Selective Roles of Normal and Mutant Huntingtin in Neural Induction and Early Neurogenesis

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Abstract

Huntington’s disease (HD) is a neurodegenerative disorder caused by abnormal polyglutamine expansion in the amino-terminal end of the huntingtin protein (Htt) and characterized by progressive striatal and cortical pathology. Previous reports have shown that Htt is essential for embryogenesis, and a recent study by our group revealed that the pathogenic form of Htt (mHtt) causes impairments in multiple stages of striatal development. In this study, we have examined whether HD-associated striatal developmental deficits are reflective of earlier maturational alterations occurring at the time of neurulation by assessing differential roles of Htt and mHtt during neural induction and early neurogenesis using an in vitro mouse embryonic stem cell (ESC) clonal assay system. We demonstrated that the loss of Htt in ESCs (KO ESCs) severely disrupts the specification of primitive and definitive neural stem cells (pNSCs, dNSCs, respectively) during the process of neural induction. In addition, clonally derived KO pNSCs and dNSCs displayed impaired proliferative potential, enhanced cell death and altered multi-lineage potential. Conversely, as observed in HD knock-in ESCs (Q111 ESCs), mHtt enhanced the number and size of pNSC clones, which exhibited enhanced proliferative potential and precocious neuronal differentiation. The transition from Q111 pNSCs to fibroblast growth factor 2 (FGF2)-responsive dNSCs was marked by potentiation in the number and size of dNSC clones, whereas their multi-lineage potential was unaltered. These abnormalities in neural induction were associated with differential alterations in the expression profiles of Notch, Hes1 and Hes5. These cumulative observations indicate that Htt is required for multiple stages of neural induction, whereas mHtt enhances this process and promotes precocious neurogenesis and oligodendrocyte progenitor cell elaboration.

Introduction

HD is a neurodegenerative disorder caused by abnormal polyglutamine expansion in the amino-terminal end of huntingtin protein (Htt) and characterized by preferential striatal and cortical cellular dysfunction and death associated with late-onset neuropsychiatric and motor disabilities [1]. The molecular basis underlying the selective cellular vulnerability in HD and HD pathogenesis in general remains largely elusive. Htt is a large cytosolic protein of ~340 kDa with ubiquitous cellular localization and adult functional pleiotropism, involving cellular processes ranging from transcriptional regulation to cell survival, whereas mutant Htt (mHtt) causes selective and progressive striatal and cortical neuronal dysfunction and subsequent cell death [2–6]. These observations suggest that Htt may mediate a distinct, selective and previously uncharacterized set of developmental functions, and the pathogenic mutation may therefore have the potential to deregulate these maturational processes and predispose to neurodegeneration. Identifying and characterizing this potential developmental diathesis may have important implications for defining an earlier HD pathogenic window, for explaining the occurrence of a protracted prodromal phase of the disease and for developing novel disease modifying therapies. An increasing number of reports have begun to implicate Htt in seminal early neural developmental processes. For example, conditional deletion of the huntingtin gene (Htt) in the whole brain (Hdh<sup>lox<sup>+</sup>/?;Camk2α<sup>Ccre<sup>+</sup></sup>) from as early as embryonic day 14.5 (E14.5) causes widespread neurodegeneration that mimics several HD phenotypes [7]. Hypomorphic expression of Htt (Hdh<sup>+/−Q50</sup> homozygotes) leads to severe malformation of fore- and mid-brain

Citation: Nguyen GD, Gokhan S, Molero AE, Mehler MF (2013) Selective Roles of Normal and Mutant Huntingtin in Neural Induction and Early Neurogenesis. PLoS ONE 8(5): e64368. doi:10.1371/journal.pone.0064368

Editor: Domingos Henrique, Instituto de Medicina Molecular, Portugal

Received February 26, 2013; Accepted April 12, 2013; Published May 14, 2013

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Funding: M.F.M. is supported by grants from the National Institutes of Health (NS071571, HD071593, MH66290), as well as by the F.M. Kirby, Alpern Family, Ruth L. and David S. Gottesman Institute for Stem Cell and Regenerative Medicine Molecular, Portugal

Competing Interests: The authors have declared that no competing interests exist.

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regions, whereas the complete ablation of Htt results in embryonic lethality as early as E6.5 with a range of severe neuronal developmental defects, including impairments in the formation of the neural plate and absence of head-folds [9–12]. KO embryos also display a shortened primitive streak and absence of the embryonic organizer, which are essential for neural development [13]. These cumulative observations suggest that Htt has additional roles in earlier stages of neural induction and early neurogenesis. In line with these observations, a spectrum of neural developmental deficits have recently been demonstrated by our group in a HD knock-in mouse model (Q111) as early as E13.5, including impairments in striatal neural stem cell (NSC) maintenance, and NSC-mediated medium spiny neuron (MSN) specification and maturation [13]. Other reports of aberrant profiles of adult neurogenesis, including enhanced NSC proliferation, have also been documented in HD models and human pathological specimens [14–16]. Another recent study reported significantly reduced neuronal differentiation in HD knock-in embryonic stem cell (ESC)-derived NSCs with enhanced cell death [17]. In concert with these overall findings, there is increasing evidence of abnormalities of brain morphology, alterations in synaptic and neural plasticity, the presence of subtle neuropsychological impairments and other HD-associated manifestations occurring long before the advent of overt clinical deficits in HD patients and mouse models [18–27].

In the present study, we examined the roles of Htt during neural induction and early neurogenesis, and also assessed whether the HD pathogenic mutation (mHtt) may affect the integrity of these essential developmental processes. To accomplish this goal, we utilized an established ESC neural induction culture model that recapitulates the progressive stages of neural induction and early neurogenesis occurring in vivo.

