Inhibition of DNA methyltransferases blocks mutant huntingtin-induced neurotoxicity

Yanchun Pan  
*Washington University School of Medicine in St. Louis*

Takuji Daito  
*Washington University School of Medicine in St. Louis*

Yo Sasaki  
*Washington University School of Medicine in St. Louis*

Yong Hee Chung  
*Washington University School of Medicine in St. Louis*

Xiaoyun Xing  
*Washington University School of Medicine in St. Louis*

*See next page for additional authors*

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**Recommended Citation**

Pan, Yanchun; Daito, Takuji; Sasaki, Yo; Chung, Yong Hee; Xing, Xiaoyun; Pondugula, Santhi; Swamidass, S. Joshua; Wang, Ting; Kim, Albert H.; and Yano, Hiroko, "Inhibition of DNA methyltransferases blocks mutant huntingtin-induced neurotoxicity." *Scientific Reports*. 6, 31022. (2016).  
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Authors
Yanchun Pan, Takuji Daito, Yo Sasaki, Yong Hee Chung, Xiaoyun Xing, Santhi Pondugula, S. Joshua Swamidass, Ting Wang, Albert H. Kim, and Hiroko Yano
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Yanchun Pan1, Takuji Daito1, Yo Sasaki2, Yong Hee Chung1, Xiaoyun Xing2, Santhi Pondugula3, S. Joshua Swamidass4, Ting Wang2, Albert H. Kim1,5,6,7 & Hiroko Yano1,2,5,7

Although epigenetic abnormalities have been described in Huntington’s disease (HD), the causal epigenetic mechanisms driving neurodegeneration in HD cortex and striatum remain undefined. Using an epigenetic pathway-targeted drug screen, we report that inhibitors of DNA methyltransferases (DNMTs), decitabine and FdCyd, block mutant huntingtin (Htt)-induced toxicity in primary cortical and striatal neurons. In addition, knockdown of DNMT3A or DNMT1 protected neurons against mutant Htt-induced toxicity, together demonstrating a requirement for DNMTs in mutant Htt-triggered neuronal death and suggesting a neurodegenerative mechanism based on DNA methylation-mediated transcriptional repression. Inhibition of DNMTs in HD model primary cortical or striatal neurons restored the expression of several key genes, including Bdnf, an important neurotrophic factor implicated in HD. Accordingly, the Bdnf promoter exhibited aberrant cytosine methylation in mutant Htt-expressing cortical neurons. In vivo, pharmacological inhibition of DNMTs in HD mouse brains restored the mRNA levels of key striatal genes known to be downregulated in HD. Thus, disturbances in DNA methylation play a critical role in mutant Htt-induced neuronal dysfunction and death, raising the possibility that epigenetic strategies targeting abnormal DNA methylation may have therapeutic utility in HD.

Huntington’s disease (HD) is a progressive and invariably fatal, autosomal-dominant neurodegenerative disease characterized by progressive loss of selective neurons in the striatum and cortex, leading to movement, cognitive, and psychiatric disorders1,2. Although HD is known to be caused by an abnormal expansion of polyglutamine repeats in the huntingtin (Htt) protein3, how the toxic mutant protein drives neuronal dysfunction and death remains poorly understood, and currently, no curative treatment exists for this disease.

Transcriptional dysregulation is an early abnormality in the course of HD progression and has been suggested to represent an underlying pathogenic mechanism for this disease4–6. Previous RNA profiling studies using human and mouse HD brains found a number of critical genes are dysregulated, including downregulation of brain-derived neurotrophic factor (Bdnf), dopamine D2 (Drd2) receptor, and protein phosphatase 1 regulatory inhibitor subunit 1B (Ppp1r1b or Darpp-32)7–12. BDNF is a major neurotrophic factor implicated in HD pathogenesis and plays important roles in a wide range of neuronal functions, from survival and differentiation to synaptic transmission and learning and memory in the adult brain13,14. The reduction of BDNF-mediated trophic support from cortical neurons to the medium spiny neurons in HD striatum has been suggested to be a major contributor to the striatal neurodegeneration observed in HD14–17.

Epigenetic mechanisms, including covalent chemical modifications of DNA and histones, play an important role in defining gene expression through modulation of chromatin structure and function18–20. Recent genome-wide studies using cell and animal models of HD have identified changes in the patterns of several epigenetic modifications, including DNA methylation and histone methylation, ubiquitination, and acetylation, suggesting that epigenetic dysregulation may be responsible for altered gene transcription observed in HD12,21–24.

1Department of Neurological Surgery, Washington University School of Medicine, St. Louis, MO 63110, USA.
2Department of Genetics, Washington University School of Medicine, St. Louis, MO 63110, USA.
3Department of Pediatrics, Washington University School of Medicine, St. Louis, MO 63110, USA.
4Department of Immunology and Pathology, Washington University School of Medicine, St. Louis, MO 63110, USA.
5Department of Neurology, Washington University School of Medicine, St. Louis, MO 63110, USA.
6Department of Developmental Biology, Washington University School of Medicine, St. Louis, MO 63110, USA.
7Hope Center for Neurological Disorders, Washington University School of Medicine, St. Louis, MO 63110, USA. Correspondence and requests for materials should be addressed to H.Y. (email: yanoh@wudosis.wustl.edu)
However, it is currently not known which epigenetic alterations in HD play causal roles in death of striatal and cortical neurons as well as disease progression.

