* (male) cross (Table 7.68). Islam, Shepherd, and Sparrow (1981) observed 1.3% seed set in a *Triticum aestivum* cv. Chinese Spring (2n = 42) (female) × *Hordeum vulgare* cv. Betzes (2n = 14) (male) cross, while 15.4% seed set was obtained with the reciprocal cross.

Occasionally, direct interspecific or intergeneric crosses are not successful. In each case, bridge crosses may be utilized (Sears, 1953a; DeVerna, Chetelat, and Rick, 1987). Sears (1953a) added seven chromosomes of *Haynaldia villosa* to hexaploid wheat by using a tetraploid (2n = 4x = 28) wheat (*T. dicoccoides*) as a bridging species. The amphidiploid (AABBVV) was crossed to *T. aestivum* and back crossed to *T. aestivum*. Two of the 53 back-cross progenies were AABBDDV. Subsequently, Hyde (1953) isolated five of the possible seven DAALs and six of the possible seven MAALs. Friebe, Cermeño, and Zeller (1987) examined *T. aestivum-Dasypyrum villosum* (formerly *H. villosa*) addition lines of Sears using C-banding technique and identified six DAALs.

				AALs	(nos)	
Recipient Species	n	Donor Species	n	MAALs	DAALs	Authority
Allium cepa	8	Allium fistulosum	8	4	_	Peffley et al., 1985
Allium fistulosum	8	Allium cepa	8	8	—	Barthes and Ricroch, 2001; Shigyo et al., 1996
Avena sativa	21	Avena barbata	14	1	1	Thomas, Leggett, and Jones, 1975
Avena sativa	21	Avena hirtula	7	6	4	Thomas, 1968
Avena sativa	21	Avena strigosa	7	1	—	Dyck and Rajhathy, 1963
Avena sativa	21	Zea mays	10	_	5	Ananiev et al., 1997
Beta vulgaris	9	Beta patellaris	9	1	_	Heijbroek, Roelands, and De Jong, 1983; Speckmann, De Bock, and De Jong, 1985
Beta vulgaris	9	Beta patellaris	9	9	—	Mesbah, deBock, and Sandbrink, 1997
Beta vulgaris	9	Beta procumbens	9	1	—	Savitsky, 1975
Beta vulgaris	9	Beta procumbens	9	9	_	Lange et al., 1988
Beta vulgaris	9	Beta procumbens	9	3	—	Speckmann, De Bock, and De Jong, 1985
Beta vulgaris	9	Beta webbiana	9	9	—	Reamon-Ramos and Wricke, 1992
Beta vulgaris	9	Beta corolliflora	19	9	—	Gao, Guo, and Jung, 2001
Brassica campestris	10	Brassica alboglabra	9	4	_	Cheng et al., 1997
Brassica campestris	10	Brassica oleracea	9	8	8	Quiros et al., 1987
Brassica napus	19	Brassica campestris	10	1	_	McGrath and Quiros, 1990
Brassica napus	19	Brassica nigra	8	?		Jahier et al., 1989
Brassica napus	19	Brassica nigra	8	6		Chévre et al., 1991
Brassica oleracea	9	Brassica nigra	8	5		Chévre et al., 1997
Raphanus sativus	9	Brassica oleracea	9	7	—	Kaneko, Matsuzawa, and Sarashima, 1987
Cucurbita moschata	20	Cucurbita palmata	20	6	—	Graham and Bemis, 1979
Diplotaxis erucoides	7	Brassica nigra	8	7	—	This, Ochoa, and Quiros,1990
Glycine max	20	Glycine tomentellla	39	22	—	Singh, Kollipara, and Hymowitz, 1998b
Gossypium hirsutum	26	Gossypium sturtianum	13	4	—	Rooney, Stelly, and Altman, 1991
Hordeum vulgare	7	Hordeum bulbosum	7	2	—	Thomas and Pickering, 1988
Lycopersicon esculentum	12	Solanum lycopersicoides	12	6	—	DeVerna, Chetelat, and Rick, 1987
Lycopersicon esculentum	12	Solanum lycopersicoides	12	12	_	Chetelat et al., 1998
Lolium multiflorum	14	Festuca drymeja	7	4	—	Morgan, 1991
Nicotiana tabacum	24	Nicotiana glutinosa	12	1	1	Gerstel, 1945
Nicotiana tabacum	24	Nicotiana paniculata	10	1	1	Lucov, Cohen, and Moav, 1970
Nicotiana tabacum	24	, Nicotiana plumbaginifolia	10	3	3	Cameron and Moav, 1957
Nicotiana plumbaginifolia	10	Nicotiana sylvestris	12	12	—	Suen et al., 1997
Oryza sativa	12	Oryza officinalis	12	12	—	Shin and Katayama, 1979
Oryza sativa	12	Oryza officinalis	12	12	_	Jena and Khush, 1989
	12	Oryza australiensis	12	8		Multani et al., 1994

Table 7.67 A Partial List of Alien Addition Lines

continued

Recipient SpeciesnDonor SpeciesnAALs (nos)Triticum durum14Agropyron elongatum776Triticum durum14Aegilops umbellulata772Triticum durum14Dasypyrum villosum76	Authority Mochizuki, 1962
Triticum durum14Agropyron elongatum776Triticum durum14Aegilops umbellulata772	
elongatum Triticum durum 14 Aegilops 7 7 2 umbellulata	WOCHIZUKI, 1902
umbellulata	
Triticum durum 14 Dasvpvrum villosum 7 6 —	Makino, 1976, 1981
	Blanco, Simeone, and Resta, 1987
Triticum durum 14 Secale cereale 7 2 —	Sadanaga, 1957
Triticum durum 14 Triticum tauschii 7 6 —	Makino, 1981
Triticum durum 14 Triticum tauschii 7 7 —	Dhaliwal et al., 1990
Triticum turgidum 14 Hordeum chilense 7 7 —	Fernandez and Jouve, 1988
Triticum turgidum 14 Triticum distichum 14 8 —	Fominaya et al., 1997
Triticum aestivum 21 Aegilops caudata 7 — 6	Friebe et al., 1992
Triticum aestivum21Aegilops geniculata14113	Friebe, Tuleen, and Gill, 1999
Triticum aestivum 21 Aegilops markgrafii 7 — 5	Peil et al., 1998
Triticum aestivum 21 Aegilops ovata 14 — 4	Landjeva and Ganeva, 1999
Triticum aestivum21Aegilops searsii777	Pietro, Tuleen, and Hart, 1988
Triticum aestivum 21 Aegilops speltoides 7 — 7	Friebe et al., 2000
Triticum aestivum 21 Aegilops uniaristata 7 — 5	Miller et al., 1997
Triticum aestivum 21 Aegilops variabilis 14 – 14	Friebe et al., 1996b
Triticum aestivum 21 Aegilops 7 1 1 sharonensis	Miller et al., 1982a
Triticum aestivum21Aegilops comosa711	Riley, Chapman, and Macer, 1966a
Triticum aestivum 21 Agropyron ciliare 14 1 6	Wang et al., 2001
Triticum aestivum 21 Agropyron 7 7 — elongatum	Dvořák and Knott, 1974
Triticum aestivum 21 Agropyron 7 — 1 elongatum	Konzak and Heiner, 1959
Triticum aestivum 21 Haynaldia villosa 7 6 5	Hyde, 1953
Triticum aestivum 21 Hordeum chilense 7 6 —	Miller et al., 1982a
Triticum aestivum21Hordeum vulgare756	Islam, Shepherd, and Sparrow, 1981
Triticum aestivum 21 Roegneria ciliaris 14 1 6	Wang et al., 2001
Triticum aestivum 21 Secale cereale 7 3 3	O'Mara, 1940
Triticum aestivum21Secale cereale734	Riley and Chapman, 1958b
Triticum aestivum 21 Secale cereale 7 — 7ª	Riley and Macer, 1966
Triticum aestivum21Secale cereale775	Evans and Jenkins, 1960
Triticum aestivum21Secale cereale77	Driscoll and Sears, 1971
Triticum aestivum21Secale cereale766	Bernard, 1976
Triticum aestivum 21 Thinopyrum 21 1 1 intermedium	Wienhues, 1966
Triticum aestivum 21 Thinopyrum 21 — 6 intermedium	Forster et al., 1987
Triticum aestivum 21 Thinopyrum 21 — 6 intermedium	Larkin et al., 1995
Triticum aestivum 21 Thinopyrum 21 — 6 bessarabicum	William and Mujeeb- Kazi, 1995
Triticum aestivum 21 Triticum 7 7 7ª longissimum	Feldman and Sears, 1981
Triticum aestivum 21 Triticum ovatum 14 4 —	Mettin et al., 1977

Table 7.67 (continued) A Partial List of Alien Addition Lines

				AALs	(nos)	
Recipient Species	n	Donor Species	n	MAALs	DAALs	Authority
Triticum aestivum	21	Triticum umbellulatum	7	1	1	Sears, 1956b
Triticum aestivum	21	Triticum umbellulatum	7	6	6	Kimber, 1967
Triticum aestivum	21	Triticum umbellulatum	7	1	5	Friebe et al., 1995a
Triticum aestivum	21	Triticum variabile	14	9	—	Jewell and Driscoll, 1983

Table 7.67 (continued) A Partial List of Alien Addition Lines

^a Six distinct DAALs by C-banding technique.

Source: From Hueros et al., Genome, 34: 236-240, 1991. With permission.)

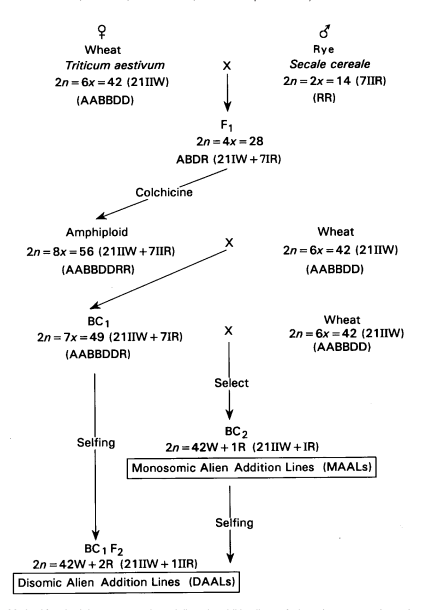


Figure 7.56 Method for obtaining monosomic and disomic addition lines of wheat (2n = 6x = 42)-rye (2n = 2x = 14).

	Numberof		Proportion	of Plants	with Chro	mosome N	lumbers	
Generation	Plants	42	42 + telo	43	44	45	46	47
BC ₂	189	0.63	0.01	0.26	0.06	0.04	0.0	0.0
BC ₁ -selfed	39	0.18	0.02	0.33	0.28	0.13	0.03	0.03

Table 7.68 Frequency of Avena Plants with Various Chromosome Numbers in BC₂ Population and Progeny of Selfed BC₁ Plants

Source: From Thomas, H., Can. J. Genet. Cytol., 10, 551–563, 1968. With permission.

Transfer of useful genes of economic importance to wheats from alien species has been successful compared to other important crops. Shepherd and Islam (1988) list the following sequence of 24 species utilized for introgressing desired genes by producing alien additions, substitutions, and translocations: (1) *Secale cereale*, (2) *S. montanum*, (3) *Aegilops umbellulata*, (4) *Ae. variabilis*, (5) *Ae. comosa*, (6) *Ae. mutica*, (7) *Ae. longissima*, (8) *Ae. sharonensis*, (9) *Ae. bicornis*, (10) *Ae. searsii*, (11) *Ae. ventricosa*, (12) *Agropyron elongatum*, (13) *Ag. intermedium*, (14) *Ag. glaucum*, (15) *Ag. trichophorum*, (16) *Haynaldia villosa*, (17) *Elymus trachycaulus*, (18) *Hordeum vulgare*, (19) *H. chilense*, (20) *Triticum urartu*, (21) *T. thaoudar*, (22) *T. monococcum*, (23) *T. timopheevi*, and (24) *T. aestivum* added to *S. cereale*.

2. Tetraploid Species

A complete set of the seven possible MAALs was obtained from crosses of tetraploid wheat, *T. durum* (2n = 4x = 28, AABB) with *Agropyron elongatum* (2n = 2x = 14) (Mochizuki, 1962) and from *T. durum* with *Ae. umbellulata* (2n = 2x = 14, C^u C^u) (Makino, 1976) crosses based on principles utilized at the hexaploid level. Makino (1976, 1981) obtained allotriploid F₁ hybrids ($2n = 3x = 21 \text{ ABC}^{u}$) between *T. durum* and *Ae. umbellulata*. The allotriploid hybrid plants were back crossed with pollen from *T. durum*, and the pentaploid plants (2n = 5x = 35, AABBC^u), obtained by the fertilization of unreduced female gametes (ABC^u) with normal male gametes (AB), were again back crossed to *T. durum*. In the BC₂, seven possible MAALs (2n = 29) were isolated. In a similar way, Blanco, Simeone, and Resta (1987) produced six of the possible seven MAALs from crosses of *Dasypyrum villosum* (2n = 2x = 14, VV) to *T. durum*.

3. Diploid Species

Monosomic alien addition lines (allotrisomics) have been produced in *Beta vulgaris* (2n = 2x = 1 8) (Savitsky, 1975; Heijbroek, Roelands, and De Jong, 1983; Speckmann, De Bock, and De Jong, 1985; Lange et al., 1988), *Oryza sativa* (2n = 2x = 24) (Shin and Katayama, 1979; Jena and Khush, 1989) and *Lycopersicon esculentum* (DeVerna, Chetelat, and Rick, 1987). Savitsky (1975) added a single chromosome of *B. procumbens* (2n = 2x = 18) to *B. vulgaris*. The added *B. procumbens* chromosome carried a gene for nematode (*Heterodera schachtii*) resistance.

DeVerna, Chetelat, and Rick (1987) produced the 12 possible MAALs (2n = 25) from crosses of *Solanum lycopersicoides* (2n = 2x = 24, SS) to *Lycopersicon esculentum* (2n = 2x = 24, LL) by using *L. pennellii* (2n = 2x = 24, PP) as a bridging species.

Monosomic alien addition lines of *Oryza sativa* (AA) and *O. officinalis* (CC) were produced by Shin and Katayama (1979) and by Jena and Khush (1989). The initial cross made by Shin and Katayama (1979) was between a tetraploid (2n = 4x = 48) *O. sativa* (AAAA) and a diploid (2n = 24) strain of *O. officinalis* (CC). The allotriploid (2n = 3x = 21, AAC) F₁ plants were back crossed to *O. sativa*. The progeny segregated plants with chromosome numbers ranging from 2n = 24 to 2n = 54 (Table 7.69).

Jena and Khush (1989) crossed diploid *O. sativa* and *O. officinalis*. The F_1 (AC) plants were obtained through embryo rescue technique. An amphidiploid (AACC) was not produced, but F_1

Chromosome Numbers	Number of Plants (%)ª	Number of Plants (%) ^b
24	51 (12.6)	25 (26.6)
25	33 (8.1)	40 (42.5)
26	33 (8.1)	11 (11.7)
27	60 (14.8)	10 (10.6)
28	81 (20.0)	4 (4.3)
29	69 (17.0)	3 (3.2)
30	50 (12.3)	1 (1.1)
31	12 (3.0)	
32	11 (2.7)	
33	3 (0.7)	
36	1 (0.2)	
54	1 (0.2)	
Total	405	94

Table 7.69	Chromosome Numbers in AAC ×
	AA Crosses of Oryza sativa and O.
	officinalis

^a Shin, Y.-B. and Katayama, T., *Jpn. J. Genet.*, 54, 1–10, 1979.

^b Jena, K.K. and Khush, G.S., *Genome*, 32, 449–455, 1989.

plants were back crossed to *O. sativa*. A growth hormone mixture, gibberellic acid (GA3) + naphthalene acetic acid (NAA) + kinetin (K) in the proportion of 100 + 25 + 5 mg/L, was sprayed on with an atomizer 24 h postpollination, two times a day for 5 days. Of the 41,437 spikelets pollinated, only 539 (1.3%) set seeds. A total of 367 BC₁ plants were recovered through culture (one-quarter strength MS medium). Of the 367 BC₁ plants, 357 were allotriploids, and ten were hypotriploids. This result indicates that some of the unreduced female gametes contained all the A and C genome chromosomes and produced allotriploid zygotes with AAC constitution when fertilized with gametes carrying an A chromosome. Allotriploid F₁ plants were back crossed to *O. sativa*, and 94 plants were recovered. Chromosome numbers ranged from 2n = 24 to 30, much narrower than those recovered (2n = 24 to 54) by Shin and Katayama (1979) (Table 7.69). Twelve morphological types, each containing a complete chromosome complement of *O. sativa* and a single different chromosome of *O. officinalis* were isolated and identified.

The wealth of untapped genetic diversity of wild perennial *Glycine* species has not been exploited in the soybean breeding programs because of their extremely low crossability with G. max and a need to employ in vitro embryo rescue methods to produce F_1 , BC₁, and BC₂ plants (Singh et al., 1998). Wide hybrids produced in the genus *Glycine* are few compared with those in cereals. Most soybean researchers were unable to produce beyond the amphidiploid stage. Singh et al. (1998) produced, for the first time, BC₁, BC₂, and BC₃, and monosomic alien addition lines (MAALs) from G. max (2n = 40; GG) female \times G. tomentella (2n = 78; DDEE) male \rightarrow F₁ (2n= 59; GDE) \rightarrow Colchicine treatment \rightarrow amphidiploid (2n = 118; GGDDEE) female \times soybean cv. Clark 63 male \rightarrow BC₁ [2n = 76 (expected 2n = 79); GGDE] female × soybean cv. Clark 63 male \rightarrow BC₂, BC₃, BC₄ \rightarrow produced plants with 2n = 40, 40 + 1, 40 + 2, 40 + 3. Singh et al. (1998) distinguished morphologically 22 MAALs. The main hurdle was to produce BC₁ plants. It consumed 5 years of hybridization, perseverence, and motivation to break the unthinkable barrier. Similar observation was recorded by Jacobsen et al. (1994) to obtain BC_1 plant from potato and tomato somatic hybrids. Their experience is informative to readers. The first back cross is the most difficult step in starting a successful back-cross program. The addition of an extra chromosome of G. to the 2n soybean complement modified several morphological traits, including flowering habit, plant height, degree of pubescence, seed fertility, number of seed per pod, and plant, pod, and seed color and seed yield. Disomic alien addition lines (DAALs) were isolated, but plants died prematurely. The female transmission rate of an extra *G. tomentella* chromosome in MAALs averaged 36.5%, and male transmission averaged 11.7%.

C. Identification of Alien Addition Lines

Alien addition lines can be identified by gross morphology, karyotype of the added chromosomes, meiotic chromosome pairing in F_1 plants obtained by intercrossing two DAALs, isozyme analysis, and by genetic tests.

1. Morphological Identification

Monosomic and disomic alien addition lines can be identified morphologically from the recipient parent and among themselves, because the extra added chromosome(s) modifies the specific vegetative and spike morphological features. The alterations may be qualitative and quantitative. Most of the modifications are due to the interaction between genes of the recipient and donor parents, and each chromosome has a specific effect on plant morphology and fertility.

Each disomic wheat-rye addition expresses diagnostic morphological traits such as stature of plants, color of leaves, compact and lax spikes, tapered head, hairy-neck, awned, and seed fertility. (O'Mara, 1940; Riley and Chapman, 1958b; Evans and Jenkins, 1960). Figure 7.57 shows spikes of seven "Chinese Spring" wheat-"Imperial" rye disomic alien addition lines together with parents and amphiploids. Based on morphological features of spikes, seven DAALs can be easily identified and distinguished. Similarly, spikes of ditelosomic alien addition lines of "Chinese Spring" wheat-"Imperial" rye exhibit contrasting distinguishing features. It should be noted that long arms of rye chromosomes alter spike morphology more than that of short arms (Figure 7.58).

The wheat-barley disomic addition lines produced by Islam, Shepherd, and Sparrow (1981) differed morphologically from "Chinese Spring" and from each other. An extra pair of barley chromosomes expressed diagnostic characters in hexaploid wheat background identical to the effect of an extra chromosome in primary trisomic condition. Primaries Bush (Triplo 1) and Slender (Triplo 2) were similar to DAALs 1 and 2, respectively. DAAL 5 was not isolated because of complete seed sterility. In contrast, primary Pseudonormal (Triplo 5) was quite fertile (Tsuchiya, 1960a, 1967).

The addition of single chromosomes (monosomic = MAAL) from wild diploid wheats to tetraploid (2n = 4x = 28) wheat alters morphological features of plants more drastically than those recorded in MAALs of hexaploid wheat. Several sets have been produced in *T. durum* (Table 7.67).

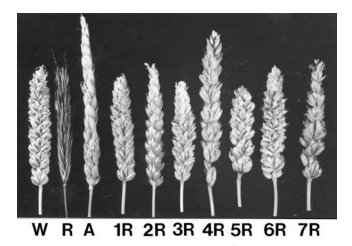


Figure 7.57 Spikes of "Chinese Spring" wheat (W), "Imperial" rye (R), amphiploid (A), and seven disomic (2n = 42W + 2R) addition lines. (Courtesy of B. Friebe.)



Figure 7.58 Spikes of 11 ditelosomic "Chinese Spring" wheat—"Imperial" rye (2*n* = 42 + 2telo) addition lines. L = long arm; S = short arm. (Courtesy of B. Friebe.)

Makino (1976) produced and identified the seven possible monosomic additions (C^u1 to C^u7) derived from *T. durum* and *Ae. umbellulata*. The C^u1 line was distinct from others in that it had a waxless spike. MAALs C^u2 and C^u3 carried the satellite chromosomes; C^u2 expressed reduced culm and top internode length, reduced ear density, and increased rachis length, while C^u3 had only reduced culm and top internode lengths. MAAL C^u4 was characterized by red seed and carried a dense spike. The culm length of line C^u5 was short (84.7 cm), as was MAAL C^u2 (81.9 cm), suggesting that genes for short culm of *Ae. umbellulata* are probably located on Chromosomes C^u2 and C^u5. MAAL C^u6 expressed thicker culm (3.2 mm) than *T. durum* (2.9 mm), but C^u7 did not show specific characteristics.

The plants of a DAAL (2n = 4x = 50) from *Nicotiana tabacum* by *N. glutinosa* crosses were slightly reduced in overall length, were later in maturity, and carried greater compactness of the inflorescence and shorter internodes than normal plants (Gerstel, 1945).

The addition of an alien chromosome to a diploid chromosome complement modifies the morphological features in the same way as observed in autotrisomics. Jena and Khush (1989) produced 12 possible MAALs from *Oryza sativa* by *O. officinalis* crosses. The addition lines differed from their diploid sibs by an array of morphological traits and resembled the 12 simple primary trisomics of *O. sativa*.

2. Cytological Identification

The alien chromosomes in addition lines can be identified from observations of chromosome morphology (total length, relative length, arm ratio) and cytological chromosome markers (heterochromatic and euchromatic distribution, heterochromatic knobs, kinetochore position, presence of satellite).

O'Mara (1940) and Riley and Chapman (1958b) designated wheat-rye addition lines arbitrarily. Evans and Jenkins (1960) designated DAALs derived from wheat-rye crosses based on karyotype analysis. For example, DAAL I carried the longest chromosome and DAAL VII the satellited chromosome. Gupta (1971, 1972) suggested designating wheat-rye addition lines on the basis of homoeologous relationships between rye and wheat chromosomes related mainly to substitution-compensation. The rye chromosomes in wheat-rye addition lines were numbered 1R to 7R. This nomenclature is far from being complete and is occasionally controversial due to the lack of correspondence at the gene level and in naturally occurring translocations (Sybenga, 1983). Thus, the ability of an alien chromosome to be matched against the wheat homoeologue grouping depends on the genetic equivalence of the alien chromosome to the wheat group.



Figure 7.59 Giemsa C-banded somatic metaphase of a disomic wheat ("Holdfast")–rye ("King II") addition line showing 2n = 42 wheat + 2 satellited rye (V = 1R) chromosomes (arrow). (From Singh, R.J. and Röbbelen, G., Z. Pflanzenzüchtg., 76, 11–18, 1976. With permission.)

By utilizing the Giemsa C-banding technique, Darvey and Gustafson (1975) identified rye chromosomes in four sets of wheat–rye disomic addition lines (Imperial, Dakold, King II, Petkus) and verified the homoeologous chromosome (1R to 7R) relationships established by genetic studies of seed shriveling in wheat and triticale (Darvey, 1973). Singh and Röbbelen (1976) examined Holdfast–King II wheat–rye disomic addition lines by the Giemsa C-banding technique. The addition line V was found to carry a pair of nucleolus organizer rye (1R) chromosomes (Figure 7.59), and addition lines II and VI each showed a deletion in its short arm.

Giemsa C- and N-banding and genomic *in situ* hybridization (GISH) techniques have helped identify wheat–rye addition lines precisely, and it has been determined that Chinese Spring–Imperial additions are complete. Furthermore, Giemsa C-banding also identified ditelosomic additions of Chinese Spring–Imperial. Telocentric chromosomes showed characteristic telomeric and intercalary heterochromatic bands (Figure 7.60).

3. Meiosis in Alien Addition Lines

It is expected that a pair of alien chromosomes in DAALs should form a bivalent at metaphase-I. However, it is not always recorded (Gerstel, 1945; Hyde, 1953; Riley and Chapman, 1958b; Bernard, 1976; Kimber, 1967; Islam, Shepherd, and Sparrow, 1981; Orellana, Cermeño, and Lacadena, 1984).

Orellana, Cermeño, and Lacadena (1984) examined meiotic pairing in "Chinese Spring"–"Imperial" and "Holdfast"–"King II" addition and substitution lines using the C-banding technique. They observed a decrease in pairing of wheat and rye homologous chromosomes and suggested that chromosome pairing in DAALs and substitution lines is a complex process influenced by factors such as genes controlling meiotic pairing, constitutive heterochromatin, and cryptic wheat–rye interactions.

In wheat–barley DAALs, chromosome pairing was influenced by the added barley chromosomes (Islam, Shepherd, and Sparrow, 1981). Synapsis was reduced in DAAL A (76.9%) and D (84.8%), while it was almost normal (22II) in other DAALs (B = 91.3%; C = 90.2%; E = 98.6%; F = 90%). The wheat cultivar Chinese Spring showed 21II in 95.9% of the sporocytes. It should be noted that MAALs A and B showed a much higher frequency of univalents than Chinese Spring, and the difference was significant with MAAL A. This observation indicates that barley chromosome A (Chromosome 4) causes significantly increased asynapsis of at least one pair of wheat chromosomes

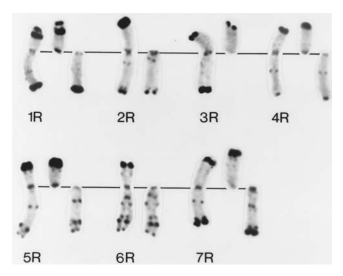


Figure 7.60 Giemsa C-banded somatic metaphase of 11 ditelosomic "Chinese Spring" wheat–"Imperial" rye addition lines showing characteristic telomeric and intercalary heterochromatic bands. (From Mukai, Y., Friebe, B., and Gill, B.S., *Jpn. J. Genet.*, 67, 71–83, 1992. With permission.)

in monosomic addition, and presumably also in disomic addition. They found no evidence of any meiotic pairing between wheat and barley chromosomes.

The lagging alien chromosomes generally misdivide in the first meiosis and may generate telocentric chromosomes. Ditelosomic addition lines have been isolated and identified in the progeny of "Chinese Spring" wheat–"Imperial" rye (Figures 7.58 and 7.60) and wheat–barley DAALs (Islam and Shepherd, 1990).

An alien chromosome generally remains as an univalent in MAALs, and the chromosomes from the recipient parent synapse regularly. However, a low frequency of trivalent formation was occasionally seen (Savitsky, 1975; Shin and Katayama, 1979; Jena and Khush, 1989). Shin and Katayama (1979) recorded 0–1 trivalents in ten of the 12 MAALs of *Oryza sativa* and *O. officinalis*, while Jena and Khush (1989) observed a trivalent in three sporocytes of MAAL 3. Furthermore, desynapsis was found in MAALs G and H of Shin and Katayama, but it was not recorded by Jena and Khush (1989).

In *Raphanobrassica* MAALs, chromosome configurations of 9II + 1I predominated, and no cell with 1III + 8II was observed (Kaneko, Matsuzawa, and Sarashima, 1987).

4. Intercrossing Disomic Alien Addition Lines

Positive identification of addition lines can also be determined by intercrossing among DAALs. Meiotic chromosome pairing in F_1 hybrids is analyzed. Alien addition lines carrying the same two chromosomes generally form a bivalent, while the occurrence of two univalents indicates the DAALs are different.

5. Biochemical Identification

Biochemical characters such as isozyme banding patterns are useful in identifying alien addition lines of wheat-rye (Miller, 1984; Benito et al., 1991a,b; Wehling, 1991), wheat-barley (Hart, Islam, and Shepherd, 1980; Islam and Shepherd, 1990), *Lycopersocon esculentum–Solanum lycopersicoides* (Chetelat, Rick, and DeVerna, 1989), *Cucurbita moschata–C. palmata* (Weeden, Graham, and Robinson, 1986), *Allium cepa–A. fistulosum* (Peffley et al., 1985), *Beta vulgaris–B. procumbens* (Lange et al., 1988), *Brassica campestris–B. oleracea* (Quiros et al., 1987), *Triticum turgidum conv.*

Barley		
Chromosome		Gene Locus
1	<i>Amy 2</i> (α-Amylase)	Est 3, 5 (Esterase)
	CM a,c (A hordeins)	Pgd 1 (Phospho gluconate dehydrogenase)
	Enp 1 (Endopeptidase)	Prx 4 (Peroxidase)
2	Est 7 (Esterase)	Isa 1 (Inhibitor subtilisin, amylase)
	Gpd 1 (Glucose-6-phosphate dehydrogenase)	Ndh 4 (NADH dehydrogenase)
	Idh 2 (Isocitrate dehydrogenase)	Prx 2 (Peroxidase)
3	Aat 3 (Aspartate aminotransferase)	<i>Est 1,2,4,10</i> (Esterase)
	CMEa [A hordeins (CM proteins)]	Mdh 2 (Malate dehydrogenase)
	Itc 1a (Inhibitor trypsin, chymotrypsin)	Tpi 1 (Triose phosphate isomerase)
4	Acp 2 (Acid phosphatase)	CM b,d [A hordeins(CM proteins)]
	Adh 1 (Alcohol dehydrogenase)	Ibf1 (lodine binding factor)
	Adh 2 (Alcohol dehydrogenase)	Ndh 1 (NADH dehydrogenase)
	β-Amy 1 (β-Amylase)	<i>Paz 1</i> (Protein Z4 in endosperm = Antigen 1a in beer)
	Cat 1 (Catalase)	Pgm 1 (Phospho gluco mutase)
5	Aco 2 (Aconitase hydratase)	Hor 3 (D hordeins)
	Gdh 1 (Glutamate dehydrogenase)	<i>Ica 1,2</i> (Inhibitor chymotrypsin, <i>Aspergillus</i> protease)
	Gpi 1 (Glucose phosphate isomerase)	Mdh 1 (Malate dehydrogenase)
	Hor 1 (C,B hordeins)	Pgd 2 (Phospho gluconate dehydrogenase)
	Hor 2 (C,B hordeins)	Adh 3 (Alcohol dehydrogenase)
6	Aat 2 (Aspartate aminotransferase)	Amp 1 (Amino peptidase)
	α -Amy 1 (α -Amylase)	Dip 1 (Dipeptidase)
	Aco 1 (Aconitate hydratase)	<i>Dip 2</i> (Dipeptidase)
	Acp 3 (Acid phosphatase)	
7	Est 9 (Esterase)	<i>Paz 2</i> (Protein Z7 in endosperm = Antigen 1b in beer)
	Sdh 1 (Shikimate dehydrogenase)	
	Gpi 2 (Glucose phosphate isomerase)	Tpi 2 (Triose Phosphate isomerase)
	lsozyme (plain type)	
	Seed protein (bold type)	

Table 7.70 Association of Isozyme Markers and Seed Protein Characters of Barley Using Wheat-Barley Alien Addition Lines

Source: From Islam, A.K.M.R. and Shepherd, K.W., In: *Biotechnology in Agriculture and Forestry, Wheat*, Y.P.S. Bajaj, Ed., Springer-Verlag, Berlin, 1990, 128–151. With permission.

V. *durum–Hordeum chilense* (Fernandez and Jouve, 1988), wheat–*H. chilense* (Miller, Reader, and Chapman, 1982b). Lange et al. (1988) assigned at least one isozyme marker to eight of the nine MAALs of *Beta vulgaris–B. procumbens*.

By using wheat–barley addition lines, Islam and Shepherd (1990) summarized the assigned genes controlling at least 58 isozymes to specific barley chromosomes or chromosome arms. Thus, wheat–barley DAALs are useful material for determining the chromosome arm location of protein and isozyme structural genes in barley (Table 7.70). Furthermore, these lines are helping to construct a restriction fragment length polymorphism (RFLP) map for barley (Heun et al., 1991).

D. Breeding Behavior of Alien Addition Lines

1. Monosomic Alien Addition Lines

In MAALs, an alien chromosome remains as an univalent at meiotic metaphase-I, and during gametogenesis, it can be eliminated or included in one of the gametes. Thus, transmission of the extra chromosome is expected to be lower than that observed in autotrisomics. The male

transmission rates of the extra chromosome in MAALs with an extra *Oryza officinalis* chromosome were not observed in six MAALs (Chromosomes 2, 3, 5, 6, 7, 8, 11). The highest male transmission rate was 14.6% in MAAL 6. In autotrisomics, male transmission was not recorded in Triplo 1, 2, 3, 6, and 7, and the highest male transmission was 27.3% in Triplo 9, a major nucleolus organizer chromosome (Table 7.71).

Blanco, Simeone, and Resta (1987) examined the transmission rate of the extra chromosome after selfing in six MAALs of *Triticum durum–Dasypyrum villosum* (Table 7.72). The transmission of the extra chromosome through the female was variable, ranging from 5 (MAAL A) to 15.7% (MAAL B). Plants with 2n = 28 + 1 telo and 2n = 29 + 1 telo were also identified. The occurrence of telocentric chromosomes indicates misdivision of a univalent that occurred at a higher rate than in *Triticum durum–Ae. umbellulata*, where monotelotrisomic plants were not found (Table 7.73).

The frequency of DAALs in the selfed progeny of MAALs in tetraploid wheat was low, because the extra alien chromosome causes more unbalance than those recorded at the 6*x* level. Therefore, transmission of the extra alien chromosome through the male is too low (Makino, 1981; Blanco, Simeone, and Resta, 1987; Dhaliwal et al., 1990), and generally, the plants are sterile. Joppa and McNeal (1972) reported six D-genome disomic addition lines of *Durum* wheat. Three DAALs (1D, 3D, 6D) were male sterile, while 4D and 5D were stable and were partially male fertile.

2. Disomic Alien Addition Lines

DAALs are expected to breed true. However, this is not always observed. Differences in meiotic cycles between recipient and donor chromosomes lead to the elimination of donor chromosomes. For example, in wheat–rye DAALs, the rye chromosomes fail to function normally in the wheat nucleus, and the tendency for asynapsis may be due to the absolute homozygosity of the rye chromosomes achieved during back crossing and selfing (O'Mara, 1940).

Miller (1984) summarized the breeding behavior of "Chinese Spring" wheat–"Imperial" rye DAALs grown over several years at Cambridge, England. An average of 86.8% plants carried expected 2n = 44 chromosomes, and the range was 86 (4R) to 95% (2R). Plants with 2n = 42, 43, and 45 chromosomes in low frequencies were also identified (Table 7.74).

Dvořák and Knott (1974) studied the stability of wheat–*Agropyron elongatum* DAALs. The majority of segregating plants contained the expected 2n = 44 chromosome number, ranging from 96.6% for DAAL VII to 85.5% for DAAL IV. The MAAL (2n = 43) plants were identified for all the lines, but the frequency was low. Plants with 2n = 43 + 1 telo and 2n = 45 were also recorded. This result indicates that DAALs are not stable and throw aneuploids in segregating populations. In contrast, the only DAAL of *N. tabacum–N. glutinosa* reported was meiotically normal and chromosomally fairly stable (Gerstel, 1945). The majority of the self-pollinated progeny of DAALs of wheat–*Ae. umbellulata* (*T. umbellulatum*) carried the expected 2n = 44 chromosomes. However, plants with 2n = 43 were also found in which the alien chromosomes failed to form a bivalent.

E. Fertility in Alien Addition Lines

Seed fertility in MAALs and DAALs is influenced by the ploidy level of the recipient parent, the genomic affinity between the receipient and donor species, and the genetic constitution of the added chromosomes.

Seed fertility in MAALs at 2x level is similar to the seed set observed in autotrisomics. MAAL 4 of *Oryza sativa–O. officinalis* was completely sterile (Jena and Khush, 1989); likewise, Triplo 4 of autotrisomics of *O. sativa* was totally sterile (Khush et al., 1984). Analogous results were found in MAALs of *Beta vulgaris–B. procumbens* (Speckmann, De Bock, and De Jong, 1985).

Seed set in MAALs of *T. durum–Ae. umbellulata* ranged from 32 (C^u5) to 84% (C^u4), while the selfed seed set in the recipient *T. durum* was 100% (Makino, 1976). Similarly, seed set was

		(2 <i>n</i> +	1) × 2 <i>n</i>				2n × ($2n \times (2n + 1)$		
MAAL	Total	2n	2 <i>n</i> + 1	% (2 <i>n</i> + 1)	% Autotrisomic ^a	Total	2 <i>n</i>	2 <i>n</i> + 1	% (2 <i>n</i> + 1)	% Autotrisomic ^a
	35	32	e	8.6	15.5					0.0
0	76	69	7	9.2	31.6	32	32	0	0.0	0.0
ო	72	67	5	6.9	17.8		I		I	0.0
4	323	267	56	17.3	43.9	144	134	10	6.9	0.5
ß	151	141	10	6.6	32.7	50	50	0	0.0	1.6
9	156	145	11	7.0	37.9	35	35	0	0.0	0.0
7	267	235	32	12.0	31.1	40	40	0	0.0	0.0
8	125	108	17	13.6	25.5	49	49	0	0.0	14.3
6	316	270	46	14.5	35.5	19	17	N	11.7	27.3
10	472	363	109	23.1	27.4	76	74	0	2.6	1.5
ŧ	375	326	49	13.1	39.7	16	16	0	0.0	2.7
12	655	479	176	26.8	37.1	89	76	13	14.6	5.6

^a Khush, G.S. et al., *Genetics*, 107, 141–163, 1984. With permission. Khush, G.S. and Singh, R.J., In: *Chromosome Engineering in Plants: Genetics, Breeding, and Evolution, Part* A, P.K. Gupta and T. Tsuchiya, Eds., 1991, 577–598. With permission.

Source: From Jena, K.K. and Khush, G. S., Genome, 32, 449–455, 1989. With permission.

	Number of			% Fre	% Frequency (2n)	(r	
Lines	Plants Examined	B	28	28+ telo	59	29 + telo	30
⊲	119		91.6	2.5	5.0	0.9	0.0
m	159		67.9	12.0	15.7	1.9	2.5
G	130		89.2	1.5	8.5	0.0	0.8
0	31		77.4	9.7	9.7	0.0	3.2
	137		76.6	11.0	11.0	0.7	0.7
	29		72.4	6.9	13.8	3.4	3.4
Total	605	(Mean)	79.8	7.3	10.6	1.0	1.3
Source: Fro	Source: From Blanco, A., Simeone, R., and Resta, P., Theor. Appl. Genet., 74, 328–333, 1987. With	Simeone, R.,	and Resta, P	, Theor. Appl.	Genet., 74,	328-333,	1987. With

Table 7.72 Breeding Behavior of the Selfed Monosomic Addition Lines of Dasvorrum villosum in

	milloum dure				
	Number of	Number of	% Frequency (2 <i>n</i>)		
Line	Progenies Examined	Plants Examined	28	29	30
C ^u 1	1	87	79.3	20.7	
C ^u 2	2	121	81.8	16.5	2.5
C ^u 3	2	99	85.7	13.1	1.0
C ^u 4	1	91	90.1	9.9	
C ^u 5	1	66	95.5	4.5	
C ^u 6	1	27	74.1	25.9	
C ^u 7	4	248	93.1	6.9	

Table 7.73 Breeding Behavior of the Selfed Monosomic Addition Lines of Aegilops umbellulata Chromosomes in Triticum durum

Source: From Makino, T., Can. J. Genet. Cytol., 18, 455–462, 1976. With permission.

Table 7.74 Breeding Behavior of the Selfed Disomic Alien Addition Lines of "Imperial" Rye Chromosomes in Chinese Spring Wheat

Chromosome	42	43	44	45
1R	0.05	0.05	0.90	
2R	—	0.05	0.95	_
3R	_	0.07	0.93	_
4R	_	0.14	0.86	_
5R	0.07	0.23	0.63	0.07
6R	0.04	0.07	0.89	_
7R	0.04	0.04	0.92	_
Mean	0.029	0.093	0.868	0.010

Source: From Miller, T.E., Can. J. Genet. Cytol., 26, 578–589, 1984. With permission.

lower in MAALs of *T. durum–Dasypyrum villosum* compared to the recipient parent (Blanco, Simeone, and Resta, 1987).

Seed set in MAALs and DAALs at the 6x level is expected to be better than at the 4x and 2x levels, because the extras cause more imbalance at 2x and 4x than at 6x. The extent of seed fertility depends upon the degree of taxonomic closeness between wheat and the alien species. MAALs and DAALs derived from crosses of *T. aestivum* and *Secale cereale*, *Agropyron*, *Haynaldia*, or *Aegilops* are relatively more fertile than the wheat–barley additions. DAAL 5 of wheat–barley was not isolated, because barley Chromosome 5 causes total male and female sterility in the presence of wheat chromosomes (Islam, Shepherd, and Sparrow, 1981). Despite complete homology and almost normal meiosis in wheat and alien species disomic additions, seed fertility varied from line to line.

V. ALIEN SUBSTITUTION LINES

A. Introduction

Exotic germplasm (the alien species), relative to cultigens, is a rich reservoir for economically valuable traits and has been proven to be a useful source for increasing genetic variability in many crops, by conventional plant breeding methods or by molecular techniques. Exotic germplasm includes all germplasm that does not have immediate usefulness without hybridization

and selection for adaptation for a given region (Hallaur and Miranda, 1981). Alien genes are introgressed into cultigens by hybridization followed by several back crossings to the recurrent parent or by genetic transformation. However, the majority of examples of alien substitution lines are available from common wheat. These lines are of two types: (1) substitution of a total genome of a cultigen into the cytoplasm of an alien species (genome substitution) and (2) substitution of a single chromosome or a pair of chromosomes into the chromosome complement of a cultigen (chromosome substitution). Sometimes, substitution of a segment(s) of alien chromosome into a cultigen occurs spontaneously or is induced by a mutagen to produce chromosomal interchange-segmental substitution (Gupta, 1971).

Shepherd and Islam (1988) listed 176 entries of alien addition lines, 84 entries of alien substitution lines, and 58 entries of translocation lines involving alien and wheat chromosomes. Such an endeavor is lacking for other crop plants. During the past decade, considerable efforts in improving tomato, potato, Alliums, Brassicas, soybean, rice, and maize by using alien germplasm. The genetic transformation has revolutionized the production of pest-resistant (Bt maize and cotton) and herbicide-tolerant (roundup ready soybean) crops.

B. Types of Alien Substitution

1. Genomic Substitution

Cytoplasms from alien species are an excellent source of cytoplasmic male sterility (CMS) (Virmani and Edwards, 1983). CMS lines are produced by alien species (female) \times cultigen (male) crosses, followed by a recurrent back-cross method to eliminate the undesirable traits donated by the alien species (Figure 7.61). Thus, a modified cultivar known as an alloplasmic line is derived.

In the tetraploid *Triticum durum* (2n = 4x = 28) and the hexaploids *T. aestivum* (2n = 6x = 42), alloplasmic lines expressing CMS are derived from related genera (Kihara, 1951; Fukasawa, 1953; Maan, 1973a, 1973b; Sasakuma and Maan, 1978). The taxonomy of genera *Triticum* and *Aegilops* is changing frequently, however, a revised and accepted classification is summarized in Table 7.75 that contains nuclear genome symbols and synonyms (B.S. Gill, personal communication).

Alloplasmic lines are not easy to produce. Examples are numerous, but only one is cited here. Smutkupt (1968) observed 61% seed set in a cross of common wheat cv. Chinese Spring (2n = 6x = 42) female × rye cv. Petkuser (2x = 14) (male), while the reciprocal cross set only 1% seed. This difference may be attributed to relatively slow growth of wheat pollen tubes in the styles of rye. In rye (female) × wheat (male) crosses, 36% of the seeds were shriveled and required embryo culture to rescue alloplasmic F₁ plants. In contrast, the reciprocal cross produced 94% fully developed seeds.

Occasionally, alloplasmic wheat lines show a reduction in seed germination, alteration in growth habit (normal to reduced growth), variable seed fertility (full fertility to total sterility), maturity changes (early to delayed), and seed variability (normal plump to shriveled) (Tsunewaki, Yoshida, and Tsuji, 1983; Virmani and Edwards, 1983).

Alloplasmic lines of common wheat and triticale exhibit multiple sporocytes (syncytes) in anthers. Tsuji, Mukai, and Tsunewaki (1983) examined 11 alloplasmic lines of wheat cv. "Rosner." The ploidy level ranged from 3x to 40x, and the modal level was 8x. However, the frequency of syncytes was not high (range of 0.2 to 4.6%). The occurrence of syncytes was mostly correlated with male sterility.

2. Chromosome Substitution and Translocation Lines

When alien chromosomes compensate completely for the absence of cultigen chromosomes in sporophytes and gametophytes, the phenomenon is known as chromosome substitution. Substitution is only possible when there is a close genetic relationship between the substituting alien

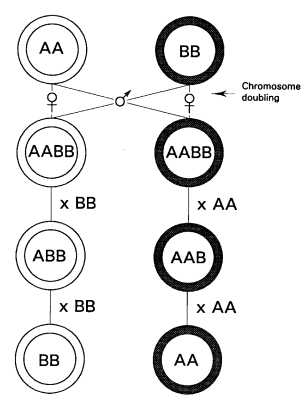


Figure 7.61 Substitution of genome complements after doubling of chromosomes through colchicine treatment. (From Kihara, H., *Cytologia*, 16, 177–193, 1951. With permission.)

chromosome and the replaced recipient (cultigen) chromosome. A considerable amount of research has been done in producing alien substitution and translocation lines in common wheat, with the objective to breed wheats carrying resistance to pathogens and pests (Friebe et al., 1996a). Sometimes, substitution of a complete chromosome donates several undesirable traits from the wild species. In such cases, segmental substitutions are produced (Sears, 1956b).

a. Production of Chromosome Substitution and Translocation Lines

Common wheat, being an allopolyploid (2n = 6x = 42; AABBDD), tolerates a considerable degree of an euploidy. It hybridizes rather easily with its wild relative genera, such as *Secale*, *Aegilops*, *Agropyron*, and *Hordeum*, and useful traits have been introgressed into tetraploid and hexaploid wheats (Sharma and Gill, 1983; Shepherd and Islam, 1988). Of all the substitution lines studied and available, wheat–rye substitutions have been the most extensively examined.

Wheat (W)–rye (R) substitutions of spontaneous origin are routinely found in nature; several widely grown European wheat cultivars contain mainly the 1RS (1BL) substitution, or the 1R chromosome is involved in translocations (1RS.1BL; 1RS.1AL) with genomes of wheat (Zeller, 1973; Zeller and Baier, 1973; Mettin, Blüthner, and Schlegel, 1973; Münzer, 1977; Zeller and Hsam, 1983). By using the Giemsa C-banding technique, Lukaszewaski (1990) examined 207 entries in six major 1989 U.S. Wheat nurseries and 30 entries in the 21st International Winter Wheat Performance nurseries. Among entries of the United States, 4.3% of lines contained 1RS.1AL, and 7.1% of lines contained 1RS.1BL translocations. The international entries predominantly carried 1RS/1BL translocations, and the frequency was higher (38.1%). Javornik et al. (1991) recorded 36 out of 59 Yugoslav wheat cultivars carrying 1BL.1RS translocations.

Species	Genome Symbol	Synonyms
Diploid	s (2 <i>n =</i> 2 <i>x =</i> 1	14)
Triticum monococcum subsp. aegilopoides	А	T. monococcum subsp. boeoticum, T. aegilopoides,
Triticum monococcum subsp. monococcum	А	T. monococcum
Triticum urartu	A	T. monococcum
Aegilops speltoides var. speltoides	S	<i>T. speltoides, Ae. speltoides</i> subsp. <i>speltoides</i>
Aegilops speltoides var. lingustica	S	T. speltoides,Ae. speltoides subsp lingustica
Aegilops bicornis var. bicornis	Sb	T. bicorne
Aegilops bicornis var. anathera	Sb	T. bircorne
Aegilops longissima	S ¹	T. longissimmum
Aegilops sheronensis	S ^{sh}	T. sharonensis (invalid)
Aegilops searsii	Ss	T. searsii (invalid)
Aegilops mutica var. mutica	Т	T. tripsacoides
Aegilops mutica var. loliacea	Т	T. tripsacoides
Aegilops tauschii	D	Ae. squarrosa, T. tauschii
Aegilops comosa var. comosa	М	T. comosum
Aegilops comosa var. subventricosa	М	Ae.comosa subsp. heldreichii
Aegilops uniaristata	Ν	T. uniaristatatum
Aegilops caudata	С	T. dichasians (invalid), Ae. markgraf
Aegilops umbellulata	U	T. umbellulatum
Tetraploi	ds (2 <i>n</i> = 4 <i>x</i> =	- 28)
Triticum turgidum subsp. carthlicum	AB	T. turgidum
Triticum turgidum subsp. dicoccoides	AB	T. turgidum
Triticum turgidum subsp. dicoccum	AB	T. turgidum
Friticum turgidum subsp. durum	AB	T. turgidum
Triticum timopheevii subsp. timopheevii	AG	T. timopheevii
Aegilops crassa	DM	T. crassaum
Aegilops cylindrica	CD	T. cylindricum
Aegilops geniculata	UM	T. ovatum
Aegilops neglecta	UM	T. neglecta
Aegilops biuncialis	UM	T. macrochaetum
Aegilops columnaris	UM	T. columnare (invalid)
Aegilops triuncialis var. triuncialis	UC	T. triunciale
Aegilops ventricosa	DN	T. ventricosum
Hexaploi	ds (2 <i>n =</i> 6 <i>x =</i>	- 42)
Triticum aestivum subsp. aestivum	ABD	
Triticum aestivum subsp. compactum	ABD	
Triticum aestivum subsp. compactum	ABD	
Triticum aestivum subsp. macha	ABD	
<i>Triticum aestivum</i> subsp. <i>spelta</i>	ABD	
Triticum aestivum subsp. sphaerococcum	ABD	
Triticum zhukovskyi	AAG	
Aegilops neglecta	UMN	T. neglectum
Aegilops crassa	DDM	T. crassum
Aegilops vavilovii	DMS	T. syriacum
Aegilops juvenalis	DMU	T. juvenale

 Table 7.75 Taxonomy of the Genus Triticum and Aegilops, 2n Chromosome, Genomes, and Their Synonyms^a

Source: From Kimber, G. and Tsunewaki, K., In: *Proc. 7th Int. Wheat Genet. Symp.*, T.E. Miller and R.M.D. Koebner, Eds., Bath Press, Avon, U.K., 1988, 1209–1211. With permission.

