Huntington’s disease is caused by expanded polyglutamine (polyQ) sequences in huntingtin, which procures its aggregation into intracellular inclusion bodies (IBs). Aggregate intermediates such as soluble oligomers are predicted to be toxic to cells yet because of a lack of quantitative methods, the kinetics of aggregation in cells remains poorly understood. We used sedimentation velocity analysis to define and compare the heterogeneity and flux of purified huntingtin with huntingtin expressed in mammalian cells under non-denaturing conditions. Non-pathogenic huntingtin remained as hydrodynamically elongated monomers in vitro and in cells. Purified polyQ-expanded pathogenic huntingtin formed elongated monomers (2.4 S) that evolved into a heterogeneous aggregate population of increasing size over time (100-6,000 S). However in cells, mutant huntingtin formed three major populations: monomers (2.3 S), oligomers (mode $s_{20,w} = 140$ S) and IBs (mode $s_{20,w} = 320,000$ S). Strikingly, the oligomer population in proportion of total huntingtin did not change over 3 days and had an unchanging size distribution despite continued monomer conversion to IBs, suggesting oligomers are rate-limiting intermediates to IB formation. We also determined how a chaperone known to modulate huntingtin toxicity, Hsc70 influences in-cell huntingtin partitioning. Hsc70 decreased the pool of 140 S oligomers but increased the overall flux of monomers to IBs, suggesting Hsc70 reduces toxicity by facilitating transfer of oligomers into IBs. Together, our data suggests huntingtin aggregation is streamlined in cells, and is consistent with the 140 S oligomers, which remain invariant over time, as a constant source of toxicity to cells irrespective of total load of insoluble aggregates.

The misfolding and aggregation of proteins is thought to underlie pathogenesis in a range of amyloid-related neurodegenerative diseases including Alzheimer’s, Creutzfeldt-Jakob and Parkinson’s diseases (1). Huntington’s disease (HD) is one example of a fatal neurodegenerative condition caused by mutations in the huntingtin gene that induce the huntingtin protein (Htt) to oligomerize into larger forms (2-5). The mutations lead to an expansion in an N-terminal polyglutamine (polyQ) sequence to beyond a HD threshold of 37 residues (6). Longer polyQ sequences correlate with faster aggregation rates and earlier ages of disease onset, which implicates the intrinsic aggregation potential as central to the mechanisms causing pathogenesis.

A key marker of pathology is the coalescence of mutant Htt into punctate intracellular reservoirs of aggregates known as inclusion bodies (IBs). Yet despite intensive research over the last 20 years, it is still not well understood how this process, nor aggregation more generally, relates to cellular dysfunction. While IB co-sequestration of other cellular components has been postulated as a mode of toxicity in HD (7,8), more recent evidence suggests that IBs are not themselves toxic, but are...
an active response by cells to sequester more toxic soluble forms (3).

While it now well established that SDS-insolubility of mutant Htt loosely correlates to inclusions, there is a dearth of knowledge to the kinetics of aggregation of native Htt complexes expressed directly in the cellular environment (9). Knowledge of this process is fundamentally important to understanding earliest molecular steps of pathogenesis and for determining how the cellular machinery engages with the mutant forms of Htt. Several studies examining mutant Htt in cellular contexts have unequivocally identified the presence of oligomers in the range of 5 – 80 nm (10-12). Other work has shown that SDS-insoluble oligomers accumulate in cell culture and transgenic mouse models of HD prior to visible inclusions (9). Yet insight into the full heterogeneity, quantity and flux of the soluble forms of Htt under non-denaturing conditions have been hampered by a lack of high resolution and unbiased approaches to directly characterize and quantify these forms (13).

We sought to investigate the aggregation kinetics of mutant huntingtin expressed in mammalian cells using fluorescence adapted sedimentation velocity (SV) analysis in the analytical ultracentrifuge. SV analysis is a powerful approach for defining complex molecular size distributions and hydrodynamic properties of soluble proteins including that of aggregating proteins (14,15). However, only recently have advances in analytical ultracentrifugation hardware enabled green fluorescent protein (GFP)-tagged proteins to be uniquely tracked in complex solutions by virtue of the GFP fluorescence (16). In principle, this allows selective detection of the GFP-tagged protein from all other cellular components and extends the utility of SV analysis from purified proteins into more natural biological complexes such as those in cell lysates. Recent studies have used this approach to measure the size and heterogeneity of fluorescently-labelled purified proteins in the presence of other macromolecules including serum, yet the approach is yet to be examined on endogenous proteins expressed directly in mammalian cells or on proteins that are known to aggregate intracellularly in neurodegenerative diseases, such as HD (17,18).

This technique was applied to determine the kinetics of aggregation of the exon 1 fragment of huntingtin (Httex1), which has been commonly used to model HD, expressed directly in neuroblastoma cells (19-22). The proteins were examined in non-denaturing buffers and examined with no pre-purification steps that may selectively remove molecules, such as large aggregates, from the sample. In comparison, we also investigated the aggregation kinetics of purified mutant Httex1 to investigate how the cellular machinery and environment mediates aggregate size and flux. We extended our analysis to probe how the cognate heat shock protein 70 chaperone, Hsc70, which has been implicated in mediating Httex1 toxicity and aggregation, binds to Httex1 and influences the heterogeneity of the intracellular pools of Httex1 when co-expressed with Httex1 in neuroblastoma cells (23).