Methylation of the fifth carbon position of cytosine (5-methylcytosine, 5mC) in CpG dinucleotide is an epigenetic modification that affects multiple biological processes, including embryonic development, X chromosome inactivation, genomic imprinting, and diseases such as cancers20,25,26. DNA methylation in the gene promoter generally leads to genomic silencing. Recently, the role of this epigenetic modification in the nervous system has begun to attract considerable attention. Emerging evidence suggests that DNA methylation-mediated transcriptional regulation is critically involved in normal brain function as well as brain diseases, including learning and memory, synaptic transmission, and psychiatric disorders27–34. But little is known about the involvement of DNA methylation in the pathogenesis of neurodegenerative diseases, including HD. DNA methylation is catalyzed by members of the DNA methyltransferase (DNMT) family of enzymes, including DNMT1, DNMT3A, and DNMT3B. Whereas DNMT1 is essential for the maintenance of DNA methylation patterns in the newly synthesized strand during DNA replication, DNMT3A and DNMT3B are required for de novo DNA methyla-
tion20,36. Interestingly, DNMTs are also highly expressed in non-dividing neurons; DNMT1 and DNMT3A are the major DNMTs expressed in postmitotic neurons of the brain35–37. How these DNMTs modulate transcription of neuronal genes and contribute to the function and survival of postmitotic neurons remains poorly understood in the context of healthy and diseased brains. Given the altered patterns of several epigenetic modifications in mouse and cell models of HD32–34, identification of the key epigenetic pathway(s) that dominantly drive mutant Htt-induced transcriptional alterations and cytotoxicity in neurons remains an important unanswered question.

Here we demonstrate, through an unbiased epigenetic drug library screening using a primary neuron model of HD, that pharmacological inhibitors of DNMTs effectively protect neurons from mutant Htt-induced toxicity. We also demonstrate that knockdown of DNMT1 or DNMT3A inhibits mutant Htt-induced neuronal toxicity in culture. Inhibition of DNMTs in primary neuron models of HD restored the expression of critical genes known to be downregulated in HD, including Hdh. Consistent with this finding, increased levels of DNA methylation were found in the promoter of Hdh in mutant Htt-expressing neurons. Furthermore, pharmacological inhibition of DNMTs in HD transgenic mice upregulated the transcription of key striatal genes in vivo. These findings demonstrate that DNA methylation plays a functionally important role in mutant Htt-induced transcriptional dysregulation and neurotoxicity, raising the possibility that manipulations of the DNA methylation pathway might represent an attractive new therapeutic strategy to attenuate HD neurodegeneration.

Results
DNA demethylating agents protect neurons from mutant Htt-induced cytotoxicity. To identify the critical epigenetic pathways that contribute to the death of mutant Htt-expressing neurons, we performed an epigenetic drug screen using a library composed of 84 epigenetic compounds with known targets, including writers, erasers, and readers of the epigenetic code (Fig. 1A and Table S1). As epigenetic gene regulation is a cell-type specific mechanism, we used a physiologically relevant, postmitotic cortical neuron culture system for this screen. In this system, lentivirus-mediated expression of the exon 1-encoded N-terminal fragment of mutant Htt (Htt-72Q with a 72 glutamine repeat), but not wild-type (WT) Htt (Htt-25Q), induces neurotoxicity39. The N-terminal short fragments of mutant Htt, which can be generated in cells by proteolytic cleavage of the full-length Htt or alternative splicing, is known to be more cytotoxic than the full-length protein and is expressed in HD patients39–44. In the drug screen, the viability of Htt-expressing cortical neurons was determined by the resazurin (Alamar Blue) assay, a quantitative measurement of mitochondrial metabolic activity, which correlates with cell viability. Following validation assays of possible screen "hits" using the MTS assay, we identified the cytosine nucleoside-analog DNA methyltransferase (DNMT) inhibitor decitabine, as the most effective mutant Htt-induced neurotoxicity screen (Fig. 1B and S1A). Remarkably, decitabine exhibited nearly full neuroprotection at 0.2 μM in our HD system (Fig. 1B). Decitabine, also known as 5-aza-2′-deoxycytidine or DacogenTM (DAC), is a U.S. Food and Drug Administration (FDA)-approved drug and has been used clinically for the treatment of cancers, including myelodysplastic syndrome (MDS) and acute myeloid leukemia (AML)26,45, but not for neurodegenerative diseases. We further verified the protective effects of decitabine in mutant Htt-expressing primary cortical neurons using two additional assays: neurite degeneration by quantifying the loss of neurofilament (NF) immunofluorescence intensity, an early marker of neuronal toxicity (Fig. 1C,D), and cell death by scoring condensed or fragmented nuclei (Fig. 1E). Since decitabine is an epigenetic agent that affects gene expression, we confirmed that decitabine does not decrease the expression of Htt-72Q in our system compared to vehicle control by qRT-PCR (Fig. S2A). We then tested if treatment with decitabine affects the burden of mutant Htt aggregates—an HD pathological hallmark—in primary cortical neurons using an antibody that preferentially detects mutant Htt aggregates (Fig. S2B–D). By both immunofluorescence and immunoblot analyses, we found that decitabine could decrease the levels of mutant Htt aggregates in Htt-72Q-expressing neurons. Given that misfolded and aggregated Htt may interfere with several important biological functions in neurons, its reduction may contribute to neuroprotection by decitabine.

To verify the effect of DNMT inhibition against mutant Htt toxicity, we next performed similar experiments with three other well-characterized nucleoside-analog DNMT inhibitors, 5-fluoro-2′-deoxycytidine (FdCyd), 5-azacytidine (azacitidine, 5-AC, VidazaTM), and zebularine. The latter two drugs are ribonucleoside analogs, which target primarily RNA rather than DNA, and small fractions of these drugs can be converted to their deoxyribose form in cells, thereby leading to inhibition of DNA methylation46,47. 5-azacytidine, like decitabine, is a FDA-approved, potent anti-cancer drug that has been used for the treatment of MDS and AML. FdCyd was developed by the National Cancer Institute and is currently being investigated in ongoing clinical trials in solid tumors. Interestingly, the deoxyribonucleoside analog FdCyd, but not the ribonucleoside analogs, zebularine and 5-azacytidine, demonstrated neuroprotective effects against mutant Htt-induced toxicity in primary cortical neu-
rons in cell viability and neurite degeneration assays (Fig. 1E,G, and S3A–C), suggesting that the deoxyribonucleo-
side form of DNMT inhibitors, which act directly through DNA, exerts neuroprotective activity in HD neurons.
To test if DNMTs play a role in the HD striatal neurons, one of the most severely affected cell types in the disease, we established a cultured striatal neuron model, in which the N-terminal exon-1 fragment of mutant or WT Htt was expressed by lentiviral infection. Strikingly, treatment with decitabine or FdCyd also attenuated mutant Htt-induced striatal neurite degeneration (Fig. 1H,I). Similar to cortical neurons, 5-azacytidine did not inhibit mutant Htt-induced neurite degeneration in striatal neurons (Fig. S3D). Together, results utilizing two disease-relevant neuronal cell types suggest that DNMTs play an important role in mutant Htt-induced neurodegeneration.
Reduced DNMT3A or DNMT1 expression protects neurons from mutant Htt-induced toxicity.