Results

Htt is required for the specification, self-renewal and proliferation of LIF-responsive pNSCs, whereas mHtt enhances these processes and promotes precocious neurogenesis

Utilizing the ESC paradigm for neural induction, pNSCs can be identified with a colony-forming assay in the presence of leukemia inhibitory factor (LIF) in which they form clonally derived primitive neurospheres (pNSs) [28]. The pNSs express nestin and proneural genes while suppressing expression of the ESC marker, SSEA1 and alternate endodermal/mesodermal lineage genes [29]. To determine whether neural induction can normally occur in the absence of Htt, we compared ESC-derived pNSCs generated from Hdh^ex157/Hdh^ex4/5 ESCs, hereafter referred to as KO ESCs, and control ESCs, hereafter referred to as CTL ESCs [10]. Both the size and number of the KO pNSs were significantly smaller than the CTL pNSs (size: 1.5×10^6 vs 3.0×10^6 μm^3, respectively; p=0.0002; number: 9 vs 44 respectively, p-value<0.0001; Fig. 1A). In addition, KO pNSs were composed of significantly fewer KI67+ and phosphorylated histone H3 (pHisH3)+ cells, markers for dividing cells and G2/M-phase cells, respectively, as compared to CTL pNSs (34.4 vs 50.4%; 13.1% vs 22.7%, p-values<0.0001, respectively; Fig. 1C and D). Moreover, the percentage of dying cells, defined by TUNEL expression, was significantly higher in the KO pNSs (27.0 vs 9.4%, p-value<0.0001, respectively; Fig. 1C and Fig. S1A). Lineage analysis of the KO pNSs as compared to CTL pNSs also revealed persistent expression of SSEA1 and reduced expression of nestin, marker for ESCs and NSCs, respectively, thereby suggesting that the loss of Htt resulted in impairment of the ESC transition to

Figure 1. Htt is required for the elaboration of LIF-responsive pNSs, whereas mHtt differentially deregulates this process. (A, B) Quantification of the size and number of KO and Q111 pNSs as compared to CTL and Q18 pNSs, respectively. Error bars represent ±SEM; unless otherwise stated, *p-value<0.05. (C, D) Quantification of the percentage of positive cells for the proliferation markers, KI67 and pHisH3, and for the cell death marker, TUNEL, in KO pNSs as compared to CTL pNSs, and in Q111 pNSs as compared to Q18 pNSs, respectively. (E) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO and Q111 pNSs. (F, G) Quantification of the percentage of positive cells for the ESC marker, SSEA1, and the NSC marker, Nestin, in KO pNSs as compared to CTL pNSs, and in Q111 pNSs as compared to Q18 pNSs, respectively. (H) Immunofluorescence micrographs of SSEA1 and Nestin immunoreactive cells in CTL, KO, Q18 and Q111 ESCs. (I) Quantification of pNSs expressing the early neuronal marker, β-Tubill, from CTL, KO, Q18 and Q111 ESCs. (J) Immunofluorescence micrographs of β-Tubill immunoreactive in CTL, KO, Q18 and Q111 pNSs. Error bars represent ±95% CI; unless otherwise stated, *p-value<0.05. All scale bars = 25 μm. doi:10.1371/journal.pone.0064368.g001

pNSCs (SSEA1: 55.4 vs 41.9%; Nestin: 20.5 vs 50.9% for CTL and KO respectively, all p-values<0.0001; Fig. 1F and H).

However, when we compared Q111 ESCs, which has an expansion of 111 CAG repeats [30], with control Q18 ESCs, which express 18 CAG repeats, we observed an aberrant enhancement in the specification of pNSCs. Both the size and the number of Q111 pNSs were significantly increased as compared to Q18 pNSs (size: 2.7×10^6 vs 2.1×10^6 μm^3; number: 62 vs 28, respectively, all p-values<0.0001, respectively; Fig. 1B). In addition, there was an increase in the percentage of
proliferating cells that were Ki67 and pHisH3 positive, while no differences were observed in the percentage of TUNEL positive cells in Q111 pNSCs as compared to Q18 pNSCs (Ki67: 48.2% vs 49.5%; p-value =<0.0001; pHisH3: 24.8 vs 24.7%, p-value = 0.127; TUNEL: 13.7 vs 12.4%, p-values = 0.0912, respectively; Fig. 1D and E; Fig. S1A). Moreover, compared to Q18 pNSCs, there was a significant increase in the percentage of nestin+ cells and a concomitant reduction in SSEA1+ cells in Q111 pNSCs (SSEA1: 33.7 vs 43.7%; nestin: 33.9 vs 36.1%, respectively, all p-values=<0.0001; Fig. 1G and H). It has previously been shown that only about 1% of all pNSCs express β-TubIII; however, about 20% of all Q111 pNSs displayed expression of β-TubIII, which indicates the presence of precocious neurogenesis in the presence of mHtt (Fig. 1I and J). These overall observations suggest that Htt is required for the incipient program of neural induction as well as for the self-renewal, proliferation and survival capacity of LIF-responsive pNSCs. Furthermore, the mutation in Htt enhances ESC-derived neural induction and leads to precocious neuronal lineage specification.

Htt is required for the specification, self-renewal and proliferation of FGF2- and EGF-responsive dNSCs, whereas mHtt differentially alters these processes and further promotes precocious neurogenesis in FGF2-responsive dNSCs

We next examined the role of Htt in the program of neural specification of dNSCs and assessed whether the presence of mHtt alters this developmental program. Individual LIF-responsive pNSCs were dissociated and re-propagated in the presence of FGF2 to generate FGF2-responsive dNSCs [29,31]. These dNSCs are equivalent to their in vivo counterparts that exist from E8.5 through adulthood. Analogous to the developmental profiles observed with KO pNScs, both the size and number of the KO FGF2-responsive dNSCs were significantly decreased as compared to CTL FGF2-responsive dNSs (size: 3.0 × 10^4 vs 5.7 × 10^4 μm^3; number: 4 vs 29, respectively, all p-values=<0.0001, respectively; Fig. 2A). Consistently, KO FGF2-responsive dNSCs were composed of significantly fewer numbers of Ki67+ and pHisH3+ cells than CTL FGF2-responsive dNSCs, whereas the percentage of TUNEL+ cells remained significantly higher (Ki67: 22.7 vs 46.2%; 9.1 vs 17.3%; TUNEL: 26.7 vs 11.7%, respectively, all p-values=<0.0001; Fig. 2C and E; Fig. S1B). Immunofluorescence lineage analysis of KO FGF2-responsive dNSCs also revealed a significantly lower percentage of nestin+ NSCs and β-TubIII+ neuronal precursors with a higher percentage of SSEA1+ ESCs as compared to CTL FGF2-responsive dNSs (SSEA1: 81.1 vs 1.0%; Nestin: 8.9 vs 37.9%; β-TubIII: 10.7 vs 19.3%, respectively, all p-values=<0.0001; Fig. 2F, H and I). In contrast to the findings with the KO pNSCs, the presence of mHtt resulted in significantly higher numbers of Q111 FGF2-responsive dNSCs than those of Q18 FGF2-responsive dNSCs even though there was no difference in their respective sizes (number: 27.9 vs 11.9, respectively, p-value=<0.0001; size: 3.7 × 10^4 vs 3.9 × 10^4 μm^3, respectively, p-value = 0.0321; Fig. 2B). In addition, as compared to Q18 FGF2-responsive dNSs, there was an increase in the percentage of Ki67+ and pHisH3+cells in Q111 FGF2-responsive dNSs, whereas the percentage of TUNEL+ cells was unchanged (Ki67: 45.7 vs 33.3%, p-value =<0.0001; pHisH3: 16.8 vs 14.0%, p = 0.0019; TUNEL: 12.3 vs 11.7%, p-value = 0.5760, respectively; Fig. 2D and F; Fig. S1B). Further lineage analysis revealed that Q111 FGF2-responsive dNSs also displayed significantly higher proportions of nestin+ NSCs and β-TubIII+ neuronal precursors as compared to the Q18 FGF2-responsive dNSs (nestin: 65.4 vs 56.7%; β-TubIII: 49.3 vs 31.6%, relatively, all p-values=<0.0001; Fig. 2G-J). The increase in the percentage of β-TubIII+ cells suggests an enhanced specification of committed neuronal progenitors. These cumulative observations suggest that Htt is required for the transition of LIF-responsive pNSCs to FGF2-responsive dNSCs and for the promotion of self-renewal, proliferation and neuronal lineage fate of FGF2-responsive dNSCs. Conversely, mHtt enhanced the transition from LIF-responsive pNSCs to FGF2-responsive dNSCs with alterations in proliferative potential and precocious neurogenesis.