**Experimental Procedures**

The protein expression and purification, Western blotting and confocal microscopy procedures are included in the **Supplemental material, Procedures.**

**Cloning and mutagenesis**- Httex1-EGFP cDNA was amplified by PCR from the pGW1-Httex1-EGFP template(3) with primers *Nde*I-Htt forward (5’ aaaccccatatgccacccctggaaaagctgatg) and *Xho*I-GFP end reverse (5’ gggtttctcgagttacttgtacagctcgtccatgcc). The PCR product was cloned into the *Nde*I/*Xho*I restriction sites of a modified pET28b vector (pet28-HMT) that encodes a 6×histidine–maltose binding protein (MBP)–TEV site multiple cloning site (gift of Dan Minor and Rebecca Howard, University of California, San Francisco). The EGFP was converted to monomeric CFP variant Cerulean by mutating residues Y66W, S72A, Y145A, N146I, M153T, V163A, H148D, and A206K (24) with the QuickChange Multi-site directed mutagenesis kit (Stratagene).

To generate mammalian expression constructs, the cDNA sequences of Httex1-EGFP was amplified by PCR from the pET28-HMT
constructs with primers Htt TOPO forward (5’ caccatggcgacccctggaaaagct) and EcoRI-GFP reverse (5’ aaaaagaattcttacttgtacagctcgtccatg). EGFP was converted to monomeric Emerald by mutations S72A, N149K, M153T, I167T, A206K using the QuickChange mutagenesis kit (Stratagene) (25). The PCR product was inserted into pcDNA3.2/V5/GW/D-TOPO via TOPO cloning (Invitrogen). The Httex1-EGFP construct was then transferred first to pDONR-221 vector and then to the pT-REx-Dest30 vector via Gateway recombination (Invitrogen). To create an expression vector for Emerald alone, Emerald cDNA was amplified from pT-REx Httex1-Emerald with primers SacII GFP forward (5’ aggctccgcggccgcccccttcaccatgctggtgagcaagggcgag gagctgttc) and EcoRI-GFP reverse and religated back into the pT-REx Httex1-Emerald vector via the SacII/EcoRI sites. Vectors expressing the Httex146Q-Cherry and Httex125Q-Cherry were gifts of S. Finkbeiner and A. Tsvetkov (Gladstone Institutes, San Francisco). A vector expressing β-galactosidase, pT-REx/GW-30/ lacZ was obtained from Invitrogen.

pCMV6-XL5-HSPA8 (Hsc70 gene; NM_006145) was purchased from Origene. The Hsc70 cDNA was amplified by PCR with the primers SacII Hsc70 forward (5’ aggtcctcggcgccgttcctacctatgtccagggacagggagt g) and BamHI link Hsc70 rev (5’ ggtcaaggagatccagtgtcggcagctttggtccagtcgttgc) and cloned into the pGEM T-vector (Promega). The Hsc70 cDNA was cloned into the SacII/BamHI sites of pDONR Httex1-Emerald via the SacII/BamHI sites and then transferred to the pT-REx-Dest30 vector by Gateway recombination (Invitrogen). This construct contains a similar linker region between Hsc70 and Emerald to a previously described construct that was validated to have normal chaperone activity (26).

The Y66L mutation was introduced into Emerald using the QuickChange site directed mutagenesis kit. This mutation removes a critical residue involved in fluorophore formation but does not alter the overall structure of the protein (27). All vectors were sequenced and verified for accuracy within the cDNA regions. The resultant protein sequences expressed off these vectors is shown in Supplemental material Fig S1.

Preparation of cells and lysates for analytical ultracentrifugation- 2.5×10^6 Neuro2a cells were plated on a T25 culture flask and maintained in Opti-Mem (Invitrogen) containing 10% fetal calf serum, 1 mM glutamine, 100 µg/mL penicillin/streptomycin. The following day, the cells were transfected with the 10 µg total plasmid DNA and 25 µL Lipofectamine 2000 according to the manufacturers instructions (Invitrogen). Media was replaced daily. For the timecourse studies, cells were rinsed with Dulbecco’s Phosphate Buffered Saline (PBS) and imaged on an AF6000 fluorescence microscope (Leica) with a HCX PL FLUOTAR 10.0×0.30 DRY objective. Data were collected with the GFP filter cube and recorded in 12-bit with a DFC360FX camera. Cells were harvested by rinsing in PBS before removal from the flask with a cell scraper in 4 mL PBS. Cells were pelleted (1,000 g; 2.5 min), resuspended in 1 mL PBS, transferred to a 2 mL microcentrifuge tube and pelleted again (1,800 g; 4 min). The pellet was snap frozen in liquid nitrogen and stored overnight at −80 °C. The pellet was resuspended in 1 mL chilled lysis buffer (20 mM Tris pH 8.0, 2 mM MgCl2, 1% (v/v) Triton X-100, EDTA-free protease inhibitor cocktail tablets (Roche; 1 tablet per 10 mL solution), 20 U/ml Benzonase (Novagen)) and extruded through a 27 G syringe needle 25 times on ice. NaCl concentration was adjusted to 150 mM. Single-use aliquots were snap frozen in liquid nitrogen and stored overnight at −80 °C. Total cellular protein concentrations were determined in triplicate by a bicinchoninic acid (BCA) protein assay (Thermo Scientific) with bovine serum albumin as the standard.

Analytical ultracentrifugation- For the recombinant proteins, proteins were prepared at 0.7 µM in diluent buffer (50 mM Tris, 150 mM NaCl, 12.5 mM imidazole, 1 mM tris(2-carboxyethyl)phosphine, 0.1% (w/v) sodium azide). For the lysates, samples were adjusted to 0.2 or 0.5 mg/mL total protein concentration and zero or 2 M sucrose concentrations with lysis buffer (containing 150 mM NaCl) and freshly prepared 3 M sucrose, added by weight. Sucrose
stock solutions were made by heating sucrose with water and stirring until dissolved and were calibrated for concentration and density by refractive index at 20 °C (28). Final sucrose concentrations were verified in the sample by refractive index.