Because inhibition of DNMTs by decitabine and FdCyd rescued neurons from mutant Htt-induced toxicity, we next determined if genetic inhibition of DNMTs by RNA interference (RNAi) attenuates neuronal death in the mutant Htt context. Among members of the DNMT family, postmitotic neurons in the brain are known to highly express DNMT3A and DNMT1. Knockdown of either DNMT3A or DNMT1 protein by lentiviral delivery of two distinct short hairpin RNAs (shRNAs) for each DNMT significantly increased the viability of mutant Htt-expressing neurons (Fig. 2A–D). Knockdown of DNMT3A or DNMT1 did not decrease the levels of the other DNMT’s mRNA or protein (Fig. 2A,B, and S4A,B), demonstrating specificity of the shRNAs used. These findings indicate that both DNMT3A and DNMT1 are required for mutant Htt-induced neuronal death. Taken
Figure 3. Inhibition of DNMTs restores the expression of Bdnf exon IV and VI transcripts in primary cortical neurons. (A) DIV 5 cortical neurons were infected with Htt lentivirus. RNA was harvested 5 days later and subjected to qRT-PCR for total Bdnf (coding exon IX) using β-actin and 18S rRNA as reference genes. Htt-72Q decreased the expression of total Bdnf transcripts (Mann-Whitney U test, *P = 0.008 vs. Htt-25Q, n = 5).

(B) Cortical neurons transduced as in (A) were cultured with recombinant BDNF (50 ng/ml) and subjected to MTS assay at DIV 14. BDNF increased the viability of Htt-72Q-expressing neurons (ANOVA, *P < 0.0001 vs. Htt-72Q with vehicle, n = 9–15).

(C) (Top) Schematic of the mouse Bdnf locus. White boxes, non-coding exons; gray box, coding exon. (Bottom) qRT-PCR was performed as in (A) using exon-specific Bdnf primers. Htt-72Q decreased the expression of exon IV and VI transcripts (Mann-Whitney U test, *P = 0.008 vs. Htt-25Q, n = 5).

(D–F) Cortical neurons transduced with Htt lentivirus were treated with indicated DNMT inhibitor or vehicle and processed as in (C). qRT-PCR was performed using β-actin and Hprt as reference genes. Both decitabine and FdCyd increased the expression of Bdnf exon IV, VI, and IX transcripts in Htt-72Q-expressing neurons (ANOVA,
progressive motor deficits and late-onset selective neuropathology in the cortex and striatum. Inhibition of Bdnf increased by DNMT inhibition (Fig. 3D,E,G–I), we next test the hypothesis that mutant Htt stimulates DNA CaRE2, and CRE (schematic in Fig. 4A) and is activated in response to various extracellular stimuli in vivo.

Promoter IV harbors several transcription factor binding sites, including calcium responsive elements, CaRE1, Bdnf DNMTs by decitabine in BACHD mouse cortical neurons increased its neuroprotective effects in the context of mutant Htt in part through the upregulation of Bdnf.

Inhibition of DNMTs restores Bdnf gene expression in mutant Htt-expressing cortical neurons. BDNF is a major neurotrophic factor involved in fundamental brain processes, including neuronal survival, synaptic plasticity, and learning and memory. Bdnf mRNA and protein levels were found to be decreased in the brains of human HD patients and mouse models, which is thought to contribute to HD pathology. Consistent with these observations, Bdnf expression was reduced by mutant Htt expression in primary cortical neurons (Fig. 3A). Addition of recombinant BDNF protein in the culture medium was sufficient to rescue cortical neurons from mutant Htt-induced toxicity (Fig. 3B), suggesting an important role for BDNF in the survival of mutant Htt-expressing neurons. Using Bdnf as a model gene, we next focused on determining if Bdnf transcriptional repression could be rescued by manipulating DNA methylation in mutant Htt-expressing neurons. As the Bdnf gene has a complex structure with multiple noncoding exons and a common protein coding exon, we first examined the differential expression of major exon-specific Bdnf transcripts in primary cortical neurons. Each noncoding exon has an independent promoter, and the expression of the exon-specific transcript is differentially regulated in response to diverse extracellular stimuli and signaling events (schematic in Fig. 3C). Mutant Htt-expressing cortical neurons exhibited decreased expression of Bdnf mRNA at a time before neurons begin to die, specifically Bdnf exon IV- and VI-containing transcripts, compared to control neurons expressing WT Htt-25Q or the empty vector (Fig. 3C, data not shown), consistent with a previous observation. The reduction of these transcripts is clinically relevant since they have also been found to be decreased in HD model mouse brains and human HD postmortem brain.

Next, to test the hypothesis that abnormal DNA methylation contributes to the downregulation of Bdnf mRNA, we examined if pharmacological inhibition of DNMTs could rescue the expression of Bdnf exon IV and VI-containing mRNAs in mutant Htt-expressing cortical neurons by qRT-PCR analysis (Fig. 3D,E). Intriguingly, both decitabine and FdCyd restored the levels of Bdnf exon IV and VI transcripts at doses effective for neuroprotection (Fig. 3D,E). These DNMT inhibitors also increased the levels of the common coding exon IX transcript (total Bdnf mRNA) in mutant Htt-expressing cortical neurons (Fig. 3F). Consistent with the effects of DNMT inhibitors on Bdnf transcription, knockdown of DNMT3A or DNMT1 in mutant Htt-expressing cortical neurons using two shRNAs targeting each DNMT reversed the mutant Htt-triggered decrease in Bdnf exon IV and VI mRNAs (Fig. 3G,H). These results suggest that both DNMTs contribute to downregulation of Bdnf mRNA in HD neurons.

