Indirect inhibition of 26S proteasome activity in a cellular model of Huntington’s disease

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Pathognomonic accumulation of ubiquitin (Ub) conjugates in human neurodegenerative diseases, such as Huntington’s disease, suggests that highly aggregated proteins interfere with 26S proteasome activity. In this paper, we examine possible mechanisms by which an N-terminal fragment of mutant huntingtin ( htt; N-htt) inhibits 26S function. We show that ubiquitinated N-htt—whether aggregated or not—did not choke or clog the proteasome. Both Ub-dependent and Ub-independent proteasome reporters accumulated when the concentration of mutant N-htt exceeded a solubility threshold, indicating that stabilization of 26S substrates is not linked to impaired Ub conjugation. Above this solubility threshold, mutant N-htt was rapidly recruited to cytoplasmic inclusions that were initially devoid of Ub. Although synthetically polyubiquitinated N-htt competed with other Ub conjugates for access to the proteasome, the vast majority of mutant N-htt in cells was not Ub conjugated. Our data confirm that proteasomes are not directly impaired by aggregated N-terminal fragments of htt; instead, our data suggest that Ub accumulation is linked to impaired function of the cellular proteostasis network.

Introduction

Ubiquitin (Ub) has been recognized for >20 yr to be a prominent and invariant feature of intracellular inclusion bodies (IBs) in the vast majority of neurodegenerative diseases, and Ub immunohistochemistry is considered to be the definitive marker in postmortem neuropathological diagnosis of these disorders (Lehman, 2009). In the absence of disease, Ub is diffusely distributed in brain tissue, whereas in diseased brain and in cell culture models thereof, cellular Ub becomes visibly concentrated together with aggregated proteins within IBs (Mayer et al., 1989), and markedly elevated levels of poly-Ub conjugates can be detected by mass spectrometry (MS) in tissue lysates (Bennett et al., 2007). Although the linkage of disrupted Ub homeostasis and protein aggregation to disease is undisputed, the functional significance of these correlations remains unresolved, and the mechanistic implications for pathogenesis remain controversial.

One prominent hypothesis holds that aggregated proteins inhibit protein degradation by the 26S proteasome (26S; Taylor et al., 2002; Ciechanover and Brundin, 2003; Valera et al., 2005; Ortega et al., 2007). Poly-Ub chains are best known for their role in targeting proteins for degradation by 26S, and enzymatic disassembly of these chains by 26S-associated deubiquitinating enzymes is essential to allow entry of substrates into the 20S proteolytic chamber and to “recycle” Ub to maintain cellular-free Ub levels (Finley, 2009). Experimental impairment of proteasome function leads to accumulation of Ub conjugates and to broad dysregulation of cellular Ub homeostasis, and mice harboring loss-of-function mutations in genes required for maintenance of Ub homeostasis develop neurodegenerative disorders (Saigoh et al., 1999; Wilson et al., 2002; Ryu et al., 2008). However, despite the well-documented accumulation of Ub conjugates in the neurodegenerative disease brain, there is no consistent evidence documenting an accompanying decrease in levels of free Ub or disruption of Ub pool dynamics in neurodegenerative disease or models thereof, and Ub conjugate accumulation may be a benign but robust indication of an underlying lesion in 26S function. In addition to degrading aggregation-prone, folding-defective...
neurodegenerative disease-linked proteins (Bence et al., 2001; Illing et al., 2002; Petrucelli et al., 2002; Mandrusiak et al., 2003; Bennett et al., 2005; van Tijn et al., 2007; Deriziotis and Tabrizi, 2008). Although these studies are consistent with the hypothesis that protein aggregates can inhibit UPS function, direct measurement of proteasome function in extracts from brains of patients or animal models of neurodegenerative disease has yielded equivocal results, with decreased (Kabashi et al., 2004), increased (Bett et al., 2006), or unchanged (Diaz-Hernández et al., 2003; Gillardon et al., 2007) levels of proteasome activity being reported in different diseases and model systems.

In the present study, we have investigated the temporal and mechanistic relationship between aggregation of an N-terminal exon 1 fragment of huntingtin (htt; N-htt) and UPS function using a quantitative flow cytometry assay and live-cell time-lapse imaging to simultaneously interrogate N-htt expression, proteasome control the levels of critical cellular regulators, and impaired 26S function has severe and pleiotropic consequences for cell function and survival (Kawazoe et al., 1998).

The hypothesis that protein aggregates impair 26S function is supported by the demonstration that endogenous short-lived 26S substrates accumulate in models of Huntington’s disease (HD; Jana et al., 2001; Zemskov and Nukina, 2003) and by studies using synthetic reporters consisting of GFP fused to destabilizing degrons (Dantuma et al., 2000; Bence et al., 2001; Bennett et al., 2005). In cells expressing these reporters, steady-state levels of GFP fluorescence should—assuming constant synthesis rates—provide a readout of Ub–proteasome system (UPS) functional capacity. Multiple studies have confirmed that the levels of these reporters, when expressed in mammalian cell culture, increase significantly upon coexpression of the aggregation-prone mutant but not wild-type, nonaggregating variants of neurodegenerative disease-linked proteins (Bence et al., 2001; Illing et al., 2002; Petrucelli et al., 2002; Mandrusiak et al., 2003; Bennett et al., 2005; van Tijn et al., 2007; Deriziotis and Tabrizi, 2008). Although these studies are consistent with the hypothesis that protein aggregates can inhibit UPS function, direct measurement of proteasome function in extracts from brains of patients or animal models of neurodegenerative disease has yielded equivocal results, with decreased (Kabashi et al., 2004), increased (Bett et al., 2006), or unchanged (Diaz-Hernández et al., 2003; Gillardon et al., 2007) levels of proteasome activity being reported in different diseases and model systems.

In the present study, we have investigated the temporal and mechanistic relationship between aggregation of an N-terminal exon 1 fragment of huntingtin (htt; N-htt) and UPS function using a quantitative flow cytometry assay and live-cell time-lapse imaging to simultaneously interrogate N-htt expression,
Ub distribution, and UPS function in single cells. Our data establish that N-htt—whether aggregated or not—does not choke or clog 26S and lead us to propose a model in which Ub accumulation is the consequence of simple competition for limited 26S capacity by cellular proteins that become diverted to the UPS in the face of disrupted protein folding homeostasis, which occurs when one or more components of the cell’s proteostasis network becomes overwhelmed by folding-impaired, aggregation-prone N-htt.