FGF2-responsive dNSCs are the direct precursors of EGF-responsive dNSCs [31]. To further investigate the role of Htt and the effects of mHtt in the specification of EGF-responsive dNSCs from FGF2-responsive dNSCs, we dissociated and re-propagated FGF2-responsive dNSs in EGF to generate EGF-responsive dNSNs. Both the number and size of the KO EGF-responsive dNSNs were significantly decreased as compared with CTL EGF-responsive dNSs (size: 0.1 × 10^4 vs 3.5 × 10^4 μm^2; number: 0 vs 9, respectively, all p-values=<0.0001; Fig. S1C). Upon differentiation after 7 days in vitro (DIV), few irregularly shaped EGF-responsive dNSNs were formed from KO ESCs and these clones failed to differentiate into neurons and glia (Fig. S1E). In contrast, the size of Q111 EGF-responsive dNSNs was comparable to Q18 EGF-responsive dNSNs, whereas the number of Q111 EGF-responsive dNSNs was significantly decreased as compared to those of Q18 EGF-responsive dNSNs (2.0 × 10^4 vs 1.8 × 10^4 μm^2, p-values = 0.5550; 1 vs 2; p-values=<0.0001, respectively, Fig. S1D). Nonetheless, the elaboration of β-TubIII+ neuronal species and GFAP+ astrocytes in Q111 EGF-responsive dNSNs were comparable to those in the Q18 EGF-responsive dNSNs under differentiating conditions. (Fig. S1E). These observations indicate that Htt is required for the developmental transition of FGF2-responsive dNSCs to EGF-responsive dNSNs and the subsequent differentiation into neurons and glia, whereas mHtt selectively impairs the specification of EGF-responsive dNSNs but does not alter their neural lineage potential.

Htt is required for the expression of ectodermal and neural genes, and the repression of genes specifying alternate endodermal cell fate in LIF-responsive pNSCs, whereas mHtt selectively enhances ectodermal and neuronal gene expression

To investigate the roles of Htt and mHtt in mediating lineage potential during the process of early neural induction, we assessed expression levels of genes involved in promoting neural and non-neural lineage decisions from pNSCs, and further examined NSC maintenance and lineage potential under differentiating conditions. KO pNSCs exhibited significant downregulation in the expression level of the primitive ectoderm gene, FGF3 (0.333, p-value = 0.001), the proneural genes, Ngn2 and Mash1 (0.31 and 0.035, respectively; p<0.001), the neurogenic gene, NeuroD1 (0.207, p-value =<0.001) and the early patterning and gliogenic gene, Nkx2.2 (0.329, p-value = 0.012), as compared to CTL pNSCs (Fig. 3A). Additionally, contrary to previous studies reporting that wild type pNSCs do not express any endodermal or mesodermal genes [28], the expression of the endodermal gene, Gata4, was significantly increased in KO pNSCs as compared to CTL pNSCs (7.387, p-value =<0.001), thereby suggesting Htt modulates the repression of endodermal lineages during neural induction in ESCs (Fig. 3C). Furthermore, immunofluorescence lineage analysis after 7DIV under differentiating conditions revealed that in contrast to CTL pNSCs, KO pNSCs retained SSEA1 expression and failed to express nestin and the neuronal precursor marker, doublecortin (DCX) (Fig. 3E). Conversely, gene expression
analysis in Q111 pNSs revealed a significant upregulation in the expression levels of the primitive ectodermal gene, FGF5 (2.989, \(p\)-value = 0.05). (C, D) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs, respectively. (E) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. (F, G) Quantification of the percentage of total positive cells for the primitive ectodermal gene, Nestin-immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs, respectively. (H) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. (I) Quantification of the percentage of positive cells for proliferation markers, KI67 and pHisH3, and for the cell death marker, TUNEL, in KO FGF2-responsive dNSs as compared to CTL FGF2-responsive dNSs, and in Q111 FGF2-responsive dNSs as compared to Q18 FGF2-responsive dNSs, respectively. (J) Quantification of the percentage of positive cells for the ESC marker, SSEA1, and the NSC marker, Nestin, in KO FGF2-responsive dNSs as compared to CTL FGF2-responsive dNSs, and in Q111 FGF2-responsive dNSs as compared to Q18 FGF2-responsive dNSs, respectively. (K) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. Error bars represent ± SEM; unless otherwise stated, *\(p\)-value < 0.05. All scale bars = 25 \(\mu\)m. doi:10.1371/journal.pone.0064368.g002

