

Early postnatal behavioral, cellular, and molecular changes in models of Huntington disease are reversible by HDAC inhibition

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Huntington disease (HD) is an autosomal dominant neurodegenerative disorder caused by expanded CAG repeats in the huntingtin gene (HTT). Although mutant HTT is expressed during embryonic development and throughout life, clinical HD usually manifests later in adulthood. A number of studies document neurodevelopmental changes associated with mutant HTT, but whether these are reversible under therapy remains unclear. Here, we identify very early behavioral, molecular, and cellular changes in preweaning transgenic HD rats and mice. Reduced ultrasonic vocalization, loss of prepulse inhibition, and increased risk taking are accompanied by disturbances of dopaminergic regulation in vivo, reduced neuronal differentiation capacity in subventricular zone stem/progenitor cells, and impaired neuronal and oligodendrocyte differentiation of mouse embryo-derived neural stem cells in vitro. Interventional treatment of this early phenotype with the histone deacetylase inhibitor (HDACi) LBH589 led to significant improvement in behavioral changes and markers of dopaminergic neurotransmission and complete reversal of aberrant neuronal differentiation in vitro and in vivo. Our data support the notion that neurodevelopmental changes contribute to the prodromal phase of HD and that early, presymptomatic intervention using HDACi may represent a promising novel treatment approach for HD.

neurodegeneration | development | experimental therapy | animal model | multiomics

Pathological CAG repeat expansions in the human *Huntingtin* (*HTT*) gene on chromosome 4p16.3 cause autosomal dominant Huntington disease (HD), which is clinically characterized by the triad of chorea, cognitive dysfunction, and psychiatric symptoms (1, 2). After clinical onset the disease inevitably progresses, leading to severe disability and premature death with no effective disease-modifying treatments available.

The CNS pathology in advanced stages of HD is characterized by a loss of striatal medium spiny neurons (3) and cortical structures (4) as well as the presence of HTT⁺ and ubiquitin⁺ cytoplasmic and nuclear protein aggregates in neurons (5–7). The lengths of pathological CAG repeat expansions, which extend the polyglutamine stretch within HTT, correlate with earlier disease onset and greater severity (1, 8, 9). Over the past decades, the field has shifted from considering HD solely as a late-onset neurodegenerative disease: Instead HD is

more recognized as a developmental disorder with various effects of mutant HTT (mHTT) on early neurodevelopment (10, 11).

HTT is expressed in preimplantation stages of the embryo (12, 13), and loss of HTT expression in mice is embryonic lethal (14–16), underlining its developmental importance. Several studies document mHTT-dependent changes during development, both

Significance

In Huntington disease (HD) gene carriers the disease-causing mutant *Huntingtin* (m*HTT*) is already present during early developmental stages, but, surprisingly, HD patients develop clinical symptoms only many years later. While a developmental role of *Huntingtin* has been described, so far new therapeutic approaches targeting those early neurodevelopmental processes are lacking. Here, we show that behavioral, cellular, and molecular changes associated with m*HTT* in the postnatal period of genetic animal models of HD can be reverted using low-dose treatment with a histone deacetylation inhibitor. Our findings support a neurodevelopmental basis for HD and provide proof of concept that pre-HD symptoms, including aberrant neuronal differentiation, are reversible by early therapeutic intervention in vivo.

The authors declare no conflict of interest

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in vivo and in stem cell-derived culture paradigms (11, 17–21). Specifically, mHTT affects striatal development in the Hdh-Q111 embryo (22), and conditional mHTT expression during development is sufficient to generate HD-like symptoms (23, 24). In animal models, early behavioral, cellular, and metabolic abnormalities are discernible in 4-wk-old tgHD rats and R6/2 mice, and changes in myelination were found in 14-d-old Hdh-Q250 mice (25–28). These studies indicate that HD pathology includes a neurodevelopmental phase that is present long before the onset of HD-like neurodegeneration and which not only contributes to the classical disease symptoms but is also likely to shape prodromal stages of HD.

In contrast to the later stages of symptomatic HD, prodromal manifestations are much less clearly defined. Previous studies indicate that emotional dysfunction and minor motor signs may precede the characteristic symptomatology (29, 30). Longitudinal studies of HD carriers aim to define the spectrum of early psychiatric symptoms in HD (29, 31). Additionally, differences in cranial volume and growth have been observed in young, presymptomatic gene carriers (32). Thus, the clinical and molecular delineation of the early stages of HD is a central issue in the implementation of future targeted treatment strategies and the identification of new biomarkers.

Here, we provide evidence that changes in neuronal differentiation continue postnatally in two mouse and rat HD animal models and that molecular, cellular, and behavioral alterations may be reversible through early therapeutic intervention with histone deacetylase inhibitors (HDACi). HDACi are promising compounds that are neuroprotective in HD mice (33), alleviate motor symptoms in the R6/2 model (34, 35), and were in clinical trials for HD (36).

A previous study showed that mHTT affects neural development pathways in HD-derived induced pluripotent stem cells (iPSCs) (21), and we extend these findings to the systems level, revealing cellular and behavioral correlates of neurodevelopmental geneexpression pathways. Importantly, these mHTT-induced changes are reversible by low-dose therapy with the histone deacetylase inhibitor LBH589 (Panobinostat), which restores neural differentiation phenotypes in vitro and in vivo.

Results

Transgenic Rodent HD Models Display a Postnatal Behavioral Phenotype with Decreased Anxiety. We began by investigating if the presence of mHTT in the developing CNS causes measurable changes in early postnatal HD animals. Therefore, we employed the earliest quantifiable behavioral assays and tested transgenic tgHD (51 CAG repeats, adult onset) and BACHD mouse (97 CAG repeats, juvenile onset) pups for emotional behavior (ultrasonic vocalization, USV) at postnatal day (P) 10, sensorimotor gating (prepulse inhibition, PPI) at P17, and risk-taking behavior [novel cage test, NCT (37)] at P21. USV analysis revealed a decreased number and shorter duration of ultrasonic calls from both tgHD rat and BACHD transgenic pups separated from their dam compared with WT littermates (Fig. 1 A and B), indicating changes in perception (reduced awareness), emotionality (reduced anxiety), and/or executive functions (reduced pattern to vocalize as an indicator of emotional and/or motor functions). We employed PPI to test for a reduction in sensorimotor gating and observed a loss of PPI in tgHD rats but not in BACHD mice (Fig. 1 C and D). In a test for anxiety and risk-taking behavior, transgenic pups from both models showed greater mobility and remained in the center area of a new cage for significantly longer durations than WT pups, suggesting reduced anxiety and increased risk-taking behavior (Fig. 1 E and F). These combined tests indicate decreased anxiety levels in early postnatal HD rodents. Fig. 1 G and H summarize the behavioral phenotypes of the tgHD rat model and the BACHD mouse model, respectively.



Fig. 1. Early behavioral phenotyping in transgenic models of Huntington disease. Rat (SPRDtgHD) and mouse (BACHD) pups were screened for a behavioral phenotype between P10 and P21. (A and B) The number and total duration of ultrasonic calls in WT, hemizygous (HET), and homozygous (HOM) rat pups (n > 30) (A) and WT and hemizygous (BACHD) mouse pups (n > 7) (B) were determined at P10. Transgenic pups emitted significantly fewer calls of shorter duration. (C and D) Startle response and PPI were measured at P17. Transgenic HD rat pups showed a significant loss of PPI at a prepulse of 80 dB white noise (n > 8) (C), while no differences were observed in BACHD mice at this age (n = 10)(D). (E and F) Altered exploration and risk behavior were detected in transgenic preweaning rats (E) as well as in transgenic mice at P21 (F) using the NCT. The time spent in the center of the cage was significantly increased in tgHD homozygous rat pups (n = 5) (E) and BACHD mouse pups (n = 8) (F), while the time spent in the wall area was significantly decreased. (G and H) A time-course graph illustrates the development of the behavioral phenotype of the transgenic tgHD rats (G) and BACHD mice (H). Data represent means \pm SEM. Statistical analyses using ANOVA: *P < 0.05, **P < 0.01, and ***P < 0.001 vs. WT pups.