**Results**

**UPS substrates accumulate above a critical concentration threshold of mutant htt expression**

To precisely assess the quantitative relationship between the amount of N-htt expressed in cells and accumulation of a UPS reporter, we devised a two-color flow cytometry–based assay to simultaneously analyze UPS function using an unstable green fluorescent UPS reporter, Ub^{G76V}-GFP (Dantuma et al., 2000), and N-htt expression using a red (mCherry) fluorescent N-htt fragment, N-htt(Qn)–chFP (Fig. 1). Control experiments established that the presence of chFP does not alter the aggregation properties of N-htt(Qn) (Fig. S1 A) or its impact on the UPS (Fig. S1 B). Because N-htt(Q25)–chFP or N-htt(Q91)–chFP differ only in the length of the polyglutamine tract, the red fluorescence signal can be used to directly compare the expression levels of the two proteins. Expression of N-htt(Q25)–chFP in cells stably expressing Ub^{G76V}-GFP resulted in the appearance of a population of doubly labeled cells in which the distribution of green fluorescence was not substantially different from that of untransfected reporter cells (Fig. 1 A, bottom). In contrast, expression of N-htt(Q91)–chFP resulted in the appearance of a distinct subpopulation of cells displaying high levels of both green and red fluorescence. Transformation of the data in Fig. 1 A to plot the concentration of N-htt(Qn)–chFP versus Ub^{G76V}-GFP fluorescence (Fig. S2) revealed that although N-htt(Q25)–chFP had only a minimal effect on the level of green fluorescence, N-htt(Q91)–chFP induced accumulation of the UPS reporter at high expression levels (Fig. 1 B).

The Q91-specific mean reporter fluorescence, determined by subtracting green fluorescence in cells expressing N-htt(Q25)–chFP from that of cells expressing N-htt(Q91)–chFP, shows that Ub^{G76V}-GFP accumulates significantly only in cells expressing N-htt(Q91)–chFP above a threshold (~22 arbitrary units [a.u.]; Fig. 1 B). This threshold concentration of N-htt(Qn)–GFP at which the UPS reporter chFP-CL1 accumulated was inversely related to polyglutamine length (Fig. 1 C), suggesting a strict relationship between N-htt aggregation and UPS impairment. We further found that the concentration of N-htt(Q91)–chFP at which we observed stabilization of Ub^{G76V}-GFP was strongly affected by the length of time that the cells had been expressing mutant N-htt (Fig. 1 D). These data suggest that chronic expression of mutant N-htt decreases the intrinsic capacity of cells to degrade UPS substrates.

**Htt aggregation is associated with accumulation of diverse 26S substrates**

Stabilization of Ub^{G76V}-GFP by high levels of mutant N-htt could be caused by interference by aggregated or oligomeric protein with 26S function, with some aspect of the Ub conjugation cascade, or both. To discriminate between these possibilities, we generated HEK293 cell lines stably expressing GFP molecules that are destabilized by degrons that target proteins to 26S by different Ub-dependent mechanisms. These reporters include, in addition to Ub^{G76V}-GFP (Dantuma et al., 2000), a substrate of the Ub fusion degradation pathway that recognizes proteins with an uncleavable N-terminal Ub (Johnson et al., 1995), Ub-Arg-GFP (Dantuma et al., 2000), a substrate of the N-terminal end rule pathway that recognizes proteins with destabilizing N-terminal amino acids (Varshavsky, 1997), and GFP-CL1 (Bence et al., 2001), which is destabilized by a short amphipathic motif that, in yeast, targets proteins for ubiquitination by a distinct set of E2 and E3 enzymes (Gilon et al., 2000).

![Figure 2](image_url)
Figure 3. Ubiquitinated polyglutamine aggregates do not choke the 26S proteasome in vitro. (A) In vitro ubiquitination of radiolabeled Sic1 substrate. \(^{35}\)S-PY-Sic1 purified from E. coli was ubiquitinated in vitro as described in Materials and methods. Aliquots of the reaction were removed at the indicated times, separated on a 4–20% gradient gel, and visualized by autoradiography. Mobilities of unmodified Sic1 and polyubiquitinated Sic1 are indicated.
In addition, we examined the effect of N-htt(Qn)–chFP expression on the levels of cODC-GFP, a 26S substrate that is destabilized by a Ub-independent degron consisting of 37 C-terminal amino acids from ornithine decarboxylase (ODC; Zhang et al., 2003). All four reporters, stably expressed in clonal HEK293 cell lines, exhibit low steady-state fluorescence in the absence of stress and time-dependent increases in diffuse cytoplasmic fluorescence after challenge with MG132 (Fig. S3, A and B). Control experiments using ts20 Balb/C 3T3 fibroblasts harboring a thermolabile E1 (Kulka et al., 1988) verified that the cODC-GFP fusion protein degradation is indeed subject to Ub-independent degradation by 26S (Fig. S3 C).

Expression of N-htt(Q91)–chFP, but not N-htt(Q25)–chFP, resulted in an N-htt(Q91)–chFP dose-dependent increase in the fluorescence of all four reporters (Fig. 2 A), strongly suggesting that expression of N-htt(Q91)–chFP impairs the UPS downstream of any Ub-dependent targeting steps, most likely at the level of 26S itself. To further test this hypothesis, we assessed the effect of N-htt(Q91)–chFP expression on the degradation of substrates of the ER-associated degradation (ERAD), a quality control pathway that delivers misfolded or unassembled proteins in the early secretory pathway to the UPS (Vembar and Brodsky, 2008). We found that N-htt(Q91)–chFP, but not N-htt(Q25)–chFP, expression resulted in the accumulation of a folding-defective mutant cytisic fibrosis transmembrane conductance regulator, GFP-ΔF508 (Ward et al., 1995), and of unassembled TCR-α–GFP, a subunit of the T cell receptor (Fig. 2 B; Yu et al., 1997). Two additional ERAD substrates, an amyloidogenic mutant of the nonglycosylated protein trans-thrytin (Sekijima et al., 2005), and GluR1, an unassembled subunit of the n-methyl-d-aspartate receptor, exhibited similar behavior (unpublished data). Remarkably, all of these ERAD substrates accumulate at precisely the same threshold of N-htt(Q91)–chFP expression as found for the four cytosolic UPS reporters (Fig. 2, A and B). It is possible that the elevated fluorescence of all of these unstable GFP conjugates could be caused by increased transcription of the reporter cDNAs in response to N-htt(Q91)–chFP expression (Bowman et al., 2005; Alvarez-Castelao et al., 2009). However, we found no evidence of increased levels of GFP in response to the highest levels of N-htt(Q91)–chFP expression (Fig. S3 D). Because GFP is a very stable protein in mammalian cells (Bence et al., 2001), small changes in its rate of synthesis are integrated over time to generate robust changes in steady-state levels. The observation that all of the unstable reporters accumulate above the same level of N-htt(Q91)–chFP expression strongly suggests that mutant N-htt expression above a critical concentration threshold has pleiotropic effects on the activity or capacity of 26S. The finding that a Ub-independent 26S substrate, cODC-GFP, along with substrates that depend on different Ub conjugation pathways, accumulates in response to expression of N-htt(Q91)–chFP above a critical concentration threshold suggests that this pleotropic effect is not caused by an impaired Ub metabolism.