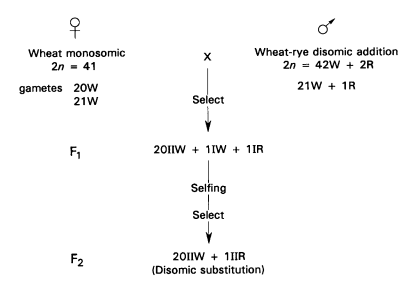


Figure 7.62 Production of wheat-rye substitution line.

Substitution lines are produced experimentally. A prerequisite to isolate substitution lines is to have available monosomic alien addition lines (MAALs) and disomic alien addition lines (DAALs). In producing wheat-rye substitution lines, the DAALs (2n = 42W + 2R) are used as a pollen parent and wheat monosomics (2n = 41W) as a female parent. The F₁ plants containing 2n = 20IIW +1IW + 1IR are selfed or pollinated onto DAALs. It is expected that the F_1 plants will produce four kinds of female and male spores: 20W + OR, 20W + 1R, 21W + OR, 21W + 1R in the ratio of 9:3:3:1 assuming both rye chromosomes are excluded in 75% of gametes (Sears, 1944). Because nullisomic (20W + OR) male spores will not function, and the missing wheat chromosome is fully compensated by a rye chromosome, the functional male spores will be in the ratio of 0:3:3:1. However, in a noncompensating combination, male spores will segregate in a ratio of 0:0:3:1. In the F_1 , plants with 2n = 41W + 2R (20IIW + 1IW + 1IIR) are selected cytologically, and after selfing, these plants' disomic substitution lines (2n = 40W + 2R) are selected, because 1W univalent is usually eliminated during gametogenesis (Figure 7.62). This technique can also be used to produce ditelocentric addition lines. The telocentric chromosomes are easy to identify cytologically and can be used for locating genes on a particular arm of a chromosome. The same procedure was used to produce wheat-Agropyron substitution lines (Dvořák 1980).

Merker (1979) felt that the above procedure for producing wheat-rye substitution lines requires a considerable amount of cytological work. His procedure is as follows: Pollinate monosomic wheats (2n = 41W) by octoploid triticales (2n = 8x = 56; AABBDDRR). The F₁ plants are expected to segregate two cytotypes: (1) 2n = 5x = 49 (42W + 7R) and (2) 2n = 5x - 1 = 48 (41W + 7R). Substitution lines are expected after selfing 2n = 48 chromosome plants, where a missing wheat chromosome (in question) has been substituted by a rye chromosome that possesses the best compensating ability. The deficiency of this method is that another rye chromosome may substitute for the questionable wheat chromosome. In contrast, if substitution lines are produced from DAALs, a rye chromosome is forced to substitute for a specific wheat chromosome, because the rye chromosome has no preference. However, both methods require cytological work.

Friebe and Larter (1988) produced a complete set of isogenic wheat–rye D-genome substitutions by crossing an inbred line of spring rye cv. "Prolific" to a tetraploid wheat. The A- and B-genomes were extracted from the hexaploid wheat cv. "Thatcher" and selection for wheat–rye substitution lines was carried out in BCF_3 to BCF_6 families.

Joppa and Williams (1977) isolated 14 possible substitution monosomics by crossing the appropriate Chinese Spring nullisomic-tetrasomic with *T. turgidum durum* "Longdon" (2n = 28,

AABB). The F₁ plants were allowed to self-pollinate. Selection for disomic substitution for a Dgenome chromosome pair (homoeologous A- or B-genome chromosome pair) was conducted in F_2 and F_3 generations. In another study, Joppa and Maan (1982) isolated a substitution line in the F_2 population from a cross of durum wheat selection 56-1 and T. boeticum. One T. boeticum chromosome compensated for the loss of a durum chromosome 4B. Present day durum and T. aestivum wheats are structurally different from the originally synthesized tetraploid and hexaploid wheats because of chromosomal interchanges among A B D genomes of wheats (Okamoto and Sears, 1962). For example, T. urartu is considered to be an A-genome donor of the AA genome of cultivated wheat (Chapman, Miller, and Riley, 1976). However, it has been established that only six A genome chromosome pairs are present in tetraploid emmer wheat and hexaploid bread wheat. It is now known that Chromosome 4A is not from the A genome donor species but from an unknown species and appears to have multiple structural rearrangement in comparison to 4A chromosome of T. monococcum and T. urartu (Naranjo, 1990; Zeller, Cermeño, and Miller, 1991). Based on isozyme and molecular homoeoloci, Liu et al. (1992) furnished evidence that the present 4AL chromosome constitutes chromosome segments of 7BS and 5AL, and telomere of 7BL came from 5AL. These observations suggest that wheat genomes underwent considerable structural changes during the course of evolution of present day wheats, and still these changes are occurring.

b. Wheat–Rye Substitutions and Homoeologous Relationships

Homoeologous relationships among the three genomes (ABD) of common wheat cv. "Chinese Spring" were established and classified into seven homoeologous groups by Sears (1966b), based on the knowledge of nullisomic tetrasomic series, where a pair of extra chromosomes of one genome could successfully compensate for the loss of two chromosomes from the other two genomes. The three genomes of common wheat are closely related and originated from a common progenitor.

Several nomenclature systems have been used to designate rye chromosome (Table 7.76; Sybenga, 1983). However, to avoid confusion and to adopt an universal nomenclature, rye chromosomes are being assigned to the respective homoeologous wheat chromosomes based on morphological and cytological markers such as characteristic C-bands (Figure 7.63), meiotic chromosome pairing, telocenterics, interchanges, substitution and compensation ability, and genetic tests (O'Mara, 1947; Sears, 1968; Gupta, 1971; Zeller and Hsam, 1983; Miller, 1984; Naranjo and Fernández-Rueda, 1991). The Giemsa C-karyogram of "Imperial" rye in "Chinese Spring" wheat is considered standard. The results are briefly summarized below.

i. Chromosome 1R — It is a nucleolus organizing chromosome, substitutes completely for wheat chromosomes 1A, 1B, and 1D, and carries genes for resistance to pests and diseases. The 1A (1R) and 1B (1R) substitutions and 1B.1R translocations are common in many wheat cultivars (Table 7.76).

Table 7.76 The Equivalence of the Original Nomenclature to the Homoeological Nomenclature of the Lines of the Major Wheat–Rye Addition Series

	Homo	eologous	Groups a	nd Rye C	hromosor	nes Desig	gnation
	1, 1R	2, 2R	3, 3R	4, 4R ^a	5, 5R	6, 6R	7, 7 R ª
"Holdfast""King II"	V	111		IV	I	П	VII
"Kharkov"-"Dakold"	VII	11	_	V	VI	IV	111
"Chinese Spring"-"Imperial"	E	В	G	С	А	F	D
"Chinese Spring"-"King II"	1R	2R	3R	4R	5R	6R	7R
"Chinese Spring"-S. montanum	1R	2R	_	4R	5R	6R	_
"Fec 28"-"Petkus 10"	F	D	С	А	Е	—	В

^a Partial homoeology to group 4 and group 7.

Source: From Miller, T.E., Can. J. Genet. Cytol., 26, 578-589, 1984. With permission.

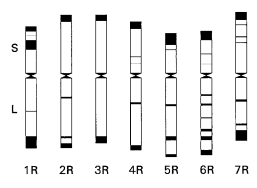


Figure 7.63 An idiogram of Imperial rye in Chinese Spring wheat addition line, represents karyogram from Figure 7.60. (Redrawn from Mukai, Y., Friebe, B., and Gill, B.S., *Jpn. J. Genet.*, 67, 71–83, 1992.)

ii. Chromosome 2R — It is the longest chromosome in the rye chromosome complement, and both arms have prominent heterochromatic bands. Chromosome 2R substitutes reasonably well for wheat chromosomes 2B and 2D. Interchanges involving 2RL.2AS, 2BL.2RS, and 2BS.2RL (Figure 7.64) are rather common. However, interchange 4A.2R and location of homoeoloci *Gli*-2 on 2RS, 6AS, 6BS, and 6DS suggest small structural differences in 2R. Based on the location of gene *Gli*-2 on chromosome 6R^m (m = *S. montanum*), Shewry, Parmar, and Miller (1985) postulated the presence of an interchange between 2R and 6R^m, probably also present in wheat. Chromosome 2R contains genes for resistance to powdery mildew, Hesian fly and leaf rust.

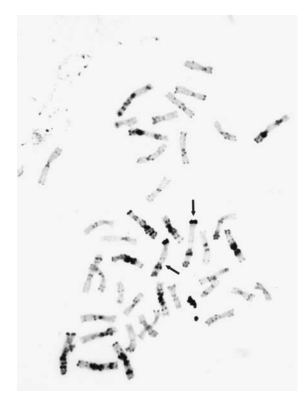


Figure 7.64 Giemsa C-banded mitotic metaphase of Hessian fly-resistant wheat–rye translocation line, 2BS.2RL (arrows). (From Friebe, B. et al., *Theor. Appl. Genet.*, 79, 385–389, 1990a. With permission.)

iii. Chromosome 3R — It contains a terminal heterochromatic prominent band on both arms, substitutes for wheat chromosomes 3A and 3D. The 3RS (short arm) shows homoeology with 3AS, 3BS, and 3DS, but 3RL (long arm) contains little homoeology with 3AL, 3BL, and 3DL. Furthermore, the distal region of 3RL has homoeology with 6AL, 6BL, and 6DL. Chromosome 3R contains genes for resistance to wheat mildew.

iv. Chromosome 4R — Chromosome 4R carries a prominant telomeric heterochromatic band on the short arm and a medium size telomeric band on the long arm (Figure 7.63) and substitutes for wheat chromosomes 4A, 4B, and 7B (Miller, 1984). Chromosome pairing results suggest homoeology of 4RS to 4BS and 4DS and no affinity with 4A, but to some extent, it is homoeologous to 4AL; chromosome 4RL contains some homoeology to 6AS, 6BS, and 6DS and also wheat group 7 chromosomes, which is in agreement with the detection of the

4RL.7RS translocation (Koller and Zeller, 1976). Thus, Chromosome 4R includes chromosome segments of homoeologous groups 4 and 7. According to Liu et al. (1992), present 4R is composed of 4RL and 7RS.

v. Chromosome 5R — This chromosome contains a large terminal heterochromatic band on the short arm. Genetically, Chromosome 5R is identified by the "hairy neck" gene HP (controls peduncle pubescence) and substitutes for wheat Chromosomes 5A, 5B, and 5D. As far as homoeologous relations of arms are concerned, 5RS is homoeologous to 5AS, 5BS, and 5DS (chromosome pairing and genetic data). However, substitution 5R (4A) and interchanges 4A/5R suggest that 5R has some homoeology with wheat groups 4.

vi. Chromosome 6R — It possesses a prominent telomeric heterochromatic band on the short arm and substitutes for wheat chromosomes 6A, 6B, and 6D. Chromosome 6RL shows homoeology with 6AL, 6BL, and 6DL and some homoeology to 3AL, 3BL, and 3DL (genetic data). This relation may be due to a reciprocal translocation involving 3RL and 6RL in rye relative to wheat. Pairing between 6RL and the long arm of wheat group 7 chromosomes suggests that an additional interchange involving 6RL and 7RL could have occurred during the evolution of rye (Naranjo and Fernández-Rueda, 1991).

vii. Chromosome 7R — This chromosome carries telomeric heterochromatic bands on both arms and shows partial homoeology with wheat groups 4 and 7 chromosomes. The 7RS does not pair with 7AS and 7DS but pairs with 5BL, 5DL, and 7BS (the terminal segment of 7S is homoeologous to the long arm of group 5 chromosomes and the 7RS.5RL translocation in rye relative to wheat). The 7RL shows homoeology to 7AL, 7BL, and 7DL as demonstrated by genetic and RFLP data and shows affinity with 2AS, 2BS, and 2DS. Thus, Chromosome 7 is a modified chromosome containing several rearrangements with respect to wheat chromosomes. Liu et al. (1992) recorded that 7R is modified due to interchanges; 7RS constitutes segments of 7RS (the region proximal to kinetochore), 4RL, and telomere from 5RL.

Incomplete homoeology of 4R, 5R, and 7R to the corresponding homoeologous D-genome chromosomes is due to the occurrence of reciprocal translocations in these chromosomes (Taketa et al., 1991).

In addition to wheat–rye addition and substitutions, attempts are being made to produce addition and substitution lines from more distantly related wild species of wheats. Wheat-Agropyron substitution lines have been produced in order to transfer disease resistance genes from Agropyron to common wheat (Knott, 1964; Knott, Dvořák, and Nanda, 1977; Dvořák, 1980; Wienhues, 1966, 1971,1973) and emmer wheat (Tsujimoto, Shinotani, and Ono, 1984). The scientific botanical name of *A. elongatum* has a confused history. This particular species is also named, *Elytrigia elongata*, *Thinopyrum elongatum*, and *Lophopyrum elongatum*.

Examples of wheat-Aegilops substitutions are a few, and no systematic effort has been made to produce all possible substitution lines. However, two classical examples of gene introgression from Aegilops species to common wheat need to be mentioned. Sears (1956b) transferred a small segment of Ae. umbellulata chromosome conferring resistance to leaf rust to wheat cv. "Chinese Spring." The translocation was a substitution with 21II recovered following irradiation. Riley, Chapman, and Johnson (1968) transferred a gene for yellow rust resistance from Ae. comosa to wheat cv. "Chinese Spring" through genetically induced homoeologous recombination.

Miller, Hutchinson, and Chapman (1982a) found an *Ae. sharonensis* chromosome that substitutes for homoeologous group 4 (ABD) of Chinese Spring.

C. Practical Application of Substitution Lines

Knowledge of substitution lines has enhanced our understanding of homoeologous relationships among three wheat genomes and allied genera such as *Secale*, *Agropyron*, and *Aegilops*. Useful traits for economic importance have been introgressed into several major crops (Table 7.77).

Iable 1.11 Alleli Cili Ollos			ITOIII I IIIEII AIIIEU Species anu	dellera
Recipient	Donor	Procedure	Gene Transferred	Authors
Beta vulgaris	Beta patellaris	Substitution	Nematode resistance	Salentijn et al., 1994
Beta vulgaris	Beta procumbens	Substitution	Nematode resistance	Salentijn et al., 1994
Brassica napus	Brassica nigra	Substitution	Blackleg	Chevre et al., 1996
Brassica napus	Sinapsis arvensis	Substitution	Blackleg	Snowdon et al., 2000
Cicer arietinum	Cicer reticulatum	Homoeologous pairing	Cyst nematode	Di Vito et al., 1996
Glycine max	Glycine tomentella	Homoeologous pairing	Soybean cyst nematode	Riggs et al., 1998
Hordeum vulgare	Hordeum bulbosum	Irradiation induced	Powdery mildew	Pickering et al., 1995
Lycopersicon esculentum	Lycopersicon chilense	Homoeologous pairing	Cucumber mosaic virus	Stamova and Chetelat, 2000
Lycopersicon esculentum	Lycopersicon pennellii	Homoeologous pairing	Total soluble solids	Eshed and Zamir, 1994
Lycopersicon esculentum	Lycopersicon pennellii	Homoeologous pairing	Potato aphids	Hartman and St. Clair, 1999b
Lycopersicon esculentum	Lycopersicon pennellii	Homoeologous pairing	Beat army worm	Hartman and St. Clair, 1999a
Lycopersicon esculentum	Lycopersicon peruvianum	Homoeologous pairing	Root-knot nematode	Doganlar, Frary, and Tanksley, 1997
Lycopersicon esculentum	Solanum lycopersicoides	Homoeologous pairing	Male fertility	Chetelet et al., 1997
Oryza sativa	Oryza australeinsis	Homoeologous pairing	Bacterial blight	Multani et al., 1994
Oryza sativa	Oryza australeinsis	Homoeologous pairing	Brown plant hopper	Multani et al., 1994
Oryza sativa	Oryza brachyantha	Homoeologous pairing	Bacterial blight	Brar and Khush, 1997
Oryza sativa	Oryza latifolia	Homoeologous pairing	Bacterial blight	Multani et al., 1994
Oryza sativa	Oryza latifolia	Homoeologous pairing	Brown plant hopper	Multani et al., 1994
Oryza sativa	Oryza latifolia	Homoeologous pairing	White-backed plant hopper	Multani et al., 1994
Oryza sativa	Oryza longistaminata	Homoeologous pairing	Bacterial blight	Brar and Khush, 1997
Oryza sativa	Oryza minuta	Homoeologous pairing	Bacterial blight	Amante-Bordeos et al., 1997
Oryza sativa	Oryza minuta	Homoeologous pairing	Rice blast	Amante-Bordeos et al., 1997
Oryza sativa	Oryza minuta	Homoeologous pairing	Brown plant hopper	Amante-Bordeos et al., 1997
Oryza sativa	Oryza nivara	Homoeologous pairing	Grassy stunt	Brar and Khush, 1997
Oryza sativa	Oryza officinalis	Homoeologous pairing	Bacterial blight	Brar and Khush, 1997
Oryza sativa	Oryza officinalis	Homoeologous pairing	Brown plant hopper	Brar and Khush, 1997
Oryza sativa	Oryza officinalis	Homoeologous pairing	White-backed plant hopper	Brar and Khush, 1997
Oryza sativa	Oryza perennis	Homoeologous pairing	Cytoplasmic male sterility	Dalmacio et al., 1995
Triticum aestivum	Secale cereale	1B (1R) Substituions and	Powdery mildew	Mettin, Blüthner, and Schlegel, 1973;
		translocations	resistance	Zeller, 1973
Triticum aestivum	Secale cereale	1B (1R) Substituions and	Stem and leaf rust	Mettin, Blüthner, and Schlegel, 1973;

Table 7.77 Alien Chromosome Substitutions and Transfer of Useful Traits to Cultigen from Their Allied Species and Genera

Zeller, 1973

Stripe rust resistance

1B (1R) Substituions and translocations

translocations

Secale cereale Secale cereale

Triticum aestivum Triticum aestivum

Stem and leaf rust resistance

Mettin, Blüthner, and Schlegel, 1973; Zeller, 1973

Table 7.77 (continued)	Alien Chromosome Substitut	Alien Chromosome Substitutions and Transfer of Useful Traits to Cultigen from Their Allied Species and Genera	o Cultigen from Their Allied	Species and Genera
Recipient	Donor	Procedure	Gene Transferred	Authors
Triticum aestivum	Secale cereale	1B (1R) Substituions and translocations	Powdery mildew	Zeller and Hsam, 1983
Triticum aestivum Triticum aestivum	Secale cereale Secale cereale	1A.1R translocation 1A.1R translocation	Green bug resistance Powdery mildew resistance	Zeller and Fuchs, 1983 Zeller and Fuchs, 1983
Triticum aestivum	Secale cereale	1A.1R translocation	Stem and leaf rust resistance	Zeller and Fuchs, 1983
Triticum aestivum	Secale cereale	4A.2R translocation ^a	Powdery mildew resistance	Driscoll and Jensen, 1964
Triticum aestivum	Secale cereale	4A.2R translocation ^a	Leaf rust resistance	Driscoll and Jensen, 1964
Triticum aestivum Triticum aestivum	Secale cereale Secale cereale	3A.3R translocation 3A.3R translocation	Stem rust resistance Powdery mildew	Stewart et al., 1968 Lind, 1982
Triticum aestivum	Sanala naraala	6BS 6BL transformation	Wheat mildew resistance	l ind 1082
Triticum aestivum	Secale cereale	2BS.2RL translocation	Hessian fly resistance	Eriebe et al. 1990a
Triticum aestivum	Secale cereale	2BS.2RL translocation	Stripe rust resistance	Friebe et al., 1990a
Triticum aestivum	Secale cereale	1BL.1RS translocations	Powdery mildew resistance	Friebe, Heun, and Bushuk, 1989
Triticum aestivum	Secale cereale	Irradiation-induced translocations	Hessian fly resistance	Friebe et al., 1991a
Triticum aestivum	Secale cereale	4BL.5RL translocation	High copper tolerance	Schlegel, Werner, and Hülgenhof, 1991
Triticum aestivum	Secale cereale	2A.2R translocation	Green bug resistance	Friebe et al., 1995b
Triticum aestivum	Triticum monococcum	Bridge cross	Stem rust resistance	McIntosh et al., 1984
Triticum aestivum	Agropyron elongatum	Irradiation-induced translocation	Leaf rust resistance	Wienhues, 1966; Sharma and Knott, 1966
Triticum aestivum	Agropyron elongatum	Irradiation-induced translocation	Leaf and stem resistance	Dvořák and Knott, 1977
Triticum aestivum	Agropyron elongatum	Homoeologous pairing	Leaf rust resistance	Sears, 1973
Triticum aestivum	Agropyron elongatum	Substitution	Wheat streak mosaic virus	Larson and Atkinson, 1973
Triticum aestivum	Agropyron elongatum	Substitution	Salt tolerance	Omielan, Epstein, and Dvořák, 1991
Triticum aestivum	Agropyron intermedium	Translocations	Leaf rust resistance	Wienhues, 1973
Triticum aestivum	Agropyron intermedium	Irradiation-induced translocation	Leaf rust resistance	Friebe et al., 1993
Triticum aestivum	Aegilops comosa	Homoeologous pairing	Yellow rust resistance	Riley, Chapman, and Macer, 1966a; Chapman and Johnson, 1968
Triticum aestivum	Aegilops longissimum	Homoeologous pairing	Powdery mildew resistance	Ceoloni et al., 1988
Triticum aestivum	Aegilops sharonensis	Irradiation induced	Male fertility	King et al., 1991
Triticum aestivum	Aegilops speltoides	Homoeologous pairing	Leaf rust resistance	Dvořák, 1977; Dvořák and Knott, 1980
Triticum aestivum	Aegilops umbellulata	Irradiation-induced translocation	Leaf rust resistance	Sears, 1956b

Spetsov et al., 1997	Delibes et al., 1993 Miller et al., 1997 Marais et al. 1988 Chen et al., 1998 Larkin et al., 1995 Larkin et al., 1995 Sharma et al., 1995	Dyck, 1992 Shi, Leath, and Murphy, 1998 Dyck, 1994 Liu et al., 1988	Friebe et al., 1991b Friebe et al., 1991b Kerber and Dyck, 1990 Aung and Thomas, 1978 Sharma and Forsberg, 1977 Hussain et al., 1997	
Powdery mildew resistance	Cereal cyst nematode Aluminum tolerance Leaf rust resistance Wheat streak mosaic virus Barley yellow dwarf Rust resistance Barlev vellow dwarf	Stem rust resistance Powdery mildew resistance Leaf rust resistance Powdery mildew resistance	Wheat streak mosaic virus Green bug resistance Leaf rust resistance Mildew resistance Crown rust resistance Clover cyst nematode	latum are one species.
Homoeologous pairing	Homoeologous pairing Homoeologous pairing Translocation Translocation Addition Addition Substitution	Homoeologous pairing Homoeologous pairing Homoeologous pairing substitution	Substitutions and translocations Substitutions and translocations Homoeologous pairing Irradiation-induced translocation Irradiation induced translocations Polyploidization	ation). elongata, Thinopyrum elongate, and Lophopyrum elongatum are one species.
Aegilops variabilis	Aegilops ventricosa Aegilops uniaristata Thinopyrum distichum Thinopyrum intermedium Thinopyrum intermedium Thinopyrum intermedium	Trifticum araraticum Trifticum monococcum Trifticum turgidum Haynaldia vilosa	Agropyron intermedium × Tritticum monococcum Aegilops speltoides × Tritticum monococcum Avena barbata Avena abyssinica × Avena strigosa Trifolium nigrescens	ial communication). um, Elytrigia elongata, Thinopyr
Triticum aestivum	Triticum aestivum Triticum aestivum Triticum aestivum Triticum aestivum Triticum aestivum Triticum aestivum	Triticum aestivum Triticum aestivum Triticum aestivum Triticum aestivum	Triticum aestivum Triticum aestivum Avena sativa Avena sativa Trifolium repens	^a 4B.5R (B. Friebe, personal communication). Note: Agropyron elongatum, Elytrigia elonge

Transfer of useful genetic traits from alien species to cultigens is not limited to only wheat, but it is being explored for several diploid and polyploid species (Table 7.77). Jena and Khush (1990) transferred genes for resistance to insects and diseases from *Oryza officinalis* to *O. sativa*. Singh, Kollipara, and Hymowitz (1990, 1993) produced for the first time intersubgeneric hybrids with fertile plants from a cross of soybean, *Glycine max*, and a wild perennial species *G. tomentella*. It took a decade to isolate monosomic alien addition lines (Singh et al., 1998), and a gene for resistance to soybean cyst nematode has been introgressed from *Glycine tomentella* to the soybean (Riggs et al., 1998).

"The wider crosses not only take patience but also determination, commitment, and, frequently, a good deal of skill and ingenuity. Not surprisingly, the easy materials are more likely to be used" (Harlan, 1976). It is an extremely valuable and encouraging statement for cytogeneticists involved in wide hybridization.

D. Gene Pool Concept

A prerequisite for the exploitation of wild species to improve cultivars is to have complete comprehension and understanding of the taxonomic and evolutionary relationships between cultigen and its wild allied species (Hawkes, 1977). Harlan and de Wet (1971) proposed the concept of three gene pools based on hybridization among species. These are primary (GP-1), secondary (GP-2), and tertiary (GP-3) (Figure 7.65).

1. Primary Gene Pool (GP-1)

The primary gene pool consists of biological species, and crossing within this gene pool is easy. Hybrids are vigorous, exhibit normal meiotic chromosome pairing, and possess total fertility. The gene segregation is normal, and gene exchange is generally easy. GP-1 is further subdivided into Subspecies A, which includes cultivated races, and Subspecies B, which includes spontaneous

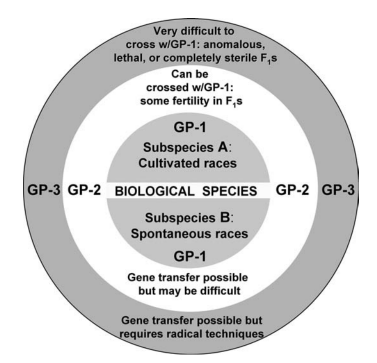


Figure 7.65 Gene pool concept in plants established based on hybridization. (Redrawn from Harlan, J.R. and de Wet, J.M.J., *Taxon*, 20, 509–517, 1971.)

races. Soybean cultivars and land races and their wild annual progenitor, *Glycine soja*, are included in GP-1. The GP-1 in allotetraploid cultivated cotton (*Gossypium hirsutum*, *Gossypium barbadense*) includes the wild tetraploid species *Gossypium tomentosum*, *Gossypium darwinii*, and *Gossypium mustelinum*. Interspecific hybrids are fertile. Introgression between *Gossypium hirsutum* and *Gossypium barbadense* occurs (Percival, Wendel, and Stewart, 1999).

A race is a biological unit with some integrity, originated in some geographical region at some time in the history of the crop, it is not clearly separable as a species but has a distinct cohesion of morphology, geographical distribution, ecological adaptation, and frequently, of breeding behavior (Harlan and de Wet, 1971).

Zeven (1998) defined race as: "an autochthonous landrace is a variety with a high capacity to tolerate biotic and abiotic stress, resulting in high yield stability and an intermediate yield level under a low input agricultural system."

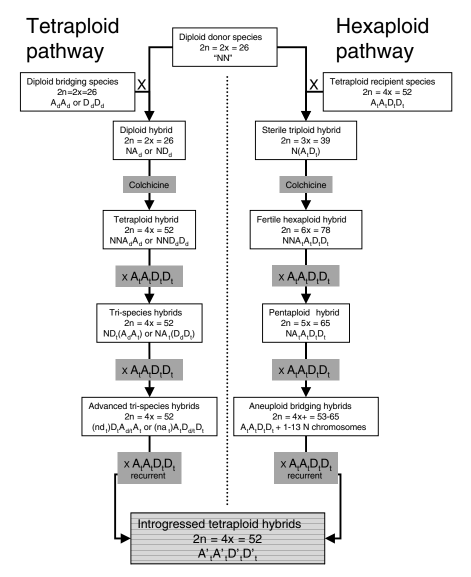


Figure 7.66 Diagrammatic flow pathways for transferring genes from GP-3 to cultivated cotton. (Courtesy of C. L. Brubaker.)

2. Secondary Gene Pool (GP-2)

The secondary gene pool includes all species that can be crossed with GP-1 with at least some fertility in F_1 . The gene transfer is possible but may be difficult. Based on this criterion, soybean does not have a GP-2. Also, barley does not have a GP-2 and has a small GP-3. The secondary gene pool in wheat is large and includes all species of *Aegilops, Secale*, and *Haynaldia*, plus at least *Agropyron elongatum*, *A. intermedium*, and *A. trichophorum*. Rice has substantial GP-2 and small GP-3. The secondary gene pool in cotton is large and includes A, D, B, and F genome diploid species.

3. Tertiary Gene Pool (GP-3)

Tertiary gene pool is the extreme outer limit of potential genetic resource. Hybrids between GP-1 and GP-3 are difficult to produce, require *in vitro* technique to rescue F_1 plant, and F_1 plants are anomalous, lethal, or completely sterile. Based on this definition, GP-3 includes all the wild perennial species of the subgenus *Glycine* for the soybean. GP-3 for wheat includes *Agropyron* and several *Elymous* species.

In cotton, GP-3 includes species with C, G, and K (Australian) and E (African-Arabian) genome species. The GP-3 species may or may not be hybridized easily with upland cotton, and introgression of traits presents difficulties that are not easily resolved (Percival, Wendel, and Stewart, 1999).

Brubaker et al. (1999) modified the proposed two procedures for the utilization of cotton GP-3 germplasm to broaden the germplasm base of cultivated tetraploid (2n = 4x = 52) cotton. A diagrammatic flow scheme, modified from Brubaker et al. (1999), is shown in Figure 7.66. One is a hexaploid pathway, and the other is a tetraploid pathway. Intergeneric hybrids were totally sterile in both types of crosses. In hexaploid strategy, hybridize the diploid (2n = 2x = 26) wild species carrying economically useful traits to a tetraploid cotton. Produce hexaploid (2n = 6x = 78) by treating F₁ with the colchicine. Back cross to recurrent parent, and isolate modified tetraploid cotton after BC₂ to BC₄ generations. They were able to isolate derived fertile lines by hexaploid pathway. In tetraploid pathway, hybridize wild diploid (2n = 2x = 26) species to an A or D genome diploid bridging species. Produce tetraploid (2n = 4x = 52) by treating F₁ with the colchicine. Back cross to recurrent parent, and isolate two populations of modified tetraploid cotton or intercross two populations. However, they have not been able to isolate derived fertile lines from the tetraploid pathway. Thus, difficult materials are not easily approachable and require considerable efforts.

CHAPTER 8

Genome Analysis

I. INTRODUCTION

Understanding the genomic relationships among species is important to systematists, evolutionary biologists, cytogeneticists, molecular biologists, and plant breeders. The taxonomic nomenclature of species and their evolutionary relationships can be refined by cytogenetic evidences such as chromosome morphology (karyotypes), crossability, hybrid viability, meiotic chromosome pairing, and molecular (isozymes, RFLP, RAPD, chloroplast and mitochondrial DNA, and other methods) approaches. Phylogenetic relationships among species can be understood more precisely by a multidisciplinary approach rather than through reliance on a single technique (Jauhar, 1990, 1996; Jauhar et al., 1991; Singh, Kollipara, and Hymowitz, 1992a). For example, van der Maesen (1986) combined the genus *Atylosia* with the genus *Cajanus* based on morphological, cytological, and chemotaxonomic data. However, the chromosome pairing and molecular methods to establish genomic relationships among species will be the main theme of this chapter.

II. CLASSICAL TAXONOMY AND GENOME DESIGNATIONS

A taxonomic nomenclature of taxa based on morphological traits is the traditional and valuable starting point for cytogeneticists as well as for molecular geneticists. The genus *Glycine* Willd. has been selected as an example to elucidate the importance of taxonomy.

Based on classical taxonomy, the genus *Glycine* has been divided into two subgenera, *Glycine* and *Soja* (Moench) F.J. Herm. Subgenus *Glycine* currently consists of 22 wild perennial species, and all the diploid (2n = 40) species are indigenous to Australia and associated areas (Table 8.1). *Glycine tabacina* (Labill.) Benth., with 2n = 40, 80 chromosomes, has been found in Australia (New South Wales, Queensland, Australian Capital Territory), Taiwan, South Pacific Islands (New Caledonia, Fiji, Tonga, Vanuatu), and West Central Pacific Islands (Mariana, Ryukyu). All accessions of *G. tabacina* collected outside of Australia are tetraploid, and within Australia, tetraploid *G. tabacina*'s predominate with an occasional diploid. *Glycine tomentella* Hayata has four cytotypes (2n = 38, 40, 78, 80). Aneuploids (2n = 38, 78) are found in New South Wales and in the adjoining regions of Queensland. Diploid (2n = 40), and tetraploid (2n = 80) cytotypes have been collected from various islands off Queensland, Australia, and Papua New Guinea. *Glycine tomentella* accessions from the Philippines, Indonesia, and Taiwan are tetraploid. *Glycine dolichocarpa* contains 2n = 80 chromosomes and is distributed in Taiwan. Morphologically, it looks similar to 80-chromosome *G. tomentella*. *Glycine hirticaulis* has extremely narrow distribution in Australia and contains 2n = 40 and 80 diploids and tetraploids, respectively (Tindale and Craven, 1988).

		Genome S	ymbol	
Species	2 <i>n</i>	N ^a	Cp	Distribution
	Subgen	us <i>Glycine</i>		
G. albicans Tind. and Craven	40	I	А	Australia
G. aphyonota B. Pfeil	40	?	?	Australia
<i>G. arenaria</i> Tind.	40	Н	А	Australia
<i>G. argyrea</i> Tind.	40	A ₂	Α	Australia
<i>G. canescens</i> F. J. Herm.	40	А	Α	Australia
<i>G. clandestina</i> Wendl.	40	A ₁	Α	Australia
<i>G. curvata</i> Tind.	40	C ₁	С	Australia
<i>G. cyrtoloba</i> Tind.	40	C	С	Australia
G. falcata Benth.	40	F	А	Australia
G. hirticaulis Tind. and Craven	40	H,	А	Australia
G. hirticaulis Tind. and Craven	80	?	?	Australia
G. lactovirens Tind. and Craven	40	I,	А	Australia
G. latifolia (Benth.) Newell and Hymowitz	40	B ₁	В	Australia
G. latrobeana (Meissn.) Benth.	40	A	А	Australia
G. microphylla (Benth.) Tind.	40	в	В	Australia
G. peratosa B. E. Pfeil & Tind.	40	?	?	Australia
<i>G. pindanica</i> Tind. & Craven	40	H,	А	Australia
<i>G. pullenii</i> B. Pfeil, Tind. & Craven	40	?	?	Australia
<i>G. rubiginosa</i> Tind. & B. E. Pfeil	40	?	?	Australia
G. stenophita B. Pfeil and Tind.	40	B ₃	B?	Australia
<i>G. dolichocarpa</i> Tateishi & Ohashi	80	?	?	South East Coast of Taiwa
<i>G. tabacina</i> (Labill.) Benth.	40	B ₂	В	Australia
	80	(complex)	?	Australia, West Central and South Pacific Islands
G. tomentella Hayata	38	E	А	Australia, Papua New Guinea
	40	D, D ₁ , D ₂ , D ₃	А	Australia
	78	(complex)	?	Australia, Papua New Guinea
	80	(complex)	?	Australia, Papua New Guinea, Philippines, Taiwa
Subgen	us <i>Soja</i> (I	Moench) F. J. H	lerm.	
<i>G. soja</i> Sieb. and Zucc.	40	G	G	China, Russia, Taiwan, Japan, Korea
<i>G. max</i> (L.) Merr.	40	G	G	Cultigen

Table 8.1 Taxonomy of the Genus Glycine Willd.ª

^a N = nuclear; Kollipara, K.P., Singh, R.J., and Hymowitz, T., Genome, 40, 57–68, 1997. With permission.

^b C = chloroplast; Doyle, J.J. et al., *Syst. Bot.*, 25, 437–448, 2000. With permission.

Subgenus *Soja* contains *G. max* (L.) Merr., a cultigen, and its wild annual progenitor *G. soja* Sieb. and Zucc., both of which carry 2n = 40 chromosomes. *Glycine soja* is found in the Peoples' Republic of China, Russia, Korea, Japan, and Taiwan. Table 8.1 shows species, 2n chromosome number, nuclear and plastome genomes, and geographical distribution of *Glycine* species.

Classical taxonomy has played a major role in the identification and nomenclature of new species in the genus *Glycine*. *Glycine clandestina* (2n = 40) has been observed to be morphologically a highly variable species (Hermann, 1962). The curved and short pod forms have been segregated from *G. clandestina sensu lato* as *G. cyrtoloba* (Tindale, 1984) and *Glycine microphylla* (Tindale, 1986), respectively. The separation of both morphological forms is logical, because both forms differ genomically from other species (Singh and Hymowitz, 1985), c; Singh, Kollipara, and Hymowitz, 1988). Recently, two more morphotypes have been segregated from

G. clandestina and have been identified taxonomically as G. rubiginosa and G. peratosa (Pfeil et al., 2001).

The genomes of diploid species are assigned capital letter symbols according to the degree of chromosome homology between species in F_1 hybrids (Kihara and Lilienfeld, 1932). Similar letter symbols are designated for species with interspecific F_1 hybrids that show normal chromosome pairing. Minor chromosome differentiation is indicated by placing a subscript after the letter. Highly differentiated species are designated by different letter symbols, because their hybrids exhibit highly irregular chromosome pairing (see *Brassica* — U, 1935; *Glycine* — Singh and Hymowitz, 1985b; *Gossypium* — Beasley, 1940; *Oryza* — Nezu, Katayama, and Kihara, 1960; *Triticum* — Kihara and Nishiyama, 1930).

Genome designations of the *Glycine* species have been made arbitrarily, assuming the diploid chromosome number to be 2n = 40. Beginning with G. canescens (A genome), the genomes of the other species were designated based on chromosome pairing, hybrid seedling lethality, and hybrid seed inviability in interspecific hybrids. The same genome symbol designations were assigned to species possessing similar genomes, but the genomes were differentiated by placing a subscript after the letter, such as A1 (G. clandestina), A2 (G. argyrea), and A3 (G. latrobeana); B (G. microphylla), B_1 (G. latifolia), B_2 (G. tabacina), and B_3 (G. stenophita). Genome symbols were assigned to species as soon as their relationships were established: C = G cyrtoloba; $C_1 = G$. curvata; D = G. tomentella; E = G. tomentella, 2n = 38; F = G. falcata (Singh, Kollipara, and Hymowitz, 1988, 1992a). It should be pointed out that cytogenetic studies, isozyme banding patterns, and molecular investigations have demonstrated at least three distinct genomic forms in diploid G. tomentella (Brown, 1990; Doyle, Doyle, and Brown, 1990a; Singh, Kollipara, and Hymowitz, 1988, 1992a). The genomic designations are broadly functional and primarily based on chromosome pairing rather than evolutionary definition. The frequency of bivalent formation is the primary direct measure of chromosome homology, whereas crossability (generating fertile hybrids) is an indirect measure of chromosome homology. Genomic groups typically correlate with evolutionary groups; they do not reflect speciation per se.

III. GENOMIC RELATIONSHIPS AMONG DIPLOID SPECIES

A. Crossing Affinity

Crossability rate is an excellent indirect measure for estimating the degree of genomic relationship between parental species. Interspecific crosses involving parental species with similar genomes usually set normal pods and seeds, while in crosses between genomically dissimilar species, seed abortion is common, or the hybrid is sterile. Certain hybrid combinations do not set seed, and sometimes, crosses are successful only in one direction and often are genotype dependent (*Cajanus* — Dundas et al., 1987; *Capsicum* — Pickersgill, 1988; *Cuphea* — Ray, Gathman, and Thompson, 1989; *Glycine* — Singh, Kollipara, and Hymowitz, 1988; *Gossypium* — Skovsted, 1937; *Hibiscus* — Menzel and Martin, 1970; *Medicago* — McCoy and Bingham, 1988; *Vigna* — Chen, Baker, and Honma, 1983).

In the genus *Glycine*, intragenomic crosses set mature pods, while pod abortion is common in intergenomic hybrids (Table 8.2). In *Avena*, Baum and Fedak (1985) reported no internal barriers to gene flow among the diploid species of Strigosa group (*A. hirtula*, A. *wiestii*, *A. strigosa*, *A. atlantica*).

The failure of intragenomic crosses can be attributed to a wide range of phenomena (e.g., genetically determined species isolation mechanism, pollen tube dysfunction on alien stigmas, genetic incompatibilities that produce physiologically unfit embryos) but is often correlated with long periods of reproductive isolation. Menzel and Martin (1970) attempted to cross *Hibiscus cannabinus* (2n = 36) and *H. asper* (2n = 36) and were not successful (*H. cannabinus* × *H. asper*)

Cross	Genome	Pod Set/Total Florets	% Pod Set
G. argyrea × G. canescens	$A_2 \times A$	6/26	23.3
G. microphylla \times G. latifolia	$B \times B_1$	3/40	7.5
G. latifolia \times G. tabacina	$B_1 \times B_2$	18/147	12.2
G. canescens $ imes$ G. clandestina	$A \times A_1$	9 + 1ª/86	10.5 + 1.2ª
G. canescens × G. microphylla	A×B	1ª/20	5.0ª
G. canescens × G. latifolia	$A \times B_1$	14ª/462	4.9 ^a
G. canescens $ imes$ G. tabacina	$A \times B_2$	0/395	0.0
G. canescens ×G. tomentella	A × D	21/150	14.0
G. latifolia $ imes$ G. tomentella	$B_1\timesD$	8ª/280	2.9ª

 Table 8.2
 Pod Set in F₁ Hybrids between Wild Perennial Glycine Species

^a Aborted pods.

Source: From Singh, R.J., Kollipara, K.P., and Hymowitz, T., *Genome*, 29, 490–497, 1987b. With permission.

158 crosses; reciprocal, 130 crosses). Both species possess a similar genome but are isolated geographically. *Hibiscus asper* is a native of West Africa, whereas *H. cannabinus* is found in East Africa. However, reproductive isolation is not always a predictor of incompatibility. *Oryza sativa* (2n = 24), a native of Asia, and *O. glaberrima* (2n = 24), an inhabitant of Africa, are isolated geographically, but they hybridize readily and produce vigorous hybrids with complete meiotic chromosome pairing (12II). However, the hybrids are sterile (Nezu, Katayama, and Kihara, 1960; Bouharmont, Olivier, and Dumonte de Chassart, 1985).

B. Chromosome Pairing

The chromosome pairing (chiasmata frequency) method "Analysartoren-Methode" established by Kihara and his associates (Kihara, 1924, 1930; Kihara and Nishiyama, 1930; Kihara and Lilienfeld, 1932; Lilienfeld, 1951; Kihara, 1963) to ascertain the genomes of hexaploid wheats (2n = 6x = 42) and their relatives, originally founded by Rosenberg (1909), is still a reliable technique (Kimber and Feldman, 1987; Jauhar and Crane, 1989; Wang, 1989), despite a few objections (de Wet and Harlan, 1972; Baum, Estes, and Gupta, 1987).

A prerequisite for Kihara's method is to ascertain chromosome pairing relationships among all available diploid species in order to establish diploid analyzers. Based on this principle, genomic relationships among diploid species have been assessed for several important genera, such as *Arachis* (Smartt, Gregory, and Pfluge Gregory, 1978; Wynne and Halward, 1989), *Avena* (Rajhathy and Thomas, 1974; Ladizinsky, 1974; Nishiyama, Yabuno, and Taira, 1989), *Brassica* (U, 1935; Röbbelen, 1960; Prakash and Hinata, 1980), *Bromus* (Armstrong, 1981), *Cajanus* (Subrahmanyam, Kumar, and Faris, 1986), *Capsicum* (Pickersgill, 1988; Egawa and Tanaka, 1984), *Dactylis* (Lumaret, 1988), *Eleusine* (Chennaveeraiah and Hiremath, 1974), *Glycine* (Singh and Hymowitz, 1985b,c; Singh, Kollipara, and Hymowitz, 1988, 1992a), *Gossypium* (Skovsted, 1937; Beasley, 1940; Stephens, 1947, 1950; Phillips, 1966; Endrizzi, Turcotte, and Kohel, 1985; Percival and Kohel, 1991), *Helianthus* (Chandler, Jan, and Beard, 1986), *Hibiscus* (Menzel and Martin, 1970), *Hordeum* (Bothmer, Flink, and Landström, 1986), *Nicotiana* (Goodspeed, 1954; Smith, 1968), *Oryza* (Nezu, Katayama, and Kihara, 1960; Nayar, 1973), *Paspalum* (Burson, 1981a,b), *Pennisetum* (Jauhar, 1981; Pantulu and Rao, 1982), *Secale* (Khush, 1962; Stutz, 1972; Singh and Röbbelen, 1977), and *Triticum* and its allied genera (Lilienfeld, 1951; Dewey, 1984; Kimber and Feldman, 1987).

The degree of chromosome pairing in interspecific hybrids provides an important cytogenetic context for inferring phylogenetic relationships among species, enhances our understanding of evolution of the genus, and provides information about the ancestral species. Smith (1968) recognized five categories of chromosome pairing in F_1 hybrids:

- · Complete or almost complete pairing
- "Drosera scheme" pairing
- High, variable pairing
- Low, variable pairing
- Minimal pairing

In category one intragenomic hybrids, the pairing of chromosomes is by definition, complete in 75 to 100% of the sporocytes, although occasionally, species will differ by chromosomal interchanges. The classical examples of chromosomal interchanges differentiating species are the species of *Oenothera* (Cleland, 1962), *Datura* (Avery, Satina, and Rietsema, 1959), and *Secale* (Schiemann and Nürnberg-Krüger, 1952; Riley, 1955). Sometimes, genomically similar species differ by inversions (Singh, Kollipara, and Hymowitz, 1988). "*Drosera scheme*" pairing (2xII + 1xI) occurs in tetraploid (4x) × diploid (2x) crosses and has been named after chromosome pairing in *Drosera longifolia* (2n = 4x = 40) × *Drosera rotundifolia* (2n = 2x = 20) (Rosenberg, 1909). High and low variable chromosome pairing occurs in intergenomic hybrids, where some but not all chromosomes retain some level of genetic and structural homology. The degree of pairing between chromosomes depends upon genomic affinity of the parental species. Minimal chromosome pairing (all chromosomes remain univalent in over 50% of the sporocytes) occurs in F₁ hybrids, where two species have diverged sufficiently to obscure any ancestral chromosome homology.

Several numerical methods have been developed to interpret chromosome pairing data (Gaul, 1959; Driscoll, Bielig, and Darvey, 1979; Driscoll, Gordon, and Kimber, 1980; Kimber, Alonso, and Sallee, 1981; Alonso and Kimber, 1981; Kimber and Alonso, 1981). Menzel and Martin (1970) used a genome affinity index (GAI) for comparing the degree of homology among parental genomes. The mean number of "bivalent-equivalents" (II-equiv.) is divided by the base chromosome number. Because base chromosome numbers have been debated for a majority of the crop species, it is suggested to use, for diploid species, gametic chromosome numbers instead of a base chromosome number of parents to compute GAI. For example, a base chromosome x = 10 has been suggested for the genus *Glycine* (2n = 40) (Darlington and Wylie, 1955). However, *Glycine* species with 2n = 2x = 20 have not been identified so far (Singh and Hymowitz, 1985b; Kumar and Hymowitz, 1989). A bivalent and trivalent are each equal to one II-equiv., but a quadrivalent with chiasmata in at least three arms is counted as two II-equivalent.

1. Intragenomic Chromosome Pairing

Generally, species with similar genomes exhibit complete or almost complete chromosome pairing in their hybrids (Figure 8.1A). Sometimes, species differ by chromosomal interchanges or by paracentric inversions (Figure 8.1B).

In the genus *Triticum*, A-genome species (2n = 2x = 14), *T. monococcum* and *T. urartu*, are found in the wild in the same geographical area. They exhibit seven bivalents at metaphase-I, and mainly, the Drosera scheme (7II + 7I) of pairing in the *T. turgidum* (2n = 4x = 28) × *T. urartu* cross (Johnson, 1975), suggesting that *T. turgidum* is an allotetraploid carrying an A-genome derived either from *T. monococcum* or *T. urartu*. Furthermore, *T. monococcum* or *T. urartu* possess identical karyotypes (Giorgi and Bozzini, 1969) and insignificantly different nuclear DNA content (Furuta, Nishikawa, and Yamaguchi, 1986), suggesting that they carry similar genomes.

