



# Overexpression of the Heterochromatinization Factor BAHD1 in HEK293 Cells Differentially Reshapes the DNA Methylome on Autosomes and X Chromosome

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BAH domain-containing protein 1 (BAHD1) is involved in heterochromatin formation and gene repression in human cells. BAHD1 also localizes to the inactive X chromosome (Xi), but the functional significance of this targeting is unknown. So far, research on this protein has been hampered by its low endogenous abundance and its role in epigenetic regulation remains poorly explored. In this work, we used whole-genome bisulfite sequencing (BS-seq) to compare the DNA methylation profile of HEK293 cells expressing low levels of BAHD1 (HEK-CT) to that of isogenic cells stably overexpressing BAHD1 (HEK-BAHD1). We show that increasing BAHD1 levels induces de novo DNA methylation on autosomes and a marked hypomethylation on the X chromosome (chrX). We identified 91,358 regions that have different methylation patterns in HEK-BAHD1 compared to HEK-CT cells (termed "BAHD1-DMRs"), of which 83,850 mapped on autosomes and 7508 on the X chromosome (chrX). Autosomal BAHD1-DMRs were predominantly hypermethylated and located to satellites, interspersed repeats, and intergenic regions. In contrast, BAHD1-DMRs on chrX were mainly hypomethylated and located to gene bodies and enhancers. We further found that BAHD1-DMRs display a higher-order organization by being clustered within large chromosomal domains. Half of these "BAHD1-Associated differentially methylated Domains" (BADs) overlapped with lamina-associated domains (LADs). Based on these results, we propose that BAHD1-mediated heterochromatin formation is linked to DNA methylation and may play a role in the spatial architecture of the genome.

Keywords: cytosine methylation, DNA methylation, whole genome bisulfite sequencing, heterochromatin, epigenetics, LAD, Xi

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# INTRODUCTION

DNA methylation is a heritable epigenetic mark that has critical roles in the regulation of genome structure and transcription in eukaryotes (for reviews see Suzuki and Bird, 2008; Laird, 2010). This modification, catalyzed by DNA methyltransferases (DNMTs), is essential for embryonic development and a number of key processes, such as X chromosome inactivation, genomic imprinting, chromosome stability, and silencing of repetitive elements (Li et al., 1993; Jones and Laird, 1999; Baylin et al., 2001; Geiman and Muegge, 2010). Consistent with these important roles, changes in DNA methylation are associated with several human diseases (Robertson, 2005).

In somatic mammalian cells, DNA methylation mainly occurs at CpG dinucleotides, with ~70% of CpG being methylated (mCpG; Lister et al., 2009). This modification is dynamically regulated in a developmental context across multiple cell types (Ziller et al., 2013) and is commonly associated with gene silencing, especially within CpG islands (CGIs). CGIs in promoters are predominantly unmethylated and can be methylated in a tissue-specific manner (Straussman et al., 2009; Deaton and Bird, 2011). CGI methylation contributes for instance to the mechanism of X chromosome inactivation (Sharp et al., 2011), which results in one inactive X chromosome (Xi) and one active X chromosome (Xa) in mammalian female cells (Nora and Heard, 2010; Pollex and Heard, 2012). Methylation of CGI promoters on Xi, in contrast to Xa, correlates with silencing of ~85% of all Xi genes (Carrel and Willard, 2005; Zhang et al., 2013). However, several studies indicate that DNA methylation is not only functionally linked to gene repression (Suzuki and Bird, 2008; Ndlovu et al., 2011). For instance, a positive correlation between active transcription and gene body methylation has been detected on the Xa and evidence suggest that X-linked gene bodies are less methylated on Xi than on Xa (Viegas-Pequignot et al., 1988; Prantera and Ferraro, 1990; Weber et al., 2005; Hellman and Chess, 2007). A link between active transcription and gene body methylation has also been detected at highly expressed genes on autosomes (Aran et al., 2011). Conversely, in cancer cells, formation of repressive domains coincides with global hypomethylation (Hon et al., 2012).

Mechanisms controlling which DNA sequences become methylated involve recruitment of DNMTs at specific loci, though the driving mechanism is not well understood. There is a complex interplay between DNA methylation and histone modifications. This relationship can be partially mediated by interactions of DNMTs with histone lysine methyltransferases (KMT, such as SETDB1, SUV39-H1, and G9a) and by action of mCpG-binding proteins, such as MECP2 and MBDs, that are capable of recruiting histone deacetylases (HDACs) to the methylated region (for reviews see Bernstein et al., 2007; Cedar and Bergman, 2009; Buck-Koehntop and Defossez, 2013). However, how this scenario unfolds at specific genome loci according to time, cell type and stimuli, is not well characterized. Hence, it is crucial to identify components of chromatinremodeling complexes that bridge DNA methylation, histone modifications and transcription factors.

One such component could be BAHD1, a vertebrate-specific protein that we previously identified as a repressor involved in chromatin compaction (Bierne et al., 2009). Overexpression of BAHD1 in human cells is sufficient to stimulate de novo formation of heterochromatic foci that lack acetylated histone H4. Also, exogenous BAHD1 localizes to the heterochromatic Xi in female cell lines. This heterochromatin factor co-purifies with HDAC1/2, HP1 (α, β, γ), and KAP1/TRIM28 (Lebreton et al., 2011) and interacts with the methyl-CpG-binding protein MBD1 and the KMTs SETDB1 (Bierne et al., 2009) and SUV39-H1 (Weimann et al., 2013) suggesting that it belongs to a multiprotein chromatin-repressive complex ("the BAHD1 complex"; Bierne et al., 2012). Several BAHD1-associated partners are known to interact with DNMT3A (Fuks et al., 2001, 2003; Lehnertz et al., 2003; Li et al., 2006) and KAP1 plays a role in DNA methylation (Quenneville et al., 2011). Collectively, these findings suggest an attractive hypothesis wherein BAHD1 plays a role in the relationship between DNA methylation and chromatin compaction in higher eukaryotes. However, research on this protein has been hampered by its absence or low abundance in mammalian tissues, making it difficult to study its effect on the epigenome.

In this study, we searched for a connection between BAHD1 and DNA methylation by using, as an experimental model, Human Embryonic Kidney 293 (HEK293) cells stably expressing BAHD1. In the widely used HEK293 line, our previous studies showed that BAHD1-associated heterochromatin and multiprotein complexes are formed when BAHD1 expression increases (Bierne et al., 2009; Lebreton et al., 2011). Using whole genome bisulfite sequencing (BS-seq; Lister et al., 2009), we generated methylomes at single-nucleotide resolution of HEK293 cells with endogenous or stable overexpression of BAHD1. Comparison of the methylation profiles of these two lines suggests that BAHD1 contributes to the differential patterning of DNA methylation, both at local (kb) and large (Mb) genome scales, with distinct effects on autosomes and on the X chromosome (chrX). In addition, gene expression data suggest that BAHD1-associated differential methylation could be associated with transcriptional repression.

### MATERIALS AND METHODS

# Cell Lines, Plasmids, Antibodies, and Immunostaining

The stable cell line with constitutive BAHD1 expression (HEK-BAHD1) and its isogenic control (HEK-CT) are described in the Supplementary Material. The HPT-BAHD1 inducible cell line and isogenic HPT-HEK293 control are described in Lebreton et al. (2011). Cells were grown at  $37^{\circ}$ C in a humidified 10% CO<sub>2</sub> incubator, in Dulbecco's modified Eagle's medium with GlutaMAX TM (Gibco) supplemented with 10% FBS (Gibco) and Hygromycin (HygroGold, Invivogen 200 µg/ml). To induce BAHD1 expression in HPT-BAHD1 cells, tetracycline was added to cell culture media ( $11 \mu g/ml$ ) 30 h before cell recovery. Plasmid pYFP-BAHD1, which was used for transient overexpression of BAHD1-YFP, is described in Bierne et al. (2009). Antibodies used

in the study were raised against BAHD1 (Abcam, 46573) and HDAC1 (Abcam, ab7028). Preparation of chromatin extracts is described in Lebreton et al. (2011). Immunofluorescence and XIST FISH assays were carried out as described in Bierne et al. (2009).