Ubiquitinated polyglutamine aggregates do not choke or clog the 26S proteasome

The most parsimonious model to explain why all UPS substrates accumulate in response to N-htt aggregation would predict that aggregated N-htt interacts directly with and inhibits 26S activity. Although we have previously shown that neither soluble nor aggregated polyglutamine peptides nor N-htt fragments inhibit 20S or 26S proteasomes in vitro (Bennett et al., 2005), it is conceivable that if polyubiquitinated, these aggregates could antagonize 26S proteasomes by virtue of their affinity for Ub receptors, but resistance to ATP-dependent unfolding associated with the 19S ATPases. To directly test the possibility that 26S proteasomes are inhibited in such a manner, we established an in vitro assay for 26S proteasome activity to measure the effect of polyubiquitination on the ability of aggregated polyglutamine-containing proteins to interfere with substrate degradation in trans. A purified 35S-labeled substrate (PY-Sic1) was efficiently and quantitatively converted to a high molecular weight polyubiquitinated PY-Sic1(Ub6) form in vitro by the E3 Ub ligase Rsp5 (Fig. 3 A; Saeki et al., 2005). Addition of purified proteasomes to this preparation resulted in efficient degradation that was dependent on ATP (unpublished data) and inhibited by MG132 and by pretreatment of the PY-Sic1(Ub6) substrate with the catalytic core of the deubiquitinating enzyme
Usp2 (USP2-cc; Fig. S4, A and B; Catanzariti et al., 2004). Michaelis–Menten analysis of \(^{35}S\)-PY-Sic1(Ub\(_n\)) degradation yielded a Michaelis constant (\(K_m\)) of 46.9 ± 3.5 nM (Fig. 3 B), in good agreement with a previous study with other 26S proteasome substrates in vitro (Thrower et al., 2000) and confirming that this model substrate can be used to quantify the activity of 26S in vitro. To assess inhibition of 26S activity in vitro, we used the same enzymatic strategy to generate high molecular weight polyubiquitinated forms of an unlabeled chimeric protein (MPC) consisting of an N-terminal maltose binding protein fused to a fragment of Xenopus laevis cyclin (N100; Chen and Fang, 2001) and a T7 epitope tag (Fig. 3 C). The initial rate of \(^{35}S\)-PY-Sic1(Ub\(_n\)) degradation by purified 26S rat liver proteasomes was strongly inhibited (inhibition constant \(K_i\) of 150 nM) by MPC(Ub\(_n\)) but not by nonubiquitinated (Fig. 3 D) or deubiquitinated (not depicted) MPC, confirming that ubiquitinated proteins can inhibit 26S function and establishing the utility of this assay for the evaluation of proteasome inhibition by ubiquitinated N-htt aggregates.

To assess the ability of ubiquitinated protein aggregates to inhibit 26S activity, we used a similar approach to append poly-Ub chains on GST–PY-N-htt(Qn) purified from Escherichia coli. A previous study has established that the presence of GST retards the aggregation of N-htt fusions containing intermediate (~Q50) length polyglutamine repeats and that aggregation of Q > 40 fusions can be efficiently induced upon cleavage of GST with a site-specific protease (Scherzinger et al., 1997). Both GST–PY-N-htt(Q18) and GST–PY-N-htt(Q51) were efficiently converted to high molecular weight Ub-conjugated species. Treatment with Usp2-cc resolved both fusion proteins to their native, unmodified molecular weights, indicating that their reduced mobility was caused by Ub conjugation rather than to aggregation (Fig. 3 E). Hydrolysis of the GST moiety with tobacco etch virus (TEV) protease rapidly converted GST–PY-N-htt(Q51) into high molecular weight aggregate that did not migrate beyond the stacking gel on SDS-PAGE and were efficiently retained on a 0.2-µm cellulose acetate filter trap (Fig. S4 C; Wanker et al., 1999), confirming that the presence or absence of GST can be used to switch this fusion protein from a soluble nonaggregated protein into an aggregated state. We found that GST–N-htt(Q18)(Ub\(_n\)) inhibited degradation of \(^{35}S\)-PY-Sic1(Ub\(_n\)) by 26S in a dose-dependent manner (Fig. 3 F), with a \(K_i\) of ~250 nM, similar to the value obtained for inhibition by MPC(Ub\(_n\)). Importantly, soluble GST–N-htt(Q51)(Ub\(_n\)) and aggregated PY-N-htt(Q51)(Ub\(_n\)) (but not nonubiquitinated forms; not depicted; Bennett et al., 2005) also inhibited \(^{35}S\)-PY-Sic1(Ub\(_n\)) degradation with inhibitory constants of 150–215 nM (Fig. 3 F), demonstrating that 26S inhibition is associated with the presence of a poly-Ub chain on these proteins but is independent of their aggregation state. TEV-cleaved GST–N-htt(Q51)(Ub\(_n\)) aggregates, retained on a 0.2-µm cellulose acetate filter, were immunoreactive with antibodies to Ub and S protein, confirming the presence of polyubiquitinated N-htt aggregates (Fig. 3 G). The similar \(K_i\) values for MPC(Ub\(_n\)), GST–PY-N-htt(Q51)(Ub\(_n\)), and aggregated PY-N-htt(Q51)(Ub\(_n\)) support the conclusion that neither the presence of a polyglutamine tract nor the aggregation state influences the ability of a polyubiquitinated protein to inhibit the degradation of other ubiquitinated proteins by 26S in vitro. Therefore, our data lead to the conclusion that, at least in vitro, proteasomes are neither choked nor clogged by N-htt but leave open the possibility that ubiquitinated N-htt aggregates can compete with other ubiquitinated substrates for 26S if these species accumulate to sufficient levels relative to other 26S substrates.

Stabilization of UPS reporters is not caused by direct competition for 26S proteasome by ubiquitinated N-htt

It is possible that ubiquitinated forms of N-htt could accumulate to sufficiently high concentrations in cells expressing polyglutamine-expanded N-htt(Qn)–chFP to be able to outcompete other poly-Ub–tagged substrates, including highly expressed UPS reporters. However, if ubiquitinated N-htt were to act as a simple competitor for binding of other ubiquitinated proteins to proteasomes, the concentration dependence of Ub\(^{76V}\)-GFP stabilization would not be expected to exhibit such a steep threshold as is observed in Fig. 1. To test this, we examined the effect of expression of one short-lived fluorescent proteasome substrate on the accumulation of another (Fig. 4 A). We found that expression of GFP-CL1 or cODC-GFP resulted in accumulation of chFP-CL1 in a concentration-dependent manner but exhibited no sharp threshold, as reflected by the nearly linear relationship of the fluorescence profiles of the two proteins when represented on a double logarithmic plot (Fig. 4 A, inset). This suggests that these two UPS substrates must compete with each other for one or more limiting UPS component. Because the two CL1-tagged proteins share the same degron, they might compete with each other for limiting Ub conjugation machinery, but the observation that a Ub-independent proteasome substrate (cODC-GFP) also competes for degradation of a Ub-dependent substrate (chFP-CL1) strongly suggests that the competition between these two reporters is at the level of proteasome binding, as the ODC degron functions as a molecular mimic that competes directly with Ub chains for binding to the 19S cap (Zhang et al., 2003). Therefore, competition for limited substrate binding capacity is a potential mechanism by which 26S activity might be reduced by a highly expressed protein, such as N-htt(Qn)–chFP, only to the extent that such a protein was modified with poly-Ub chains or a Ub mimic, such as the ODC degron, and whether this Ub-conjugated species was present at sufficient levels relative to other 26S substrates to be effective.