To verify these findings using an alternative HD model system, we next determined if decitabine could upregulate Bdnf mRNA expression in primary cortical neurons derived from bacterial artificial chromosome (BAC)-mediated HD transgenic (BACHD) mice, which express full-length mutant Htt. BACHD mouse cortical neurons exhibit progressive motor deficits and late-onset selective neuropathology in the cortex and striatum. Inhibition of DNMTs by decitabine in BACHD mouse cortical neurons increased Bdnf exon IV- and VI-containing transcripts as well as total Bdnf (exon IX) mRNAs by qRT-PCR (Fig. 3I), supporting the findings obtained using neurons expressing the N-terminal fragment of mutant Htt (Fig. 3D). Collectively, these results suggest that DNMT inhibition exhibits neuroprotective effects in the context of mutant Htt in part through the upregulation of Bdnf.

Mutant Htt increased the levels of DNA methylation in the Bdnf exon IV regulatory region in primary cortical neurons. Because Bdnf exon IV and VI transcripts in mutant Htt-expressing cortical neurons are increased by DNMT inhibition (Fig. 3D,E,G–I), we next test the hypothesis that mutant Htt stimulates DNA methylation in the promoter regions of these two exons, leading to repression of these transcripts. The Bdnf promoter IV harbors several transcription factor binding sites, including calcium responsive elements, CaRE1, CaRE2, and CRE (schematic in Fig. 4A) and is activated in response to various extracellular stimuli in vivo. The levels of DNA methylation in the Bdnf exon IV regulatory region that contains 13 CpG sites (base pairs −148 to +65 relative to the transcriptional start site (TSS)) was assessed by bisulfite conversion followed by DNA sequencing. A widely used technique to measure levels of cytosine methylation on specific genomic regions with single CpG resolution. Among the 13 CpG sites examined, methylation of eight CpGs between base pairs −87 to +65 was increased (1.5- to 3.2-fold) by mutant Htt expression (Fig. 4B and S5A). Three CpG sites located between base pairs −148 and −109, including the previously reported methyl CpG binding protein 2 (MeCP2) binding site at the position −148, exhibited robust methylation with no significant difference between WT and mutant Htt-expressing neurons (Fig. 4B). The bisulfite sequencing results revealed that mutant Htt-expressing neurons exhibit an overall increase in the levels of cytosine methylation, compared to WT Htt-expressing neurons, in the regulatory region of the Bdnf exon IV surrounding the TSS (Fig. 4B and S5A). The increased levels of cytosine methylation (5-mC) in the region was independently confirmed by methylated DNA immunoprecipitation assay (MeDIP), which uses a specific antibody against 5-mC (Fig. 4C). These data suggest that mutant Htt
Figure 4. Mutant Htt increases the levels of DNA methylation at Bdnf exon IV regulatory region in primary cortical neurons. (A) Schematic of the mouse Bdnf exon IV regulatory region near the TSS. The positions of CpG sites are indicated relative to the TSS. (B) DIV 5 primary cortical neurons were infected with lentivirus expressing Htt-25Q or Htt-72Q exon 1 fragment; 5 days later, genomic DNA was purified and subjected to bisulfite sequencing analysis. The data show percentage of cytosine residues that were methylated in Htt25Q- and Htt-72Q-expressing neurons. Increased DNA methylation was found in mutant Htt-expressing
expression downregulates Bdnf exon IV transcript via increased DNA methylation of the promoter. In contrast, similar bisulphite sequencing analysis for the regulatory region of Bdnf exon VI containing 17 CpG sites displayed little if any DNA methylation in both WT Htt-25Q and mutant Htt-72Q-expressing cortical neurons (Fig. SSB), indicating that Bdnf promoter VI is not a direct target of DNA methylation but is indirectly suppressed by events initiated by aberrant DNA methylation in mutant Htt-expressing neurons.

DNA methylation-mediated gene repression is generally associated with a closed chromatin structure, which is induced by cooperation with altered histone modifications[57]. By chromatin immunoprecipitation (ChIP) analysis we found that the mutant Htt-triggered increase in DNA methylation is associated with decreased trimethylation at lysine 4 of histone H3 (H3K4me3), a transcriptionally active histone mark, in the promoter region of Bdnf exon IV in mutant Htt-expressing primary cortical neurons compared to WT Htt-expressing neurons (Fig. 4D). Together, these results illustrate that mutant Htt-induced increased DNA methylation are associated with loss of active open chromatin in this region, consistent with mutant Htt-induced transcriptional repression of Bdnf exon IV.

Next, to further support the hypothesis that decreased Bdnf exon IV transcription by mutant Htt is the consequence of changes in DNA methylation at this locus, we examined whether inhibition of DNMTs in mutant Htt-expressing primary cortical neurons decreases DNA methylation in the regulatory region of Bdnf exon IV by MeDIP-qPCR. We found that inhibition of DNMTs by decitabine or FdCyd reversed the increase in DNA methylation triggered by mutant Htt (Fig. 4C,E,F). Consistently, knockdown of either DNMT3A or DNMT1 decreased the levels of DNA methylation in this region (Fig. 4G,H). Together, our results suggest that altered DNA methylation drives the repression of Bdnf transcription in HD neurons and demonstrate that two distinct DNMTs in neurons are both required for the mutant Htt-induced increase in DNA methylation in the Bdnf regulatory region.