Kerby and Kuspira (1987) identified five diploid (2n = 2x = 14) species in the section *Sitopsis*: *Aegilops speltoides* (S), *Aegilops sheronensis* (S^{sh}), *Aegilops longissima* (S¹S¹), *Aegilops bicornis*(S^b), and *Aegilops searsii* (S^s). Hybrids between *Aegilops longissima* and *Aegilops sheronensis* display univalents (0–2), bivalents (5–7), trivalents (0–1), and quadrivalents (0–1), and almost normal pollen (81.3–87.1%) and seed (81.0–95.0%) fertility. This suggests that both species carry similar genomes and differ only by a reciprocal translocation (depends on the type of quadrivalents — alternate, ring). Although *Aegilops bicornis* is given a different superscript genome symbol (S^b),

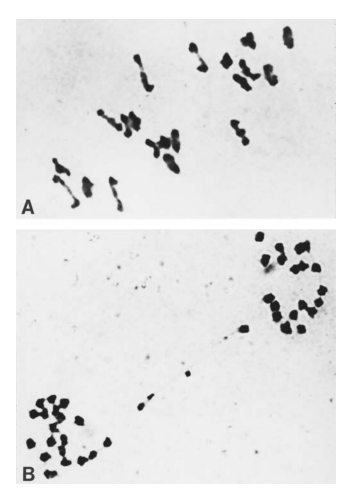


Figure 8.1 Meiosis in *Glycine latifolia* (B₁B₁) × *Glycine microphylla* (BB): A, metaphase-I showing 20 bivalents. (From Singh, R.J., unpublished results). B, anaphase-I showing a chromatin bridge and an acentric fragment in interspecific hybrid of *Glycine clandestina* (A₁A₁) × *Glycine canescens* (AA). (Figure 8B from Singh, R.J., Kollipara, K.P., and Hymowitz, T., *Genome*, 30, 166–176, 1988. With permission.)

it shows almost the same meiotic behavior, pollen, and seed set in crosses with *Aegilops sheronensis* and *Aegilops longissima* (Feldman, Strauss, and Vardi, 1979).

Hybrids Aegilops speltoides (S) × Aegilops bicornis (S^b) and Aegilops sheronensis (S^{sh}) × Aegilops speltoides (S) showed three to seven bivalents and set no seed. Furthermore, crosses Aegilops speltoides × Aegilops longissima and Aegilops searsii × Aegilops longissima showed a quadrivalent configuration together with univalents, bivalents, and trivalents (Feldman, Strauss, and Vardi, 1979). This indicates a considerable amount of diversity among the B-genome species of the genus Aegilops.

In the genus *Glycine*, all F_1 plants from crosses among A- and B-genome species display 20 bivalents in a majority of sporocytes. However, *G. microphylla* (B) × *G. tabacina* (B₂) F_1 s showed an 0.83 GAI value, suggesting some differences between genomes of B and B₂ (Table 8.3). A chromatin bridge and an acentric fragment at anaphase-I were clearly seen in A × A₁, B × B₁, B₁ × B₂, and B × B₂ hybrids.

Based on classical taxonomy, *G. soja* and *G. max* are different species (Hermann, 1962). Both species carry 2n = 40 chromosomes, hybridize readily, produce viable, vigorous, and fertile hybrids, and sometimes differ by a reciprocal translocation (Palmer et al., 1987; Singh and Hymowitz, 1988) or by a paracentric inversion (Ahmad, Britten, and Byth, 1977). Therefore, *G. soja* and *G. max* possess similar genomes (G).

Hybrids (Genomes)					
	_	=	≥ + III	GAIª	Authority
1. Glycine $(n = 20)$					
rescens $(A_2 \times A)$	0.8 (0-4)	19.6 (18–20)		0.99	Singh, Kollipara, and Hymowitz, 1988
(1	0.1 (0–2)	19.9 (19–20)		1.0	Singh, Kollipara, and Hymowitz, 1988
	6.8 (0–12)	16.6 (14–20)		0.83	Singh, Kollipara, and Hymowitz, 1988
latifolia \times microphylla (B ₁ \times B) 2.	2.0 (0-4)	19.0 (18–20)		0.95	Singh and Hymowitz, 1985c
	0.6 (0-4)	19.7 (18–20)		0.99	Singh and Hymowitz, 1985c
arboreum $ imes$ herbaceum (${\sf A}_2 imes {\sf A}_1$)	0.2	11.1	0.9	0.99	Gerstel, 1953
anomalum $ imes$ triphyllum (${ m B_1} imes { m B_2}$)	0.10	12.90		1.0	Douwes, 1953
raimondii $ imes$ gossypioides ($D_{5} imes D_{6}$)	1.14	12.43		0.96	Menzel and Brown, 1955
stocksii $ imes$ areysianum (E $_1 imes$ E $_2$)	0.16	12.20		0.97	Douwes, 1953
somalense $ imes$ areysianum (E $_2$ $ imes$ E $_3$)	0.20	12.90		0.99	Douwes, 1953
3. Oryza (n = 12)					
sativa $ imes$ perennis (A $ imes$ A)	0.0	12.0		1.0	Nezu, Katayama, and Kihara, 1960
sativa $ imes$ glaberrima (A $ imes$ A $^{ m g}$)	0.0	12.0		1.0	Nezu, Katayama, and Kihara, 1960
glaberrima $ imes$ perennis (A $^{\mathfrak{g}}$ $ imes$ A)	0.0	12.0		1.0	Nezu, Katayama, and Kihara, 1960
officinalis $ imes$ eichingeri (C $ imes$ C)	0.0	12.0		1.0	Ogawa and Katayama, 1973
4. Hordeum $(n = 7)$					
	0.24 (0–2)	6.81 (5–7)	0.05 (0–1)	0.98	Bothmer, Flink, and Landström, 1986
Ĥ	0.19 (0-4)	6.90 (5–7)		0.99	Bothmer, Flink, and Landström, 1986
	0.40 (0–6)	6.76 (4–7)		0.97	Bothmer, Flink, and Landström, 1986
patagonicum \times flexuosum (H \times H) 0.1	.79 (0–4)	6.60 (5–7)		0.94	Bothmer, Flink, and Landström, 1986
patagonicum \times stenostachys (H \times H) 0.4	0.43 (0–4)	6.79 (5–7)		0.97	Bothmer, Flink, and Landström, 1986

Table 8.3 Interspecific Hybrids with Complete or Almost Complete Chromosome Pairing at Diakinesis and Metaphase-I

GENOME ANALYSIS

Another approach for determining the chromosome homology between species is to analyze the chromosome pairing of F_1 hybrids at pachynema. This technique, however, is feasible only for species with analyzable pachytene chromosomes. By utilizing pachytene chromosomes, Singh and Hymowitz (1988) identified 20 pachytene bivalents of *G.* max × *G.* soja F_1 hybrids. Heteromorphic regions were observed only in Chromosomes 6 and 11, while pairing was completely normal all along the length of the long and short arms of the remaining chromosome pairs.

A close genomic relationship between a cultigen and its wild progenitor suggests that they belong to one biological species or Primary Gene Pool, the GP-1 of Harlan and de Wet (1971). Diploid (2n = 26) cultivated cottons (G. herbaceum-A₁, G. arboreum-A₁) are closely related to the B-genome (G. anomalum-B₁, G. typhyllum-B₂, G. capitis-viridis-B₃) wild species (Endrizzi, Turcotte, and Kohel, 1985). Hybrids $A_1 \times B_1$ and $A_2 \times B_1$ exhibit almost complete chromosome pairing (Skovsted, 1937). However, more recent phylogenetic studies, suggest that the two Gossypium A genome species are more closely related to the F-genome species, Gossypium longicalyx (Cronn et al., 2002). Cultivated rice (O. sativa, 2n = 24) and its wild annual progenitor, O. nivara (2n = 24) 24) are cross compatible and show essentially normal chromosome pairing in F_1 , indicating that O. sativa and O. nivara have the same genomic (A) constitution (Dolores, Chang, and Ramirez, 1979). Dundas et al. (1987) recorded almost complete pairing involving Cajanus cajan (2n = 22) with 2n = 22 chromosome C. acutifolius and C. confertiflorus. In the classical study of Nicotiana, 90% of the intrasectional hybrid combinations displayed complete or almost complete chromosome pairing (Goodspeed, 1954). In Hordeum, North and South American species have the same (H) genome designation, although complete genome homology (GAI, range = 0.52-0.82) is lacking (Bothmer, Flink, and Landström, 1986). For reasons of clarity, it is suggested that genome symbol H_1 be designated to North American species and H_2 to South American species. In general, chloroplast DNA variation results support isoenzyme, molecular, cytological, and crossing data (Doebley, von Bothmer, and Larson, 1992). This suggests that evolutionary divergence within a group occurred because of geographical isolation and accumulation of chromosomal structural changes during the time (Stebbins, 1950; Grant, 1971).

A wrong conclusion can be reached if genomic relationship is based on fertility or sterility of F_1 hybrids without analyzing the chromosome pairing. One might expect to obtain fertile F_1 hybrids between two genomically similar species, but find the hybrids to be sterile. This could be due to cryptic structural hybridity, complementary lethal genes, or differentiation in genes and chromosomal structures (Stebbins, 1950). A classical example of cryptic structural hybridity is the *Primula verticillata* (2n = 18) × *Primula floribunda* (2n = 18) hybrid. Chromosome pairing was almost normal (nine loosely associated bivalents), but the hybrid was completely sterile (Newton and Pellew, 1929). In *Oryza*, intragenomic hybrids between the A- (Nezu, Katayama, and Kihara, 1960) and C-genome species (Ogawa and Katayama, 1973) exhibit cryptic structural hybridity. Hybrids of *Avena canariensis* (AcAc) × *A. damascena* (AdAd) showed seven normal bivalents but almost complete sterility after selfing (Leggett, 1984; Nishiyama, Yabuno, and Taira, 1989). Likewise F_1 s from a *Triticum monococcum* × *Triticum urartu* cross exhibited complete chromosome pairing (6.97II), yet the hybrids was so regular that it was assumed that reduction in hybrid fertility was caused by genic or cryptic structural hybridity (Lumaret, 1988).

One expects to observe normal chromosome pairing in morphologically similar species. However, there are some exceptions. For example, aneudiploid (2n = 38) and diploid (2n = 40) *G. tomentella* are morphologically indistinguishable, yet at metaphase-I, a majority of the chromosomes remain as univalents (GAI 0.31), while bivalents are loosely connected and rod shaped (Singh, Kollipara, and Hymowitz, 1988). Similarly, *Paspalum notatum* $(2n = 20) \times P$. *vaginatum* (2n = 20) F₁s exhibited 20I in 57% of the sporocytes and 1II + 18I (weakly synapsed) in 30% of the sporocytes. These species belong to different taxonomic groups but have similar morphological features (Burson, 1981a). Likewise, *Cuphea procumbens* (2n = 18) and *Cuphea crassiflora* (2n = 24) are morphologically similar, but their hybrids predominantly exhibit univalents (Ray, Gathman, and Thompson, 1989).

2. Intergenomic Chromosome Pairing

The genomic relationships among genomically dissimilar species can be determined by the degree of chromosome pairing in their hybrids. The extent of chromosome association in the hybrids elucidates structural homology in the parental chromosomes, and hence, furnishes evidence regarding the progenitor species (Phillips, 1966; Smith, 1968).

Usually the F_1 hybrids generated from genomically unlike parents (different biological species) are germinated through *in vitro* techniques. In general, hybrids are weak, slow in vegetative and reproductive growth, and sterile. The lack of seed set is attributed to the reduced amount of chromosome pairing that results in the formation of chromosomally unbalanced gametes. In the subgenus *Glycine*, A- and B-genome species show intergenomic chromosome association (Figure 8.2). Hybrid seed inviability, seedling lethality, and vegetative lethality are common occurrences in distant intergenomic crosses (Stebbins, 1958; Hadley and Openshaw, 1980). The reduced chromosome pairing between distantly related species is attributed to structural differences between chromosomes (chromosomal) perpetuated during speciation, rather than to asynaptic or desynaptic genes found in certain genetic stocks. According to Stebbins (1958), doubling chromosomes of chromosomally sterile F_1 hybrids restores fertility by providing each genome a set of perfect homologs with which to pair at meiosis. Chromosome pairing is regular, and often, fertility is restored. In contrast, desynapsis persists in giving genic sterility after doubling the chromosomes of asynaptic or desynaptic genetic stocks.

3. Variable and Minimum Chromosome Pairing

Variable (semihomologous–homoeologous) and minimum chromosome pairing are common in intergenomic F₁ hybrids (Tables 8.4 and 8.5). The classical examples reported are from the genera *Brassica* (U, 1935; Mizushima, 1950a,b; Attia and Röbbelen, 1986a,b), *Glycine* (Grant et al., 1984; Singh, Kollipara, and Hymowitz, 1988, 1992a), *Gossypium* (Skovsted, 1937; Endrizzi and Phillips, 1960; Phillips, 1966); *Hordeum* (Bothmer, Flink, and Landström, 1986), *Nicotiana* (Goodspeed, 1954), *Oryza* (Nezu, Katayama, and Kihara, 1960; Shastry and Ranga Rao, 1961; Katayama, 1982), *Paspalum* (Burson, 1981a,b), and *Triticum* (Kihara and Lilienfeld, 1932; Dewey, 1984).



Figure 8.2 Meiosis in *Glycine latifolia* (2n = 40) × *Glycine canescens* (2n = 40) hybrid; metaphase-I showing 20 univalents + 10 bivalents. (From Singh, R.J. and Hymowitz, T., *Theor. Appl. Genet.*, 71, 221–230, 1985b. With permission.)

	Average Chr	Average Chromosome Pairing (Range)	(Range)		
Hybrids (Genomes)	_	=	2 + 1	GAI	Authority
1. Glycine $(n = 20)$					
clandestina $ imes$ microphylla (A $_1 imes B$)	23.6 (18–30)	8.2 (5–11)		0.41	Singh, Kollipara, and Hymowitz, 1988
latifolia $ imes$ argyrea (${ m B_1} imes { m A_2}$)	19.7 (12–30)	10.2 (5–14)		0.51	Singh, Kollipara, and Hymowitz, 1988
canescens × microphylla (A × B)	22.2 (14–26)	8.9 (7–13)		0.45	Singh and Hymowitz, 1985b
latifolia \times canescens (B ₁ \times A)	20.9 (12–32)	9.5 (4–14)		0.48	Singh and Hymowitz, 1985b
tomentella $ imes$ clandestina (D $ imes$ A ₁)	13.6 (4–22)	13.3 (9–18)		0.67	Singh and Hymowitz, 1985b
argyrea \times tomentella (D \times A ₂)	18.6 (8–24)	10.7 (8–16)		0.54	Singh, Kollipara, and Hymowitz, 1993
argyrea × cyrtoloba ($A_2 \times C$)	30.8 (20–38)	4.6 (1–10)		0.23	Singh, Kollipara, and Hymowitz, 1988
latifolia $ imes$ cyrtoloba ($B_1 imes C$)	29.9 (26–36)	5.05 (2–7)		0.25	Singh, Kollipara, and Hymowitz, 1988
tomentella $ imes$ canescens (E $ imes$ A)	30.0 (25–37)	4.5 (1–7)		0.23	Singh, Kollipara, and Hymowitz, 1988
tomentella \times tomentella (E \times D)	26.6 (19–33)	6.2 (3-10)		0.31	Singh, Kollipara, and Hymowitz, 1988
2. Gossypium $(n = 13)$					
herbaceum $ imes$ sturtianum ($A_{I} imes C_{I ext{-n}}$)	10.53	7.46	0.17	0.59	Phillips, 1966
arboreum × raimondii ($A_2 \times D_5$)	13.79	5.90	0.13	0.47	Endrizzi and Phillips, 1960
arboreum $ imes$ stocksii ($A_2 imes E_1$)	14.90	5.50		0.42	Skovsted, 1937
stocksii × arboreum ($E_1 \times A_2$)	18.40	3.80		0.29	Beasley, 1942
anomalum $ imes$ sturtianum (${\sf B}_1 imes{\sf C}_{1-n}$)	9.23	8.09	0.16	0.64	Phillips, 1966
anomalum $ imes$ davidsonii (${\sf B}_1 imes {\sf D}_{3-d}$)	19.90	3.05		0.23	Skovsted, 1937
anomalum $ imes$ klotzschianum (${\sf B}_{ extsf{1}} imes {\sf D}_{ extsf{3} extsf{k}}$)	17.45	4.26	0.01	0.33	Phillips, 1966
anomalum $ imes$ stocksii ($B_1 imes E_1$)	20.57	2.70		0.21	Douwes, 1951
sturtianum $ imes$ armourianum (C $_{1-n} imes$ D $_{2-1}$)	8.45	8.2	0.35	0.66	Skovsted, 1937
3. <i>Oryza</i> $(n = 12)$					
sativa $ imes$ officinalis (A $ imes$ C)	14.31(16–24)	4.8 (0-4)	0.022 (0–1)	0.40	Shastry et al., 1961
punctata × eichingeri (B × C)	18.52	2.74		0.23	Katayama and Ogawa, 1974
punctata $ imes$ officinalis (B $ imes$ C)	17.02	3.49		0.29	Katayama and Ogawa, 1974
4. Hordeum $(n = 7)$					
<i>bogdani × patagonicum (</i> I × H)	7.38(0–13)	3.33(1–7)		0.48	Bothmer, Flink, and Landström, 1986
brevisubulatum $ imes$ brachyantherum (I $ imes$ H)	10.24 (6–14)	1.88 (0-4)		0.27	
brevisubulatum $ imes$ brachyantherum (I $ imes$ H)	3.20 (0–8)	5.24 (3–7)		0.75	Bothmer, Flink, and Landström, 1986
5. Brassica $(n = 8, 9, 10)$					
oleracea $ imes$ campestris (C $ imes$ A)	2.3	7.3	0.61 ^a		Attia and Röbbelen, 1986a
oleracea \times nigra (C \times B)	13.6	1.9	0.02		and Röbbelen,
campestris $ imes$ nigra (A $ imes$ B)	8.75	4.36	0.71		Attia and Röbbelen, 1986a
^a Includes V. VI (0.02).					

Table 8.4 Interspecific Hvbrids with Variable (Semi homologous) Chromosome Pairing at Diakinesis and Metaphase-I

^a Includes V, VI (0.02).

In intergenomic hybrids, most of the bivalents are rod shaped and are weakly associated, suggesting a weak chromosomal (genomic) affinity. Thus, the higher the number of univalents, the lesser the genomic affinity. In *Nicotiana*, approximately 90% of the intersectional diploid hybrids showed semihomologous chromosome pairing (Goodspeed, 1954). A considerable amount of chromosome homology exists among A, B, and D genomes of *Triticum aestivum* (2n = 6x = 42) (Dewey, 1984). Chromosome association at metaphase-I ranged from 1II + 12I to 7II in *Triticum monococcum* (2n = 14, A) × *Aegilops speltoides* (2n = 14, B) (Kihara and Lilienfeld, 1932).

Chromosome pairing among three elementary species of *Brassica*, namely, *Brassica campestris* (AA, 2n = 20), *Brassica oleracea* (CC, 2n = 18), and *Brassica nigra* (BB, 2n = 16), showed a high degree of chromosome association in A × C (mean 0.02V,VI + 0.29IV + 0.30III + 7.3II + 2.3I) and a low degree of affinity in B × C (mean 0.02III + 1.9II + 13.6I) and A × B (0.01IV + 0.7III + 4.36II + 8.75I) crosses (Attia and Röbbelen, 1986a). This suggests that *Brassica nigra* is distantly related to *Brassica oleracea* and *Brassica campestris*.

Minimum chromosome pairing is usually observed in intersubgeneric and intergeneric hybrids. For example, an intersubgeneric hybrid of *Glycine max* (2n = 40, G) and *Glycine clandestina* (2n = 40, A₁) exhibits 40 univalents in 40.9% of the sporocytes, and one to six are loosely paired rodshaped bivalents (Singh, Kollipara, and Hymowitz, 1987a). Burson (1981a,b) observed little genome homologies among five diploid (2n = 20) *Paspalum* species (*Paspalum intermedium*, I; *Paspalum jurgensii*, J; *Paspalum vaginatum*, D; *Paspalum setaceum*, S; *Paspalum notatum*, N), as a majority of the sporocytes carried 20 univalents. In a *Nicotiana glauca* (2n = 24) × *Nicotiana plumbaginifolia* cross, over 75% of the sporocytes showed zero bivalents (Goodspeed, 1954). Williams and Pandey (1975) recorded only zero to two rod-shaped bivalents in a *Nicotiana glauca* (2n = 24) × *Nicotiana glauca* (2n = 24) cross. In certain Eurasian and South American *Hordeum* species hybrids, Bothmer, Flink, and Landström (1986) observed negligible chromosome pairing (Table 8.5). Diploid (2n = 2x = 14) species hybrids of *Hordeum* and *Secale* exhibited 14 univalents in a majority of the sporocytes (Gupta and Fedak, 1985).

IV. GENOMES OF GLYCINE

In the genus Glycine, genome A shows partial genome homology with genome B and genome D but a stronger genomic affinity with D than with B. Genome B shows no similarity with D, because hybrids are seedling lethal (Singh, Kollipara, and Hymowitz, 1988). Genome C from *Glycine cyrtoloba* shows slightly stronger (may be insignificant) affinity with B ($B \times C = 6.9II$) + 26.2I; $B_1 \times C = 5.05II + 29.9I$) than with A ($A_2 \times C = 4.6II + 30.8I$). Moreover, $A \times C$ and $A_1 \times C$ are vegetatively lethal. Likewise, genome E from an eudiploid *Glycine tomentella* (2n = 38) exhibits limited chromosome pairing with genomes A and D (Table 8.4). In A \times E at metaphase-I, the average chromosome associations (range) were 4.5II(1-7) + 30.0I(25-37), and in D \times E, they were 6.2II (3–10) + 26.6I (19–33). In these crosses, bivalents are usually rod shaped and are loosely connected at metaphase-I; univalents are scattered in the cytoplasm (Figure 8.3A). In the subgenus *Glycine*, *Glycine falcata* is a unique species, because it differs from the other species in several morphological traits (Hermann, 1962; Hymowitz and Newell, 1975), seed protein composition (Mies and Hymowitz, 1973), oil and fatty acid contents (Chavan, Hymowitz, and Newell, 1982), ribosomal gene variation (Doyle and Beachy, 1985), and phytoalexin production (Keen, Lyne, and Hymowitz, 1986). Chromosome pairing results support the uniqueness of genome (F) of Glycine falcata, because they show minimum chromosome synapsis with A and B genomes (Figure 8.3B). Metaphase-I in F_1 s A \times F and B \times F showed 40 univalents in a majority of the sporocytes. An average chromosome association (range) of 1.12II(0-6) +37.8I (28–40) was found in $B_1 \times F$. (Table 8.5).

Information on chromosome pairing (translated into genomic affinity index) in intergenomic hybrids may shed some light regarding which genome is most likely to be the progenitor genome

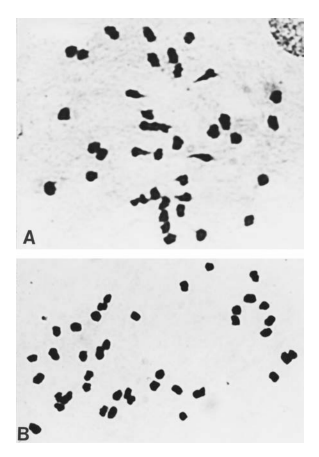


Figure 8.3 Meiosis in interspecific *Glycine* hybrids: A, metaphase-I showing 31 univalents + 4 bivalents in *Glycine tomentella* (2n = 38, EE) × *Glycine canescens* (2n = 40, AA); B, metaphase-I showing 40 univalents in (*Glycine clandestina* × *Glycine canescens*) × *Glycine falcata.* (From Singh, R.J., Kollipara, K.P., and Hymowitz, T., *Genome*, 30, 166–176, 1988. With permission.)

of the genus. In the genus *Glycine*, it is proposed that *Glycine falcata* is the ancestral species. The absence of chromosome homology of F with A and B genomes suggests that A and B genome species diverged into two evolutionary pathways from a common gene pool; which one differentiated first is uncertain. Divergence within A and B genomes occurred because of geographical isolation. Chromosomal structural changes and morphological variants were maintained due to the autogamous nature of pollination. In further divergence from A genome species, *Glycine tomentella* evolved, and it is likely that *G. clandestina* (A₁A₁) may have played a role, because D has closer affinity (16.1II + 7.9I) with *G. clandestina* than those observed with *G. canescens* (AA) (13.0II + 14.0I) or with *G. argyrea* (A₂A₂) (12.6II + 14.8I). Aneudiploid (2n = 38) *Glycine tomentella* (E) originated from D, because both cytotypes are morphologically indistinguishable, and E has a higher genomic affinity with D (0.31) than that recorded with A (0.23). Genome E could not be derived from B, C, and F genomes because there is no genomic affinity. It is suggested that *Glycine cyrtoloba* (C) probably originated from B genome species and *Glycine curvata* (C₁C₁) is a variant of *Glycine cyrtoloba*.

Darlington and Wylie (1955) proposed that x = 10 is the basic chromosome number for the cultivated soybean. Based on this proposal, Singh, Kim, and Hymowitz (2001) hypothesized a putative ancestor with 2n = 20 chromosomes for the genus *Glycine* and carrying at least a pair of nucleolus organizer regions (NORs). Although such a progenitor is currently unknown, it would be most likely found in South East Asia (Cambodia, Laos, Vietnam) and has not been collected

Average Chromosome Pairing (Bange)	Average Chro	Average Chromosome Pairing (Bange)	ia (Bande)		
Hybrids (Genomes)	_	=		GAI	Authority
1. Glycine $(n = 20)$					
microphylla $ imes$ falcata (B $ imes$ F)	38.6 (32–40)	0.7 (0-4)		0.04	Singh, Kollipara, and Hymowitz, 1992a
latifolia $ imes$ falcata ($B_1 imes F$)	37.8 (28–40)	1.12 (0–6)		0.06	Singh, Kollipara, and Hymowitz, 1988
tomentella $ imes$ microphylla (E $ imes$ B)	38.2 (35–39)	0.40 (0–2)		0.02	Singh, Kollipara, and Hymowitz, 1992a
max \times clandestina (G \times A ₁)	37.8 (28–40)	1.15 (0-6)		0.06	Singh, Kollipara, and Hymowitz, 1987a
2. Gossypium $(n = 13)$					
somalense × australe ($E_2 \times C$)	23.78	1.11		0.09	Phillips, 1966
somalense $ imes$ bickii (E $_2 imes$ G $_1$)	25.58	0.21		0.02	Phillips, 1966
3. Oryza $(n = 12)$					
sativa $ imes$ australiensis (A $ imes$ E)	23.6	0.2 (0–1)		0.02	Nezu, Katayama, and Kihara, 1960
sativa \times officinalis (A \times C)	24.0	0.0 (0-1)		00.0	Nezu, Katayama, and Kihara, 1960
officinalis $ imes$ australiensis (C $ imes$ E)	23.0	0.5 (0–3)		0.04	Nezu, Katayama, and Kihara, 1960
4. Hordeum $(n = 7)$					
marinum $ imes$ bogdani (X $ imes$ H)	13.71 (12–14)	0.14 (0–1)		0.02	Bothmer, Flink, and Landström, 1986
marinum $ imes$ brachyantherum (X $ imes$ H)	13.10 (10–14)	0.45 (0–2)		0.06	Bothmer, Flink, and Landström, 1986
bulbosum $ imes$ patagonicum (I $ imes$ H)	13.90 (12–14)	0.06 (0–2)		0.01	Bothmer, Flink, and Landström, 1986
marinum $ imes$ muticum (X $ imes$ H)	13.00 (10–14)	0.50 (0–2)		0.07	Bothmer, Flink, and Landström, 1986
roshevitzii $ imes$ patagonicum (I $ imes$ H)	13.42 (11–14)	0.18 (0–1)	0.03 (0–1)	0.03	Bothmer, Flink, and Landström, 1986
5. Paspalum ($n = 10$)					
jurgensii $ imes$ intermedium (J $ imes$ I)	16.48 (10–20)	1.70 (0–5)		0.17	Burson, 1981a
jurgensii $ imes$ vaginatum (J $ imes$ D)	18.22 (8–20)	0.88 (0–6)		0.09	Burson, 1981a
jurgensii $ imes$ setaceum (J $ imes$ S)	17.73 (10–20)	1.14 (0–5)		0.11	Burson, 1981a
intermedium \times notatum (I \times N)	18.43 (12–20)	0.78 (0-4)		0.08	Burson, 1981b
notatum $ imes$ vaginatum (N $ imes$ D)	18.83 (12–20)	0.59 (0-4)		0.06	Burson, 1981b

Table 8.5 Interspecific Hybrids with Minimum Chromosome Pairing at Diakinesis and Metaphase-I

and identified so far. Whether tetraploidization (2n = 4x = 40) involved auto- (spontaneous chromosome doubling) or allo- (interspecific hybridization followed by chromosome doubling) polyploidy of the progenitor species and whether it occurred prior to dissemination or after, cannot be substantiated experimentally, because we do not know where the progenitor of the genus *Glycine* originated. The progenitor of the wild perennial species of the subgenus *Glycine* radiated out into several morphotypes depending on the growing conditions in the Australian subcontinent. These species have never been domesticated and remained as wild perennials. In contrast, the pathway of migration from a common progenitor to China is assumed as: wild perennial (2n = 4x = 40;unknown or extinct) \rightarrow wild annual $(2n = 4x = 40; Glycine soja) \rightarrow$ soybean (2n = 4x = 40;cultigen). All known species of the genus *Glycine* exhibit diploid-like meiosis and are inbreeders (Singh and Hymowitz, 1985a).

Allopolyploidization probably played a key role in the speciation of the genus *Glycine*. This implies that the 40-chromosome *Glycine* species and the 80-chromosome *Glycine tabacina* and *Glycine tomentella* are tetraploid and octoploid, respectively. The expression of four rDNA loci in *Glycine curvata* and *Glycine cyrtoloba* (Singh, Kim, and Hymowitz, 2001) strongly supports the hypothesized allotetraploid origin suggested by cytogenetic (Singh and Hymowitz, 1985b; Xu et al., 2000b) and molecular studies (Shoemaker et al., 1996).

V. GENOMES OF GOSSYPIUM

The genus *Gossypium* consists of 45 diploid (2n = 2x = 26) and five allotetraploid (2n = 4x = 52) species (Fryxell, 1992). Diploid species constitute eight genomic groups (A, B, C, D, E, F, G, K) and have a wide geographical distribution. Allotetraploids are limited to the New World (Table 8.6; Figure 8.4).

Genomic relationships among *Gossypium* species (2n = 26) and their genome designations have been based on chromosome pairing in intra- and interspecific hybrids, and they correlate strongly with geographical distribution, morphological features, karyotype analysis, seed protein banding patterns, and phylogenetic analysis of gene sequences (Stephens, 1947, 1950; Endrizzi, Turcotte, and Kohel, 1985; Cronn et al., 2002).

The A-genome species (*Gossypium herbaceum* = A_1 ; *Gossypium arboreum* = A_2) are the cultigens of Africa and Asia and differ by a reciprocal translocation (Gerstel, 1953). Of the three B-genome species, *Gossypium anomalum* (B₁) has been studied extensively, has disjunct distribution in northern and southern Africa, and shows almost complete pairing with B₂ (*Gossypium triphyllum* — Southern Africa) and B₃ (*Gossypium capitis-viridis* — Cape Verde Islands). The C, G, and K genomes are confined to Australia. Edwards and Mirza (1979) designated *Gossypium bickii*, an Australian wild species, genome symbol G based on karyotype comparison with *Gossypium herbaceum* (A₁) and *Gossypium sturtianum* (C₁). Total chromosome length of *Gossypium bickii* was found to be shorter (55.54 µm) than C₁ (74.04 µm) and A₁ (66.54 µm) genome species. Furthermore, they did not observe nucleolus organizer regions in *Gossypium bickii*. However, this study needs to be repeated.

The D-genome species are found in the New World (Arizona, Mexico, Galapagos Islands, Peru). It is interesting to note that despite a wide geographical distribution, D-genome species are morphologically variable but carry similar genomes. The E-genome species, *Gossypium stocksii* (E_1), *Gossypium somalense* (E_2), *Gossypium areysianum* (E_3), and *Gossypium incanum* (E_4) are distributed from eastern Africa, north and east through the southern tip of the Arabian peninsula, to Pakistan (Phillips, 1966). Three species, *G. benadirense*, *G. bricchettii*, and *G. vollesenii* have also been assigned to the E genome but are poorly known (Percival, Wendel, and Stewart, 1999). All the E-genome species possess similar genomes.

Phillips and Strickland (1966) suggested that *Gossypium longicalyx* from Africa carries the F genome, as it differs from the D-genome (21.60I in $D \times F$ cross). Schwendiman, Koto, and Hau (1980) confirmed the F-genome designation to *Gossypium longicalyx*, because it differs from

	Species	Genome	Distribution
	Diploid (2 <i>n</i> = 26)		
	genus Sturtia (R. Brown) Todaro [All the indigenous stralian species]		
1.	Section \rightarrow <i>Sturtia</i> [Species do not deposit terpenoid		
	aldehydes in the seeds (gossypol)]		
	i. G. sturtianum J.H. Willis	C ₁	Australia
	ii. G. robinsonii F. Mueller	C ₂	Australia
2.	Section \rightarrow <i>Grandicalyx</i> Fryxell [Unusual perennial, thick underground root-stock, fat bodies on seeds]		
	i. <i>G. costulatum</i> Todaro	K	Australia
	ii. <i>G. cunninghamii</i> Todaro	K	Australia
	iii. G. exiguum Fryxell, Craven, and Stewart	К	Australia
	iv. G. rotundifolium Fryxell, Craven, and Stewart	К	Australia
	v. G. enthyle Fryxell, Craven, and Stewart	К	Australia
	vi. G. nobile Fryxell, Craven, and Stewart	к	Australia
	vii. <i>G. pilosum</i> Fryxell	ĸ	Australia
	viii. <i>G. pulchellum</i> (C.A. Gardner) Fryxell	ĸ	Australia
	ix. <i>G. londonderriense</i> Fryxell, Craven, and Stewart	ĸ	Australia
	x. <i>G. marchantii</i> Fryxell, Craven, and Stewart	K	Australia
	xi. <i>G. populifolium</i> (Bentham) F. Mueller ex Todaro	K	Australia
	xii. <i>G. "anapoides</i> " (not yet formally described)	K	Australia
0	Section \rightarrow <i>Hibiscoidea</i> Todaro [Species do not deposit	ĸ	Australia
3.	terpenoid aldehydes in the seeds]		
	i. <i>G. australe</i> F. Mueller	G	Australia
			Australia
	, , ,	G	
<u> </u>	iii. <i>G. bickii</i> Prokhanov	G	Australia
Me	genus <i>Houzingenia</i> (Fryxell) Fryxell [New World → primarily xico; large shrubs or small trees]		
1.	Section \rightarrow Houzingenia		
	a. Subsection: Houzingenia	_	
	i. <i>G. thurberi</i> Todaro	D ₁	Mexico, Arizona (United States)
	ii. <i>G. trilobum</i> (DC) Skovsted	D ₈	Mexico
	b. Subsection: Integrifolia (Todaro) Todaro		
	i. <i>G. davidsonii</i> Kellog	D_{3-d}	California (United States)
	ii. G. klotzschianum Anderson	D _{3-k}	California (United States)
	c. Subsection: Caducibracteolata Mauer		
	i. G. armourianum Kearney	D ₂₋₁	Mexico
	ii. <i>G. harknessii</i> Brandegee	D ₂₋₂	Mexico
	iii. <i>G. thurberi</i> Fryxell	D ₋₁₀	Mexico
2.	Section \rightarrow <i>Erioxylum</i> (Rose and Standley) Prokhanov		
	a. Subsection: Erioxylum		
	i. G. aridum (Rose and Standley ex Rose) Skovsted	D_4	Mexico
	ii. G. lobatum H. Gentry	D ₇	Mexico
	iii. <i>G. laxum</i> Phillips	D ₉	Mexico
	iv. G. schwendimanii Fryxell and S. Koch	D ₁₁	
	b. Subsection: <i>Seler</i> a (Ulbrich) Fryxell		
	i. <i>G. gossypioides</i> (Ulbrich) Standley	D_6	Mexico
	c. Subsection: <i>Austroamericana</i> Fryxell	0	
	i. <i>G. raimondii</i> Ulbrich	D_5	Peru
Suk	genus Gossypium L.	D_5	i ciu
5uu 1.	• ,,		
ι.	Section \rightarrow <i>Gossypium</i>		
	a. Subsection: Gossypium	٨	Africo Aria
	i. <i>G. herbaceum</i> L.	A ₁	Africa, Asia

Table 8.6	Snacias	Genomic S	vmhole	and	Distribution	of the	Genue	Gossypium L.
	Species,	Genomic 3	ymbols,	anu	Distribution	or the	Genus	Gussypium ∟.

			Species	Genome	Distribution		
		ii.	G. arboreum L.	A ₂	Asia		
	b.	Subs	ection: Anomala Todaro				
		i.	G. anomalum Wawra and Peyritsch	B ₁	Africa		
		ii.	<i>G. triphyllum</i> (Harvey and Sonder) Hochreutiner	. B ₂	Africa		
		iii	G. capitis-viridis Mauer	B ₃	Cape Verde Islands		
	с.	Subs	ection: Longiloba Fryxell				
		i.	G. longicalyx J.B. Hutchinson and Lee	F ₁	Africa		
	d.	Subs	ection: Pseudopambak (Prokhanov) Fryxell				
		i.	G. benadirense Mattei	E	African-Arabian		
		ii.	G. bricchettii (Ulbrich) Vollesen	E	African-Arabian		
		iii.	G. vollesenii Fryxell	E	African-Arabian		
		iv.	G. stocksii Masters ex. Hooker	E1	African-Arabian		
		V.	G. somalense (Gürke) J.B. Hutchinson	E ₂	African-Arabian		
		vi.	G. areysianum Deflers	E ₃	African-Arabian		
		vii.	G. incanum (Schwartz) Hillcoat	. E ₄	African-Arabian		
			Allotetraploid $(2n = 4x = 52)$				
D.	Subgen	us <i>Kar</i> j	pas Rafinesquåe [The allotetraploid cottons]				
		i.	G. hirsutum L.	AD ₁	Central America		
		ii.	G. barbadense L.	AD ₂	South America		
		iii.	G. tomentosum Nuttall ex Seemann	AD ₃	Hawaiian Islands		
		iv.	G. mustelinum Miers ex Watt	AD_4	NE Brazil		
		V.	G. darwinii Watt	AD_5	Galapagos Islands		

Table 8.6 (continued) Species, Genomic Symbols, and Distribution of the Genus Gossypium L.

Source: From Percival, A.E., Wendel, J.F., and Stewart, J.M., In: Cotton: Origin, History, Technology, and Production, W.C. Smith, Ed., John Wiley & Sons, New York, 1999, 33–63. With permission.

Gossypium stocksii karyologically. According to Schwendiman, Koto, and Hau (1980), *Gossypium longicalyx* appears to be a "mixed" genome related to all the other genomes (except D). Recent phylogenetic analysis suggests a close relationship between *G. longicalyx* and the two A-genome species (Cronn et al., 2002).

Considering karyotype reduction during the course of speciation (Stebbins, 1950), the relative chromosome size of *Gossypium* species reveals the following karyotypic (in decreasing order) relationships: C (largest) \rightarrow EF \rightarrow B \rightarrow A \rightarrow G \rightarrow D (smallest) (Stephens, 1947; Katterman and Ergle, 1970; Edwards and Mirza, 1979). Thus, species with the C-genome should be considered as ancestral. However, by utilizing the frequency of univalents observed in the intergenomic hybrids, Phillips (1966) postulated that the E genome is ancestral in the genus, because it has negligible genomic affinity with A, B, C, D, and G genomes. However, the E genome has a greater affinity with B and A than with C and G genomes. The evolutionary pathways of the genus *Gossypium* shown in Figure 8.4. are based on relationships among diploid species. The origin of the allotetraploids (New World AD-genome) following intergenomic hybridization is described. More recent data (Cronn et al., 2002) highlight the difficulty in pinpointing the basal divergence in the genus *Gossypium*. The early stages of evolution are characterized by a rapid radiation, and thus, no genome can be regarded as "ancestral." The traditional conception of the genus being replaced by a rapid radiation of the proto-*Gossypium* milieu into the D, ABF, E, CGK lineages, begins, most likely, with the divergence between the New World (D genome) and Old World/Oceanic lineages (ABCEGK).

VI. ANCESTORS OF ALLOPOLYPLOID SPECIES

Interspecific hybridization between morphologically and genomically distinct diploid species followed by chromosome doubling (amphidiploidy–allopolyploidization) has played a prominent role

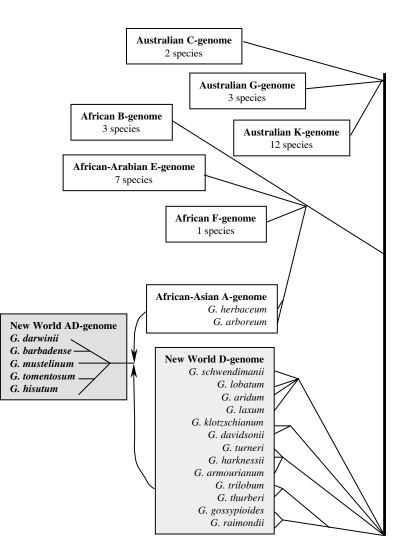


Figure 8.4 Evolutionary history of the *Gossypium*. Genomic relationships among diploid species. The origin of allotetraploids following intergenomic hybridization. (From Percival, A.E., Wendel, J.F., and Stewart, J.M., In: *Cotton: Origin, History, Technology, and Production*, W.C. Smith, Ed., John Wiley & Sons, 1999, 33–63. With permission.)

in the speciation of plants. A large number of our agriculturally important crops such as Brassicas, coffee, cotton, oat, sugarcane, tobacco, and wheat are natural allopolyploids. The characteristic features of allopolyploids are vigorous and aggressive growth habit, diploid-like meiosis (allosyndesis), normal fertility, and true breeding.

Traditionally, ancestors of allopolyploids were inferred based on geographical distribution, morphological features, chromosome count, karyotype analysis, and isozyme banding patterns. Molecular phylogenetics studies have confirmed many of these early hypotheses, supplemented by data from interspecific hybridizations, resynthesis of allopolyploids, and comparison of the morphological traits, meiotic pairing, and fertility in the allopolyploids and their hybrids. These procedures have been used to explain the ancestors of natural allopolyploid species of genera *Brassica* (Prakash and Hinata, 1980), *Glycine* (Singh, Kollipara, and Hymowitz, 1989; Singh et al., 1992b), *Gossypium* (Beasley, 1940; Harland, 1940), *Nicotiana* (Goodspeed and Clausen, 1928; Clausen, 1932; Greenleaf, 1941), and *Triticum* (McFadden and Sears, 1946). These examples are briefly outlined as follows.

A. Brassica (B. carinata 2n = 4x = 34; B. juncea 2n = 4x = 36; B. napus 2n = 4x = 38)

The genus *Brassica* consists of three elementary or basic genome species (*B. campestris* n = 10 = A; *B. oleracea* n = 9 = C; *B. nigra* n = 8 = B), and the species are secondary polyploids from an extinct species with a base chromosome number x = 6. This was elegantly demonstrated by Röbbelen (1960) based on pachytene chromosome analysis.

It was observed that *Brassica nigra* (B) is double tetrasomic (genome A B C DD E FF), *Brassica oleracea* (C) is triple tetrasomic (genome A BB CC D EE F), and *Brassica campestris* (A) is doubly tetrasomic and a hexasomic (AA B C DD E FFF). Each letter designates a particular chromosome. Thus, three basic genomes are derived from a basic chromosome number x = 6. Natural allotetraploids *Brassica napus* (n = 19, AC), *B*rassica *juncea* (n = 18, AB), and *Brassica carinata* (n = 17, BC) exhibit diploid-like meiosis and disomic inheritance.

U (1935) determined the ancestral species of *Brassica napus*, *Brassica juncea*, and *Brassica carinata* by studying the meiotic chromosome pairing in triploid hybrids of *Brassica napus* × *Brassica campestris*, *Brassica napus* × *Brassica oleracea*, *Brassica carinata* × *Brassica oleracea*, and *Brassica carinata* × *Brassica nigra* (Figure 8.5). The chromosome association of 2x bivalent + x univalent predominated in triploid hybrids. Further confirmation of the ancestral genomes of allotetraploids is provided by synthesizing amphidiploids. Mizushima (1950b) synthesized *Brassica carinata* (2n = 34, BBCC) by hybridizing *Brassica nigra* (2n = 16, BB) and *Brassica oleracea* (2n = 18, CC), and subsequently doubled the chromosomes by colchicine. Amphidiploid (2n = 4x = 34, BBCC) exhibited 0–4IV and 9–17II at metaphase-I. Anaphase-I and second meiosis were regular in 80% of the cells, and pollen was nearly fertile. In contrast, *B. carinata* synthesized by

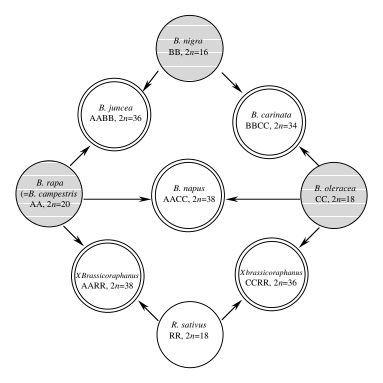


Figure 8.5 The Brassica-Raphanus diamond (extension of U's triangle), showing the species relationships of the major cruciferous crops. [Brassica nigra (black mustard); Brassica juncea (Indian or brown mustard); Brassica rapa (Oriental vegetables); Brassica carinata (Ethiopian mustard); Brassica oleracea (various types of cole crops); Brassica napus (oil-seed rape, fodder rape); x Brassic-oraphanus, AARR (Cultivar group Raparadish); CCRR, Radicole; and Raphanus sativus (Fodder radish, oil-seed radish). (From Lange, W. et al., Euphytica, 40, 1–14, 1989. With permission.)

Pearson (1972) showed normal meiosis (17II) and high seed fertility. Similarly, *Brassica juncea* (2n = 36, AABB) and *Brassica napus* (2n = 38, AACC) have been generated artificially by crossing *Brassica campestris* $(2n = 20, AA) \times Brassica nigra$ (2n = 16, BB) and *Brassica campestris* $(2n = 20, AA) \times Brassica nigra$ (2n = 16, BB) and *Brassica campestris* $(2n = 20, AA) \times Brassica$ nigra (2n = 16, BB) and *Brassica campestris* $(2n = 20, AA) \times Brassica$ nigra (2n = 16, BB) and *Brassica campestris* $(2n = 20, AA) \times Brassica$ nigra (2n = 18, CC), respectively. Synthesized *Brassica juncea* and *Brassica napus* exhibit normal meiosis, resemble morphologically the respective natural allotetraploid species, and are fertile (Prakash and Hinata, 1980).

It has been observed that the established allotetraploids have a more stable meiotic behavior than the newly synthesized amphidiploids of the same species. Such stabilization is based on genetic control (Attia and Röbbelen, 1986b).

Studies on seed protein electrophoresis (Uchimiya and Wildman, 1978), chloroplast DNA variation (Palmer et al., 1983; Yanagino, Takahata, and Hinata, 1987), mitochondrial DNA variation (Palmer, 1988), and nuclear restriction fragment length polymorphisms (RFLPs) (Song, Osborn, and Williams, 1988) have confirmed the established cytogenetic studies that three allotetraploid species are derived from the interspecific hybridization of three elementary diploid species. Yanagino, Takahata, and Hinata (1987) observed significant correlation between the species relationships as seen with the chloroplast DNA and cytogenetic studies, but not with those based on numerical morphological taxonomy. Another example is for the genus *Oryza*, where molecular approaches simply verified the phylogenetic relationships among species established based on taxonomy, hybridization, isozymes, and cytogenetics (Wang, Second, and Tanksley, 1992).

Lange et al. (1989) extended the *Brassica* triangle of U (1935) to the *Brassica-Raphanus* diamond (Figure 8.5), representing the genomic relationships among species of the majority of the Cruciferous crops. The introduction of *Raphanus sativus* in interspecific hybridization in *Brassica* created two new hybrids: (1) Radicole = *Raphanus sativus* (female) × *Brassica oleracea* (male) \rightarrow amphidiploid \rightarrow x *Brassicoraphanus* (2n = 36; CCRR) and (2) Raparadish = *Raphanus sativus* (female) × *Brassica rapa* (male) \rightarrow amphidiploid \rightarrow x *Brassicoraphanus* (2n = 36; CCRR) and (2) Raparadish = *Raphanus sativus* (female) is the major disadvantage in radicole and raparadish.

B. Glycine (G. tomentella 2n = 4x = 80)

In the genus *Glycine*, 80-chromosome *G. tomentella* has wide geographical distribution, aggressive and vigorous growth habit, and diploid-like meiosis in its accessions. Meiotic pairing in intra- and interspecific hybrids suggested that the 80-chromosome *Glycine tomentella* evolved through allopolyploidization (Singh and Hymowitz, 1985a; Singh, Kollipara, and Hymowitz, 1987b). Morphologically, 40- and 80-chromosome *Glycine tomentella* are indistinguishable, indicating that one of the ancestors of 80-chromosome tomentellas is a 40-chromosome *Glycine tomentella*.

Triploid F₁ hybrids of *Glycine tomentella* (2n = 80-Queensland = Qld.) × *Glycine canescens* (2n = 40, AA) and *Glycine tomentella* (2n = 80-Qld) × *Glycine tomentella* (2n = 40, DD) exhibit the "Drosera Scheme" (20 bivalents + 20 univalents) of pairing in a majority of the cells (Table 8.7; Figure 8.6A). Therefore, *Glycine canescens* and *Glycine tomentella* (2n = 40) are the probable ancestors of 80-chromosome *Glycine tomentella* from Qld, Australia (Putievsky and Broué, 1979; Singh and Hymowitz, 1985a,b; Singh, Kollipara, and Hymowitz, 1989).

The artificially synthesized amphidiploid (AADD) of *Glycine tomentella* $(2n = 40, DD) \times Glycine canescens (2n = 40, AA)$ is morphologically almost identical to the natural 80-chromosome *Glycine tomentella*. Average chromosome associations (ranges) in AADD were 0.13 IV (0–1) + 39.6 II (38–40) + 0.3 I (0–2). A total of 75% of the sporocytes showed 40 bivalents (Figure 8.6B), and 90% of the sporocytes exhibited 40–40 chromosome migration at anaphase-I. Pollen fertility was 84.8%, and the plants set normal pods and seeds (Table 8.8).