We therefore used absolute quantitative (AQUA)–MS (Gerber et al., 2003; Kirkpatrick et al., 2005) to determine the extent to which N-htt is ubiquitinated and the fraction of cellular poly-Ub conjugated to N-htt in FACS-sorted cells expressing N-htt(Q91)–chFP at levels above (high sort [HS]) or below (low sort [LS]) the threshold at which we observe accumulation of Ub\(^{76V}\)-GFP (Fig. S5 A). Total N-htt levels in lysates of these cells reflected this enrichment, increasing nearly 20-fold from 4.64 ± 1.04 pmol/mg in the LS population to 85.14 ± 17.59 pmol/mg in HS cells (Fig. 4 B). Poly-Ub conjugates were isolated by hP2UBA affinity capture (Bennett et al., 2007) from the same lysates and analyzed by AQUA-MS to determine the abundance of N-htt(Q91)–chFP present in the poly-Ub–associated
levels of N-htt in which we observe elevated levels of Ub chains are covalently linked to N-htt in the cells expressing the highest data). Therefore, only a minor fraction of total poly-Ub chains analysis in cell lines and R6/2 transgenic mice (unpublished data). In these experiments or in MS/MS we have never detected any N-htt peptides containing a diglycin covalent Ub signature in these experiments or in MS/MS. This technique does not distinguish between the amount of covalent N-htt–Ub conjugates. Indeed, the Ub AQUA-MS approach to assess the fraction of the total soluble N-htt pool that is associated with Ub, which, as in the previous paragraph, overestimates the actual fraction of covalent N-htt–Ub conjugates. This analysis revealed that of the 85.14 ± 17.59 pmol/mg total N-htt, at most 0.57± 0.08 pmol/mg (0.7%) was Ub associated (Fig. 4 B), indicating that soluble N-htt, in cells sorted to express the highest levels of this protein, is not ubiquitinated to a significant extent.

**N-htt and Ub are recruited to IBs independently**

To reconcile the persistent presence of elevated poly-Ub chains associated with N-htt inclusions in this and previous studies (Bence et al., 2001; Bennett et al., 2005), with the AQUA-MS data indicating little or no covalent Ub modification of N-htt, we used real-time fluorescence video microscopy to assess the temporal and spatial relationship between N-htt aggregation and Ub recruitment to IBs. Because HEK293 cells are not optimal for live-cell imaging experiments, we performed these experiments in U2OS cells. Control experiments in U2OS cells stably expressing UbG76V-GFP verified that the relationship between UPS reporter accumulation and N-htt expression in these cells is similar to that observed in HEK293 cells (Fig. S5 B). N-htt(Q91)-chFP fluorescence was initially diffusely distributed throughout the cytoplasm, and the overall fluorescence intensity increased linearly after transfection (Fig. 5 A and Video 1). The IB appeared as an initially dim spot of chFP fluorescence detectable above the background of diffuse fluorescence. Once it appeared, the diameter and intensity of this spot increased rapidly, and within 25 min, the vast majority of diffuse chFP fluorescence had coalesced into these IBs could also be visualized as phase-dense spots that became evident in parallel with recruitment of fluorescent N-htt. Quantification of total cellular chFP fluorescence in individual N-htt(Q91)-chFP–expressing cells at the time point immediately before the first appearance of detectable fluorescent puncta revealed that puncta formation occurred when total N-htt(Q91)–chFP concentration exceeded a threshold of total abundance of 431 ± 49 (mean ± SEM; Fig. 5 B). In cells expressing N-htt(Q47)-chFP, a similar threshold behavior was observed; but the threshold occurred at significantly (P < 0.001) higher total N-htt concentration (1,510 ± 81 a.u.), consistent with the relationship between N-htt concentration and Ub association with pure N-htt fragments in vitro. IBs were never observed in cells expressing N-htt(Q25)-chFP. The UbG76V-GFP reporter (Fig. S5 A), making it unlikely that polyubiquitinated N-htt represents a sufficient fraction of total poly-Ub conjugates to effectively outcompete other substrates.

We also used the AQUA-MS approach to assess the fraction of the total soluble N-htt pool that is associated with Ub, which, as in the previous paragraph, overestimates the actual fraction of covalent N-htt–Ub conjugates. This analysis revealed that of the 85.14 ± 17.59 pmol/mg total N-htt, at most 0.57± 0.08 pmol/mg (0.7%) was Ub associated (Fig. 4 B), indicating that soluble N-htt, in cells sorted to express the highest levels of this protein, is not ubiquitinated to a significant extent.

**Fig. 4. Stabilization of proteasome reporters by aggregation-prone N-htt is not caused by direct competition for 26S.** (A) Short-lived proteasome substrates, but not N-htt, compete for degradation of chFP-C1 degradation. HEK293 cells stably expressing chFP-C1 were transfected with GFP-C1, cODC-GFP, GFP, N-htt(Q25)-GFP, or N-htt(Q91)-GFP and analyzed by two-color flow cytometry. (Inset) Double logarithmic plot shows that cODC-GFP and GFP-C1 data fit well with a linear least-squares regression (r² > 0.98), whereas N-htt(Q91)–GFP does not. The data shown are from a single representative experiment out of two independent repeats. (B) Concentration of Ub, N-htt, and Ub-associated N-htt in HEK293 cells sorted on the basis of N-htt(Q91)–chFP fluorescence intensity. HEK293 cells stably expressing UbG76V-GFP and transfected with N-htt(Q91)–chFP were sorted based on high and low N-htt(Q91)–chFP fluorescence intensity followed by Ub AQUA-MS analysis to determine the amount of each species in the two sorted populations. All data are represented as means ± SEM (n = 3).
puncta (Fig. 6 A, 6- and 9-min time points). YFP-Ub became recruited to the nascent IBs only after a delay (average time of 43 min)—a time at which the IB had reached its mature size. These data demonstrate the N-htt molecules that are recruited into IBs are not conjugated to Ub and that Ub conjugation is not required for IB formation.

UPS reporter accumulation is delayed relative to IB formation

To assess the spatial and temporal relationship between IB formation and stabilization of UPS reporters, we used real-time single-cell analysis in a U2OS cell line stably expressing YFP-Ub to monitor changes in Ub distribution during the process of N-htt IB formation (Fig. 6 A and Video 2). Ub molecules bearing N-terminal epitope tags or protein fusions can functionally complement deletion of chromosomal Ub in yeast (Ellison and Hochstrasser, 1991), and in mammalian cells, GFP-Ub fusions have been used to monitor the localization and trafficking of intracellular Ub conjugates (Qian et al., 2002; Dantuma et al., 2006). We observed that YFP fluorescence was diffusely distributed in cells before IB formation but, strikingly, was distinctly excluded from the initial N-htt(Q91)–chFP puncta (Fig. 6 A, 6- and 9-min time points). YFP-Ub became recruited to the nascent IBs only after a delay (average time of 43 min)—a time at which the IB had reached its mature size. These data demonstrate the N-htt molecules that are recruited into IBs are not conjugated to Ub and that Ub conjugation is not required for IB formation.