Figure 2. Htt is required for the elaboration of FGF2-responsive dNSs, whereas mHtt differentially deregulates this process. (A, B) Quantification of the size and number of KO and Q111 pNSs as compared to CTL and Q18 FGF2-responsive dNSs, respectively. Error bars represent ± SEM; unless otherwise stated, *\(p\)-value < 0.05. (C, D) Quantification of the percentage of positive cells for proliferation markers, KI67 and pHisH3, and for the cell death marker, TUNEL, in KO FGF2-responsive dNSs as compared to CTL FGF2-responsive dNSs, and in Q111 FGF2-responsive dNSs as compared to Q18 FGF2-responsive dNSs, respectively. (E) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. (F, G) Quantification of the percentage of positive cells for the ESC marker, SSEA1, and the NSC marker, Nestin, in KO FGF2-responsive dNSs as compared to CTL FGF2-responsive dNSs, and in Q111 FGF2-responsive dNSs as compared to Q18 FGF2-responsive dNSs, respectively. (H) Immunofluorescence micrographs of KI67 and pHisH3 immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. (I) Quantification of the percentage of total positive cells for the early neuronal marker, b-TubIII, in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. (J) Immunofluorescence micrographs of b-TubIII-immunoreactive cells in CTL, KO, Q18 and Q111 FGF2-responsive dNSs. Error bars represent ±95% CI; unless otherwise stated, *\(p\)-value < 0.05. All scale bars = 25 \(\mu\)m.
addition, as compared to Q18, Q111 FGF2-responsive dNSs exhibited a modest increase in the proportion of O4+ OL progenitor species (0.07% vs 0.3%, respectively, p-value<0.0001), potentially due to the increased expression of Nkx2.2 as observed previously (Fig. 4F). There was no significant difference in the proportion of unipotent and bipotent clones between Q18 and Q111 FGF2-responsive dNSs. Interestingly, as compared to Q18 FGF2-responsive dNSs, Q111 dNSs exhibited a large proportion of multipotent clones (0% vs. 35.3%, respectively). These experimental findings indicate that Htt is required for the expression of ectodermal and pro-neural genes and for the repression of genes associated with mesodermal fate during the transition from LIF-responsive pNSCs to FGF2-responsive dNSCs. This is associated with corresponding reductions in the elaboration of neuronal and glia lineages, as well as significant but selective reductions in unipotent neuronal and bipotent neuronal-astrocyte clonal potential. By contrast, mHtt enhances ectodermal and selective pro-neural gene expression, and enhances the elaboration of neuronal and glial lineages with a selective increase in multilineage potential.

Htt is associated with Notch signaling pathways during the specification and maintenance of pNSCs and dNSCs, whereas mHtt differentially deregulates this developmental signaling cascade.

Notch signaling pathways play pivotal roles in cell fate diversification during development. In particular, the Notch/Hes pathway is essential for the transition of pNSCs to dNSCs and NSC proliferation and maintenance [32,33]. To determine whether Htt is required for the integrity of Notch signaling during the process of neural induction, we analyzed the expression...
profiles of Notch, Hes1 and Hes5 in both KO pNSCs and dNSCs. Gene expression analysis showed that both Notch (RQ = 0.603, p-value < 0.001) and Hes5 (RQ = 0.062, p-value < 0.001) expression were significantly downregulated in KO pNSCs, whereas the expression of Hes1 (RQ = 2.119, p-value < 0.001) was significantly upregulated as compared to control CTL pNSCs (Fig. 5A). However, the expression levels of Notch (RQ = 0.995, p-value = 0.233) and Hes5 (RQ = 0.923, p-value = 0.097) in KO FGF2-responsive dNSCs were comparable to CTL FGF2-responsive dNSCs, whereas the expression of Hes1 (RQ = 0.609, p-value < 0.001) was significantly downregulated (Fig. 5B). By contrast, for Q111 pNSCs, Notch (RQ = 1.352, p-value < 0.001) and Hes5 (RQ = 1.705, p-value < 0.001) expression levels were significantly upregulated, whereas the level of expression of Hes1 (RQ = 0.296, p-value < 0.001) was significantly downregulated as compared to Q18 pNSCs (Fig. 5C). Although the expression level of Hes1 (RQ = 0.662, p-value = 0.012) remained significantly downregulated as Q111 LIF-responsive pNSCs transitioned to FGF2-responsive dNSCs, Notch expression levels became significantly upregulated (RQ = 3.291, p-value < 0.001; Fig. 5D). These observations suggest that Htt modulates Notch signaling pathways during the specification and maintenance of pNSCs and dNSCs.
and mHtt differentially disrupts Notch/Hes1/Hes5 signaling during both pNSC and dNSC developmental stages.

**Discussion**

In this study, we employed a specialized ESC clonal culture paradigm to characterize the entire program of neural induction and early neurogenesis [28,29], and demonstrated the essential roles of Htt in the program of neural induction, progressive specification of neural progenitor cell types and the subsequent elaboration of neural lineage species. Our study also revealed that the HD pathogenic mutation aberrantly enhanced ESC-derived neural fate specification, resulting in precocious neurogenesis of pNSCs, and enhanced the elaboration of neuronal and glial lineages from dNSCs.

The development of the central nervous system (CNS) begins with the early program of neural induction within the anterior region of the epiblast. It has been shown that neural fate specification in the pre-gastrula epiblast exists as a ‘default’ state and FGF signaling from the organizer antagonizes the inhibitory effects of bone morphogenetic proteins (BMPs) on anterior neural fate [34]. However, following gastrulation, the organizer further ‘induces’ the elaboration and patterning of neural tissue by antagonizing other neural inhibitory signals, such as Nodal and FGF signaling from the organizer further impair the survival of the KO pNSCs. Interestingly, these adapter molecules are also part of the FGF receptor-signaling pathway that is essential for the specification and proliferation of pNSCs and their transition to dNSCs [28,37].