# **BS-seq Procedures**

Isolated DNA was truncated into 100-300 bp fragments by sonication followed by DNA-end repair, 3'-dA overhang addition and ligation of methylated sequencing adapters. Samples underwent bisulfite treatment with the ZYMO EZ DNA Methylation-Gold kit. Desalted, size-selected, PCR amplified fragments were size-selected again. Quality controlled libraries were selected for Illumina sequencing by the Beijing Genomics Institute (BGI). Data filtering included removing adaptor sequences, contamination and low-quality reads from raw reads. These reads were filtered by custom BGI scripts, as described in Li et al. (2010). There were two types of low-quality reads removed from the data: (1) when the ratio of N in whole read was over 10%; (2) when the ratio of base whose quality was <20 was over 10%. Observed cytosines on the forward read of each read pair were in silico replaced by thymines, and observed guanines on the reverse read of each read pair were in silico replaced by adenines (Xiang et al., 2010). The "alignment form" reads were then mapped to the "alignment form" reference genome by SOAP aligner. Every hit with a single placement with minimum numbers of mismatches and a clear strand assignment was defined as unambiguous alignment (uniquely mapped reads) and was used for methyl-cytosine determination. Only the uniquely mapped reads were used to estimate the copy numbers of the local region (Xiang et al., 2010). Genomic bases with a copy number larger than 1.5 were not used to call methylcytosines and not used in any subsequent analysis to avoid errors caused by misalignment following the protocol described in Li et al. (2010). For each of the replicates, 1.03 billion (HEK-CT) and 1.05 billion (HEK-BAHD1) paired-end reads of 90 bp length were generated and aligned to the female human reference sequence (NCBI build 37/hg19), yielding 93.0 and 94.3 gigabases (Gb), with an average of 23.4 and 26.1-fold genome coverage. In total,  $\sim$ 38 and  $\sim$ 37 million methylcytosines were detected in HEK-CT and HEK-BAHD1, respectively, where >95% of the queried CpGs had at least one read, and ~85% had more than four reads. Methylcytosines amounted for 3.35 and 3.45% of the cytosines with sequence coverage and ~97% of them were in the CpG context. Rare non-CpG methylation (CHH and CHG) was present throughout all chromosomes (Supplementary Figure S1). Whole-genome BS-seq reads were subjected to data import, smoothing, and DMR analysis with the Bioconductor package bsseq (Hansen et al., 2012). Replicates were tiled over 300 bp windows for statistical analysis where Fisher's exact test was applied for inference with multipletesting adjusted values (Benjamini-Hochberg procedure, p < 0.1). DMRs were defined by subsetting the results based on a regional methylation difference of at least 25% across the 300 bp windows. To investigate a higher order organization, BAHD1specific DMRs were first counted and grouped over windows of 0.5 Mb, defining "BAHD1-DMR clusters." Then, the mapping of

DMR-rich regions was refined by defining contiguous domains of DMR clusters with more than the 75% quantile of DMR counts (i.e., representing the top 25% quantile). These domains were referred to as "BAHD1-Associated differentially methylated Domains" (BADs). The overlap between BADs and LADs was calculated in the R environment using the data from Guelen et al. (2008) accessible via the UCSC genome browser. Full datasets have been deposited in Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo) and are accessible through GEO series accession number GSE51867.

# **Native ChIP-seq Procedures**

The immunoprecipitation of the HPC-BAHD1 protein in HPT-BAHD1 cells with anti-Protein C antibodies is described in Lebreton et al. (2011). A native ChIP (NChIP) of isogenic HPT-HEK293 control cells was performed in parallel. 200 μL of eluted fractions from the first HPC4 affinity column, or  $10\,\mu L$ of input chromatin fractions diluted in 200 µL, were used for DNA extraction from two independent biological HPT-BAHD1 replicates or one HPT-HEK293 sample. DNA was purified by two extractions with 200 µL of phenol:chloroform:isoamylalcohol (25:24:1, pH = 8). Residual phenol was eliminated from the aqueous phase by extraction with chloroform, after which DNA was precipitated by addition of 900 μL of ethanol, 100 μL of ammonium acetate 7.5 M, and 0.2 μL of glycogen (20 μg/μL), incubation at  $-80^{\circ}$ C for 30 min and centrifugation at  $4^{\circ}$ C,  $20,000 \times g$  for 15 min. The pellet was washed once in  $100 \,\mu L$ of ethanol, then suspended in 50 µL of pure water for inputs samples, while 25 µL was used for NChIP samples.

For high-throughput sequencing, libraries were constructed by Fasteris SA (Geneva, Switzerland) from Input and NChIP samples, according to the Illumina guidelines. Independent libraries were produced for two biological replicates of the experiment, R1 and R2, input and control NChIP. The DNA colony template library was sequenced as single reads of 50 bp. Sequencing was multiplexed at three libraries per channel. We acquired a minimum of  $\sim$ 37 million aligned unique reads for HPT-BAHD1 and HPT-HEK293 NChIP and inputs, and used stringent cut-offs for peak calling and statistical analysis of replicate DNAs. Trimmed fastq files were mapped with Bowtie version 0.12.7 to the female human genome (UCSC hg19). Only uniquely mapping reads were used for further analysis; clonal reads were removed. Initial peak calling was performed with CCAT version 3.0 (Xu et al., 2010) with the following parameters: fragmentSize = 150, slidingWinSize = 500, movingStep = 50, isStrandSensitiveMode = 0, minCount = 4, outputNum = 10000, randomSeed = 123456, minScore = 5.0, bootstrapPass = 50. A minimum of five-fold change of NChIP over input was required for peak calling. Further analysis was performed on the regions identified by CCAT with edgeR (Robinson et al., 2010), by binning the genome into 200 and 400 bp non-overlapping windows. Reads were extended of 200 bp to reflect the original size of the fragment. Empty windows and outliers (>400 counts per bin) were excluded from the analysis. edgeR analysis was performed with a generalized linear model, excluding the effect of input and HPT-HEK293 control NChIP from the model. Full datasets have been deposited in the

Gene Expression Omnibus (GEO, http://www.ncbi.nlm.nih.gov/geo) and are accessible through GEO series accession number GSE53372.

## **Data Annotation**

In order to investigate qualitative differences between samples, data were subset for regions corresponding to genomic features of interest. Annotations from Ensembl (version 72) were retrieved using the interface biomaRt, which accesses ensembl via BioMart, a federated system of databases. Annotation was performed using custom scripts and functions built in the IRanges, GenomicFeatures and ChIPpeakAnno. The UCSC CpG island track was imported into the R environment using the package rtracklayer. CpG island shores were defined as those regions up to 2000 bp upstream and downstream of CpG islands. The annotation for the human genome repeat regions was downloaded from the repeat masker website hosted at the Institute for Systems Biology (http://www.repeatmasker.org/cgibin/AnnotationRequest). The lncRNA annotation (UCSC track: lincRNAsTranscripts) was obtained from Cabili et al. (2011).

# **Gene Expression Microarrays**

RNA quality was monitored using Agilent RNA Pico LabChips (Agilent Technologies, Palo Alto, CA). One hundred nanogram of RNA from HEK-CT or HEK-BAHD1 cells, was used as templates for the synthesis of hybridization probes for Affymetrix GeneChip Microarrays (Genechip HuGene 1.0 ST). Hybridization was carried out with three biological replicates according to the expression analysis technical manual with wash and stain kit (Affymetrix). Gene-level expression values were derived from the CEL file probe-level hybridization intensities using the model-based Robust Multichip Average algorithm (RMA; Bolstad et al., 2003). RMA performs normalization, background correction and data summarization. An analysis was performed using the LPE test (Jain et al., 2003) and a p-value threshold of p < 0.05 was used as the criterion for expression. The estimated false discovery rate (FDR) of this analysis was calculated using the Benjamini-Hochberg (BH) procedure in order to correct for multiple comparisons. Results were further subset on absolute median difference >0.2. All data is MIAME compliant and the raw data have been uploaded to the Gene Expression Omnibus database (GEO series accession number GSE51868).

# RNA Extraction, Reverse Transcription, and Quantitative PCR (RT-QPCR)

Total RNA from HEK-CT and HEK-BAHD1 cells was extracted using the RNeasy Mini Kit (Qiagen), from three biological replicates. Genomic DNA was removed by treatment with TURBO DNA-freeTM kit (Ambion). cDNAs were generated from 1 to  $2\mu g$  total RNA using the RT2-HT first strand kit (Qiagen/SABiosciences). Quantitative PCR was performed on Bio-Rad MyIQ or cfx384 devices (Biorad), using SsoFast Evagreen supermix (Biorad), as specified by the supplier, using described BAHD1, IGF2, and GAPDH primers (Bierne et al., 2009). Each reaction was performed in triplicate. Data were analyzed by the  $\Delta\Delta$ Ct method. Target gene expression data were

normalized to the relative expression of the *GAPDH* reference gene.