Samples were loaded into 2-channel Sapphire-window charcoal/epon sedimentation velocity cells (Beckman-Coulter) with 50 μL FC-43 heavy oil (Fluorinert). Cells were placed in a pre-chilled 8-hole AnTi rotor (Beckman-Coulter) and equilibrated to 7 °C in an XL-A Analytical Ultracentrifuge (Beckman-Coulter) fitted with a Fluorescence Detection System (Aviv) (16). Radial fluorescence scans were collected continuously at 50,000 rpm or 3,000 rpm using a 488 nm laser for excitation, and 520 nm cut-off emission filter and with the photomultiplier voltage kept constant in each experiment.

Data were analysed using SEDFIT (29). High speed data sets were fitted to c(s) distributions (29) and van Holde-Weischet analysis (30). Low speed data sets were fitted to ls-g*(s) distributions (14). For c(s) and ls-g*(s) (except the 2 M sucrose data), data were fitted with radial independent and in some cases, time independent noise, and with confidence levels (F-ratio) of 0.95. For samples containing 2M sucrose, data were fitted similarly except without radial or time dependent noise and with the baseline floated to account for the non-sedimenting material, and with the confidence level of 0.99. The meniscus and bottom positions were fixed. Sedimentation coefficients for samples lacking sucrose were corrected to $s_{20,w}$ using Sednterp (31). For samples containing 2 M sucrose, $s_{20,w}$ was calculated as described previously (32) using previously reported values for density and viscosity of concentrated sucrose solutions (33).

Statistics- Differences in IB frequency or Htt partitioning between cells expressing Htt$_{ex1}$46Q-Emerald/β-galactosidase with cells expressing Htt$_{ex1}$46Q-Emerald/Hsc70-Emerald(Y66L) over three days transient expression were evaluated using two-way analysis of variance (ANOVA) in SigmaPlot 11 software. Significance was evaluated using the Holm-Sidak method with p=0.05.

RESULTS

Aggregation kinetics of recombinant Htt exon 1. As a first step to understanding the aggregation kinetics, we examined recombinant Htt$_{ex1}$ fused to the monomeric cyan fluorescent protein, Cerulean, which is a derivative of the enhanced green fluorescent protein, EGFP (24). Non-pathogenic Htt$_{ex1}$-Cerulean (Htt$_{ex1}^{25Q}$) and a pathogenic form (Htt$_{ex1}^{46Q}$-Cerulean) were fused to maltose binding protein (MBP), which prevents aggregation until the MBP moiety is removed with TEV protease (Supplemental material Fig S2A). Upon cleavage with TEV, Htt$_{ex1}^{25Q}$-Cerulean did not aggregate, whereas the Htt$_{ex1}^{46Q}$-Cerulean formed sodium dodecyl sulphate (SDS)-insoluble, fibrillar aggregates over 72 h (Supplemental material Fig S2B).

Samples of recombinant protein were assessed using the analytical ultracentrifuge at different time points subsequent to TEV cleavage at a high angular velocity of 50,000 rpm, which is optimal for the detection of smaller mass species such as monomers and low-order oligomers (34). Freshly-TEV cleaved Htt$_{ex1}^{25Q}$-Cerulean and material incubated for 72 h formed a highly monodisperse population with a sedimentation coefficient centered at 2.3 S (Fig 1A,B). Sedimentation coefficients are proportional to the size, shape and density of the sedimenting species. Hence, 2.3 S best describes a monomer (37.4 kDa for Htt$_{ex1}^{25Q}$-Cerulean) with a frictional ratio ($f/f_0$) of 1.71, which suggests a reasonably elongated average conformation. This result is consistent with recent crystal structures of Htt$_{ex1}$ suggesting a high level of flexibility and conformational heterogeneity (35). Htt$_{ex1}^{46Q}$-Cerulean was also initially highly monodisperse as monomers upon TEV cleavage from MBP ($s_{20,w}$ =2.4 S; which for a 40 kDa monomer mass gives an $f/f_0$ of 1.72). After incubation at 37 °C for 72 h, Htt$_{ex1}^{46Q}$-Cerulean had aggregated into much larger mass species that mostly pelleted before data could be acquired (Fig 1A, inset). These data confirm that 25Q and 46Q counterparts of Htt$_{ex1}$-Cerulean both initially exist mostly as elongated monomers, however, only the
46Q variant assembles into large aggregates over 72 h incubation that sedimented too rapidly to be quantitatively analyzed under these experimental conditions.

We next determined whether we could define the molecular size distribution of the aggregated Httex146Q-Cerulean using SV analysis at lower centrifugal force. At 3,000 rpm, which we predicted would slow the rate of sedimentation of larger aggregates to within a detectable range as has been previously described for other amyloid fibrils (15), we found the Httex146Q-Cerulean to produce a family of sedimenting boundaries upon centrifugation (Fig 2A). The initial scan taken within 5 min of centrifugation (shown in red on Fig 2A) revealed no material had accumulated at the bottom of the cell prior to data acquisition, which indicated that the sedimenting boundaries encompass all the huntingtin molecules present in the sample. Fitting the boundaries to a sedimentation coefficient size distribution model optimal for large, non-diffusing particles, ls-g*(s), indicated a heterogeneous population of species ranging from \( s_{20,w} = 100-6000 \) S, slightly larger than the range of values observed with another amyloid-forming protein apoC-II (15). These sedimentation values correspond to amyloid-like fibrillar structures detected by electron microscopy (Supplemental material Fig S2B). Controls of freshly TEV-cleaved 25Q and 46Q Httex1-Cerulean and for the Httex125Q-Cerulean incubated for 72 h at 37 °C each showed no sedimentation, which confirmed an absence of aggregates and hence is consistent with these samples comprising just monomers (Fig 2A).