Singh, Kollipara, and Hymowitz (1989) hybridized the synthesized amphidiploid (AADD) with four *Glycine tomentella*-Qld (2n = 80) accessions (Table 8.8). Hybrid seed germinated normally. Seedlings grew vigorously, carried the expected chromosome numbers of 2n = 80, and were highly

	Origin of	Origin of Number		Chromosome	Chromosome Association ^b		Total	Seed
Hybrid	E ¹ a	F, Plants	2 <i>n</i>	Univalent	Bivalent	Trivalent	PMC	Set
CAN $(2n = 40) \times TOM (2n = 80)$								
440928×441005	SC	ო	60	24.9 (18–34)	24.9 (18–34) 16.9 (13–21) 0.44 (0–1)	0.44 (0–1)	25	ST ^{c,d}
TOM $(2n = 80) \times CAN (2n = 40)$								
441005 × 440932	თ	ო	60	23.5 (20–30)	18.3 (15–20)	I	12	ST®
446958 × 440928	SC	0	60	23.0 (19–30)	18.2 (15–20)	0.2 (0–1)	10	ST⁺
TOM $(2n = 40) \times TOM (2n = 80)$								
505267 imes 505214	SC	0	60	20.4 (18–24)	19.8 (18–21)	I	18	ST
505222×505256	SC	ო	60	20.8 (16–26)	19.6 (17–22)	I	51	ST
505267 imes 441005	SC	-	60	20.0 (18–24)	20.0 (18–21)		30	ST
^a S = seed: SC = seed culture.								

ൽ

^b Means with range in parentheses. ^c ST = sterile.

^d Singh, R.J. and Hymowitz, T., *Theor. Appl. Genet.*, 71, 221–230, 1985b. With permission
 ^e Singh, R.J. and Hymowitz, T., *Z. Pflanzenzüchtg.*, 92, 289–310, 1985c. With permission
 ^f Singh, R.J., Kollipara, K.P., and Hymowitz, T., *Genome*, 29, 490–497, 1987b. With permission. *Note:* CAN = Glycine canescens; TOM, Glycine tomentella.

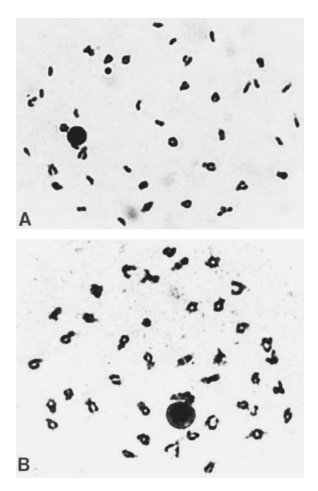


Figure 8.6 Meiosis in (A) an intraspecific hybrid of *Glycine tomentella* (2n = 40, DD) × *Glycine tomentella* (2n = 80), diakinesis showing 20 univalents + 20 bivalents; (B) synthesized amphidiploid [*Glycine tomentella* (2n = 40, DD) × *Glycine canescens* (2n = 40, AA) F₁ 2n = 40 AD = colchicine treatment = 2n = 80, AADD = H321CT) diakinesis showing 40 bivalents. (From Singh, R.J., Kollipara, K.P., and Hymowitz, T., *Genome*, 32, 796–801, 1989. With permission.)

fertile. A majority of the sporocytes showed complete pairing. The conclusion drawn from these results is that *Glycine canescens* (AA) and *Glycine tomentella* (DD) are the ancestors of *Glycine tomentella*-Qld (2n = 80). A similar approach was used to determine the putative diploid ancestors of tetraploid (2n = 80) *Glycine tabacina* (Singh et al., 1992b). By using the chloroplast DNA polymorphism technique, Doyle et al. (1990b) reached the same conclusion.

C. Gossypium (G. hirsutum, G. barbadense, 2n = 4x = 52)

The genus *Gossypium* carries five allotetraploid (2n = 4x = 52) species, and all are distributed in the Americas (New World). It has been demonstrated that allotetraploid cottons are of an amphidiploid origin between diploid (2n = 2x = 26) cultivated Asiatic (Old World) cottons and diploid wild American cottons (Skovsted, 1934, 1937; Wendel, 1989).

Skovsted (1934, 1937) reported that the American allotetraploid species (*Gossypium hirsutum*; *Gossypium barbadense*) contains a set of 13 large and 13 small chromosomes. The small chromosomes are contributed by the American diploid (D-genomes) species, while large chromosomes are donated by cultivated Asiatic (A-genome) cotton. Triploid hybrids between Amer-

			ancoren							
				Chre	Chromosome Association ^b	sociation ^b			Pollen	
	Origin	No. F,						Total	Fertility	Seed
Hybrid	of F ₁ ª	Plants	2n	Univalents	Bivalents	Trivalents	Trivalents Quadrivalents	PMC	(%)	Set°
TOM $(2n = 40) \times [CAN \times CAN (2n = 40)]CT, 2n = 80,446993 \times 146993 \times 146933 \times 146033 \times 146033 \times 146033 \times 146033 \times 146033 \times 1460333 \times 146033 \times 146033 \times 146033 \times 1460333 \times 1460333 \times 146033 {{10003333 \times 1400333 \times 146033 \times 140033 \times 1400333 \times 1400333 \times 14003		-	80	0.3 (0–2)	39.6 (38–40)	0.0	0.13 (0–1)	40	84.8	ш
[440936 × 440928] = H321C1 H321CT × 441005 (2 <i>n</i> = 80)	S	N	80	4.7 (0–14)	37.2 (33_40)	0.0	0.2 (0–1)	42	89.5	ш
H321CT \times 505214 (2 n = 80)	S	N	80	0.4 (0–2)	39.8	0.0	0.0	30	99.4	ш
H321CT \times 505215 (2 n = 80)	S	N	80	2.0 (0–6)	(39-40) 38.6 (35 40)	0.0	0.2 (0–2)	33	81.3	ш
H321CT \times 505256 (2 n = 80)	S	4	80	3.6 (0–10)	37.6 37.6	0.0	0.3 (0–2)	85	86.0	ш
					(04-40)					

^a S, seed ^b Means with ranges in parentheses. ^c F, fertile *Note:* TOM, *G. tornentella*; CAN, *G. canescens*; CT, colchicine treatment; H, hybrid.

Source: From Singh, R.J., Kollipara, K.P., and Hymowitz, T., Genome, 32, 796-801, 1989. With permission.

	Average Chromosome Pairing					
Hybrids $(4x \times 2x)$	I	I	III			
G. barbadense $(AD)_2 \times G$. aridum (D_4)	13.99	12.28	0.15			
G. hirsutum $(AD)_1 \times G$. aridum (D_4)	13.15	12.40	0.35			
G. barbadense (AD) ₂ × G. armourianum (D ₂ -1)	14.75	12.05	0.05			
G. darwinii (AD) ₅ × G. armourianum (D ₂ -1)	13.20	12.00	0.60			
G. barbadense $(AD)_2 \times G$. trilobum (D_8)	12.70	12.40	0.50			
G. hirsutum $(AD)_1 \times G$. trilobum (D_8)	13.50	12.45	0.20			
G. barbadense $(AD)_2 \times G$. anomalum (B_1)	33.80	2.60				
G. barbadense $(AD)_2 \times G.$ stocksii (E_1)	37.90	0.55				

Table 8.9 Chromosome Pairing in Triploid (2n = 39) Interspecific Hybrids of Cotton

Source: From Skovsted, A., J. Genet., 34, 97-134, 1937. With permission.

ican tetraploids × Asiatic diploids (Skovsted, 1934; Beasley, 1942) and American tetraploids × American diploids exhibit the "Drosera Scheme" (13II + 13I) of chromosome association (Table 8.9). This suggests that the New World tetraploid cottons are allotetraploid with only the A and D genome species being the donor parents, because triploids F_1 s from *Gossypium barbadense* (AD₂) × *Gossypium anomalum* (B₁) and *Gossypium barbadense* × *Gossypium stocksii* (E₁) exhibit mainly univalents (Table 8.9).

Beasley (1940) and Harland (1940) independently synthesized amphidiploid $(A_2A_2D_1D_1)$ by doubling the chromosome number in *Gossypium arboreum* (cultivated Asiatic cotton) × *Gossypium thurberi* (wild American cotton). Synthesized allotetraploid plants were female fertile but were male sterile, because boll and seed set were normal when pollinated with the American *n* = 26 chromosome pollen. These studies demonstrate convincingly that the cultivated Asiatic cotton-A genome and wild American cotton-D genome are the ancestors of the tetraploid American cottons. The question of where hybridization and amphidiploidy occurred are still being debated (Valiček, 1978; Endrizzi, Turcotte, and Kohel, 1985; Wendel, 1989). Seed protein electrophoresis also revealed that the genomes similar to A₁ (*Gossypium herbaceum*) and D₅ (*Gossypium raimondii*) are the probable ancestors of the natural allotetraploid genomes (Cherry, Kotterman, and Endrizzi, 1970).

Another approach toward elucidating the relationship between the A and D genome species and the tetraploid American cotton species is to synthesize amphidiploids involving genomes other than A and D, to hybridize the newly synthesized allotetraploids with natural tetraploid American cottons, and to analyze crossability rate, hybrid seed viability, meiotic chromosome pairing, and pollen and seed fertility of hybrids. Brown (1951) identified a spontaneously induced amphidiploid of *Gossypium davidsonii* (D_{3-d}) and *Gossypium anomalum* (B_1). Meiotic chromosome pairing in the amphidiploid was 26 bivalents in nine out of the 23 sporocytes studied. Univalents ranged from two to four, and some sporocytes showed a trivalent or a quadrivalent. Most bivalents were normally synapsed with two chiasmata, and the number of bivalents averaged 25.0. Amphidiploid *davidsonii-anomalum* set aborted bolls when crossed with *G. hirsutum*, and seeds failed to germinate. This shows that *Gossypium anomalum* (B_1) is not a genome donor species to allotetraploid American cottons.

D. Nicotiana (N. tabacum 2n = 4x = 48)

The genus *Nicotiana* contains species with n = 9, 10, 12, 16, 19, 20, 21, 22, and 24 chromosomes from a base x = 6 chromosome species that probably is now extinct. *Nicotiana tabacum* is an allotetraploid (2n = 4x = 48) cultivated species, belonging to the genus *Nicotiana*, subgenus *tabacum*, and section *Genuinae*. Because of its commercial value, *Nicotiana tabacum* has been studied extensively cytogenetically (Goodspeed, 1954; Smith, 1968). The allotetraploid nature of *Nicotiana tabacum* was demonstrated by Goodspeed and Clausen (1928) by studying a series of triploid F_1 hybrids involving *Nicotiana sylvestris* (2n = 2x = 24; S^1 genome), *Nicotiana tomentosa* (2n = 2x = 24; T^1 genome), and *Nicotiana tabacum* (SSTT) triangle. The F_1 hybrids of *Nicotiana sylvestris* × *Nicotiana tabacum* and *Nicotiana tomentosa* × *Nicotiana tabacum* exhibited classical "Drosera Scheme" (12II + 12I) chromosome pairing, suggesting that *Nicotiana sylvestris* and *Nicotiana tomentosa* are the probable ancestors of *N. tabacum*, which most likely originated in Central South America. Both diploid species are found there today, and haploid *Nicotiana tabacum* (2n = 2x = 24) displays a complete lack of meiotic chromosome pairing similar to that in *Nicotiana sylvestris* × *Nicotiana tomentosa* F_1 hybrids (Goodspeed, 1954).

Clausen (1932), on the basis of morphological features of *Ncotiana sylvestris* × *Nicotiana tomentosiformis* and *Nicotiana sylvestris* × *Nicotiana tomentosa*, suggested that *Nicotiana tomentosiformis* satisfies more nearly the requirements of a progenitor of *Nicotiana tabacum* than does *Nicotiana tomentosa*. However, *Nicotiana tomentosa* and *Nicotiana tomentosiformis* are closely related and are differentiated by a reciprocal translocation (Goodspeed, 1954).

Greenleaf (1941) synthesized amphidiploids *Nicotiana sylvestris* × *Nicotiana tomentosa* and *Nicotiana sylvestris* × *Nicotiana tomentosiformis*. Both amphidiploids showed normal meiosis and over 90% pollen fertility but were completely female sterile, although megasporogenesis was quite normal. The sterility was due to genetic factors. Morphologically, the *Nicotiana sylvestris* × *Nicotiana tomentosiformis* amphidiploid closely resembled *Nicotiana tabacum*.

Gerstel (1960) used a genetic approach to learn whether the chromosomes of *Nicotiana tomentosiformis* or those of *Nicotiana otophora* are more nearly homologous with one genome of *Nicotiana tabacum*. Several recessive marker stocks of *Nicotiana tabacum* (SSTT) were crossed to both diploid species carrying the corresponding dominant allele (T¹T¹). The chromosome number of interspecific triploid (STT¹) F₁ hybrids was doubled by treating emerging seedlings with 0.1% aqueous colchicine solution for 3 h at room temperature. The segregation ratios were obtained by back crossing the amphidiploids (SSTTT¹T¹) to recessive tester stocks of *Nicotiana tabacum*. The amphidiploids (TTT¹T¹) had the duplex genotype (ZZzz). The observed segregation ratios directly reflect the genetic output of the amphidiploid parents. A ratio of 5:1 (random chromosome segregation) or 3.7:1 (chromatid segregation) are expected if chromosomes of T and T¹ genomes form multivalents. However, the observed ratios were smaller even than the 3.7:1 ratio expected with random chromatid segregation but might fit the maximum equational (3.5:1) segregation (Table 8.10). Segregation ratios were higher in hexaploid (*Nicotiana tabacum* × *Nicotiana otophora*) crosses. This indicates that *Nicotiana tomentosiformis* is more closely related than to *Nicotiana otophora* to *Nicotiana tabacum* in chromosome homology.

A similar approach was used by Gerstel (1963) to show that *N. sylvestris* has a close genomic affinity with a genome of *Nicotiana tabacum*. However, it was concluded that "some chromosomes

Morphological	(N. tab	acum × N.	tomentos	iformis)	(N. tabac	um × N.	otophora)	
Traits	++	-	Ratio	6x as	++	-	Ratio	6 <i>x</i> as
fs (Fasciated)	87	29	3.0:1	Male	195	56	3.5:1	Male
Pb (Purple bud)	147	44	3.3:1	Female	329	39	8.4:1	Female
,	381	114	3.3:1	Male	342	77	4.4:1	Male
<i>ws</i> (White seedling)	762	95	3.5:1	Female	1887	98	9.1:1	Female
	2481	309	3.5:1	Male	644	44	6.8:1	Male
yb (Early yellow leaves)	314	100	3.1:1	Female	159	19	8.4:1	Female
yg (Yellow green)	107	23	4.7:1	Female	785	53	14.8:1	Female

Table 8.10 Segregation of Morphological Traits in 6x Amphidiploids of Nicotiana

Note: ++ = dominant. - = recessive.

Source: From Gerstel, D.U., Genetics, 45, 1723-1734, 1960. With permission.

of the two species have remained completely homologous while others have become differentiated to some degree during evolution."

According to Gerstel (1960), "It is quite likely that none of the now living species of that taxonomic group is identical with the parent that once went into ancestral *Nicotiana tabacum*, but perhaps it is not entirely futile to attempt to designate the modern species closest to that ancestor." I concur with his opinion.

E. Triticum (T. aestivum, 2n = 6x = 42)

On the basis of plant morphology and nuclear genomic constitution, species of the genus *Triticum* can be classified into three main groups: einkorn (2n = 2x = 14), emmer (2n = 4x = 28), and dinkel (2n = 6x = 42) (Sears, 1975). The emmer and dinkel wheats are allopolyploids. The mode of chromosome pairing in triploid (2n = 3x = 21) and pentaploid (2n = 5x = 35) hybrids helped Kihara to uncover the ancestral species of the allopolyploid wheats. Triploid (*Triticum turgidum*, $2n = 4x = 28 \times Triticum monococcum$, 2n = 2x = 14) and pentaploid (*Triticum turgidum* × *Triticum aestivum*) hybrids exhibit 7II + 7I and 14II + 7I chromosome configurations, respectively. This indicates that diploid (*Triticum monococcum*), tetraploid (*Triticum turgidum*), and hexaploid (*Triticum aestivum*) have one (AA), two (AABB), and three (AABBDD) sets of seven chromosomes each, and that the A genome is present in tetraploid and hexaploid wheats (Kihara and Nishiyama, 1930). This suggests that hexaploid wheats are allohexaploids and originated from the hybridization of one species of the emmer group (AABB) and *Aegilops tauschii* (DD) (Figure 8.7).

1. The Source of the A-Genome

It has been demonstrated, as shown above, and also verified by Giemsa C-banding (Gill and Kimber, 1974; Gill, Friebe, and Endo, 1991), and sequences of purothionins — small basic proteins (Jones et al., 1982) — that *Triticum monococcum* is the A-genome donor. Another diploid species, *Triticum urartu*, which was suggested as the B-genome donor (Johnson, 1975), is similar genomically to *Triticum monococcum* (Dvořák, 1976; Chapman, Miller, and Riley, 1976). Indications are that *Triticum urartu* rather than *Triticum monococcum* is the source of the A-genome (Kerby and Kuspira, 1987).

2. The Source of the B-Genome

The B-genome donor species is a puzzle for wheat cytogeneticists, and it is still an unsettled issue (Kerby and Kuspira, 1988). Based on geographical distribution, morphological features, karyotype analysis, Giemsa C- and N-banding patterns, meiotic chromosome pairing, nuclear DNA content, in situ hybridization studies, restriction fragment patterns of rRNA genes, DNA:DNA hybridization, seed protein banding pattern, and cytoplasmic compatibility and incompatibility studies, six diploid wild species have been proposed as the possible source of the B-genome (Table 8.11). All of the species, except Triticum urartu, belong to the Sitopsis section, are cross compatible, and exhibit almost normal meiosis and various degrees of pollen and seed sterility (Feldman, Strauss, and Vardi, 1979). By comparing the possible evidence reported so far, Kerby and Kuspira (1987, 1988) suggest, "Aegilops searsii is the source of the B-genome, if this set of chromosomes is monophyletic in origin." However, ctDNA reveals that the cytoplasm of emmer and common wheats originated from Aegilops longissima and not from Aegilops speltoides, Aegilops sharonensis, Aegilops bicorne, Aegilops searsii, or Triticum urartu (Tsunewaki and Ogihara, 1983). Meiotic chromosome pairing of Triticum aestivum involving Aegilops speltoides, Aegilops sharonensis, and Aegilops longissima revealed that none of these three Aegilops are the B-genome donor of the wheat (Fernández-Calvín and Orellana, 1994).

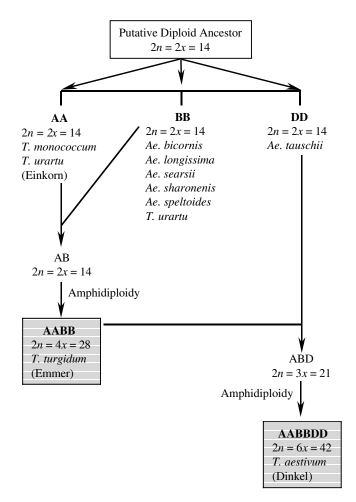


Figure 8.7 The origin of hexaploid wheat.

Table 8.11 Species, Genome Symbols, and Authority Reported Genomes of Hexaploid Wheat

Species	Genome $n = 7$	Authority
Triticum monococcum	А	Kihara, 1924
Aegilops speltoides	B?	Pathak, 1940; Sarkar and Stebbins, 1956; Daud and Gustafson, 1996
Aegilops longissima	B?	Tanaka, 1956
Aegilops sheronensis	B?	Kushnir and Halloran, 1981
Aegilpos bicornis	B?	Sears, 1956a
Aegilops searsii	B?	Feldman and Kislev, 1977
T. urartu	B?ª	Johnson, 1975
Aegilops tauschii	D	Pathak, 1940; Kihara, 1944; McFadden and Sears, 1944

^a *T. urartu* = *T. monococcum*.

3. The Source of the D-Genome

Pathak (1940), on the basis of morphological traits, geographical distribution, karyotype analysis, and susceptibility to rust, proposed that *Aegilops squarrosa* (*A. tauschii*), a diploid wild species, was the donor of the D-genome of hexaploid wheat. This was verified experimentally by Kihara (1944, 1947) and McFadden and Sears (1944, 1946) independently. McFadden and Sears (1944,

1946) synthesized allohexaploids by doubling the chromosomes of F_1 triploid hybrids of *T. turgidum* (*Aegilops dicoccoides*) × *A. tauschii*. The synthesized hexaploid plants resembled *Triticum aestivum* for several morphological traits, formed 21 bivalents, and were fertile. Furthermore, hybrids between synthetic hexaploid wheat and cultivated hexaploid wheat also showed 21 bivalents and showed 96.3% seed set. This suggests that synthesized and cultivated wheats have similar genomes, and hence, *A. tauschii* is the D-genome donor species.

The genomic affinities among species established on the basis of the chromosome pairing method is valid, as has been shown from several classical examples, and is being verified by biochemical and molecular approaches. It is suggested that classical taxonomy is a foundation for genome analysis and should not be disturbed.

VII. GENOME ANALYSIS BY MOLECULAR METHODS

During the past decade, literature on genomic relationships (plant phylogenetic relationships) has been dominated by molecular data, including nuclear (RFLP, AFLP, RAPD, SSR, sequence variation in the gene such as ITS regin of rDNA), extranuclear (chloroplast and mitochondrial DNA) DNA variation, and genomic *in situ* hybridization (GISH) by multicolor FISH. This latter approach is extremely powerful, where production of interspecific or intergeneric hybrids is not feasible by conventional methods (Heslop-Harrison, 2000).

Molecular methods so far have verified the genomic relationships established by cytogenetics in the genus Arachis, Avena, Glycine, Gossypium, Hordeum, Lycopersicon, Oryza, Phaseolus, Pisum, Secale, Vigna, Zea, Triticum, and many more. The intention here is to use Glycine as an example, where classical taxonomy is clear, genomic relationships based on cytogenetics, molecular methods, and chloroplast DNA have been exceedingly examined.

A. Genome Analysis by Nucleotide Sequence Variation

Kollipara, Singh, and Hymowitz (1997) determined phylogenetic relationships among 18 species of the genus *Glycine* from nucleotide sequence variation in the internal transcribed spacer (ITS) region of nuclear ribosomal DNA (Figure 8.8). Of a total of 648 characters used in the analyses, 16, 215, 168, 199, and 50 belonged to the 5S, ITS1, 5.8S, ITS2, and LS sequence, respectively. The mean length of ITS1 was 16.4 nucleotides longer than that of the ITS2 sequence. The alignment assumed a total of 18 indels, excluding the ambiguous regions. Only nine of these 18 indels were phylogenetically informative. The length of the gaps ranged from one (in most cases) to five (indel 5) nucleotides.

The sequence divergences obtained from the pairwise comparisons of unambiguous characters ranged from 0.2% between *Glycine max* and *Glycine soja* to 8.6% between *Glycine hirticaulis* and *Glycine falcata*. The highest divergence was recorded between *Glycine hirticaulis* and *Glycine*

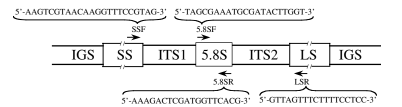


Figure 8.8 The general structure of the nrDNA showing the PCR and sequencing strategy. The relative positions of the small subunit (SS), large subunit (LS), 5.8S, internal transcribed spacers (ITS1 and ITS2), the intergenic spacer (IGS), and the primers (with sequences) are indicated. (Redrawn from Kollipara, K.P., Singh, R.J., and Hymowitz, T., *Genome*, 40, 57–68, 1997. With permission.)

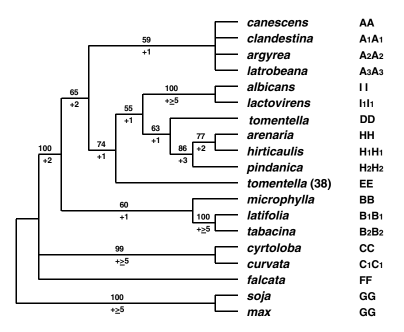


Figure 8.9 Strict consensus of the 16 most parsimonious trees with equal length of 176 steps obtained from equally weighted parsimony analysis, using the Branch-and-Bound option of the PAUP program, of the entire unambiguously aligned ITS region (CI excluding the uninformative characters = 0.632; RI = 0.735). The numbers above the branches indicate the number of times a monophyletic group occurred in 100 bootstrap replicates. The numbers shown below the branches indicate the additional number of steps over 176 (the total number of steps in the shortest tree) needed to collapse that branch (decay values). Decay analysis with tree length ≥5 steps longer than the most parsimonious trees could not be performed owing to the computational constraints. The taxa and the genome symbols are indicated; tomentella (38) indicates *Glycine tomentella* with 2*n* = 38. (From Kollipara, K.P., Singh, R.J., and Hymowitz, T., *Genome*, 40, 57–68, 1997. With permission.)

falcata, both in the ITS1 (24.9%) and ITS2 (10.3%) sequences. *Glycine falcata* showed maximum divergence (2.4%) from *Glycine lactovirens*, *Glycine cyrtoloba*, and *Glycine curvata* in its 5.8S sequence. *Glycine soja* and *Glycine max* differed by one nucleotide in the entire sequence.

Analysis of the entire sequence, excluding the ambiguous sequence, resulted in 16 maximally parsimoneous tree with equal length of 176 steps. The strict consensus trees with bootstrap and decay values are shown in Figure 8.9. The CI (consistency index) of these trees was 0.632 when uninformative characters were excluded. The RI (retention index) of these most parsimonious trees was 0.735.

The *Glycine* species with the same letter genome symbols, assigned based on cytogenetics, were resolved as monophyletic groups on the strict consensus tree (Figure 8.9). The newly described species (Tindale, 1986; Tindale and Craven, 1988; 1993), *Glycine albicans, Glycine arenaria, Glycine hirticaulis, Glycine lactovirens,* and *Glycine pindanica* formed two distinct clades. The clade containing *Glycine arenaria, Glycine hirticaulis,* and *Glycine pindanica,* supported by a bootstrap value of 86 and decay value of +3, was assigned an H-genome. The other clade consisting of *Glycine albicans* and *Glycine lactovirens* supported by bootstrap value of 100 and decay value of $+ \ge 5$ was assigned I-genome. The evolutionary tree estimated by the maximum likelihood method was essentially similar in its topology to the maximally parsimonius trees. The only difference was with respect to the placement of *Glycine falcata* in relation to the outgroup taxa and the C-genome clade.

The ITS region (nrDNA) is a multigene family. However, in the soybean, the nrDNA is mapped to a single locus on the short arm of Chromosome 13 based on the location of the

nucleolus organizer region by pachytene chromosome analysis (Singh and Hymowitz, 1988) and also by fluorescent *in situ* hybridization using ITS as a probe (Singh, Kim, and Hymowitz, 2001). The wild perennial *Glycine* species also contain one pair of NOR chromosome, like those in the soybean, except for *Glycine curvata* and *Glycine cyrtoloba*, which have two NOR chromosomes (Singh, Kim, and Hymowitz, 2001). The ITS region appeared highly homogenous from this study, because direct sequencing of the PCR fragments did not show polymorphisms in the nucleotide sequences.

Comparison of strict consensus trees based on sequences revealed that the ITS1 and ITS2 sequences complemented each other in resolving the major clades, the A- and B-genome species, *Glycine microphylla, Glycine latifolia*, and *Glycine tabacina*, as a sister group to the clade with A-, D-, H-, and I-genome species, while the ITS2 resolved the A-genome species, *Glycine argyrea, Glycine canescens, Glycine clandestina*, and *Glycine latrobeana*, from the clade containing B-, D-, E-, H-, and I-genome species. The phylogenetic tree was found to be more robust owing to the complimentary nature of both regions.

The robustness of a hypothesis can be tested by assessing its congruence with phylogenetic hypothesis developed from different methods, like morphological, geographical, biochemical, molecular, etc., and by it yielding the identical trees. In the genus *Glycine*, extensive cytogenetic and limited biochemical investigations supported by classical taxonomic data contributed to the establishment of intergenomic relationships among 13 of the 20 species. The phylogeny established from molecular data helped assign genome symbols: H to *Glycine arenaria*, H₁ to *Glycine hirticaulis*, H₂ to *Glycine pindanica*, I to *Glycine albicans*, and I₂ to *Glycine lactovirens*. The H- and I-genome groups were strongly supported by the higher bootstrap and decay values, suggesting the close relationship among the species within these clades (Figure 8.9). However, cpDNA phylogeny was found to be congruent with the inference based on cytogenetic studies with respect to B-, C-, and G-genome species, but the remaining species were classified as an A-plastome group (Doyle et al., 2000). It was concluded that cpDNA is not the perfect phylogenetic tool for establishing genomic relationships among species of the subgenus *Glycine* as a whole, as plastomes of species within groups are poorly differentiated. Its utility is often limited at lower taxonomic levels among closely related taxa (Doyle, 1991; Doyle et al., 1998).

A multidisciplinary approach (cytogenetics and ITS of rDNA) was applied by Singh, Kollipara, and Hymowitz (1998a) to uncover the genomic diversity in *Glycine canescens* and diploid *Glycine tomentella* of Western Australia. Cytogenetic results demonstrated that *Glycine canescens* from Western Australia are genomically similar, however, differ by a paracentric inversion from the standard *Glycine canescens* from South Australia. On the other hand, 40-chromosome *Glycine tomentella* accessions, morphologically similar, from Western Australia are highly diverse genomically. Cytogenetics and ITS investigations supported the four isozyme groupings of Doyle and Brown (1985). This demonstrates that diploid *Glycine tomentella* of Western Australia is complex, and from an evolutionary viewpoint, is actively radiating out into several genomic variants.

B. Genome Analysis by Multicolor FISH

Genome analysis in polyploid plants can be determined by multicolor genomic *in situ* hybridization (Heslop-Harrison and Schwarzacher, 1996; Mukai, 1996; Shishido et al., 1998). Mukai (1996) described labeling-detection combinations to discriminate genomes of polyploid *Aegilops* and *Triticum* (Table 8.12). Multicolor FISH expresses genomes much clearer in synthetic amphidiploids than observed in the natural amphidiploids. This may be attributed to chromosomal interchanges among different genomes during evolution. Although the multicolor FISH technique is a powerful cytogenetic tool to determine the genome of polyploid crops, this technique has some limitations. If more than three probes, FITC (green), rhodamine (red), and AMCA (blue), are applied, chromosomes cannot be counterstained by propidium iodide (red) or DAPI (blue).

Table 8.12 Labeli	Table 8.12 Labeling-Detection Combinations in Multicolor FISH Using Total Genomic DNA Probes	is in Multicolor FISH U	sing Total Genomi	c DNA Probes
	Labeling of Probe	Detection	on	
	DNA	Reagent	Fluorescence	Application
			Tetraploid	
<i>Method 1</i> First genome Second genome	Biotin-dUTP Digoxigenin-dUTP	Avidin-FITC Anti-digrhodamine	Yellow Orange	<i>Aegilops</i> (CCDD, CCUU, DDNN), Synthetic amphidiploid (AASS)
<i>Method 2</i> First genome Second genome	Biotin-dUTP Cold (blocking)	Avidin-FITC Propidium iodide	Yellow Red	Tetraploid wheat (AABB, AAGG), <i>Aegilops</i> (DDMM, UUMM, UUSS)
Method 3				
First genome Second genome	Biotin-dUTP None	Avidin-FITC Propidium iodide	Yellow Red	Millium montianum Tobacco
			Hexaploid	
First genome Second genome Third genome	Biotin-dUTP Digoxigenin-dUTP Cold (blocking)	Avidin-FITC Anti-digrhodamine	Yellow Orange Brown	Common wheat Synthetic amphidiploid (AABBNN, AAGGUU)
			Octoploid	
First genome Second genome	Biotin-dUTP Digoxigenin-dUTP	Avidin-FITC Anti-digrhodamine	Yellow Orange	Synthetic amphidiploid (AABBDDEE, AABBDDRR)
Fourth genome	Cold (blocking) 2/3 bio. + 1/3 dig.	Avidin-FITC + Anti- digrhodamine	Yellow	
Courses Erom Mult	Course: Erom Multai V Mathada in Ganama	a Analysia in Dlanta DD Jaylar Ed		

Source: From Mukai, Y., Methods in Genome Analysis in Plants, P.P. Jauhar, Ed., CRC Press LLC, 1996, 181–192. With permission.

CHAPTER 9

Chromosomal Aberrations in Cell and Tissue Culture

I. INTRODUCTION

Chromosomal aberrations are a common occurrence in cell and tissue culture-derived calluses and their regenerants. In general, numerical and structural changes in chromosomes are attributed to spindle failure that causes endoreduplication, c-mitosis, nuclear fragmentation (amitosis), multipolar configurations, and lagging chromosomes. These changes are induced by media composition; age of the callus; nature of the callus, morphogenic versus nonmorphogenic; genetic background of the explants; and kinds of media, solid versus liquid (Bayliss, 1973, 1980; Evans and Reed, 1981; Wersuhn and Dathe, 1983; D'Amato, 1985; Ogura, 1990; Greier, 1991).

II. CHROMOSOMAL ABERRATIONS IN CALLUS

A. Media Composition

Chromosomal aberrations induced in cultures are the result of direct influence of chemical substances present in the medium. Medium compositions that can produce more organized and meristematic states cause less somaclonal variation (Choi, Lemaux, and Cho, 2000a,b; Bregitzer et al., 2002). However, the majority of reports agree that chromosome aberrations are generated during culture by growth hormones such as 2,4-D (Venketeswaran, 1963; Shimada and Tabata, 1967; Torrey, 1959, 1967; Mohandas and Grant, 1972; Nuti Ronchi, Martini, and Buiatti, 1976; Kar and Sen, 1985; Ziauddin and Kasha, 1990), IAA (Huskins and Steinitz, 1948; Naylor, Sander, and Skoog, 1954; Nishiyama and Taira, 1966), NAA (Dermen, 1941; Berger and Witkus, 1948; Zhang et al., 1987), and kinetin (Torrey, 1959; Van't Hof and McMillan, 1969; Nuti Ronchi, Martini, and Buiatti, 1976). The medium containing yeast extract and coconut water is also responsible for chromosomal aberrations (Straus, 1954; Inomata, 1982; Kar and Sen, 1985). Yeast extract contains many substances, including nucleic acid, that cause chromosome stickiness and breakage (Kodani, 1948).

Venketeswaran (1963) examined cytologically the cell aggregates of *Vicia faba* (2n = 2x = 12) grown in a medium containing 2,4-D and observed different ploidy levels with a predominance of aneuploids. Naylor, Sander, and Skoog (1954) reported that 2 mg/L IAA induced mitosis without cytokinesis in tobacco pith tissues grown in a modified White's agar medium that resulted in multinucleate cells and an abnormally lobed nucleus. Dermen (1941) observed 4x, 8x, and 16x cells in bean (2n = 2x = 22) by NAA treatment and suggested that a mixture of diploid and polyploid tissues may be induced in some parts of treated plants.

Van't Hof and McMillan (1969) studied the pea root (2n = 2x = 14) callus and observed that the callus was smooth and spherical in shape when kinetin was added to the medium, while it was rough and irregular in medium without kinetin. Chromosome counts indicated that in medium with kinetin, the mitotic index in polyploid cells increased rapidly during 0 to 7 days in culture (mixed-cell populations), and a slow phase occurred from 7 to 14 days (consisting of approximately 50% 2x and 50% 4x cells). In contrast, they observed only diploid cells in callus grown in medium without kinetin. Matthysse and Torrey (1967a,b) recorded that diploid cells do not require kinetin or kinetinlike substances. Zhang et al. (1987) recorded variable frequencies of haploids (2n = x = 6), tetraploids (2n = 4x = 24), and breakage in chromosomes in the cotyledon attachment region in induced calluses of *Vicia faba* (2n = 2x = 12), cultured in a medium containing various concentrations of NAA and kinetin. In *Haplopappus gracilis*, the increase in frequency of polyploid cells was associated with a decrease in supply of nutrient (Singh and Harvey, 1975b; Singh et al., 1975).

Chromosomal aberrations induced in culture are dependent on the kind of growth hormone used. Published results suggest that 2,4-D induces greater amounts of polyploidy than NAA (Jacobsen, 1981; Jha and Roy, 1982; Kar and Sen, 1985). Libbenga and Torrey (1973) recorded in pea cortical nuclei that both auxin and cytokinin were needed to induce chromosome doubling.

Investigations by Singh and Harvey (1975a) and Bayliss (1975) contradict the above results. Their results showed that the higher the concentration of 2,4-D in the culture medium, the lower the frequency of mitotic irregularities.

Preferential selection of haploid cells from mixed populations of cells of varying ploidy levels has been demonstrated cytologically by Gupta and Carlson (1972), following parafluorophenylalanine (PFP) treatment. They used pith calluses of tobacco (2n = 48) that consisted of haploid and diploid cells. The growth of diploid cells was inhibited at the level of 9.0 g/mL PFP, and diploid cells turned black and died at 15.0 g/mL of PFP. In contrast, growth of haploid cells was not affected.

Do chromosome aberrations arise during culture or do they pre-exist in the cultured tissues? To answer this question, Singh (1986) analyzed chromosomes of five calluses, each after 1, 2, 3, 6, 7, 8, 10, and 15 days of inoculation of barley immature embryos, although embryos were not callused in the beginning. It is evident from Table 9.1 that chromosome aberrations, mainly tetraploid, initiated in cultured embryos after 6 days. After 8 days, all five cultured embryos showed low mitotic index. The interphase nuclei were enlarged, with condensed chromatin (endoreduplication). Tetraploid cells began to increase gradually after 10 days. These observations suggest that chromosome aberrations are induced during the culture (Cooper et al., 1964; Shimada and Tabata, 1967; Singh and Harvey, 1975b; Balzan, 1978; Armstrong, Nakamura, and Keller, 1983; Sree Ramulu et al., 1984b; Natali and Cavallini, 1987; Pijnacker et al., 1989). However,

Days after		2 <i>n</i> Chro	mosome	Number		Tot	al	
Embryo Cultured	14	14 + 1telo	15	28	13 + 2telo	Fragments	Cells Studied	% Diploid
1	182	_	_	_	_	_	182	100.0
2	130	_	_	_	_	_	130	100.0
3	248	_	_	_	_	_	248	100.0
6	202	_	1	1	_	_	204	99.0
7	255	1	_	2	_	_	258	98.8
8	50	_	_	3	1	_	54ª	92.6
10	372		_	9	2	2	385	96.6
15	149	_	_	11	_	_	160	93.1

Table 9.1 Chromosome Count in Immature Embryo-Derived Callus of Barley cv. Himalaya after 1–15 d in Culture

^a Low mitotic index, condensation of chromatin in the interphase cells.

Source: From Singh, R.J., Theor. Appl. Genet., 72, 710-716, 1986. With permission.

Period (Year)	Number of Cells Examined	Chromosome Numbers ^a
0 ^b	53	2 <i>n</i> = 48 (25), 96 (28)
0°	44	2n = 48 (4), 96 (31), 192 (7), aneuploids (2); 182, 184
1	15	2n = 96 (4), 192 (3), aneuploids (8): 54, 74, 86, 88, 92, 156, 158, 304
6	12	2n = aneuploids (12): 108, 122, 124, 140, 146, 148 (2), 152 (2), 154, 160, 174

 Table 9.2
 Chromosome Numbers in Tobacco Pith and Calluses

^a (0) denotes number of cells. Normal chromosome (2n = 48).

^b Pith freshly excised from region of the stem 3.5-10.5 cm below apex.

° Pith freshly excised from region of the stem 15.5-22.5 cm below apex.

Source: From Murashige, T. and Nakano, R., Am. J. Bot., 54, 963-970, 1967. With permission.

variable chromosome numbers in root tips of *Allium cepa* 2n = 2x = 16 (Partanen, 1963), and *Pisum sativum*, 2n = 2x = 14 (Torrey, 1959), pith tissues of *Nicotiana tabacum*, 2n = 4x = 48 (Shimada and Tabata, 1967; Murashige and Nakano, 1967) (Table 9.2), and hypocotyl tissues of *Haplopappus gracilis*, 2n = 2x = 4 (Singh and Harvey, 1975b) suggest that chromosome aberrations pre-exist. If these tissues are grown *in vitro*, polyploid and aneuploid cells are expected to be generated more frequently than tissues lacking initial chromosome abnormalities.

Enhanced tolerance to NaCl in tobacco is attributed to chromosome number. Kononowicz et al. (1990a) found cells adapted to 428 mM NaCl were predominantly hexaploid (2n = 6x = 72), while unadapted cells contained normal (2n = 4x = 48) chromosome numbers. Enrichment of the cell population for hexaploid cells occurred only after exposure to higher NaCl (428 mM) or at a lower level of NaCl (171 mM). Hexaploidy induced by higher NaCl bred true for at least 25 cell generations after being removed from NaCl exposure. The hexaploidy could not be correlated with several phenotypic changes associated with plants regenerated from adapted cells, including male sterility and increased salt tolerance (Kononowicz et al., 1990b).

B. Age of Callus

Generally, the older calluses lose the capacity to regenerate plants, because they carry high ploidy in the tissues or accumulate an increasing number of aneuploid cells that lead to the loss of balanced chromosomal constitution of the cells (Murashige and Nakano, 1965, 1967; Yazawa, 1967; Torrey, 1967; Shimada and Tabata, 1967; Heinz, Mee, and Nickell, 1969; Shimada, 1971; Novák, 1974; Ogihara and Tsunewaki, 1979; Roy, 1980; Orton, 1985; Kar and Sen, 1985; Lavania and Srivastava, 1988).

Murashige and Nakano (1965, 1967) observed the complete loss of morphogenic potentialities in 1- to 6-year-old-calluses of tobacco (*Nicotiana tabacum* cv. Wisconsin), and that was associated with increased polyploidy $(2x \rightarrow 4x \rightarrow 8x)$ and highly variable aneuploidy (Table 9.2). Similarly, Shimada (1971) found only aneuploid cells in 2-year-old-calluses of tobacco, though only ten cells were studied, while 64% of the cells carried expected 2n = 48 chromosomes in 3- to 7-month-old-calluses.

Novák (1974) studied changes of karyotype in 92-, 157-, 254-, and 339-day-old leaf calluses of *Allium sativum* (2n = 2x = 16). Older calluses showed high frequencies of polyploid and aneuploid cells, and chromosomes ranged from haploid to hypertetraploid (Table 9.3). Chromatin bridges, laggards, and chromosome fragments at anaphase were also recorded.

Twelve-month-old calluses derived from immature petioles of celery carried normal chromosome complements in 2.5% of the cells and completely lost the plant regenerability trait. However, 6-month-old calluses showed the normal karyotype of 2n = 2x = 22 in 84% of cells, and plant regenerability was normal (Orton, 1985). In contrast, aging did not influence chromosomal aberrations in maize (Gresshoff and Doy, 1973; Edallo et al., 1981; McCoy and Phillips, 1982). McCoy and Phillips (1982) observed a high frequency of diploid (2n = 2x = 20) cells in 4-month-old (96 to 97%) and 8-month-old (94 to 95%) immature embryo derived calluses of *Zea mays*. As expected, these calluses were highly morphogenic.

		Age of Cu	lture (Days)	
Chromosomal Type	92	157	254	339
Haploid $(2n = x = 8)$	0.75	1.51	1.59	2.03
Hypodiploid (>8<16)	1.49	3.79	8.73	4.73
Diploid $(2n = 2x = 16)$	53.73	46.21	42.86	26.35
Hyperdiploid (>16<32)	20.89	6.82	9.52	12.16
Tetraploid $(2n = 4x = 32)$	14.18	34.85	33.33	45.27
Hypertetraploid (>32)	8.96	6.82	3.97	9.46

Table 9.3 Frequency (%) of Chromosomal Aberrations in AS-1 Strain of *Allium* sativum (2n = 16) Callus as Related to the Length in Culture

Source: From Novák, F.J., Caryologia, 27, 45–54, 1974. With permission.

C. Nature of Callus (Morphogenic vs. Nonmorphogenic)

The morphogenic ability of a callus depends upon the chromosome constitution of the cells. Balanced chromosome number in a callus is a prerequisite for regenerating plants. This is in agreement with the results reported in *Daucus carota* (Smith and Street, 1974), *Hordeum vulgare* (Scheunert, Shamina, and Koblitz, 1978; Singh, 1986; Gaponenko et al., 1988), *Haworthia setata* (Ogihara and Tsunewaki, 1979), *Nicotiana tabacum* (Murashige and Nakano, 1965, 1967; Mahfouz, De Boucaud, and Gaultier, 1983; Wersuhn and Sell, 1988), *Pisum sativum* (Torrey, 1959, 1967), *Triticum aestivum* (Ahloowalia, 1982), and *Zea mays* (Balzan, 1978).

Singh (1986) examined cytologically ten morphogenic calluses of barley (Table 9.4). The expected diploid (2n = 2x = 14) chromosome cells predominated in all morphogenic calluses (Figure 9.1A). The percentage of diploid cells ranged from 74.1 to 100% (Table 9.4), however, haploid (2n = x = 7) (Figure 9.1B), tetraploid (2n = 4x = 28), octoploid (2n = 8x = 56) (Figure 9.1C), and triploid (2n = 3x = 21) cells were also recorded in low frequencies. The occurrence of these chromosomal types varied from callus to callus. The calluses with the higher number of tetraploid cells were not as morphogenic as those that showed largely diploid cells. In contrast, diploid cells were comparatively lower in nonmorphogenic calluses (Table 9.4). The majority of aneuploid cells also carried chromosome structural changes such as ring, acrocentric, dicentric, telocentric, and fragment chromosome(s) (Figure 9.2). Similar results have been recorded in several plant species (Torrey, 1967; Sacristán and Wendt-Gallitelli, 1971; Novák, 1974; Orton, 1980; Singh, 1981; Murata and Orton, 1984).

It is interesting to note that an euploid cell chromosome counts were around haploid, diploid, triploid, and tetraploid numbers. The uneven chromosome separation during cell division (Figure 9.3A) may have contributed to the origin of cells with haploid, triploid, and uncountable microchromosomes (Figure 9.3B). This supports the observations of Chen and Chen (1980), who suggested that triploid cells might have originated from 4x cells through reductional grouping of chromosomes, accompanied by multipolar formation. Thus, the loss of regenerability of a callus is attributed to an increased frequency of cells with polyploid and an euploid chromosome numbers.

Tissue culture may play an important role in restructuring the chromosome after interspecific and intergeneric hybridization. Once a desired chromosome combination is accomplished, efforts should be directed to bring the chromosome number to a genetic balance level by media modifications (Torrey, 1967; Evans, Sharp, and Medina-Filho, 1984). Multiplication of tetraploid cells in pea root callus was favored by the medium supplements yeast extract and kinetin, but by nutrient modifications, Torrey (1967) obtained uniformly diploid cells. Similarly, Evans, Sharp, and Medina-Filho (1984) established liquid cell cultures of the SU/SU genotype of *N. tabacum* (2n =48) that maintained stable chromosome numbers for 6 years in culture.

	Callus			2n Chro	2n Chromosomes Number	s Number				
Callus Types	Numbers	7	14	15	21	28	56	Others ^a	Total Cells	Diploid (%)
Morphogenic	-		190			I		I	190	100.0
	0	Ι	170	Ι	Ι	Ι	Ι	I	170	100.0
	ო	0	198	Ι	Ι	Ι	Ι	I	200	0.06
	4	Ι	200	Ι	Ι	0	I		202	0.06
	5 2	I	100	I	I	-	I		101	0.06
	9	I	118	I	I	e			121	97.5
	7	I	147	I	I	17	I	I	164	89.6
	80	Ι	100	Ι	2	10	2	I	114	87.7
	6	I	60	I	I	16	0		78	76.9
	10	I	117	I	ო	33	Ŋ	I	158	74.1
Nonmorphogenic	-	Ι	49	0	2	49	2	44	148	33.1
	0	I	10	-	ო	6	I	59	82	12.2
	3-7 ^b									
^a Others (number of cells observed are in parenthesis): 13 + 2 telocentrics (5), 14 + 1 telocentrics (7), 14 + 2 metacentrics (2), 18 (2), 19 (5), 19 + 1 acrocentric (1) 21 + 1 telocentric (3) 21 + 1 dicentric (1) 22 (6)	of cells observed	d are in pa telocentri	arenthesis)	: 13 + 2 te	locentrics	(5), 14 + 1	telocentri	cs (7), 14 +	2 metacentrics	(2), 18 (2), 19

Himalaya
of Barley cv.
Calluses of
Embryo-Derived
Immature E
Analysis in
Chromosome
Table 9.4

22 + 1 ring (1), 22 + 2 dicentrics (4), 23 (1), 24 (6), 24 + 1 fragment (5), 24 + 1 acrocentric (1), 25 (2), 25 + 1 telocentric (3), 27 (4), 27 + 1 telocentric (1), 7 + 29 telocentrics (1), fragment chromosomes (3), clumped chromosomes (14), uneven cell division (5), and high uncountable ploidy chromosome number (5).
 ^b Lacked mitotic cell division.

Source: From Singh, R.J., Theor. Appl. Genet., 72, 710–716, 1986. With permission.

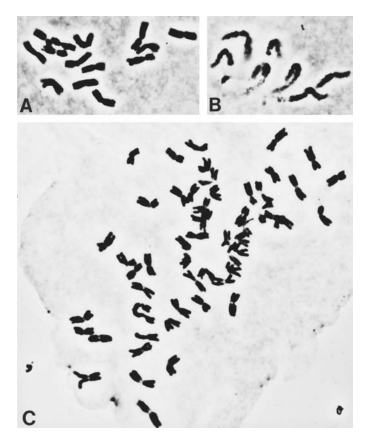


Figure 9.1 Cells observed in morphogenic callus of barley. (A) 2n = 14 (From Singh, R.J., unpublished results.). (B) 2n = 7; and (C) 2n = 56. (From Singh, R.J., *Theor. Appl. Genet.*, 72, 710–716, 1986. With permission.)



Figure 9.2 A cell observed in nonmorphogenic callus of barley showing seven complete + 29 telocentric chromosomes. (From Singh, R.J., *Theor. Appl. Genet.*, 72, 710–716, 1986. With permission.)