Decitabine reactivates expression of key striatal genes in a primary striatal neuron model of HD. Given the neuroprotective effect of decitabine in mutant Htt-expressing striatal neurons (Fig. 1H), we next tested if DNMT inhibition with decitabine restores the expression of other genes that are known to be downregulated in HD. Consistent with previous gene expression studies using human and mouse HD striatum[7,8,12,58,59], qRT-PCR analyses demonstrated that mutant Htt expression in primary striatal neurons triggers robust changes in gene expression, including downregulation of dopamine receptor D2 (Drd2), protein phosphatase 1, regulatory (inhibitor) subunit 1B (Ppp1r1b, also known as Darpp-32), preproenkephalin (Penk), Purkinje cell protein 4 (Pcp4), and RASD family, member 2 (Rasd2, also known as Rhes) (Fig. 5A). Thus our culture system faithfully reproduces key gene expression changes observed in HD in vivo. These transcriptional changes were detected before mutant Htt neurons exhibit significant neurite degeneration (data not shown), suggesting that mutant Htt-induced transcriptional changes contribute to striatal neurodegeneration. Inhibition of DNMTs by decitabine dramatically increased the expression of these downregulated transcripts (Fig. 5A). This reactivation of gene expression was specific because the mRNA levels of lysine (K)-specific demethylase 8 (Kdm8) were unchanged by decitabine treatment. Together these results demonstrate that the inhibition of DNA methylation can restore gene expression, which is deficient in HD neurons, suggesting that abnormal DNA methylation plays a critical role in transcriptional dysregulation in HD striatal and cortical neurons.

Pharmacological inhibition of DNMTs in HD mouse brains upregulates the expression of key striatal genes in vivo. We next determined if DNMT inhibition could restore the expression of genes downregulated in HD in vivo using R6/2 HD mouse, a well-characterized transgenic mouse model expressing an N-terminal mutant Htt fragment[60]. This mouse model exhibits robust phenotypes with early disease onset and short life span and recapitulates the altered expression of a number of genes observed in HD patients, including Drd2 and Ppp1r1b in the striatum early in the course of disease progression[7,12,60,61]. Although decitabine has been reported to cross the blood-brain barrier[62,63], the cytosine nucleoside analog DNMT inhibitors, including decitabine and FdCyd, are known to be degraded rapidly by cytidine deaminase in the liver (in vivo half life of decitabine <20 min)[64], indicating that systemic administration may not be an effective strategy for drug delivery to the brain. We therefore chose intracerebroventricular (icv) administration using an Alzet osmotic pump, which provides continuous infusion of drug at a consistent rate from a subcutaneous pump (Fig. 5B). Although FdCyd when tested structurally similar to decitabine, we found that, whereas decitabine lost its in vitro neuroprotective activity after 11 days of pre-incubation at 37°C in saline, FdCyd fully maintained its neuroprotective activity even after 45
days of pre-incubation (Figure S6A,B), suggesting that FdCyd is chemically more stable than decitabine at 37 °C in vitro and is better suited for drug delivery with osmotic pumps. The instability of decitabine in vitro has been reported previously.64,65. We therefore used FdCyd to determine the effect of DNMT inhibition on gene expression in R6/2 brain. One week after the implantation of Alzet osmotic pumps filled with FdCyd or saline in R6/2 mice or WT littermates, RNA was prepared from the striatum (Fig. 5B). Striatal expression of several key mRNAs, Drd2, Ppp1r1b, Rasd2, Adora2a, and Penk mRNAs, was found to be downregulated in HD mouse striatum compared to
control animals (Fig. 5B), consistent with previous reports in human and mouse HD striata as well as in our HD model striatal neurons (Fig. 5A)7,8,12. Infusion of FdCyd in R6/2 brains significantly upregulated the expression of striatal Drd2, Ppp1r1b, Rasd2, and Adora2a mRNAs and also showed a trend towards increasing Penk mRNA (Fig. 5B), indicating that pharmacological inhibition of DNMTs can correct transcriptional deficiencies in HD mouse brain. Together, these results suggest that DNA methylation plays an important role in transcriptional alterations in HD and potentially, neuronal dysfunction and death in vivo (See a model in Fig. 6).

Discussion

In this study, we have demonstrated that pharmacological or genetic inhibition of DNMTs substantially attenuates mutant Htt-induced transcriptional dysregulation and neurotoxicity in primary cortical and striatal neurons. We have also provided evidence that aberrant promoter methylation contributes to a reduction in Bdnf expression in mutant Htt-expressing cortical neurons. Given the neuroprotective effects of exogenous BDNF in HD model cortical neurons, blockade of DNMTs may protect neurons from mutant Htt-induced death in part through upregulation of Bdnf gene expression. Remarkably, in vivo experiments demonstrated that treatment of HD mice with DNMT inhibitor FdCyd could reverse the transcriptional repression of key striatal genes in HD mouse brain. Together, we provide evidence that DNA methylation in HD is a critical epigenetic mechanism, which underlies mutant Htt-induced transcriptional alterations and neurodegeneration, raising the possibility that the DNA methylation pathway might represent a new therapeutic target for HD.

Previous studies have shown that a large number of genes are dysregulated in the brains of HD patients and various mouse models5–12. Recent genome-wide analysis as well as candidate gene approaches using cell and mouse models of HD have found altered patterns of several epigenetic modifications66,67, including acetylation, ubiquitination, and methylation (H3K4me3 and H3K9me3) of histones12,21,23,24,68–70 as well as methylation (5-mC) and hydroxymethylation (5-hmC) of DNA22,71. Therefore, abnormal chromatin state may be a critical driver for neurodegeneration in HD. Although the causal role of these epigenetic modifications in vulnerable neurons in HD remains unknown, our unbiased drug library screen with 84 chemical compounds, which target known epigenetic pathways, suggests that DNA methylation-mediated gene silencing plays a dominant role in triggering neuronal death.

In primary cortical neuron models, we found that mutant Htt induces increased DNA methylation in the regulatory region of Bdnf exon IV, which is associated with transcriptional repression and a reduction in the transcriptionally active H3K4me3 mark. Additionally, inhibition of DNMTs by pharmacological inhibitors or RNAi could rescue the expression of Bdnf exon IV mRNA in mutant Htt-expressing primary cortical neurons (Fig. 3D,E,G–I), suggesting that the mutant Htt-triggered increase in DNA methylation in this region directly causes transcriptional repression. Multiple previous studies have suggested that DNA methylation and histone modifications exhibit crosstalk and cooperate in the regulation of gene expression72,73. Understanding the epigenetic hierarchy downstream of mutant Htt in neurons in relation to RNA expression represents an important future direction.