We next examined how the molecular size and heterogeneity changed for Httex146Q-Cerulean during the aggregation kinetics. Samples incubated at 37 °C for different time points revealed a systemic increase in the fraction of the material that partitioned into the sedimenting boundary of aggregates, as anticipated for aggregation (Fig 2B). The mode sedimentation coefficient and size heterogeneity of the sedimenting material also increased over 72 h, which is consistent with a systematic increase in fibril size and/or tangling (15) (Fig 2C).

A notable feature of the recombinant protein is the apparent lack of species between monomers and ~50 S oligomers, which roughly translates to a MW of 3,500,000 Da or about 80 Httex146Q-Cerulean subunits. A similar phenomenon has been observed in the SV analysis of another amyloid forming protein apoC-II, and these data can be explained by the small oligomers being highly transient, dynamic and/or infrequent under steady-state conditions (36).

Aggregation kinetics of Httex1 in cells. We next determined the molecular size of Httex1 expressed directly in the mouse neuroblastoma cell line Neuro2a. For the cell culture experiments, we replaced the Cerulean fluorescent protein moiety with Emerald, another monomeric EGFP variant that has a greater signal-noise than Cerulean with the 488 nm laser used in SV experiments and which we found necessary to distinguish Httex1 sufficiently from non-specific cellular autofluorescence. After two days transient transfection, Httex146Q-Emerald formed SDS-insoluble material and was distributed into IBs in approximately 35% of the cells as examined by confocal microscopy (Fig 3A,B). In contrast, the Httex125Q-Emerald did not form SDS-insoluble material and remained evenly distributed within the cytosol of all cells expressing Httex125Q-Emerald.

Lysates were prepared in solutions containing mild detergents to minimize structural perturbations of Htt conformations or intermolecular interactions. At the high angular velocity of 50,000 rpm to examine more specifically the lower-order oligomers, the Httex125Q-Cerulean formed a highly monodisperse distribution centered at \( s_{20,w} = 2.2 \) S, which indicated a population of extended monomers, similar to that observed with the recombinant counterpart (Fig 3C,D). The slightly lower value obtained in the lysate relative to recombinant protein (2.3 S versus 2.4 S) could reflect true differences in conformation or arise from differences in solution non-ideality from solvation, electrostatic interactions or protein concentration differences (16). Hence, as a control we also performed SV analysis of Emerald (without Httex1) expressed in the Neuro2a cells. Emerald had a
monodisperse population centered at $s_{20,w} = 2.3$ S, which for its predicted mass of 26.8 kDa, gives an $f/f_0$ of 1.37, indicating that in the same lysate conditions, Emerald has a more compact conformation than Httex1-Emerald (Fig 3D). The relative “compactness” can be related to the known crystal structure of the Emerald parent protein, EGFP, which describes a reasonably compact β-barrel (37), hence confirming the elongated conformation of Httex1-Emerald monomers in the cell.

The Httex146Q-Emerald displayed two discernable populations when analysed at the angular speed of 50,000 rpm. The first scan showed a rapidly moving boundary and a fraction of material already pelleted at the bottom of the column volume indicative of large molecular mass species (Fig 3C). Subsequent scans revealed the remaining material to be highly monodisperse, with a mode sedimentation coefficient of $s_{20,w} = 2.3$ S, indicative of an extended monomeric conformation ($f/f_0 = 1.79$; Fig 3D). For both the 25Q and 46Q counterparts, we noted a very minor fraction of the protein (less than 5%) to have a sedimentation coefficient at ~3–8 S, which likely represent complexes of Htt with other cellular proteins or ligands (Fig 3D, inset).

To assess the molecular size and heterogeneity of the larger aggregates of Httex146Q-Emerald, we performed SV analysis at the low angular velocity of 3,000 rpm. The first scan (red line, Fig 4A, left panel) indicated that a fraction of the material had immediately pelleted prior to data acquisition. This suggests the presence of very large macromolecular aggregates, far larger than that observed with the recombinant protein, where all material could be visualized sedimenting at 3,000 rpm. In addition to the large material immediately pelleting, approximately one third of the remaining Httex146Q-Emerald further evolved into a sedimenting boundary, which when fitted to an $ls$-$g^*(s)$ size distribution revealed a mode $s_{20,w}$ of 140 S. This population remained constant in size and heterogeneity between 2 and 3 days of expression (Fig 4C, left panel). SV analysis of Httex125Q-Emerald showed a lack of sedimentation altogether at 3,000 rpm, consistent with it comprising more than 95% as monomers (Fig 4B).

In an attempt to characterise the largest aggregates present, we decreased the rates of sedimentation at 3,000 rpm by increasing the solution viscosity from 0.01 P to 0.64 P with 2 M sucrose. Under these conditions, the largest material in the Httex146Q-Emerald sample sedimented as a well-defined boundary, whereas no material sedimented for the Httex125Q-Emerald as expected (Fig 4A,B, right panels). A notable feature of the sedimenting material was its high apparent noise contribution relative to the other species present in solution, which could be attributed to particles having a sufficiently large size to move in and out of the laser beam path during data acquisition. Nevertheless, the data could be fitted to an $ls$-$g^*(s)$ size distribution, which gave a broad size distribution with a mode $s_{20,w}$ of 320,000 S over 3 days expression (Fig 4C, right panel). While it is difficult to predict accurate masses from such large sedimentation coefficients, this value corresponds to a spherical proteinaceous particle of radius 0.62 μm assuming a partial specific volume typical of proteins (0.73 ml/g), and hence likely reflects the IBs. While the IBs in our Neuro2a cells had an average radius of 3.1 μm (± 1.0 s.d.; n=280 IBs; Supplemental material Fig S3), the estimate of the particle size from the sedimentation coefficient is likely a gross underestimate considering the IBs are likely far more asymmetrical and less-dense than our modelling of a proteinaceous sphere. To determine whether the high sucrose concentration might induce artifactual aggregation, we also examined the sedimentation of recombinant Httex146Q fibrils and Httex146Q monomers in 2 M sucrose at 3,000 rpm. Under these conditions, we saw no sedimentation, further consistent with the conclusion that that the sedimentation boundaries obtained in the lysate truly reflects the inclusion population. Similar experiments were performed in 0.5 M sucrose, and fibrils gave a similar sedimentation coefficient distribution ($s_{20,w}$) to samples without sucrose indicating that fibrils are not noticeably dissociated or altered in size under the experimental conditions and time frame of the experiments.

