Post-translational modifications clustering within proteolytic domains decrease mutant huntingtin toxicity

Huntington’s disease (HD) is an autosomal dominant neurodegenerative disorder caused by a CAG triplet repeat expansion in the huntingtin (Htt) protein. Post-translational modifications (PTMs) control and regulate many protein functions and cellular pathways, and PTMs of mutant Htt are likely important modulators of HD pathogenesis. Alterations of selected numbers of PTMs of Htt fragments have been shown to modulate Htt cellular localization and toxicity. In this study, we systematically introduced site-directed alterations in individual phosphorylation and acetylation sites in full-length Htt constructs. The effects of each of these PTM alteration constructs were tested on cell toxicity using our nuclear condensation assay and on mitochondrial viability by measuring mitochondrial potential and size. Using these functional assays in primary neurons, we identified several PTMs whose alteration can block neuronal toxicity and prevent potential loss and swelling of the mitochondria caused by mutant Htt. These PTMs included previously described sites such as serine 116 and newly found sites such as serine 2652 throughout the protein. We found that these functionally relevant sites are clustered in protease-sensitive domains throughout full-length Htt. These findings advance our understanding of the Htt PTM code and its role in HD pathogenesis. Because PTMs are catalyzed by enzymes, the toxicity-modulating Htt PTMs identified here may be promising therapeutic targets for managing HD.

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1 To whom correspondence may be addressed: CMSC 8-121, 600 N. Wolfe St., Baltimore, MD 21287. E-mail: narbez1@jhmi.edu.
2 To whom correspondence may be addressed: CMSC 8-121, 600 N. Wolfe St., Baltimore, MD 21287. Fax: 410-614-0013; E-mail: caross@jh.edu.
3 The abbreviations used are: HD, Huntington’s disease; PTM, post-translational modification; aa, amino acid(s); TMRRM, tetramethyl rhodamine methyl ester; DIV, day in vitro.

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Palmitoylation of Htt by HIP14 at cysteine 214 mediates membrane attachment (15), and a reduced palmitoylation of mutant Htt can be responsible for Htt mislocalization and aggregation. The acetylation of lysine 444 has been shown to improve clearance of the mutant protein by macroautophagy by facilitating trafficking of mutant Htt into autophagosomes and reduced Htt toxicity in cell and Caenorhabditis elegans models (16).

In a previous study of Htt PTMs, we identified several phosphorylation sites within the first 586-amino acid (aa) (N586) fragment of Htt (17). Using toxicity assays and other functional measurements, we confirmed sites previously described, and we identified serine 116 as a critical site to control Htt toxicity. Unlike for the other previously identified sites, the phospho-null alteration of Ser\textsuperscript{116} was protective for neurons against mutant Htt toxicity. Other alterations previously shown to reduce Htt toxicity include serines 421, 536, 1181, and 1201 (17). At serine 421, phosphorylation by Akt and SGK improves axonal transport and protects neurons (18, 19). At serine 536, the cleavage of Htt by calpain reduces mutant Htt toxicity (20). Phosphorylation by cyclin-dependent kinase 5 at serines 1181 and 1201 is induced by DNA damage and is neuroprotective (21).

Many of the PTMs studied in these previous experiments were identified in a context of Htt fragments and in cell lines. There has been relatively little study of the PTMs on full-length Htt. In a recent study (22), we analyzed by mass spectrometry PTMs on full-length Htt immunoprecipitated from human postmortem and mouse HD brains. We identified close to 40 PTMs, including 17 novel sites, throughout full-length Htt.

In the current study, we characterized the functional effects of PTM site modifications on mutant Htt. We introduced in full-length Htt alterations preventing PTMs of the amino acids that we have found modified in human or mouse brain, to determine whether abolition of single PTM sites could reduce mutant Htt-induced toxicity and mitochondrial dysfunction. We were able to identify several PTM sites that can modulate expanded Htt toxicity and its effects on mitochondria. These sites are located outside of the HEAT repeat domains and appear to cluster in the proteolysis-prone regions. This study is the first step toward deciphering the PTM code of the full-length Htt and validating PTMs on Htt as potential therapeutic targets for HD.