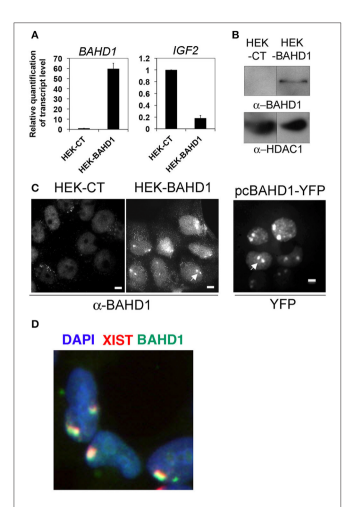
# **RESULTS**

# Generation of a Human Cell Line Stably Producing BAHD1

In order to investigate the effect of BAHD1 overexpression on the dynamics of DNA methylation, we generated a HEK293 cell line with stable expression of BAHD1 (referred to as HEK-BAHD1) by integration of a single copy of the BAHD1 coding sequence under the control of the human cytomegalovirus (CMV) promoter in the HEK293 genome. We also produced an isogenic control line (referred to as HEK-CT), by integration of the empty vector. The increase in BAHD1 mRNA levels by  $\sim$ 60 fold in HEK-BAHD1 cells (Figure 1A) enabled the detection of the BAHD1 protein in HEK-BAHD1 cells, in chromatin extracts (Figure 1B), as well as in nuclei by immunofluorescence microscopy, using BAHD1 antibodies (Figure 1C). In contrast, endogenous BAHD1 was undetectable in control HEK-CT chromatin (Figure 1B), consistent with the low expression of BAHD1 in HEK293 cells and many other cell lines (Bierne et al., 2009). We have previously shown that BAHD1 represses expression of the IGF2 gene in HEK293 cells transiently expressing BAHD1 from a plasmid (Bierne et al., 2009). In agreement with these data, IGF2 mRNA levels decreased by 6 fold in HEK-BAHD1 cells, when compared to control cells (Figure 1A). HEK293 are female cells with an atypical karyotype, with often two copies of the inactive X chromosome (Xi). The two Xi are visible as large heterochromatic bodies (Gilbert et al., 2000) to which BAHD1 is recruited (Bierne et al., 2009). Accordingly, we observed that BAHD1 was enriched at the Xi in HEK-BAHD1 cells, as shown by labeling of BAHD1 and XIST RNA (Figure 1D). The BAHD1 staining was less intense in HEK-BAHD1 cells than in cells transiently transfected with an YFP-BAHD1-expressing plasmid (Figure 1C). Thus, HEK-BAHD1 cells stably expressing BAHD1 can be used as a model system to study whether increasing BAHD1 cellular levels affect DNA methylation.

# **BAHD1** Overexpression Differentially Changes the DNA Methylation Landscape of Autosomes and chrX

In order to obtain high-resolution DNA methylation profiling, we performed whole-genome BS-seq (MethylC-seq/BS-seq Lister et al., 2009) of two HEK-BAHD1 replicates and of the HEK-CT control DNA, generating complete reference methylomes of these cell lines at single base resolution (see Materials and Methods and Supplementary Figure S1A for detailed analytical procedures). We verified that our BS-seq results were in agreement with partial methylomes previously released for HEK293 cells in ENCODE RRBS and HEK293 chromosome 21 datasets (Zhang et al., 2009; ENCODE Project Consortium, 2012; Supplementary Figure S2). Comparison of these methylomes highlighted a significant gain of methylation (on average  $\sim 2 \pm 0.66\%$ ) in autosomes of HEK-BAHD1 cells, when compared



**FIGURE 1 | Constitutive expression of BAHD1 in HEK-BAHD1 cells.** (A) RT-QPCR measurement of BAHD1 and IGF2 mRNA levels in HEK-BAHD1 relative to that in HEK-CT. Bars represent mean $\pm$ SD of three replicates. (B) Immunoblot of chromatin extracts from HEK-CT and HEK-BAHD1 with BAHD1 or HDAC1 antibodies. Endogenous BAHD1 is undetectable in the control line. (C) Immunofluorescence studies of BAHD1 location in HEK-CT, HEK-BAHD1, or in HEK293 cells transfected with a plasmid expressing BAHD1-YFP. Scale bars, 5 μm. BAHD1 localizes to Xi in cells overexpressing BAHD1 from a chromosomal integration or from a plasmid (arrows). (D) Overlay image of an immunoFISH assay with anti-BAHD1 antibodies (green), combined with Xist RNA FISH (red), and staining of nuclei with DAPI (blue).

to autosomes of HEK-CT cells, corresponding to methylation of  $\sim$ 478,000 CpG. In contrast, the average methylation level on chrX decreased by  $\sim$ 4  $\pm$  0.05% (Figure 2A; Table 1). Differential analysis of HEK-CT and HEK-BAHD1 methylome replicates (Fisher's exact test, p < 0.1, BH-corrected, minimum methylation difference 25%) identified 91,358 regions of 300 bp that became reproducibly differentially methylated in the two biological replicates of the HEK-BAHD1 DNA, when compared to the isogenic control. 83,850 of these differentially methylated regions (herein referred to as "BAHD1-DMRs") mapped on autosomes and 7508 on chrX (Figure 2B; Table 2). Relative to chromosome size, the highest enrichment of BAHD1-DMRs was on chrX (Figure 2C; Table 2). In agreement with global methylation levels, 89.5% of BAHD1-specific DMRs identified

on autosomes showed a gain of methylation (referred to as "hyper-DMRs"), whereas 81.8% DMRs on chrX showed a loss of methylation ("hypo-DMRs"). These results indicate that upregulation of BAHD1 induces *de novo* cytosine methylation on autosomes and loss of methylation on chrX.

# Distribution of BAHD1-specific DMRs on Autosomes

We next examined the location of BAHD1-DMRs in different genomic elements on autosomes. The majority of DMRs were principally outside of CpG islands and mapped on intergenic regions, introns and interspersed repeats (LINEs, LTR, SINEs; Figure 2D). Relative to the size of each genomic element, hyper-DMRs were particularly enriched at satellites and other repeat sequences, intergenic regions and lncRNAs, whereas they rarely mapped in 5'UTRs, CGIs, and promoters (Figure 2E; Table 3A). Subsets of hypermethylated BAHD1-DMRs were observed at enhancers (4579) and CGI-shores (1175), which like CGIs, are cis-regulatory modules that play important role in regulation of gene expression (Schilling and Rehli, 2007; Irizarry et al., 2009). The % of mCpG can be divided into low levels (0-25%), "partially methylated domains" (PMD) and high levels (75–100%; Lister et al., 2009; Hon et al., 2012). The directional change in methylation in response to overexpression of BAHD1 went toward a higher methylation (Table 4A).

We searched for the presence of binding sites for BAHD1associated partners in hypermethylated BAHD1-DMRs on autosomes by examining the overlap of DMRs and binding sites for transcription regulators listed in the ENCODE TFBS (Transcription Factor Binding Sites) cluster track. Remarkably, the two first transcription regulators of this analysis (Supplementary Table S1) were SETDB1 and KAP1, two known partners of BAHD1. HP1-y, HDAC2 and the transcription factor SP1, which also co-precipitate with BAHD1 in HEK293 cells (Bierne et al., 2009; Lebreton et al., 2011) were also in this list. Additionally, there were binding sites for EZH2, the H3K27 KMT that induces H3K27me3, a mark associated with BAHD1 heterochromatic foci (Bierne et al., 2009), and STAT transcription factors (STAT1, STAT3, and STAT5A) that can be functionally related to BAHD1-mediated repression of interferon-stimulated genes (Lebreton et al., 2011). These findings support the hypothesis that BAHD1, in association with its partners within macromolecular complexes, is mechanistically linked to the establishment of DNA methylation patterns on autosomes.

# Distribution of BAHD1-specific DMRs on chrX

In contrast to what was observed on autosomes, comparison of methylation profiles on chrX in HEK-CT and HEK-BAHD1 DNA revealed a puzzling loss of methylation on all genomic elements, from the high level to partially methylated level (**Table 4B**), with the exception of satellites that gained methylation like in autosomes. DMR analysis emphasized unique patterns of differential methylation on chrX in BAHD1-overexpressing cells. Relative to the size of each genomic element, DMR counts at enhancers, gene bodies (3'-UTR, introns, exons)

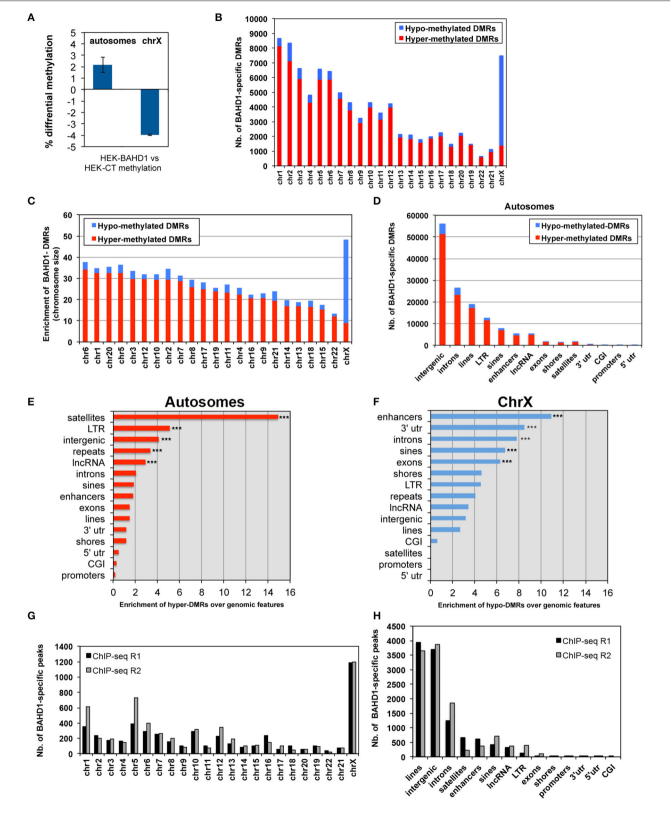


FIGURE 2 | BAHD1-induced differential methylation among chromosomes and genomic elements. (A) Global difference in the percentage of CpG methylation level between HEK-CT and HEK-BAHD1 DNA in autosomes and chrX (methylation level = 100 \* total mC reads/all reads). Results are mean±SD of two (Continued)