In context of the 320,000 S pool of Httex146Q-Emerald pool likely being the IBs, we hence
concluded that the pool of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald sedimenting at ~ 140S are oligomeric intermediates to IBs. 140 S corresponds to a spherical protein assembly of MW 7,700,000 Da, which is equivalent to approximately 200 Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald molecules or particles with radius 13 nm. The oligomer pool, while vastly smaller in absolute size than the IB pool, still has a large tail after the 140 S mode, which indicates a high degree of complexity in the range of oligomer assembly states. For simplicity, we refer herein to this population as the “140 S” oligomers.

We next examined how the proportion of the three pools of Htt\textsubscript{ex1}\textsuperscript{46Q} changed over 3-days transient expression in Neuro2a cell culture. Of the cells expressing Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald, the percentage containing IBs steadily increased over this time period (from 11% to 57%; p<0.001 two-way analysis of variance (38)) consistent with a progressive shift in the cellular localization from a diffuse cytosolic pattern to IBs (Fig 5A). Lysates were prepared and assessed using the combination of SV experiments described above to characterize the partitioning of the total population of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald molecules in the cells. The mass proportion of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald as monomers significantly decreased from 82% on day 1 to 28% on day 3 (Fig 5B; p<0.001; two-way ANOVA). Concurrently the proportion in IBs significantly increased from 1% to 53% (p<0.001; two-way ANOVA). Somewhat unexpectedly, the proportion of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald in the 140 S pool (16-19%) did not significantly change over the three day time course (p=0.546; two-way ANOVA).

\textbf{Chaperone-mediated transport of monomers into IBs.} Using our SV experimental platform we investigated how the cellular machinery might be involved in processing the Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald by co-expressing the cognate member of the heat shock protein 70 (Hsp70) family, Hsc70. Hsp70 proteins are components of the protein quality control network and prevent proteins from misfolding and aggregating by binding transiently to unfolded hydrophobic motifs in an ATP-dependent manner (39). Hsp70 members have been previously reported to co-localize with polyQ-related IBs and play a role in reducing toxicity of the polyQ-expanded proteins in animal and cell culture model systems (23,40-45). While some of these studies have reported Hsp70 members to suppress the formation of IBs, others found no effect on IBs, which led us to speculate that mediation of toxicity might lie in Hsc70 altering the relative population of oligomers or monomers in the cell regardless of the total number of IBs formed.

To detect Hsc70 fluorometrically, we fused Emerald to the C-terminus using the same paradigm as previously described, which has been validated to not interfere with its normal chaperone activity (26). Confocal microscopy revealed that Hsc70-Emerald was enriched at the periphery of IBs formed by Htt\textsubscript{ex1}\textsuperscript{46Q} fused to the red fluorescent protein Cherry, while it remained evenly diffuse through the cytosol when co-transfected with Htt\textsubscript{ex1}\textsuperscript{25Q}-Cherry (Fig 5D). To determine the effect of Hsc70 on rate of IB formation, we co-transfected Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald with Hsc70-Emerald(Y66L) whereby the Y66L mutation in Emerald renders Emerald non-fluorescent without altering its structure (27). When compared to cells co-expressing Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald with β-galactosidase as a negative control, we found a small but significant increase in the proportion of cells that contained IBs over three days (p<0.001 two way ANOVA; Fig 5A). Expression levels of the Htt\textsubscript{ex1} and Hsc70 moieties were equivalent in all the samples when examined by Western Blot probed with an anti-GFP antibody indicating that this difference was not due to variation in expression load (Fig 5D).

To examine the effect of Hsc70-Emerald(Y66L) on Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald molecular partitioning in detail, we performed SV analysis of the samples over a 3 day co-transfection time course. In accordance with the number of cells containing IBs, we found Hsc70-Emerald(Y66L) to significantly increase the proportion of Htt molecules in IBs compared to β-galactosidase over three days (Fig 5B; p=0.044, two way ANOVA). There was an apparent trend to Hsc70-Emerald(Y66L) decreasing the amount of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald as monomers over three days, yet this was not statistically significant (Fig 5B; p=0.154; two-way ANOVA). However, the proportion of Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald in the 140 S pool...
was slightly, but significantly, lower in cells expressing Hsc70-Emerald(Y66L) compared to β-galactosidase over the three days (a decrease in average proportion from 17.8% to 15.0%; ± 0.8 s.e., p=0.030, two-way ANOVA). The differences in 140 S partitioning between days was not significant for cells expressing β-galactosidase or Hsc70-Emerald(Y66L) (p=0.546; two-way ANOVA), indicating that while Hsc70 lowered the amount of Htt ex1 46Q-Emerald in oligomers, the oligomer population still remained static over time.