Results

Neuronal toxicity of expanded full-length Htt

As our primary screen, we have used a previously established cell death assay based on the measurement of nuclear condensation during neuronal cell death using fragments of Htt, Htt Exon 1 (23), and Htt aa1–586 (17). In this study, to evaluate the effects of post-translational modifications identified in the full-length Htt, we first have established the toxicity of full-length Htt in our neuronal model. This new model allows comparison of the toxicity levels between the fragments and full-length Htt. First, we transfected neurons with equal amounts of each construct. After 24 h of transfection, neurons transfected with full-length constructs express detectable levels of protein. Htt is distributed throughout the cell and does not present signs of aggregation (Fig. 1A). When measured by nuclear condensation, full-length normal Htt-23Q induced only a low level of toxicity, comparable in this system with levels observed in cells transfected with Htt fragments, whereas transfection with expanded Htt-82Q elicited a significant amount of cell death (Fig. 1B). However, the total amount of toxicity that is induced by full-length Htt is significantly lower than the toxicity induced by Htt aa 1–586 or by exon 1 containing the same number of glutamines.

Expanded full-length Htt PTM alterations

As a first step toward understanding the complexity of PTM of full-length Htt and identifying the potential site for therapeutic intervention on Htt, we studied whether an alteration of a single PTM site may affect toxic and functional properties of mutant Htt. Similar to our previous study with Htt aa 1–586 (17), we developed a series of constructs expressing full-length Htt-82Q with single point mutations at PTM sites. In addition to modification-null alterations done for all the selected sites (Ser to Ala and lys to Arg), we have also generated modification-mimetic mutations (Ser to Asp, Ser to Glu, and Lys to Glu) for some PTMs of particular interest. To produce these PTM alterations throughout full-length Htt, we used a multistep approach (described under “Experimental procedures”) because of the large size of Htt and the presence of the CAG repeats. All constructs were shown by Western blot analysis in HEK cells (supplemental Fig. S1) and by immunocytochemistry in primary cortical neurons (supplemental Fig. S2) to produce comparable levels of full-length Htt. The list of sites selected for this study was based on the PTM sites identified by mass spectrometry on human and mouse brains (22), as well as some of the previously identified PTMs within the N terminus of Htt and in full-length cell models (summarized in Ref. 7), and is detailed in Table 1. Many of these N-terminal sites have not been previously characterized in the context of full-length Htt.

Alterations of PTM modulate Htt toxicity

Toxicity measurements of the point mutation PTMs constructs in primary cortical neurons is shown on Fig. 2A. Similar to previous reports, we found that phospho-mimetic alteration of the Thr\textsuperscript{3} and Ser\textsuperscript{13/16} sites ameliorated expanded Htt toxicity (8, 9, 13) and confirmed our previous findings with Htt aa 1–586 (17) that S116A alteration has a neuroprotective effect against the toxicity of expanded Htt to primary neurons. Contrary to previous studies with the 1–480 fragment of Htt (18, 25) but consistent with our previous data obtained using the 1–586 fragment of Htt (17), we found that both phospho-null (S421A) and presumably phospho-mimetic (S421D and S421E) alterations elicited neuroprotection in our system.

The acetylation mimetic Htt K444Q alteration showed no protection, whereas the acetylation-null alteration K444R ameliorated expanded Htt toxicity measured by nuclear condensation. Previously, acetylation at this site has been shown to be protective (16). Other modification-null alterations that reduced toxic properties of expanded Htt in cortical neurons are K1190R, S1201A, K2548R, and S2652A. Notably, an absence of Htt phosphorylation at position Ser\textsuperscript{1201} (within 1301
amino acids fragment of Htt with normal poly(Q) range) was previously found to make Htt toxic at levels similar to those of expanded protein, whereas it had no effect on poly(Q)-Htt-induced cell death in striatal neurons (21).