How mutant Htt promotes DNA methylation at specific gene loci at the molecular level remains a significant open question. Possible mechanisms include: 1) mutant Htt expression in neurons increases the levels of DNMT expression, 2) mutant Htt enhances the activity of DNMTs, 3) mutant Htt facilitates the recruitment of the DNA methylation machinery to specific genomic regions, and/or 4) mutant Htt increases 5-mC levels by decreasing DNA demethylation activity in neurons. The first mechanism, however, is unlikely since we have found that
mutant Htt does not significantly increase the mRNA or protein levels of DNMT1 or DNMT3A in primary cortical neurons (Y.P. and H.Y., unpublished data). The second and third mechanisms are reasonable possibilities and may be caused by aberrant protein–protein interactions and/or abnormal posttranslational modifications of DNMTs downstream of mutant Htt. Regarding the fourth possible mechanism, the recent discovery of the ten-eleven translocation (TET) family of enzymes that promote DNA demethylation by converting 5-mC to 5-hmC and the abundance of TET proteins and 5-hmC in brain has highlighted DNA methylation as a dynamically regulated process important for brain function. Therefore, whether mutant Htt increases 5-mC levels on repressed genes by inhibiting the DNA demethylation pathway in HD is an interesting question.

BDNF deficiency in the brains of HD patients and mouse models has been suggested to play a crucial role in the development of the disease. However, the mechanisms underlying downregulation of exon-specific Bdnf transcripts, in particular, exon IV and VI in HD neurons remains largely unknown. We focused on Bdnf as a model gene to test the hypothesis that mutant Htt represses neuronal gene expression through promoter hypermethylation. Our results show that mutant Htt expression increases cytosine methylation in the regulatory region of Bdnf exon IV and that inhibition of DNMTs decreases the methylation and reactivates exon IV transcription, supporting the idea that increased DNA methylation plays a causal role in repression of Bdnf transcription in HD. In contrast, the regulatory region of Bdnf exon VI, appears not to be directly regulated by DNA methylation, suggesting instead that indirect mechanisms are initiated by aberrant DNA methylation in the control of the Bdnf exon VI expression. Several approaches to increase BDNF-mediated trophic support in HD brain are being performed in ongoing clinical trials and preclinical studies. Our results suggest that manipulation of DNA methylation may offer a new therapeutic approach to increase neuronal BDNF expression in HD brain.

The reduction of either DNMT1 or DNMT3A by RNAi is sufficient to block transcriptional changes and neuronal death induced by mutant Htt (Fig. 2A–D and S5LH), suggesting that both DNMTs are required to exert mutant Htt-mediated toxicity. Although in dividing cells, the roles of DNMT1 and DNMT3A as maintenance and de novo DNMTs, respectively, are known, the specific roles of the two DNMTs in postmitotic neurons remain undefined, necessitating exploration in future studies. An intriguing previous study demonstrated that double conditional knockout mice lacking DNMT3A and DNMT1 in postnatal forebrain excitatory neurons, but not single ablation of either DNMT3A or DNMT1, exhibited a deficit in synaptic function and learning and memory, suggesting redundant or overlapping roles. Targeting either DNMT1 or DNMT3A in adult neurons therefore may attenuate mutant Htt-induced neurotoxicity with minimal side effects in regard to normal synaptic functions in the brain. It has been demonstrated that the nucleoside analog DNMT inhibitors, such as decitabine and FdCyd, must first be incorporated into DNA to exert their DNMT inhibitory activity. In dividing cells, drug incorporation occurs during DNA synthesis. The mechanism of action of these DNMT inhibitors in non-dividing postmitotic neurons, however, still remains unclear, although it is possible that the base excision repair pathway contributes to the incorporation of nucleoside-analog DNMT inhibitors. Improved molecular understanding of the action of decitabine and FdCyd may identify potential "hot spots" of incorporation in the neuronal genome, providing relevant information regarding specific gene targets undergoing active methylation in the HD epigenome. Integrating genome-wide DNA methylation and transcriptional changes associated with DNMT inhibition in future studies will identify the key gene targets of DNMT inhibition-induced demethylation in HD neurons.

Finally, the findings from the current study immediately suggest that inhibition of DNMTs might ameliorate HD phenotypes in vivo, which will be the subject of important future experiments. Improved understanding of the epigenetic gene regulation in HD neurons will provide important foundational knowledge for the development of therapeutic strategies targeting DNA methylation abnormalities in HD.

**Methods**

**Antibodies and reagents.** Mouse monoclonal anti-neurofilament (NF) (165kDa) (clone 2H3, Developmental Studies Hybridoma Bank) was used for immunofluorescence. Mouse monoclonal anti-β-actin (sc-47778, Santa Cruz Biotechnology), rabbit monoclonal anti-DNMT1 (D63A6, Cell Signaling Technology, Inc.), and rabbit polyclonal anti-DNMT3A (sc-20703, Santa Cruz Biotechnology) antibodies were used for immunoblotting. Mouse monoclonal anti-Htt (EM48) antibody was used for immunofluorescence and immunoblotting. Decitabine was purchased from Cayman Chemical (11166) and LC laboratories (D-3899). 5’-fluoro-2’-deoxycytidine (FdCyd) was purchased from Sigma (F5307) and Santa Cruz Biotechnology (sc-252267). These drugs were confirmed to exhibit similar effects regardless of the source.