Dopaminergic and Glutamatergic Imbalance in Postnatal tgHD Rat Striatum. We then analyzed gene expression in the striatum of homozygous tgHD P10 rat pups and age-matched WT littermates using Affymetrix arrays to identify molecular changes that may be the underlying cause for the early behavioral symptoms. This revealed a high number of differentially regulated genes (Fig. 2*A* and Dataset S1). Using the Ingenuity Pathway Analysis software to classify genes according to function, we identified 17 aberrantly regulated candidate genes associated with "behavior" (Fig. 2 *B* and *C*). qRT-PCR validation confirmed aberrant regulation of seven candidates: angiotensinogen (*Agt*), ATPase, Ca²⁺ transport (*Atp2A2*), Forkhead box G1B (*FoxG1B*), hypocretin (orexin) receptor (*Hcrtr2*), potassium channel, member 1 (*Kcnc1*), solute carrier family 6, 3 (dopamine transporter) (*Slc6A3*), and tyrosine hydroxylase (*Th*) (Fig. 2*B*). Since the two most up-regulated of these seven candidates were related to dopaminergic signaling (Fig. 2*C*), we analyzed striatal

dopaminergic circuits in more detail and further quantified the mRNA levels of DARPP-32, the dopamine receptor D1A, and protein kinase A (PKA) by qRT-PCR. In contrast to the observed upregulation of *Slc6A3* and *Th*, we found that all three genes were significantly down-regulated in the striatum of tgHD P10 pups (Fig. 2D).

Western blot analyses of total protein extracts from striatal tissue at P10 confirmed significantly reduced protein levels of DARPP-32 (Fig. 2*E*). Of note, there was a trend toward increased phosphorylation at Thr^{75} and Thr^{34} in DARPP-32.

These studies were complemented by autoradiography for striatal expression of D1A, D2, NMDA, 5-HT2, adenosine A1,



Fig. 2. Molecular analyses of tgHD P10 striata. (*A*) Microarray analysis of rat striata at P10 (WT vs. HOM, n = 5 each) demonstrated clusters of up- (yellow) and down- (blue) regulated gene expression. (*B*) Ingenuity Pathway Analysis revealed 17 target mRNAs associated with behavior, 7 of which (#) were validated by qRT-PCR (*P* values are from ANOVA vs. WT; n > 8). (*C*) Network analysis indicated deregulation of dopaminergic signaling; relevant pathways are highlighted in yellow). Rat P10 striatal mRNA expression of DARPP-32, PKA, and dopamine receptor D1A is down-regulated, whereas the expression of the dopamine transporter Slc6a3 is up-regulated. (*D*) Data are displayed as mean normalized expression (percentage of WT; ANOVA; n > 8). (*E*) DARPP-32 down-regulation two-tailed *t* test; n > 5) could be confirmed in Western blot studies. Phosphorylation levels of DARPP-32 (Thr³⁴ and Thr⁷⁵) in P10 striata were not significantly different in WT and transgenic pups. (*F*) Receptor autoradiography revealed a significant reduction of dopamine receptor D1A density and NMDAR density in the striata of transgenic rat pups (two-tailed *t* test; n = 5). Representative autoradiographs of coronal sections of P10 rats are presented. (G) Immunohistochemistry confirmed the reduced expression of DARPP32 and TH in the striatum of P10 tgHD pups (shown are representative images from n = 10 pups). Data represent means \pm SEM: **P* < 0.05, ***P* < 0.01, and ****P* < 0.001 vs. WT pups.

and adenosine A2A receptors in P10 rat pups. We found decreased levels of the dopaminergic receptor D1A (Fig. 2F) and NMDA receptor (NMDAR) density (Fig. 2F and SI Appendix, Fig. S1A). Further Western blot analyses of NMDAR subunits revealed down-regulation of NR1, NR2B, and NR2C (SI Appendix, Fig. S1B). Immunohistochemical analysis of P10 tgHD striata confirmed reduced expression of DARPP32 and TH (Fig. 1G).

As mRNA expression changes do not always translate into altered protein levels, and because other regulatory mechanisms may also affect protein synthesis, we performed proteomic analysis using 2D electrophoresis and subsequent MALDI-TOF of P10 pup striata as well. In the R6/2 model, large-scale alterations of protein levels precede the onset of classical symptoms (38). Similarly, we found 156 proteins to be significantly altered in pre-HD tgHD rat brains. These could be classified into six Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways, confirming the observed alterations in metabolic pathways (38) and further identifying changes in axon sprouting and neurodegenerative pathways (Dataset S2). These studies revealed an additional down-regulated regulatory subunit of protein phosphatase 1 (PPP1R7) (39, 40). Taken together, these data reveal imbalances in dopaminergic and glutamatergic pathways in the early postnatal period of tgHD rats sufficient to explain the behavioral phenotype.

Altered Neuronal and Oligodendroglial Differentiation Capacity in HD. Olfactory dysfunction is among the earliest discernible symptoms in neurodegenerative diseases, including Parkinson, Alzheimer, and Huntington disease (41). Because of the wellestablished link between subventricular zone (SVZ) neurogenesis and the olfactory system, the observed association of aberrant SVZ neurogenesis and olfactory dysfunction in neurodegenerative diseases (42), the reported regulatory functions of dopaminergic and glutamatergic signaling for neurogenesis (43), and the potential implications of the olfactory system for our observations of animal behavior, we investigated SVZ neurogenesis in our model.

While neurodegeneration is characteristic of HD, previous studies observed aberrant neuronal differentiation during embryonic development in vitro and in vivo in the Hdh-Q111 model (22, 23), as well as in iPSC-derived neural precursors from HD patients (21). Therefore it is likely that any aberrations on the cellular and molecular level underlying behavioral changes are the product of aberrant neurodevelopmental rather than neurodegenerative processes. Consequently, we asked whether mHTT affected neuronal differentiation capacities of SVZ neural stem/progenitor cells (NSCs) during postnatal development. We cultured NSCs from P10 WT, tgHD+/-, and tgHD+/+ pups and differentiated them into glial and neuronal lineages. This resulted in a significant, gene dose-dependent reduction in neuronal numbers at all time points (Fig. 3 A-C). Analysis of differentiation to mature neuronal phenotypes revealed a reduction in bIII tubulin⁺ neurons at 7 d in vitro (DIV) (Fig. 3B), as well as lower numbers of NeuN⁺ neurons at 14 and 28 DIV (Fig. 3B). Likewise, Darpp32⁺ cell numbers were reduced in tgHD^{+/+} cultures at 14 and 28 DIV, while we observed an increase at 14 DIV and a decrease at 28 DIV in tgHD^{+/-} cultures (Fig. 3C). Astrocyte numbers were unchanged, but oligodendrocyte counts were significantly increased after differentiation (Fig. 3 A and B). These findings demonstrate an increase in oligodendroglial differentiation at the expense of neuron production from SVZ-derived NSC cultures in heterozygous and homozygous tgHD pups. Differentiation of tgHD cultures was accompanied by an increase in apoptotic cells identified by active caspase 3 staining. This is also reflected in reduced total cell counts in $tgHD^{+/-}$ and $tgHD^{+/+}$ cultures compared with WT cultures (Fig. 3D).