The Notch pathway is an active signaling cascade regulating the early program of neural induction. Our study has shown that the absence of Htt disrupts expression of Notch as well as Hes5, an essential Notch effecter, in KO pNSCs. By contrast, the ablation of Notch (Notch−/−) in mouse embryos has been demonstrated to only reduce Hes5 expression, but does not disrupt the generation of pNSCs [33]. This strongly indicates that the requirement of Htt for the specification of pNSCs is independent from its putative role in modulating Notch signaling pathways. Alternatively, high Hes1 in KO pNSCs can suppress proliferation as it has been shown that high Hes1 levels in neural progenitors can repress cyclin D1 and result in G1 phase retardation [38]. Additionally, and consistent with our observations, high Hes1 expression levels have also been shown to promote preferential mesodermal differentiation over neural differentiation, possibly via repression of Notch signaling [39,40]. Interestingly, Notch−/− embryonic brains as well as Notch−/− ESCs are severely impaired in the generation of FGF2- and EGF-responsive dNSCs [33]. Similarly, knockdown of RBP:Jk, a downstream mediator of Notch signaling, has also been shown to deplete early dNSCs [32]. Thus, the reduced Notch/Hes5 expression we observed in KO pNSCs may have resulted in impairment in their transition to dNSCs. However, some KO pNSCs were capable of undergoing early neural developmental transition to form FGF2-responsive dNSCs, which also displayed comparable levels of expression of Notch and Hes5 as the controls. As appropriate Notch expression levels are important to direct ESC differentiation to neuronal lineages [41], the apparent normal levels of expression of Notch and Hes5 in KO FGF2-responsive dNSCs suggest that Htt may not play a role in the regulation of Notch signaling in dNSCs, and that alterations in neuronal and glial

Furthermore, the presence of severe mesodermal impairments in KO embryos prevents definitive assessment of the direct roles of Htt in neural development, as these mesodermal structures play critical inductive roles for neural development [35]. The in vitro clonal ESC neural induction model recapitulates the in vivo program of neural induction that follows the “default” pathway in the absence of confounding extrinsic factors [29,29]. Thus this experimental paradigm provides an important alternative approach that circumvents many of the aforementioned experimental limitations. However, control conditions for the KO and Q111 ESC lines differed in several lineage parameters. These observations are likely due to the fact that the appropriate controls differed by the presence of mouse (R1 ESC) and human (Q18) 5’ sequences within the huntingtin gene, which did not exhibit complete homology. The utilization of separate controls for the KO and Q111 ES cell conditions was necessary because the mutant huntingtin ESC line was constructed with the humanized expansion repeat sequence.

The early stage of LIF-responsive pNSC induction in vitro, however, is a particularly vulnerable developmental phase due to the enhanced sensitivity of apoptosis signaling pathways to caspase-mediated cell death [29,29]. Htt has been shown to display primary anti-apoptotic functions that are mediated, in part, through direct inhibition of activation of caspase 3 and 9, and therefore the absence of Htt may enhance the cellular vulnerability of the KO pNSCs [5]. The presence of LIF has also been demonstrated to have important pro-survival roles in pNSCs, and Htt is known to interact with Grb2 and RasGAP, two adapter molecules of the LIF receptor [36]. Thus, the absence of Htt may disrupt the LIF receptor-mediated pro-survival pathway and further impair the survival of the KO pNSCs. Interestingly, these adapter molecules are also part of the FGF receptor-signaling pathway that is essential for the specification and proliferation of pNSCs and their transition to dNSCs [28,37].

It is also likely that Htt, as an integral component of the cytoskeleton, is responsible for maintaining the integrity of nuclear membrane during neuronal and glial differentiation, which is otherwise compromised in KO pNSCs. Notch signaling in dNSCs is known to regulate cell survival via the LIF receptor-mediated pro-survival pathway [36]. Thus, the absence of Htt may be detrimental to the survival of the KO pNSCs [5]. The presence of LIF has also been demonstrated to have important pro-survival roles in pNSCs, and Htt is known to interact with Grb2 and RasGAP, two adapter molecules of the LIF receptor [36]. Thus, the absence of Htt may disrupt the LIF receptor-mediated pro-survival pathway and further impair the survival of the KO pNSCs. Interestingly, these adapter molecules are also part of the FGF receptor-signaling pathway that is essential for the specification and proliferation of pNSCs and their transition to dNSCs [28,37].

The Notch pathway is an active signaling cascade regulating the early program of neural induction. Our study has shown that the absence of Htt disrupts expression of Notch as well as Hes5, an essential Notch effecter, in KO pNSCs. By contrast, the ablation of Notch (Notch−/−) in mouse embryos has been demonstrated to only reduce Hes5 expression, but does not disrupt the generation of pNSCs [33]. This strongly indicates that the requirement of Htt for the specification of pNSCs is independent from its putative role in modulating Notch signaling pathways. Alternatively, high Hes1 in KO pNSCs can suppress proliferation as it has been shown that high Hes1 levels in neural progenitors can repress cyclin D1 and result in G1 phase retardation [38]. Additionally, and consistent with our observations, high Hes1 expression levels have also been shown to promote preferential mesodermal differentiation over neural differentiation, possibly via repression of Notch signaling [39,40].

Interestingly, Notch−/− embryonic brains as well as Notch−/− ESCs are severely impaired in the generation of FGF2- and EGF-responsive dNSCs [33]. Similarly, knockdown of RBP:Jk, a downstream mediator of Notch signaling, has also been shown to deplete early dNSCs [32]. Thus, the reduced Notch/Hes5 expression we observed in KO pNSCs may have resulted in impairment in their transition to dNSCs. However, some KO pNSCs were capable of undergoing early neural developmental transition to form FGF2-responsive dNSCs, which also displayed comparable levels of expression of Notch and Hes5 as the controls. As appropriate Notch expression levels are important to direct ESC differentiation to neuronal lineages [41], the apparent normal levels of expression of Notch and Hes5 in KO FGF2-responsive dNSCs suggest that Htt may not play a role in the regulation of Notch signaling in dNSCs, and that alterations in neuronal and glial
lineage elaboration may be due, in part, to additional non-redundant developmental signaling pathways. Thus far, Htt has not been shown to have any direct interaction with components of the Notch signaling cascade, with the exception of a single study reporting an indirect functional association between Huntingtin interacting protein 1 (HIP-1) and deltex-dependent Notch signaling in *Drosophila* that plays a role in neurogenesis [42]. Additional studies are required to elucidate the mechanisms underlying this regulatory function.

Htt may also play an important role at the intersection of neural and non-neural fate decisions during the incipient program of neural induction as both KO pNSCs and dNSCs displayed preferential increases in mesodermal and endodermal gene expression over pro-neural gene expression. These specialized roles of Htt in cell fate decisions may be orchestrated by modulating the functions of the neuron-restrictive silencing factor/RE1-silencing transcription factor (NRSF/REST) by normally sequestering it within the cytoplasm [43]. REST is a transcriptional and epigenetic regulator of both neural and non-neural cell fate specification programs [43]. It has previously been demonstrated that overexpression of REST in ESCs can promote early differentiation of ESC-derived embryoid bodies to primitive endoderm and also disrupt specification of the epiblast [44]. Further studies are required to show whether the loss of Htt may enhance aberrant accumulation of REST in the nucleus and contribute to the preferential acquisition of endodermal over ectodermal fates during the program of neural induction.