To further investigate the interaction of Hsc70 with Htt ex1 46Q, we measured the effect of Hsc70-Emerald(Y66L) on the molecular size and heterogeneity of Htt ex1 46Q-Emerald. Using a high-velocity (50,000 rpm) SV experiment, we found no effect of Hsc70-Emerald(Y66L) on the hydrodynamic properties of Htt ex1 46Q-Emerald monomers, which suggested that either there was no direct binding between these species, or the interaction was too transient to detect (Supplemental material Fig S4). Examination of lysates co-expressing Hsc70-Emerald with Htt ex1 46Q-Emerald(Y66L) also showed no evidence of Htt monomers, oligomers or IBs altering the hydrodynamic properties of Hsc70-Emerald, further consistent with Hsc70-Emerald at best binding highly transiently to all forms of Htt ex1 46Q and at worst not binding at all (Fig 5E−F). In light of the confocal data showing an enrichment of Hsc70 with the periphery of Htt ex1 46Q IBs, these findings point to Hsc70 facilitating the transfer of Htt ex1 46Q monomers to IBs by binding in a highly dynamic fashion.

**DISCUSSION**

Using analytical ultracentrifugation for quantifying molecular size and heterogeneity under non denaturing conditions, we identified several key findings. First, we found that purified mutant Htt self-associates from an expanded monomeric conformation into a heterogeneous population of aggregates that systematically increased in modal size over three days (50-6000 S). This is consistent with other studies showing that Htt ex1 forms amyloid-like fibrils and also that analytical ultracentrifugation can be used to measure fibrillization rates and fibrillar size heterogeneity (15,34,46). The finding that huntingtin monomers adopt an expanded conformation is also consistent with findings of others (35,47). However, our work extends this knowledge into the cellular environment whereby monomeric Htt ex1 remained in a remarkably similar expanded conformation, and did not appear to bind appreciably to other intracellular ligands (less than 5% of the total “monomeric” pool of mutant and non-mutant Htt appeared in complex with other species). It is noteworthy that these results contrast with recent reports that polyQ fragments form mechanically collapsed disordered structures (48,49), and the apparent discrepancy may be explained by the flanking residues of the polyQ sequence in Htt (notably the pro-rich region C-terminal to the polyQ sequence) promoting more extended, non-compact conformations. Another possibility we cannot exclude is that the Emerald/Cerulean fusion tag alters the structural properties of the Htt exon 1 sequence. Alternatively, it remains possible that a subset of “collapsed” conformations remain undetected if these forms cause the Cerulean/Emerald moiety to not fold properly.

A striking and novel finding from our study was that oligomers formed by mutant Htt in neuroblastoma cells were static in proportion of total Htt molecules and in their size heterogeneity despite continued conversion of monomers to IBs. Hence the 140 S oligomers may present a rate limiting step in the trafficking of small huntingtin aggregates to IBs through a series of highly compartmentalized intermediates (with a sharp mode of 140 S followed by an extended tail in the distribution extending up to approximately 1600 S). These species may denote specific complexes with different cellular machineries, or simply be larger oligomers and small fibrils that are constantly moved into the IBs via a rate limited mode of action. This result is supported by recent electron micrograph images of IBs in human brain, showing oligomeric structures of a similar size to the oligomers in our study (approximately 30-60 nm) located at the periphery of an IB (50). Similar sized structures ~30 nm in diameter have also been purified from Htt ex1-transfected cells or brain tissue of transgenic mouse models of HD (12,51). Larger fibrillar structures have also been reported
in IBs from \textit{ex vivo} HD brain samples (52), which could reflect the larger species (>500 S) within the 140 S pool that more closely match the sedimentation coefficients of recombinant fibrils.

Genetic and biochemical evidence have shown that IB formation is a regulated process driven by dynein-mediated retrograde transport to the microtubule organizing center, consistent with our results suggesting that small oligomers are formed in a regulated fashion (38,53,54). It remains to be examined whether the different sized oligomers within the 140 S pool contain components of trafficking or quality control machinery or indeed have the same structural architecture relative to the other forms within the pool and with small fibrils formed by purified Htt. An alternative possibility is that the oligomers are off-track to IB formation and present a parallel partition which may be handled by distinct cellular machineries. For example, it remains an intriguing possibility that a subset of monomers indeed do adopt a collapsed coil conformation as proposed (48,49), but that these forms are rapidly recruited into larger macromolecular complexes comprising other cellular machinery.

An important consequence of our results in light of studies showing IBs to correlate poorly with cellular toxicity, is that oligomers are clearly uncorrelated to the presence of IBs or monomers (3). More poignantly, if oligomers are indeed the most toxic molecular form as previously postulated (4-5,13,55), then this implicates whole populations of cells under a steady and constant duress from mutant Htt toxicity. Hence specifically mediating the pool of oligomers offers an attractive therapeutic target to modulate toxicity, and provides more specificity than current approaches targeting total inclusion/SDS insoluble protein levels or monomer levels.

Our results also showed that Hsc70 facilitated the flux of Ht\textsubscript{ex1}\textsuperscript{46Q} monomers to IBs, a result consistent with other findings in that Hsp70-family proteins alone do not prevent the aggregation of Htt\textsubscript{ex1} despite suppressing Htt toxicity (40-42). The binding appeared to be highly dynamic and transient, which is consistent with Hsp70-family proteins binding substrates by dynamic on-off cycling (56). Hsc70 has been suggested to bind polyQ-mediated IBs in a highly transient fashion (26,57). Thus, Hsc70 seems to act as a carrier in the traffic of Htt\textsubscript{ex1} to the IBs, yet we cannot determine from our studies whether this is through direct binding to monomers or is independent to the 140 S population. The efficacy of Hsc70 in the traffic and binding to Htt\textsubscript{ex1}\textsuperscript{46Q} may be limited by other components of the quality control network, such as Hsp40 co-chaperones, which have been shown to suppress IBs and to work in tandem with Hsp70 proteins to alter the morphology of Htt\textsubscript{ex1} aggregates (23,58).