To determine if neuronal, oligodendroglial, and glial differentiation capacity was also affected in BACHD mice, we generated neurospheres from BACHD and WT littermates at E13.5. We detected significant reductions in the total number of cells, MAP2-immunopositive neurons, and 2'3'-cyclic-nucleotide 3'phosphodiesterase (CNPase)-immunopositive oligodendrocytes in BACHD cultures compared with WT cultures at 7 DIV, while astrocyte numbers remained unchanged (Fig. 3*E*).

Because of the profound changes in differentiation capacity between WT and tgHD NSC cultures, we were interested in whether in vivo neurogenesis was affected by the presence of mHTT. To address this, we crossbred tgHD rats with a model expressing the reporter dsRed2 under control of the Dcx promoter (dcx::dsRed2,tgHD) (44). Dcx is a marker of newborn, migratory neurons and is expressed up to 14 d after the birth of new neurons (45). We analyzed sagittal sections of dsRed2⁺ WT and tgHD pups, revealing minor differences in rostral migratory stream (RMS) and olfactory bulb (OB) volumes and a significant reduction in the content of Dcx⁺ neurons (proportional to the relative fluorescence intensity; see heatmaps in Fig. 3*F*). This corroborates the reduced capacity for SVZ neurogenesis in the tgHD model.

Of note, diffusion tensor imaging (46) revealed a trend toward an increase in axial diffusivity, radial diffusivity, and mean diffusivity and a reduction in fractional anisotropy in tgHD pups compared with WT pups (*SI Appendix*, Fig. S1C).

Low-Dose LBH589 Treatment Restores the Early Behavioral, Cellular, and Molecular Phenotype in HD. The observed mHTT-associated aberrations at the molecular, cellular, and behavioral levels in these HD models extend previously reported alterations in embryonic development to the early postnatal phase, a time of neuronal maturation and gliogenesis. We next tested whether therapeutic intervention can restore these changes.

HDACi have a long history of use for their mood-stabilizing and antiepileptic effects. In models of HD, previous studies have demonstrated that the HDACi SAHA (Vorinostat) could reverse motor symptoms and improve phenotypes in the R6/ 2 model (34, 35). Because of the neuroprotective effects of LBH589 in mouse models of HD (33) and its ability to cross the blood-brain barrier (Dataset S3) (47), we treated tgHD pups with this HDACi. TgHD rat pups and WT littermates were subjected to four different dose regiments, administered every other day between P8 and P20 (Fig. 4A and SI Appendix, Fig. S2 A and B). We analyzed USV at P11, startle response and PPI at P17, and the NCT at P21. Administration of 0.001 mg/kg LBH589 led to significantly increased USV in transgenic rat pups (Fig. 4B). LBH589 had a suppressive effect on PPI and startle response independent of the individual genotype at all doses (Fig. 4C). In addition, performance in the NCT was improved in tgHD rats: Duration in the center was reduced, and duration in the wall areas increased to match values of vehicletreated WT animals (Fig. 4D). In contrast, LBH589 had no effect on WT animals in this test. Furthermore, the average velocity and track length in tgHD pups were restored to WT levels (SI Appendix, Fig. S2A), while in the 0.01 mg/kg WT group the average velocity and track length were significantly decreased compared with vehicle-treated WT animals. The improved performance of LBH589-treated tgHD pups in USV and NCT tests indicates that this compound alleviates the reduced anxiety and increases risk-taking behavior in these animals.

We have previously demonstrated that HDACi increase the neuronal differentiation capacities of NSCs (48). Therefore, we tested whether exposure to LBH589 could restore neuronal differentiation of tgHD NSCs in vitro. After testing for optimal dose regimens (*SI Appendix*, Fig. S2C), we found that treatment of differentiating NSC cultures with 10 nM LBH589 for 24 h restored neuronal differentiation of tgHD cultures to WT levels.



Fig. 3. (*A, Left*) Cellular analysis of tgHD P10 neurogenesis. Quantitative analysis of P10 rat neurosphere differentiation demonstrated reduced neuronal (bll tubulin, green, *Upper*) and increased oligodendroglial (CNP, green, *Lower*) differentiation, while astroglial (GFAP, red, *Upper*) fates were unaffected (Scale bar, 20 μ m applies to all images.). (*Right*) Graphical representation indicates percentages \pm SEM of individual cell types. (*B*) After 7 d in vitro (DIV) differentiation, neuron numbers were significantly decreased, while oligodendrocyte counts were increased. Astrocyte numbers were not affected significantly. Numbers of mature, NeuN-expressing neurons were significantly reduced at 14 DIV and 28 DIV. (C) Similarly, DARPP32⁺ cell counts were significantly reduced at both time points. (*D*) The differentiation phenotype was accompanied by an increase in apoptosis, which is reflected in lower cell counts after differentiation. In *A–D*: ANOVA; *n* = 3 independent experiments with 10 replicates. (*E*) Differentiation of BACHD embryonic (E13.5) neurospheres revealed significantly lower numbers of total cells and MAP2⁺ and CNPase⁺ cells compared with WT cultures (ANOVA; *n* = 8). GFAP⁺ astrocyte counts were unchanged. (*F*) In vivo changes in neuronal differentiation are reflected in the RMS and OB of P10 Dcx-dsRed2-tgHD double transgenic pups (Mann–Whitney *U* test, *n* = 7). (*Upper Left*) Inverted fluorescence images. (*Lower Left*) Heatmaps of fluorescence intensity. (*Right*) While the volume of RMS and OB was not significantly different, the mean fluorescence intensity (proportional to neuronal numbers) was significantly reduced in both structures. Data represent means \pm SEM. **P* < 0.05; ***P* < 0.01; ****P* < 0.001.

Comparing bIII tubulin⁺ cells at 7 DIV and NeuN⁺ and Darpp32⁺ cells at 14 DIV, we observed a significant increase in neurons at all time points (Fig. 4*E*), indicating improved regenerative capacities of SVZ stem/progenitor cells after treatment. LBH589 treatment had no effect on total cell numbers in WT controls but increased tgHD total cell numbers (Fig. 4*F*).

Similarly, treatment of E13.5-derived BACHD NSC cultures with 10 nM LBH589 increased the percentage of MAP2⁺ neurons in both WT and BACHD cultures at 7 DIV (Fig. 4G).

LBH589 exerted a positive effect in WT cultures at 1 nM, whereas BACHD cultures required 10 nM to increase the percentage of MAP2 cells. Furthermore, treatment increased the percentage of CNPase⁺ oligodendrocytes at 7 DIV. There were no differences in GFAP⁺ astrocyte numbers between any treatment conditions in BACHD and WT cultures (*SI Appendix*, Fig. S2 D–G).