**Plasmids.** Lentiviral expression plasmids containing Htt exon1-25Q (Htt-25Q) and Htt exon1-72Q (Htt-72Q) constructs under the control of the mouse PGK (Pkg1) promoter (mpPGK-Htt exon1-25Q and mpPGK-Htt exon1-72Q) were kindly provided by D. Krainc (Northwestern University, Chicago, IL). Lentivirus-based Dnmt3a RNAi and Dnmt1 RNAi constructs (pLKO.1-puro), developed at the Broad Institute of MIT and Harvard, were obtained from Sigma-Aldrich. The oligo sequences in the shRNA vectors targeted Dnmt3a and Dnmt1 are as follows: pLKO.1-Dnmt3a#1 (TRCN000039034): CCGGCCAGATGTTCCTTCATTTCAACACTCTGGTTTG; pLKO.1-Dnmt3a#2 (TRCN000039035): CCGGCCAGACCAACATCGAATCCATCTCGAGATGGATTCGATGTTGGTCTGCTTTTTG; pLKO.1-Dnmt1#1 (TRCN000019081): GTACCGGATCTATGGAGGTTTGATATTACACTCGAGTATTAATACCACCTTCCATAGATTTTTTTG; pLKO.1-Dnmt1#2 (TRCN0000225698): CCGGTATATGAAGACCTGATCAATACTCGAGTATTGATCAGGTCTTCATATA

**Primary neuron cultures, lentiviral transduction drug treatments.** Mouse primary cortical and striatal neurons from embryonic day (E) 15.5 Swiss Webster mouse fetuses (Taconic) were first plated in the minimal essential medium (MEM) containing 10% FBS, 0.37% glucose, 1 mM sodium pyruvate, 2 mM glutamine,
20 U/ml penicillin and 20 μg/ml streptomycin, for 3 h and then maintained in serum-free Neurobasal medium (Life Technologies) containing NeuroCult™ SM1 neuronal supplement (STEMCELL Technologies), 0.5 mM glutamine and 25 μM glutamate for the first 3 d in a humidified incubator (37 °C in 5% CO2). Half of the medium was replaced with Neurobasal medium with SM1 and 0.5 mM glutamine every 3 days. Primary cortical neurons plated on 96-well flat clear bottom black plates (Corning #3904) were infected with Htt exon1 expression lentivirus (Htt-25Q or Htt-72Q) or control empty vector lentivirus at 5 days in vitro (DIV 5). Primary striatal neurons plated on 96-well plates at 1 × 104 cells/well were infected with Htt exon1 expression lentivirus at DIV 4. Viral copy number was adjusted for transduction of neurons on the basis of titers measured using the Lenti-X qRT-PCR titration kit (Clontech), and equal numbers of viral particles of Htt-25Q and Htt-72Q expressing lentiviruses were used for transduction. For the experiments to test effects of DNMT inhibitors, neurons were treated with inhibitors six hours after Htt lentiviral infection. One half of the media was changed every 3 days with media containing new drug. In knockdown experiments in Htt-expressing neurons, primary cortical neurons were cotransduced with Htt-expressing lentivirus and Dnmt shRNA or control shRNA lentivirus at DIV 5. pLKO.1-TRC1-luciferase (Luci) and pLKO.1-TRC2-LacZ were used as control for RNAi with pLKO.1-TRC1-Dnmt3a and pLKO.1-TRC2-Dnmt1, respectively. Lentiviral particles were prepared by transfecting 293 T.E cells with the lentiviral plasmid of interest along with packaging plasmid psPAX2 and envelope plasmid pCMV-VSVG as described previously69. Four days after transfection, viruses in the conditioned media were collected and purified using Lenti-X Concentrator (Clontech). Primary cortical neurons from BACHD mouse embryos (E15.5) were individually plated into separate wells and treated at DIV 4 with decitabine or vehicle for 3.5 days.

**Measurements of cell viability/cytotoxicity in primary neurons.** Primary cortical neurons grown in a 96-well plate were transduced with Htt-expressing lentiviruses at DIV 5 and assessed for mitochondrial metabolic activity at 9 days post-infection (DIV 14) using 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulphonylphenyl)-2H-tetrazolium (MTS) (Promega) per manufacturer’s instructions. MTS-reducing activity was normalized for each condition to Htt-25Q lentiviruses treated with vehicle or cotransduced with control RNAi lentivirus (=1). Experiments were performed in 3 or more wells per experiment in three to five independent experiments.

For the measurement of neurofilament (NF) immunofluorescence intensity, cortical and striatal neurons cultured in a 96-well plate were fixed in 4% paraformaldehyde (PFA) in PBS for 20 min and nine and seven days after Htt lentiviral infection, respectively, permeabilized with 0.1% Triton X-100 in PBS for 15 min at room temperature, and subjected to indirect immunofluorescence with anti-NF (2H3) primary antibody and Alexa Fluor 568-conjugated goat anti-mouse IgG secondary antibody (Life Technologies). Images of Alexa Fluor 568-labeled were captured (nine random fields per well) using an Operetta high-content imaging system (PerkinElmer) with a 20 × objective lens. Following image background subtraction, the NF immunofluorescence intensity was quantified using an ImageJ-based macro. Image capture and quantification of Htt (EM48) immunofluorescence intensity were performed as described for those of NF. In this quantification analysis, we confirmed that the number of cells in a cultured well are similar among mutant Htt-expressing neurons with or without DNMT inhibitor treatment, by counting the number of nuclei (Hoechst 33342) in the images used for quantification: Htt-72Q neurons treated with vehicle (299 ± 7.8 cells), decitabine (299 ± 4.6 cells), n = 18 wells from 6 independent experiments, and therefore the data reflect EM48 intensity per cell. For the quantification of cell death, primary cortical neurons grown in a 96-well plate were infected with Htt lentivirus at DIV 5 and fixed nine days after infection as described above. Cell nuclei were labeled with Hoechst 33342 (Life Technologies), and neurons were assessed in a blinded fashion for cell death by scoring condensed or fragmented nuclei. Experiments were performed in 4 to 6 wells per experiment in three independent experiments. About 300 nuclei from three random fields in a well were counted.

**Drug library screen.** Epigenetic drug screen was performed using a primary cortical neuron model of HD with a drug library composed of 84 compounds (Table S1), among which 80 drugs are purchased from Cayman Chemical (Epigenetic Screening Library Item No 11076) and four drugs, SGC0946, EPZ5676, EPZ6438, and GSK126, were obtained from Xcessbio Biosciences Inc. Mouse primary cortical neurons were plated on 96-well flat clear bottom black plates (Corning #3904). WT or mutant Htt exon1 fragment (Htt-25Q or Htt-72Q)-expressing lentiviruses are infected at DIV 5 as described above. The 84 compounds were added to the media at DIV 6 one day after Htt lentiviral infection with three different doses (0.02, 0.2, 2 μM) for each compound at triplicates. DMSO was used as control. One half of the media containing compounds or DMSO were changed every three days to test the compound’s activity. At DIV 14, the medium of neurons was determined by resazurin (Alamar blue) assay, a quantitative measurement of mitochondrial metabolic activity. The screen was fully automated and performed in the High-Throughput Screening Center in Washington University School of Medicine. Any possible plate effects were determined using control plates treated with DMSO and used for normalization. Screen hits were validated by MTS assay.