Figure 5. IBs form rapidly in a concentration- and repeat length–dependent manner. (A) U2OS cells were transiently transfected with N-htt(Q91)–chFP, and formation of IBs was followed by time-lapse fluorescence video microscopy at 6-min intervals. Note the absence of diffuse mCherry fluorescence outside of the IB at the 30-min time point [Video 1]. The arrowheads indicate the site of a forming IB. Bars, 25 µm. (B) Threshold for IB formation. Cells transiently transfected with N-htt(Q91)–chFP or N-htt(Q47)–chFP were monitored by time-lapse fluorescence video microscopy as in A. Total chFP fluorescence intensity in individual cells in the frame just before the first detectable IB was quantified and plotted. The data shown are from a single representative experiment out of two independent repeats.
Figure 6. **Ub recruitment and UPS substrate stabilization are delayed after IB formation.** (A) U2OS cells stably expressing YFP-Ub were transiently transfected with N-htt(Q91)–chFP and followed by time-lapse two-color fluorescence video microscopy (Video 2). Sequential frames show the formation of the N-htt IB (left hand column) and the complete recruitment of chFP fluorescence by 33 min. Note that during the initial formation of the IB (6- and 9-min time points), YFP fluorescence is excluded from the IB and is recruited there only after a ~40-min delay. The arrowheads indicate the site of the forming IB. (B) U2OS cells stably expressing UbG76V-GFP were transiently transfected with N-htt(Q91)–chFP and monitored by time-lapse two-color fluorescence video microscopy at 20-min intervals. Selected frames of the recorded image series (Video 3) are presented. The outlined cell formed an IB at t = 0, and UbG76V-GFP became detectably stabilized in the cell after an ~17-h delay. Bars, 25 μm.
Ub<sup>G76V</sup>-GFP. These cells were transiently transfected with N-htt(Q91)–chFP, and double-positive cells were continuously monitored at 20-min intervals for >72 h (Table S1). Ub<sup>G76V</sup>-GFP stabilization was never observed in cells lacking IBs. However, we observed Ub<sup>G76V</sup>-GFP stabilization in 8/33 (24%) cells that formed IBs (Fig. 6 B, Video 3, and Table S1). This stabilization occurred on average 22.8 ± 4 h after IB formation, revealing a significant delay between IB formation and UPS substrate stabilization. The cells that never exhibited Ub<sup>G76V</sup>-GFP stabilization died, on average, within 12 h after IB formation, suggesting that the low frequency of cells that accumulate Ub<sup>G76V</sup>-GFP is a reflection of the poor survival of cells burdened with N-htt above the critical concentration at which IBs form. Thus, stabilization of Ub<sup>G76V</sup>-GFP and Ub-R-GFP (unpublished data) is a late event after IB formation in cells expressing high levels of aggregated N-htt; most cells experiencing a high burden of N-htt aggregation die before accumulating this reporter.

Discussion

Proteasomes are not directly impaired by aggregated N-htt

The data in the present study establish that polyglutamine fragments with covalently attached poly-Ub can inhibit 26S activity in vitro. However, because the <i>K<sub>i</sub></i> for N-htt fragments is independent of polyglutamine length and aggregation state and is similar to the <i>K<sub>i</sub></i> of an unrelated polyubiquitinated protein, MPC, these data suggest that ubiquitinated N-htt, whether aggregated or not, does not choke or otherwise noncompetitively inhibit 26S.

Thus, N-htt expression can directly inhibit proteasome activity only to the extent that ubiquitinated forms of the protein are able to accumulate to concentrations approaching or in excess of other polyubiquitinated species with which they compete for binding to 26S. Although some N-htt exon 1 can be recovered from metal affinity chromatography of extracts of cultured HeLa cells or striatal neurons transfected with His<sub>T</sub>-tagged Ub, the fraction of N-htt that was Ub associated (which does not necessarily reflect covalent ubiquitination of N-htt) was exceedingly low (Steffan et al., 2004), suggesting that the vast majority of N-htt exon 1 expressed in cells is not modified with poly-Ub, a finding that is confirmed by our AQUA-MS results and by the distinctly different kinetics with which N-htt and Ub are recruited to IBs and the complete exclusion of Ub during IB formation. We conclude that N-htt is neither a direct competitive nor a noncompetitive inhibitor of 26S.

Why do Ub conjugates accumulate in neurodegenerative disease?

The molecular genetics of dominantly inherited neurodegenerative diseases like HD strongly support a gain-of-function mechanism in which the disease-causing mutation—for example, expansion of the Gln repeat in N-htt—causes the disease-encoded protein to adopt one or more nonnative conformations that are toxic to cells (Hatters, 2008; Finkbeiner, 2011). In our experiments, the sharp transition between diffuse localization of cytoplasmic N-htt(Qn)–chFP and its rapid recruitment to IBs is observed when total N-htt(Qn)–chFP accumulates above a critical concentration threshold. Dependence of this critical concentration upon the length of the Gln tract argues strongly that this transition is caused by the intrinsic capacity of this protein to form organized, fibrillar aggregates. However, because molecular chaperones, including Hsp70/Hsp40, which constitute part of the cell’s proteostasis machinery, are able to suppress the aggregation of polyglutamine-expanded proteins in vitro and in vivo (Warrick et al., 1999; Chan et al., 2000; Jana et al., 2000; Kazemi-Esfarjani and Benzer, 2000; Howarth et al., 2007), the critical concentration for N-htt(Qn)–chFP aggregation in any given cell must be determined by both the absolute concentration of the aggregation-prone species and the capacity of the complement of molecular chaperones to resist this thermodynamically driven process. Therefore, the point at which diffusely localized N-htt transitions to IBs must represent the point at which the production of aggregation-prone N-htt species can no longer be offset by the cellular proteostasis network. At this point, preoccupation of one or more critical chaperones by the ever-increasing burden of N-htt production erodes the capacity of the cellular machinery to sustain the folding of its normal complement of cellular substrates. This proteostatic collapse is observed in <i>Caenorhabditis elegans</i>, in which expression of aggregation-prone, disease-associated proteins, including polyglutamine-expanded N-htt fragments (Gidalevitz et al., 2006) or mutant Cu and Zn superoxide dismutase (SOD1) linked to amyotrophic lateral sclerosis (Gidalevitz et al., 2009), leads to pleiotropic disruption of the protein-folding environment, indicated by the emergence of temperature-sensitive mutant phenotypes in unrelated genes at permissive temperatures. Conversely, the presence of temperature-sensitive polymorphic variants of cellular genes dramatically enhances the aggregation and toxicity of N-htt and SOD1 mutants in worms (Gidalevitz et al., 2006, 2009).