Transcriptomic and proteomic profiling (Fig. 2, *SI Appendix*, Fig. S3, and Dataset S2) identified alterations in several pathways relevant to HD pathogenesis and neurodevelopment (*SI*



Fig. 4. (A) Treatment of early phenotype using HDAC inhibitor LBH589. tgHD HOM pups and WT controls were treated with the HDACi LBH589 from P8 to P20 (n = 10 each). Behavioral readouts were USV (P11), startle response and PPI (P17), and NCT (P21). Four different doses (0.001, 0.01, 0.1, and 1.0 mg/kg body weight) were administered i.p. every other day. (B) The numbers of ultrasonic calls were significantly increased in transgenic animals treated with LBH589 compared with vehicle controls. The increase in the total duration in transgenic animals did not reach significance except for the 0.1 mg/kg group (0.001 mg/kg: P = 0.2561; 0.01 mg/kg: P = 0.1949; 0.1 mg/kg: P = 0.0016; ANOVA; n > 10). (C) PPI was suppressed in animals treated with LBH589 in all doses tested compared with WT (ANOVA; n > 3). (D) LBH589 treatment significantly decreased the time spent in center areas and increased the time spent in wall areas in transgenic rats compared with corresponding vehicle-treated transgenic animals (ANOVA; n > 4). LBH589 had no significant effect on WT animals. (E) LBH589 treatment of differentiating neurosphere cultures restored tqHD HOM neuron numbers to WT levels. Shown are representative images of cultures treated with 10 nM LBH589 for 24 h. (Scale bar, 20 µm in both images.) TuJ⁺ neurons were counted after 7 d, and NeuN⁺ and Darpp32⁺ neurons were quantified after 14 d. TuJ⁺, NeuN⁺, and Darpp32⁺ cells were within the range of WT cultures (ANOVA; n = 3 independent experiments with 10 replicates). (F) LBH589 treatment resulted in increased cell counts in transgenic but not in WT cell cultures. (G, Right) The percentage of MAP2-immunopositive neurons was significantly increased with 10 nM LBH589 in WT and BACHD E13.5 neurosphere cultures (ANOVA; n = 8). (Left) A representative photomicrograph showing MAP2⁺ neurons (green) and DAPI (blue) from WT and BACHD cultures. (Scale bar, 100 μm.) (H) LBH589 treatment (0.001 mg/kg) increased striatal RasGRP2 gene expression and DARPP32 and PPP1R7 protein levels in P10 tgHD pups but not in WT littermates (ANOVA; n > 6). (I, Left) Heatmaps of dcx-stained immunofluorescence micrographs from P21 tgHD pups highlight SVZ-RMS neurogenesis in vivo. (Scale bar, 100 µm.). (Right) LBH589 treatment restored both RMS volume and dcx fluorescence intensity in tgHD pups to WT levels (seven injections of 0.001 mg/kg compared with vehicle control; two-way ANOVA; n > 3). Data represent means ± SEM. Significant effects vs. WT (*P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001) and treatment effects vs. vehicle control $(^{\#}P < 0.05; ^{\#\#}P < 0.01; ^{\#\#\#}P < 0.001).$

Appendix, Fig. S4). We therefore tested whether in vivo LBH589 treatment affected expression of a target gene of the dopamine pathway in medium spiny neurons (RasGRP2), which is down-regulated in HD (49). LBH589 treatment increased RasGRP2 expression and restored protein levels of two additional markers of striatal development, DARPP-32 and PPP1R7 (Fig. 4*H* and *SI Appendix*, Fig. S5).

To determine whether LBH589 was capable of restoring neurogenesis in vivo, we quantified RMS volume and Dcx staining intensity in vehicle- and LBH589-treated WT and tgHD pups at P21. Continuous administration of 0.001 mg/kg LBH589 from P8 to P20 (seven doses) could restore in vivo neuronal differentiation and neurogenesis in tgHD pups to WT levels (Fig. 4*I*).

Thus, early postnatal behavioral, cellular, and molecular alterations of HD can be alleviated by low-dose therapy with LBH589.

Discussion

Here we describe behavioral, cellular, and molecular changes during a developmental phase of gliogenesis and neuronal maturation that is comparable to the third trimester of human gestation (50). Importantly, these changes can be reversed by treatment with HDACi, which fully restores alterations in neuronal differentiation in HD pups. Hence, these data contribute to the increasing evidence of neurodevelopmental aberrations in genetic animal models of HD and suggest that HD should be considered a neurodevelopmental disorder.

We identify reduced separation-induced USV at P10 and increased risk-taking behavior at P21 as key features of an early behavioral phenotype in transgenic HD mice and rats. These features may correspond to an anxiolytic phenotype similar to the reduced anxiety-like behavior found in transgenic HD rats at 1 mo of age (26) or may be attributable to depression-like features as described in 2-mo-old BACHD mice (51). Dopaminergic and glutamatergic signaling regulate neuronal differentiation during development, and deregulation of these pathways may provide a molecular basis for HD-associated behavioral changes. These species-independent behavioral abnormalities provide a readout for the prodromal phase of HD, which is characterized by neurodevelopmental rather than neurodegenerative mechanisms. mHTT causes aberrant neuronal differentiation during embryonic development and from iPSCs (21-24). Our findings link alterations in neuronal differentiation and molecular pathways to behavior on the systems level.

These observations support the notion that HD pathology consists of at least three phases: (i) an early phase with successful adaptation to the effects of accumulating mutant Htt, likely to be outbalanced by changes in early neurodevelopment. This early phenotype presents with signs of overcompensation and is followed by an intermediate and largely silent phase, which, depending on the number of CAG repeats at mid-age, shifts into a third phase of overt decompensation and neurodegeneration. The notion that this early phase of HD may be viewed as a slowly progressive neurodevelopmental disorder that ultimately transforms into a neurodegenerative disease is documented in many studies identifying symptoms long before the onset of neurodegeneration in animal models (25-28, 52). In view of this evidence for a "pre-HD syndrome," it should be mentioned that such changes may not be "disease-like" but instead may reflect a whole spectrum of signs and symptoms attributable to higher glutamine availability in the CNS and/or successful compensation and repair of the very early damage associated with mHTT. The former would be in line with the hypothesis of polyglutamine diseases arising from evolutionary search mechanisms (53) attempting to increase central glutamine storage in the brain, while the latter would imply that neurotrophic factors are induced, which may affect development in all its dimensions of growth, maturation, and learning. Whether mHTT causes developmental aberrations in gene carriers is still unclear, but recent studies have found differences in growth and development as well as in cranial volume in both adult (32) and juvenile (54) premanifest gene carriers.

Microarray analyses detected deregulation of the dopaminesignaling pathway in postnatal tgHD rats. This was corroborated by findings of deficits in the D1/cAMP/PKA/DARPP-32 signaling cascade of P10 tgHD rats. These findings in the postnatal tgHD rat brain mirror previous imaging studies in asymptomatic HD gene carriers (55, 56), which revealed a reduction of striatal D1 and D2 receptors. Evidence of dopaminergic deregulation and receptor imbalance has been further substantiated in molecular studies of postmortem HD brains (57-59) and imaging studies of symptomatic HD patients (60, 61). Previous studies in 4-wk-old HD mice also demonstrated a reduction of total levels of DARPP-32 and D1 receptor protein levels (62). We previously showed that a significant loss of striatal D1 receptor expression and DARPP- 32^+ cells is present in 14-mo-old male tgHD animals. Beyond deregulation of dopaminergic-signaling pathways, we observed significantly reduced NMDAR density in the striatum of tgHD P10 rat pups. NMDAR numbers are reduced, particularly in the basal ganglia, in both symptomatic (63, 64) and presymptomatic (65) HD patients. Early changes in extrasynaptic NMDAR signaling and expression have also been reported in 1-mo-old presymptomatic YAC72 and YAC128 mice before the onset of motor dysfunction and neuronal loss (66). Furthermore, there is evidence indicating that mHTT increases NR2B-containing extrasynaptic NMDARs in the striatum of young YAC72 and YAC128 mice (67). By contrast, our own data in P10 rat tissue show a downregulation of the NR1, NR2B, and NR2C subunits of NMDARs. A pathogenic link between dopamine and glutamate signaling is provided by studies showing that striatal dopamine modulates ionotropic glutamate receptors by altering glutamate currents and by modifying glutamate receptor surface localization (68, 69). Since D1 receptors and NMDARs are capable of direct interaction, which affects receptor activity (70), it is possible that the aberrant striatal receptor expression observed in P10 tgHD rats is due to abnormal glutamate/dopamine receptor interactions. Alternatively, down-regulation of D1A receptors and NMDARs could also suggest a developmental delay in tgHD pups.