On the other hand, in Q111 pNSCs the presence of mHtt enhanced *Notch* and *Hes5* expression levels. Interestingly, enhanced FGF receptor signaling in dNSC can also potentiate Notch signaling and enhance neurogenesis [45,46]. Constitutive Notch activation (NotchIC) has been shown to not only upregulate Hes5 expression levels but also more importantly to enhance the generation of dNSCs, which is consistent with our observations in Q111 dNSCs [33]. These dNSCs then progressively become Notch/He5-dependent and undergo asymmetric cell division to modulate the balance between the maintenance of NSC populations and neural lineage commitment [47,48]. Thus, the sustained increase in the expression of *Notch* in Q111 dNSCs may differentially enhance asymmetric cell divisions resulting in premature specification of committed neural progenitors, which is consistent with our observation of enhanced generation of neuronal and glial lineages [47,49–51]. Furthermore, this may also lead to the premature depletion of Q111 FGF2-responsive dNSCs, and thus to deficits in the generation of Q111 EGF-responsive dNSCs. Conversely, *Hes1* expression levels were significantly downregulated in Q111 pNSCs and dNSCs and may have contributed to the preferential expression of both neuroectodermal and neurogenic genes, to enhanced proliferative capacity and to precocious neurogenesis. Indeed, low *Hes1* expression in ESCs has been shown to preferentially enhance neural differentiation, whereas the complete ablation of Hes1 further promoted premature neurogenesis [40,49]. High *Hes1* expression has also been reported to have suppressive effects on the maturation of NG2+ /O4− OL precursors, which is consistent with the enhanced elaboration of O4 + OL progenitors in the low *Hes1*-expressing Q111 dNS culture condition [51]. The latter observation may have important implications for defining the mechanistic underpinnings of previous findings of increased oligodendrocyte density reported in the caudate nucleus in HD patients [52]. Remarkably, Notch signaling, particularly with respect to Notch1/3, has been shown to play pivotal roles in the developmental stage-specific regulation of neural progenitors in the ventricular zone that contribute to striatal development [53]. These findings suggest that mHtt alters Notch signaling cascades during neural induction, and these and related molecular pathways may have important implications for explaining the regional striatal developmental deficits previously reported by our group in the HD knock-in Q111 mouse model [13].

Our findings of significant alterations in proliferative potential, self-renewal as well as neural and non-neural lineage potential in Q111 pNSCs and dNSCs have important implications for HD. First, these cellular alterations may result in impairments in neural lineage specification in neurogenic zones that, in part, is consistent with several reports of enhanced self-renewal and precocious neurogenesis in the subventricular zone (SVZ) of R6/2 and Hdh-Q150 KI HD mouse models [14,54]. Second, highly proliferative Q111 pNSCs undergo enhanced DNA replication and are therefore at increased risk for accelerated DNA damage and repair responses, which have been shown to promote mutational instability of CAG repeats and potentially contribute to the pathogenesis of HD [55,56]. Putative DNA instability may persist in mutant pNSCs and subsequently in their progeny, thereby promoting the propagation of developmental mutation length-mediated cellular and functional impairments into adult life. These pathogenic possibilities are consistent with several reports of increased DNA instability and CAG expansion mosaicism in the brains of HD patients and mouse models [53,57].

A recent study by Conforti and colleagues reported that the loss of Htt and the presence of mHtt (NS-HdhQ111/7 NSCs and NS-HdhQ111/7, respectively) did not disrupt the *in vitro* derivation of ESC-derived NSCs or impair their self-renewal and proliferative properties. In addition, the HdhQ111/7 NSCs were shown to display reduced neurogenesis and increased cell death [17]. The differences observed between these findings and those of the present study may stem from the use of alternate experimental protocols. Importantly, the current study extends our previous published observations in Q111 mice that mHtt deregulates cell cycle parameters of NSCs and results in aberrant expansion of intermediate progenitors in the absence of increased cell death [13]. Furthermore, the previous work from our group [13] and the current findings strongly suggest that HD-associated abnormalities in adult life (reviewed in [50]) may stem from early and cumulative neurodevelopmental impairments, and may therefore support the notion that HD represents a primary neurodevelopmental disorder in addition to a neurodegenerative disease [39]. Equally important is the concept that semial impairments occurring during early stages of the neural developmental program can potentially lead to multiple foci of regional cellular vulnerabilities along the entire neuraxis, observations increasingly shown to be associated with HD and other neurodegenerative disease phenotypes [26,27,60–63].

It is imperative to corroborate the observations in this study with other *in vivo* HD models to better refine our understanding of the potential contributions of the HD pathogenic mutation and of differing numbers of pathogenic expansion repeats during incipient stages of embryonic and neuronal development. Moreover, it is also important to define key molecular impairments occurring along the continuum of developmental and adult stages in affected individuals and in robust HD animal models by identifying potentially unique developmental protein partners of Htt, such as regulators of transcriptional, epigenetic and additional diverse cellular processes. These essential initiatives will open up new possibilities for innovative and efficacious diagnostic, therapeutic and preventative strategies for HD.
Materials and Methods

Embryonic Stem Cell Culture Paradigms

The KO, Q18 and Q111 ESCs (Hdh<sup>+/y</sup>/Hdh<sup>x/y</sup>, Hdh-Q18 and Hdh-Q111, respectively), were previously generated and graciously supplied by MacDonald et al. [10,30]. The R1 ESC line from ATCC was used as the control (CT). ESCs were maintained on mouse embryonic fibroblast (MEF) feeder layers that had previously been inactivated with Mitomycin C (Sigma, M4287). Prior to use for specific experimental protocols, ESCs were plated and maintained on 0.1% gelatin-coated tissue culture plates in ES cell media consisting of knockout Dulbecco’s minimal essential medium (Invitrogen, DMEM, 10313) supplemented with 1000 U/ml of leukemia inhibitory factor (LIF/ESGRO; Chemicon, ESG1106), 10% ES-qualified FBS (ATCC, SCRR-30-2020), 1X MEM nonessential amino acids (from 100x stock, Invitrogen, 11140), 1X L-glutamine and antibiotics (from 100x stock, Invitrogen 10378-016), and 0.1 mM 2-mercaptoethanol (Sigma, M7522).