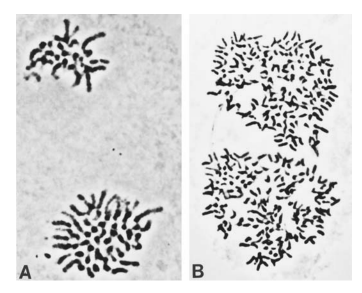


Figure 9.3 Cells observed in nonmorphogenic callus of barley. (A) Telophase with unequal chromosome on each pole. (B) Two daughter nuclei with numerous microchromosomes. (From Singh, R.J., *Theor. Appl. Genet.*, 72, 710–716, 1986. With permission.)

D. Genetic Background of the Explants

Chromosomal instability induced in the culture and plant regenerability are often influenced by the genotype of the explants (Murashige and Nakano, 1967; Heinz, Mee, and Nickell, 1969; Kao et al., 1970; Sacristán, 1971; Okamoto et al., 1973; Asami et al., 1975; Novák, Ohnoutková, and Kubaláková, 1978; Scheunert, Shamina, and Koblitz, 1978; Jacobsen, 1981; Browers and Orton, 1982a,b; Wersuhn and Dathe, 1983; Bajaj and Gill, 1985; Sree Ramulu et al., 1985; Rhodes, Phillips, and Green, 1986; Lee and Phillips, 1987; Pijnacker and Sree Ramulu, 1990; Ohkoshi et al., 1991; Ruíz et al., 1992). Kao et al. (1970) analyzed the chromosomes of the suspension cultures of *Triticum monococcum* (2n = 2x = 14), *T. aestivum* (2n = 6x = 42), *Glycine max* (2n = 2x = 40), *Melilotus alba* (2n = 2x = 16), and *Haplopappus gracilis* (2n = 2x = 4). All the cultures except *H. gracilis* carried chromosome aberrations.

Asami et al. (1975) examined chromosomes in 6-month-old calluses of four an euploid stocks (nulli-5B, tetra-5D, ditelo 5A^L, ditelo 5B^L, ditelo 5D^L) and a disomic control (2n = 6x = 42) of *Triticum aestivum* cv. Chinese Spring. The disomic control showed normal chromosome complements of 2n = 42 in 97% of the cells, and only one cell carried 2n = 40. By contrast, an euploid stocks showed a wide range of chromosome distribution, as shown below:

Stocks	Chromosome (Range)	Frequency of 2 <i>n</i> = 42 (%)
	(0)	
Disomic	40-42	97
Nulli-5B tetra-5D	24–86	22
Ditelo 5A ^L	27–126	52
Ditelo 5B [⊥]	28–85	58
Ditelo 5D ^L	35–84	68

Browers and Orton (1982a) recorded a clear-cut genotypic response on chromosome stability in a 9-week-old suspension culture of celery (2n = 2x = 22) Apium graveolens var. rapeceum. Keeping the culture conditions unchanged, PI 169001 contained 80% diploid and 20% nondiploid cells and was highly morphogenic. In contrast, suspension culture of PI 171500 was 31% diploid and 69% nondiploid and failed to regenerate plants. Sree Ramulu et al. (1985) studied the chromosomes in calluses and suspension cultures of monohaploid (2n = x = 12), dihaploid (2n = 2x = 24), and tetraploid (2n = 4x = 48) potatoes (*Solanum tuberosum*) after 7 and 14 days of culture. Polyploidization ($1x \rightarrow 2x \rightarrow 4x \rightarrow 8x$) was more rapid in monohaploid than in dihaploid and tetraploid potatoes. After 14 days of culture, chromosome numbers ranged from 12 to 96 for monohaploid, 24 to 48 for dihaploid, and 45 to 200 for tetraploid. The occurrence of aneuploid cells between two polyploid levels (2x and 4x; 4x and 8x) could be attributed to chromosome elimination or addition to the nearest ploidy level due to unequal chromosome separation and chromosome fragmentations, because aneuploid cells are physiologically and genetically more unbalanced than the cells with the exact multiple of basic chromosomes.

E. Kind of Media (Solid vs. Liquid)

The extent of chromosomal aberrations generated in solid (agar) and liquid (suspension) media is a controversial issue. Mitra, Mapes, and Steward (1960) examined chromosomes of carrot (2n = 2x = 18) cells cultured in liquid and solid media. Liquid culture showed haploid (2n = x = 9), diploid (2n = 2x = 18), tetraploid (2n = 4x = 36), and octoploid (2n = 8x = 72) cells, while diploid cells were recorded in the solid medium. Demoise and Partanen (1969), working with garden peony (*Paeonia suffruticosa*), 2n = 2x = 10, suggested that liquid medium favors higher polyploids than solid medium, though results were not conclusive. Singh and Harvey (1975a) studied the chromosome constituents of solid and liquid cultures of *H. gracilis* (2n = 2x = 4). Polyploid cells decreased (28 days = $88\% \rightarrow 46$ days = $75.9\% \rightarrow 122$ days = $90\% \rightarrow 147$ days = $66.7\% \rightarrow 220$ days = 61.7%) in solid medium with time, whereas the frequency of polyploid cells increased (14 days = $56.9\% \rightarrow 94$ days = $59.2\% \rightarrow 175$ days = $84.7\% \rightarrow 258$ days = 92.9%) in liquid culture.

A failure to regenerate plants from suspension and protoplasts is attributed to the extensive chromosome damage generated in culture. Karp et al. (1987a) examined the chromosomes of suspension cells and protoplasts derived from embryogenic callus of two wheat cultivars. Chromosome numbers ranged from 20 to 100, and the majority of cells carried numbers between 30 and 39. Chromosome aberrations such as deletions, dicentrics, fragments, telocentrics, and megachromosomes were recorded. These chromosome aberrations created cells with unbalanced chromosomes, making them physiologically and genetically unfit to regenerate normal plants. Stable karyotype was observed in bulk of *Brachycome dichromosomatica* (2n = 2x = 4) suspension culture, and only one cell line carried 2n = 5 chromosomes after 3 years (Nagl and Pfeifer, 1988).

III. MECHANISM OF CHROMOSOMAL ABERRATIONS IN CULTURE

It is expected that all the cells of somatic tissues of plants are genetically identical, and their chromosome constitutions are similar. However, it is not universally accepted. In pea roots and pith cells of tobacco, diploid and polyploid cells (polysomaty) have been observed in various frequencies, and it occurs during tissue maturation.

Cell division in somatic tissues follows regularly the path of DNA synthesis \rightarrow mitosis \rightarrow cytokinesis. When cytokinesis is delayed, cells fail to divide equationally, resulting in doubling somatic chromosomes. This is called endomitosis (endoreduplication). The majority of reports suggest that polyploidization in cultures is generated through endoreduplication (D'Amato, 1985).

The occurrence of tetraploid (2n = 4x = 28) and octoploid (2n = 8x = 56) cells in morphogenic and nonmorphogenic calluses in the barley suggests that polyploidization followed in a progressive fashion $(2x \rightarrow 4x \rightarrow 8x)$. This suggests that chromosome doubling occurred during the culture, because initially cultured embryos carried only diploid (2n = 2x = 14) cells. The observation of anaphase cells revealed that chromosomes separated during mitotic anaphase but failed to reach their respective poles. This may have been due to disturbed spindle formation (C-mitosis). Therefore, such cells will have a doubled chromosome number in the next mitotic division, and the cycle will keep on repeating (Singh, 1986). It is assumed that some media components act like colchicine (Levan, 1938) and inhibit formation of the spindle during cell division, causing chromosome doubling.

Unequal chromosome separation at mitotic telophase may be the cause of the occurrence of haploid cells. Haploid cells in appreciably low frequency have been recorded in *Daucus carota* (2n = 2x = 18) suspension cultures (Mitra, Mapes, and Steward, 1960), callus cultures of *Allium sativum* (2n = 2x = 16) (Novák, 1974), *Hordeum vulgare* (2n = 2x = 14) (Singh, 1986), *Triticum aestivum* (2n = 6x = 42) (Novák, Ohnoutková, and Kubaláková, 1978), and protoplast-derived calluses of *Solanum tuberosum* (2n = 4x = 48) (Sree Ramulu et al.,1984b). However, haploid cells may not compete with diploid or balanced polyploid cells during mitosis, and they will be eliminated eventually.

The quantitative trait loci (QTLs) controlling callus growth and subsequent shoot regeneration have been identified by 222 markers in doubled haploid (DH) lines derived from the cross between Steptoe × Morex barleys (Mano et al., 1996). Morex contributed two QTLs for callus growth (QcgI, Qcg 2), and Steptoe contributed three QTLs for shoot regeneration (Qsr I, Qsr 2, Qsr 3), while Morex contributed Qsr 4. In the multiple genic model, two QTLs for callus growth and four QTLs for shoot regeneration accounted for 47.8% and 49.8% of the total variation in the barley population, respectively. Chromosomal regions associated with green plant regeneration in anther culturederived wheat plants have been identified (Torp, Hansen, and Andersen, 2001). Four QTLs for 80% green plant were on 2AL and 2BL and one QTL explaining 31.5% of the genetic variation for green plant formation were detected on 5BL.

Chromosome breakage in culture is a nonrandom phenomenon that occurs preferentially in the heterochromatic region or at euchromatin/heterochromatin junctions (Michaelis and Rieger, 1958; Döbel, Schubert, and Rieger, 1978; Jørgensen and Anderson, 1989). Barley calluses produced high frequencies of telocentric, acrocentric, and fragment chromosomes (Singh, 1986). All the chromosomes in barley carry centromeric heterochromatin, and a few have them at intercalary regions (Singh and Tsuchiya, 1982b). The occurrence of acrocentric chromosomes suggests that breaks occur at interband regions and also at heterochromatic regions. That has been observed in the cultures of *Crepis capillaris* (Sacristán, 1971; Ashmore and Gould, 1981), maize (McCoy, Phillips, and Rines, 1982; Lee and Phillips, 1987; Benzion and Phillips, 1988; Fluminhan, de Aguiar-Perecin, and Dos santos, 1996), oat (Johnson, Phillips, and Rines, 1987a,b), wheat (Fedak, Armstrong, and Handyside, 1987), and wheat–rye hybrids (Armstrong, Nakamura, and Keller, 1983; Lapitan, Sears, and Gill, 1984).

Sacristán (1971) studied the extent of preferential chromosome aberrations in 3-month-old calluses of *Crepis capillaris* (2n = 2x = 6). The nucleolus organizer chromosome showed the highest (82.3%) chromosomal rearrangement followed by the shortest chromosome (64.6%). The longest chromosome showed the least (47.0). It was reasoned that in the case of the SAT-chromosome, the break region corresponds to a region of late DNA synthesis.

In this respect, the mode of breakage in chromosomes induced in culture is not different from that produced by x-ray treatment or any other mutagen (Khush and Rick, 1968a; Natarajan and Ahnström, 1969; Jancey and Walden, 1972; Gill et al., 1980; Inomata, 1982; Nuti Ronchi, Bonatti, and Turchi, 1986a; Nuti Ronchi et al., 1986b). Khush and Rick (1968a), working with tomato, found that about 60% of the breaks induced by x-ray treatment occurred in heterochromatin. Thus, late-replicating DNA in heterochromatic regions is more vulnerable to breakage than the euchromatic regions (Sacristán, 1971; Döbel, Schubert, and Rieger, 1978; McCoy, Phillips, and Rines, 1982; Armstrong, Nakamura, and Keller, 1983; Murata and Orton, 1984).

The decrease in the frequency of heritable variation with increasing exposure to 2,4-D in regenerated soybean plants expressed partial sterility, complete sterility, curled and wrinked leaves, dwarfism, chlorophyll deficiency, chlorophyll chimaera, indeterminate to determinate flowering, lack of unifoliates, yellow-edged cotyledons, and isozyme variants (Shoemaker et al., 1991). These mutants were not examined cytologically, and an inheritance study was not conducted.

IV. CHROMOSOMAL ABERRATIONS IN REGENERATED PLANTS

Phenotypic variability among cell and tissue culture-derived regenerants may be attributed to epigenetic, genetic, and chromosomal changes induced by the culture conditions (Evans and Reed, 1981; Sibi, 1984; Evans, Sharp, and Medina-Filho, 1984; D'Amato, 1985; Karp, 1986; Vasil, 1988; Stelly et al., 1989; Wersuhn, 1989; Oono, 1991; Skirvin et al., 2000). The culture-induced variants have been termed "Calliclones" (Skirvin and Janick, 1976; Skirvin, 1978), "Protoclones" (Shepard, Bidney, and Shahin, 1980), and a widely used term "Somaclones" (Larkin and Scowcroft, 1981). The frequency of somaclonal variation is at a higher rate (up to 10% per cycle of regeneration) than chemical- or radiation-induced mutation. This makes somaclonal variation a viable alternative to mutagenesis and a valuable tool for a plant breeder to introduce variation into breeding programs (Skirvin et al., 2000).

Epigenetic variations are due to the results of culture stress, and these variations are not transmitted from generation to generation. Thus, these changes are acquired traits and are not genetically controlled.

The genetic variations are induced during culture due to single nuclear gene mutations. The mutants exhibit Mendelian inheritance. A large number of plant species have been regenerated from cell and tissue cultures carrying somaclonal variation; the nature of mutation has been elucidated in only a few cases (Table 9.5). van den Bulk (1991) summarized application of cell and tissue culture for disease-resistance breeding. A total of 14 crops were listed. Inheritance was not determined in 19 studies. In banana, peach, potato, and sugarcane, disease resistance was transmitted after vegetative reproduction, and resistance was transmitted to the progeny in tomato, sugarcane, rice, rapeseed, maize, and celery. Moreover, heritable isozyme mutants have been recorded in the regenerants of broccoli (Orton and Browers, 1985), wheat (Ryan and Scowcroft, 1987), wild barley (Breiman et al., 1987), and blackberry (McPheeters and Skirvin, 2000). Dominant and recessive mutants isolated from culture are similar to those mutants obtained spontaneously in nature or induced by mutagens (Neuffer and Sheridan, 1980; Evans and Sharp, 1983; 1986; Gavazzi et al., 1987, Ullrich et al., 1991). Genetic variations for quantitative traits (maturity, height, lodging, seed yield, seed weight, protein and oil contents) were derived from tissue culture of the soybean, however, the magnitude of the genetic variation was relatively small (Hawbaker et al., 1993).

Crops	Mutants	Authority
	Recessive Mutants	
Oryza sativa	Early heading, albino, short culm, sterility	Fukui, 1983
	Dwarf mutants	Sun et al., 1983
Lactuca sativa	Seedling mutants	Engler and Grogan, 1984
Lycopersicon esculentum	Fruit color, male sterility, seedling mutants	Evans and Sharp, 1983
	Disease resistance	Evans, Sharp, and Medina Filho, 1984
	Chlorophyll mutants	Buiatti et al.,1985
Brassica napus	Yellow seed color	George and Rao, 1983
Medicago sativa	Flower color	Groose and Bingham, 1984
Zea mays	Shrunken kernel	McCoy and Phillips, 1982
Nicotiana sylvestris	Streptomycine sulfate resistant	Maliga, 1981
Glycine max	Male sterility	Graybosch, Edge, and Delannay, 1987
Sorghum bicolor	Male sterility	Elkonin, Gudova, and Ishin, 1994
	Dominant Mutants	
Nicotiana tabacum	Herbicide resistance	Chaleff and Parsons, 1978

Table 9.5 Recessive and Dominant Gene Mutations Observed in Regenerated Plants

A majority of morphological variants observed among the regenerated plants are due to numerical (aneuploidy, polyploidy) and structural (deletions, duplications, interchanges, inversions) chromosome changes induced during the culture. Generally, a high frequency of regenerants from diploid species carries normal chromosome complements. On the other hand, regenerants from polyploid species such as sugarcane, wheat, oat, triticale, potato, and tobacco have a comparatively higher frequency of plants with aberrant chromosome numbers. This is due to the fact that polyploid species can tolerate, to a greater extent than true diploid species, aneuploidy, because of the buffering capacity of the polyploid condition.

Despite many potential uses claimed for somaclonal variation, and substantial efforts by scores of individuals, the fact remains that thus far, there is not a single example of any significantly important new variety of any major crop species developed as a result of somaclonal variation (Vasil, 1990). It may not be true for the horticultural crops where tissue culture-derived somaclones were released as cultivars (Skirvin, McPheeters, and Norton, 1994).

A. Diploid vs. Polyploid Species

Chromosomal variations among regenerated plants of several diploid species are shown in Table 9.6. It is clearly seen that a high proportion of regenerated plants carried normal chromosome complements. Gould (1979) observed no chromosomal and phenotypically aberrant *Brachycome dichromosomatica* (2n = 4) regenerated plants from a year-old culture; this suggests again that diploid species are highly stable in culture. The exceptions are *Pisum sativum* and *Brassica oleracea*. In *Pisum sativum*, nine of the 20 regenerants were mixoploid, and in *Brassica oleracea* (2n = 18), of the 71 regenerants from stem pith explants, six plants were diploid, 54 plants tetraploid, and 11 octoploid. The occurrence of polyploid plants may be attributed to the source of explants. Pith cells carry diploid and polyploid cells (polysomaty) (Murashige and Nakano, 1967). This suggests that during shoot and root morphogenesis in diploid species, cells with unbalanced chromosome numbers cannot compete with balanced diploid cells in cell division. This enhances the regeneration of a large number of plants with normal diploid chromosome complements.

In contrast, morphogenic capability in polyploid species is not influenced by an euploidy, polyploidy, and chromosome structural changes. A few polyploid species such as potato (2n = 4x = 48), wheat (2n = 6x = 42), and triticale (2n = 6x = 42) will be cited here as examples.

1. Potato (Solanum tuberosum)

Cultivated potato is vegetatively propagated as an autotetraploid crop. Several authors have regenerated plants by mesophyll protoplasts and shoot tip cultures of potato and have recorded a considerable amount of genetic and chromosomal variability (Table 9.7).

Shepard, Bidney, and Shahin (1980) isolated several morphological variants among regenerants from leaf mesophyll protoplasts of the potato cultivar Russet Burbank. The variants showed compact growth habit, an earliness for tuber set, smooth and white skin, high yield, a requirement for less light (13 h) to flower than the parent (16 h), and resistance to early and late blight. The inheritance of these important traits was not determined. Gill, Kam-Morgan, and Shepard (1987) studied these variants cytologically at pachynema of meiosis. All the variants carried the expected 2n = 4x = 48 chromosomes. The high yielding variant did not carry chromosomal aberrations, but in the remaining mutants, the phenotypic alterations were due to interchanges, deletions, inversions, and duplications.

It is evident from Table 9.7 that tetraploid potato regenerants can tolerate a high number of chromosomes; for example, Gill, Kam-Morgan, and Shepard (1986) identified 27.8% of the regenerated plants with 2n = 72-96 chromosomes; Creissen and Karp (1985) found 21.5% plants with 2n = 73-96 chromosomes. Furthermore, the addition and deletion of one or two chromosomes around a mode of 2n = 48 showed little morphological change, or sometimes plants were indistin-

Decies $2n$ $2n=2x$ $2n=4x$ $2n=8x$ Mixoploid Aneuploids veolens 22 44 0 0 0 0 2 veolens 22 44 0 0 0 2 2 veolens 14 42 0 0 0 2 2 vulgare 14 52 1 0 0 0 0 0 itifiorum 14 52 0 <				Frequen	Frequency of Regenerants with	ants with			
veolens 22 44 0 0 0 2 1 $uulgare$ 14 42 0 0 0 0 2 1 $uulgare$ 14 42 0 0 0 0 0 1 $uulgare$ 14 52 0 0 0 0 1 $uitiforum$ 14 52 0 0 0 0 1 $uulaus$ 24 91 0 3 6 0 1 1 $uulaus$ 24 91 0 0 0 0 1 <	Species	2n	2n = 2x	2n = 4x	2n = 8x	Mixoploid	Aneuploids	% 2 <i>n</i> = 2 <i>x</i>	Authority
ulgare 14 42 0 0 0 0 1 $lifilorum$ 14 52 1 0 0 0 0 0 1 $lifilorum$ 14 52 1 0 0 0 0 0 1 $lifilorum$ 14 52 0 0 0 0 0 0 1 $iculatus$ 24 91 0 3 6 0 0 1 $n glaucum$ 14 11 0 0 0 0 0 1 1 vum 14 11 0 0 0 0 0 1 1 vum 20 10 0 0 0 0 0 1 1 1 $vundinaceum$ 20 10 0 0 0 0 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 0 0	Apium graveolens	22	44	0	0	0	2	95.7	Orton, 1985
Itiflorum14521000Itiflorum14520000Itiflorum14520000iculatus249103600iculatus249103600n glaucum141100000n glaucum141100001vum1411000001eale1411000001eale1411000001eale110000012011900000121119000001	Hordeum vulgare	14	42	0	0	0	0	100.0	Karp et al., 1987b
Itifiorum14520001iculatus249103601 n glaucum141100001 vum 141100001 vum 141400001 vum 201000001 $eale$ 141100001 $zundinaceum$ 2010810001 20 119000322	Lolium multiflorum	14	52	-	0	0	0	98.1	Jackson, Dalton, and Dale, 1986
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Lolium multiflorum	14	52	0	0	0	0	100.0	Jackson and Dale, 1988
n glaucum 14 30 0 0 0 0 1 vum 14 11 0 0 9 0 1 vum 14 14 0 0 0 9 0 1 vum 14 14 0 0 0 0 1 1 $eale$ 14 14 0 0 0 0 1 <	Lotus corniculatus	24	91	0	ო	9	0	91.0	Damiani et al., 1985
vum 14 11 0 0 0 9 0 eale 14 14 0 0 0 0 0 1 1 eale 20 10 0 1 20 10 1 20 10 0 1 20 10 0 0 1 20 119 0 0 0 1 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Pennisetum glaucum	14	30	0	0	0	0	100.0	Swedlund and Vasil, 1985
eale 14 14 0 0 0 0 0 1 arundinaceum 20 10 0 1 20 10 20 10 20 11 20 120 120	Pisum sativum	14	11	0	0	0	0	55.0	Natali and Cavallini, 1987
arundinaceum 20 10 0 0 0 0 1 20 108 1 0 0 1 20 119 0 0 3 2 2	Secale cereale	14	14	0	0	0	0	100.0	Lu, Chandler, and Vasil, 1984
20 108 1 0 0 1 20 119 0 0 3 2	Sorghum arundinaceum	20	10	0	0	0	0	100.0	Boyes and Vasil, 1984
20 119 0 0 3 2	Zea mays	20	108	-	0	0	÷	98.2	Edallo et al., 1981
	Zea mays	20	119	0	0	ო	0	96.0	McCoy and Phillips, 1982

Diploid Species	
s in I	
Plants	
g Regenerated	
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Variation	
Chromosomal	
Table 9.6	

Table 9.7 Chromosomal Variation (%) among Regenerated Plants in Potato (*Solanum tuberosum*, 2n = 4x = 48)

Species	48	48±	48++	Authority
S. tuberosum cv. Majestic	57.0	21.5	21.5	Creissen and Karp, 1985
S. tuberosum cv. Maris Bard	63.6	8.4	28.0	Fish and Karp, 1986
S. tuberosum cv. Bintje	32.6	55.8	11.6	Sree Ramulu, Dijkhuis, and Roest, 1983
S. tuberosum cv. Bintje	71.0	21.0	7.9	Sree Ramulu, Dijkhuis, and Roest, 1984a
S. tuberosum cv. Bintje	39.3	46.1	14.6 ^a	Sree Ramulu et al., 1986
S. tuberosum cv. Russet Burbank	61.1	11.2	27.8	Gill, Kam-Morgan, and Shepard, 1986
^a Mixonloid (included)				

* Mixopioid (includeu). Note: $\pm = 1$ or 2 chromosome and ++ = higher than 49 chromosomes.

guishable from plants carrying normal 2n = 48 chromosomes. Similar results have been reported in tobacco (Sacristán and Melchers, 1969; Nuti Ronchi, Nozzolini, and Avanzi, 1981).

Rietveld, Bressan, and Hasegawa (1993) produced about 1000 tuber disc-derived potato plants from cultivars Kennebec, Russet Burbank, and Superior. Plant regenerability trait was genotype dependent. Of the three cultivars, Russet Burbank somaclones expressed the greatest variability for most traits, however, only from 1.0 to 1.3% somaclonal populations exhibited morphological aberrations.

2. Wheat (Triticum aestivum)

Hexaploid wheat (2n = 6x = 42) is an allohexaploid and contains three genomes (A B D). Karp and Maddock (1984) studied chromosomes of 192 regenerated plants derived from immature embryo callus of four hexaploid wheat cultivars. A total of 71% of the regenerants carried the expected 2n = 42 chromosomes, and 29% of the plants were aneuploid (2n = 38-45). The most frequently observed numbers were 2n = 41 or 43. Chromosome interchanges were recorded; however, no plants with chimaerism, polyploidy, or inversion were found. In contrast, all the regenerated plants carried normal chromosome complements of 2n = 42 in the studies of Shimada and Yamada (1979), Ahloowalia (1982), and Ozias-Akins and Vasil (1982). However, they studied only a few plants cytologically.

It is essential to study the chromosomes of regenerated plants at mitosis and meiosis. A wrong conclusion can be drawn unless both stages are analyzed. Fedak, Armstrong, and Handyside (1987) regenerated four plants from suspension cultures of wheat cultivar Chinese Spring. All four plants had 2n = 42 chromosomes, but they carried chromosomal interchanges, while meiosis was normal in the parent.

Chromosomal aberrations were also higher among the regenerated plants of tetraploid wheat (2n = 4x = 28). Bennici and D'Amato (1978) recorded a wide range of chromosomes in root tips (2n = 6-30) and shoot tips (2n = 6-756). However, a majority of root and shoot somatic tissues carried 2n = 28 chromosomes.

3. Triticale (2n = 6x = 42)

Like hexaploid wheat, triticale is an allohexaploid (A B R) and regenerates plants from mature and immature embryos by organogenesis and embryogenesis (Sharma et al., 1981; Nakamura and Keller, 1982; Lazar et al., 1987; Bebeli, Karp, and Kaltsikes, 1988). Armstrong, Nakamura, and Keller (1983) studied the chromosomes of regenerants derived from 1- to 6-month-old calluses of triticale cultivar Welsh. Plants regenerated from 1-month-old callus carried 2n = 42 chromosomes in 61.1% plants, while regenerants from 6-month-old callus showed 42 chromosomes in 19.6% of the plants (Table 9.8). Chromosome aberrations such as telocentrics, duplications, deletions, and interchanges were recorded. It is evident that chromosome breakage occurred near or adjacent to

Table 9.8	Chromosome Constitution in Somatic Cells of Reg from Immature Embryo Derived Calluses of Tritica	
	Number of Dianto with On	Total

	Ν	lumber of P	lants with 2 <i>n</i>	=	Total		
Age of Culture	42	41	Others	% 2 <i>n</i> = 42	Number of Plants		
1 month	33	7	14	61.1	54		
6 months	10	12	31	19.6	51		
Control 1 Control 2	43 12	5 5	1 3	87.8 60.0	49 20		

Source: From Armstrong, K.C., Nakamura, C., and Keller, W.A., *Z. Pflanzenzüchtg.*, 91, 233–245, 1983. With permission.

heterochromatic regions near centromeres, and chromosome aberrations occurred during culture. Chromosome breakage at heterochromatic regions was also demonstrated by Lapitan, Sears, and Gill (1984) in a study of ten regenerated amphidiploid (2n = 56) plants of wheat × rye (AABBD-DRR) by utilizing Giemsa C-banding technique. They recorded three wheat/rye and one wheat/wheat chromosome translocation, seven deletions, and five amplifications of heterochromatic bands of rye chromosomes. Twelve of the 13 breakpoints in chromosomes involved in translocations and deletions occurred in heterochromatin.

B. Age of Callus

Based on chromosome analysis of the calluses, one should expect to obtain a higher frequency of regenerants with altered chromosomes from the older cultures (Choi, Lemaux, and Cho, 2000b, 2001). This has been demonstrated in maize and oat. Lee and Phillips (1987) determined the chromosomes of 78 regenerants from 3- to 4-month-old cultures of maize (2n = 2x = 20). All the plants carried normal chromosome constitution. In contrast, 189 plants regenerated from 8- to 9-month-old cultures showed 91 plants (48%) cytologically abnormal. Interchanges were the most frequent alterations (38/45); a high frequency (35%) of deficient chromosomes was also detected. A low frequency of tetraploids and a trisomic plant were also identified. In a similar study, McCoy and Phillips (1982) observed fewer cytologically abnormal plants (4/59) in 8-month-old cultures of maize than those reported by Lee and Phillips (1987).

McCoy, Phillips, and Rines (1982) conducted an extensive study on chromosome analysis, at meiosis, of plants regenerated from 4-, 8-, 12-, 16-, and 20-month-old cultures of the oat cultivars Lodi and Tippecanoe. Oat is an allohexaploid (2n = 6x = 42) with A, B, and C genomes. They demonstrated that the frequency of chromosomal aberrations was increased with the culture age and was also genotype dependent. After 4 months in culture, 49% of the Lodi regenerants carried altered chromosomes, and 12% of the Tippecanoe regenerates were abnormal. But after 20 months in culture, 88% of the Lodi and 48% of the Tippecanoe regenerated plants were cytogenetically abnormal. Chromosomal aberrations such as interchanges, monosomics, trisomics, and deletions were frequently observed. Long-term (4 to 11 years) culture-derived somaclones of rice contained deletion in plastid genome, and that was associated with the accumulation of starch granules (Kawata et al., 1995).

C. Source of Explant Materials

It has been demonstrated that regenerants from protoplast culture generally carry a higher degree of an euploidy and phenotypic variability than those plants regenerated from immature embryo or shoot tip culture. Newell, Rhoads, and Bidney (1984) reported in *Brassica napus* (2n = 38) that regenerated plants carried 87% 2n = 38, 7.8% monosomics, 2.6% trisomics, and 1.3% tetraploids. In contrast, protoplast-derived regenerates were 44% 2n = 38, 20% hypodiploid, and 36% tetraploid or hypotetraploid.

Kanda et al. (1988) studied chromosomes in seven plants regenerated from 7- to 10-month-old protoplast callus of rice. They observed one diploid (2n = 24), four tetraploids (2n = 48), and two aneuploids (2n = 46) plants. Contrarily, Kobayashi (1987) regenerated 25 plants from orange (*Citrus sinensis*) protoplasts and recorded no significant variations among somaclones in leaf and flower morphology, leaf oil, isozyme banding pattern, and chromosome numbers.

D. Culture Conditions

Chromosomal aberrations among regenerated plants are influenced by the media components and also by the way cultures are maintained. Fish and Karp (1986) obtained a higher frequency of euploid potato plants by media modifications. They cytologically screened 178 protoplast-derived

and observed 63.6% of the regenerants that carried 2n = 48 chromosomes, but in their earlier studies (Karp et al., 1982), only 4% of the regenerants were euploids.

Ogihara (1981), working with *Haworthia setata*, observed an average of 89.2% diploid (2n = 14), 6.2% tetraploid (2n = 28), 0.4% monosomic (2n = 13), and 4.1% modified chromosomes among regenerants from culture maintained by subculturing. Chromosome aberrations were higher in plants from the culture maintained by cloning: diploid = 42.2%; tetraploid = 25.8%; and modified chromosomes = 34.0%.

V. CHROMOSOMAL ABERRATIONS IN HAPLOID CALLUS AND THEIR REGENERANTS

Chromosomal variability is comparatively higher in anther-derived calluses and their regenerated plants than from plants regenerated from somatic tissues. Chromosome numbers of anther-derived plants are often doubled (Bennici, 1974; Mix, Wilson, and Foroughi-Wehr, 1978; Chen and Chen, 1980; Wenzel and Uhrig, 1981; Sacristán, 1971, 1982; Keathley and Scholl, 1983; Santos, Handro, and Floh, 1983; Toriyama, Hinata, and Sasaki, 1986; Pohler et al., 1988; Wersuhn and Sell, 1988; Kudirka, Schaeffer, and Baenziger, 1989).

Sacristán (1971) studied karyotypic changes in callus cultures from haploid (2n = 3) and diploids (2n = 6) *Crepis capillaris*. The degree of polyploidization and chromosomal interchanges in haploid culture was considerably higher than in diploid culture.

In anther culture, if plants regenerate through embryogenesis, the majority of them are haploid, while plants regenerated from callus are diploid. It has been suggested that usually many cells take part in the formation of a bud, while single cells give rise to somatic embryos (Constantin, 1981). In *Brassica napus* anther culture, Sacristán (1982) observed that plants derived through embryogenesis were haploid, while most of the plants regenerated from callus were diploid, and no haploid plants were obtained. Toriyama, Hinata, and Sasaki (1986) regenerated 15 mature plants from protoplasts that were isolated from cell suspensions of anther callus in rice. Four plants were haploid (2n = 12), and 11 plants were diploid (2n = 24). Diploids were uniform morphologically, but seed set varied from 0 to 95%. Pohler et al. (1988) studied chromosomes of callus tissues and regenerated plants from anther culture of hexaploid triticale. Chromosome variation was the highest (2n = 18-43) in the calluses, and it was the least pronounced in plants (2n = 20-43) derived from embryoids. In anther calluses, 24.8% cells carried 2n = 21 and 15.6% cells 2n = 42 chromosomes. A total of 184 plants was analyzed by root tip count. Plants derived from embryoids showed a higher frequency (83.3%) of euhaploids (2n = 3x = 21) than plant regenerants from calluses (31.6%).

Utilizing liquid culture medium, Uhrig (1985) regenerated 313 plants from anther culture of diploid (2n = 2x = 24) potato. The majority of plants (80.2%) retained diploid chromosome numbers; however, low frequencies of haploid (0.6%) and triploid (0.3%) plants were obtained. Chromosome doubling occurred in 13.7% of the plants, and mixoploidy was recorded in 5.2% of plants. A large number of embryoids can be generated with fewer chromosomal aberrations by suitable genotype and media modifications.

Tempelaar et al. (1985) recorded that monohaploid potato (2n = 12) polyploidized at a faster rate than dihaploid (2n = 24) and tetraploid (2n = 48). Of 36 plants studied cytologically from monohaploid culture, 14 were dihaploid, tetraploid, or mixoploid, and only two plants were monohaploids. Karp et al. (1984) scored nearly 100% doubled monohaploids (2n = 2x = 24) from a single leaf regeneration cycle of monoploid (2n = x = 12) potatoes, and 50% of the regenerants from doubled monohaploid leaves were tetraploids (2n = 4x = 48). Few mixoploids and aneuploids were found. Wenzel and Uhrig (1981) regenerated 6000 androgenic clones from microspores (n =12) of dihaploid potato (2n = 24). The chromosome numbers of about 90% of these plants doubled spontaneously and yielded fertile plants, and only 10% of the plants were monohaploid. Pijnacker and Ferwerda (1987) studied karyotypic changes in cell suspension, calluses, and their regenerants of an S-(2-amino ethyl) cystein resistant cell line of a dihaploid (2n = 24) potato. Chromosome numbers in cell suspension varied between 2n = 33 to 151, with peaks near 40 and 75 chromosomes. In a comparison of a 2-year-old callus culture and a morphogenic callus, chromosome numbers ranged from 36 to 217 and 33 to 130, respectively. However, the regenerates carried 2n = 34 or 35 chromosomes.

Bennici (1974) studied chromosomes in roots from old and young calluses of haploid (2n = x = 9) *Pelargonium*. In roots from young calluses, 71.42% of cells were diploid (2n = 18), while roots from old calluses had only 13.81% diploid cells, and the remaining cells possessed polyploid and aneuploid chromosome numbers.

Wersuhn and Sell (1988) recorded mainly tetraploid plants from anther culture of *N. tabacum* cv. Samsun (2n = 48). Plants regenerated from anther culture were more highly aberrant cytologically than plants regenerated from seed.

Qiren, Zhenhua, and Yuanhua (1985) regenerated 1715 plants from anther culture of rice. Chromosome counts revealed: diploid (2n = 24), 35.7%; haploid (2n = 12), 49.1%; polyploids, 5.1%; and aneuploids, 10.2%. A high frequency of regenerants from anther culture are albino in cereals, such as wheat (Marsolais, Séfuin-Swartz, and Kasha, 1984), barley (Mix, Wilson, and Foroughi-Wehr, 1978), and rye (Wenzel, Hoffmann, and Thomas, 1977). Of the 390 anther-derived plants in rye, 97 plants were green, and 293 were albino. A chromosome count of 100 plants showed 31 plants with 2n = x = 7, 63 plants with 2n = 2x = 14, five plants with 2n = 1x + 2x, and one plant with 2n = 4x = 28 (Wenzel, Hoffmann, and Thomas, 1977). Similarly, in barley, Mix, Wilson, and Foroughi-Wehr (1978) obtained 600 green plants among 4000 regenerants, and only 20% of the green plants were haploid. The remaining plants were diploid, triploid, tetraploid, and aneuploid. Diploid plants were completely fertile.

VI. CHROMOSOMAL ABERRATIONS IN SOMATIC HYBRIDS

Somatic hybridization by protoplast fusion is an alternative approach to transfer alien genes, where sexual crosses are not successful in distantly related and incompatible species (Bhojwani, Evans, and Cocking, 1977; McComb, 1978; Kumar and Cocking, 1987; Wolters et al., 1994a; Jacobsen et al., 1995; Waara and Glimelius, 1995). Several interspecific somatic hybrids involving species of the genera *Brassica, Datura, Daucus, Medicago, Nicotiana*, and *Petunia* have been reported (Table 9.9), and only a few intergeneric somatic hybrids have been produced (see D'Amato, 1985). Somatic hybridization technique helped the resynthesis of allopolyploid crops to create their genetic variability and restore ploidy level and heterozygosity after breeding at reduced ploidy level in polyploid crops (Waara and Glimelius, 1995). It is essential that one of the fusants should regenerate complete plants through protoplast.

Although considerable efforts have been made to identify somatic hybrids based on isozyme banding patterns and molecular approaches, the information on systematic chromosome analysis is lacking. Somatic hybrid plants are expected to carry the somatic chromosome (2n) constitutions of both parents (Table 9.9). For example, the somatic hybrid of tetraploid potato (2n = 4x = 48) and tomato (2n = 2x = 24) should possess 2n = 6x = 72 chromosomes.

The chromosomal stability of a somatic hybrid depends upon the degree of closeness between the genomes of the fusion parents. Somatic hybrids can be divided into three categories: (A) stable, (B) partially stable, or (C) unstable.

A. Stable Somatic Hybrids

Carlson, Smith, and Dearing (1972) generated stable somatic hybrid plants (2n = 42) by protoplast fusion of *Nicotiana glauca* (2n = 24) and *N. langsdorffii* (2n = 18). All the regenerated plants

Somatic Hybrids	Expected 2n	Observed 2n	Authority
Arabidopsis thaliana $(2n = 8x = 40) +$ Brassica campestris $(2n = 20)$	60	35–45	Hoffmann and Adachi, 1981
Brassica oleracea $(2n = 18) + B$. campestris $(2n = 20)$	38	18–54	Schenck and Röbbelen, 1982
Brassica napus $(2n = 38) + Arabidopsis$ thaliana $(2n = 10)$	48	38–86	Forsberg, Landgren, and Glimelius, 1994
Medicago sativa $(2n = 32) + M$. coerulea $(2n = 16)$	48	48	Pupilli et al., 1995
Lycopersicon esculentum $(2n = 24) + L$. peruvianum $(2n = 48)$	72	72	Parokonny et al., 1997
Lycopersicon esculentum $(2n = 24) +$ Solanum lycopersicoides $(2n = 24)$	48	48–68	Handley et al., 1986
Nicotiana glauca $(2n = 24) + N$. langsdorffii $(2n = 18)$	42	42	Carlson, Smith, and Dearing, 1972
Nicotiana glauca $(2n = 24) + N$. langsdorffii $(2n = 18)$	42	60–66	Morikawa et al., 1987
Nicotiana tabacum $(2n = 48) +$ Nicotiana nesophila $(2n = 48)$	96	96	Evans et al., 1982
Nicotiana sylvestris $(2n = 24) + N$. plumbaginifolia $(2n = 20)$	44	33–54	Parokonny et al., 1994
Oryza sativa $(2n = 24) + Porteresia$ coarctata $(2n = 48)$	72	72	Jelodar et al., 1999
Petunia parodii $(2n = 14) + P$ inflata $(2n = 14)$	28	14–36	Schnabelrauch, Kloc- Bauchan, and Sink, 1985
Solanum melongena $(2n = 24) + S$. sisymbriifolium $(2n = 24)$	48	38–48	Gleddi, Keller, and Setterfield, 1986
Solanum tuberosum $(2n = 24) + S$. brevidens $(2n = 24)$	48	45–89	Fish, Karp, and Jones, 1988
Solanum tuberosum $(2n = 24) +$ Lycopersicon esculentum $(2n = 24)$	48	50–72	Melchers, Sacristán, and Holder, 1978
Solanun tuberosum $(2n = 24) + Solanum brevidens (2n = 24)$	48	48–72	Gavrilenko et al., 2002

Table 9.9 Chromosome Variation in Regenerated Plants by Protoplast Fusion

were similar to the sexually synthesized amphidiploids. Evans et al. (1982) recovered stable somatic hybrids (2n = 96) of *N. tabacum* (2n = 48) and *N. nesophila* (2n = 48). Variation in clones was attributed to cytoplasmic segregation and mitotic recombinations. Schenck and Röbbelen (1982) synthesized for the first time rapeseed (*Brassica napus*, 2n = 38) by fusion of protoplast from *B. oleracea* (2n = 18) and *B. campestris* (2n = 20). Hybrid plants flowered and set seed. Schnabelrauch, Kloc-Bauchan, and Sink (1985) obtained stable and unstable somatic hybrids in *Petunia parodii* and *P. inflata*. Fish, Karp, and Jones (1988) generated somatic hybrids of dihaploid *Solanum tuberosum* (2n = 24) and *S. brevidens* (2n = 24). Tetraploid (2n = 48)plants were completely stable (Table 9.9).

Heath and Earle (1995) produced fertile 51 amphiploids (aacc) and one putative hexaploid (aacccc) *Brassica napus* by somatic hybrids through protoplast fusion of *Brassica oleracea* var. *botrytis* and *Brassica rapa* var. *oleifera* (high erucic content in seed oil; 22:1). An erucic acid content as high as 57.4% was found in the seed oil of one regenerated plant. The hybrids recovered carried large seed size, lodging resistance, and nonshattering seed pods.

B. Partially Stable Somatic Hybrids

Chromosome elimination during callus growth and also in regenerants occurs in some somatic hybrids (Dudits et al., 1977; Krumbiegel and Schieder, 1979; Hoffmann and Adachi, 1981; Lazar, Dudits, and Sung, 1981; de Vries et al., 1987). Krumbiegel and Schieder (1979) selected 13 somatic hybrids of *Datura inoxia* (2n = 24) and *Atropa belladona* (2n = 48). Chromosome

numbers ranged from 84 to 175. Chromosome instability was recorded, but no evidence of chromosome elimination was observed. Hoffmann and Adachi (1981) studied somatic hybrids of *Arabidopsis thaliana* (2n = 40) and *Brassica campestris* (2n = 20) and observed elimination of chromosomes from both species. Chromosome analysis of ten plants showed 2n = 35-45 chromosomes; the somatic hybrid should have consisted of 2n = 60 (Table 9.9). Chromosome numbers in different vegetative organs were stable. Chromosome structural changes were observed. This suggests that chromosome instability arose later in the stage of the plant growth, because initial somatic hybrid plants were stable (Gleba and Hoffmann, 1978). de Vries et al. (1987) recorded no preferential loss of species-specific chromosomes in the *Nicotiana plumbaginifolia* (2n = 48) × *Solanum tuberosum* (2n = 12) somatic hybrid.

C. Unstable Somatic Hybrids

Preferential loss of chromosomes occurs in those somatic hybrids where parents are genomically incompatible and distally related (Binding, 1976; Binding and Nehls, 1978; Maliga et al., 1978; Zenkteler and Melchers, 1978; Wetter and Kao, 1980; Pental et al., 1986). This may be attributed to the asynchronous mitotic cycle (Bennett, Finch, and Barclay, 1976). Uniparental chromosome elimination has been found in sexual hybrids of *Hordeum vulgare* (2n = 14) and *H. bulbosum* (2n = 14, 28), where *H. bulbosum* chromosomes are eliminated during early embryo development, and only haploid (2n = x = 7) embryos with *H. vulgare* chromosomes are recovered (Kasha and Kao, 1970; Subrahmanyam and Kasha, 1973). A similar phenomenon was observed in wheat × maize (Laurie and Bennett, 1989), wheat × pearl millet (Laurie, 1989), and oat (2n = 6x = 42) × maize (2n = 2x = 20) (Rines and Dahleen, 1990) crosses. In these crosses, maize and pearl millet chromosomes were eliminated during the first few cell-division cycles in most of the embryos. Isolated oat–maize addition lines are excellent genetic stock for studying the expression of maize genes in oat (Muehlbauer et al., 2000).

Maliga et al. (1978) analyzed chromosome numbers in somatic hybrids of *Nicotiana tabacum* (2n = 48) and *N. knightiana* (2n = 24) and recorded variation in chromosome numbers within individual plants. Variegation in leaf and flower color and segregation for morphological traits in vegetatively multiplied plants were attributed to segregation of chromosomes in the somatic cells, a result of numerical chromosomal instability.

Wetter and Kao (1980) recorded preferential loss of *Nicotiana glauca* chromosomes in the *Glycine max* (2n = 40) + N. *glauca* (2n = 18) somatic hybrid. Similarly, Pental et al. (1986) observed the elimination of tobacco chromosomes in N. tabacum (2n = 48) + Petunia hybrida (2n = 14) somatic hybrids.

Binding and Nehls (1978) investigated cytologically somatic hybrid calluses of *Vicia faba* (2n = 12) and *Petunia hybrida* (2n = 14). The putative fusants carried predominantly nuclei or chromosomes of one or the other species and a few chromosomes of the other species. If a cell carried predominantly *Vicia faba* chromosomes, *Petunia hybrida* chromosomes were eliminated, and *vice versa*.

Melchers, Sacristán, and Holder (1978) identified somatic hybrids of potato (2n = 24) and tomato (2n = 24) based on morphological features, isozyme banding patterns, and chromosome counts. Somatic hybrid plants are expected to carry 2n = 48 chromosomes. Chromosome numbers in three plants ranged from 50 to 56, and a fourth plant carried 2n = 72 (Table 9.9). The fourth plant was developed by triple fusion (2 potato + 1 tomato protoplast), because this particular regenerant contained significantly fewer staining bands of the tomato small subunit. Triple fusion is rather common in protoplast fusion (Schenck and Röbbelen, 1982; Morikawa et al., 1987). Ninnemann and Jüttner (1981) analyzed volatile patterns of potato + tomato hybrids of Melchers and found no evidence of preferential elimination of chromosomes. However, they did not study the chromosomes. This contradicts the statement of Melchers, Sacristán, and Holder (1978) that some of the plants are chimeras containing tissues with different chromosome numbers.

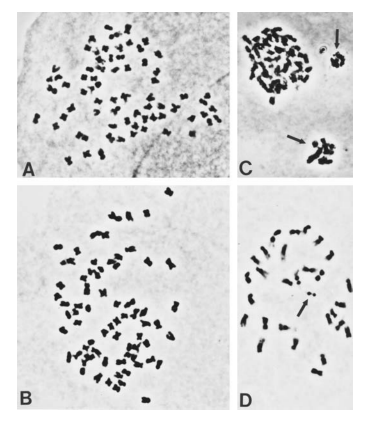


Figure 9.4 Cells observed in regenerants of potato (2n = 48) and tomato (2n = 24) somatic hybrids. (A) 2n = 72; (B) 2n = 68; (C) a cell showing the chromosome elimination (arrows); (D) 2n = 29 (note a minute chromosome — arrow). (From Singh, R.J., unpublished results.)

A detailed chromosome analysis of potato (2n = 48) and tomato (2n = 24) somatic hybrids developed by Shepard et al. (1983) was conducted by the author. Somatic hybrid plants are expected to carry 2n = 72, and the initial hybrid carried 2n = 72 chromosome (Figure 9.4A). Chromosome numbers in vegetative cuttings of potato + tomato somatic hybrids differed from cell to cell, root to root, cutting to cutting, plant to plant, and hybrid to hybrid. Chromosome numbers ranged from 2n = 24 and 2n = 120. It appears that each organ is a mixture of cells composed of various chromosome numbers. The deviation from the expected 72 chromosomes in all the cuttings of somatic hybrid is due to chromosome elimination (Figure 9.4B,C,D). The occurrence of 48 (+4) (Figure 9.5A) and 24 (Figure 9.5B), chromosome cells suggests that these cells probably carry tomato and potato (plus a few tomato) genome chromosomes, respectively. It is evident that chromosome elimination is gradual and occurs in each mitotic cycle. In contrast to these observations, Shepard et al. (p. 687, 1983) reported that somatic hybrid cuttings displayed chromosome numbers ranging from 62 to 72, depending on the cutting. The most frequent encountered chromosome number for the "Rutgers" hybrid was 70, with greater variability observed for the "Nova" hybrids. These data indicate a degree of mitotic instability and some chromosome segregation in vegetative cuttings but not wholesale chromosome elimination. This statement is mere speculation. Wolters et al. (1994b) cytologically examined a total of 107 somatic hybrids of tomato + potato. Most (79%) hybrids were aneuploid, some were hyperploid, and others were hypoploid. Chromosome numbers ranged from 34 to 72. Preferential tomato chromosome elimination in the tomato + potato fusion hybrids and variable number of tomato chromosomes (6–11) in BC₁ were recorded by Garriga-Calderé et al. (1997). This contradicts the assumption of Shepard et al. (1983). An extensive chromosome analysis of the potato +

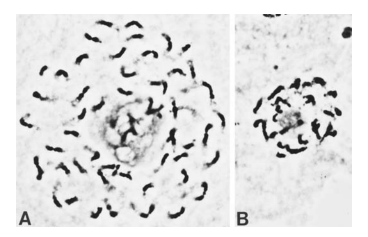


Figure 9.5 Cells observed in regenerants of potato (2n = 48) and tomato (2n = 24) somatic hybrids. (A) A prophase cell with 2n = 52. (B) A prophase cell with 2n = 24. (From R.J. Singh, unpublished results.)

tomato (2n = 72) somatic hybrid will identify derived potato and tomato lines. After BC₁ and BC₂ generations, genomic *in situ* hybridization (GISH) revealed homoeologous chromosome pairing between tomato and potato chromosomes (Wolters et al., 1994b).