SVZ neurogenesis has been shown to affect social behavior (71), and aberrations in olfaction and SVZ neurogenesis are among the earliest symptoms in neurodegenerative diseases (41, 42). SVZ neurogenesis is modulated by dopamine and glutamate (43); thus it is likely that the observed deregulation of both pathways extends to neural stem cells as well. We demonstrate that SVZ-derived NSC cultures show a prominent reduction of their neuronal differentiation in both tgHD and BACHD models, which is corroborated by a reduction of NeuN⁺ neurons at 14 and 28 DIV in tgHD. The different results in oligodendrocyte differentiation between tgHD and BACHD models may be explained by their different ages at isolation. Oligodendroglial maturation and myelination start around P14 in vivo; thus embryonic-derived BACHD cultures may be too early to reflect mHtt effects on the oligodendrocyte lineage. Altered SVZ neurogenesis could be confirmed in vivo in P10 tgHD rats. Others have shown the relevance of mHTT during development (22, 23) as well as its impact on the differentiation capacities of pluripotent SC-derived neural precursors (21, 72). mHTT has been shown to affect the proliferation and cell fate of cortical progenitor cells (18). Consequently, the reduced neuronal differentiation observed here is likely due to direct effects of mHTT in neural precursors.

Bystander microenvironmental effects, e.g., from microglia, may contribute as well, as it has been shown that these cells affect SVZ neurogenesis in vitro (73). The specific effects of mHTT on postnatal neurogenesis in symptomatic HD models are somewhat controversial. While some studies did not find changes in SVZ proliferation or differentiation in R6/2 or YAC128 mice (74, 75), others detected reduced levels of neurogenesis in the R6/2 model (76, 77) and aberrant neuronal differentiation in tgHD rats (78). In HD patients, cell proliferation in the SVZ is increased, which correlates with disease grade (79). Importantly, our observations are made long before the onset of classical symptoms. Thus, it is conceivable that mHTT reduces the generation of new neurons during earlier stages, while SVZ cell proliferation as a reaction to neurodegenerative processes dominates later stages of the disease.

Reduced neurogenesis may ultimately contribute to disease progression, i.e., lack of regeneration of striatal cell loss and dysfunction. This hypothesis is further supported by the observation of reduced numbers of striatal neurons (80) and reduced numbers of medium spiny neurons (81) in the tgHD rat model.

Transcriptional deregulation is known to be part of the pathogenesis of HD (82, 83). A number of studies showed that HDACi ameliorate neurodegeneration in cell systems and fly models (84) as well as motor deficits in a mouse model of HD (34, 85). Here we treated postnatal tgHD rats with the HDACi LBH589 (86) to intervene with neurodevelopmental changes in a preclinical setting. LBH589 is known to cross the blood–brain barrier efficiently and has already been tested in preclinical studies (33, 47). Lowdose treatment with LBH589 led to a partial reversal of behavioral symptoms and restored aberrant neuronal differentiation in tgHD and BACHD cultures in vitro and in tgHD pups in vivo. We conclude that very early LBH589 treatment may be a promising treatment approach in HD.

Methods

Animals. We used several tgHD litters derived from mating male and female tgHD^{+/-} rats carrying a truncated human *HTT* fragment of 51 CAG repeats under control of the native rat HTT promoter (87). BACHD mouse pups were derived from mating hemizygous male BACHD mice (expressing floxed *HTT* with 97 CAG repeats) with female FVB/N dams (88). All animal experiments presented here were approved by local ethical boards of the District Government of Middle Franconia, Bavaria, Germany (approval no. 54-2532.1-16/08) and the University of Florida Institutional Animal Care and Use Committee, and were conducted according to local, NIH, and ARRIVE guidelines (89).

Behavioral Phenotyping.

USV. P10 rat and mouse pups were placed into a recording cage in a soundisolation box. The detection sessions were initiated directly thereafter and were always performed by the same experienced experimenter between 8 and 10 AM. A maximum of two pups per litter were tested for 5 min each. For P10 rats and mice signals were recorded at a 35–250 kHz range using an Avisoft Ultra Sound Gate 116Hb (Avisoft Bioacoustics). Incoming signals were displayed as a time–event plot by Avisoft recorder software. For further statistical analysis, the parameters number of events (*n*) and total duration of cues (s) were analyzed using one-way ANOVA.

Acoustic startle response and PPI. Startle response and PPI were measured in homozygous (+/-; $n \ge 10$), hemizygous (+/-; $n \ge 10$), and WT (-/-; $n \ge 10$) tgHD rats and hemizygous (+/-; $n \ge 10$) and WT (-/-; $n \ge 10$) BACHD mice at P17. The test point was validated in preliminary experiments. Pups were placed in a startle chamber, and the protocol was conducted as described previously

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(90). Results were analyzed using one-way ANOVA and are displayed as the percentage of startle-response inhibition.

NCT. The NCT was performed in P21 preweaning homozygous (^{+/+}; $n \ge 5$), hemizygous (^{+/-}; $n \ge 5$), and WT pups analyzed using one-way ANOVA.

Treatment with HDACi LBH589. Pups were genotyped 3 d after birth, divided into groups of 10 according to genotype, and marked daily throughout the experiment. The animals were treated with LBH589 (Novartis) using the following doses: 0.001 mg salt/kg body weight, 0.01 mg salt/kg body weight, 0.1 mg salt/kg body weight, and 1 mg salt/kg body weight. Control animals were treated with vehicle: 0.03 M lactic acid/5% dextrose in water buffered to pH 4.2–4.5 with 0.1 M NaOH. LBH589 was administered i.p. every 48 h from P8 to P20. The behavioral tests were performed at P11 (USV), P17 (startle response and PPI), and P21 (NCT). Pups were killed at day 21 for histology.

Double-Transgenic Model and in Vivo Neurogenesis. tgHD rats were crossbred with a rat model expressing dsRed2 under control of the dcx promoter (Dcx:: dsRed2) (91). The resulting offspring were interbred to generate hemizygous tgHD rats that were either homozygous or WT for dcx-dsRed2 (tgHD^{+/-} dsRed2^{-/-} or tgHD^{+/-} dsRed2^{+/+}). Interbreeding these animals generated tgHD WT and homozygous offspring that were heterozygous for dcx-dsRed2 $(tgHD^{-/-} dsRed^{+/-} or tgHD^{+/+} dsRed2^{+/-})$. Animals from these litters were genotyped at P3 and were used for experiments at P10. Pups were anesthetized on ice and then were transcardially perfused using 4% paraformaldehyde (PFA), and their brains were removed and postfixed in 4% PFA overnight. Fixed brains were washed twice in PBS and then were sectioned at 100 microns in the sagittal plane on a Leica VT-1000S vibratome. Serial sections from one hemisphere were mounted onto glass slides, airdried, and coverslipped using VECTASHIELD with DAPI (Vector Laboratories). Low-power fluorescent images were taken on a Leica DMLB epifluorescence microscope equipped with a CCD camera (Spot Imaging Solutions). To obtain full images of brain sections, multiple gray-scale images were acquired per section using Spot Advanced software (Spot Imaging Solutions) and were merged into a full image and inverted into black-on-white images using Photoshop CS4 (Adobe Systems). Merged images were imported into ImageJ (NIH), and area size and mean pixel values of RMS and OB were analyzed.

Data Analysis. All data were analyzed with GraphPad Prizm 7.0 (GraphPad Software), and statistical significance was determined using appropriate statistical tests, as indicated in the figure legends. A *P* value of < 0.05 was deemed significant for all analyses. We used the D'Agostino–Pearson method to test for normal distribution of data points.

Additional methods are listed in SI Appendix.

