Huntingtin aggregation kinetics and their pathological role in a Drosophila Huntington’s disease model

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ABSTRACT Huntington’s disease is a neurodegenerative disorder resulting from expansion of a polyglutamine tract in the Huntingtin protein. Mutant Huntingtin forms intracellular aggregates within neurons, although it is unclear whether aggregates or more soluble forms of the protein represent the pathogenic species. To examine the link between aggregation and neurodegeneration, we generated Drosophila melanogaster transgenic strains expressing fluorescently tagged human huntingtin encoding pathogenic (Q138) or nonpathogenic (Q15) proteins, allowing in vivo imaging of Huntingtin expression and aggregation in live animals. Neuronal expression of pathogenic Huntingtin leads to pharate adult lethality, accompanied by formation of large aggregates within the cytoplasm of neuronal cell bodies and neurites. Live imaging and Fluorescence Recovery After Photobleaching (FRAP) analysis of pathogenic Huntingtin demonstrated that new aggregates can form in neurons within 12 hr, while preexisting aggregates rapidly accumulate new Huntingtin protein within minutes. To examine the role of aggregates in pathology, we conducted haplo-insufficiency suppressor screens for Huntingtin-Q138 aggregation or Huntingtin-Q138–induced lethality, using deficiencies covering ~80% of the Drosophila genome. We identified two classes of interacting suppressors in our screen: those that rescue viability while decreasing Huntingtin expression and aggregation and those that rescue viability without disrupting Huntingtin aggregation. The most robust suppressors reduced both soluble and aggregated Huntingtin levels, suggesting toxicity is likely to be associated with both forms of the mutant protein in Huntington’s disease.

Huntington’s disease (HD) is an autosomal dominant neurodegenerative disorder and one of the first characterized members of a family of neurological diseases that result from expansion of a polyglutamine [poly(Q)] tract within the causative protein (Orr and Zoghbi 2007). HD is characterized by neurodegeneration and formation of neuronal intracellular inclusions, primarily in the striatum and cortex, leading to motor impairment, personality disorders, dementia, and ultimately death (Vonsattel et al. 1985; Portera-Cailliau et al. 1995). Currently, HD has no known cure and treatments focus on delaying HD-associated symptoms. The causative mutation in HD is expansion of a CAG tract beyond 35 repeats in exon 1 of the IT15 gene encoding Huntingtin (Htt) (Huntington’s Disease Research Collaboration 1993). Similar to other poly(Q)-repeat neurological disorders, abnormal protein conformation(s) secondary to poly(Q) expansion are central to HD pathogenesis (Scherzinger et al. 1997; Persichetti et al. 1999). The expanded poly(Q) Htt protein can exist in multiple states (Hoffner et al. 2005; Nagai et al. 2007), including aberrantly folded monomeric forms, oligomeric microaggregates, fibrill states, and larger inclusion body aggregates. It is currently unclear which form(s) of mutant Htt are pathogenic and how the abnormally folded protein causes neuronal toxicity.

Poly(Q) expansion leading to aggregation is a common theme in neurodegenerative disorders. Spinocerebellar ataxias (SCA1, SCA2, SCA3/MJD, SCA6, SCA7, and SCA17), spinal bulbar muscular atrophy (SBMA), and dentatorubral pallidoluysian atrophy (DRPLA) all involve poly(Q) expansion, aggregation, and neurodegeneration (Kimura et al. 2007). Evidence that aggregates are toxic is mostly correlative for...
these diseases, but several studies support the aggregation-toxicity hypothesis. The threshold of poly(Q) repeat number required for the in vitro aggregation threshold is similar to that required for disease manifestation (Davies et al. 1997; Scherzinger et al. 1999). Longer poly(Q) tracts have faster in vitro aggregation kinetics and result in earlier disease onset (Scherzinger et al. 1999). Similarly, treatments that suppress aggregation, including chaperone overexpression (Carmichael et al. 2000) and administration of small molecule aggregation inhibitors (Chopra et al. 2007), have been shown to decrease neurodegeneration. Live imaging demonstrates that Htt aggregates can sequester and alter kinetics of trafficked organelles and proteins such as synaptic vesicles (Sinadinos et al. 2009) and transcription factors (Chai et al. 2002). However, there is also evidence that aggregates may be inert or even neuroprotective. Medium spiny projection neurons of the striatum exhibit fewer Htt aggregates than striatal interneurons, yet are more vulnerable to neuronal cell death associated with transient expression of mutant Htt in cultured striatal neurons is inversely proportional to Htt aggregate formation (Arrasate et al. 2004), suggesting that inclusion body formation may decrease levels of other toxic forms of Htt and promote neuronal survival. There is also evidence suggesting that oligomers precede aggregate formation and are the toxic species in HD (Lam et al. 2008; Lajoie and Snapp 2010). These contradictory results in different cellular contexts and HD models have led to confusion over the toxicity of aggregates and, subsequently, over whether therapeutic approaches in HD should focus on reducing or enhancing aggregate formation.