**HD transgenic mice and drug administration.** R6/2 mice, which carry the promoter sequence and exon 1 of a mutant human HTT gene, were obtained from JAX (Stock No: 002810) (Bar Harbor, ME), and a colony was maintained by breeding R6/2 males with B6CBAF1 females (JAX). PCR genotyping was performed using a primer set (CGGCTGAGGCCAGCCGGCTGT and GCAGCAGCAGCAACACCGCCACCGCC) as described previously69. To maintain mice carrying the same number of CAG repeats, a second PCR analysis was also conducted using a primer set amplifying across the CAG repeats (ATGAAAGCCTTCGAGTC CCTCAAGTCTTC and GGCGGCTGAGGAAGCTGAGGA). BACHD mice on the C57BL/6J background, which were generated by the laboratory of X. William Yang (University of California, Los Angeles)73,75, were obtained from the CHDI Foundation. All live vertebrate experiments were performed in compliance with the US National Institutes
μ5-mC-containing DNA fragments were enriched from one

References

Statistical analysis. Statistical differences were tested using XLSTAT and GraphPad Prism 6.0. Two-tailed unpaired Student t test for two group comparisons or one-way ANOVA with post-hoc tests, the Fisher's least significant difference (LSD) for comparison among three groups or the Bonferroni analysis for comparison among three or more than three groups. The Mann-Whitney U test was used for nonparametric test for comparing two groups. The data presented are from at least three independent experiments.

Bisulfite conversion and sequencing. Genomic DNA was extracted from cells using QIAamp DNA Mini Kit (QIAGEN) and subjected to bisulfite conversion using EZ DNA Methylation-Lightning™ Kit (Zymo Research) according to the manufacturer's instructions. Gene regulatory regions for Bdnf exons IV and VI were PCR amplified using ZymoTaq™ DNA Polymerase (Zymo Research) from the bisulfite-converted DNA templates. The PCR fragments were subcloned into the pCR2.1-TOPO vector using TOPO TA cloning kit (Life Technologies) and sequenced with M13 primer (Genewiz). The primers used for PCR amplification of the bisulfite-converted genomic DNA are listed in Table S2B.

Methylated DNA immunoprecipitation (MeDIP). Genomic DNA was isolated from primary cortical neurons using QIAamp DNA Mini Kit (QIAGEN) and fragmented by sonication using Bioruptor (Diagenode). 5-mC-containing DNA fragments were enriched from one μg of the sonicated genomic DNA by immunoprecipitation (IP) with mouse monoclonal anti-5mC antibody (Eurogentec, # BI-MECY-0100) as described previously. IP and 10% input DNA samples were purified using MinElute PCR Purification Kit (QIAGEN) and subjected to qPCR with Bdnf promoter IV and Gapdh primers to measure the enrichment of the DNA fragment containing the Bdnf promoter IV region. Primer sequences are provided in Table S2C. The percentage input was calculated by first normalizing IP to input DNA using the formula $(2^{(Ct(10\%\ input) - 3.32)} - Ct(IP)) \times 100$ as described previously. Gapdh was used as an internal normalization control.

Chromatin immunoprecipitation (ChIP). ChIP assays from mouse primary neurons were performed using Magna ChiP kit (Millipore) and anti-H3K4me3 antibody (Millipore, 17–614). The percentage input was calculated as $(2^{(Ct(10\%\ input) - 3.32)} - Ct(IP)) \times 100$ and compared between WT and mutant Htt-expressing neurons. Sequences of the primers used to amplify the BDNF promoter IV fragment are listed in Table S2D.

Quantitative reverse transcription PCR (qRT-PCR). RNAs were isolated from cultured neurons 5 days after infection of Htt lentiviruses and mouse brain using the RNeasy Plus Mini Kit (QIAGEN) and RNeasy Plus Universal Mini Kit (QIAGEN), respectively. Reverse transcription was performed with High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems). qPCR was performed using Power SYBR Green PCR Master Mix (Applied Biosystems) on CFX Connect Real-Time System (Bio-Rad). β-actin and hypoxanthine phosphoribosyltransferase I (Hprt) or 18S rRNA were used as reference genes for data normalization unless otherwise stated. Relative mRNA levels were calculated using the ΔΔCq method. Sequences of the primers used for qRT-PCR analysis are listed in Table S2A.

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How to cite this article
The authors declare no competing financial interests.

Acknowledgements
We thank the members of the H.Y. and A.H.K. for helpful discussions and technical support. We thank Dr. Zhongqiu Zhao for technical support. We thank for Dr. Jeff Milbrandt Dr. Alykhan Shamji, and Dr. Chuan He for helpful advice. This work was supported by US National Institutes of Health grants K01 AG033724 (to H.Y.) and K08 NS081105 (to A.H.K.), the Brain & Behavior Research Foundation (NARSAD Young Investigator Award to H.Y.), the McDonnell Center for Cellular and Molecular Neurobiology and the Hope Center for Neurological Disorders at, Washington University, the Uehara Memorial Foundation (Research Fellowship to T.D.), and the Hope Center Animal Surgery Core and the High-throughput Screening Center at Washington University School of Medicine.

Author Contributions
Y.P., T.D., Y.H.C., and H.Y. performed and analyzed experiments. Y.S., X.X., S.P., S.J.S., T.W., and A.H.K contributed to experiments and analysis. A.H.K edited the manuscript. H.Y. conceived the research project, analyzed the data, and wrote the manuscript.

Additional Information
Supplementary information accompanies this paper at http://www.nature.com/srep

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Pan, Y. et al. Inhibition of DNA Methyltransferases Blocks Mutant Huntingtin-Induced Neurotoxicity. Sci. Rep. 6, 31022; doi: 10.1038/srep31022 (2016).