Because striatum appears to be most affected in HD (26–28), we tested whether primary striatal neurons are as sensitive to acute toxicity of full-length Htt as cortical neurons. Overexpression of full-length Htt-82Q with and without PTM alterations in striatal neurons induced levels of toxicity comparable with the levels observed in cortical neurons transfected with same constructs (Fig. 2B). These results show that PTM alterations that ameliorated toxicity of expanded Htt in cortical neurons also protected striatal neurons. The only exception is the S1201A substitution, which, unlike in cortical neurons, did not result in a protective effect in striatal neurons, consistent with a previous report (21).

In addition to the nuclear condensation assay, we also conducted a direct observation of cell death using previously established (17) time-lapse video imaging of primary neurons (Fig. 3). We followed cells over time by video microscopy, beginning at hour 24 and ending at hour 34 post-transfection. Morphology of every neuron was quantified blindly using ImageJ software. The cells were considered dead when the majority of the neurites detached from the soma or were fragmented or when the soma became totally round. Morphological changes of neurons transfected with expanded full-length Htt started to appear after 3 h of imaging (27 h post-transfection). Cellular extensions retracted, fragmented, and detached from the soma. The cell soma progressively got rounder and in the most extreme cases disappeared. The morphology of the neurons showed a strong and steady decline over the course of the experiment (Fig. 3A). By the end of the imaging, most cells transfected with Htt-82Q were considered dead according to morphological criteria. The viability of control neurons transfected with normal full-length Htt showed a relatively small change in morphology by the end of the experiment. We previously showed that this late-stage decline is not specifically due to Htt because cells transfected with GFP alone undergo the same process (17). Examples of morphological changes of single transfected with the Ser13/Ser16 constructs and the controls observed over time are shown on Fig. 3A.

Consistent with the results of nuclear condensation assay, we observed that S13D/S16D phospho-mimetic alteration slows down the progression of the neuronal death induced by expanded Htt (Fig. 3B), whereas a phospho-null S13A/S16A mutation renders expanded Htt even more toxic to the neurons than unmodified full-length Htt-82Q protein. Thus, unlike the nuclear condensation assay, the time-lapse imaging allows
detection of a potential increase in toxic properties of expanded Htt with certain PTM alterations. We also tested the effect of the alterations of Ser421 in the same system (Fig. 3C). Because both phospho-null and phospho-mimetic mutations at Ser421 proved to be protective in our nuclear condensation assay, we investigated the possibility of a different effect on neuronal morphology. Both S421A and S421D alterations appeared to ameliorate expanded Htt toxicity when assayed by time-lapse imaging, in agreement with the nuclear condensation results. The other PTM alterations, K1190R, K2548R and S2652A, that showed a neuroprotective effect in nuclear condensation assay, also significantly decreased toxicity of expanded Htt, as measured by time-lapse imaging (Fig. 3C).

### Mitochondrial abnormalities caused by expanded Htt are modulated by PTMs

Metabolism and mitochondrial dysfunction have been implicated in HD pathogenesis (29–34). Mutant Htt was shown to directly associate with mitochondria, impair mitochondrial dynamics, and decrease mitochondrial function in affected in HD brain regions (35, 36).

Mitochondrial potential integrity is important for cell survival, and depolarization is a sign of cell toxicity (37). We have established an assay to measure the mitochondrial potential in primary neurons using tetramethyl rhodamine methyl ester (TMRM), a live mitochondrial dye sensitive to the potential (examples of staining appear in Fig. 4A). Using high magnification live microscopy and the Volocity software, we measured the average intensity of TMRM staining in primary neurons transfected with full-length Htt constructs with and without PTM alterations.