To further analyze the link between aggregation and toxicity in a model system, we generated transgenic Drosophila that express an N-terminal poly(Q)-Htt fragment and displays aggregate formation, but no neuronal dysfunction or degeneration (Slow et al. 2005). Indeed, neuronal cell death associated with transient expression of mutant Htt in cultured striatal neurons is inversely proportional to Htt aggregate formation (Arrasate et al. 2004), suggesting that inclusion body formation may decrease levels of other toxic forms of Htt and promote neuronal survival. There is also evidence suggesting that oligomers precede aggregate formation and are the toxic species in HD (Lam et al. 2008; Lajoie and Snapp 2010). These contradictory results in different cellular contexts and HD models have led to confusion over the toxicity of aggregates and, subsequently, over whether therapeutic approaches in HD should focus on reducing or enhancing aggregate formation.

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Materials and Methods

Generation of Htt constructs
cDNAs for mRFP-HttQ15, mRFP-HttQ138, eGFP-HttQ15, and eGFP-HttQ138 were subcloned into EcoRI (blunt-end ligation) and KpnI sites of the pUAST expression vector. cDNA for eGFP-HttQ138-mRFP was subcloned into the XbaI site of the pUAST vector. HttQ15 and HttQ138 cDNAs were kindly provided by Ray Truant (Department of Biochemistry, McMaster University, Hamilton, ON, Canada). cDNA for HttQ96-GFP was kindly provided by David Housman (Center for Cancer Research, Massachusetts Institute of Technology) and subcloned into the KpnI and XbaI sites of the pUAST vector. Microinjection of constructs was performed by Genetics Services (Cambridge, MA).

S2 cell transfection and analysis
cDNAs for mRFP-HttQ15 and mRFP-HttQ138 were subcloned into the BamHI and EcoRI (blunt-end ligation) sites of the pSR11 S2 transformation vector. To generate constructs expressing mRFP-HttQ15-eGFP and mRFP-HttQ138-eGFP, mRFP-HttQ15 and mRFP-HttQ138 cDNAs were PCR amplified with a forward primer containing an EcoRI restriction site and a reverse primer containing a 3′ SalI site and subcloned into the pPL17 vector. Constructs were transfected with 50 μl cytofectene (Bio-Rad, Hercules, CA) into Drosophila S2 cells using the Bio-Rad liposome mediated transfection protocol. After 72 hr, 20-μl cell suspensions were fixed with 3.7% formaldehyde in PBT and mounted on slides with 50% glycerol. Images were captured with a Zeiss Pascal laser scanning confocal microscope (Carl Zeiss MicroImaging), using the accompanying Zeiss PASCAL software.

Glue secretion assay
Pupae reared at 25°C were isolated shortly after pupariation and placed on slides with the ventral side facing up, using double-sided tape. Visualization was performed on a Pascal confocal microscope (Zeiss).

Drosophila genetics and deficiency screen
Drosophila were maintained on standard medium at 25°C. The deficiency collection used in the screen was obtained from the Bloomington Stock Center. Males from deficiency lines on chromosomes II and III were crossed to C155; A37/CyO-GFP or C155; Df(3)3450/TM6 virgins, respectively. For the viability screen, F1 C155/y; Df/Balancer males were mated to homozygous mRFP-HttQ138 high-expression females and the number of F2 males and females was scored. For the

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aggregation screen, F1, C155/y; Df/Balancer males were mated to homozygous mRFP-HttQ138 low-expression virgins and live wandering third instar larvae expressing mRFP-HttQ138/Df were viewed under a fluorescent stereoscope (Zeiss) to assay changes in aggregation in larval salivary glands. For aggregation formation analysis, the CCAP-Gal4 driver was recombined with a tubulin-Gal80\(^a\) repressor to drive high-expression mRFP-HttQ138 in a temporally restricted manner. mRFP-HttQ138 expression was repressed at 19\(^\circ\) until the second instar stage, at which point larvae were moved to 30\(^\circ\) to induce expression.