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Supplementary Information for

Early postnatal behavioral, cellular and molecular changes in models of Huntington disease are reversible by HDAC inhibition

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Other supplementary materials for this manuscript include the following:

Datasets S1 to S3

Supplementary Methods

RNA preparation and microarray analysis

Isolation of RNA from whole rat brain was performed using RNAeasy[®] Kit (Qiagen, Germany) according to the manufacturer's conditions and processed as previously described(1). Labeled, fragmented cRNA (15 μ g) was hybridized to a GeneChip Rat Genome 230 2.0 Array (Affymetrix) as previously described (1). In total, 12 cRNA samples were analyzed (whole brain from three wild type control P10 pups and three homozygous transgenic P10 pups and striatum from three wild type control P10 pups and three homozygous transgenic P10 pups). For analyses, signals were first filtered for an absolute change in signal level of 1.5-fold. The remaining transcripts were subjected to statistical analysis using a T-test. Transcripts with a fold change of 1.5 and a p-value<0.05 were considered as statistically significant regulated. Categorization was based on the netAffx annotations

(<u>https://www.affymetrix.com/analysis/netaffx/index.affx</u>). Affected gene regulation networks were determined using Ingenuity pathways analysis software (Ingenuity[®] systems, <u>www.ingenuity.com</u>).

qRT-PCR

Transcript levels of behavior-associated genes identified in gene array studies and genes involved in dopaminergic processes in the striatum were analyzed by quantitative real-time PCR as described previously (2). Primer sequences are presented in Table S1.

Receptor Autoradiography

Samples were collected and processed as described previously (3). Briefly, whole brains were rapidly removed and snap-frozen. Cryostat sections were stained for receptors of interest, placed on phosphor imaging plates (Raytest-Fuji, Straubenhardt, Germany) along with industrial tritium activity standards (Amersham Biosciences, Freiburg, Germany). Exposed imaging plates were scanned with a high-performance imaging plate reader (Raytest-Fuji). Digital receptor autoradiographs were processed using AIDA 2.31 (Raytest-Fuji).

Proteomic analysis

Samples were collected and processed as described (4). Briefly, total protein extracts from tgHD and WT P10 striata were separated using 2-DE. Protein spot patterns were evaluated using Delta2D imaging software (DECODON, Greifswald, Germany). Relative spot volume intensities were used for quantitative protein expression analysis. Data sets were analyzed using Student's *t* test. Protein spots of interest were excised from 2-D gels, trypsin-digested in gel, and fragments were analyzed by nanoflow HPLC (Dionex/LC Packings, Amsterdam, Netherlands)/ESI-MS and -MS/MS on an LCQ Deca XP ion trap instrument (Thermo Finnigan, Waltham, MA). To investigate an enrichment of specific pathways in the altered protein expression data set, we used the "Web-based gene set analysis toolkit" (WEBGESTALT) tool supplied by Vanderbilt University.

Western Blot

Dissected brain samples were snap-frozen and processed as previously described (5). Briefly, the samples were sonicated in 1% SDS and boiled for 10 min. 25 μ g of each sample was separated by SDS-PAGE using a 12% running gel and transferred to an Immobilon-P transfer membrane (Millipore). The membranes were incubated for 1 h at room temperature with 5% (w/v) dry milk in TBS-Tween 20 containing the following antibodies: against Ser⁸⁴⁵-GluR1, Ser⁸⁹⁷-NR1, Ser¹³⁰³-NR2B, Ser¹⁹ and Ser³¹-TH, Thr³⁴-DARPP-32, Thr⁷⁵-DARPP-32, against total GluR1, NR1, NR2B, NR2C, TH, CDK5 and DARPP-32 (Cell Signaling Technology, Danvers, USA). Membranes were washed three times and incubated with secondary HRP anti-rabbit antibody for 1 h at room temperature. Membranes were washed six times with TBS-Tween 20 and immunoreactive bands were detected by chemiluminescence using ECL reagents (Perkin Elmer). The autoradiograms were scanned and quantified using NIH Image 1.63 software. Levels of phosphorylated proteins were normalized to total (phosphorylated and non-phosphorylated) levels. The data were analyzed with two-tailed unpaired Student's *T*-test to evaluate statistical differences.

MRI

MRI acquisition

Diffusion weighted (DW) Imaging was performed on a horizontal 9.4 Tesla MR system (Biospec 94/20 USR, Bruker Biospin, Germany) and was acquired with a 4-shot spinecho EPI sequence (orientation invariant icosahedral encoding scheme with 15 directions and 7 diffusion sensitization or b-values (400 - 2800 s/mm²)). Additional image parameters were: image orientation = axial, TR/TE = 3000 ms/25 ms, diffusion gradient pulse duration (δ) = 5 ms, diffusion gradient separation (Δ) = 12 ms, NEX = 4, acquisition matrix = 128 x 128, slice thickness = 1 mm, number of slices = 18, in plane spatial resolution of 0.154 x 0.154 mm² (6).

MRI processing

The calculations of the invariant diffusion parametric maps, was done using custom written Matlab routines (The Mathworks, Natick, Massachusetts, USA). Standard diffusion metrics, including the three eigenvalues, mean diffusivity (MD), radial diffusivity (RD) and fractional anisotropy (FA) were calculated (7).

MRI analyses

Regions of interest (ROI) (Caudate Putamen) were manually defined on each individual and T_2 anatomical image and parametric maps (MD and FA), generated using AMIRA software (AMIRA; Template Graphics). The definition of the ROI was based on a rat brain atlas (8) and was done by a single examiner as previously described (9).

tgHD neurosphere culture

tgHD rat pups and wild type littermates were sacrificed at postnatal day 10, their brains removed and a single cell suspension prepared, cultured and maintained as described previously (10). Neurospheres after passage 2 were used for subsequent experiments. For differentiation experiments, spheres were dissociated using Accutase (PAA, Cölbe, Germany), counted and plated at 50,000 cells/ml on laminin/poly-L-ornithine (Sigma) coated coverslips. Differentiation was induced in N2 medium containing 10% fetal calf serum (Atlanta Biologicals, Lawrenceville, GA) and no mitogens. Each differentiation experiment was performed in triplicate. Some cultures were treated with different concentrations of LBH589 one hour after plating. In these cultures, medium was changed 24 hours after treatment. At various time points after plating, differentiated cultures were fixed with 4% paraformaldehyde in PBS (Sigma), washed and immunostained.

Immunocytochemistry

Fluorescence immunostaining was performed as described (10). The following primary antibodies were used: mouse anti beta III tubulin (TuJ, 1:1000, Promega), mouse anti NeuN (1:500, Chemicon), rabbit anti Darpp32 (1:500, Epitomics), rabbit anti GFAP (1:1,000, Dako), mouse anti GFAP (1:500, Sigma), mouse anti CNPase (1:100, Chemicon). Appropriate secondary antibodies were used. No signal was detected when omitting primary antibodies. Hoechst33342 (Invitrogen) was used for nuclear counterstaining. Images were taken from 10 random visual fields per coverslip on an Olympus IX-81 microscope equipped with a CCD camera. Images were imported in ImageJ and numbers of total cells (nuclei) and positively stained cells were manually counted. Only cells with neuronal morphology (compact nucleus and cell body with one prominent process longer than others) and marker expression were counted as neurons.