CHAPTER 10

Transgenic Crops

I. INTRODUCTION

The progress made in creating genetically modified organisms (GMO) during the past 16 (1986–2002) years is substantial (www.isb@vt.edu). For successful genetic engineering of crops, an alien gene (donor species) linked to a reporter gene and other DNA sequences essential for insertion and expression is inserted into single and totipotent cells of the recipient species. The transformed cells proliferate *in vitro* during selection and regeneration, creating a novel cultivar. Genetic transformation has improved many economically important crops. One such example is the soybean, a leguminous crop widely used for oil, feed, and soyfood products (Singh and Hymowitz, 1999).

II. PRESENT STATUS OF TRANSGENIC CROPS

To date, 109 transgenic organisms consisting of 392 phenotypes have been evaluated and released in the U.S. (www.isb@vt.edu). Many transformants carry valuable traits: resistance to pests and pathogens, tolerance to herbicides, quality traits (protein, oil, carbohydrate, and fatty acids including amino acids composition), modified reproductive capacity, photosynthetic enhancement and yield increase, delayed senescence, enhanced flavor and texture, longer shelf life, and more healthful produce (Dunwell, 2000). The status of all notifications and release permits by year of patents is shown in Table 10.1. The number of approved trials in the U.S. has increased from 18 (1987) to 1117 (2001), and by April 2002, notifications for 536 permits have been issued. The production of GMO has been revolutionized for the last decade by private and public institutions. The Monsanto company, for example, has played a major role in applying for the maximum (3105) permits, followed by Dupont-Pioneer (Table 10.2).

III. PRODUCTION OF TRANSGENIC CROPS

Several procedures are available for producing transgenic crops. Foreign DNA is delivered into recipient species through *Agrobacterium*, particle bombardment, electroporation of protoplast, polyethylene glycol (PEG), and microinjection of protoplasts (Songstad, Somers, and Griesbach, 1995). Recently, Joersbo (2001) proposed a three-selection system using glucuronic acid, mannose, and xylose that resulted in higher transformation frequencies compared with the frequently used kanamycine selection.

The development of gene transfer methodology is crop dependent. For example, alien genes of economic importance can be delivered into soybean by *Agrobacterium*-mediated transformation of cotyledonary explants (Hinchee et al., 1988; Clemente et al., 2000) and by particle bombardment,

Year	Received	Approved	Delayed Approved ^a	Total Approved ^ь	Denied	Withdrawn	Void	Pending
1986	0	0	0	0	0	0	0	0
1987	9	5	4	9	0	0	0	0
1988	18	12	6	18	0	0	0	0
1989	38	24	14	38	0	0	0	0
1990	58	37	21	58	0	0	0	0
1991	107	69	38	107	0	0	0	0
1992	161	122	28	150	0	11	0	0
1993	374	277	29	306	0	68	0	0
1994	608	555	38	593	6	9	0	0
1995	706	666	14	680	2	18	5	0
1996	654	590	35	625	8	20	0	0
1997	808	730	13	743	33	28	3	0
1998	1206	1061	25	1086	108	10	2	0
1999	1061	960	25	985	46	23	6	0
2000	1012	898	37	935	57	16	1	2
2001	1189	1040	77	1117	39	20	2	8
2002	725	536	0	536	12	1	0	176
Total	8734	7582	404	7986	311	224	19	186

Table 10.1 Status of All Notifications and Release Permits of Transgenic Crops by Year (www.isb.vt.edu)

^a Approval the next calender year from year received.
 ^b Sum of approved and delayed approved.

Institution	Total	Acknowledged	Issued	Pending	Denied	Withdrawn	Void
Monsanto	3105	2759	157	60	72	51	6
Pioneer	658	541	55	7	32	22	1
AgrEvo	344	312	14	0	14	4	0
Du Pont	321	305	15	0	0	1	0
ARS	214	137	42	12	12	8	3
DeKalb	192	172	9	0	8	3	0
Seminis Vegetable Seeds	179	156	18	1	3	1	0
Calgene	175	90	74	0	2	8	1
Avantis	128	98	4	12	6	8	0
lowa State University	97	71	6	7	11	2	0
University of Idaho	94	72	14	0	6	1	1
Rutgers University	92	71	12	5	4	0	0
ProdiGene	92	57	25	5	2	3	0
DNA Plant Tech	91	74	15	0	0	2	0
Northrup King	88	69	11	0	3	5	0
Upjohn	85	10	63	0	0	12	0
Dow	84	70	3	8	1	2	0
Stine Biotechnology	81	71	0	6	4	0	0
Asgrow	81	49	26	0	1	5	0
Novartis Seeds	79	74	3	0	2	0	0
Cargill	71	54	11	0	1	5	0
Harris Moran	63	44	11	3	4	1	0
Agracetus	61	57	3	0	1	0	0
Agritope	60	47	6	0	6	1	0
Stanford University	60	50	0	1	9	0	0

Table 10.2 Status of Notifications and Release Permits by Company (www.isb.vt.edu)

invented by Klein et al. (1987) (Christou, 1997). Hinchee et al. (1988) successfully isolated glyphosate-tolerant transformants using the *Agrobacterium*-mediated gene transfer method, and Padgette et al. (1995) reported the stability of the transformants. The soybean is marketed as Roundup Ready[®] soybean and is widely grown in the United States and is spreading into other soybean-growing countries of the world. Genetic transformation created Bollgard[®] cotton, Yield-Gard[®] maize (www.biotechbasics.com), FLAVR SAVRTM tomato (Kramer and Redenbaugh, 1994), and golden rice (Potrykus, 2001). Bollgard[®] cotton is commercially grown in the United States, Australia, China, Mexico, South Africa, Argentina, and was recently approved for India. The FLAVR SAVRTM tomato containing superior consumer quality, shelf life, and flavor failed to attract consumers, as it was more expensive than nontransgenic tomato.

IV. CHROMOSOMAL ABERRATIONS IN TRANSGENIC CROPS

Chromosomal aberrations (numerical and structural changes) are common in cell and tissue culturederived calluses and their regenerants (Singh, 1993). Selection and osmotic treatment exacerbate cytological aberrations in transformed barley cultures, but bombardment did not appear to affect the frequency of cytological aberrations in cells of callus cultures (Choi, Lemaux, and Cho, 2001). Chromosomally abnormal soybean cultures used in transformation experiments regenerate mostly sterile plants (Singh et al., 1998). Thus, soybean is one of the most difficult large-seeded legumes to regenerate plants from cell and tissue cultures. The earlier reports suggest that reproducible methods of regeneration depend upon type and age of explants, media composition (basic salts and growth hormone combinations), and genotypes (Christianson, Warnick, and Carlson, 1983; Barwale, Meyer, and Widholm, 1986; Lazzeri, Hildebrand, and Collins, 1987a,b; Wright et al., 1987; Bailey, Boerma, and Parrott, 1993). The degree of sterility in transgenic soybean plants is related to the period the cultures are nurtured on 2,4-D prior to transformation, posttransformation stresses, and the genetic background of the explant.

A. Time in Culture with 2,4-D Prior to Transformation

Generally, older cultures lose the capacity to regenerate plants due to high ploidy or an increasing frequency of an euploidy that lead to a chromosomal imbalance (Singh, 1993). Thus, these cultures will generate a low frequency of morphologically deformed plants. Singh et al. (1998) examined chromosome counts (mitotic prometaphase and metaphase) in embryogenic suspension cultures (nontransgenic) and in roots from developing somatic embryos (transgenic or nontransgenic) from T_0 , T_1 , and T_2 transgenic two Asgrow genotypes A2242 and A2872. Seeds of A2242 (control) carried the expected 2n = 40 chromosomes. The chromosome counts of suspension cultures and germinating somatic embryos from culture 817 are useful (Table 10.3). Tetraploidy occurred even in the germinating somatic embryos from 7.83 months on 2,4-D; three samples showed 2n= 80 normal chromosomes, and two samples, in addition to tetraploidy, carried 2n = 79 + 1dicentric chromosomes. Cells after 11.26 months on 2,4-D suspension cultures showed 2n = 80chromosomes, and transgenic plants recovered from these cultures were tetraploid. Embryo suspensions of culture 826, which were on 2,4-D for 4.20 months, displayed cells with 2n = 80chromosomes. Suspension culture 825 (6.86 months on 2,4-D) and somatic embryo culture 828 (only 2.86 months on 2,4-D) showed diploid cells. The germinating embryos derived from culture 817 ranging from 7.17 to 16.73 months on 2,4-D possessed imbalanced chromosomes (Table 10.3). The older cell lines, therefore, are not desirable for transformation, because chromosomal and genetic abnormalities occur with the age. This may lead to difficulty in recovery of regenerable and fertile plants.

In order to ensure morphologically normal transformants with complete fertility, Stewart et al. (1996) bombarded 3-month-old globular-stage embryos of soybean cv. "Jack" and used postbombardment selection for transgenic line on a solidified medium instead of liquid medium.

Culture ID	R0 Phenotype	Origin of Roots	Months on 2,4-D	Number of Samples	2 <i>n</i>	Karyotype
A2242	Control			5	40	Normal
22–1	Diploid	T ₁	6.43	7	40	Normal
22–1	Tetraploid	T ₁	6.90	5	40;41	39 + (1);38 + (3) ^a
22–1	Diploid	T ₁	6.90	2	40	Normal
22–1	Diploid	T ₁	6.96	9	40	Normal
22–1	Diploid	T ₁	6.96	2	80	Normal
22–1	Diploid	T ₁	7.00	8	40	Normal
22–1	Diploid	T ₁	7.30	9	80	Normal
22–1	Diploid	T ₁	7.96	8	80	Normal
22–1	Diploid	T ₁	9.00	1	80	Normal
22–1	Tetraploid	To	11.47	3	80	Normal
22–1	Tetraploid	To	15.36	1	80	Normal
22–1	—	Embryo	15.36	5	80	3,80;2,79 + 1 ^b
22–1	—	Embryo	16.73	4	80	3,80;1,40 + 80°
817	—	Embryo	7.17	4	40	3,40;1,39 + 1 ^d
817	—	Embryo	7.33	5	40	4,40;1,39 + 1 ^d
817	—	Embryo	7.83	5	80	3,80;2,79 + 1°
817	—	Embryo	8.70	1	40	Normal
817	—	Τ _o	9.13	1	40	Normal
817	—	Suspension	11.26	1	80	Normal
817	—	Embryo	12.43	2	80	Normal
825	—	Suspension	6.86	1	40	Normal
828	—	Embryo	2.86	1	40	Normal
826	_	Suspension	4.20	1	80	Normal

Table 10.3 Chromosome Analysis at Somatic Metaphase in the Transgenic Asgrow Soybean Genotype A2242

^a Three small metacentric chromosomes.

^b One megachromosome.

° Chimaera 40 + 80 chromosomes.

^d Long chromosome.

 $^{\rm e}$ One sample with 79 + 1 dicentric chromosomes and the other sample with 79 + 1 fused centromeric chromosomes.

Source: From Singh, R.J. et al., Theor. Appl. Genet., 96, 319–324, 1998. With permission.

Transgenic plants were not analyzed cytologically, but all plants were fertile and morphologically normal. Maughan et al. (1999) isolated four normal and fertile β -casein transgenic soybean cv. "Jack" plants via particle bombardment of 3- to 4-month-old somatic embryos nurtured on solid medium. Hazel et al. (1998) examined growth characteristics and transformability of embryogenic cultures of soybean cv. "Jack" and Asgrow A2872. The most transformable cultures were constituted of tightly packed globular structures and cytoplasm-rich cells in the outermost layers of the tissues with the highest mitotic index. On the other hand, the outer layers of the less transformable culture had more lobed cells with prominent vacuoles. Santarém and Finer (1999) used 4-week-old proliferative embryogenic tissue of soybean cv. "Jack" maintained on semisolid medium for transformation by particle bombardment. They produced fertile transgenic soybean 11 to 12 months following culture initiation. They claim a significant improvement over bombardment of embryogenic liquid suspension culture tissue of soybean, where transformability was recorded at least 6 months after culture initiation (Hazel et al., 1998). Clemente et al. (2000) produced 156 primary transformants by Agrobacterium-mediated transformation. Glyphosate-tolerant shoots were identified after 2 to 3 months of selection. The R_0 (T_0) plants were fertile (262 seeds per plant) and did not express gross phenotypic abnormalities. This suggests that transformation in the soybean must be conducted using young (1 to 3 months) cultures.

The plants regenerated from cultures also inherited chromosomal abnormalities (Table 10.3). Chromosomes of nine T_1 populations from culture 22–1 were examined. The parental cultures

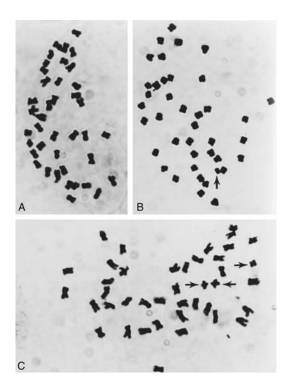


Figure 10.1 Mitotic chromosomes in root tips of seedlings from Asgrow soybean line A2242 R₁ generation.
(A) 2n = 40 showing normal karyotype; (B) 2n = 39 + 1 small metacentric chromosomes (arrow); and (C) 2n = 38 + 3 small metacentric chromosomes (arrows). (From Singh, R.J. et al., *Theor. Appl. Genet.*, 96, 319–324, 1998. With permission.)

were in contact with 2,4-D for 6.43 to 9.00 months prior to transformation. Although T_0 plants from eight cultures expressed normal, diploid morphological features, T₁ progenies from four populations carried 2n = 80 chromosomes, and four populations had 2n = 40 chromosomes (Figure 10.1A). The T_0 phenotype of one T_1 population of culture 22–1 was similar to that of tetraploid. These plants showed dark-green leathery leaves and produced mostly one-seeded pods. Chromosome counts from five T_1 seedlings showed 2n = 39 + 1 small metacentric chromosomes (Figure 10.1B) in three plants, and one plant each contained 2n = 38 + 3 small metacentric chromosomes (Figure 10.1C) and 2n = 40 chromosomes. The 40-chromosome plant may have had a small deletion, which could not be detected cytologically, or may have carried desynaptic or asynaptic genes. Four T_0 -derived plants from culture 22-1 (three plants from cultures after 11.47 months on 2,4-D and one plant after 15.36 months on 2,4-D) were morphologically tetraploid and were cytologically confirmed to have 2n = 80 chromosomes. Three transformants had 2n = 80 (Figure 10.2A), and two contained 79 + 1 megachromosome (Figure 10.2B). Most reports agree that structural and numerical changes in chromosomes induced during culture are caused by 2,4-D (Singh, 1993). Thus, it is advised to examine cytologically the chromosomes in pretransformed material to ensure that it is devoid of structural and numerical changes (Poulsen, 1996).

It has been established that various stresses, such as medium composition, age of culture, nature of culture — morphogenic versus nonmorphogenic, genotype of the explant, media, and *in vitro* culture — during the passage of plant regeneration, induce chromosomal instability (Singh, 1993; D'Amato, 1995). Chromosomal constitution in transgenic versus nontransgenic barley (2n = 14) demonstrated that the extent of ploidy changes in transgenic plants was intensified, presumably due to additional stresses that occurred during transformation. The delivery of foreign DNA into plant cells involves several stressful events, such as vacuuming,

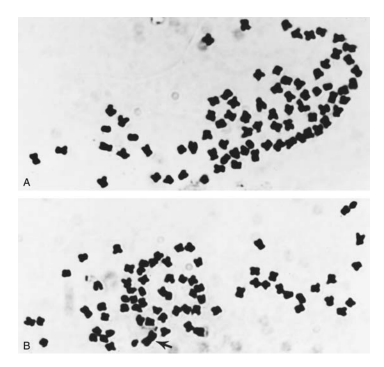


Figure 10.2 Mitotic metaphase chromosomes in root tips of seedlings from Asgrow soybean line A2242 grown in agar. (A) 2n = 80 showing normal karyotype; (B) 2n = 79 + 1 mega (monocentric) chromosome (arrow). (From Singh, R.J. et al., *Theor. Appl. Genet.*, 96, 319–324, 1998. With permission.)

cellular damage by microprojectile bombardment, selection process, and growth of dying transformed cells for prolonged time during recovery, that cause chromosomal aberrations (Choi, Lemaux, and Cho, 2000a, 2001). Chromosomal aberrations are attributed to spindle failure by growth hormones that cause endoreduplication, c-mitosis, nuclear fragmentation (amitosis), multipolar configurations, and lagging chromosomes (Singh, 1993).

B. Genetic Background of the Explant

Chromosomal aberrations induced during culture are genotype dependent (Singh, 1993). Singh et al. (1998) observed that tissues and primary transformants from soybean genotype A2872 did not display chromosomal abnormalities (Table 10.4), while soybean genotype A2242 was cytologically highly unstable (Table 10.3). Nine selfed seeds from A2872 (control) showed 2n = 40 chromosomes with normal karyotype. Culture age on 2,4-D ranged from 8.26 to 32.30 months. Embryo suspensions, germinating embryos, and T₀ and T₁ plants expressed normal diploid-like phenotypes, and, as expected, all plants carried 2n = 40 chromosomes. However, five T₁ plants from a 29.43-month-old culture of 5–2 had tetraploid morphological features with one-seeded pods but showed 2n = 40 chromosomes (Table 10.4). These abnormalities may be genetic (desynaptic or asynaptic, male sterile and female fertile). These morphotypes often express tetraploid phenotypes (slow growth, dark-green leathery leaves, clustered flowers, empty pods) with partial to complete sterility, carrying one seed per pod (Palmer and Kilen, 1987). Interestingly, the primary transformants from A2872 contained 2n = 40 chromosomes though the culture was on 2,4-D for 32.30 months. This shows that a genotype may be highly responsive to culture conditions but may be prone to chromosomal aberrations (Hermsen, 1994).

Aragão et al. (2000) developed a soybean transformation protocol that produced a high frequency of fertile transgenic soybeans that was genotype independent. Their technique includes microparticle

Culture ID	R0 Phenotype	Origin of Roots	Months on 2,4-D	Number of Samples	2 <i>n</i>	Karyotype
A2872	Control			9	40	Normal
821	_	Suspension	8.26	1	40	Normal
821	_	Embryo	9.23	5	40	Normal
2–1	Diploid	T,	22.76	1	40	Normal
2–2	Diploid	T ₁	11.20	1	40	Normal
2–2	Diploid	T ₁	18.03	1	40	Normal
2-4	Diploid	T ₁	13.16	1	40	Normal
2–4	Diploid	T ₁	20.00	1	40	Normal
5–1	Diploid	T ₁	20.50	2	40	Normal
5–1	Diploid	T ₁	21.50	2	40	Normal
5–2	Diploid	T ₁	19.06	4	40	Normal
5–2	Diploid	T ₁	20.20	1	40	Normal
5–2	Diploid	T ₁	20.73	1	40	Normal
5–2	Diploid	T ₁	21.20	2	40	Normal
5–2	Diploid	T ₁	29.00	3	40	Normal
5–2	Diploid	T ₁	29.43	3	40	Normal
5–2	Tetraploid	T ₁	29.43	5	40	Normal
90–2	Diploid	T ₁	19.06	1	40	Normal
90–2	Diploid	T ₁	32.30	4	40	Normal
801	·	Embryo	9.23	2	40	Normal

Table 10.4 Chromosome Analysis at Somatic Metaphase in the Transgenic Asgrow Soybean Genotype A2872

Source: From Singh, R.J. et al., Theor. Appl. Genet., 96, 319-324, 1998. With permission.

bombardment (*ahas* gene: a selectable marker gene isolated from *Arabidopsis thaliana* that contains a mutation at position 653 bp) of the soybean meristematic region and culturing in selection medium with imazapyr herbicides followed by multiple shooting induction. They claimed a 200-fold increase in the recovery of transgenic soybean plants over the methods of Christou (1997).

C. Seed Fertility in the Transgenic Crops

Morphological variants, particularly seed sterility, in transgenic crops are attributed to chromosomal aberrations (Austin et al., 1995; El-Kharbotly et al., 1995; Fütterer and Potrykus, 1995; Lynch et al., 1995; Schulze et al., 1995; Shewry et al., 1995; Hadi, McMullen, and Kiner, 1996; Liu et al., 1996; Choi, Lemaux, and Cho, 2000a; Cho et al., 2002). *Agrobacterium*-mediated transformation produced diploid (2n = 24) (El-Kharbotly et al., 1995) and tetraploid (2n = 2x =48) potato (Conner et al., 1994). The diploid transgenic potato that conferred resistance to *Phytophthora infestans* inherited reduced male fertility, while female fertility was not so markedly influenced. All transformed tetraploid potatoes contained normal chromosome complement, but morphological changes including low yield and small tubers were observed in the field-grown plants compared to the control.

The *Agrobacterium*-mediated transformation has been extremely effective in dicotyledonous crops. However, Ishida et al. (1996) developed an *Agrobacterium*-mediated transformation protocol to produce transgenic maize in high frequency. They produced 120 morphologically normal transformants, and 70% of them set normal seeds. Plants were not examined cytologically.

Christey and Sinclair (1992) obtained kale, rape, and turnip transformants through *Agrobacterium rhizogenes*-mediated transformation. Transgenic plants were successfully produced from hairy roots. Morphological changes with an increase in leaf edge serration, leaf wrinkling, and plagiotropic roots were observed in some plants, while in other lines, phenotypic alterations were barely noticeable. Transgenic plants were not examined cytologically.

Toriyama et al. (1988) produced five transgenic rice plants after direct gene transfer into protoplasts through electroporation-mediated transformation. One plant was diploid (2n = 24), three

plants were triploid (2n = 3x = 36), and one plant was unidentified. Ghosh-Biswas et al. (1994) produced 73 transgenic rice cv. IR 43 plants by direct transfer of genes to protoplasts, and 29 plants reached maturity in the greenhouse. Eleven plants flowered but did not produce seed. However, two protoplast-derived nontransgenic plants set seeds. Protoplast-derived plants (transgenic and nontransgenic) had fewer tillers, narrower leaves, and were shorter in height than seed-derived plants. Fertile transgenic rice plants through electroporation-mediated transformation have been produced (Shimamoto et al., 1989; Xu and Li, 1994).

Another method used to insert foreign DNA into the protoplast is by using a polyethylene-glycol-mediated DNA transformation. Hall et al. (1996) reported a high-efficiency procedure for the generation of transgenic sugar beets from stomatal guard cells. They examined ploidy level by flow cytometry and recorded 75% diploid transformants. Seed production was normal, and the average frequency of germination was 96%. Lin et al. (1995) transferred 61 rice plants (via polyethyleneglycol-mediated transformation) to the greenhouse, and 28 of them were fertile. Seed set per plant ranged from 10 to 260. The cause of seed sterility was not determined.

Microprojectile-bombardment-mediated primary transformants (T_0) in soybean genotype A2242 exhibited a range of fertility that depended upon culture medium and duration of time on 2,4-D (Table 10.5). Chromosome counts of ten seeds from randomly selected T_0 plants revealed normal chromosomes (2n = 40) from the young cultures, regardless of seed set. Tetraploidy (2n = 80) and aneuploidy (near diploidy) were predominant in plants obtained from the older cultures (Table 10.5). Culture 828–2 after 11 months on 2,4-D produced 12 T_0 plants. Of the ten plants examined, five plants were diploid, and five plants were tetraploid. Seed fertility in nine plants was more than 101 seeds per plant, while seed set in three plants ranged from 21 to 100. Occasionally, seed set in T_0 diploid plants was low (2-12; 0-19; 7-36). Thus, seed sterility may be attributed to culture conditions (epigenetic), desynaptic and asynaptic gene mutations, or minor chromosomal deletions that cannot be detected cytologically. Schulze et al. (1995) observed fruit development after selfing transgenic cucumber, however, none of the harvested fruits contained seeds. Simmonds and Donaldson (2000) produced fertile transgenic soybeans from young proliferative cultures via particle bombardment, while sterile plants were recovered from 12- to 14-month-old cultures. The cause of sterility was not established. Choi, Lemaux, and Cho (2000a) recorded 75% (15/20 normal lines) fertile lines from microprojectile-bombardment-mediated transgenic oat, while 36% (10/28) lines had low seed set. Thus, transformation procedure and age of culture play an important role in producing completely fertile, stable, and normal transformants.

V. CYTOLOGICAL BASIS OF GENE SILENCING

The loss or low expression and unexpected segregation (gene silencing) of foreign genes are routinely observed in transformed crops (Fromm et al., 1990; Fütterer and Potrykus, 1995; Meyer, 1999; Senior, 1998; Kooter, Matzke, and Meyer, 1999). Foreign DNA in transgenic common bean plants, produced from particle bombardment, was not expressed in 12 (44%) plants, and two plants showed poor transmission of the inserted gene (1:10) in R_1 (T_1) generation, although all plants had normal phenotype and were fertile (Aragão et al., 1996). This may be due to chimaerism in T_0 plants.

An extensive review on genetic transformation and gene expression in the Poaceae by Fütterer and Potrykus (1995) reveals that the expressions of transgenes in the progeny of transgenic plants are quite unpredictable. The departure from the Mendelian inheritance (unstable transgenes) in transformants occurs when inserts are located in highly repetitive sequences (heterochromatic regions of chromosomes) or in the extra chromosomal DNA (mitochondrial or chloroplast genomes). Several genomic factors, such as aberrant crossing over during meiosis; spontaneous or induced mutations; ploidy, aneuploidy, sex chromosomes and transposable elements (Maessen, 1997); and nucleolar dominance (Pikaard, 1999) cause deviations in the Mendelian inheritance of the transgenes. Table 10.5 Culture Identification (ID), Plant ID, Months in Culture with 2,4-D before Transformation, Total Number of Transformants Recovered, Seed Set Range^a, and Chromosome Number in Selected Primary Transformants in Soybean Cultivar Asgrow 2242

Culture		Month	Total Number	Seed Set		2 <i>n</i>	
ID	Plant ID	2,4-D	Plants	Range	40	80	41
828–1	671–3–4	3.20	6	0–363	14	_	_
828–1	668-8-1	5.50	4	5-50	9	_	—
828–1	668–1–3	5.60	9	14–219	10	_	_
828–1	668–1–6	5.60	5	6–65	9	—	1
828–1	668–1–5	5.60	2	15–24	7	—	3
828–1	668–1–8	5.60	2	85–101	7	_	—
828–1	668–1–12	5.60	9	75–185	18	—	—
828–1	668–1–13	5.60	5	59–178	10	—	—
828–1	668–2–3	5.60	10	6–53	—	20	—
828–1	668–2–7	5.60	7	6–111	—	10	—
828–1	668–4–3	5.60	6	2–12	4	—	—
828–1	668–4–15	5.60	5	8–51	1	—	—
828–1	668-1-1	6.00	4	0–225	4	—	—
22–1	549–2–2	6.43	1	25	3	—	1
22–1	549-4-6	6.43	2	63–206	7	—	3
22–1	549–4–10	6.43	5	90–321	20	_	_
22–1	557-1-6	7.30	3	9–65	—	10	—
22–1	557–2–3	7.30	7	90-321	—	15	—
817	610–5–1	7.33	10	10–495	54	—	_
817	610–6–1	7.83	5	56-190	23	—	15
817	610–7–1	7.83	10	52-329	—	4	_
22–1	557–2–7	7.96	10	33–350	—	30	—
22–1	557–5–2	7.96	9	65–210	—	9	_
22–1	566–5–1	9.50	1	27	—	10	—
828–2	677–3–1	11.00	12	50-268	5	5	—
817	647–1–1	12.43	5	0–19	5	_	—
817	647–2–1	12.43	5	25–66	—	3	—
817	647–15–7	12.43	5	30–133	—	10	—
817	647–15–4	12.43	2	7–36	10	_	—
817	652–6–1	13.30	12	0–235	—	9	—
828–1	668–7–2	13.80	2	28–38	4	_	2
22–1	587–5–1	15.36	2	7–18	—	2	—
22–1	587–3–1	15.36	3	4–8	—	6	—
22–1	609–6–1	16.00	1	9	—	3	—

^a T.M. Klein, personal communication.

In transformation experiments, genes may be physically present, but gene activity may be poorly expressed or totally lost in subsequent generations. This phenomenon is known as co-suppression (Matzke and Matzke, 1995; Matzke et al., 2000; Stam, Mol, and Kooter, 1997). In cosuppression, foreign genes (transgenes) cause the silencing of endogenous plant genes if they are sufficiently homologous (Stam, Mol, and Kooter, 1997). An excellent example of co-suppression was shown in tobacco by Brandle et al. (1995). They produced a transgenic tobacco line carrying the mutant *A. thaliana* acetohydroxy acid synthase gene csr1-1 that expressed a high level of resistance to the sulfonylurea herbicide chlorsulfuron. The instability of herbicide resistance that was observed during subsequent field trials was not anticipated from the initial greenhouse screening. Hemizygous plants from this line were resistant, but homozygous plants (59%) were damaged by the herbicide. Damage was correlated with co-suppression of the csr1-1transgene and the endogenous tobacco AHAS genes, *sur A* and *sur B*. The difference in the performance of glyphosate tolerance in sugar beet transformants between greenhouse and the field were also reported by Mannerlöf et al. (1997). The disparity was attributed to differences between the environments, variation in gene expression caused by copy number, position, or

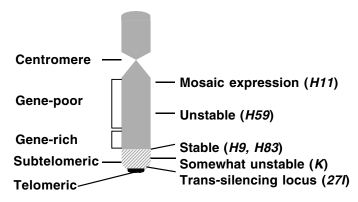


Figure 10.3 Influence of chromosomal location on transgene expression. The chromosomal locations of genetically well-characterized transgene loci (in italics, right) in tobacco as determined by FISH are shown relative to landmarks (left) on a model chromosome. (From Matzke, A.J.M. and Matzke, M.A., *Current Opinion Plant Biol.*, 1, 142–148, 1998. With permission.)

methylation effects. Based on this information, they suggested that co-suppression was triggered by agroclimatic conditions, and the initial greenhouse study was not predictable.

The stability of transformants is associated with the insertion of transgene(s) in the regions of chromosomes. Stable inserts have been located in the vicinity of telomere, as they have a tendency to integrate toward the distal end of chromosome arms which are gene-rich regions (Matzke and Matzke, 1998) (Figure 10.3). According to Matzke et al. (2000), different regions of the genomes vary in their ability to tolerate foreign DNA, resulting in erratic expression. Iglesias et al. (1997) demonstrated by fluorescence *in situ* hybridization (FISH) that two stably expressed inserts in tobacco were present in the vicinity of telomeres, while two unstably expressed inserts were located at intercalary and paracentromeric regions. In contrast, by using FISH in transgenic barley, Choi, Lemaux, and Cho (2002) recorded integration of insert was preferred distal positions on barley chromosomes: seven (37%) out of 19 lines examined had integration sites in distal positions of the chromosomes; four (21%) in telomeric regions, three (16%) in centromeric regions, three (16%) in satellite regions, and two (10%) had subtelomeric regions.

It has been demonstrated that gene silencing in transgenic crops is homology-dependent (Matzke et al., 2000). Expression of transgenes driven by barley endosperm-specific promoters was much more stable in its inheritance pattern than that driven by the constitutive promoters such as rice actin promoter and maize ubiquitin promoter (Cho et al., 1999, 2002; Choi, Lemaux, and Cho, 2002). The recipient plants often have effective defense mechanisms against the uptake, integration, and faithful maintenance of the foreign genes or their transcribed products. DNA methylation is widely associated with gene activation (Kumpatla et al., 1998).

Singh et al. (1998) presented a cytological clue that may help explain aberrant segregation ratios or loss of transgene sequences which may be applicable in some cases. For example, the selfed population of a plant with 2n = 39 + 1 metacentric chromosomes identified in soybean genotype A2242 was expected to segregate plants in a ratio of 1 (2n = 40):2 (2n = 39 + 1 metacentric):1 (2n = 38 + 2 metacentrics). Diploid plants will be normal and fertile and may not express the introgressed genes if this gene is in the deleted chromosomes. Matzke et al. (1994) attributed an erratic inheritance in a transgenic tobacco line to aneuploidy (2n = 49 or 50). Transgne silencing is common in diploid barley (M.-J. Cho, personal communication).

An early chromosome count prior to pretransformation of young cells, callus, and embryo suspension should be used as a transformation target to reduce the possibility of somaclonal variation and regenerability problems. Stability of inserts in a crop should be field tested in wide agro-eco-geo-climatic conditions (day length, drought, heat, irregular weather) for several years before releasing as a commercial cultivar.

APPENDIX 2-I

Sources of Chemicals

Catalog Price Product Company **Phone Number** Number Quantity (\$US) Slide Poly-Prep slides Sigma (800) 325-3010 P 0425 72 slides 31.00 dUTPs Biotin-16-dUTP Roche (800) 262-1640 1 093 070 50 nmol 263.00 Dig-11-dUTP Roche (800) 262-1640 1 558 706 125 nmol 573.00 Antibodies Fibers Green Detection (for Dig) FITC anti-dig (from Roche (800) 262-1640 1 207 741 200 µg 133.20 sheep) FITC anti-sheep Roche (800) 262-1640 605 340 1 ml 88.00 (rabbit) Fibers Red Detection (for Biotin) Texas Red SA-5006 65.00 Vector Labs (800) 227-6666 1 mg streptavidin Biotinylated anti-Vector Labs (800) 227-6666 BA-0500 500 µg 70.00 streptavidin Anti-fade Prolong Anti-fade Molecular (800) 438-2209 P-7481 Kit 135.00 probes Staining YoYo stain Molecular (800) 438-2209 Y-3601 200 uL 225.00 probes (1 mM)

Note: dUTPs come from Roche at 50 μL of 1mM. The working concentration should be 0.5 mM, with a dTTP:dUTP ratio of 2:1. Therefore, dilute dUTPs with 50 μL water, then mix with 200 μL 0.5 mM dTTP.

Two-Color Fiber-FISH

Product	Company	Phone Number	Catalog Number	Quantity	Price (\$US)
		Antibodies			
Fibers Green Detect	tion (for Biotin)				
FITC-Avidin Biotin-anti-avidin	Vector Labs Vector Labs	(800) 227–6666 (800) 227–6666	A–2001 BA-0300	5 mg 500 μg	70.00 60.00

One-Color Fiber-FISH

Source: From S.A. Jackson, R.M. Stupar, and J. Jiang. Personal communication.

APPENDIX 2-II

Materials for Flow Cytometry

- 1. Cell Trics (Partec): For filtration of the sample after chopping
- 2. Eye protection against UV
- 3. Gilson-pipettes and tips: 1000 μ L Gilson-pipette (blue tips) exclusively for PI; for nontoxic: 1000 μ L Gilson (blue tips) and an 100 μ L Gilson (yellow tips)
- 4. Flow tubes: Provided by Partec or from MERCK
- 5. Wash bottles: For distilled water
- 6. Distilled water: Glass distilled water
- 7. Water bath or dry incubator: Need to maintain 37°C for RNase digestion
- 8. Sample tubes: Partec
- 9. Mercury high-pressure lamp: Partec
- 10. Hypochlorite: Normal bleach
- 11. Immersion gel: Optical gel; R.P. Cargille Labs, Cedar Grove, NJ
- 12. Glycerine: Mix it with the immersion gel, when the gel becomes dry
- 13. Grease for laboratories: Small amount needed to maintain the rubber fitting between flow cytometer and sample tube
- 14. Normal razor blade: Gillette super silver
- 15. Pincers: For preparation of leaves
- 16. Small Petri dishes
- 17. Eppendorf tubes: 2 mL to store the frozen RNase

Source: From Obermayer, R., diploma thesis, 2000. With permission.

APPENDIX 2-III

Reagents and Solutions*

A. AGAROSE GEL, 1.5% (W/V)

- 3 g electrophoresis-grade agarose
- + 200 mL $1 \times TAE$ electrophoresis buffer
- Soak for 20 min at RT. Boil in a microwave oven to dissolve completely. Adjust volume to 200 mL with dH₂O. Pour gel into a casting form, and let solidify.

B. AMIPROPHOS-METHYL TREATMENT SOLUTION

- Prepare a 20 mM stock solution by dissolving 60.86 mg amiprophos-methyl in 10 mL cold acetone.
- Store up to 1 year at -20°C in 1 mL aliquots.
- Prepare the treatment solution immediately before use by combining amiprophos-methyl stock solution and $0.1 \times \text{or } 1 \times \text{Hoagland's nutrient solution specified below:}$

Preparation of Amiprophos-Methyl Treatment Solution

	Hoagland's Solution		20 mM Amiprophos-Methyl	
Species	Concentration	Volume (mL)	Amount (μL)	Final Concentration (µM)
Vicia faba	1 ×	750	94.8	2.50
Pisum sativum	1 ×	750	380.0	10.00
Hordeum vulgare	0.1 ×	750	94.8	2.50
Secale cereale	0.1 ×	750	19.0	0.50

C. 4',6-DIAMIDINO-2-PHENYLINDOLE (DAPI) STOCK SOLUTION (0.1 MG/ML)

- Dissolve 5 mg DAPI in 50 mL dH_2O by stirring for 60 min.
- Pass through a 0.22 μ m filter to remove small particles.
- Store up to 1 year at -20° C in 0.5 mL aliquots.

Caution! DAPI is a possible carcinogenic. It may be harmful if inhaled, swallowed, or absorbed through the skin, and may also cause irritation. Use gloves when handling. Be careful of particulate

^{*} For more information, see www.ueb.cas.cz/olomoucl.

dust when weighing out the dye. Consult local institutional safety officer for specific handling and disposal procedures.

D. ETHIDIUM BROMIDE SOLUTION, 0.5 MG/ML

- Dissolve 5 mg EB in 10 mL dH_2O by stirring for 60 min.
- Store up to 1 year at 4°C in the dark.
- For working solution (0.5 μ g/mL): dilute 1:1000 in dH₂O. The working solution can be used several times.

Caution! Ethidium bromide is a powerful mutagen and is moderately toxic. It may be harmful if inhaled, swallowed, or absorbed through the skin. Use gloves when handling. Be careful of particulate dust when weighing out the dye. Consult local institutional safety officer for specific handling and disposal procedures.

E. FORMALDEHYDE FIXATIVE

- 0.303 g Tris base (10 mM final)
- 0.931 g Na₂EDTA (10 mM final)
- 1.461 g NaCl (100 mM final)
- 250 µL Triton X-100 [0.1% (v/v) final]

Adjust volume to 200 mL with dH_2O , and adjust pH to 7.5 using 1 N NaOH. Add 37% (v/v) formaldehyde stock solution (37%, catalog no. 1.04003, Merck, Darmstadt, Germany). Adjust the final volume to 250 mL with dH_2O . Prepare the fixative before use as concentrations shown in the following table for several species.

Preparation of Formaldehyde Fixative

	Formaldehyde			
Species	Volume of Stock (mL)	Final (%) Concentration (v/v)		
Vicia faba	27.0	4		
Pisum sativum	20.0	3		
Hordeum vulgare	13.5	2		
Secale cereale	13.5	2		

Caution! Formaldehyde is toxic and is also a carcinogenic. It is readily absorbed through the skin and is irritating or destructive to the skin, eyes, mucous membranes, and upper respiratory tract. Wear gloves and safety glasses. Always work in a chemical fume hood. Consult local institutional safety officer for specific handling and disposal procedures.

F. FRUCTOSE SYRUP

- 30 g fructose
- 20 mL dH₂O
- Incubate at 37°C overnight
- Add one crystal of thymol
- Store up to 1 year at 4°C

G. GAMBORG'S B-5

• Basal medium with minimal organics (macro- and micronutrients and vitamins); G 5893, Sigma Chemical Co.

H. 5 N HCI

- 419.8 mL of 35% (v/v) HCl
- Adjust volume to 1000 mL with dH₂0

I. HOAGLAND'S SOLUTION

- Basal salt mixture (with macro- and micronutrients); H 2395, Sigma Chemical Co.
- · Prepare 500 mL according to manufacturer's direction
- · Autoclave, and store at RT

J. HYDROXYUREA TREATMENT SOLUTION

Prepare treatment solution immediately before use by combining hydroxyurea and $0.1 \times \text{or } 1 \times \text{Hoagland}$ nutrient solution shown below:

Preparation of Hydroxyurea Treatment Solution

	Hoagland's Nutrient Solution		Hydroxyurea	
Species	Concentration	Volume (mL)	Amount (mg)	Final Concentration (mM)
Vicia faba	1 ×	750	71.3	1.25
Pisum sativum	1 ×	750	71.3	1.25
Hordeum vulgare	0.1 ×	750	114.0	2.00
Secale cereale	0.1 ×	750	142.6	2.50

K. LB01 LYSIS BUFFER

- 0.363 g Tris base (15 mM)
- 0.149 g Na₂EDTA (2 mM)
- 0.035 g spermine 4HCl (0.5 mM)
- 1.193 g KCl (80 mM)
- 0.239 g NaCl (20 mM)
- 200 µL Triton X-100 (0.1%)

Adjust volume to 200 mL with dH₂O, and adjust final pH to 7.5 using 1 *N* HCl. Filter through a 0.22 μ m filter to remove small particles. Add 220 μ L β -mercaptoethanol, and mix well. Store up to 1 year at -20°C in 10 mL aliquots.

Caution! β -mercaptoethanol may be fatal if inhaled or absorbed through the skin and is harmful if swallowed. High concentrations are extremely destructive to the skin, eye, mucous membranes, and upper respiratory tract. Wear gloves and safety glasses and work in a chemical fume hood.

L. LB01 LYSIS BUFFER (FOR COLLECTING SORTED CHROMOSOMES)

- 0.545 g Tris (22.5 mM)
- 0.223 g Na₂ EDTA (3 mM)
- 0.052 g spermine 0.4 HCl (0.75 mM)
- 1.790 g KCl (120 mM)
- 0.351 g NaCl (30 mM)
- 300 µL Triton X-100 (0.15%)

Adjust to 200 mL with dH₂O, adjust final pH to 7.5 using 1 HCl. Filter through 0.22 μ m filter to remove small particles, add 330 μ L β -mercaptoethanol, and mix well. Store at -20°C in 5 mL aliquots.

M. LOADING BUFFER

- 2 mL 0.5 M EDTA, pH 8.0
- 100 mg SDS (1% w/v final)
- 5 mg bromphenol blue (0.05% w/v final)
- 5 mg xylene cyanol (0.05% w/v final)
- 5 mL 85% glycerol (42.5 v/v final)

Adjust volume to 10 mL with dH_2O , store up to 1 year at RT.

N. MAGNESIUM SULFATE STOCK SOLUTION

- Dissolve 1.23 g MgSO₄·7 H₂O in 50 mL dH₂O.
- Filter through a 0.22 µm filter to remove small particles.
- Store at 4°C.

O. MITHRAMYCINE STOCK SOLUTION

- Dissolve 50 mg mithramycine A in 50 mL dH₂O by stirring for 60 min.
- Filter through a 0.22 μ m filter to remove small particles.
- Store up to 1 year at -20° C in 0.5 mL aliquots.

Caution! Mithramycine is a possible carcinogen. It may be harmful if inhaled, swallowed, or absorbed through the skin. Use gloves when handling. Be careful of particulate dust when weighing out dye. Consult local institutional safety officer for specific handling and disposal procedures.

P. PCR PREMIX

- 5 μ L 10 × Taq DNA polymerase buffer (1 × final; buffer does not contain MgCl₂)
- 3 µL 25 mM MgCl₂ (1.5 mM final)
- 1 μ L 10 mM 4 dNTP mix (final 0.2 mM each dATP, dCTP, dGTP, dTTP)
- 1 μ L 50 μ M forward primer (1 μ M final)
- 1 μ L 50 μ M reverse primer (1 μ M final)
- 0.5 µL 5 U/µL Taq DNA polymerase (2.5 U/50 µL final)
- 18.5 μL sterile dH₂O

Mix well, and microcentrifuge briefly (5 to 10 sec at $2000 \times g$ at RT). Prepare on ice shortly before use.

Q. PRINS BUFFER

- 0.605 g Tris base (10 mM final)
- 1.864 g KCl (50 mM final)
- 0.203 g MgCl₂·6 H₂O (2 mM final)

Adjust volume to 500 mL with dH_2O , adjust pH to 8.0 using 1 N HCl, and store up to 6 months at 4°C.

R. PRINS REACTION MIX

- 5 μ L 10 \times Taq DNA polymerase buffer (1 \times final, containing 1.5 mM MgCl₂)
- $5 \ \mu L \ 25 \ mM \ MgCl_2$ (4 mM final)
- 2.5 µL 2 mM dCTP/dGTP (0.1 mM each final)
- + 2 μ L 0.2 mM fluorescein 12-dUTP (8 μ M final)
- + 2 μ L 0.2 mM fluorescein 15-dATP (8 μ M final)
- 4.25 μL 0.2 mM dTTP (17 μM final)
- 4.25 μL 0.2 mM dATP (17 μM final)
- + 5 μL 20 μM forward primer (2 μM final)
- 5 μ L 20 μ M reverse primer (2 μ M final)
- 0.6 μL 5 U/μL Taq DNA polymerase (3 U/50 μL final)

Adjust to 55 μ L with sterile dH₂O (includes 5 μ L for evaporation), prepare shortly before use. Actual composition of the mix should be optimized for a given primer pair and species.

S. PROPIDIUM IODIDE (PI) STOCK SOLUTION

- Dissolve 50 mg (1 mg/mL) PI in 50 mL dH_2O .
- Filter through a 0.22 μ m filter to remove small particles.
- Store at -20°C in 0.5 mL aliquots.

T. RNase STOCK SOLUTION

- Dissolve 25 mg (1 mg/mL) RNase in 25 mL dH₂O.
- Filter through a 0.22 μ m filter to remove small particles, and heat to 90°C for 15 min to inactivate DNase.
- Store at -20°C in 0.5 mL aliquots.

U. SHEATH FLUID SF 50

- 7.31 g NaCl (50 mM final)
- Adjust volume to 2500 mL with dH_2O .
- Sterilize by autoclaving.

V. STOP BUFFER FOR PRINS

- 2.923 g NaCl (0.5 M final)
- 1.861 g Na₂EDTA (0.05M final)

Adjust volume to 100 mL with dH_2O , adjust pH to 8.0 using concentrated NaOH, sterilize by autoclaving, and store up to 6 months at 4°C.

W. TAE ELECTROPHORESIS BUFFER, 50X AND 1X

- 242 g Tris base (2 M)
- 57.1 mL glacial acetic acid (1 M acetate final)
- 100 mL 0.5 M EDTA, pH 8.0 (50 mM final)

Adjust volume to 1 L with dH_2O , store up to 1 year at RT, dilute 1:50 for working solution in dH_2O before use. Final 1 × concentrations: 40 mM Tris, 20 mM acetate, and 1 mM EDTA.

Caution! Glacial acetic acid is volatile. Concentrated acids must be handled with great care. Wear gloves and safety glasses, and work in a chemical fume hood.

X. TRIS BUFFER

- 0.606 g Tris base (10 mM final)
- 1.871 g Na $_2$ EDTA (10 mM final)
- 2.922 g NaCl (100 mM final)

Adjust volume to 500 mL dH₂O, adjust pH to 7.5 using 1 N NaOH, and store up to 6 months at 4° C.

Y. WASH BUFFER FOR PRINS

- 1.161 g maleic acid (0.1 M final)
- 0.876 g NaCl (150 mM final)
- 0.5 µL Tween 20 (0.05% final)

Adjust volume to 100 mL with dH_2O , adjust pH to 7.5 using concentrated NaOH, sterilize by autoclaving, and store up to 4 months at 4°C.

APPENDIX 2-IV

Preparation of Media

Constituent	Medium A	Medium B	Medium C	Medium D
	Macron	utrients (g/L)		
KNO₃	30.00	25.00	20.00	0.80
(NH ₄) ₂ SO ₄	1.34	1.34	—	_
NH ₄ NO ₃	16.50	—	10.00	—
NaH ₂ PO ₄ ·H ₂ O	1.50	1.50	1.00	0.165
	1.70	_	3.00	_
MgSO ₄ ·7 H ₂ O	2.50	2.50	3.75	_
MgSO ₄ ·H ₂ O	_	_	_	5.76
Na ₂ SO ₄	_	_	_	2.00
CaCl ₂ ·2 H ₂ O	1.50	1.50	6.00	
Ca(NO ₃) ₂	_	_	_	4.32
	Micronu	trients (mg/L)		
			2.00	1 50
H ₃ BO ₃	3.00	3.00	3.00	1.50
MnSO ₄ ·H ₂ O	10.00	10.00	10.00	7.00
ZnSO ₄ ·7 H ₂ O	2.00	2.00	2.00	3.00
KI	0.75	0.75	0.75	0.75
CuSO ₄ ·5 H ₂ O	0.025	0.025	0.025	0.10
Na ₂ MoO ₄ ·2 H ₂ O	0.25	0.25	0.25	0.01
COCl₂·6 H₂O	0.025	0.025	0.025	_
KCI	—	—	—	65.00
EDTA	—	—	26.10	—
Na ₂ ·EDTA	37.30	—		
Fe·NaEDTA	—	73.40	—	73.40
FeSO ₄ ·7 H ₂ O	27.80	_	24.90	_
КОН	—	—	15.035	—
	Vitamins and	Hormones (m	g/L)	
Inositol	100.00	100.00	250.00	100.00
Nicotinic acid	1.00	1.00	1.00	1.00
Pyridoxine·HCl	1.00	1.00	1.00	1.00
Thiamine·HCI	10.00	10.00	10.00	1.00
Glycine	2.00	2.00	2.00	4.00
Ascorbic acid	100.00	50.00		
L-Glutamine	730.00			_
Coumarin	700.00	_	_	9.00
IAA ^a	0.20	0.30	—	9.00 1.00
NAA ^b		0.50		1.00
	2.00	0.75	_	_
Kinetin	1.28	0.75		_
BAP°	0.50	_	0.25	_

Constituent	Medium A	Medium B	Medium C	Medium D
Sucrose (g/L)	100.00	30.00	25.00	10.00
Bacto-agar (g/L)	8.00	8.00	6.00	8.00
pН	5.8	5.8	5.6	6.0

^a IAA = indol-3-acetic acid.

^b NAA = 1-naphthalene acetic acid.
^c BAP = 6-benzylaminopurine.

Source: From Singh et al., 1987. Theor. Appl. Genet., 74, 391–396. With permission.