**Adult viability analysis**

*Drosophila* viability assays were performed on white/C155, HttQ96-GFP/C155, mRFP-HttQ15/C155, mRFP-HttQ138/C155, and mRFP-HttQ138B/C155 flies by daily quantification of lethality for 100 females of each genotype. Flies were aged and UAS-eGFP-HttQ138-mRFP to C155 GAL4 driver strains mental Studies Hybridoma Bank) at 1:1000. To assay for icon) at 1:1000 or anti-Actin (JLA20 antisera; Develop-

Blots were incubated with mouse anti-Htt MAb2166 (Chem-

buffer and proteins were separated on 10% SDS

gels and transferred to nitrocellulose. Blots were incubated

with rabbit anti-GFP sc8334 (Santa Cruz Biotechnology) at

for 60 min for FRAP recovery curves at long timescales. An

Aggregates were photobleached using 100% laser power at 546 nm for
to two to five iterations through the entire z-plane until they
were at 35% of the original fluorescence intensity. Z-stack
recovery images were recorded at a rate of one per minute
for 60 min for FRAP recovery curves at long timescales. An
imaging rate of one frame per 10 sec was used for analysis of
how individual puncta interact with aggregates and for character-
izing aggregate formation events.

**Results**

A 588-aa N-terminal fragment of pathogenic human Htt reduces *Drosophila* life span

To explore pathogenic mechanisms in HD, we generated transgenic *Drosophila* that express 588-aa N-terminal fragments of human Htt with either a pathogenic poly(Q) tract of 138 repeats (HttQ138) or a nonpathogenic tract of 15 repeats (HttQ15). While several models of HD have focused on expression of the poly(Q)-containing first exon of Htt alone, the 588-aa fragment is truncated near a number of well-characterized caspase cleavage sites important in the generation of aggregate-forming Htt fragments (Kim et al. 2001; Wellington et al. 2002; Graham et al. 2006). Additionally, many sites of protein interaction that are lost in exon 1 constructs are conserved in the longer 588-aa fragment, including a region of well-conserved HEAT repeats thought to be involved in Htt binding to interaction partners such as HIP1, HAP1, and HIP14 (Harjes and Wanker 2003). Htt was fluorescently tagged with mRFP or eGFP at the N terminus or tagged at both ends with eGFP at the N terminus and mRFP at the C terminus (Figure 1A). For comparison, we generated a transgenic strain expressing exon 1 (81 aa) of reverse transcription kit (Applied Biosystems) according to the manufacturer’s protocol. PCR was carried out in triplicate for each of two independent total RNA samples per genotype in optical 96-well plates (Applied Biosystems). The reaction mixtures were as follows: 25 \(\mu\)l of 2\(\times\) Quantitec SYBR Green PCR Master Mix (QIAGEN), 300 \(nM\) forward primer, 300 \(nM\) reverse primer, and 5 \(\mu\)l of single-stranded cDNA (see above) in a total volume of 50 \(\mu\)l. A final dissociation step was carried out to evaluate product integrity, and reaction samples were run on a 1.2% agarose gel and stained with ethidium bromide. The primer sequenc-
es were as follows: Act88F (actin) forward 5’-ACTTCT GCTGGAAGGTGGAC-3’ and reverse 5’-ATCCGCAAGGATC TGATGC-3’.
the human Htt protein with a pathogenic 96Q repeat, fused to GFP at the C terminus (HttQ96-GFP). All constructs were expressed using the UAS-GAL4 system, allowing for temporal and tissue-specific control of transgene expression. To confirm transgene expression, strains were crossed to the neuronal driver elav-GAL4 (C155), and Htt expression in offspring was assessed through Western blot analysis with anti-human Htt antibodies (Figure 1B). No Htt expression is detected in control white strains crossed to C155, while mRFP-Htt, eGFP-Htt, and HttQ96-GFP lines all demonstrate abundant Htt expression. As expected, the product detected in HttQ15 strains lacking the expanded poly(Q) tract is smaller than that in HttQ138 or HttQ96 strains. These transgenic lines allow imaging of Htt aggregation in live Drosophila, providing a resource for following Htt dynamics in vivo.