After quantification using Volocity, as shown on Fig. 4C, there is a significant decrease of the mitochondrial potential of the neurons transfected with expanded full-length Htt-82Q compared with cells transfected with normal repeat Htt-23Q. We found that several alterations of previously studied sites within the N-terminal domain (T3A, S13A/S16A, S13D/S16D, S116A, S271A, S421A, and S432/S434A) were able to partially rescue the loss of mitochondrial potential, observed in primary cortical neurons transfected with expanded Htt. A few modifications within the middle part of protein (K1190R, S1201A, and K1204R) and within its C-terminal domain (S2342A, S2489A, S2342/S2346, S2548R, and K2615R) may also modulate a drop in mitochondrial potential induced by expanded Htt (Fig. 4C). When the survival of the cell and the mitochondrial potential values are
normalized to the controls (scale from 0 for Htt-82Q to 100 for Htt-23Q), the resulting correlation is moderate ($r = 0.5714$; supplemental Fig. 3A).

Mitochondrial swelling is also an early sign of cell toxicity (37). We used MitoTracker Red dye and confocal microscopy to measure mitochondrial size in primary neurons transfected with full-length Htt constructs with and without PTM alterations. We found significant mitochondrial swelling in neurons transfected with expanded Htt-82Q, compared with cells transfected with normal repeat Htt (examples of staining appear in Fig. 4B), whereas little to no fragmentation of the mitochondria was observed. Some of the PTM alterations were able to protect the mitochondria from the toxic effect of expanded Htt (Fig. 4D). Unlike changes in potential, there was a better correlation between the alterations that were protective against the mitochondrial swelling and those that ultimately protected from cell death. When the survival of the cell and the mitochondrial size values are normalized to the controls (scale from 0 for Htt-82Q to 100 for Htt-23Q), the resulting correlation is moderate ($r = 0.7677$; supplemental Fig. 3B).

Overall, with changes in both morphology and potential, our results suggest an early involvement of the mitochondria in the toxic mechanisms of expanded Htt. We observed a partial overlap between PTM alterations in expanded Htt that may modulate mitochondrial function with the sites that appear to affect neuronal toxicity of expanded Htt, as measured by nuclear condensation assay or by direct imaging of neuronal death.
Notably, most of PTM alterations that rescued toxicity also improved the mitochondrial phenotype.

Discussion

In our previous studies, we demonstrated the toxicity of mutant exon 1 and Htt aa 1–586 fragments of Htt in primary cortical neurons measured by nuclear condensation assay (23, 17). Here, we extended the study of Htt toxicity and the effects of PTM site modifications to full-length Htt. In accordance with mouse models of HD that show a faster and stronger phenotype in models expressing shorter fragments of mutant Htt (38), we find greater toxicity of mutant Htt N-terminal fragments compared with full-length mutant Htt. The Htt protein is very large and likely serves a scaffold, with many protein interaction partners. It has a complex structure, with intramolecular interactions and multiple sites of PTMs, with potential for cross-talk that may affect the structure Htt and its interactions with other proteins, as well as influencing Htt subcellular...
localization. Thus, it is essential to characterize and study PTMs within the context of the full-length protein, as we have done in this study.

Our toxicity experiments using full-length Htt expression in primary neurons support previous reports that some phospho-mimetic alterations of expanded Htt (T3D, S13D/S16D, and S421D) are neuroprotective. However, we observed several differences compared with published studies that used fragments of Htt. In contrast with earlier reports (18, 19), but in line with our previous data (17), we found neuroprotection with both phospho-null (S421A) and phospho-mimetic (S421D and S421E) alterations of S421, suggesting the possibility that Asp acts as a phospho-null rather than phospho-mimetics at this site and that phosphorylation may enhance toxicity.