Therefore, proteins that depend on the factors that become limiting as a result of the burden of maintaining N-htt solubility will be delayed in their ability to fold, and some of them will be targeted to the UPS for degradation. We propose that the increased flux of normal cellular chaperone clients that are diverted to the UPS eventually will lead to a “queue” of ubiquitinated proteins waiting their turn to be degraded by 26S, and it is these proteins, collectively, that compete with each other and with artificial fluorescent UPS reporters for degradation. Some of the excess undegraded Ub-conjugated proteins would be recruited to IBs, composed of a Ub-free core of aggregated N-htt. This model provides a plausible explanation for why the ability of cells to degrade unstable GFP reporters declines with increasing time of exposure to aggregation-prone mutant N-htt and why IBs are enriched in so many different unrelated proteins, including Ub, transcription factors, cytoskeleton proteins, nuclear pore complex components, kinases, enzymes, and cell cycle regulators (Suhr et al., 2001; Leverenz et al., 2007; Xia et al., 2008).

Our model is consistent with previous findings from a longitudinal study of primary striatal neurons in culture, which reveal that cell survival is negatively correlated with the level of diffuse N-htt and positively correlated with IB formation (Arrasate et al., 2004). Our observation that IBs form abruptly, recruiting
by amplification of the C-terminal 37 amino acids of ODC by PCR and cloned into pEGFP1 (Tokara Bio Inc.) using the HindIII and BamHI sites. Plasmids expressing N- htt(Q25)–chFP and N- htt(Q91)–chFP were created by inserting chFP (a gift from R. Tsien, University of California, San Diego, La Jolla, CA) into the BamHI site of the N- htt exon 1 plasmids described in this paragraph. The chFP–CL1 plasmid was created by insertion of chFP and the CL1 sequence into pcdNA3.1 (Invitrogen). pET3a-Ub and pET15b–Ub were provided by C. Pickart (Johns Hopkins University, Baltimore, MD) and D. Ratin (The Hospital for Sick Children, Toronto, Canada), respectively. pHUE and pHUsp2-cc were gifts from R. Baker (Australian National University, Canberra, Australia). Plasmids for GST-Rsp5 and GST−ΔC2Rsp5 were obtained from J. Hiubregtse (University of Texas, Austin, TX). The ΔC2Rsp5 coding sequence was PCR amplified and cloned into a pET28a vector (EMD) to yield a His6–tagged construct (pET28a–ΔC2Rsp5). The plasmid for expressing His6–PY–Sic1 was obtained from R. Deshaies (Howard Hughes Medical Institute, California Institute of Technology, Pasadena, CA). The PY–Sic1 coding sequence was PCR amplified and cloned into the pMAL-C2X plasmid (New England Biolabs, Inc.). Site-directed mutagenesis was used to introduce a Pro-Pro-Pro-Tyr sequence (PY motif) in front of the cyclin N100 sequence and a 17 'tag at the end of cyclin N100 to create pMAL–PY–cyclinN100–T7 (MPC). Plasmids expressing the original GST–N- htt fusion constructs containing either Q18 or Q51 were obtained from E. Wanker (Max Delbrück Center, Berlin, Germany). These constructs were modified by the addition of a TEV protease cleavage site between the GST and htt exon 1 coding regions (GST–N- htt(Qn)–S) or by the addition of a TEV protease cleavage site between the GST and htt exon 1 coding regions and the addition of an S tag C terminal to the htt exon 1 coding region (GST–N–htt(Qn); Bennett et al., 2005). Within these constructs, site-directed mutagenesis was used to insert a PY motif immediately after the TEV protease site to create plasmids encoding for GST–PY–N- htt(Q18)–S and GST–PY–N- htt(Q51)–S.

**Cell lines**

The GFP–CL1 (Bence et al., 2001), CFPΔf508-GFP (Johnston et al., 1998), and TCRα–GFP (DeLaBarre et al., 2006) cell lines were previously described. These cell lines were prepared by transfection of HEK293 cells followed after 48 h by selection of transformed cells by growth in G418. Also, the other stable HEK293 cell lines expressing the constructs described in this study were created by transfection, selected with G418, and cloned by limiting dilution. The temperature-sensitive ts20 Bally/C3T3 clone A31 fibroblast cell line (Kulka et al., 1988) was a gift from D.T. Madden (The Buck Institute for Research on Aging, Novato, CA). All cells were grown in DME with 10% animal serum complex, l-glutamine, and antibiotics.

**Flow cytometry and cell sorting**

Unless indicated, cells were harvested 72 h after transfection and analyzed with a flow cytometer (LSR II; BD) with a 488- and 535-nm laser (BD). To ensure a sufficient number of cells with elevated levels of the transfected protein, >200,000 cells were analyzed per condition in a typical experiment. To plot the level of the reporter protein versus the level of the transfected protein (described in Fig. S2), a set of 41 gates of equal width (on a logarithmic scale) was set up in the channel for the transfected protein. The mean compensated fluorescence of the reporter protein in each of these gates was calculated and plotted on the ordinate with the gate number (corresponding to the log of fluorescence intensity of the transfected protein) plotted on the abscissa. Each construct and condition in singly transfected HEK293 cells was expressed as a single-color control to compensate the spillover between chFP and GFP individually for each gate. Gates with <100 events were not included in the analysis. The data shown in Figs. 1, 2, and 4 are from single representative experiments out of a minimum of two independent repeats. Raw flow cytometry data were analyzed using FlowJo (version 8.8.6; Tree Star) software.

**Materials and methods**

**Plasmids**

The plasmids expressing GFP–CL1, N- htt–GFP, and N- htt exon 1 were described previously (Bence et al., 2001). In brief, the GFP–CL1 plasmid was created by ligating on oligonucleotide encoding ACKNWFSSHVFHIH into the GFP–CL1 plasmid (Tokara Bio Inc.). N- htt–GFP encodes htt exon 1 and contains mixed CAG/GCAA repeats fused to a C-terminal GFP tag in pcdNA3.1 (Invitrogen). N- htt exon 1 was created by removing GFP from N- htt–GFP and ligating a linker between the BamHI and XbaI sites. Plasmids expressing Ub–R-GFP and Ub–GFP were gifts from N. Dantuma (Karolinska Institutet, Stockholm, Sweden). The cDCD–GFP-expressing plasmid was created by E. Bennett (University of California, San Diego, San Diego, CA)
Protein purification
Recombinant human E1 enzyme was purchased from Boston Biochem. Recombinant Ub was expressed and purified as previously described (Kaiser et al., 2011). In brief, Ub was expressed from the pET30a-Ub expression vector in BL21 (DE3) cells upon induction with IPTG. Bacterial cells were harvested, resuspended in lysis buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 10% [vol/vol] glycerol, 1% [vol/vol] Triton X-100, 1 mM EDTA, and protease inhibitor cocktail [Complete, Roche]), lysed by french press, and clarified by centrifugation (SS-34 rotor; Sorvall; 18,500 rpm for 60 min). Glacial acetic acid was added dropwise with mixing on ice until the solution reached pH 4. The precipitate was pelleted by centrifugation (SS-34 rotor; 13,000 rpm for 20 min), and the supernatant was dialyzed against 25 mM sodium acetate, pH 4.5. The dialysate was purified using a column (HiTrap SP XL; GE Healthcare) and eluted with a 0–500 mM NaCl linear gradient. Fractions containing Ub were pooled and further purified by gel filtration chromatography on a column (Sephacryl S-200; GE Healthcare) with 50 mM Tris, pH 7.5, 500 mM NaCl, and 1 mM EDTA. Eluted fractions were dialyzed into 25 mM Hepes, pH 7.5, aliquoted, and stored at −80°C.