ESC-derived Primitive and Definitive NSC Assays

These assays were carried out as previously described [28]. Briefly, all culture conditions were carried out in serum-free media that consisted of DMEM/F-12 (Invitrogen, 11330) supplemented with 25 ug/ml insulin (Sigma, I6634), 100ug/ml transferrin (Sigma, T1147), 20 nM progesterone (Sigma, P7556), 60 μM putrescine (Sigma, P5780), 30 nM sodium selenium (Sigma, S5261), 1x L-glutamine and antibiotics (100x stock, Invitrogen 10378-016), 5 mM Hepes (Sigma, H3375) and 3mM NaHCO3 (Sigma, S5761). ESCs were plated as single-cell suspensions at densities <= 10 cells/μl on uncoated culture plates using the above media formulation supplemented with LIF (1000U/ml; Chemicon, ESG1106) for 7 days in vitro (DIV) to form LIF-responsive pNSs. The pNSs were dissociated into single cells by trypsin/0.04% EDTA (Invitrogen, 25300-054) and re-plated in the same media supplemented with 10ug/ml FGF2 (BD; 354060) and 2 μg/ml Heparin (Sigma-Aldrich) for another 7 DIV to form FGF2-responsive dNSs. The FGF2-responsive dNSs were further dissociated and re-plated in the presence of 20ng/ml EGF for an additional 7 DIV to form EGF-responsive dNSs. For differentiation paradigms, individual clonal spheres were plated onto Matrigel-coated plates in same media supplemented with 1% FBS.

Immunofluorescence Analysis

NSs were collected by centrifugation at 300 rpm for 5 minutes, washed once in PBS and fixed in 4% PFA for 20 minutes at room temperature. NSs were then collected in 20% sucrose until they became totally submerged and then frozen in M-1 Embedding Matrix (Thermo) for cryo-sectioning. Immunofluorescence analysis was carried out as previously described [13,64] (See Table S1 for the list of utilized antibodies). TUNEL analysis was performed according to the manufacturer’s protocols (Roche, 11684795910). BrdU analysis was carried out as previously described [13].

Quantitative Real-Time PCR (QPCR)

Harvesting of RNA from samples was carried out using TRI reagents® (Molecular Research Center Inc, Cincinnati, OH, USA) according to manufacturer’s protocol. The quantification of total RNA concentration was determined using the Qubit® RNA assay kit and Qubit® 2.0 Fluorometer (Invitrogen). Single strand cDNA synthesis was performed using the High Capacity RNA Reverse Transcription Kit® (Applied Biosystems, 4368814) following the manufacturer’s recommendations. TaqMan primers were purchased from PE Applied Biosystems and SYBR Green probes were generated using the Invitrogen service (See Table S1). We utilized either TaqMan Universal PCR Master Mix® or SYBR Green Master Mix and ran samples in triplicate in the Model 7000 Real Time PCR system® (Applied Biosystems, CA, USA). The housekeeping gene employed was hypoxanthine guanine phosphoribosyl transferase 1 (HPRT). Data collection and quality assessment were performed utilizing the 7000 SDS 1.1 RQ Software (Applied Biosystems, CA, USA). The analysis was accomplished with the 2-ΔΔC(T) relative quantification method with the Relative Expression Software Tool (REST) developed by Corbett Research [65–66]. Gene expression levels were reported using the relative RQ values with ±95% Confidence Interval (CI).

Statistical Analysis

Statistical comparisons were evaluated according to the type of data analyzed: proportions were compared with Chi-square test or Fisher’s Test. The means of samples were analyzed with either Mann-Whitney U test or t-test. Statistically significant differences between samples were considered using a probability of at least <0.05.

Supporting Information

Figure S1 TUNEL assays and the roles of Htt and mHtt in the elaboration of EGF-responsive dNSs. (A, B) Immunofluorescence micrographs of TUNEL-immunoreactive cells contained within CTL, KO, Q18 and Q111 pNSs and FGF2-responsive dNSs. (C, D) Quantification of the size and number of CTL, KO, Q18 and Q111 EGF-responsive dNSs. (E) EGF-responsive dNSs were cultured under differentiating conditions for 7DIV and analyzed by immunofluorescence microscopy for the expression profiles of the neuronal and astrocyte markers, β-TubIII and GFAP, in CTL, KO, Q18 and Q111 clones. Error bars represent ±95% CI; unless otherwise stated, *p-value<0.05. All scale bars = 25 μm.

Table S1 List of antibodies, TaqMan probes and SYBR Green probes utilized in the study. All antibodies are listed with manufacturers’ names, catalogue numbers, as well as concentration used. All TaqMan probes are listed with catalogue numbers from Applied Biosystems. All SYBR Green probes are listed with forward and reverse sequences.

Acknowledgments

We are grateful for Dr. Marcy MacDonald, MGH/Harvard, for kindly supplying the cell lines utilized in these studies.

Author Contributions

Conceived and designed the experiments: GDN SG AM MFM. Performed the experiments: GDN SG. Analyzed the data: GDN SG AM MFM. Contributed reagents/materials/analysis tools: GDN SG AM MFM. Wrote the paper: GDN SG AM MFM.
References

1. Gil JM, Rengo AC (2008) Mechanisms of neurodegeneration in Huntington's disease. The European journal of neuroscience 27: 2803–2820.

2. Harjes P, Wanker EE (2003) The hunt for huntingtin function: interaction analysis in Xenopus laevis. Trends in bio sciences 28: 425–433.

3. Kaltenbach LS, Romero E, Becklin RR, Chettier R, Bell R, et al. (2007) Huntington interacting proteins are genetic modifiers of neurodegeneration. PLoS genetics 3: e82.

4. Li SH, Li XJ (2004) Huntington-protein interactions and the pathogenesis of Huntington's disease. Trends in genetics 20: 146–154.

5. Rigamonti D, Sipione S, Goffredo D, Zuccato C, Fossale E, et al. (2001) Huntington's neurodegenerative activity occurs via inhibition of proapoptotic processing. The Journal of biological chemistry 276: 14545–14548.

6. Cattaneo E, Zuccato C, Tartari M (2005) Normal huntingtin function: an alternative approach to Huntington's disease. Nature reviews Neuroscience 6: 919–930.