### FIGURE 2 | Continued

biological replicates. **(B)** Number of BAHD1-specific DMRs in autosomes and chrX of HEK-BAHD1 cells relative to that in HEK-CT cells. Hypermethylated DMRs: red. Hypomethylated DMRs: blue. **(C)** Enrichment of BAHD1-DMRs relative to autosome size; chrX is shown apart. **(D)** Number of BAHD1-DMRs in genomic elements. **(E,F)** Relative enrichment in genomic elements of BAHD1-hyper-DMRs on autosomes **(E)** or BAHD1-hypo-DMRs on chrX **(F)**. The proportion of DMRs was normalized by the length and abundance of the queried element in the genome. Regions significantly over-represented in differentially methylated regions are indicated (\*\*\*p < 0.001). **(G,H)** Number of BAHD1-specific peaks in HPT-BAHD1 NChIP replicates in chromosomes **(G)** or genomic elements **(H)**.

TABLE 1 | Average percentage of methyl-cytosines per chromosome in HEK-CT and HEK-BAHD1 cells.

Chromosome	HEK-CT	HEK-I	BAHD1	HEK-BAHD1	
		Replicate1	Replicate2	Mean	sd
chr1	70.09	72.06	72.47	72.27	0.29
chr2	67.45	69.2	70.32	69.76	0.79
chr3	67.34	69.01	69.75	69.38	0.52
chr4	59.67	61.31	61.68	61.5	0.26
chr5	61.81	61.66	61.48	61.57	0.13
chr6	68.21	70.46	71.4	70.93	0.66
chr7	67.21	69.13	70.23	69.68	0.78
chr8	60.9	63.21	63.85	63.53	0.45
chr9	71.45	73.1	74.42	73.76	0.93
chr10	65.97	67.84	67.33	67.59	0.36
chr11	66.45	68.06	68.92	68.49	0.61
chr12	66.03	68	68.88	68.44	0.62
chr13	63.38	65.22	65.63	65.43	0.29
chr14	67.08	68.29	69.75	69.02	1.03
chr15	71.58	73.15	74.24	73.7	0.77
chr16	57.3	57.39	60.03	58.71	1.87
chr17	75.16	76.47	77.8	77.14	0.94
chr18	63.89	65.24	66.41	65.83	0.83
chr19	70.6	71.93	73.17	72.55	0.88
chr20	71.71	73.71	75.27	74.49	1.1
chr21	63.06	67.94	67.97	67.96	0.02
chr22	76.1	77.47	79.45	78.46	1.4
Mean autosomes	66.93	68.63	69.57	69.1	0.66
chrX	42.91	38.97	38.9	38.94	0.05

Bold values highlight methylation level in each chromosome in HEK-CT (one BS-seq) and HEK-BAHD1 (mean of two independent BS-seq). These data have been processed as described in Li et al. (2010).

and SINEs showed the highest pattern of hypomethylation on chrX (**Figure 2F**). There were only 4 BAHD1-DMRs on CGIs (all hypo-methylated) and no DMRs on 5'-UTR and promoters (**Table 3B**). Thus, in contrast to autosomes, most BAHD1-DMRs on chrX are enriched in enhancers and gene bodies and are hypomethylated in comparison with the same regions in control cells.

# Relationship Between BAHD1-associated Differential DNA Methylation and Gene Expression

With the aim of investigating the potential association between BAHD1-induced DNA methylation changes and

gene expression, we generated transcriptome datasets of HEK-CT and HEK-BAHD1 lines, using Affymetrix DNA arrays. Analysis of differential gene expression between the two lines yielded 1304 up-regulated and 1137 down-regulated transcripts upon BAHD1 overexpression (LPE, BH-adjusted p < 0.05). Given the role of BAHD1 in transcriptional repression, our analysis principally focused on genes that were down-regulated in HEK-BAHD1 cells. We found BAHD1-specific DMRs in 701 (61%) repressed genes, within a window encompassing 10 kb upstream to 10 kb downstream of each gene. In most autosomes, there was a significantly higher proportion of hyper-DMRs than hypo-DMRs associated with repressed genes (**Figure 3**), suggesting that BAHD1-associated *de novo* DNA methylation on autosomes is mainly associated with gene repression.

On chrX, analysis of the transcriptome datasets identified 119 genes that were differentially expressed in HEK-BAHD1 when compared to HEK-CT cells, of which 72 (60%) were down-regulated. In contrast to autosomes, the majority of chrX genes that were repressed in HEK-BAHD1 cells were associated with hypomethylated DMRs in both BS-seq replicates (i.e., 58/72; Supplementary Table S2; Figure 3). Since BAHD1 targets the Xi (Figure 1), we searched for evidence linking BAHD1associated repression to known methylation changes on the Xi. As mentioned above, there was no BAHD1-specific hyper-DMR on CGIs of chrX (Table 3B), ruling out an effect of BAHD1 in CGI methylation on the Xi. We investigated whether chrX genes that were down-regulated in HEK-BAHD1 were known to be always inactivated on Xi, or could escape X inactivation, by using published data on the status of Xi genes (Zhang et al., 2013). Thirty-two repressed genes were present in this list, of which 13 were classified as "always inactive" and 19 as "heterogeneous" (Supplementary Table S2). Taken together, these results suggest that repression of a set of chrX genes in HEK293 cells overexpressing BAHD1 might be due to loss of methylation on Xi, opening the possibility that BAHD1 might be involved in heterogeneous repression on Xi-linked genes.

# BAHD1-specific DMRs are Clustered into Large Chromosomal Domains

We previously noticed that high-level overexpression of *BAHD1* from a plasmid in HEK293 cells triggers massive compaction of chromatin visible when observed in electron microscopy (Bierne et al., 2009). We hypothesized that BAHD1-mediated heterochromatinization might spread and that this event might coincide with BAHD1-associated DNA methylation changes on large regions. To address this hypothesis, the overall genomic distribution of BAHD1-DMRs was examined at a higher scale by binning DMRs into 0.5 Mb windows (i.e., "BAHD1-DMR clusters"). This analysis revealed that BAHD1-DMRs were

TABLE 2 | Numbers of BAHD1-specific DMRs per chromosome.

chr	Total DMRs	Hyper-DMRs	% Hyper-DMRs	Hypo-DMRs	% Hypo-DMRs	Chromosome length (bp)	Nb. of DMRs relative to chromosome length
chrX	7508	1368	18.2	6140	81.8	1.55E+08	4.84E-05
chr6	6447	5871	91.1	576	8.9	1.71E+08	3.77E-05
chr5	6587	5877	89.2	710	10.8	1.81E+08	3.64E-05
chr20	2232	2055	92.1	177	7.9	6.30E+07	3.54E-05
chr1	8676	8128	93.7	548	6.3	2.49E+08	3.48E-05
chr2	8373	7106	84.9	1267	15.1	2.43E+08	3.44E-05
chr3	6649	5902	88.8	747	11.2	1.98E+08	3.36E-05
chr10	4333	3979	91.8	354	8.2	1.36E+08	3.20E-05
chr12	4263	3988	93.5	275	6.5	1.34E+08	3.19E-05
chr7	4987	4576	91.8	411	8.2	1.59E+08	3.13E-05
chr8	4316	3783	87.7	533	12.3	1.46E+08	2.95E-05
chr17	2288	2014	88	274	12	8.12E+07	2.82E-05
chr11	3638	3157	86.8	481	13.2	1.35E+08	2.70E-05
chr19	1515	1414	93.3	101	6.7	5.91E+07	2.56E-05
chr4	4845	4287	88.5	558	11.5	1.91E+08	2.54E-05
chr21	1152	942	81.8	210	18.2	4.81E+07	2.39E-05
chr9	3259	2907	89.2	352	10.8	1.41E+08	2.31E-05
chr16	2019	1865	92.4	154	7.6	9.04E+07	2.24E-05
chr14	2121	1803	85	318	15	1.07E+08	1.98E-05
chr18	1514	1301	85.9	213	14.1	7.81E+07	1.94E-05
chr13	2151	1929	89.7	222	10.3	1.15E+08	1.87E-05
chr15	1807	1559	86.3	248	13.7	1.03E+08	1.76E-05
chr22	678	613	90.4	65	9.6	5.13E+07	1.32E-05
Total autosomes	83850	75056	89.51	8794	10.49		

This table includes counts for non-overlapping 300 bps windows, which have been found differentially methylated in HEK-BAHD1, when compared to HEK-CT BS-seq, and which localize to individual chromosomes. DMRs are divided in hyper-methylated (more methylation than control: hypor-DMRs) and hypor-methylated (less methylation than control: hypor-DMRs). The results are ranked according to the enrichment of DMRs relative to the chromosome size. Bold Letters highlight values in chrX and in all autosomes.

non-uniformly distributed along the whole human genome (**Figure 4**). In order to refine the mapping of regions with high density of DMRs, we defined contiguous domains in DMR clusters where the DMR counts were in the top quartile of the genome-wide DMR distribution. Examples of such "BAHD1–Associated differentially methylated Domains" (BADs) are shown in **Figure 5**, for chr6 and chrX. Overall, we found 839 BADs with sizes of 0.3–6.5 Mb (median 0.5 Mb), mostly hypermethylated in autosomes (792 "hyper-BADs") and hypomethylated in chrX (26 "hypo-BADs").