Recombinant Ubc4 was expressed and purified as previously described (Kaiser et al., 2011). In brief, Ubc4 was expressed from the pET15b-Ubc4 plasmid in BL21 (DE3) cells upon induction with IPTG. Bacterial cells were harvested by centrifugation, resuspended in lysis buffer (50 mM Tris, pH 7.5, 300 mM NaCl, 20 mM imidazole, and 10% [vol/vol] glycerol), and lysed by french press. The lysate was clarified by centrifugation, and the supernatant was batch bound to Ni-nitrilotriacetic acid resin for 2 h. The resin was washed five times with 10 column volumes of lysis buffer followed by 3 ml of TALON beads. The column was washed with lysis buffer supplemented with 250 mM imidazole. The eluate was dialyzed into 50 mM Tris, pH 7.6, 150 mM NaCl, 10% glycerol, and 1 mM EDTA, aliquoted, and stored at −80°C.

Recombinant Usp2-cc was expressed from the pHUsp2-cc plasmid in BL21 (DE3) cells upon induction with IPTG. Bacterial cells were harvested by centrifugation, resuspended in lysis buffer (50 mM Tris, pH 7.5, 300 mM NaCl, 20 mM imidazole, 10% glycerol, 0.2% Triton X-100, 1 mM PMSF, and EDTA-free protease inhibitor cocktail [Complete]), and lysed by french press. The lysate was clarified by centrifugation, and the supernatant was a batch bound to Ni-nitrilotriacetic acid resin for 2 h. The resin was washed five times with 10 column volumes of lysis buffer followed by 3 ml of TALON beads. The column was washed with lysis buffer supplemented with 250 mM imidazole. The eluate was dialyzed into 50 mM Tris, pH 7.6, 150 mM NaCl, 10% glycerol, and 1 mM EDTA, aliquoted, and stored at −80°C.

Recombinant Sps2-cc was expressed from the pHsps2-cc plasmid in E. coli and purified by metal affinity chromatography (TALON; Takara Bio Inc.). In brief, cell pellets were resuspended in 40 vol of 50 mM Tris-HCl, pH 7.6, 300 mM NaCl, 5 mM β-mercaptoethanol, and 1 mM EDTA, and lysed by sonication. The lysate was clarified by centrifugation, and the supernatant was bound to Ni-nitrilotriacetic acid beads. The beads were washed with buffer A without Triton X-100 and eluted with 400 mM imidazole. The eluted protein was dialyzed into 50 mM Hepes, pH 7.5, 150 mM NaCl, 30% glycerol, and 5 mM EDTA, aliquoted, and stored at −80°C.

The His6-tagged ΔC2Rsp5 protein was expressed from the pET28a-ΔC2Rsp5 plasmid in E. coli and purified by using metal affinity chromatography (TALON; Takara Bio Inc.). In brief, cell pellets were resuspended in 40 vol of 50 mM Tris-HCl, pH 7.6, 300 mM NaCl, 5 mM β-mercaptoethanol, 1 mM EDTA, and 1 mM DTT, and lysed by sonication. The lysate was clarified by centrifugation, and the supernatant was batch bound to Ni-nitrilotriacetic acid beads. The resin was washed five times with 10 column volumes of lysis buffer followed by 3 ml of TALON beads. The column was washed with 20 column vol suspension buffer without β-mercaptoethanol and eluted with 3 column vol of 50 mM Tris, pH 7.6, 300 mM NaCl, 200 mM imidazole, and 10% glycerol. The eluted protein was dialyzed into 50 mM Tris-HCl, pH 7.6, 150 mM NaCl, 30% glycerol, and 5 mM EDTA, aliquoted, and stored at −80°C.

The His6-tagged ΔC2Rsp5 plasmid in E. coli was grown in minimal media supplemented with [35S]methionine (Zhang et al., 2003). In brief, a 100-ml culture of E. coli was grown at 37°C in M9 minimal media to OD600 = 0.5, washed twice in buffer A, and resuspended in buffer A containing 0.4% glucose and 0.063% methionine assay media. After further incubation at 37°C for 30 min, protein expression was induced with 1 mM IPTG. After a 1-h incubation, 1 μCi [35S]methionine was added. After 10 min, 1 mM of unlabeled methionine was added to the culture and grown for an additional 10 min. Cells were harvested by centrifugation and lysed by 3x Freeze-thaw steps. The presence of 10 mM l-lysine and 100 μg/ml of lysozyme was measured (Sigma-Aldrich). Clarified lysate was then bound to TALON metal affinity resin for 1 h at RT. Resin-bound protein was incubated with His-tagged Usp2-cc. PY-Sic1 was released from the TALON beads upon cleavage with Usp2-cc, collected in the unbound fraction, and supplemented with 0.2 mg/ml BSA as a carrier. The final protein concentration was measured using a bicinchoninic acid protein assay (Thermo Fisher Scientific) relative to a BSA standard.

Substrate ubiquitination
0.05–0.2 mg/ml [35S]-PY-Sic1 and MPC were ubiquitinated for 16 h in the presence of 50 nM E1 (Boston Biochem), 2.4 μM E2 (His6-Ubc4), 2 μM E3 (GST-Rsp5), 300 μM Ub, and ATP buffer (40 mM Tris-Cl, pH 7.2, 2 mM ATP, 5 mM MgCl2, and 1 mM DTT). The reaction was then mixed with 0.5 vol glycerol and 0.2 ml of 20 mM NaOH and stored at 80°C. The reaction was then quenched with 3 ml of 50 mM Tris, pH 7.5, 1 mM EDTA, 50 mM NaCl, and 10% [vol/vol] glycerol. Proteasomes were eluted using a 50–800-mM NaCl linear gradient. Active fractions were pooled, enriched by sedimentation centrifugation as described in this paragraph, and finally resuspended in 20 mM Tris, pH 7.5, 1 mM EDTA, 50 mM NaCl, and 10% [vol/vol] glycerol. The concentration of 26S proteasomes was measured using a Bradford assay relative to a BSA standard.

In vitro Sic1 degradation reaction
The substrate degradation reaction was performed at RT in 50–100 μl of total volume containing 10 nM 26S, ATP buffer, and 100 nM 35S-Sic1 (Ub). 5-μl reaction aliquots were removed at various time points and separated on a 4–20% gradient gel followed by autoradiography. Alternatively, aliquots were added to 10% TCA or double-distilled H2O on ice for 30 min and 2 mg/ml BSA as a carrier. Samples were centrifuged at 12,000 g for 30 min, and the supernatant was removed for analysis by scintillation spectrometry. The percentage of degradation is determined by dividing the measured radio counts after 10 min by the radio counts before the addition of the ubiquitiqated GST-Rsp5. Similarly, 0.05–0.2 mg/ml GST-PY-Nhtt(Qn)S was ubiquitinated, except that 2 μM E3 (AC2Rsp5) was used, and 1 vol TALON metal affinity resin was needed to remove any autoubiquitiqated E3 enzyme. 35S-PY-Sic1 ubiquitination ([35S]-Sic1(Ub)) was monitored by SDS-PAGE and autoradiography. The ubiquitination of MPC (MPC(Ub)) was monitored by SDS-PAGE and detected with T7-HRP conjugate (Thermo Fisher Scientific) followed by chemiluminescence. The ubiquitination of GST–PY-Nhtt(Qn)S was monitored by separation on SDS-PAGE and detection with S-HRP conjugate (EMD) followed by chemiluminescence.