In conclusion, we have used SV analysis to provide unprecedented insight to the molecular heterogeneity and hydrodynamic properties of all Htt molecules in the cellular milieu under non denaturing conditions and without pre-purification or pre-fractionation steps. Most noteworthy is the observation of a large pool of 140 S oligomers that are invariant in size and seem to be uncorrelated to the presence of monomers or IBs. This methodology will be useful for the study of parameters that mediate the aggregation pathway and for research in other neurodegenerative diseases that have been associated with oligomers, such as Alzheimer’s disease.

\textbf{REFERENCES}


FOOTNOTES
* We thank Ken Goldie for performing the electron microscopy experiments. This work was funded by grants to AFH (National Health and Medical Research Council program grant) and DMH (NHMRC project grant 566640). DMH is a Grimwade Fellow.
Abbreviations used are: polyQ, polyglutamine; IBs, inclusion bodies; HD, Huntington’s disease; SV, sedimentation velocity; EGFP, enhanced green fluorescent protein; Htt exon 1 fragment of huntingtin; MBP, maltose binding protein; PBS, Dulbecco’s Phosphate Buffered Saline; SDS, sodium dodecyl sulphate; f/f₀, frictional ratio; Hsp70, heat shock protein 70.
FIGURE LEGENDS

**Fig. 1** Hydrodynamic properties of recombinant Htt\textsubscript{ex1}\textsuperscript{25Q}-Cerulean monomers. **A)** Sedimentation velocity analysis of Htt\textsubscript{ex1}\textsuperscript{25Q}-Cerulean and Htt\textsubscript{ex1}\textsuperscript{46Q}-Cerulean at high speed (50,000 rpm). The initial data acquisition scans are shown in red and indicate all htt\textsubscript{ex1}-Cerulean to be in solution. The remaining scans at subsequent time points are shown in incrementing grayscale (note the scans may be overlapping). The data was fitted to a c(s) continuous size distribution, which best describes the behaviour of small, diffusing particles (residuals to the c(s) fit is shown in the lower panels). The inset shows the sedimentation behaviour of samples incubated for 72 h at 37 °C prior to data analysis. **B)** Sedimentation coefficient distributions based on the fits in (A) for the c(s) analysis (top panel) – note the overlapping data for the Htt\textsubscript{ex1}\textsuperscript{25Q}-Cerulean samples. An alternate mode of analysis, the van Holde-Weischet analysis, which qualitatively describes the heterogeneity of sedimentation coefficients (lower panel) similarly indicates a population at 2-2.5 S.

**Fig. 2** Kinetic assessment of aggregate size formed by recombinant Htt\textsubscript{ex1}-Cerulean aggregates by low speed centrifugation (3,000 rpm). **A)** Under these conditions only large aggregates form sedimenting boundaries while monomers do not sediment (resulting in a plateau). The first scan is shown in red, with subsequent scans in incrementing grayscale. The data for the 72 h incubated Htt\textsubscript{ex1}\textsuperscript{46Q}-Cerulean was fitted to a ls-g*\{s\} continuous size distribution, which best describes the behaviour of very large particles that have negligible diffusion (residuals to the fit shown in the inset). **B)** Percent of Htt\textsubscript{ex1}\textsuperscript{46Q}-Cerulean partitioned in the sedimenting boundary after different incubation times, reflecting a transfer of monomers to aggregates over time. **C)** Time dependent changes in aggregate size heterogeneity and sedimentation coefficients of aggregated Htt\textsubscript{ex1}\textsuperscript{46Q}-Cerulean were revealed by fits to ls-g\*\{s\} size distributions at the different time points.

**Fig. 3** Hydrodynamic assessment of Htt\textsubscript{ex1}-Emerald monomers in Neuro2a lysates by sedimentation velocity analysis. **A)** Western blot shows the expression levels over two days. Only Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald forms SDS-insoluble material over two days, indicative of aggregates. **B)** Confocal microscopy shows Htt\textsubscript{ex1}\textsuperscript{25Q}-Emerald to remain distributed evenly through the cytosol of Neuro2a cells after 2 days. In contrast, the Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald forms punctate IBs in many cells. **C)** High speed sedimentation velocity analysis (50,000 rpm) of Htt\textsubscript{ex1}-Emerald in the lysate (0.5 mg/mL total protein) after 2 days expression. The first scan is shown in red, with subsequent scans in incrementing grayscale. The first scan shows a fast pile-up of material at the bottom for Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald for a fraction of the material. Lower panels show residuals to fits to a c(s) distribution. **D)** The sedimenting boundaries for Htt\textsubscript{ex1}-Emerald dictates a highly monodisperse population (elongated monomers) when assessed by c(s) analysis (upper panel) or van Holde-Weischet analysis (lower panel). Zoomed-in rescaling of the c(s) distributions shows a minor fraction of material to sediment at 3-8 S (inset of upper panel).