Several genome-wide analyses have revealed diverse chromatin domains that play important roles in nuclear organization and function (Hu et al., 2012; Padeken and Heun, 2014). Of these, Lamina-Associated Domains (LADs) are proposed to be dynamic heterochromatic structures located to the nuclear periphery and correlated with gene repression (Guelen et al., 2008; Luperchio et al., 2014). Interestingly, in autosomes, 60% of hyper-BADs overlapped with LADs (p < 0.001, hyperG test). These results highlight the possibility that BAHD1-induced DNA methylation is linked to *de novo* formation of heterochromatic domains at the nuclear periphery. In addition, we noticed that in agreement with the hypermethylation of satellites found in our pattern analyses

(Figure 2; Table 3), a set of hyper-BADs, including those located on chrX, mapped to pericentromeric regions (Figure 5), which are known regions of constitutive heterochromatin. Thus, BAHD1-associated chromatin repressive complexes might play a role in heterochromatinization of pericentromeric satellites by increasing their 5 mC content. The repartition of hypo-BADs on chrX was also non-random. Hypo-BADs particularly mapped in enhancers and gene bodies (introns, exons) and were more abundant on the short arm of chrX (Xp) than on the long arm (Xq), and absent from the central region, in particular on the proximal part of Xq (Figure 5). Overall, these results suggest that overexpression of BAHD1 may dynamically alter heterochromatin topology by formation of large differentially methylated domains with distinct consequences on chrX and autosomes.

# Mapping of BAHD1-occupancy Domains in Response to *BAHD1* Overexpression

We next aimed to find out whether overexpression of BAHD1 stimulates binding of BAHD1 on large chromatin domains. So far, our efforts to immunoprecipitate the endogenous BAHD1 protein using different commercial or custom antibodies has been unsuccessful. To map the genome-wide distribution

TABLE 3 | Number and relative enrichment of BAHD1-specific DMRs in genomic elements in autosomes and X chromosome.

		% Hypo-DMRs on all autosomal DMRs	Hypo-DMRs autosomes	Genomic elements	Enrichment	% Hyper-DMRs on all autosomal DMRs	Hyper-DMRs autosomes	Genomic elements
LTR         11560         90.9         5.1         Enhancers         1005         18           Intergenic         51242         91         4.1         Repeats         4397         9.7           Repeats         40871         90.3         3.4         Intergenic         5086         9           IncRNA         4935         90.3         2.9         CGI         72         55.4           Introns         23132         86.6         2.1         Shores         299         20.3           Sines         6920         88.4         1.8         Exons         285         14.8           Lines         17105         90.2         1.5         Introns         3568         13.4           Exons         1646         85.2         1.5         IncRNA         529         9.7           4'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites							MES	A. AUTOSON
Intergenic   51242	9.1 <b>0.5</b>	9.1	1151	LTR	14.9	99.1	1446	Satellites
Repeats	18 <b>0.4</b>	18	1005	Enhancers	5.1	90.9	11560	LTR
IncRNA	9.7 <b>0.4</b>	9.7	4397	Repeats	4.1	91	51242	Intergenic
Introns         23132         86.6         2.1         Shores         299         20.3           Sines         6920         88.4         1.8         3'-UTR         78         18.8           Enhancers         4579         82         1.8         Exons         255         14.8           Lines         17105         90.2         1.5         Introns         3568         13.4           Exons         1646         85.2         1.5         IncRNA         529         9.7           3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genemic         Hyper-DMRs         % Hyper-DMRs on all chrX DMR         Enrichment elements         chrX         all chrX DM           BX CHROWS         280         100	9 0.4	9	5086	Intergenic	3.4	90.3	40871	Repeats
Sines         6920         88.4         1.8         3'-UTR         78         18.8           Enhancers         4579         82         1.8         Exons         285         14.8           Lines         17105         90.2         1.5         Introns         3568         13.4           Exons         1646         85.2         1.5         IncRNA         529         9.7           3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic         Hyper-DMRs         % Hyper-DMRs on all chrX DMRs         Enrichment elements         chrX         4Hypo-DMRs         % Hypo-DMRs           B.X CHROW-VOWE         Statellites         280         100         38.7         Enhancers         1179         91.5           Shores </td <td><b>5</b>5.4 <b>0.3</b></td> <td>55.4</td> <td>72</td> <td>CGI</td> <td>2.9</td> <td>90.3</td> <td>4935</td> <td>IncRNA</td>	<b>5</b> 5.4 <b>0.3</b>	55.4	72	CGI	2.9	90.3	4935	IncRNA
Enhancers         4579         82         1.8         Exons         285         14.8           Lines         17105         90.2         1.5         Introns         3568         13.4           Exons         1646         85.2         1.5         IncRNA         529         9.7           3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic Hyper-DMRs         Myper-DMRs and the remaining and the remain	20.3 <b>0.3</b>	20.3	299	Shores	2.1	86.6	23132	Introns
Lines         17105         90.2         1.5         Introns         3568         13.4           Exons         1646         85.2         1.5         IncRNA         529         9.7           3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic         Hyper-DMRs         % Hyper-DMRs on all chrX DMRs         Enrichment elements         Genomic chrX         Hypo-DMRs         % Hypo-DMRs           elements         chrX         all chrX DMRs         Enrichment elements         ChrX         all chrX DM           Extellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic <t< td=""><td>18.8 <b>0.3</b></td><td>18.8</td><td>78</td><td>3'-UTR</td><td>1.8</td><td>88.4</td><td>6920</td><td>Sines</td></t<>	18.8 <b>0.3</b>	18.8	78	3'-UTR	1.8	88.4	6920	Sines
Exons         1646         85.2         1.5         IncRNA         529         9.7           3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic Hyper-DMRs         Hyper-DMRs on all chrX DMRs         Enrichment elements         Genomic chrX         Hypo-DMRs         % Hypo-DMRs elements         hypo-DMRs         % Hypo-DMRs elements         hypo-DMRs         % Hypo-DMRs elements         chrX         all chrX DM         hypo-DMRs         hypo-DMRs         % Hypo-DMRs         hypo-DMRs         hypo-DMRs         # Hypo-DMRs         hypo-DMRs         # Hypo-DMRs         hypo-DMRs         # Hypo-DM	14.8 <b>0.3</b>	14.8	285	Exons	1.8	82	4579	Enhancers
3'-UTR         337         81.2         1.2         Sines         904         11.6           Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic chrX         Hyper-DMRs on all chrX DMRs         Enrichment elements         chrX         all chrX DMRs           Filter on the chrX         all chrX DMRs         Enhancers         1179         91.5           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Integenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1	<b>13.4 0.3</b>	13.4	3568	Introns	1.5	90.2	17105	Lines
Shores         1175         79.7         1.2         Lines         1848         9.8           5'-UTR         34         79.1         0.5         5'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic Hyper-DMRs         Myper-DMRs on all chrX DMRs         Enrichment elements         Genomic chrX         Hypo-DMRs         % Hypo-DMRs on all chrX DMRs           B. X CHROWSOME           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Integenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138	9.7 <b>0.3</b>	9.7	529	IncRNA	1.5	85.2	1646	Exons
5'-UTR         34         79.1         0.5         6'-UTR         9         20.9           CGI         58         44.6         0.3         Satellites         13         0.9           Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic         Hyper-DMRs         % Hyper-DMRs on all chrX DMRs         Enrichment elements         Genomic chrX         Hypo-DMRs whyper-DMRs on all chrX DMRs         Genomic elements         Hypo-DMRs chrX         % Hypo-DMRs all chrX DMR           B. X CHROMOSOME         Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9 <td>11.6 <b>0.2</b></td> <td>11.6</td> <td>904</td> <td>Sines</td> <td>1.2</td> <td>81.2</td> <td>337</td> <td>3'-UTR</td>	11.6 <b>0.2</b>	11.6	904	Sines	1.2	81.2	337	3'-UTR
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Promoters         76         89.4         0.2         Promoters         9         10.6           Genomic elements         Hyper-DMRs chrX         % Hyper-DMRs on all chrX DMRs         Enrichment elements         Genomic chrX         Hypo-DMRs delements         % Hypo-DMRs delements         % Hypo-DMRs chrX DMRs           B. X CHROMOSOME           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8 <td>20.9 <b>0.1</b></td> <td>20.9</td> <td>9</td> <td>5'-UTR</td> <td>0.5</td> <td>79.1</td> <td>34</td> <td>5'-UTR</td>	20.9 <b>0.1</b>	20.9	9	5'-UTR	0.5	79.1	34	5'-UTR
Genomic elements         Hyper-DMRs chrX         % Hyper-DMRs all chrX DMRs         Enrichment elements         Genomic elements         Hypo-DMRs chrX         % Hypo-DMRs all chrX DM elements           B. X CHROWOSOME           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic	0.9 0.1	0.9	13	Satellites	0.3	44.6	58	CGI
elements         chrX         all chrX DMRs           B. X CHROMOSOME           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         <	10.6 <b>0</b>	10.6	9	Promoters	0.2	89.4	76	Promoters
B. X CHROMOSOME           Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	o-DMRs on Enrichme	% Hypo-DMRs on	Hypo-DMRs	Genomic	Enrichment	% Hyper-DMRs on	Hyper-DMRs	Genomic
Satellites         280         100         38.7         Enhancers         1179         91.5           Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	rX DMRs	all chrX DMRs	chrX	elements		all chrX DMRs	chrX	elements
Shores         39         22         1.3         3'-UTR         87         89.7           Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9							OSOME	B. X CHROM
Intergenic         970         26.9         1.2         Introns         3284         89.9           Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	91.5 <b>10.9</b>	91.5	1179	Enhancers	38.7	100	280	Satellites
Repeats         942         22.8         1.2         Sines         1024         88.8           3'-UTR         10         10.3         1         Exons         248         89.9           Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	<b>89</b> .7 <b>8.5</b>	89.7	87	3'-UTR	1.3	22	39	Shores
3'-UTR       10       10.3       1       Exons       248       89.9         Enhancers       109       8.5       1       Shores       138       78         LTR       141       17.3       0.9       LTR       672       82.7         Introns       367       10.1       0.9       Repeats       3193       77.2         Lines       318       22.1       0.8       IncRNA       191       88         Sines       129       11.2       0.8       Intergenic       2634       73.1         Exons       28       10.1       0.7       Lines       1123       77.9	<b>89.9 7.8</b>	89.9	3284	Introns	1.2	26.9	970	Intergenic
Enhancers         109         8.5         1         Shores         138         78           LTR         141         17.3         0.9         LTR         672         82.7           Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	88.8 <b>6.7</b>	88.8	1024	Sines	1.2	22.8	942	Repeats
LTR     141     17.3     0.9     LTR     672     82.7       Introns     367     10.1     0.9     Repeats     3193     77.2       Lines     318     22.1     0.8     IncRNA     191     88       Sines     129     11.2     0.8     Intergenic     2634     73.1       Exons     28     10.1     0.7     Lines     1123     77.9	<b>6.3</b>	89.9	248	Exons	1	10.3	10	3'-UTR
Introns         367         10.1         0.9         Repeats         3193         77.2           Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	78 <b>4.6</b>	78	138	Shores	1	8.5	109	Enhancers
Lines         318         22.1         0.8         IncRNA         191         88           Sines         129         11.2         0.8         Intergenic         2634         73.1           Exons         28         10.1         0.7         Lines         1123         77.9	82.7 <b>4.5</b>	82.7	672	LTR	0.9	17.3	141	LTR
Sines     129     11.2     0.8     Intergenic     2634     73.1       Exons     28     10.1     0.7     Lines     1123     77.9	77.2 <b>4.1</b>	77.2	3193	Repeats	0.9	10.1	367	Introns
Exons 28 10.1 <b>0.7</b> Lines 1123 77.9	88 <b>3.4</b>	88	191	IncRNA	0.8	22.1	318	Lines
	73.1 <b>3.1</b>	73.1	2634	Intergenic	0.8	11.2	129	Sines
IncRNA 26 12 <b>0.5</b> CGI 4 100	77.9 <b>2.7</b>	77.9	1123	Lines	0.7	10.1	28	Exons
	100 <b>0.6</b>	100	4	CGI	0.5	12	26	IncRNA
5'-UTR 0 0 <b>0</b> 5'-UTR 0 0	0 0	0	0	5'-UTR	0	0	0	5'-UTR
CGI 0 0 <b>0</b> Promoters 0 0	0 0	0	0	Promoters	0	0	0	CGI
Promoters 0 0 <b>0</b> Satellites 0 0	<b>0</b>	0	0	Satellites	0	0	0	Promoters