APPENDIX 6-I

Common Name, Scientific Name, and 2*n* Chromosome Number of Some Important Plants

Common Name	Scientific Name	2 <i>n</i>
Alfalfa	Medicago sativa	16,32,64
Almond	Prunus amygdalus	16
Anise	Pimpinella anisum	18,20
Apple	Malus domestica	34
Apricot	Prunus armeniaca	16
Artichoke	Cynara scolymus	34
Ash (white)	Fraxinus americana	46,92,138
Ashoka	Saraca indica	24
Asparagus	Asparagus officinalis	20
Aster	Aster novaeangliae	10
Autumn crocus	Colchicum autumnale	38
Avocado	Persea americana	24
Babul	Acacia arabica	52
Bahiagrass	Paspalum notatum	20,40
Bamboo	Bambusa vulgaris	72
Banana	Musa sapientum	22,33,44
Banyan tree	Ficus benghalensis	26
Barley	Hordeum vulgare	14
Bean		
broad	Vicia faba	12
castor oil	Ricinus communis	20
lima	Phaseolus lunatus	22
mung	Vigna radiata	22
string (French)	Phaseolus vulgaris	22
Beet	Beta vulgaris	18
Begonia	Begonia rex	32,33,34,42,43,44
Bitter gourd	Momordica charantia	22
Black pepper	Piper nigrum	36,48,52,54,60,65,104,128
Blackberry	Rubus alleghaniensis	14
Blueberry	Vaccinium corymbosum	48
Bluegrass (Kentucky)	Poa pratensis	28–98
Bottle gourd	Lagenaria siceraria	22
Broccoli	Brassica oleracea var. Italica	18
Brussels sprouts	Brassica oleracea	18
Buckwheat	Fagopyrum esculentum	16
Cabbage	Brassica oleracea var. Capitata	18
Cantaloupe	Cucumis melo	24
Cardamom	Elettaria cardamomum	48,52
Carnation	Dianthus caryophyllus	30,90
Carrot	Daucus carota	18

Common Name	Scientific Name	2 <i>n</i>
Cashew nut	Anacardium occidentale	24,30,40,42
Cassava	Manihot esculenta	36,72
Cauliflower	Brassica oleracea var. Botrytis	18
Cedar (eastern red)	Juniperus virginiana	22,33
Celery	Apium graveolens	22
Cherry (sour)	Prunus cerasus	32
Cherry (sweet)	Prunus avium	16
Chestnut (European)	Castanea sativa	24
Chickpea	Cicer arietinum	16
Chive	Allium schoenoprasum	16,24,32
Chrysanthemum	Chrysanthemum morifolium	54
Cinnamon	Cinnamomum zeylanicum	24
Clove	Syzygium aromaticum	22
Clover (Ladino)	Trifolium repens	32
Clover (red)	Trifolium pratense	14
Cocaine plant	Erythroxylon coca	24
Cocoa	Theobroma cacao	20
Coconut	Cocos nucifera	32
Coffee	Coffea arabica	44
Coriander	Coriandrum sativum	22
Cotton (upland)	Gossypium hirsutum	52
Cotton wood	Populus deltoides	38
Cowpea	Vigna unguiculata	22
Crabapple	Pyrus ioensis (or Malus)	34
Cranberry	Vaccinium macrocarpon	24
Crocus	Crocus susianus	12
Cucumber	Cucumis sativus	14
Cumin	Cuminum cyminum	14
Daffodil	Narcissus pseudo-narcissus	14
Daisy	Bellis perennis	18
Date palm	Phoenix dactilifera	36
Dill	Anethum graveolens	22
Douglas fir	Pseudotsuga taxifolia	26
Eggplant	Solanum melongena	24
Fennel	Foeniculum vulgare	22
⁼ig	Ficus carica	26
Flax	Linum usitatissimum	30
Foxtail millet	Setaria italica	18
Garlic	Allium sativum	16
Geranium	Pelargonium graveolens	90
Ginger	Zingiber officinale	22
Gladiolus	Gladiolus communis	90,180
Grape	Vitis vinifera	38
Grapefruit	Citrus paradisi	18,27,36
Grasspea	Lathyrus sativus	14
Guar	Cyamopsis tetragonoloba	14
Guava	Psidium guajava	22
Guayule	Parthenium argentatum	54,72
Hazelnut	Corylus americana	28
Hemp	Cannabis sativum	20
Hops	Humulus lupulus	20
Hyacinth — brown	Hyacinthus orientalis	16
ndian mustard (red)	Brassica juncea	36
ris (blue flag)		72,84,105
	Iris versicolor	72,04,100
vy	Iris versicolor Hedera helix	48
vy Iasmine		
•	Hedera helix	48
lasmine	Hedera helix Jasminum officinale	48 26

Common Name	Scientific Name	2 <i>n</i>
Kiwifruit	Actinidia chinensis	58
Lemon	Citrus limon	18,36
_entil	Lens culinaris	14
_ettuce	Lactuca sativa	18
_ily	Lilium longiflorum	24
_ime (acid)	Citrus aurantifolia	18
₋ingonberry	Vaccinium vitis-idaea	24
_otus (yellow)	Nelumbo lutea	16
_uffa	Luffa cylindrica	26
_upin	Lupinus luteus	48
Macaroni wheat	Triticum turgidum var. durum	28
Vahogany	Swietenia mahaqoni	46-48
Mahwah	Bassia latifolia	24
Vaize	Zea mays	20
Vango	Mangifera indica	40
Maple (sugar)	Acer saccharum	26
Marigold	Tagets erecta	24
Villet	lagelo electa	LT
Foxtail	Setaria italica	18
pearl	Pennisetum glaucum	14
broomcorn	Panicum miliaceum	36
sawa	Echinochloa colona	54
Jilkweed		22
	Asclepias incarnata	
Anning glory	Ipomoea purpurea	30
/luskmelon	Cucumis melo	24
	Myristica fragrans	44
Dak (pedunculate)	Quercus robur	24
Dak (white)	Quercus alba	24
Dat	Avena sativa	42
Dil palm	Elaeis guineensis	32
Okra	Hibiscus esculentus	72,144
Dlive	Olea europaea	46
Dnion	Allium cepa	16
Drange (bitter)	Citrus aurantium	18
Drange (sweet)	Citrus sinensis	18,27,36,45
Papaya	Carica papaya	18
Parsley	Petroselinum crispum	22
Parsnip	Pastinaca sativa	22
Pea	Pisum sativum	14
Peach	Prunus persica	16
Peanut (groundnut)	Arachis hypogaea	40
Pear	Pyrus communis	34
Pecan	Carya illinoensis	32
Peepal tree	Ficus religiosa	26
Pepper	Capsicum annuum	24
Peppermint	Mentha piperita	36,64,66,68,70
Petunia	Petunia hybrida	14
Pigeon pea	Cajanus cajan	22
Pine (red)	Pinus spp.	24
Pineapple	Ananas comosus	50
Pistachio nut	Pistacia vera	30
Plum (European)	Prunus domestica	48
Poinsettia	Euphorbia pulcherrima	28
Poppy (opium)	Papaver somniferum	22
Potato	Solanum tuberosum	48
Pumpkin (summer squash)	Cucurbita pepo	40
Radish	Raphanus sativus	18
	Ambrosia trifida	24
Ragweed	Amprosia imiga	24

Common Name	Scientific Name	2 <i>n</i>
Ragi, Finger millet	Eleusine coracana	36
Redbud	Cercis canadensis	12
Rhubarb	Rheum officinale	22,44
Rice	Oryza sativa	24
Rose	Rosa centifilia	28
Rose	Rosa damascena	28
Rose	Rosa multiflora	14
Rubber	Hevea brasiliensis	36
Rye	Secale cereale	14
Safflower	Carthamus tinctorius	24
Sandalwood (Indian)	Santalum album	20
Sequoia (big tree)	Sequoia gigantia	22(44)
Sesame	Sesamum indicum	26
Snapdragon	Antirrhinum majus	16
Sorghum	Sorghum bicolor	20
Soybean	Glycine max	40
Spearmint	Mentha spicata	36,48
Spinach	Spinacia oleracea	12
Squash (winter)	Cucurbita maxima	40
Sugarcane	Saccharum officinarum	80
Sunflower	Helianthus annuus	34
Sweet potato	Ipomoea batatas	60,90
Sycamore	Acer pseudoplatanus	52
Tamarind	Tamarindus indica	24
Tangerine	Citrus nobilis	18
Tea	Camellia sinensis	30
Teak	Tectona grandis	24
Thyme	Thymus vulgaris	30
Timothy	Phleum pratense	42
Tobacco	Nicotiana tabacum	42
Tomato	Lycopersicon esculentum	24
Tulip		24
Turmeric	Tulipa gesneriana	64
	Curcuma longa	20
Turnip	Brassica rapa	32
Vanilla	Vanilla planifolia	
Violet (African)	Saintpaulia ionantha	28
Walnut (English)	Juglans regia	32
Watermelon	Citrullus vulgaris	22
Wax gourd	Benincasa hispida	24
Wheat	Triticum aestivum	42
Wild rice	Zizania aquatica	30
Winged bean	Psophocarpus tetragonolobus	18
Yam	Dioscorea alata	20,30,40,50,60,70,80
Zinnia	Zinnia elegans	24

Source: From Darlington, C.D. and Wylie, A.P., *Chromosome Atlas of Flowering Plants*, George Allen & Unwin Ltd., London, 1955. With permission.

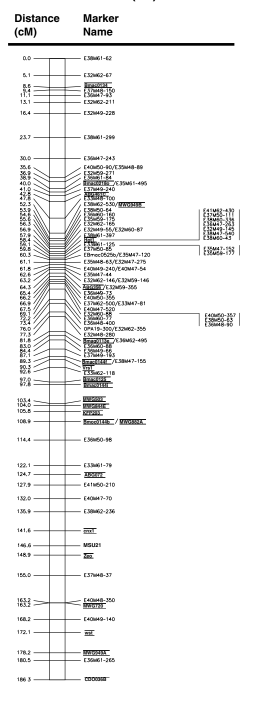
APPENDIX 7-I

Linkage Map of Barley

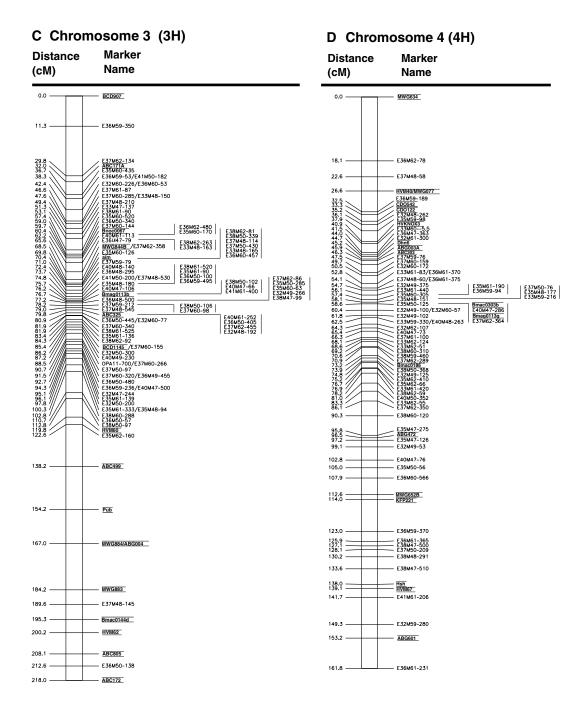
Distance (cM)	Marker Name	r	
0.0 \	, ABG704		
0.0 6.1 7.8 11.5 16.1 16.7	E37M50-88 E35M62-150 E41M62-151 E37M50-363		
16.1	E41M62-151 E37M50-363		
16.7	////, E32M48-144, E37M49-164	/E40M50-150	
20.1	E37M49-300		
22.5	Е 36М49-355, Е 36М60-157	/E41M61-350	
17.8 20.1 22.0 22.5 24.6 25.4 26.2 27.0	E37M50-363 E32M48-144, E37M49-164 E37M49-300 E32M47-67 E36M49-355, E36M60-157 E41M62-140 E36M61-121, MGC300 E36M61-121, E36M61-121, E36M61-121, E36M61-104	(570) (67)	
27.0 29.2 29.2	E36M61-121, ABG380	/E36M62-134	
31.1			
33.4 35.0	E38M61-121, E40M48-193 E37M61-66/E	/E35M61-255	
36.0	Е 37м61-66/Е	E33M48-265	
36.9	E38M59-69/8	E38M61-193 E36M62-95	
39.3 39.9 40.5 41.7 43.5 44.1 45.6	E35M59-6970 E37M61-9870 E35M48-113 E35M48-113 E35M59-217 E35M59-144 E35M59-144 E35M59-144 E35M59-243 E38M59-243 E38M59-243 E38M59-243 E41M62-142		
40.5	E35M59-217 E36M59-144		
43.5	E35M62-52 E40M48-301 E38M59-243		
45.0	E33M601 E41M62-142		
46.5	C 200049=370	/E30M39=181	
48.2	E38M62-165, E33M61-190, E35M59-74	/E38M61-118 /E38M62-192	
51.1 52.1	E 35M59-74 E 38M47-77/E	E32M48-124	
535	E 38M59-71/8	E37M59-490	
54.0 55.8 56.6	E35M59-35 E35M61-166	E35M62-400 E35M62-100	
58.9	E38M48-82	E38M48-308	
61.1 62.6	E36M59-274 Bmac0187	E41M62-270 E37M59-515	
63.9	E40M50-227 E41M62-202	E36M47-68 E32M47-62 E36M47-306	
67.6	E35M48-390	C75147 07	
54 0 556 9 566 9 567 1 62 6 65 9 67 6 6 65 9 67 6 6 71 4 71 4 73 5 2	E37M47-510 E36M59-286	E32M49-124 E37M59-490 E35M62-100 E33M48-107 E33M48-107 E33M48-107 E33M48-107 E33M48-107 E33M47-62 E35M47-62 E35M47-62 E35M47-62 E35M47-62 E35M47-62 E35M47-62 E35M47-62 E35M47-62 E35M48-70 E35M48-7	
73.6 75.2	E36M61-410 E32M49-239	E36M48-66	
76.1 78.6	E40M61-106, E36M49-210,	/E40M50-225 E36M60-471 /E32M62-58 E32M61-144 E32M62-189	
80 3	E35M48-86/E 0PA7-850 E32M49-214 Bmac0273	ZE32M62-58 E32M61-144 E32M62-188 E32M62-188 E35M48-285 E35M48-285 E35M48-285 E35M48-285 E35M50-167 E35M50-167 E35M50-167 E35M50-2191 E35M	1
81.5	E32M49-214 Bmac0273	E36M50-167 E38M60-209	
80.9 81.5 82.0 82.7 83.9	Bmac047B	E38M61-540 	E İ
85.2 //////	Bmac0303A	E38M61-540 E38M49-177 /E36M60-57 E38M61-78 E36M61-286 E36M61-286	
88.8 //////	E40M61-104 E32M61-55 E38M47-166	E36M61-286	
90.8 91.5			
027///////	E41M50-262, E36M49-550,	/E36M60-418 /E36M50-65	
93.2 /////	E35M60-117, E38M48-254	E36M47-250 E41M50-360 E36M50-240	
95.4 ////	E37M49-114 E32M49-380	E33M61-156 E40M61-290	
93.2 94.1 95.4 98.3 98.8 99.7 100.7	E38M48-1254 E38M48-1254 E37M49-114 E32M49-380 Nud E36M50-308 E36M60-370	/E37N47-500	
104.4 ////			
100.3 110.9 113.0 114.4 118.7 120.1 121.6 126.4		M60-385	
114.4	E37M60-390 F35M47-128		
120.1	E33M48-65 WG380B		
	E37M59-147 Bis44 /E37	E37M61-146 E35M61-377	
130.7	E38M50-77 Lks2 /(37) E33M48-84 E37M60-390 E33M48-65 WG3806 E33M59-147 Ris44 /(23) E41M50-270 E41M50-270	35M50-137	
139.3	Е 36м59-153		
144.7		E32M61-176	
145.6	E41M50-130		
151.9 -	- F.32M47-355		
151.9 153.0 154.0 155.3	E32M47-355 E38M62-84 E37M62-90		
155.3	ABG461A		
160.7		E41M61-185	
163.8	F 33M47-135		
165.6	Е 36м59-76		
185.3	E35M50-137		
189.2	E35M61-377		
191.9	<u>Tha1</u>		

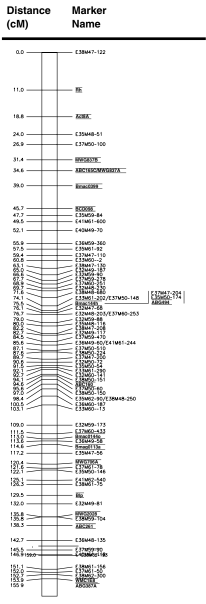
A Chromosome 1 (7H)

B Chromosome 2 (2H)

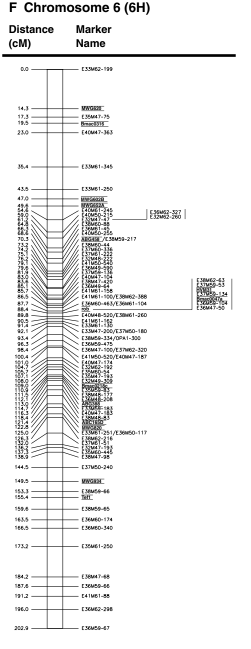


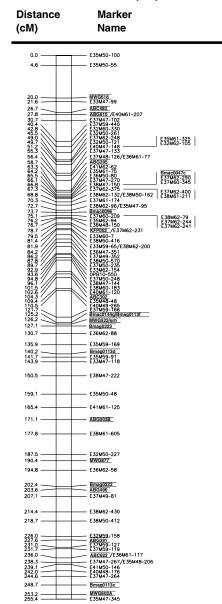






E Chromosome 5 (1H)





G Chromosome 7 (5H)

(From Costa, J.M. et al., 2001. Theor. Appl. Genet., 103: 415-424. With permission.)

Glossary*

- **Abscission layer** Cells of abscission zone disintegrate, causing the separation of plant organs (leaf, fruit, pod, branch).
- Abscission zone Layer of cells that brings about the abscission of plant organs.

Accessory chromosome Supernumerary or B-chromosome.

- Acentric fragment A chromosome fragment without a kinetochore.
- Achene Seed and pericarp attached only at the funicules, the seed usually tightly enclosed by the fruit wall, as in the sunflower and buckwheat.
- Achiasmate Meiosis without chiasma. Chromosome 4 in *Drosophila*, which is mostly heterochromatic, is without chiasma. Heterochromatin has been implicated in chromosome pairing for achiasmatic disjunction.
- Achromatic Chromosome regions that do not stain with specific dyes.
- A-chromosome One of the standard chromosomes of an organism.
- Acquired characters Changes in phenotype of an organism arise purely by environmental influences during development and are not the result of gene action (Lamarckian inheritance).
- Acrocentric chromosome A chromosome consisting of one complete arm and a small piece of the other arm.
- Acrotrisomics An individual with a normal chromosome complement plus an extra acrocentric chromosome.
- Actinomorphic Sepals and petals radiating from the center of the flower; the sepals are all similar to one another, as are the petals to one another.
- **Adaptation** The process by which an organism undergoes modifications (developmental, behavioral, anatomical, physiological) in such a way to survive perfectly in a given environment.
- **Additive** Quantitative effects of genes added together to produce phenotypes that are the sum total of the negative and positive effects of the individual polygenes.
- Adjacent-1 distribution In an interchange, heterozygote *nonhomologous* kinetochores move to opposite poles.
- Adjacent-2 distribution In an interchange, heterozygote *homologous* chromosomes move to opposite poles.
- Adventitious embryony (adventive embryony) Embryo develops from the nucellus or integument cells of the ovule.
- Adventitious roots Formation of the roots from stem nodes of a plant.
- Agamospermy See Apomixis.
- Albuminous seed A mature seed that contains endosperm.
- Aleurone One or more outermost layers of endosperm that store substantial quantities of protein, e.g., bran.
- Alien addition line A line containing an unaltered chromosome complement of one species (generally a cultigen) plus a single chromosome (monosomic alien addition lines, MAALs) or a pair of chromosomes (disomic alien addition lines, DAALs) from an alien species.

Alien chromosome A chromosome from a related species.

Allele An allele is merely a particular form of the same gene, can be dominant or recessive, and can be in homozygous and heterozygous conditions.

* Allard, 1960; Chapman and Peat, 1992; Darlington and Mather, 1949; Fahn, 1990; Fehr, 1987; Lewin, 1990; Nelson, Robinson, and Boolootian, 1970; Rieger, Michaelis, and Green, 1976; Smith, 1977; http://linkage.rockefeller.edu.

Allelic Two genes that are similar — the same gene.

Allelopathic The populations are not in physical contact and live in different geographic areas. **Allogamy** Cross-fertilization.

- **Allopolyploid** An organism containing different sets of chromosomes derived from two or more genomically diverse species, for example, allotetrapliod, allohexaploid, and allooctoploid.
- **Allosyndesis** Meiotic chromosome pairing of completely or partially homologous (homoeologous) chromosomes in allopolyploids.

Alternate leaf A plant with one leaf at a node.

- **Alternation of generation** In the life cycles of many plants, the gametophyte phase alternates with a sporophyte phase.
- Ameiotic mutants Meiosis does not ensue, and pollen mother cells degenerate.
- Amitosis Absence of mitosis.
- **Amphidiploid** An individual that contains entire somatic chromosome sets of two species produced after chromosome doubling and exhibits diploid-like meiosis.
- **Amphimixis** Seed develops by fusion of a female gamete and a male gamete normal sexual reproduction.
- **Amplification** The production of additional copies of a chromosome sequence, found as intrachromosomal or extrachromosomal DNA.
- **Anaphase (mitosis)** Anaphase is the shortest stage of mitosis, mutual attachment between homologous chromosome is ceased, sister chromatid at the kinetochore region at metaphase repulse, and two chromatids, with an equal distribution of chromosomes, move toward the opposite poles at the spindle.

Anaphase-I (meiosis) Migration of homologous chromosomes to opposite poles.

- Androecium A term that refers to all stamens of a flower; a male reproductive organ of a plant.
- **Androgenesis** Synonymous with male parthenogenesis. The progeny inherits the genotype of the male gamete nucleus.
- **Androgynophore** A stalk-like elongation of the floral axis between the perianth and stamens, which elevates the androecium and gynoecium.

Anemophily Wind pollination.

- **Aneuhaploid** The chromosome number in aneuhaploid is not an exact multiple of the basic set, but an individual is deficient for one or more chromosome(s).
- **Aneuploid** An organism or species containing chromosomes other than an exact multiple of the basic (*x*) chromosome number.
- Aneusomatic An organism having euploid and aneuploid cells.
- Annealing The pairing of complementary single strands of DNA to form a double helix.

Annual A plant completing the life cycle within a year.

- Anther The part of the stamen bearing pollen grains.
- Anthesis The time of flowering in a plant.
- **Antipodal cells** A group of three cells, each with a haploid nucleus formed during postmegasporogenesis, located at the chalazal end of the mature embryo sac.
- **Aperture** This term is for the pollen grain. A depressed area of characteristic shape in the wall; the pollen tube emerges via such an area.

Apex The terminal portion of the shoot or root in which the apical meristem is located.

Apocarpy The carpels are free in a flower and ovary.

Apogamety Embryo is formed from synergids and antipodals other than the egg without fertilization.

- **Apomeiosis** A type of gametophytic apomixis, where failure of chromosome reduction and of recombination occurs.
- **Apomict** An organism produced by apomixis.
- **Apomixis** Apomixis is a mode of asexual reproduction in plants, where seed is produced without sexual fusion of female and male gametes. Synonymous with agamospermy.
- Apospory Embryo sac is formed from somatic cells in the nucellus or chalaza of the ovule.
- **Asexual reproduction** A type of reproduction that does not involve the union of male and female gametes; vegetative propagation.
- **Astral rays** Distinctive lines forming a network between the two centrioles. It is common in animal cells.

- **Asymmetric** Nonadditivity and interactions of various kinds will cause a normal distribution curve to appear skewed or asymmetrical.
- Asymmetric cell division Division of a cell into two cells of different chromosome numbers.

Asymmetrical karyotype Characterized by the set containing some large and some small chromosomes.

Asynapsis A complete failure of homologous chromosome pairing during meiosis.

Autoallopolyploid It is confined to hexaploidy or higher levels of polyploidy.

- **Autogamy** Self-pollination predominates, and it is a common and widespread condition in angiosperms, particularly in annual herbs.
- **Automixis** Fusion of two haploid nuclei in a meiotic embryo sac without fertilization of the two egg nucleus producing maternal-like progeny.

Autonomous apomixis No dependence on pollination, e.g., Teraxacum and Hieracium.

- Autopolyploid An individual containing three or more genomically identical sets of chromosomes.
- **Autoradiography** This method detects radioactivity-labeled molecules by their effects in creating an image on photographic film.
- **Autosegregation** Spontaneous changes in genotype in the egg cells that develop without fertilization; may occur in obligate as well as facultative apomixis.

Autosome Any chromosome other than sex (X or Y) chromosome.

Autotetraploid (4x, AAAA) An individual possessing four identical sets of homologous chromosomes.

Autotriploid (3*x*, **AAA**) An individual possesses three identical sets of homologous chromosomes. **Autotroph** An organism that synthesizes its own food materials.

- **Auxin** Hormones that promote cell elongation, callus formation, fruit production, and other functions in plants.
- Auxotroph An organism that synthesizes most of its cellular constituents but ingests others.

Avirulent Parasite unable to infect and cause disease in a host plant.

B–A translocation These interchanges are obtained by breakage in the A chromosomes and in the B chromosomes and after reciprocal exchanges A^B and B^A chromosomes are produced.

Back cross The crossing of an F_1 heterozygote to one of its parents.

Back mutation Restores wild type.

Bacteria Bacteria are single-cell, microscopically small organisms with simple structures that usually reproduce asexually with very short generation times (20 min). A single bacterium produces a clone that is genetically identical. Accidental changes (mutation) lead to a new type of clone. They may be autotrophs (lost the ability to produce, usually, amino acids) or prototrophs (can produce these nutrients).

Bacteriophage Any virus that infects bacteria.

Bacterium A microscopic, single-celled prokaryotic organism.

Balanced lethal A situation in which only certain combinations of gametes survive in zygotes. For example, heterozygous velans/gaudens survive, and homozygotes die, in the *Oenothera lamarkiana*.Balbiani ring An extremely large puff at a band of a polytene chromosome.

Base pairs A partnership of A (adenine) with T (thymine) or of C (cytosine) with G (guanine) in a DNA double helix. In a RNA, thymine is replaced by uracil (A = U; C = G).

Base sequence The order of nucleotide bases in a DNA molecule.

Base sequence analysis A technique, often automated, for determining the base sequence.

Base substitution A form of mutation in which one of the bases in DNA replaces another.

Basic chromosome number The haploid chromosome number of the ancestral species represented as *x* (basic genome). For example, hexaploid wheat is identified as 2n = 6x = 42 (AABBDD).

Bidirectional replication When two replication forks move away from the same origin in different directions.

Biennial An individual living for two years, typically flowering and fruiting the second year.

Biotechnology The set of biotechnology methods developed by basic research and now applied to research (recombinant DNA technology) and product development.

Biotype A group of individuals having the same genotype, and may be homozygous or heterozygous. **Bipartitioning** Normal meiosis.

Bisexual (= perfect) A flower containing stamens and carpels.

Bivalent A pair of homologous chromosomes synapsed during meiosis.

- **bp** An abbreviation of base pairs; distance along DNA is measured in bp.
- **C-banding** Constitutive heterochromatin, exhibits bands after staining with Giemsa, is redundant, and is usually present in the proximity of kinetochore, telomere, and in the nucleolus organizer region.
- **C-mitosis** Complete inactivation of spindles producing a restitution nucleus with double chromosome number by colchicine treatment.
- C-value The total amount of DNA in a haploid genome.
- Caducous Plant parts that fall off early or prematurely.
- **Callose** A polysaccharide (β -1,3 glucan) present in sieve areas, walls of the pollen tubes.
- **Callus** A disorganized mass of undifferentiated plant cells obtained through culture from small pieces of plant tissues.
- Carpel Pistil; includes ovary, style, and stigma.
- Caryopsis A characteristic feature of plants of the Gramineae in which the pericarp and testa are fused.
- Cell The smallest, membrane-bound, living unit of biological structure capable of autoreduplication.
- **Cell culture** A process of culturing and growing cells through *in vitro* and regenerating plants by several subcultures.
- **Cell cycle** The period from one division to the next consists of G_1 , S, G_2 , and M.
- **Cell furrow** The wall formed between the two daughter nuclei at the end of telophase in animals.
- Cell plate The wall formed between the two daughter nuclei after karyokinesis.
- CentiMorgan (cM) A unit of measure of recombination frequency.
- Centric A chromosome segment having kinetochore.
- **Centriole** Small hollow cylinders consisting of microtubules that become located near the poles during mitosis.
- Centromere See kinetochore.
- **Centromeric index** (CI) = short arm length/chromosome length \times 100.
- **Chalaza** The region on an ovule where the nucellus and integuments connect with the funiculus; the opposite end to the micropyle.
- **Character** Phenotypic trait of an organism controlled by genes that may be dominant or recessive and may be governed by a single gene or multigenes.
- **Chasmogamy** Fertilization after the opening of a flower.
- **Chiasma** An exchange of chromatid segments during the first division of meiosis that results in the genetic crossing over (plural: chiasmata).
- **Chimaera** A plant composed of two or more genetically distinct types of tissues due to mutation, segregation, or irregularity of mitosis (mosaic).
- **Chlamydomonas** A unicellular haploid green alga (*Chlamydomonas reinhardi*) that contains two motile flagella and a small distinct nucleus with haploid chromosomes and numerous mitochondria. Chloroplast, constituting 50% of the cell, helps to nurture phototrophically in light, utilizing CO_2 as its carbon source. However, chlamydomonas can grow heterotrophically on an acetate culture without light. The asexual cycle takes about 6 h, and sexual cycles take less than 2 weeks.
- **Chlorenchyma** A chloroplast-containing parenchyma tissue such as the mesophyll and other green tissues.
- Chlorophyll The major photosynthetic green pigment present in the chloroplast.
- **Chloroplast** A cell organelle characterized by containing chlorophyll and enzymes required for photosynthesis. Chloroplast contains DNA and is capable of self-replication and protein synthesis.
- **Chromatid** A half-chromosome (sister chromatids) that originates from longitudinal splitting of a chromosome during mitotic anaphase or meiotic anaphase-II.

Chromatid bridge Dicentric chromatid(s) with kinetochores passing to opposite poles at anaphase. **Chromatid segregation** Double reduction, used in autopolyploidy.

- **Chromatin** The complex of DNA and protein in the nucleus of the interphase cell; originally recognized by reaction with stains specific for DNA.
- **Chromocenters** Darkly stained bodies in the interphase nuclei consisting of fused heterochromatic telomeres.

- **Chromomeres** Knob-like regions along the entire length of pachytene chromosomes that occupy specific positions on the chromosomes; larger heterochromatic chromomeres stain relatively darker than the euchromatic chromomeres.
- Chromonema The chromosome threads at leptonema.
- **Chromosomal inheritance** The inheritance of genetic traits located in the chromosomes (Mendelian inheritance).
- **Chromosomal sterility** Sterility in F_1 plants is caused by the lack of homology between the parental chromosomes.
- **Chromosome** An important, complex organelle of the nucleus, containing genes. The number is maintained constant from generation to generation by mitotic and meiotic divisions; it consists of protein, nucleic acid, and genes.
- **Chromosome arm** One of the two parts of a chromosome divided by an intercalary centromere.

Chromosome complement The total chromosome constitution of an organism.

- **Chromosome configurations** An association of chromosomes during meiosis in the form of univalent, bivalent, trivalent, quadrivalent (tetravalent), and so on.
- **Chromosome congression** Chromosomes move to the equatorial plate halfway between two poles of the spindle with the help of spindle fibers.

Chromosome drift The shift in the chromosome number.

- **Chromosome elimination** The loss of chromosomes from nuclei due to genic, chromosomal, or genomic causes.
- Chromosome fragmentation The breakage of chromosomes by radiation.
- **Chromosome map** A linear representation of a chromosome in which genetic markers belonging to particular linkage group are plotted based on their relative distances physical mapping. Chromosome maps are: (1) cytological maps according to karyotype of chromosomes; (2) genetic maps according to crossing over, recombination, and map units; (3) cytogenetic maps gene located in a particular chromosome based on cytologically identifiable chromosome aneuploids, such as deletions, duplications, translocations, inversions, trisomics, and other chromosomal aberrations; and (4) restriction fragment length polymorphism (RFLP) map a linear graph of sites on DNA cleaved by various restriction enzymes.
- **Chromosome set** A minimum complement of chromosomes derived from the gametic complement of a putative ancestor (basic chromosome).
- **Chromosome walking** Describes the sequential isolation of clones carrying overlapping sequences of DNA, allowing regions of the chromosome to be spanned. Walking is often conducted in order to reach a particular locus.
- **Class intervals** Collection of observations into groups. Class intervals may be large or small depending upon the need of the investigators.
- Cleavage Division of cytoplasmic portion in animals.
- **Cleistogamous flower** Pollination and fertilization occur before flowers open, insuring complete self-pollination.
- Climbing Ascending upon other plants.
- Clone An organism propagated asexually, vegetatively, from the same individual.
- **Cloning vector** A plasmid or phage that is used to carry inserted foreign DNA for the purpose of producing more material or a protein product.
- **Coding strand** DNA has the same sequence as mRNA.
- Codominance Genes when both alleles of a pair are fully expressed in the heterozygote.

Codon A triplet of nucleotides that represents an amino acid.

- **Coefficient of variation (c.v.)** It is defined as percent c.v. = $(s/\bar{x}) \times 100$; which is standard deviation, and 0 is mean in a single sample.
- Coenocyte A multinucleate cell resulting from the repeated division of nuclei without cytokinesis.

Cohort The group of individuals that enters the breeding program in the same season.

Colchicine An alkaloid chemical that induces polyploidy by inhibiting the formation of spindles, delaying the division of kinetochore.

Colchiploidy Polyploidy induced by colchicine.

Coleoptile A sheath surrounding the apical meristem and leaf primordia of the grass family.

Coleorrhiza A sheath of tissues surrounding the radicle of the grass embryo.

Compatible A congenial interaction between a parasite and its host in plant disease.

- **Compensating diploids** One normal chromosome that is replaced by two telocentric chromosomes representing both arms or four telocentric chromosomes that compensate for a pair of normal homologues.
- **Compensating trisomics** A missing normal chromosome is replaced by two tertiary chromosomes or by a secondary and a tertiary chromosome.
- **Compilospecies** A genetically aggressive plundering related species of their heredities, and in some cases, it may completely assimilate a species, causing extinction.
- **Complementary genes** Two nonallelic gene pairs complement each other in their effect on the same trait. For example, purple-flowered F_1 plants are produced by crossing two white-flowered parents in sweet pea. Selfing of F_1 produces nine purple and seven white; purple arose as the complementary effect of dominant alleles at two different gene pairs.
- **Composite** A mixture of several genotypes maintained in bulk from one generation to the next.
- **Concerted evolution** The ability of two related genes to evolve together as though constituting a single locus.
- **Configuration** An association of two or more chromosomes at meiosis and segregating independently of other associations at anaphase.
- **Constitutive heterochromatin** It is differentiated by darker staining during interphase and prophase. In most mammals, constitutive heterochromatin is located in the proximity of the kinetochore. It is an inert state of permanently nonexpressed sequences, usually highly repetitive DNA.
- Constitutive mutation The gene that is usually regulated to be expressed without regulation.
- **Continuous spindle fibers** Connect the two polar regions with each other and remain from early prometaphase to early prophase.
- **Controlling elements** Originally found and identified in maize by their genetic property, they are transposable units. They may be autonomous (transpose independently) or nonautonomous (transpose only in the presence of an autonomous element).
- **Convergent coorientation** In a trivalent, two chromosomes face one pole, while a third orients to the other pole.
- Coorientation Orientation of kinetochores at metaphase-I.
- **Cotyledon** One of the two halves of a dicot seed; the cotyledons develop into the first two leaves of the young dicot plants.
- **Coupling** When linked recessive alleles (ab) are present in one homologous chromosome and the other homologous chromosome carries the dominant allele (AB). In the case of repulsion, each homologue contains a dominant and a recessive gene (Ab, aB).
- Crossbreeding Outbreeding.
- **Crossing over** The exchange of chromatid segments of homologous chromosomes by breakage and reunion following synapsis during prophase of meiosis.
- Crossing-over value The frequency of crossing over between two genes.
- **Cryptic structural hybridity** Hybrids with chromosomes structurally different but not visible by metaphase-I chromosome pairing.
- ct DNA Chloroplast DNA.
- Culm Aerial stems of grasses, usually vertical but can be prostrate or spreading.
- Cultivar A category within a species of crop plants; cultivated variety.
- **Cuticle** A fatty substance that is almost impermeable to water and is on the outer walls of the epidermal cells.
- Cytochimera Tissues contain different chromosome numbers in a single plant organ.
- **Cytogenetics** A hybrid science that deals with the study of cytology (study of cells) and genetics (study of inheritance). It includes the study of the number, structure, function, and movement of chromosomes and numerous aberrations of these properties as they relate to recombination, transmission, and expression (phenotype) of the genes.
- **Cytokinesis** The division of cytoplasm and organelles in equal proportions. In plants, it occurs by the formation of a cell plate, while in animals, it begins by furrowing. In lower organisms, cytokinesis does not follow after telophase, resulting in multinucleate cells.
- Cytokinin Plant hormones that stimulate shoot in tissue culture.
- Cytological hybridization See in situ hybridization.

Cytological maps The genes located on the maps based on cytological findings such as deficiencies, duplications, inversions, and translocations.

Cytology The study of cells.

Cytoplasm A portion of the cell apart from the nucleus.

Cytoplasmic inheritance Inheritance of traits with determinants that are located in the cytoplasmic organelles, also known as non-Mendelian inheritance.

Cytosine A nitrogenous base, one member of the base pair G-C (guanine and cytosine).

Cytoskeleton Networks of fibers in the cytoplasm of the eukaryotic cells.

Cytosol The volume of cytoplasm in which organelles are located.

Cytotype Chromosome numbers other than the standard chromosome complement of that species. For example, *Glycine tomentella* contains accessions with four chromosome numbers (2n = 38, 40, 78, 80).

Daughter chromosome One of the chromatids of mitotic anaphase-I or meiotic anaphase-II.

Deciduous Plants that shed all their leaves at the end of each growing season.

- **Deficiency** The loss of a segment from a normal chromosome, which may be terminal or intercalary. The term deletion refers to an internal deficiency.
- **Dehydration** Loss of water during reactions in which large molecules are synthesized from smaller ones; the reverse of hydrolysis.
- **Denaturation of DNA or RNA** Conversion from the double-stranded to the single-stranded state accomplished by heating.
- Deoxyribose The four-carbon sugar that characterizes DNA.

Detassel Removal of male inflorescence (tassel) from maize.

Developmental genetics The study of the operation of genes during development.

- **Diakinesis (meiosis)** Chromosomes continue to shorten and thicken. This is a favorable stage in which to count chromosomes. Chromosomes lie spaced uniformly in the nucleus and often form a row near the nuclear membrane.
- **Diallel cross** Production of all possible hybrid combinations from a set of parents.
- **Diaspore** Any part of a plant dispersed from the parent plant and that functions in reproduction.
- Dicentric chromosome A chromosome with two kinetochores.
- **Dichogamous** Protandry; male and female organs mature at different times, allowing cross-pollination.
- **Differentiation** It occurs in cell, tissue, and organ cultures, where a plant develops from meristematic cells.
- **Diffuse kinetochore** Centromere activity is spread over the surface of the entire chromosome in certain insects of the order Hemiptera and in plant Luzula.
- Dihybrid cross A genetic cross involving two genes.
- **Dioecious** A plant bears staminate or pistillate flowers.
- **Diplo-haplontic** This phenomenon occurs in higher plants and in many algae and fungi. The life cycle of a typical plant contains spore-bearing (2n) and gamete-bearing (n) stages.
- **Diploid** An organism with two sets of chromosomes (two complete genomes). For example, barley is a diploid with 2n = 2x = 14.
- **Diploid-like meiosis** An allopolyploid showing bivalent chromosome pairing that, sometimes, is under genetic control.
- **Diplonema (meiosis)** A stage at which sister chromatids begin to repulse at one or more points in a bivalent.
- **Diplontic** Mature individuals are diploid by mitotic divisions and differentiations. Meiosis produces only haploid gametes. Diplophase is more prominent than haplophase in humans, higher animals, and some algae.
- Diploid cells overgrow the aneuploid cells in culture.
- Diplospory The embryo sac is formed from a megasporocyte as a result of abnormal meiosis.
- **Direct tandem inversions** It is due to two successive inversions involving adjacent chromosome segments.
- Discontinuous variation Phenotypic distinct variation caused by genes.
- **Disjunct** A genus or species with representatives that are at separate locations; once, distribution may have been continuous.

Disjunction Migration (separation) of chromosomes to opposite poles at meiotic anaphase.

Disomic inheritance It occurs by the association of chromosomes in bivalents at meiosis.

Dispermic An egg fertilized by two male gametes.

- **Distal chromosome** A part of a chromosome, which is relative to another part, is farther from the kinetochore (opposed to proximal).
- **Divergence** A pattern of evolution in which distinction among groups occurs through natural selection and environments two groups are not alike.
- **Dizygous** It is also known as fraternal twins. When two different eggs, each fertilized by a different sperm, have different genotypes.
- DNA Deoxyribose nucleic acid. All hereditary information is encoded.
- **DNase** An enzyme that attacks the bond in DNA.
- **Dominance** When one member of a pair of alleles manifests, more or less, when heterozygous, than its alternative allele.
- **Donor parent** A parent from which one or more gene-controlling economically important traits are transferred to a recurrent parent by back crossing.
- **Double cross** Crossing two single cross- F_1 hybrids.
- **Double crossing over** The production of a chromatid in which crossing over has occurred twice.
- **Double fertilization** The union of one sperm nucleus with the egg nucleus and one with polar nuclei (secondary nucleus) to form an embryo and endosperm in flowering plants.
- Double heterozygote Heterozygote in respect of two genes.
- Double reduction Chromatid segregation.
- **Drosera scheme pairing** A triploid hybrid showing 2x bivalent + 1x univalent configurations at diakinesis or metaphase-I.
- Duplex genotype AAaa.
- **Duplication** When an extra piece of chromosome segment is attached to the same chromosome (homologous) or transposed to another member (nonhomologous) of the genome.
- **Dysploid** A species with differing basic chromosome numbers (e.g., x = 5, x = 6, x = 7, etc.). Basic chromosome numbers in dysploid are stable.
- **Ear** In maize, the female is an ear from which protrude the stigma (silks).
- **Ecology** A branch of science that deals with the structure and function of nature; a study of the relationship between organisms and their environments.
- **Ectopic pairing** Chromosome pairing between an allele and nonallelic (ectopic) homologous sequence due to some homology searching mechanism.
- Egg The female gamete as opposed to the sperm.
- **Egg apparatus** Includes an egg cell and two synergid cells at the micropyle end of the embryo sac. **Electron microscope** A microscope that uses an electron beam as a light source.
- **Electrophoresis** A biochemical technique that utilizes an electrical current to separate mixtures of similar substances.
- Emasculation Removal of anthers from a flower.
- Embryo Rudimentary plant in a seed.
- **Embryo culture** The excision of young embryos under aseptic conditions, and the transfer of them on a suitable artificial nutrient media for germination or callusing.
- **Embryo sac** A megagametophyte of flowering plants that includes an egg, synergids, polar nuclei, and antipodals.
- **Embryogenic cells** Cells that have completed the transition from somatic state to one in which no further exogenously applied stimuli are necessary to produce the somatic embryos.
- Embryoid An embryo-like structure observed in cell and tissue cultures.
- Embryology To study the development of reproductive organs in an organism.
- **Endemic** A native; confined to a certain geographical area.
- **Endomitosis** Separation of chromosomes, but two sister chromatids lie side by side, and retention within a nucleus.
- Endoplasmic reticulum The outer membrane of the nuclear envelope.
- Endopolyploidy Polyploidy that arises from repeated chromosome replication without cytokinesis.
- **Endoreduplication** Chromosomes replicate time after time without condensation and mitosis and appear as bundles of multiple chromatids, e.g., polytene chromosomes.

- **Endosperm** Triploid tissue that originates from the fusion of a sperm nucleus with the secondary nucleus and nourishes the embryo during an early growth of the seedling.
- **Endothelium** The inner epidermis of the integument, next to the nucellus, where cells become densely cytoplasmatic and secretory. It is also known as integumentary tapetum.
- **Endothesium** A layer of cells situated in the pollen-sac wall situated below the epidermis.
- **Enneaploid** 9*x*.
- Entomophilous Pollination by insects.
- **Environment** The external conditions that affect growth and development of an organism.
- Ephemeral A plant completing its life cycle in considerably less than 12 months (short-lived).
- Epidermis The outermost layer of cells.
- Epigenetic A change in phenotype of an individual without altering the genotype.
- Epipetalous stamen A stamen that is congenitally attached to a petal.
- Epiphytotic An uncontrollable spread of a destructive plant disease.
- Epistasis Dominance of one gene over a nonallelic gene.
- **Equational division** A division of each chromosome into exact longitudinal halves that are incorporated equally into two daughter nuclei, also known as mitosis.
- Equational plate Metaphase plate.
- Erectoides A mutant plant with short erect stature and compact appearance.
- **EST** Expressed sequence tag.
- Ethnobotany The science that deals with the folklore knowledge of plants.
- **Euchromatin** The decondensed region of a chromosome containing the majority of the genes and which stains lightly.
- Eukaryotes An organism with true nucleus.
- Eumeiosis Normally occurring meiosis.
- **Euploid** An organism possessing chromosomes in exact multiple of the basic (x) chromosome number, such as 2x, 3x, 4x, and so on.
- Euspory The typical sporogenesis with normal flow of reduction division-reduced apomixis.
- Eutelomere A weakly staining subterminal segment adjacent to the protelomere.
- Exalbuminous seed A seed devoid of endosperm after maturity.
- **Exine** The outer wall of a mature pollen grain.
- **Exocarp** The outermost layer of the fruit wall; it may be the skin of the fruit or may be a leathery rind or may be quite hard.
- Explant An excised piece of tissue used to initiate tissue culture in vitro.
- **Expressivity** The degree of phenotypic effect of a gene in terms of deviation from the normal phenotype.
- **Extranuclear genes** Genes located in the cytoplasmic organelles, such as mitochondria and chloroplasts.
- F_1 , F_2 , etc. Filial generations. The offspring results from experimental crossing (F_1) and selfing (F_2). Facultative apomixis The plant produces seed apomictically and sexually.
- Facultative heterochromatic Chromatin that is heterochromatinized and may or may not be condensed in interphase.
- **Female** ("Venus looking glass"); female person, animal, or plant; in plants, having pistil and no stamens; capable of being fertilized and bearing fruits.
- Feral An uncultivated crop.
- Fertility Ability of a plant to produce offspring.
- Fertilization Fusion of a male sperm with an egg cell (formation of zygote).
- **Feulgen staining** A cytological colorless stain, utilizes Feulgen or Schiff's reagent. Chromosomes are colored, cytoplasm and nucleolus are colorless, and it is specific for DNA.
- Filament The stalk of a stamen.
- Filiform apparatus A complex wall outgrowth in a synergid cell.
- **Fixation** The killing of cells or tissues by chemicals in such a way that the internal structure of a cell is preserved.
- Flag leaf The last leaf before inflorescence emergence.
- **Fluorescence** *in situ* **hybridization (FISH)** A cytological technique for the detection of repetitive DNA sequences such as ribosomal DNA sequences (NOR sequences).

Fluorescence microscopy The principle of fluorescence microscopy is based on the light transmitted or reflected by the specimen. The specimen is stained with a fluorochrome (a dye that emits a longer wavelength when exposed to a shorter wavelength of light). The fluorescent region of a specimen becomes bright against a dark background.

Frequency distribution Frequencies of class intervals provide frequency distribution.

Funiculus The stalk by which the ovules are attached to the placenta.

- G_1 The period of the eukaryotic cell cycle between the last mitosis and the start of DNA replication (synthesis).
- G_2 The period of the eukaryotic cell cycle between the end of DNA replication (synthesis) and the start of the next mitosis.
- **G-band** Bands observed in the chromosomes after Giemsa staining. The chromosomes are pretreated with a dilute trypsin solution, urea, or protease. Bands have been used to identify human and plant chromosomes.

G-banding A cytological technique that exhibits a striated pattern in chromosomes.

Gamete A sex cell that contains half the chromosome complement of the zygote formed after meiosis that participates in fertilization.

Gametic chromosome number *n*.

Gametocyte A cell that produces gametes through division; a spermocyte or oocyte-germ cell.

Gametogenesis The formation of male and female gametes.

- **Gametophyte** A part of the life cycle of the plant that produces the gametes, where there is an alternation of generations.
- **Gametophytic incompatibility** A series of alleles at one locus determines whether the pollen will be functional or nonfunctional. A similar gene in pollen and stylar tissue will cause inhibition of pollen tube growth. Thus, the ability of a pollen to function is determined by its own genotype and not by the plant that produces it.
- **Gene** The unit of inheritance located mainly in chromosomes at fixed loci; contains genetic information that determines characteristics of an organism.
- Gene dosage Refers to the number of copies of a particular gene in the genome.

Gene flow The contribution of a particular gene or genes to a population through interbreeding.

Gene frequency Proportion in which alternative alleles of a gene occur in a population.

Gene interaction Modification of gene action by a nonallelic gene or genes.

Gene pool All the alleles in an inbreeding population.

- **Generative nucleus** A product (two daughter nuclei) of division of primary nucleus in the pollen grain. One usually divides into two sperm nuclei in the pollen tube, while vegetative nucleus does not divide again.
- Genetic A property obtained by an organism by virtue of its heredity.
- Genetic code The correspondence between triplets in DNA (or RNA) and amino acids in proteins.
- **Genetic combination** For *n* pairs of homologous chromosomes, only 1/2n 1 gametes will contain parental combination.

Genetic drift Random change in gene frequency in a population due to sampling error.

Genetic male sterility Failure to produce and shed viable pollen, controlled by genes.

Genetic mapping The assignment of genes to specific linkage groups and determination of the relative distance of these genes to other known genes in that particular linkage group.