Previous reports (18, 19) have suggested that phosphorylation is protective. However, the experimental systems used in the previous studies were quite different from ours and often did not directly assay neuronal toxicity of mutant Htt. Recently, the importance of the potential phosphorylation at serine 421 for modulation of pathogenesis has been confirmed in vivo by
the analysis of mutant HTT-S421D BACHD mice. However, a difference in expression levels in the mutant HTT-S421A did not allow a definitive comparison among the three constructs (39). Overall, there is generally consensus on which are the relevant PTM sites, with discrepancies among models regarding whether phosphorylation is protective or pathogenic, in part because of the limitations of using amino acid substitutions to mimic or remove a phosphorylation. More experiments, both in vitro and in vivo, will be required in the future.

In our full-length mutant Htt neuronal model, we also confirmed the activity of several phospho-null modifications. S116A, S1201A, and S2652A were protective alterations. The effect of Ser116 confirmed our previous results with Htt aa 1–586 (17). These sites for which the phospho-null mutation is protective are of particular interest, because they are potential candidates for small molecule therapeutic development targeting the potential kinases active at the site. Using a kinase prediction software, NetPhos 3.1 (40), we identified candidates kinases, which include PKC, p38 MAPK, and CKII.

Interestingly, we found that S1201A yielded different results in striatal neurons than in cortical neurons. This is consistent with the idea of cell type–specific vulnerability and protective factors. Therefore exploring mechanisms of Ser1201 phosphorylation protection could uncover potential neuronal cell type–specific protective pathways. Previously, the modification of Ser1201 was reported to have no effect on cell death in striatal neurons, while turning normal Htt into pro-apoptotic protein (21).

It has been suggested by several groups (35, 36, 41), including our own (33), that Htt may be localized to mitochondria (possibly on the outer membrane, but may also enter and interact with mitochondrial proteins). The effect of Htt phosphorylation on the mitochondrial localization and function of Htt is largely unknown and could be the subject of future studies. We did not observe a complete correlation between toxicity and mitochondrial measures, suggesting that mitochondrial pathways represent just one portion of pathogenesis or neuroprotection.

In this full-length model, we also identified several acetylation sites that can modulate Htt toxicity and effects on mitochondria. Acetylation at Lys444 has been reported to be protective (16); however, putative acetylation-mimetic Htt K444Q alteration showed no effect in our system, whereas acetylation-null alteration K444R ameliorated expanded Htt toxicity. These discrepancies of our data with some previous findings may be accounted for by using the full-length Htt system, preserving potential short-range and distant PTM interactions within the whole protein. Interestingly, each of the acetylation sites we observed having an effect on toxicity was located near one of the identified phosphorylation sites, suggesting the possibility of cross-talk between acetylation and phosphorylation of Htt, which merits further investigation.

Biophysical and cross-linking mass spectroscopy experiments indicated that Htt adopts a compact conformation with intramolecular contacts (42), both short and long range. It is likely that PTMs can modulate these interactions. The physical interaction between matching N-terminal and C-terminal fragments of Htt may be important for Htt toxicity. When the N-terminal and C-terminal fragments do not interact, there may be enhanced toxicity of the fragment containing the polyglutamine, but also it is conceivable that the C-terminal fragment may become toxic (43).

Predictions of structure of Htt based on its sequence show that there are unstructured proteolysis-prone PEST domains between highly structured α-helix supercoiled HEAT repeat domains (44). Hayden and co-workers (6) suggested that most of the PTM identified on Htt are found in these PEST domains and not in the HEAT repeat domains, and our data confirm and extend this concept.

Locations of the Htt PTMs with functional effects are shown in a schematic diagram of Htt in Fig. 5. Because we have used several approaches to functionally assess PTMs identified on
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the full-length Htt, we combined the results to summarize the properties of each site. At each PTM, a score of 1 was given for each assay where significant changes were observed (and score 0 if there were no significant changes).