This table includes counts for non-overlapping 300 bps windows, which have been found differentially methylated in HEK-BAHD1, when compared to HEK-CT BS-seq, and which localize to individual genomic elements. DMRs are divided in hyper-methylated (more methylation than control, "hyper-DMRs") and hypo-methylated (less methylation than control, "hypo-DMRs"). The results are ranked according to the enrichment relative to the size of each genomic feature, as highlighted in bold.

of BAHD1 binding sites in response to transient induction of *BAHD1* expression, we used a HEK293 cell line stably expressing tetracycline-inducible His6-Protein C (HPC)-tagged BAHD1 (referred to as HPT-BAHD1 cells) that we generated earlier (Lebreton et al., 2011). The genomic DNA present in chromatin extracts from HPT-BAHD1 cells before (input) and after immunoprecipitation (ChIP) of the HPC-BAHD1 protein with Protein C antibodies was analyzed by high-throughput sequencing. This Native ChIP-seq (NChIP-seq) enabled the analysis of the DNA to which BAHD1 binds without any requirement for formaldehyde crosslinking. To filter out non-specific DNA binding on the affinity matrix, we

also performed NChIP-seq analysis of HPT-HEK293 control chromatin extracts. A stringent analysis yielded 4936 and 5742 BAHD1-binding events ("BAHD1-peaks") across the genome in replicates 1 and 2, respectively. Notably, the distribution of BAHD1-peaks across chromosomes showed enrichment of BAHD1 binding sites on chrX, with 24% (replicate 1) and 21% (replicate 2) of all BAHD1-peaks mapping on this chromosome (**Figure 2G**). This result is in agreement with enrichment of BAHD1-DMRs on chrX and with the recruitment of BAHD1 to the Xi. Regarding the distribution on genomic elements, 80% of BAHD1-peaks preferentially mapped at interspersed repeats LINEs and SINEs (37.4  $\pm$  5.9%), intergenic regions (33.2  $\pm$  0.1%),

TABLE 4 | Average difference in global methylation levels per genomic elements between HEK-CT and HEK-BAHD1 cells.

	Average Difference between HEK-BAHD1 and HEK-CT					
	Low (0-25%)	PMD (25-75%)	High (75-100%)			
A. AUTOSON	MES					
Satellite	-6.1	-1.52	7.61			
LTR	-1.85	-4.21	6.06			
Intergenic	-1.6	-4.25	5.85			
IncRNA	-1.47	-4.35	5.82			
Lines	-2.07	-3.18	5.24			
Enhancers	-0.13	-3.26	3.38			
Introns	-0.43	-2.88	3.29			
Sines	-0.61	-2.41	3.01			
Shores	0.04	-2.6	2.54			
Exons	0.25	-2.02	1.78			
Promoters	0.27	-1.9	1.62			
3'-UTR	-0.1	-0.62	0.7			
CGI	0.53	-1.06	0.52			
5'-UTR	0.68	-0.94	0.25			
B. chrX						
Satellite	-9.46	-10.96	20.68			
LTR	0.47	5.31	-5.8			
Intergenic	0.2	4.54	-4.76			
IncRNA	2.85	4.24	-7.09			
Lines	0.31	2.79	-3.11			
Enhancers	-1.11	15.11	-14.01			
Introns	0.51	11.9	-12.41			
Sines	0.34	12.09	-12.43			
Shores	-4.94	14.8	-9.86			
Exons	-2.59	12.54	-9.96			
Promoters	-7.27	13.69	-6.43			
3'-UTR	1.15	13.42	-14.57			

The methylation level ranges were defined into low levels (0–25%), partially methylated domains (25–75%), and high levels (>75%). The average difference indicates subtraction of HEK-BAHD1 (replicates 1 and 2) from HEK-CT methylation levels, with positive and negative differences in red and blue boxes, respectively.

12.21

12.93

-1.81

-2.22

-10.41

-10.71

CGI 5'-UTR

and introns (13.6  $\pm$  3.4%; **Figure 2G**). Relative to the size of each genomic element, BAHD1-peaks were, like BAHD1-DMRs, mostly enriched at satellites (**Table 5**).