7. Dragatis I, Levine MS, Zeitlin S (2000) Inactivation of Hdh in the brain and tests results in progressive neurodegeneration and sterility in mice. Nature genetics 26: 300–306.

8. Nasir J, Lefebvre B, O’Kusky JR, Diewert VM, Richman JM, et al. (1995) Targeted disruption of the Huntington's disease gene results in embryonic lethality and behavioral and morphological changes in heterozygotes. Cell 81: 811–823.

9. Zeitlin S, Lin JP, Chapman DL, Papisbonne AU, Estradias A (1995) Increased apoptosis and early embryonic lethality in mice nullizygous for the Huntington's disease gene homologue. Nature genetics 11: 153–163.

10. Duyao MP, Auerbach AB, Ryan A, Persichetti F, Barnes GT, et al. (1995) Inactivation of the mouse Huntington's disease gene homolog Hdh. Science 269: 407–410.

11. Woda JM, Calzonetti T, Hilditch-Maguire P, Duyao MP, Conlon RA, et al. (2005) Inactivation of the Huntington's disease gene (Hdh) impairs stress response and early patterning of the mouse embryo. BMC developmental biology 5: 17.

12. White JK, Auerbach W, Duyao MP, Vonasett JP, Gasella JF, et al. (1997) Huntington is required for neurogenesis and is not impaired by the Huntington's disease CAG expansion. Nature genetics 17: 404–410.

13. Molero AE, Gokhan S, Gonzalez S, Feig JL, Alexandre LC, et al. (2009) Impairment of developmental stem cell-mediated striatal neurogenesis and pluripotency genes in a knock-in model of Huntington's disease. Proceedings of the National Academy of Sciences of the United States of America 106: 21900–21905.

14. Rabbits CM, Kippin TE, Willaine-Morawek S, Shimabukuro MK, Akamato W, et al. (2006) A progressive and cell non-autonomous increase in striatal neuronal cell number in the Huntington's disease mouse R6/2 mouse. The Journal of neuroscience: the official journal of the Society for Neuroscience 26: 10452–10460.

15. Curtis MA, Penney EB, Pearson AG, van Roon-Mom WM, Butterworth NJ, et al. (2007) Evidence for more widespread cerebellar pathology in early HD: an MR-based morphometric analysis. Neurology 60:1615–1620.

16. Kusunoki S, Troppe V, Hishida S, Shirai CM, Iwakawa H, et al. (2001) Direct neural cell fate specification from embryonic stem cells: a primitive mammalian neural stem cell stage acquired through a default mechanism. Neuron 30: 65–78.

17. Wilson SI, Edlund T (2001) Neural induction: toward a unifying mechanism. Nat Neurosci 4 Suppl: 1161–1168.

18. Levine AJ, Brivanlou AH (2007) Proposal of a model of mammalian neural induction. Developmental biology 308: 247–256.

19. Tropepe V, Hishida S, Shirai CM, Iwakawa H, Rosant J, et al. (2000) Long glutamine tracts cause nuclear localization of a novel form of huntingtin in medium spiny striatal neurons in HdhQ25 and HdhQ11 knock-in mice. Hum mol genet 9(4): 503–513.

20. Tropepe V, Sibilla M, Ciruna BG, Rossant J, Wagner EF, et al. (1999) Distinct neural cell types proliferate and differentiate in Pdgf and in the developing mouse telencephalon. Developmental biology 200: 166–.
52. Myers RH, Vonsattel JP, Paskevich PA, Kiely DK, Stevens TJ, et al. (1991) Decreased neuronal and increased oligodendroglial densities in Huntington's disease caudate nucleus. Journal of neuropathology and experimental neurology 50: 729–742.

53. Mason HA, Rakowiecki SM, Raftopoulou M, Nery S, Huang Y, et al. (2005) Notch signaling coordinates the patterning of striatal compartments. Development 132: 4247–4258.

54. Lorincz MT, Zawistowski VA (2009) Expanded CAG repeats in the murine Huntington's disease gene increases neuronal differentiation of embryonic and neural stem cells. Molecular and cellular neurosciences 40: 1–13.

55. Kennedy L, Shelbourne PF (2000) Dramatic mutation instability in HD mouse striatum: does polyglutamine load contribute to cell-specific vulnerability in Huntington's disease? Hum Mol Genet 9: 2539–2544.

56. Goula AV, Berquist BR, Wilson DM, 3rd, Wheeler VC, Trottier Y, et al. (2009) Stoichiometry of base excision repair proteins correlates with increased somatic CAG instability in striatum over cerebellum in Huntington's disease transgenic mice. PLoS genetics 5: e1000749.

57. Kennedy L, Evans E, Chen CM, Craven L, Detloff PJ, et al. (2003) Dramatic tissue-specific mutation length increases are an early molecular event in Huntington disease pathogenesis. Hum Mol Genet 12: 3359–3367.

58. Zuccato C, Valenza M, Cattaneo E (2010) Molecular mechanisms and potential therapeutical targets in Huntington's disease. Physiological reviews 90: 905–981.

59. Marler K, Mehler MF (2012) Development and neurodegeneration: turning HD pathogenesis on its head. Neurology 79: 621–622.

60. Dogan I, Eickhoff SB, Schulz JB, Shah NJ, Laird AR, et al. (2012) Consistent Neurodegeneration and Its Association with Clinical Progression in Huntington's Disease: A Coordinate-Based Meta-Analysis. Neur-odegenerative diseases.

61. Braak H, Thal DR, Ghebremedhin E, Del Tredici K (2011) Stages of the pathologic process in Alzheimer disease: age categories from 1 to 100 years. J Neuropathol Exp Neurol 70: 960–969.

62. Braak H, Del Tredici K (2009) Neuroanatomy and pathology of sporadic Parkinson's disease. Adv Anat Embryol Cell Biol 201: 1–119.

63. Vonsattel JP, DiFiglia M (1998) Huntington disease. J Neuropathol Exp Neurol 57: 369–384.

64. Abrajano JJ, Qureshi IA, Gokhan S, Zheng D, Bergman A, et al. (2009) Differential deployment of REST and CoREST promotes glial subtype specification and oligodendrocyte lineage maturation. PloS one 4: e7665.

65. Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods 25: 402–408.

66. Pfaffl MW, Hieger GW, Dampfle I. (2002) Relative expression software tool (REST) for group-wise comparison and statistical analysis of relative expression results in real-time PCR. Nucleic Acids Res 30: e36.