**Fig. 4** Quantification of in-cell aggregate size heterogeneity with low speed and high viscosity (2 M sucrose) sedimentation velocity experiments (0.5 mg/mL total protein). **A)** Sedimentation velocity analysis of the Htt\textsubscript{ex1}\textsuperscript{46Q}-Emerald at low speed (3,000 rpm). The first scan is shown in red and in the absence of sucrose indicates a proportion of material that has already piled up on the bottom. The remaining sedimenting boundary (incrementing grayscale) was fitted to an ls-g\*\{s\} size distribution (residuals to fit shown in lower panels). Sedimentation was slowed by increasing the viscosity with 2 M sucrose. **B)** Sedimentation velocity analysis of Htt\textsubscript{ex1}\textsuperscript{25Q}-Emerald at low speed (3,000 rpm). Here, there is no sedimentation in the presence or absence of 2 M sucrose, indicative of a lack of aggregates. For comparison, lysates expressing β-galactosidase showed negligible signal intensity under the same photomultiplier voltage as Htt\textsubscript{ex1}\textsuperscript{25Q}-Emerald (purple scans) indicating negligible non-specific fluorescence contributions to the samples. **C)** ls-g\*\{s\} size distributions detected at low speed (3,000 rpm) for the oligomer boundary detected in the absence of sucrose (left panel) and IBs from the boundary detected in 2
M sucrose (right panel). Data shown is the mean (± s. d.) of fits to three independent preparations of lysates.

**Fig. 5** The influence of Hsc70 on the evolution of three discrete species of Htt<sub>ex1</sub><sup>46Q</sup>-Emerald in Neuro2a cells over 3 days transient expression. A) The percent of cells expressing Htt<sub>ex1</sub><sup>46Q</sup>-Emerald containing IBs prior to lysis for assessment by SV analysis. Htt<sub>ex1</sub><sup>46Q</sup>-Emerald was co-expressed with Hsc70-Emerald(Y66L) or β-galactosidase. Values are mean ± s.d. (n = 3 separate transfections). B) The relative population of 2.3 S monomers, 140 S oligomers, and 320,000 S IBs as assessed by SV analysis (Htt<sub>ex1</sub><sup>46Q</sup>-Emerald & β-galactosidase dashed line; Htt<sub>ex1</sub><sup>46Q</sup>-Emerald & Hsc70-Emerald(Y66L) solid line). Values are mean ± s.d. (n = 3 separate transfections) C) The localization of Hsc70-Emerald and Htt<sub>ex1</sub>-Cherry in Neuro2a cells after 3-days transfection (imaged by confocal microscopy). Hsc70-Emerald is enriched at the periphery of the IBs in cells expressing Htt<sub>ex1</sub><sup>46Q</sup>-Cherry, and remains diffuse in cells expressing Htt<sub>ex1</sub><sup>25Q</sup>-Cherry. Scale bar, 20 µm. D) Western blot probed with anti-GFP (upper panel) and anti-α-tubulin (lower panel) antibodies shows the expression levels and the presence of SDS-insoluble material over 3 days for the samples in A) and B). Samples shown include Emerald & β-galactosidase (3 day expression: i); Hsc70-Emerald & β-galactosidase (3 day expression: ii); Htt<sub>ex1</sub><sup>46Q</sup>-Emerald & β-galactosidase (iii); Htt<sub>ex1</sub><sup>46Q</sup>-Emerald & Hsc70-Emerald(Y66L) (iv); Htt<sub>ex1</sub><sup>46Q</sup>-Emerald(Y66L) & Hsc70-Emerald (v). For all combinations, equivalent masses of DNA were used in the transfections. E) c(s) size distributions of high speed (50,000 rpm) SV analysis showed no detectable interactions between Hsc70-Emerald and Htt<sub>ex1</sub><sup>46Q</sup>-Emerald(Y66L). F) Is-g*(s) size distributions of low speed (3,000 rpm) SV analysis showed no co-sedimentation of Hsc70-Emerald with Htt<sub>ex1</sub><sup>46Q</sup>-Emerald(Y66L). 140 S oligomers or 320,000 S IBs. The first scan is shown in red, last scan in blue and intermediate scans in grayscale (note many are overlapping due to lack of sedimentation).
Fig. 3

A

1 day | 2 days | 1 day | 2 days

SDS-insoluble aggregates

Htt_{ex1}^{25Q}.Emerald

Htt_{ex1}^{46Q}.Emerald

α-tubulin

25Q | 46Q

B

Htt_{ex1}^{25Q}.Emerald

Htt_{ex1}^{46Q}.Emerald

25 μm

C

Htt_{ex1}^{25Q}.Emerald

Htt_{ex1}^{46Q}.Emerald

Fluorescence intensity (arbitrary units)

Fluorescence profile

Residuals to Fit

Radius (cm)

D

Boundary Fraction

s_{20,w} (S)

46Q

25Q

Emerald alone
Fig. 4
Fig. 5

Panel A: Graph showing the percent of cells versus days post-transfection for Htt & β-gal and Htt & Hsc70.

Panel B: Graph showing the percent of species versus days post-transfection for Htt & β-gal and Htt & Hsc70.

Panel C: Images illustrating Httex146Q-Cherry and Httex125Q-Cherry, showing Emerald, Cherry, and Merge.

Panel D: SDS-insoluble aggregates, Hsc70-Emerald, Httex146Q-Emerald, Emerald, and α-tubulin.

Panel E: Graph showing relative frequency versus c(s) S20,w (S) for Hsc70-Emerald & β-gal, Day 1, Day 2, & Httex146Q-Emerald(Y66L).

Panel F: Graph showing fluorescence intensity versus radius for 2 M sucrose and No sucrose.
Tracking mutant huntingtin aggregation kinetics in cells reveals three major populations including an invariant oligomer pool
Maya A. Olshina, Lauren M. Angley, Yasmin M. Ramdzan, Jinwei Tang, Michael F. Bailey, Andrew F. Hill and Danny M. Hatters

J. Biol. Chem. published online May 5, 2010

Access the most updated version of this article at doi: 10.1074/jbc.M109.084434

Alerts:
- When this article is cited
- When a correction for this article is posted

Click here to choose from all of JBC's e-mail alerts

Supplemental material:
http://www.jbc.org/content/suppl/2010/05/05/M109.084434.DC1