Binning BAHD1-specific peaks into 0.5 Mb windows revealed that they clustered into larger regions, as BAHD1-DMRs. Examples of such "BAHD1-occupancy domains" are shown for chr6, chr7, and chrX on **Figure 6A**. Surprisingly, overlapping BAHD1-occupancy and-differentially methylated regions revealed a marked difference between autosomes and chrX. On autosomes, BAHD1-binding and-hypermethylated domains often overlapped or were adjacent. In contrast, on chrX, there was an inverse correlation between the location of BAHD1-binding and hypomethylated domains (**Figure 6B**). Furthermore, BAHD1-occupancy domains were more abundant on the long

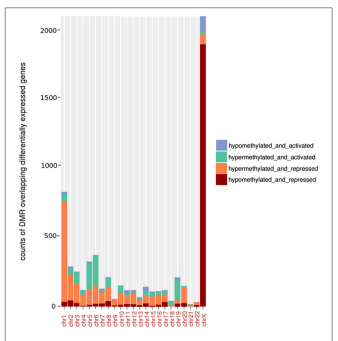


FIGURE 3 | Relationship between BAHD1-DMRs and gene expression. Histograms represent, for each autosome and chrX, the number ("counts") of hypermethylated or hypomethylated DMRs that are located within a window of 10 kb upstream to 10 kb downstream of a gene differentially expressed (activated or repressed) in HEK-BAHD1 cells compared to HEK-CT.

arm (Xq) than the short arm of chrX (Xp), particularly in the half part of Xq containing the X-inactivation center (XIC) on the Xi. Taken together, these data are in agreement with the hypothesis that BAHD1 plays distinct roles on chrX and autosomes and support a model in which BAHD1-mediated chromatin compaction coincides with binding of BAHD1 on large genomic regions.

# DISCUSSION

In this study, we report that increasing BAHD1 cellular levels in human embryonic 293 cells is sufficient to trigger profound changes in CpG methylation patterns at both the local (kb) and large (Mb) genome scale. In addition, BAHD1-associated methylation signatures are distinct depending on location, with gain of methylation on autosomes and loss of methylation on chrX. We cannot exclude that these changes are indirect consequences of BAHD1 expression at non-physiologically high levels. Yet, our experimental data provide insight into potential functions of BAHD1 in epigenetic regulation. We hypothesize a role for BAHD1 in the modulation of DNA methylation patterning and in the spatial organization of silent chromatin.

Our results indicate that BAHD1-specific hypermethylated DMRs are enriched in binding sites for the known BAHD1-associated partners SETDB1, KAP1, HP1- $\gamma$  and HDAC2, as well as for the H3K27me3 writer EZH2, which are all known to interact with *de novo* DNMTs (Fuks et al., 2001, 2003; Lehnertz et al., 2003; Li et al., 2006; Vire et al., 2006; Rush et al., 2009).

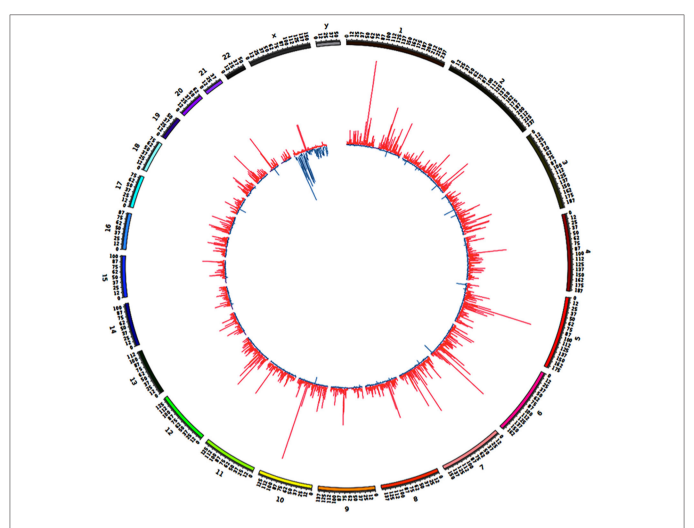


FIGURE 4 | Distribution of BAHD1-DMR clusters across the genome. DMRs present in HEK-BAHD1 cells in comparison to HEK-CT cells were binned into 0.5 Mb windows as "BAHD1-DMR clusters" represented by bars. The distribution of clusters across the genome of HEK-BAHD1 cells, with hypermethylated clusters in red and hypomethylated clusters in blue, is plotted with Circos (Krzywinski et al., 2009).

Also of interest is the presence in a subset of BAHD1-DMRs of binding sites for the transcription factor SP1, with which BAHD1 co-immunoprecipitates (Bierne et al., 2009), as well as for STAT factors that may act in synergy with BAHD1 to repress immunity gene expression during a bacterial infection (Lebreton et al., 2011). In addition, the comparison of HEK293 and HEK-BAHD1 transcriptomes suggests a relationship between the location of BAHD1-DMRs and BAHD1-associated repressed genes. It is worth mentioning that BAHD1 has similarities to the metastasis-associated proteins (MTAs) of the Nucleosome Remodeling and Deacetylation (NuRD) co-repressor complex (Lai and Wade, 2011), which has been shown to couple histone deacetylation and DNA methylation (Morey et al., 2008; Latos et al., 2012). Like MTAs, BAHD1 is a BAH domain-containing protein that binds a reader of 5-methylcytosines (i.e., MBD1, Bierne et al., 2009). BAHD1 might thus act as a scaffold protein that bridges MBD1 and HDAC1/2, as MTAs, and also connects HP1 and KMTs, which can recruit DNMTs. These data converge to a model in which BAHD1-associated multiprotein complexes act in synergy with the DNA methylation machinery to form repressive chromatin.

When transiently overexpressed from plasmid-based expression systems, BAHD1 promotes the formation of nuclear heterochromatic foci (Bierne et al., 2009). These foci seem to correspond to different heterochromatin subtypes, as they are enriched with HP1- $\alpha$ , a marker of constitutive heterochromatin, or H3K27me3, a feature of facultative heterochromatin, but HP1- $\alpha$  and H3K27me3 do not strictly overlap in these foci (Bierne et al., 2009). BAHD1 is also recruited to Xi, a paradigm of facultative heterochromatin. In addition, BAHD1 interacts with chromatin regulators that play a role in the formation of both constitutive and facultative heterochromatin (i.e., HP1 proteins  $\alpha$ ,  $\beta$  and  $\gamma$ , KAP1, HDAC1/2, SETDB1). Here we show that stable expression of BAHD1 in HEK293 cells induces *de novo* DNA methylation at different genomic elements in autosomes, predominantly at satellites, the major

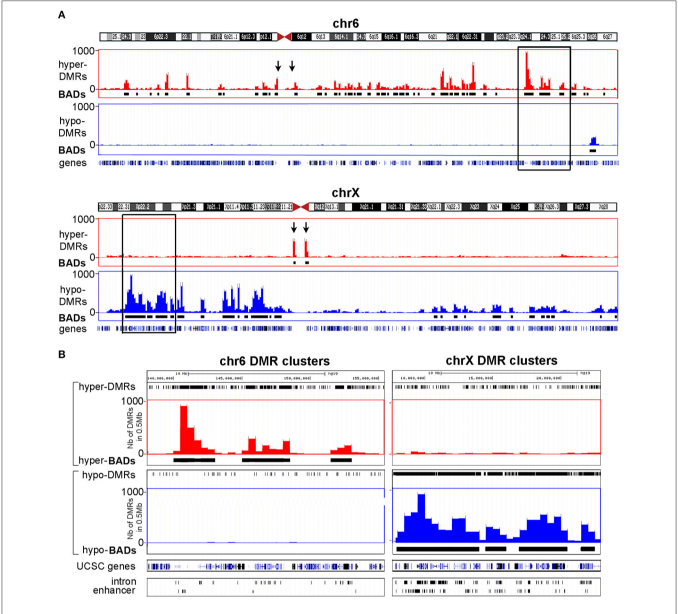


FIGURE 5 | BAHD1-specific domains (BADs). Large-scale genomic organization of DMRs present in HEK-BAHD1 cells in comparison to HEK-CT cells is illustrated for an autosome (chr6) and for chrX. Scale indicates the number of DMRs per 0.5 Mb window. Bars represent clusters of DMRs with gain of methylation (hyper-DMRs, in red) or loss of methylation (hypo-DMRs, in blue). BADs (indicated by black boxes) correspond to contiguous domains of DMR clusters where DMR counts are in the top quartile of the genome-wide DMR distribution. The position of genes is shown below. (A) Chromosome-wide representation of hypermethylation and hypomethylation domains on chr6 (top) and chrX (bottom). Arrows indicate the position of pericentromeric regions. (B) Magnification of the domain squared in A, with a chr6 hypermethylation domain (left) and a chrX hypomethylation domain (right). The position of DMRs in introns and enhancers is shown below.

component of the pericentromeric heterochromatin, and other regions prone to being packaged into heterochromatin (i.e., interspersed repeats, intergenic regions, introns and sequences transcribed in non-coding RNAs). BAHD1 overexpression also increases DNA methylation at a set of cis-regulatory modules, including enhancers and CGI shores. Several studies suggest that methylation of these modules may play a more significant role in the regulation of gene expression than CGIs (Irizarry et al., 2009; Hon et al., 2013; Long et al., 2013; Ziller et al., 2013).