Proteasome inhibition assay
The proteasome degradation inhibition assay was set up in 50 μl total volume containing 10 nM 26S proteasomes, 100 nM 35S-Sic1 (Ub), and 0–500 nM MPC(Ub) in ATP buffer. 5-μl aliquots were removed at 5, 10, and 20 min for TCA precipitation, and scintillation counting was performed as described in the previous paragraph. The initial rate of degradation as a percentage of the control reaction is reported as the percent ratio of the initial rate of degradation of 35S-Sic1(Ub) in the presence of varying concentrations of MPC(Ub) and the initial rate of degradation of 35S-Sic1(Ub) in a control reaction with no MPC(Ub). The dependence of the initial rate of degradation on substrate concentration was plotted, and the data were fit to the Michaelis–Menten equation using SigmaPlot. The dependence of the initial rate of inhibition on substrate concentration was plotted, and the data were fit to the following equation for competitive inhibition using SigmaPlot: $v_i = \frac{v_0 [S]}{K_i + [S]}$, in which $K_i = \frac{[S]}{[S]}$; $v_0$ is the concentration of uninhibited inhibitor; $v_i$ is the initial rate in the presence of inhibitor (Thower et al., 2000).
In vitro formation of ubiquitinated N htt aggregates

The TEV protease-induced cleavage and aggregation of GST-N htt (Q51)-S and filter trap analysis were performed as previously described (Bennett et al., 2005). Aggregation of 0.15 mg/ml ubiquitinated GST-FY-N htt (Q51)-S was initiated by the addition of 1:1 (wt/wt) TEV protease and 1:1 (wt/wt) GST-N htt (Q51)-S in aggregation buffer [40 mM Tris-HCl, pH 7.2, 1 mM DTT, and 0.5 mM EDTA]. The aggregation reaction was incubated at 37°C and considered complete after 24 h. Filter trap analysis of ubiquitinated aggregates was performed using both the anti-Ub antibody (clone FK2; Enzo Life Sciences) and an S tag HRP conjugate followed by chemiluminescence. The degradation of 100 nM 35S-Sic1(Ub) in the presence of 0–625 nM ubiquitinated aggregates was assessed, and the Kt was determined as described in the previous paragraph.

Microscopy

HEK293 cells were transfected with the indicated N htt-chFP plasmid and grown on polyl-lysine–coated coverslips. 72 h after transfection, cells were fixed with 4% paraformaldehyde. Cells were imaged by epi-fluorescence microscopy on an inverted microscope (Axiovert 200M; Carl Zeiss) with a 100× oil lens, NA 1.4 (Carl Zeiss). Digital (12 bit) images were acquired with a cooled charge-coupled device (CoolSNAP HQ; Photometrics) and processed using MetaMorph software (Universal Imaging).

For live-cell imaging of transiently transfected cells U2OS cells, cells were plated in DME with 10% FCS and 5% CO2, 2% fetal bovine serum (FBS), and 2% N-ethylmaleimide, 2% SDS, and EDTA-free protease inhibitors (Complete). Cells were transfected with N-htt(Q91)–chFP. Cells were then transfected with N-htt(Q91)–chFP as described in the previous paragraph. Live-cell imaging was performed on an inverted microscope (Axiovert 200M) in a periscope chamber that was heated to 37°C. Plates were placed in a smaller internal chamber that was continuously perfused with 5% CO2. Cells were imaged by epi-fluorescence microscopy on an inverted microscope (Axiovert 200M) encased in a periscope chamber that was heated to 37°C. Plates were placed in a smaller internal chamber that was continuously perfused with 5% CO2. The entire set up was controlled by MetaMorph software (Universal Imaging).

For imaging of cells expressing YFP-Ub and Ub35S-GFP, cells were stably cotransfected with pEF-YFP-Ub or Ub35S-GFP with a vector that confers G418 resistance. Drug-resistant colonies were pooled, and expressing cells were sorted by FACS and further propagated. These cells were then transiently transfected with N htt(Q91)–chFP as described in the previous paragraph. Live-cell imaging was performed on an inverted microscope (Axiovert 200M) in a periscope chamber that was heated to 37°C. Plates were placed in a smaller internal chamber that was continuously perfused with 5% CO2. The set up was controlled by MetaMorph software (Universal Imaging).

Poly-Ub affinity capture and MS

For poly-Ub affinity capture, HEK293 cells stably expressing Ub35S-GFP were transfected with N htt(Q91)–chFP. Cells were then sorted by FACS. Drug-resistant colonies were pooled, and expressing cells were sorted by FACS and further propagated. These cells were then transiently transfected with N htt(Q91)–chFP as described in the previous paragraph. Live-cell imaging was performed on an inverted microscope (Axiovert 200M) in a periscope chamber that was heated to 37°C. Plates were placed in a smaller internal chamber that was continuously perfused with 5% CO2. The set up was controlled by MetaMorph software (Universal Imaging).

For poly-Ub affinity capture, HEK293 cells stably expressing Ub35S-GFP were transfected with N htt(Q91)–chFP. Cells were then sorted by FACS. Drug-resistant colonies were pooled, and expressing cells were sorted by FACS and further propagated. These cells were then transiently transfected with N htt(Q91)–chFP as described in the previous paragraph. Live-cell imaging was performed on an inverted microscope (Axiovert 200M) in a periscope chamber that was heated to 37°C. Plates were placed in a smaller internal chamber that was continuously perfused with 5% CO2. The set up was controlled by MetaMorph software (Universal Imaging).

Online supplemental material

Fig. S1 shows that the chFP tag does not alter the aggregation or UPS impairment phenotypes of htt. Fig. S2 explains the method of transformation of FACS data used to generate the plots in Figs. 1, 2, and 4. Fig. S3 shows experiments further characterizing the reporter lines used in this study. Fig. S4 accompanies Figs. 3 and shows the kinetics of radiolabeled Sic1 degradation by 26S proteasomes and controls for the production of ubiquitinated and aggregated htt(Q51) in vitro. Fig. S5 shows MS data accompanying Fig. 4 B and characterizes the cell line used in Figs. S and 6. Video 1 shows aggregation of N htt(Q91)–chFP in U2OS cells. Video 2 shows that N htt(Q91)–chFP aggregates recruit Ub. Video 3 shows that N htt(Q91)–chFP aggregates form many hours before impaired proteasomal degradation. Table S1 shows the event log for time-lapse video microscopy. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201110093/DC1.

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