- Genetic marker An allele used to identify a gene or a chromosome or a chromosome segment.
- **Genetics** The science that deals with heredity and variation.
- Genome A complete haploid set of chromosomes of a species.
- **Genome affinity index (GAI)** Used for comparing the degree of homology among parental genomes the mean number of bivalent-equivalents is divided by the basic chromosome number.
- **Genome segregation** The segregation of sets of chromosomes (genomes) during mitosis in the somatic tissues that occurs in polyploid species and also in some wide crosses.
- **Genomic** *in situ* **hybridization (GISH)** A cytological technique by which a genome-specific probe is used. A DNA sequence specific to a particular genome serves as a marker to identify genome, translocations, and alien addition and substitution lines.
- **Genotype** The description of the genes contained by a particular organism for the character or characters the entire genetic constitution.

Genotypic ratio Proportion of different genotypes in a population.

Genus The first element of the scientific name. It includes a taxonomic group of related species. **Germ cell** Gamete.

Germ plasm The total hereditary materials available in a species and genus.

Germination The emergence of root and shoot from a seed; may be epigeal (elongation of hypocotyls) or hypogeal (little elongation of hypocotyl or no elongation of hypocotyls).

Golgie apparatus All golgie bodies in a given cell are not like staining bodies usually engaged in secretion.

Golgie bodies A cell organelle made up of closely packed broad cisternae and small vesicles.

Guanine A nitrogenous base, one member of the base pair G-C (guanine and cytosine).

- Gynoecium The female portion of the flower stigma, style, and ovary.
- **Gynomonoecy** A plant having female and hermaphrodite flowers.
- **Gynophore** An elongation of the floral axis between the stamens and the carpels, forming a stalk that elevates the gynoecium.
- **Haploid** An organism containing a gametic (*n*) chromosome number.
- **Haploid-initiator gene** A partially dominant haploid-initiator gene (*hap*) in barley that controls the abortion or the survival of abnormal embryos and endosperms. Plants homozygous for the *hap* gene produce progenies that include 10 to 14% haploids.
- **Haplontic** In most primitive unicellular or filamentous algae and protozoa, haplophase is more prominent than diplophase. Haplophase occurs from meiosis to fertilization.
- **Haplotype** The particular combination of alleles in a defined region of some chromosome, in effect, the genotype in miniature.

Haustorium A specialized organ that draws nutrients from another organ.

Helobial endosperm A type of endosperm that develops in a way intermediate between that by which a cellular and a nuclear endosperm develop.

Hemigamy Sometimes, the sperm nucleus does not unite with the egg nucleus but divides independently and simultaneously. The embryo sac nuclei are unreduced. Chimeric embryos (somatic + half the somatic chromosome) develop.

Hemizygous A gene allele opposite to a deficiency.

- **Heredity** The process that brings about the biological similarity and dissimilarity between parents and progeny.
- Hermaphroditic Flowers include male and female sexes.
- Heterocarpy The production of two or more types of fruits by a plant.

Heterocaryosis The presence of two or more genetically different nuclei within single cells of a mycellium.

- **Heterochromatin** The region of a chromosome that takes differential stain. These regions contain condensed chromatin.
- Heterogametic sex An organism has different sex chromosomes (X and Y, Z and W) and autosome chromosomes.
- **Heterogenetic chromosome pairing** The F_1 hybrids of genomically divergent species lack chromosome affinity and form only a small number of loosely synapsed bivalents.
- **Heteromorphic bivalent** A bivalent in which one of the chromosomes differs structurally. The chromosomes may differ in size or kinetochore positions.

Heteroplasma Extranuclear genes that are not alike (analogous to heterozygote for nuclear genes).

- **Heteroploid** An individual containing a chromosome number other than the true monoploid or diploid number.
- **Heteropycnosis** Differential spiralized regions of chromosomes that are positive at interphase or early prophase and negative at late prophase or metaphase.
- Heterosis Heterozygote superiority of an F_1 hybrid over parents, with respect to some character or characters.
- **Heterozygous** A zygote derived from the union of gametes unlike in respect of the constitution of their chromosomes or from genes in a heterozygote (e.g., Aa). Also, used in case of hybrid for interchanges, duplications, deficiencies, and inversions opposite of homozygotes.
- Hexaploid A polyploid with six sets of basic chromosomes.

- **Highly repetitive DNA** It is equated with the satellite DNA that is composed of many tandem repeats (identical or related) of a short basic repeating unit (= simple sequence DNA).
- **Hilum** The scar on the surface of seed revealed by abscission of the funicle; the point of attachment of the seed to the pod.
- **Histology** Study of tissues.
- Holandric genes The genes located on Y chromosomes.
- **Holocentric** Diffused kinetochore. Every point along the length of the chromosome exhibits centromeric activity.
- **Homoeobox** A short stretch of nucleotides with a base sequence that is virtually identical in all genes that contain it. It determines when particular groups of genes are expressed during development.
- **Homoeologous chromosomes** Partially homologous chromosomes; for example, in wheat, ABD genome chromosomes are known as homoeologous.
- Homogametic sex An organism has two X chromosomes and autosome.
- Homogamy The male and female parts of a flower mature simultaneously.
- Homogenetic chromosome pairing Synapsis of homologous chromosomes of the same genome.
- **Homologous chromosomes** Like chromosomes showing synapsis along their length during pachynema. For *n* pairs of homologous chromosomes, only 1/2n 1 gametes will contain parental combination.
- **Homomorphic incompatibility** It is due to relative length of the stamens and the style.
- Homoplasmon Extranuclear gene alike (analogous to homozygote for nuclear genes).
- **Homozygote** An organism having like allelles at corresponding loci on homologous chromosomes (e.g., AA, aa).
- **Hot spot** A site in the chromosome at which the frequency of mutation (or recombination) is greatly increased.
- Hybrid The progeny from mating parents of different genotypes, strains, species, and genera.
- **Hybrid inviability** The death of F_1 hybrid plants due to aberrant development processes.
- **Hybridization** The production of a hybrid by crossing two individuals of unlike genetic constitution. **Hybrid vigor** See heterosis.
- Hyperploid A diploid organism with an extra chromosome.
- **Hypertriploid** An individual containing chromosome numbers higher than 3*x*.
- Hypoploid A diploid organism lacking an extra chromosome.
- Ideotype An ideal plant form formulated to assist in reaching selection goals.
- Idiogram A diagrammatic representation of chromosome morphology of an organism.
- **Immunity** A plant that does not permit infection by a parasitic organism.
- Inbred A line originated by continued self-fertilization accompanied by selection.
- Inbreeding Mating of closely related plants or animals.
- **Included inversions** A segment that is part of an inverted segment is inverted once again.
- **Incompatibility** Inability of a male gamete to fertilize a female gamete (egg), thus preventing the zygote formation.
- **Indehiscent** A fruit that does not open by sutures, pores, or caps, the seeds being released by the rotting away of the fruit wall.
- **Independent assortment** The random distribution to the meiotic products (gametes) of genes located on different chromosomes. A dihybrid individual with RrYy genotype will produce four gametic genotypes (RY, Ry, rY, ry) in equal proportions.
- **Independent inversions** Such inversions occur in independent parts of the chromosomes, and the two resultant inverted segments are separated from each other by an uninverted (normal) chromosome segment.
- **Induced mutation** Induction of mutation by mutagens.
- Inflorescence Spikelets arranged on a common branching system (panicle, raceme, spike).
- *in situ* hybridization A cytological technique conducted by denaturing the DNA of cells squashed on a microscope slide so that reaction is feasible with an added single-stranded DNA or RNA; which is labeled and the following hybridization gene is located on a certain site of chromosome.
- **Integument** Layers of cells surrounding the ovule.
- **Interchanges (reciprocal translocation)** The reciprocal exchange of terminal segments of nonhomologous chromosomes.

Intergeneric cross A cross between two species from two different genera.

Interkinesis The time gap between cell division I and II. It is short or may not occur at all.

- Internode The interval on a stem or rachis between two nodes.
- **Interphase** The period between mitotic cell division (or intermitotic period) that is divided into G_1 , S, and G_2 .
- Interspecific cross The cross between two parents from different species.

Intine The inner wall of a mature pollen grain.

Intragenomic pairing Pairing of chromosomes in monohaploid.

- **Introgressive hybridization** The incorporation of a gene or genes of a species into the gene pool of another species.
- **Inversion** Reversal of gene order within a chromosome, may be paracentric, which does not include kinetochore, or pericentric, which involves kinetochore in the inverted region.
- **Inversion heterozygote** One chromosome carries a normal gene sequence, while the other chromosome contains an inverted gene sequence may be paracentric or pericentric.

in vitro Outside a living organism.

Irradiation Exposure of an organism to x-rays or other radiation to enhance mutation rates.

Irregular flower The perianth constituted in such a way that it is not possible to divide it into two equal halves, as in the canna flower.

Isogenic lines Genetically uniform, homozygous lines.

Isolation The separation of one group from another, preventing mating between and among groups.

Kappa particles Cytoplasmic kappa particles cause a killer trait in *Paramecium* that is genotypic dependent. Kappa particles reduce in cells containing the genes k, s1, and s2. Some races of killers paramecium produce paramicin, which is lethal to other races (sensitive). Certain portions of kappa particles contain proteinaceous R-bodies. These are responsible and required for the toxic action of kappa particles on sensitive types. The behavior and structure of R-bodies are similar to any other bacterial structure that has been established by chemical composition and electron microscopic studies.

Karyogamy The fusion of two gametic nuclei.

Karyokinesis The process of nuclear division.

Karyology Study of the nucleus.

Karyotype The usual chromosome number, size, and morphology of the chromosome set of a cell of an organism. The complete set of chromosomes. For example, barley contains 2n = 2x = 14.

kb An abbreviation for 1000 base pairs of DNA or 1000 bases of RNA.

Kinetochore The region of a chromosome attached to a spindle fiber during movement of the chromosomes at mitosis and meiosis.

- **Kinetochore-misdivision** Division of kinetochore crosswise instead of lengthwise, particularly during meiosis.
- **Kinetochore-orientation** The process of orientation of kinetochore during prometaphase of mitosis and meiosis.
- **Knob** A heterochromatic darkly stained region that may be terminal or intercalary. It is often used as a landmark to identify specific chromosomes.
- **Lagging** Delayed migration of chromosome from the equatorial plate to the poles at mitotic or meiotic anaphase and, consequently, excluded from the daughter nuclei producing aneuploid gametes.
- **Lampbrush chromosome** A special type of chromosome found in the primary oocyte nuclei of vertebrates and invertebrates. Such chromosomes are due to prolonged diplonema. Chromosomes exhibit a fuzzy appearance due to paired loops. These loops vary in size and shape from chromosome to chromosome and have been used to construct cytological maps.

Landrace A primitive cultivar evolved from a natural population and selected for cropping.

Leaky mutation It allows some residual expression of a gene.

Leptonema (meiosis) Chromonemata become distinct from one another and appear as long and slender threads. Nuclei contain a large nucleolus and a distinct nuclear membrane.

Lethal gene Any gene in which a lethal mutation can be obtained (deletion of the gene).

Leucoplast A plastid devoid of pigment.

Linkage Association of qualitative and quantitative traits in inheritance due to location of marker genes in proximity on the same chromosome.

- **Linkage group** It includes all loci that can be connected (directly or indirectly) by linkage relationships; equivalent to haploid chromosome. For example, barley has seven linkage groups corresponding to seven haploid chromosome numbers.
- **Linkage map** A graphical representation of the positions of genes along a chromosome.
- **Linkage value** Recombination fraction: the proportion of crossovers versus parental types in a progeny. It may vary from 0 to 50%.
- **Localized kinetochore** A chromosome carrying normal permanently localized kinetochore to which spindle fiber is attached during chromosome separation.

Locus The position of a gene in a chromosome.

- **Lysenkoism** The doctrines proclaimed in the former Soviet Union by T.D. Lysenko that do not accept the gene concept and are based on the inheritance of acquired traits.
- M_1 , M_2 , etc. Symbols used to designate first, second, etc., generations after treatment with a mutagenic agent.
- **Maceration** The artificial separation of the individual cell from a tissue by enzyme treatment.
- Male ("Mars shield and spear"); male person, animal, or plant; generally used in contrast to female.
- **Male sterility** The male sterile genes (*ms*) cause complete breakdown of microsporogenesis, but macrosporogenesis (megasporogenesis) remains uninfluenced.

Map distance It is measured at cM (centiMorgan) = percent recombination.

Marker An identifiable physical location on a chromosome, determined by inheritance.

Maternal inheritance Inheritance of a trait through a female.

Maturity group Division of soybean cultivars in North America into 12 groups based on how flowering responds to dark periods; maturity groups are selected for the latitude, so that flowering is initiated for maximum yield.

Mean Central value of a set of sample observations.

- Median The value for which 50% of the observations are on either side.
- **Megabase** (Mb) Unit length for DNA fragments equal to 1 million nucleotides and roughly equal to 1 cM.

Megagametophyte Female gametophyte; embryo sac within the ovule of angiosperms.

Megaspore A gametophyte bearing only female gametes.

- **Megaspore mother cells (MMCs)** Produce eight nuclei (megaspores) after one of four haploid macrospores becomes functional in the embryo sac. Two mitotic divisions produce eight nuclei (one egg, two synergids, three antipodals, and two polar nuclei).
- **Meiosis** A phenomenon that occurs in reproductive organs of male and female organisms, producing gametes with the haploid (n) chromosome constitution. The first division is reductional, and the second division is equational.
- **Meiotic segregation** The random orientation of bivalents at equatorial plate at metaphase-I regulates the segregation of paternal and maternal chromosomes to the daughter nuclei.
- **Meristem** An actively growing tissue that produces cells which undergo differentiation to form mature tissues.

Mesocarp The middle layer of the fruit wall; often the fleshy edible portion.

- **Mesophyll** The photosynthetic parenchymatus tissue situated between the two epidermal layers of the leaf.
- **Mesopolyploid** Individuals have undergone moderate aneuploid-induced base number shift. Apomicts may contain complete or nearly complete duplicate sets of genes.

Metacentric chromosome A chromosome with a median kinetochore (both arms are equal in length). **Metakinesis** Prometaphase.

- **Metaphase (mitosis)** Chromosomes are arranged at the equatorial plate. Complete breakdown of the nuclear membrane and disappearance of the nucleolus occur.
- **Metaphase-I** (meiosis) Bivalents are arranged at the equatorial plate with their kinetochores facing their poles. Chromosomes reach to their maximum contractions, nuclear membrane and nucleolus disappear, and spindle fibers appear.
- **Metaphase plate** Movement and arrangement of all the chromosomes takes place midway between the two poles of the spindle, at the metaphase plate.
- **Metastatis** The ability of tumor cells to leave their site of origin and migrate to other locations in the body, where a new colony is established.

Metaxenia Tissues outside the embryo sac are influenced by the pollen source.

- Microgametophyte The male gametophyte of a heterosporous plant.
- **Micronucleus** A nucleus separate from the main nucleus formed at telophase by one or more lagging chromosomes or fragments.
- Micropyle An opening of the ovule.
- Microsomes The fragmented pieces of endoplasmic reticulum associated with ribosomes.
- Microsporangium A structure that produces the microspores in a heterosporous plant.
- Microspore A gametophyte producing only male gametes a pollen grain.
- **Microspore mother cells** Diploid cells in the stamens (pollen mother cells, PMCs) give rise to haploid microspores after meiosis.
- Microsporocyte A cell that differentiates into a microspore.
- **Microtubules** Manipulation of chromosomes in eukaryotes is due to microtubules. They are organized into spindle-shaped bodies and disappear at the end of cell division.
- Minichromosomes Small chromosomes with kinetochores, totally heterochromatic.
- **Mitochondrion** A cell organelle found in the cytoplasm. Mitochondrial DNA is responsible for extranuclear inheritance. It is the site of most cellular respiration.
- **Mitosis** A process of cell division known as equational division, because the exact longitudinal division of the chromosomes into two chromatids occurs. The precise distribution into two identical daughter nuclei leads to the generation of two cells identical to the original cell from which they were derived.

Mitotic anaphase Daughter chromosomes are pulled to opposite poles by the spindle fibers.

- **Mitotic metaphase** Chromosomes are shrunk to the maximum limit, and kinetochores move onto the equatorial plate.
- **Mitotic prophase** Chromosomes begin to reappear, are uniformly distributed in the nucleus, and are more or less spirally coiled and seem longitudinally double.
- **Mitotic telophase** By the end of anaphase, spindle fibers disappear, and chromosomes at each pole form a dense boll. Nucleolus, nuclear membrane, and chromocenters re-emerge. Chromosomes lose their stainability.
- **Mixoploidy** A failure in chromosome migration results in the formation of multiploid microsporocytes.
- Mode The values that occur most frequently.
- **Modifying genes** The genes that, by interaction, affect the phenotypic expression of nonallelic genes or a nonallelic gene. The gene may be an enhancer (intensifies the expression) or reducer (decreases the expression of the other gene).
- Monocentric One centromere per chromosome.
- Monoecious Both staminate and pistillate flowers are on the same plant.

Monogenic character A character determined by a single gene.

- Monohybrid cross A genetic cross that involves only one pair of genes.
- **Monoploid** An organism having a basic (*x*) chromosome number.
- **Monosomics** Where one of the chromosomes is missing from the normal chromosome complement (2n 1).
- **Monozygous twins** Also known as identical twins; twins arise from a single zygote formed by the fertilization of a single egg by a single sperm. In this case, the developing egg cleaves at an early division, forming two separate embryos instead of the usual one.
- Morphogenesis The differentiation of structures during development.
- **Mother cell** A diploid nucleus which, by meiosis, yields four haploid nuclei the microspore or pollen mother cell (PMC) and the megaspore or embryo-sac mother cell in flowering plants.
- mtDNA Mitochondrial DNA.
- **Multiform aneuploidy** An organism contains cells with different aneuploid chromosome numbers producing the tissues with chromosome mosaicism.
- **Multiple factors** Many genes involved in the expression of any one quantitative character.
- **Multivalent** An association of more than two completely or partially homologous chromosomes during meiosis, generally observed in chromosomal interchanges and in autopolyploidy.
- **Mutagen** A physical or chemical agent that significantly raises the frequency of mutation above the spontaneous rate. These types of mutations are called induced mutations.

Mutagenic Capable of causing mutation.

- Mutant An organism carrying a gene that resulted from mutation.
- **Mutation** Any heritable change in the genetic material (sequence of genomic DNA) not caused by segregation or genetic recombination, may occur spontaneously or be induced experimentally; detectable and heritable changes in the genetic material transmitted to succeeding generations.
- **N-bands** This procedure was originally developed to stain nucleolus organizer chromosomes. Chromosomes are treated at low pH (4.2 ± 0.2) in 1 *M* NaH₂Po₄ for 3 min at a high temperature ($94 \pm 1^{\circ}$ C). Chromosomes are stained with Giemsa after incubation and washing.

Natural crossing It occurs between individuals within a population.

Necrotic Dead plant tissues; death is usually caused by disease, insect activity, or nutrient deficiency.

Neocentromeres The centromere region is replaced by a secondary center of movement; chromosome ends move first during anaphase-I of meiosis.

Neopolyploid Polyploid individuals of recent origin.

Neospecies New species.

Neurospora The common *Neurospora crassa* is a bread mold. The normal vegetative phase is a branching haploid individual with hyphae that forms a spongy pad, the mycelium. Mycelium fuses with each other, but nuclei do not (heterokaryon), although they are mixed and remain in the same cytoplasm. Asexual reproduction occurs by repeated mitosis of nuclei or by mitotic formation of special haploid spores (conidia). Conidia can form new mycelium.

Node The point at which a leaf is inserted into or attached to the stem.

- **Nonchromosomal** It shows a non-Mendelian inheritance; the genes are extrachromosomally located in chloroplasts and mitochondria.
- **Noncoorientation** In a quadrivalent, two nonhomologous centromeres are on opposite sites, and centromeres of two translocated chromosomes are stretched between the other two and not attached to the opposite poles. Noncooriented quadrivalent configuration results in a 3:1 chromosome segregation, always produces unbalanced gametes, and generates trisomics in the progenies.
- **Nondisjunction** It can occur during mitosis and meiosis. Chromosomes fail to separate at metaphase, resulting in their passage to the same pole. This may occur due to failure of chromosome synapsis or from multivalent formation.
- **Nonhomologous** Chromosomes carrying dissimilar chromosome segments and genes that do not pair during meiosis.
- **Nonlocalized centromeres** The spindle attachment is not confined to a strictly localized chromosome area.

Nonrecurrent parent The parent of a hybrid not used again as a parent is backcrossing.

Northern blotting A molecular biology technique for transferring RNA from an agrose gel to a nitrocellulose filter on which it can be hybridized to a complementary DNA.

- Nucellus Ovule tissues internal to the integuments and surrounding the embryo sac.
- **Nuclear envelope** A doubled-layer outer membrane of the nucleus that separates the nucleus and the cytoplasm of the cell in eukaryotes.
- **Nuclear pore** Holes in the nuclear envelope that are assumed to be used for the transport of macromolecules.

Nucleic acid A large molecule composed of nucleotide subunits.

- **Nucleolus** A spherical body found inside the nucleus is associated with a specific chromosomal segment (the nucleolus organizer) and is active from telophase to the following prophase and is the site of ribosomal RNA (rRNA) synthesis.
- **Nucleolus organizer region** The chromosome region associated with the nucleolus, responsible for the formation of the nucleolus, and contains ribosomes. Referred to as the NOR.
- **Nucleosome** The basic structural subunit of chromatin, consisting of ~200 bp of DNA and an octamer of histone of proteins.
- **Nucleotype** The gross physical characterization of the nucleus; its mass and, particularly, the DNA content.
- **Nucleus** A spheroidal, most important, body within the cells of eukaryotes, which contains chromosomes.

Null mutation Physical deletion of a gene.

Nulliplex In polyploid, one recessive allele is carried by all chromosomes (aaaa).

- **Nullisomics** An individual lacking one chromosome pair from the normal chromosome complement (2n 2).
- **Obligate apomixis** In plants where no sexual reproduction occurs.
- Ontogeny The development of an organism from fertilization to maturity.
- **Oöcyte** The cell that becomes the egg by meiosis.
- **Oögonium** A mitotically active cell that gives rise to oöcytes.
- **Organelle** Any structure of characteristic morphology and function in a cell, such as nucleus, mitochondrion, chloroplast, etc.
- Organogenesis Formation of organs (structural and functional units flowers).
- **Outbreeding** A mating of genetically unrelated plants.
- **Overdominance** A heterozygote (Aa) is superior in phenotype than either homozygotes (AA, aa).
- **Overlapping inversions** A part of an inverted segment being inverted a second time, together with an adjacent segment that was not included in the first inversion segment.
- **Ovum** An unfertilized egg ovule, egg.
- p Human chromosome nomenclature for the short arm (petit, French for short).
- P_1 Denotes parents of a first filial (F_1) generation.
- **P-particles in** *Paramecium* A group of about eight different particles (kappa, gamma, delta, pi, mu, lambda, alpha, and tau) are found in the cytoplasm of *Paramecium aurelia*. These are called P-particles. These particles consist of DNA, RNA, and proteins.
- **Pachynema (meiosis)** Synapsis of homologous chromosomes is completed. Chromosomes are noticeably thicker and shorter than those in leptonema. Nucleolus is clearly visible, and certain chromosomes may be attached to it.
- **Pairing** Chromosome pairing synapsis.
- **Pairing segment** The segment of chromosomes (X and Y chromosomes) that pairs and crosses over (pairing segment) and the remaining segments that do not pair, are known as differential segments.
- **Paleopolyploids** Ancient polyploids; all their 2*x*-progenitors apparently have become extinct.
- **Pangenesis** Proposed by Charles Darwin, all structures and organs throughout the body contribute copies of themselves to a sex cell.
- **Panicle** A much branched raceme in which flowers are borne on the ultimate branchlets, e.g., rice, oat. **Panmixis** Random mating in a population.
- **Paracentric inversion** A type of intrachromosomal structural change that does not include the kine-tochore of the chromosome.
- **Paramecia** Unicellular, slipper-shaped animals that maintain three individual nuclei: two are small (micronuclei) with a diploid chromosome number, and the third is large (macronucleus) and polyploid. Micronuclei divide mitotically by binary fusion. The macronucleus constricts in the middle and separates into two.
- **Parameters** Values of the larger population from which the sample has been taken are known as parameters.
- Paranemic coiling Chromatids are easily separated laterally.
- **Parasexuality** Crossing over and genetic recombination occur in viruses and fungi by means other than regular meiosis. The diploid nuclei are borne from the rare fusion of two genetically unlike nuclei in a heterokaryon. Haploid nuclei are produced from the diploid with a new combination of genes.
- **Parthenocarpy** The development of fruits without seeds as a result of lack of pollination, a failure of fertilization, or death of the embryo at an early stage of development.
- **Parthenogenesis** Embryo is produced from a haploid or diploid unfertilized egg cell with or without the stimuli of pollination and fertilization.
- Pathogen An organism causing disease in a plant.
- **PCR** (Polymerase chain reaction); a molecular technique in which cycles of denaturation, annealing with primer, and extension with DNA polymerase, are used to amplify the number of copies of a target DNA sequence by $>10^6$ times.
- **Pedicel** The stalk that supports the paired spikelet or flowers.
- Pedigree The record of the ancestry of an individual or a cultivar.
- Peduncle The stalk that supports an inflorescence including the stalk that supports a flower.

Penetrance The proportion of genotypes that exhibits an expected phenotype, and depends on genotype and the environment.

Pentaploid 5x.

Perennial Plants continue to grow through the years (two or more) and flower and fruit repeatedly. **Perfect flower** A flower possessing stamen and pistil.

Pericentric inversion A type of inversion that includes the kinetochore within the inverted segments.

Permanent heterozygote Individuals in which chromosomal heterozygosity is fixed, due to balanced lethal factors, translocations, or inversions, and that always breed true, e.g., *Oenothera*.

- **Phenocopy** The phenotype of a given genotype changed by external conditions to resemble the phenotype of a different genotype.
- **Phenotype** The observable external appearance of an organism produced by the interaction of genotype and the environment.
- **Phylogeny** The relationships of groups of organisms (taxonomic groups) as expressed by their evolutionary relationships.

Pistil The female reproductive organ (gynoecium) of a flower consisting of ovary, style, and stigma.

- **Pistillate** A female flower containing a functional gynoecium but lacking well-developed functional stamens.
- Placenta The region of attachment of the ovules to the carpel.

Plasma membrane A membrane that surrounds the boundary of every cell.

Plasmagene An extranuclear hereditary gene that shows non-Mendelian inheritance.

- **Plasmotype** All plasmagenes (total hereditary complement of a cell) constitute the plasmon or plasmotype.
- **Plastid** A self-cytoplasmic organelle of plant cells. Plastid includes chloroplasts (chlorophyll), chromoplasts (carotenoides), and leucoplasts (no visible pigments).
- Plastogene Genes located in the plastids showing non-Mendelian inheritance.

Plectonemic coiling Chromatids cannot be separated without unwinding the coil.

Pleiotropy The multiple distinct phenotypic effect of a single gene.

Ploidy The number of chromosome sets per cell of an organism as haploid, diploid, and polyploid. **Plumule** The embryonic shoot — coleoptile.

Point mutation A change in single base pairs (gene mutation).

- **Polarity** The effect of a mutation in one gene in influencing the expression of subsequent genes in the same transcription unit.
- **Pollen grain** Haploid male spores produced after meiosis male gametophyte.

Pollen mother cell (PMC) Microsporocyte that produces pollen grains as a result of microsporogenesis. **Pollen tube** A germinating pollen grain that carries male gametes to the ovum.

Pollination Natural or artificial placement of pollen on the receptive stigma of a female flower.

Pollinium A pollen mass, as in the milkweeds and orchids.

Polycentromeres Each chromosome is attached by many spindle fibers.

Polyembryony The production of several embryos in a seed.

Polygamous The plants produce perfect and imperfect flowers.

- **Polygene** A gene that individually exerts two slight effects, but together with several or many genes, control quantitative character.
- **Polyhaploid** Haploid (*n*) individuals arising from polyploid species.
- **Polymitosis** Occurence of a series of mistoses in rapid succession with or without division of chromosomes.
- **Polymorphism** The occurrence of two or more genetically different types in the population-allelic variation. In molecular terms, changes in DNA influence the restriction pattern.

Polyploid An organism containing more than two sets of chromosomes.

Polyploidization Induction of polyploids artificially or naturally from haploid and diploid plants.

- **Polyspory** Characterized by numerous cytological abnormalities, because complete sets of asynchronously expressed genes are not present. It is associated with the frequent occurrence of an an euploid series formation and stabilization (Poaceae).
- **Polytene chromosome** A giant chromosome consisting of many identical chromosomes closely associated along their entire length (endomitosis). In *Drosophila melanogaster*, salivary gland chromosomes about 5000 bands can be recognized. Homologous polytene chromosomes pair along

the length in the somatic cells (somatic pairing). Any chromosomal structural changes could be determined by differences in the pairing pattern. Changes in the appearance of bands in certain tissues occur at different times, either condensing or expanding to form "puffs."

Population A group of organisms inhabiting an area, which shares a common gene pool.

Position effect A change in the expression of a gene brought about by the translocation to a new site in the genome. A previously active gene may become inactive if placed near heterochromatin.

Primary constriction Synonymous with centromere or kinetochore.

- **Primary trisomics** An individual with normal chromosome complement plus an extra complete standard chromosome is designated as a simple primary trisomic.
- **Primer** A short sequence, often, RNA that is paired with one strand of DNA and provides a free 3'-OH end at which a DNA polymerase starts.
- **Primordium** A cell or organized group of cells in the earliest stage of differentiation.
- **Probe** Single-stranded DNA or RNA molecules of specific base sequence, labeled radioactively or immunologically, used to detect the complementary base sequence by hybridization.
- **Procumbent** A plant stem that lives on the ground for all or most of its life.
- Proembryo The earliest stages of embryo development.

Prokaryote An organism lacking nuclei — bacteria.

- **Prophase (mitosis)** The first stage of mitosis, in which cells prepare for cell division. Coiling of chromosomes (condensation) occurs, and chromosomes appear with a thread-like structure. Prophase includes the splitting of chromosomes longitudinally into two duplicates (chromatids) and the centrosome, and dissociation of each half to opposite sides of the nucleus, the synthesis of mitotic apparatus, the disappearance of nucleolus, and the beginning of the breakdown of the nuclear membrane.
- **Protandry** Maturation of anthers before pistils.
- **Protelomere** A terminal deep-staining structure with sharp limits, normally composed of one to three dark-staining large chromomeres.
- Protoclone A clone derived from a protoplast.
- **Protogyny** Maturation of pistils before anthers.
- **Protoplast** The plant cell exclusive of the cell wall.
- **Pseudogamy** Occurs in plants, when an embryo develops from a diploid or haploid egg cell with a sperm penetrated but without the occurrence of fertilization, and the egg cell nucleus remains functional.
- **Pseudogenes** These are inactive genes but are stable components of the genome derived by mutation of an ancestral active gene.
- **Pseudoisochromosome** Chromosome ends are homologues due to reciprocal translocation; chromosomes pair at meiotic metaphase-I like an isochromosome, but the interstitial segments (proximal to kinetochore) are nonhomologous.
- Pubescence A general term used for hairiness.
- **Punnett square** A checkerboard used in finding all possible zygotes produced by a fusion of parental gametes. This also results in determining the genotypic and phenotypic ratios.
- **Pure line** The descendent obtained from self-fertilization of a single homozygous parent.
- **Purine** A nitrogen-containing single-ring, basic compound that occurs in nucleic acids. The purines in DNA or RNA are adenine and guanine.
- **Pyrimidine** A nitrogen-containing, double-ring, basic compound that occurs in nucleic acids. The pyrimidines in DNA are cytosine and thymine; in RNA, they are cytosine and uracil.
- Q Human chromosome nomenclature for the long arm.
- **Q-bands** Chromosomes of many plant species stained with quinacrin mustard exhibited bright and dark zones under ultraviolet light. This method was later used for the identification of individual human chromosomes (A-T-rich region of a chromosome).
- **Quadripartitioning** During meiosis, cytokinesis does not occur after meiosis-I and is postponed until after the second division.
- **Quadrivalent** An association of four chromosomes generally observed in a reciprocal translocation or in autotetraploids.
- **Qualitative inheritance** Inheritance of qualitative traits controlled by a single gene (discontinuous inheritance).

- **Quantitative inheritance** Inheritance involving the simultaneous action of many genes in a cumulative or additive fashion, each of which produces a small effect (height, weight, and size are a few traits).
- **Quartet** The four nuclei produced at the end of meiosis (= tetrad).
- **Quasibivalent** Pseudobivalent Bivalent-like association during meiosis due to stickiness rather than to chiasmata.
- **R-bands** The lightly stained G-bands become darkly stained after R-banding staining. R-bands are obtained after treating chromosomes at low pH (4 to 4.5) at 88°C incubation in NaH₂Po₄ (1 *M*). R-bands are reversible if pH is adjusted to 5.5 to 5.6.
- Race A genetically distinct mating group within a species.
- Raceme An unbranched, elongate, indeterminate inflorescence on padicellate flowers.
- Rachilla The central axis of a spikelet is in the grass family.
- Radicle Embryonic root.
- **Radioautography** A photographic technique in which a radioisotope is taken in by a cell that can then be traced with a sensitive photographic film.
- **Radioisotope** A radioactive isotope of an element.
- **Radiomimetic** A chemical that mimics ionizing radiation in terms of damage to genes and chromosomes, resulting in gene and chromosome mutations.
- **Random mating** Occurs in a population where an individual has equal probability to mate with any individual of the opposite sex.
- **Range** The distance between two extreme values.
- **Reassociation of DNA** The pairing of complementary single strands of DNA to form a double helix.
- Recessive A member of an allelic pair of contrasting traits that are not expressed.
- Recessive lethal Recessive homozygous is lethal.
- Reciprocal cross Male and female parents are reversed.
- **Reciprocal recombination** Reciprocal exchange of one part of a chromosome with part of another chromosome.
- **Recombination** The occurrence of new combinations of genes in hybrids, not present in the parents, due to independent assortment and crossing over in crosses between genetically different parents.
- Recurrent parent In a back-crossing program, a parent used in successive back crosses.
- **Reduction** A process that halves the somatic chromosome number at meiosis; segregation of homologous includes a member of each set, but these chromosomes are no longer the unaltered parental chromosomes.

Regeneration The production of plants from cells and tissues through *in vitro* culture.

- **Regulatory gene** It codes for an RNA or protein product with the function of controlling the expression of the other gene.
- **Relational coiling** Chromatids are twisted about each other during early prophase.
- Renaturation The return of a denatured nucleic acid or protein to its native configuration.
- **Renner complexes** In *Oenothera*, several genetic factor complexes are found, and these complexes segregate as a whole in meiosis, each gamete carrying one or the other. Certain complexes are lethal in gametes (gametophytic lethals), while some combinations are lethal zygotes (zygotic lethals). Specific names were assigned to these complexes by Renner, and they are known as Renner complexes.
- **Repetitive DNA** Nucleotide sequences that are present in the genome in many copies.
- **Reproductive isolation** An isolating mechanism of evolution, where a population is isolated by several genetically controlled mechanisms that prevent gene exchange between two populations.
- **Resting stage** Interphase; any nucleus not undergoing division. However, the nucleus is very active metabolically.
- **Restitution** The spontaneous rejoining of broken ends of chromosomes to produce the original chromosome configuration.
- **Restitution nucleus** A single nucleus with unreduced chromosomes is produced through failure of one of the divisions of meiosis.
- **Restorer genes** In cytoplasmic-genetic male sterility, certain male parents carry genes that restore the pollen-producing ability of male-sterility cytoplasm.

- **Restriction enzyme** It recognizes specific short sequences of (usually) unmethylated DNA and cleaves the duplex randomly.
- **Restriction fragment length polymorphism (RFLP)** Inherited differences in sites for restriction enzymes that result in differences in the lengths of the fragments produced by cleavage with the relevant restriction enzyme. RFLPs are used in genetic mapping to link the genome directly to a conventional genetic marker.

Restriction map A linear array of sites on DNA cleaved by various restriction enzymes.

- **Reverse tandem inversions** Two inverted segments are adjacent to each other but are mutually interchanged.
- **Rhizome** An underground horizontal stem with scale leaves and auxiliary buds that serves as a means of vegetative propagation.
- **Ribosome** Small cellular components composed of specialized ribosomal RNA and protein; site of protein synthesis. Responsible for the synthesis of long-chained molecules known as proteins.
- **Ring bivalent** A ring-shaped chromosome association at diakinesis or metaphase-I of meiosis with a chiasma in both arms.
- **Ring chromosome** A physically circular chromosome, produced as a result of chromosomal structural changes, and usually unstable mitotically and meiotically.

RNA Ribonucleic acid; long chain of nucleotides connected by phosphate-to-sugar bonds.

S-period The period of DNA (replication) synthesis during interphase.

S-phase Synthesis of DNA that occurs in the eukaryotic cell cycle.

 S^1 nuclease An enzyme that specifically degrades unpaired (single-stranded) sequences of DNA.

Salivary gland chromosomes In certain tissues of insects (flies, mosquitoes, midges) of Diptera, polytene chromosomes are found in the interphase nuclei. Many extra replications of each chromosome occur in a single nucleus (endopolyploidy), and all lined up together in parallel fashion (polytene), result in thick chromosomes. Discovered and identified first in *Drosophila melanogaster*.

SAT-chromosome A chromosome with a secondary constriction, usually associated with a nucleolus.

- **Satellite** A small-terminal segment of chromosome, mostly associated with the nucleolus and known as the nucleolus organizer chromosome.
- **Satellite DNA** It is highly repititive, mostly found in heterochromatin, and is usually not transcribed. Many satellite DNA sequences are localized to the kinetochore.

Scale leaf A dry rudimentary version of a layer of greener leaves observed on the same plant.

- **Scion** A portion of an actively growing shoot of a plant is grafted onto a stock of another.
- **Scutellum** A part of the embryo of the Gramineae, considered similiar to a cotyledon. It serves as an organ that transfers nutrients from the endosperm to other parts of the germinating embryos.
- **Second division segregation** First meiotic division is equational, and second meiotic division is reductional.

Secondary chromosomes A chromosome consists of both arms homologous (an isochromosome).

- **Secondary constriction** It is usually associated with the regions when the nucleolus is formed or associated (nucleolus organizer).
- **Secondary trisomic** An individual carrying an extra isochromosome in addition to its normal somatic chromosome complement.
- **Seed** The fertilized and ripened ovule with an embryo and generally with a food reserve (endosperm or cotyledons).
- **Segmental allopolyploidy** Characterized by genomes intermediate in degree of similarity and generally exhibiting preferential pairing $(B_1B_1B_2B_2)$.
- **Segregation** The migration of homologous chromosomes into daughter nuclei during meiosis, resulting in separation of the genes due to recombination in the offspring.
- Self-compatible A plant that can be self-fertilized.
- **Self-fertilization** An individual produces seeds by the fusion of male and female gametes from the same plant.
- **Self-incompatibility** An individual is self-sterile, although male and female gametes are produced at the same time and are functional.

Self-production The transfer of pollen to the stigma of the same plant.

Semigamy An embryo develops from an egg cell with sperm penetration but without fertilization.

Semisterility An organism heterozygous for a gene or chromosome (reciprocal translocation) exhibits approximately 50% male and female abortion.

Sequence-tagged sites (STSs) A short stretch of genomic sequence that can be detected by the PCR.Sequencing Determination of the order of nucleotides (base sequences) in a DNA or RNA molecule or the order of amino acids in a protein.

Sex chromosomes The chromosomes that determine the sex of an organism (X and Y chromosomes).

Sex linkage The inheritance of a sex chromosome is coupled with that of a given gene located on the X chromosome.

Sexual reproduction Requires meiosis and fertilization alternating in a cycle.

- **Short-day plant** A plant in which flowering is induced by daily exposure to less than 12 h of light. **Siblings** Progenies from the same plant (parents).
- **Sigma virus in** *Drosophila* Sigma virus is found in the cytoplasm of *Drosophila*, which is cytoplasmically inherited for CO_2 sensitivity. However, it recovers when CO_2 is removed. Sensitivity is caused by a small virus-like particle named "sigma." Maintenance of sigma is genetically controlled.

Silent mutation It does not change the product of a gene.

Simple sequence DNA It is equal to satellite DNA.

Simplex Aaaa.

Single cross Crossing between two genetically different parents.

Sister chromatids The copies of a chromosome produced by its replication during interphase.

SMC Sperm or spore mother cell.

Somatic cell Cells other than the germ cells.

- **Somatic chromosome doubling** Doubling the chromosomes of vegetative tissues of an organism experimentally, such as by colchicine treatment.
- **Somatic mutation** A mutation that occurs in the somatic tissues and is not inherited.

Somatic pairing Synapsis of homologous chromosomes in the somatic cells.

- **Somatoplasm** Tissues essential for the functioning of the organism but lacking the property of entering into sexual reproduction.
- **Somatoplastic sterility** Death of zygotes during embryonic stage due to disturbance in embryoendosperm relationships.
- **Southern blotting** A molecular biology procedure that transfers denatured DNA from an agarose gel to a nitrocellulose filter, where it can be hybridized with a complementary nucleic acid.
- **Speciation** The differentiation of two or more species from originally one species. It may be due to sympatric (without geographical isolation), allelopathic (with geographical isolation), and instantaneous (allopolyploidy) speciation.
- **Species** A kind of plant or animal, its distinctness seen in morphological, anatomical, and cytological and chemical discontinuities, presumably brought about by reproductive isolation in nature from all other organisms. Populations are capable of interbreeding and producing viable fertile offspring.
- **Spermatogonia** These cells are in testes, divide mitotically, and produce a group of diploid primary sporocytes. Primary sporocytes divide meiotically, generating two secondary spermatocytes and producing four spermatids after second mitotic division.
- **Spike** An elongated, unbranched, intermediate inflorescence in which flowers are sessile (without a stalk).
- **Spikelet** In the grass family, each flower is subtended by two bracts, called a spikelet. The central axis of a spikelet is the Rachilla.
- **Spindle** A bipolar-oriented structure made of protein fibers, organized by centrosomes or kinetochores, that functions in orientation, coorientation, and separation of chromosomes during metaphase and anaphase.
- Spindle attachment region Kinetochore.

Spontaneous mutation A normally occurring mutation.

- **Spore** A reproductive cell in plant, product of meiosis, and produces gametes (microspore, mega or macrospore) after three mitoses.
- Sporocyte A spore mother cell.

Sporogenesis Formation of haploid spores in the higher plants.

Sporophyte A part of the life cycle of plants, when spores are produced.

Sporophytic incompatibility The incompatibility relationship of pollen is determined by the plant producing it (heterostyle plants).

Sporulation Formation of spores.

- **Stamen** The pollen-bearing organ of the flower, consisting of filament and anther. It is a basic unit of androecium.
- Staminate A flower with stamens; the carpels are rudimentary or suppressed.

Statistics All values computed from the sample data are known as statistics.

- **Stigma** The region of a gynoecium, usually at the apex of the style on which compatible pollen grains germinate.
- Stolon Horizontal stems produced above ground, and roots at nodes.
- Stomata Pores on the underside of leaves that control gas exchange and water loss.
- Style A portion of gynoecium between ovary and stigma through which pollen tubes grow.
- Subculture Transfer of cultured tissues of an organism to a fresh medium.
- Submedian Kinetochore located nearer one end than the other.
- Subspecies A taxonomic subdivision of species; genomically similar and capable of interbreeding.
- **Supernumerary chromosomes** Chromosomes also known as B-chromosomes or accessory chromosomes. They are largely heterochromatic and are believed to be genetically inert or contain few coding genes.
- Symbiosis Two or more species living together; parasitic or mutualistic.
- Symmetric hybrid Stable allopolyploid hybrid.
- Symmetrical karyotype All chromosomes are about the same size.
- **Sympatric** Two or more species occupy the same geographical area or overlap in their distribution. **Synapsis** Pairing of homologous chromosomes during meiosis.
- **Synaptonemal complex** The morphological structure of synapsed chromosomes under the electron microscope.
- **Syncyte** A multinucleate cell produced by inhibition of cytokinesis in mitosis, migration of nucleus from one cell to another, and fusion of two cells.
- Syndesis Synapsis.
- Synergids The two haploid cells that lie beside the egg cell in a mature embryo sac.
- **Syngamy** The union of male and female gametes during fertilization to produce a zygote sexual reproduction.
- Synizesis Clumping of chromosomes in one side of a nucleus (= synizetic knot).

Syntenic Gene loci located on the same chromosome.

- **Systematics** The classification of an organism on the reconstruction of evolutionary relatedness among living organisms.
- **T-bands** (T = terminal bands); terminal bands are observed after treating chromosomes at high temperatures (87°C), at pH 6.7, and after Giemsa staining shows the telomere of some chromosomes' preferential staining.

Tandem duplication A fragment chromosome is inserted into the partner chromosome.

Tandem repeats Multiple copies of the same sequence.

Tandem satellites Separate constrictions in the larger satellites.

Tapetum Refers to the innermost layer of the pollen-sac wall.

Tassel The staminate inflorescence of maize.

Taxon Refers to the taxonomic group of an organism (family, genus, species).

Taxonomy Systematics; the study of the classification of living organisms.

Telocentric chromosome A chromosome having a terminal kinetochore and one complete arm of a normal chromosome.

Telomere The end of a chromosome arm.

- **Telophase-I** (meiosis) Chromosomes reach their respective poles; the nuclear membrane and the nucleolus start to develop; and eventually, two daughter nuclei each, the haploid chromosomes are generated.
- **Telotrisomic** An individual with a normal chromosome complement plus an extra telocentric chromosome is called telotrisomic.

Terminal chromosomal association The end-to-end chromosome pairing.

Terminalization The movement of chiasma toward the ends of the chromosomes.

Tertiary chromosome A chromosome formed by interchange between nonhomologous chromosomes.

- **Tertiary trisomics** An individual containing an interchanged nonhomologous chromosome in addition to the normal somatic chromosome complement.
- **Test cross** A cross of a heterozygote to the homozygous recessive to determine the heterozygosity or linkage.
- **Tetrad (meiosis)** Four uninucleate cells with a haploid chromosome number produced at the end of meiosis.
- **Tetraploid** An organism having four basic chromosome sets (4x) in the nucleus.
- Tetrasome An organism with one chromosome in the complement represented four times.
- Tetraspory Meiotic karyokinesis occurs, but cytokinesis does not occur.
- **Tissue culture** The maintenance or growth of tissues *in vitro* in a way that may allow further differentiation and preservation of architecture or function or both.
- Totipotency The capacity of a cell cultured *in vitro* to regenerate into a plant.
- **Transgenic** A plant or animal modified by genetic engineering to contain DNA from an external source is called transgenic.
- **Translocation** A chromosomal structural aberration involving the reciprocal exchange of terminal segments of nonhomologous chromosomes.
- **Transposable elements** Gene loci capable of being transposed from one spot to another within and among the chromosomes of the complement.
- **Transposition** The transfer of a chromosome segment to another position due to intra- and interchromosomal structural changes.
- Triplex AAAa.
- **Triploid** An organism with three basic sets (3x) of chromosomes in the nucleus, may be auto- or alloploid.
- Trisomic An organism containing a normal chromosome complement and one extra chromosome.
- Trivalent An association of three homologous chromosomes.
- True-breeding An organism homozygous for a trait.
- **Tuber** A swollen underground stem tip that contains stored food material and serves as a source for vegetative propagation.
- Twining A plant coiling around objects as a measure of support.
- **Twins** Two embryos developed simultaneously within an ovary. Fraternal twins originate from two fertilized eggs. Identical twins arise from only one fertilized egg.
- **Umbel** A flat-topped inflorescence, like sunflower, in which all of the flowers are borne or pedicel of approximately equal length, and all arise from the apex of the main axis.
- Uninemy Single-strandedness of DNA in the chromosome.
- Unisexual (= imperfect) A flower with stamens or carpel but not both.
- Univalent An unpaired chromosome during meiosis.
- **Univalent shift** A meiotic irregularity in monosomics that produces a monosomic plant deficient for a chromosome other than that originally deficient in the monosomic parent.
- **Variance** The extent to which values within frequency distribution depart from the mean is called variance. It is large when values are spread out and small when values are close together.
- **Variegation** Mosaic phenotype, is widespread phenomenon attributed to plastid variation, genetic instability, instability of the phenotypic expession of genes, somatic crossing over, and somatic instability of chromosomes (breakage-fusion-bridge cycle) and transposable elements.
- **Variety** In classical taxonomy, a subdivision of a species, also termed cultivars. A group of individuals distinct in form and function and gentically uniform.
- **Vegetative reproduction** Reproduction of the higher plants, either naturally (runners, bulbs, rhizomes, etc.) or artificially (cutting, grafting, etc.).
- **Viruses** Minuscule agents, of which several kinds have been identified through electron microscope. Shape ranges from round to rod-like to polyhedral with attached tail (T-even bacteriophage). Size ranges from vaccinia virus to as large as a small bacterium. The structure of a virus is like a protein coat containing a single chromosome. Bacteriophage lambda does not kill all the infected bacteria. They are called lysogenic bacteria. The inactive form of a virus is known as the prophage (temperate phage). Virulent phages always cause destruction to the bacterium.
- Vivipary Bearing seeds that germinate within a flower or inflorescence.

Weismannism A concept by August Weismann that proposes that acquired traits are not inherited, and only changes in the germ plasm are transmitted from generation to generation.

Wild type Normal phenotype of an organism.

x A symbol for the basic number of chromosomes in a polyploid species.

X-chromosome A chromosome concerned with the determination of sex.

Xenia Effect of genotype of pollen on embryo and endosperm.

Xerophyte A type of plant of arid habitats.

Xylem A specialized vascular tissue that conducts water and nutrients in plants and provides mechanical support in vascular plants.

Y-chromosome The sex chromosome found only in males in heterozygous condition.

Yeast Yeast is included in fungi that produce asci and ascospores (ascomycetes). Yeast may be haploid or diploid, a unicellular organism that can multiply by budding (*Saccharomyces cerevisiae*).

Zygomorphic The parianth constituted in such a way that only a median plane will yield two equal halves, as in most orchids or the sweet pea flower.

Zygonema (meiosis) The stage at which homologues begin to pair.

Zygote A fertilized egg.

Zygotic or somatic chromosome number 2*n*.

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