To determine the effects of pathogenic and nonpathogenic Htt expression in Drosophila, viability profiles were generated for control and HttQ138-expressing animals. Pan-neuronal expression of mRFP-HttQ138 with C155 causes pharate adult lethality with <1% viable adult escapers. Expression of pathogenic Htt with a weaker elav-GAL4 driver results in viable adults that appear behaviorally normal at the time of eclosion. However, several days after eclosion, these adults begin to exhibit motor coordination defects and abnormal grooming behaviors, worsening with age and resulting in premature death. Similar defects occur at a later time in a separate mRFP-HttQ138 insertion line (mRFP-HttQ138B) expressing pathogenic Htt at a lower level, as well as in flies expressing the pathogenic HttQ96-GFP exon 1 protein (Figure 1C). These behaviors are not observed in mRFP-HttQ15-expressing or control Drosophila. To quantify the reduction in viability of pathogenic Htt-expressing lines, life span curves were generated for control adults and adults expressing mRFP-HttQ15, mRFP-HttQ138, mRFP-HttQ138B, or HttQ96-GFP (Figure 1C). The $T_{50}$ (age at which 50% of the culture has died) for mRFP-HttQ138 lines is dramatically decreased by >70% in comparison to controls. mRFP-HttQ138B expression results in a 30% decrease in $T_{50}$, indicating that lethality is correlated with the level of expression of the pathogenic protein. HttQ96-GFP lines also demonstrate a decrease in $T_{50}$ of 50%, suggesting that expression of the expanded poly(Q)-containing first exon of Htt is also toxic. Decreases in $T_{50}$ for all lines expressing fragments of the pathogenic Htt protein, but not the normal protein, indicate that expression of pathogenic Htt results in behavioral dysfunction and reduced life span in Drosophila. Although differences in genetic background may contribute to some of the effects on life span,
poly(Q) expansion in Htt consistently reduces life span in our analysis.

**Pathogenic Htt forms cytoplasmic aggregates in neuronal and nonneuronal cells in vivo**

A hallmark of HD is the formation of intracellular aggregates immunopositive for the pathogenic Htt protein. Aggregates have been found in the nucleus, cell body, and neurites in HD (Difiglia et al. 1997). However, it is unknown whether toxic Htt activity occurs in the nucleus, in the cytoplasm, or in both. To determine the intracellular distribution of our pathogenic and nonpathogenic transgenic Htt fragments, *Drosophila* S2 cells were transiently transfected with the mRFP-HttQ15 or mRFP-HttQ138 constructs and fixed cells were imaged by confocal microscopy. While mRFP-HttQ15 demonstrated diffuse cytoplasmic localization, mRFP-HttQ138 formed large, distinct cytoplasmic aggregates (Figure 2A). Neither protein localized to the nucleus.

To assess whether intracellular aggregates are formed in vivo by transgenic Htt proteins in *Drosophila*, UAS-mRFP-Htt strains were crossed to lines expressing the elav-GAL4 driver. As observed in S2 cells, the nonpathogenic mRFP-HttQ15 remained diffuse throughout the cytoplasm and neurites of neurons in both the CNS (Figure 2B) and the PNS (Figure 2D). In contrast, distinct Htt aggregates were observed throughout the cytoplasm and neurites in lines expressing the pathogenic mRFP-HttQ138 (Figure 2, C and E). To observe subcellular localization in other cell types, mRFP-HttQ15 and mRFP-HttQ138 were driven with a ubiquitous GAL4 driver (tubulin-GAL4). In nonneuronal cells, including epidermis (Figure 2F) and salivary glands (Figure 2H), mRFP-HttQ15 localized diffusely in the cytoplasm, while mRFP-HttQ138 formed cytoplasmic aggregates (Figure 2, G and I). Nuclear aggregates were not observed in any cell type. These results suggest that Htt fragments induce pathology primarily through a cytoplasmic localization in this *Drosophila* model.