E13.5 embryo-derived neurosphere generation and differentiation

Neurospheres were generated from the lateral ganglionic eminences (LGE) of E13.5 embryos that resulted from the crossing of FVB/N BACHD male mice with FVB/N female mice. Note: each embryo was treated separately. At first, the uterine horns of the timed pregnant female mice were dissected out and placed in HBSS. E13.5 embryos were separately harvested and their two LGE (one from each hemisphere) were dissected out as shown in Fig. A, in a laminar flow hood under sterile conditions. The LGE were dissociated in medium used to expand the progenitors as neurospheres, according to a standardized protocol and as previously described (11). The tail of each animal was harvested and genotyping was performed, while the neurospheres were forming. The genotyping was performed using conventional PCR using the primers for the BACHD transgene: 5'-CCGCTCAGGTTCTGCTTTTA-3' (Forward) and 5'-AGGTCGGTGCAGAGGCTCCTC-3' (Reverse). After a week of growth, the neurospheres were mechanically dissociated to form secondary neurospheres. A week later, the neurospheres were isolated, washed in PBS containing 0.4% glucose and treated with 0.05% Trypsin-EDTA (Life Technologies) and mechanically triturated to obtain a single cell suspension. The cells were then counted with a hemocytometer and seeded onto poly-L-ornithine (100 µg/ml; Sigma-Aldrich) and laminin (15 µg/ml; Invitrogen) coated 96-well plates at a density of 15,000 cells/100µl in N2 medium containing 2% (v/v) N2 supplement (Thermo Fisher), penicillin and streptomycin (50 µg/ml each;

Thermo Fisher), 2mM L-glutamine (Thermo Fisher) added to Adv. DMEM/F12 medium (Thermo Fisher) supplemented with 10% fetal bovine serum (Thermo Fisher). Each embryo-derived culture was treated with 1, 10 or 100 nM of the HDAC inhibitor, LBH589 (Novartis, Switzerland) or 0.002% DMSO alone as vehicle for control condition. One well was used for each treatment condition for each embryo (n=8 embryos/genotype/treatment group). Differentiation medium was changed 24 hours later, for a medium that did not contain any of the treatments mentioned above. Cultures were fixed seven days later with 4% paraformaldehyde and processed for immunocytochemistry.

Immunocytochemistry

Triple-label immunofluorescence for glial fibrillary acidic protein (GFAP), 2',3'-cyclic nucleotide 3'-phosphodiesterase (CNPase) and microtubule associated protein 2 (MAP2) was performed on the Day 7 mouse differentiated cultures. Cells were incubated in the primary antibody against GFAP (rabbit IgG, DAKO; 1:1,000), MAP2 (chicken IgG; Abcam; 1:1,000) and CNPase (mouse IgG; Sigma; 1:1,000) overnight at 4°C in 10% normal donkey serum and 0.1% Triton-X in PBS, followed by incubation with the appropriate secondary antibodies. DAPI counterstaining (1:10,000) was used to detect cell nuclei.

Image acquisition and quantitative analysis

Images were randomly taken from 6-8 visual fields per well using an inverted LRI-Olympus IX-73 epifluorescence microscope equipped with an OCRA-R2 camera, with the Olympus cellSens Dimension program. Images were then imported and counted using the modules *cell scoring* and *neurite outgrowth* of the MetaMorph software (Version 7.8.6.0, Molecular Devices), where the total number of GFAP, CNPase, MAP2 was quantified out of DAPI positively-stained cells, and as described by others (12, 13). The same parameters were used on all the images with the exception of that for *intensity above background level*, which was minimally adjusted between the cultures from different animals. The *intensity above background level* was not changed for all the images within the same condition. The whole analysis was carried out by an investigator blinded to the experimental groupings.

Statistical analysis

Data are expressed as fold change to the DMSO-treated group. Statistical analysis was performed using GraphPad Prism 7.0. For each measurement, mean fold change values for each genotype were compared using a one-way ANOVA with Dunnett's multiple comparisons test and a possible interaction genotype-treatment was determined using a two-way ANOVA evaluation with uncorrected Fisher's LSD to determine changes between genotype for each treatment. Statistical significance was accepted at p<0.05.



Striatum tgHD HOM (+/+) vs wild type (-/-)



Fig. S1. (A) Receptor autoradiography data. Scintillation measurements of receptor autoradiography from P10 WT vs. tgHD pups show significant decreases in D1R, D2R and NMDAR expression, but no changes in A1, A2A and 5-HT2 receptor expression (*p<.05, **p<01). (B) Densitometry analysis for NMDAR expression. Western Blot densitometry reveals significant reduction in NR1, NR2A and NR2B subunits in P10 tgHD pups over WT littermates, as well as decreased phosphorylation of NR2B (*p<.05, ***p<0.001). (C) Diffusiontensor MRI imaging of tgHD pups. A single axial slice of a tgHD HOM (upper row) and WT animal (bottom) is shown (P15): (1) Fractional anisotropy (FA) map, (2) mean diffusivity (MD) map, (3) radial diffusivity (RD), (4) axial diffusivity (AD) map and (5) color encoded FEFA map. The anisotropy map discriminates white (bright regions, anisotropic) and grey matter (dark regions, isotropic). The FEFA map provides orientation of the anisotropy, which indicates the main direction of the underlying fibers (blue, caudal-rostral; red, left-right; and green, dorsal ventral). These different DT maps provide an image contrast, which facilitates the delineation of the Caudate Putamen. Different diffusion parameters were measured in the Caudate Putamen in WT versus tgHD HOM pups. Data are presented as mean and standard deviation. No statistical significance was found between both groups (AD (p = 0.563); MD (p = 0.232); RD (p = 0.121); FA (p = 0.130).



Fig. S2. Additional LBH589 treatment data. (A) LBH589 treatment significantly decreased average velocity and track length in transgenic rats compared to corresponding vehicle treated transgenic animals (average velocity: 0.001 mg/kg: p=0.0194; 0.01 mg/kg: p=0.0119; 0.1 mg/kg: p < 0.0001; track length: 0.001 mg/kg: p = 0.0213; 0.01 mg/kg: p = 0.0152; 0.1 mg/kg: p < 0.0001). LBH589 had no effect on WT animals except for the 0.01mg/kg group, where average velocity and track length were significantly decreased compared to vehicle treated animals (average velocity: 0.01 mg/kg: p=0.04; track length: 0.01 mg/kg: p=0.0389). (B) Single dose LBH589 treatment (i.p.) increased histone H3 (K9) acetylation in the brains of rat pups at postnatal day 10. Quantitative analysis of western blots from n=3 animals, normalized to loading control (*p<0.05, one-way ANOVA). (C) Treatment of WT-derived, differentiating neural stem/progenitor cell cultures with increasing concentrations of LBH589 (0-25 nM) did not alter their neuronal differentiation potential significantly. Of the different LBH589 concentrations tested, the greatest induction of neuronal differentiation was observed at 10 nM (p=0.0090). (D) Schematic representation of the neurosphere differentiation procedure and lateral ganglionic eminence (LGE) dissection from E 13.5 BACHD embryos. (E) Percentage of MAP2+ neurons were significantly increased with 1nM and 10nM LBH589 in WT cultures. BACHD cultures showed increased percentage of MAP2+ neurons only with 10nM LBH589 treatment. 100nM LBH589 treatment resulted in reduced percentage of MAP2 neurons in BACHD cultures. Representative photomicrograph showing MAP2+ neurons (green) and DAPI (blue) from WT and BACHD cultures at 0 and 10 nM LBH589 treatment. (F) Percentage of CNPase+ oligodendrocytes were significantly increased with 1 and 10nM LBH589 in both WT and BACHD derived cultures. Representative photomicrograph showing CNPase+ oligodendrocytes (red) and DAPI (blue) from WT and BACHD cultures at 0 and 10nM LBH589 treatment. 100nM LBH589 resulted in significantly reduced CNPase+ oligodendrocytes. (G) No change in the percentage of GFAP+ astrocytes with 1, 10 and 100nM LBH589 treatment in both WT and BACHD cultures. Representative photomicrograph showing GFAP+ astrocytes (turquoise) and DAPI (blue) from WT and BACHD cultures at 0 and 10nM LBH589 treatment. Data represent means ± SEM. Significant effects are indicated by asterisk (*p<.05, **p<.01; ***p<.001; ***p<.0001 vs NT). Scale bar corresponds to 100 µm in all images.