Based on this scoring, we found four clusters of PTMs that modulate expanded Htt functional or toxic properties: the previously well characterized N-terminal domain (cluster 1, residues 1–116), the region around Ser116 (cluster 2), the mid region (cluster 3, residues Ser2101–Lys1246), and C-terminal cluster 4 (residues Ser2342–Ser657). Notably, it appears that PTM clusters modulating Htt properties are grouped within predicted protease-sensitive regions (http://emboss.bioinformatics.nl/cgi-bin/emboss/epestfind; Fig. 5). Interestingly, three of these clusters of PTMs lie closely to the boundaries between previously suggested “distinct domains defined by protease-sensitive sites,” as described by Seong and co-workers (42, 45): our cluster 2 falls between N-terminal domains I and II; cluster 3 is close to the location of the protease-sensitive major hinge region (residues 1184–1254) between N-terminal domain II and C-terminal domain I; and cluster 4 maps to the C-terminal domain II boundary.

Proteolytic cleavage into a toxic N-terminal fragment containing the mutant polyglutamine stretch has been proposed to contribute to HD pathogenesis (6, 7, 28, 43, 46). Cleavage sites containing the mutant polyglutamine stretch has been proposed to

**Experimental procedures**

**Plasmid generation and mutagenesis**

Mammalian expression constructs encoding full-length Htt with the N-terminal PTM alterations (of Thr3, Lys6, Lys9, Ser13, Ser16, Ser116, and Ser220) were generated in two steps. As a first step, these alterations were introduced within an existing Htt-N586–82Q plasmid (obtained from D. Borchelt and described in Ref. 24), using site-directed mutagenesis and the QuikChange II XL kit (Stratagene) according to the manufacturer’s protocol, except that Stbl2™ competent cells (Thermo Fisher Scientific), specifically designed for cloning of unstable inserts, were used throughout this study. The presence of PTM mutations and the entire N586–82Q sequences were confirmed by sequencing. As a second step, the fragments comprising the first 171 aa of Htt with corresponding mutations were generated from the above N586 constructs using digestion at 5’ flanking NotI and Htt-internal XhoI restriction sites. These N-terminal fragments were subcloned into a vector comprising the rest of the downstream sequence of full-length Htt. This vector was constructed by subcloning Xhol/Sall downstream Htt fragment (of the original synthetic plasmid containing the entire full-length Htt-23Q) in the XhoI site of pcDNA vector. The synthetic FL-Htt-23Q plasmid was generated (DNA 2.0) previously and was checked for expression of full-length Htt protein in our laboratory. PTM alterations of Thr271, Ser221, Ser213, and Lys444 were also introduced within an existing Htt-N586–82Q plasmid, as described above. As a second step, the fragments containing N586–82Q with corresponding mutations were generated by PCR of the above Htt-586–82Q constructs using the primers incorporating KpnI restriction sites on both 5’ and 3’ ends. Purified PCR products were digested with the above enzyme and subcloned into a vector comprising the rest of the downstream sequence of full-length Htt (described above), prepared upon digestion with KpnI (utilizing 5’ flanking and Htt-internal KpnI sites). The rest of the PTM alterations (C-terminal of aa 586) were generated using a newly synthesized big C-terminal Htt fragment (downstream of aa 586) with incorporated convenient unique restriction sites (DNA2.0), allowing isolation of five domains separated by these restriction sites. These domains were subcloned into pJ cloning vectors (DNA2.0) and used to introduce alterations by site-directed mutagenesis and the QuikChange II XL kit (Stratagene). These mutagenized domains were subcloned into the big C-terminal Htt fragment using unique restriction sites (swapping with corresponding domains without mutations). At the final step we reintroduced the N-terminal 586–aa fragment with poly(Q) into mutagenized C-terminal fragment (for each PTM alteration).