We hypothesize that BAHD1-associated chromatin repressive complexes may recruit DNMTs to non-coding DNA repeats, thus contributing to the stabilization of constitutive heterochromatin, and/or to cis-regulatory modules, and thereby compacting these key regulatory elements into facultative heterochromatic domains leading to gene silencing. However, it is also possible that upon BAHD1 overexpression, BAHD1-interacting proteins are delocalized from various endogenous complexes, which could indirectly alter the DNA methylome.

TABLE 5 | Number and relative enrichment of BAHD1-specific peaks in genomic elements in HPT-BAHD1 NChIP-seq replicates.

Genomic element	Nb. of BAHD1 peaks	Nb. of BAHD1 peaks	Replicate 1	Replicate 2	Mean
	Replicate 1	Replicate 2	enrichment	enrichment	enrichment
Satellites	675	234	2.183	0.757	1.47
Lines	3940	3647	0.151	0.14	0.1455
Intergenic	3691	3859	0.087	0.09	0.0885
IncRNA	326	369	0.064	0.072	0.068
Enhancers	613	382	0.08	0.05	0.065
Introns	1240	1864	0.038	0.058	0.048
Sines	416	724	0.019	0.032	0.0255
LTR	131	391	0.011	0.034	0.0225
Promoters	11	24	0.009	0.02	0.0145
Exons	34	116	0.006	0.02	0.013
3'-UTR	4	20	0.004	0.018	0.011
Shores	13	31	0.004	0.011	0.0075
5'-UTR	1	5	0.002	0.011	0.0065
CGI	1	0	0.001	0	0.0005

Mean of enrichement in biological replicates is highlighted in bold.

NChIP-seq analysis of BAHD1-binding sites suggests that BAHD1 spreads on chromatin in BAHD1-overexpressing cells (Figure 6). This spreading might coincide with propagation of DNA methylation, as we identified regions with high densities of BAHD1-DMRs. Indeed, BAHD1-DMRs are non-uniformly distributed along the genome and group into large chromosomal domains, termed BADs. We find that 60% of BADs coincide with LADs, which are large genomic segments anchored at the nuclear periphery. LADs are proposed to be nuclear compartments of silent chromatin formed in response to development stimuli (Guelen et al., 2008; Luperchio et al., 2014; Padeken and Heun, 2014). The size distribution (0.1-10 Mb) and the median sequence length (0.5-0.8 Mb) of LADs are very similar to that of BADs (0.3-6.5 Mb; median 0.5 Mb). Furthermore, our previous electron microscopy experiments highlighted BAHD1-induced heterochromatin at the nuclear periphery (Bierne et al., 2009). From this, we propose that BAHD1 may mediate chromatin reorganization into de novo heterochromatic regions through DNA methylation changes, thereby contributing to the threedimensional organization of chromatin within the nucleus, in response to developmental, physiological or environmental stimuli. Interestingly, a recent study compiling data from a 100 human epigenomes highlights the dynamics of mCpG distribution across tissue and cell types and emphasizes the importance of examining this dynamics at the megabase-scale (Roadmap Epigenomics et al., 2015). Taken together, these results call for future research to investigate whether BADs are formed in response to activation of the BAHD1 endogenous gene by external stimuli. Such stimuli, and their effect on BAHD1 expression level in different tissues, are presently unknown. It is therefore, necessary to characterize the cellular signaling pathways that control BAHD1 expression and association with other proteins.

In this work, we also show that in HEK293 female cells overexpressing BAHD1, chrX undergoes a significant loss of DNA methylation, while being a binding hotspot for BAHD1.

We cannot assign these methylation changes to specific Xi or Xa territories because the data were derived from mixed cell and chromosome populations. However, since BAHD1 is recruited to Xi, as shown by microscopy, it is tempting to speculate that overexpression of BAHD1 triggers a loss of DNA methylation on the Xi. This observation must be put into perspective within the specific context of the differential methylation of Xi compared to Xa chromosomes in female cells. Different pieces of evidence suggests that DNA methylation patterns are inverted on Xi and Xa, with CGIs and gene bodies being more and less methylated, respectively (Viegas-Pequignot et al., 1988; Prantera and Ferraro, 1990; Weber et al., 2005; Hellman and Chess, 2007). Up to now, a mechanistic explanation of the role of gene body methylation and gene silencing on Xi is still missing. We found that BAHD1specific DMRs on chrX are clustered into hypomethylated domains on gene bodies, particularly on the short arm of chrX (Xp), in contrast with BAHD1-occupancy domains, which are abundant on the long arm (Xq). This puzzling observation opens the possibility, among other hypotheses, that overexpressed BAHD1 may sequester DNMTs at certain sites of Xq, altering X-gene body methylation in cis on Xp. Interestingly, two recent studies point to a possible role for BAHD1 at Xi: BAHD1 co-purifies with CDYL, a new partner of Xi (Escamilla-Del-Arenal et al., 2013) and BAHD1 interactors MBD1 and SETDB1 were found to contribute to the maintenance of X inactivation in somatic cells (Minkovsky et al., 2014). Overall, this work establishes a basis for future studies aimed at exploring the role of BAHD1 in transcription/maintenance/stability of the Xi.

# CONCLUSIONS

De novo changes in DNA methylation patterns induced by the stable overexpression of BAHD1 in HEK293 cells suggest that BAHD1 may play a role in the formation of distinct types of heterochromatic domains by controlling the setting and distribution of mCpG. We did not demonstrate that the

Role of BAHD1 in DNA Methylation

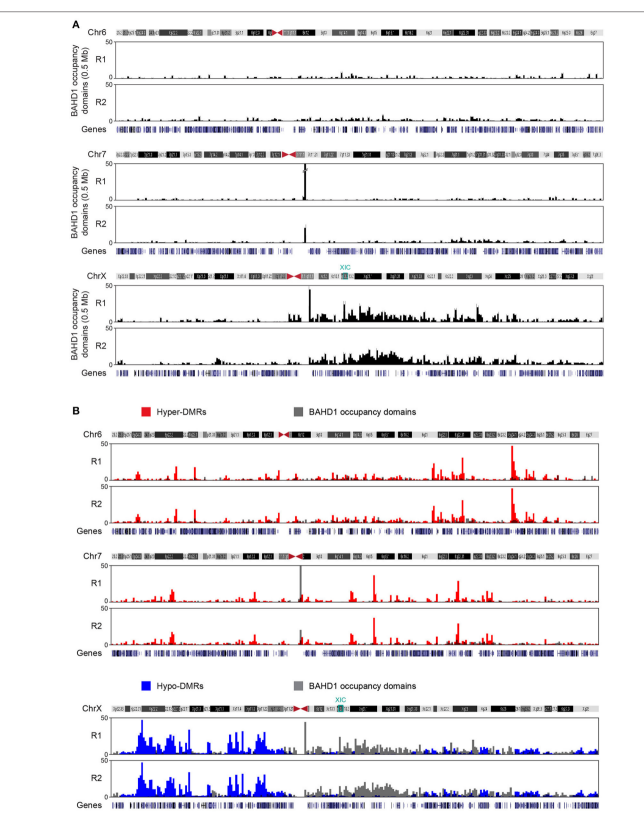


FIGURE 6 | BAHD1-specific binding domains. (A) Large-scale genomic organization of BAHD1-binding sites ("occupancy domains") upon induction of BAHD1 expression for 30 h in HPT-BAHD1 cells. Sequencing DNA from NChIP of Protein-C-tagged BAHD1 generated peaks that were clustered in 0.5 Mb windows (black bars). Results are shown for two autosomes (chr6, chr7) and chrX. Scale indicates the number of peaks per bar. (B) Overlap of BAHD1-DMR clusters and BAHD1-occupancy domains on chr6 (hypermethylated DMRs) and chrX (hypomethylated DMRs). The position of XIC and genes is indicated.

DNA methylation changes observed in this study are a direct consequence of BAHD1 binding to the chromatin. To confirm these hypotheses, it is now essential to study whether the DNA methylome is remodeled upon inactivation of the endogenous *BAHD1* gene. The use of novel techniques to generate knockout cells or animals, such as the CRISPR/Cas9 technology, should facilitate the achievement of these goals, opening an exciting new area of future research on this epigenetic regulator.

# **AUTHOR CONTRIBUTIONS**

EL analysed the BS-seq, ChIP-seq, and transcriptome data and wrote the manuscript. AL participated in sample collections and analyzed and drafted the ChIP-seq data. GL generated the cellular model and participated in sample collections and QPCR. MD and JC participated in data analysis and drafted the BS-seq data. SB and PC participated in data analysis and obtained funding. HB conceived the study, analyzed the BS-seq, ChIP-seq, and transcriptome data, obtained funding and wrote the manuscript. All authors read and approved the final manuscript.

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# SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: http://journal.frontiersin.org/article/10.3389/fgene. 2015.00339

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