Fig. S3. Quantitative real-time PCR of HD striata. qPCR validation of transcripts SLC6A3, ATP2A2, TH, FOXG1, AGT, KCNC1, and HCRTR2 in striatal preparations of P10 tgHD pups. Significant effects are indicated by asterisk (*p<.05, **p<.01; vs HOM, ANOVA, n>6).



Fig. S4. Molecular alterations in early postnatal HD pups. Light green arrows indicate positive actions, while light red arrows indicate inhibitory actions. Strong red arrows indicate proteins downregulated in P10 HD pups, and strong green arrows show proteins upregulated in these animals. Black boxes indicate transcripts or proteins validated in this study.



Fig. S5. Additional transcriptional changes after LBH589 treatment. tgHD pups were treated with vehicle (V) or different doses of LBH589 at postnatal day 8, and brain tissue was harvested at postnatal day 10. Gene expression was profiled in striatal preparations by qPCR. Numbers in bars represent number of samples for each genotype/treatment group; * represents genotype differences (compared to WT; * p<0.05; ** p<0.01; *** p<0.001); # represents treatment differences (compared to V; # p<0.05; ## p<0.01; ### p<0.001; two-way ANOVA).

Additional dataset S1 (separate file)

Microarray data of P10 WT vs tgHD rats (striatum and whole brain). The Excel spreadsheet provides descriptors, gene names and symbols, as well as p values for expression profiles of whole brain and striatal tissue of P10 WT vs. tgHD pups.

Additional dataset S2 (separate file)

Proteomic analysis of P10 striata. The Excel spreadsheet provides GI numbers, protein names, gene names, SwissProt IDs, p-values and ratios (downregulated in green, upregulated in red) for protein samples differently expressed between P10 tgHD and WT pups.

Additional dataset S3 (separate file)

PK/PD data for LBH589. The Excel spreadsheet provides dosing regimens, blood and brain levels for LBH589 in tgHD rat pups, as well as the brain/blood ratio, and mean concentrations in both tissues.

Additional data table S1

Primers and annealing temperatures used in this study.

s – sense, as - antisense

Gene	Primer sequence (5'-3')	Annealing [°C]
GAPDH_s	GATGACATCAAGAAGGTGGTGA	58
GAPDH_as	ACCAGGAAATGAGCTTCACAAT	58
AGT_s	ACACCCCTGCTACAGTCCAC	56
AGT_as	ACCCCCTCTAGTGGCAAGTT	56
ATP2A2_s	TGGAGAACGCTCACACAAAG	57
ATP2A2_as	TGCTCGATCACAAGTTCCAG	57
Cort_s	TCTGACTTTCCTTGCCTGGT	59
Cort_as	TACTTGCACGAGGAGAAGG	59
FoxG1B_s	GAACGGCAAGTACGAGAAGC	57
FoxG1B as	TCACGAAGCACTTGTTGAGG	57
GABArb3_s	CTTGACAATCGAGTGGCTGA	56
GABArb3 as	ATTTCCAGGGTGCAGTTTTG	56
GCH1 s	GGGAAGGGTCCATATTGGTT	59
GCH1 as	ACCTCGCATGACCATACACA	59
JunB_s	ATGTGCACGAAAATGGAACA	60
JunB as	CCTGACCCGAAAAGTAGCTG	60
HCRTR2 s	GCCTTTTCTTGCTGTCTTGG	58
HCRTR2 as	GTGAAGATGGCACTCCCTGT	58
HPRT s	GCAGACTTTGCTTTCCTTGG	56
HPRTas	CCGCTGTCTTTTAGGCTTTG	56
KCNC1 s	CCTGGTCTCCATCACAACCT	58
KCNC as	ACTCCACCTTGTTGGGACAG	58
Nrg1 s	GGCACATCCATCCAAATACC	58
Nrg1_as	ACTTGGTGCAGTTTCCTGCT	58
OPRL1 s	TGAGCGTAGACCGCTATGTG	55
OPRL1_as	CATGATGGCAACAGGAACAC	55
$Slc18A\overline{2}$ s	CCACTCTGCTTTTTCCTTCG	59
Slc18A2 as	GCGACGTTAGAGGGTCTCAG	59

Slc6A3 s	CGGAGCACTGGCTTTAGTTC	58
Slc6A3 as	ACAGGCCTGCATTTAACACC	58
TH_s	TGTGTCCGAGAGCTTCAATG	57
TH_as	GGGCTGTCCAGTACGTCAAT	57
D1A_s	CATTCTGAACCTCTGCGTGA	56
D1A_as	GTTGTCATCCTCGGTGTCCT	56
PKA_s	AGCAGGAGAGCGTGAAAGAG	58
PKA_as	GCTTCAGCTTCACCACCTTC	58
$CDK\overline{5}$ s	GCAATGATGTGGATGACCAG	56
CDK5_as	CACGACATTCACCAAGGATG	56
DARPP32 s	ACCACCCAAAGTCGAAGAGA	56
DARPP32 as	GAGGCCTGGTTCTCACTCAG	56
Ppp3ca_s	TACACGGTGGTTTGTCTCCA	57
Ppp3ca as	CAACCCCTGACTGTGTTGTG	57
Ppp3cb s	ACAGGGATGTTGCCTAGTGG	56
Ppp3cb as	TCAGTGGTATGTGCGGTGTT	56
Ppp2cb s	TGGTGGAAAATCACCAGACA	56
Ppp2cb as	GGCGTTCCCATACTTTCGTA	56
COMTs	GAGCTGGGAGCTTACTGTGG	58
COMTas	GTAGGCCTGCAAAGTTCAGC	58
FOXP1 s	CGGCCTTTCTAAAACATCTCAAC	63
FOXP1 as	TGTACTCTACATTGAGCTGTGCTTCTATC	63
FOXP2 s	TATGGAGCAGCCCTTAATGC	63
FOXP2 as	GGTTACTTAGCAAAGGCAAACTG	63
RASGRP1 s	GATGAACTGCCACAAACAGTGC	64
RASGRP1 as	TGCATGGAGCAGATCTTTGG	64
RasGRP2 s	CAAGATCCTGTTCCAGGACTATCAC	58
RasGRP2 as	ACCCACTGAGAGACACTGTTGAAG	58
HSF1 s	AGCTTCCACGTGTTTGACCAG	65
HSF1 as	AAGCTGGCCATGTTGTTGTG	65
CHI3L1 s	CTGTGCACCCATATCATCTACAGC	66
CHI3L1 as	AGTGTCTTCAGTCTGGGGGTTTCTG	66
MAPT s	GCAAGGGGAAGTCTTGAAGTGTAG	66
MAPTas	TAGCCTTATGGCACCTCTTCTCTC	66
NEFL s	TCGAGCATTCCCAGCCTACTATAC	66
NEFLas	TCTTTCTCCTTCTCCTCCTCTTCTG	66
NRGN s	TCCCGCTCTCCTTTGTTTATGC	66
NRGNas	AAAAACCTTCCAGCCACAACCC	66
NPY s	TCGTGTGTTTTGGGCATTCTG	63
NPY as	TTGATGTAGTGTCGCAGAGC	63
HCRTR1 s	TGGTGGTTCTGCTGGTTTTTGC	67
HCRTR1 as	TGGCGAAACATCCCAAACACTC	67
PPARGCIA s	ACACCGCACACATCGCAATTC	66
PPARGC1A as	TTGCTTTCTGCTTCTGCCTCTC	66

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