**Primary neuron preparation**

Primary neurons from CD1 mice were prepared from embryonic days 15–17 embryos. Cells from both cortex and striatum were independently prepared from the same embryos and plated at 10⁶ cells/mm² in Neurobasal medium with 2% B27, 2 mM GlutaMAX, and 1% penicillin/streptomycin in a 24-well plate. The cells were maintained at 37 °C with 5% CO₂ throughout the experiments. All cell culture supplies were obtained from Corning, and all the media were from Thermo Fisher Scientific.

**Transfection of primary neurons**

Neurons were co-transfected using Lipofectamine 2000 (Thermo Fisher Scientific) at day in vitro (DIV) 5 or 6. Per well of 24-well plates, 1 μg of Htt plasmid, and 0.1 μg of GFP was

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used with a ratio DNA/Lipofectamine of 1/1.75. DNA and Lipofectamine were diluted separately in 50 μl of OptiMEM/well for each and incubated for 5 min before being combined. After being combined, the transfecting solution is incubated for 90 min at room temperature. During the incubation of the transfecting solution, the plates are emptied, and the medium is replaced with 400 μl of fresh OptiMEM. After the incubation, 100 μl of transfection solution is deposited in each well. The plates are returned to the incubator for 4.5 h when the wells are empty, and the medium was replaced with fresh Neurobasal medium.

**Nuclear condensation cell death assay**

The toxicity experiments were performed in primary cortical and primary striatal neurons according to our established protocol (16). Nuclei were stained using Hoechst 33424 (Sigma; 0.2 μg/ml in PBS for 5 min). Automated picture acquisition was performed using a Zeiss Axiovert 200 inverted microscope with a 10× objective. Automatic quantification of the nuclear intensity of transfected cells was performed using Volocity. The cells were considered dead when their nuclear intensity was higher than the average intensity plus two standard deviations. Each condition was performed in quadruplicate within each experiment, and each experiment was repeated in at least four independent neuronal preparations for each construct studied.

**Time-lapse imaging of cell death**

Primary cortical neurons were transfected with full-length Htt constructs and GFP as described above. 24 h after transfection, neurons expressing GFP were randomly chosen and followed for 10 h with a picture taken every 20 min. Morphology of every neuron was quantified blindly using ImageJ software. Neurons were quantified as 100 when alive and 0 when dead for each frame. The cells were considered dead when most of the neurites detached from the soma or were fragmented or when the soma became totally round. The results are expressed as means ± S.E. (n = 200 cells analyzed in three independent experiments).

**Mitochondrial function assays**

For the potential measurement, primary cortical neurons were co-transfected at DIV5 with the full-length Htt constructs and GFP, as described above. 24 h after transfection, the cells were loaded with 20 nM TMRM stain for 45 min at 37 °C. After three washes with PBS, the medium was changed to Neurobasal + B27 without phenol red for imaging. The cells were imaged on an Axiovert 200 microscope. Average measure of the TMRM intensity of transfected cells was quantified using Volocity. For mitochondrial size measurement of the mitochondria, primary cortical neurons were co-transfected at DIV5 with the full-length Htt constructs and GFP, as described above. 24 h after transfection, the cells were loaded with 500 nM MitoTracker Red (Molecular Probes) for 45 min at 37 °C. After three washes with PBS, the cells were fixed with 4% paraformaldehyde, 1× PBS for 30 min. The cells were imaged on a laser-scanning confocal Zeiss LSM 510-meta microscope. Average sizes of individual mitochondria of transfected cells were measured using Volocity. The results were normalized to the average size of mitochondria of cells transfected with Htt-82Q.

**Author contributions**—N. A. designed, performed, and analyzed the experiments and wrote the paper. T. R., E. R., E. C., and J. C. S. designed and engineered all the plasmids, performed the expression experiments, and prepared the figures. T. R. compiled the results and designed Fig. 5. M. R. and X. W. provided technical assistance and contributed to the preparation of the neurons. D. J. L. coordinated the study. C. A. R. conceived and coordinated the study. All authors reviewed the results and approved the final version of the manuscript.

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