**Mutant Htt causes defects in salivary gland glue secretion in *Drosophila***

To determine whether HttQ138 expression might cause defects in nonneuronal tissues, we tested whether Htt expression in larval salivary glands causes cellular dysfunction. We analyzed secretion of the GFP-tagged salivary gland glue protein Sgs3 (Biyasheva et al. 2001) in controls and animals expressing mRFP-HttQ15 or mRFP-HttQ138 (Figure 3). During normal pupariation, *Drosophila* secretes ecdysteroid-induced glue granules that mediate attachment of the developing pupal case to surfaces. While glue secretion is evident in both control and mRFP-HttQ15–expressing pupae (Figure 3, A and B), secretion is decreased in pupae expressing mRFP-HttQ138 (Figure 3C). Normal third instar larval salivary gland cells are filled with glue proteins (Figure 3D) that are depleted during pupariation (Figure 3E). In contrast, the pupal salivary glands of mRFP-HttQ138–expressing larvae retain glue (Figure 3F), suggesting salivary gland dysfunction mediated by the cytoplasmic accumulation of mutant Htt.

**The 588-aa fragment of pathogenic human Htt does not produce nuclear cleavage products in *Drosophila***

Mutant forms of Htt have been reported to undergo cleavage by caspases and calpains to generate smaller N-terminal fragments that can be observed in the nucleus and cytoplasm (Kim et al. 2001; Lunkes et al. 2002; Wellington et al. 2002; Gafni et al. 2004). To determine whether cleavage of the N-terminal 588 aa of Htt occurs in our *Drosophila* model to generate smaller fragments that might localize to the nucleus, S2 cells were transiently transfected with 588-aa Htt constructs labeled with eGFP at the N terminus and mRFP at the C terminus. Complete colocalization of the eGFP and mRFP signals was observed for both the normal eGFP-HttQ15-mRFP fragment (Figure 4A) and the pathogenic eGFP-HttQ138-mRFP fragment (Figure 4B). These results suggest that cleavage of Htt does not occur in the context of *Drosophila* S2 cells or, alternatively, that any cleaved N- and C-terminal fragments of Htt remain colocalized in the cytoplasm.

To assess whether cleavage and separation of N- and C-terminal fragments of Htt occur in vivo, we generated transgenic strains expressing the double-labeled pathogenic Htt fragment eGFP-HttQ138-mRFP. As observed in the S2 cell model, eGFP and mRFP signals colocalized in all tissues studied, including CNS neurons (Figure 4C), salivary gland cells (Figure 4D), and epidermal cells (Figure 4E), suggesting that pathogenic Htt is unlikely to be cleaved in vivo in *Drosophila*. Neither N- nor C-terminal fragments of Htt are observed in the nucleus. To further test whether cleavage occurs in vivo, we performed Western analysis with anti-Htt antibodies to probe for breakdown products that would result from cleavage of the protein. As shown in Figure 4F, no differential cleavage products were observed by Western analysis in head extracts prepared from animals expressing HttQ138-mRFP vs. those from animals expressing eGFP-HttQ138-mRFP. Thus, the pathogenic Htt-mediated toxicity seen in our *Drosophila* HD model does not appear to require Htt cleavage or nuclear entry and reflects an effect of the 588-aa fragment in the cytoplasm.

**Exon 1 of pathogenic Htt forms cytoplasmic and neuritic aggregates**

Many classic HD models express exon 1 of the mutant protein (Davies et al. 1997; Jackson et al. 1998; Krobitsch and Lindquist 2000; Tagawa et al. 2004), which is capable of forming inclusions postulated to play a role in HD pathology (Davies et al. 1997; Becher et al. 1998). To determine whether the in vivo subcellular localization of the 81-aa exon 1 fragment of pathogenic human Htt (HttQ96-GFP) differs from that of the 588-aa pathogenic Htt fragment in our *Drosophila* model, HttQ96-GFP–expressing third instar larvae were imaged using confocal microscopy. In both neuronal and nonneuronal cell types, HttQ96-GFP formed
Figure 2 Cytoplasmic aggregation of mRFP-HttQ138 in neuronal and nonneuronal tissues. (A) Htt localization in Drosophila S2 cells transiently transfected with mRFP-HttQ15 or mRFP-HttQ138. mRFP-HttQ15 is found diffusely throughout the cytoplasm, while mRFP-HttQ138 forms cytoplasmic aggregates. Bar, 2 μm for each panel. (B) Visualization of mRFP-HttQ15 (magenta) and GFP with a nuclear localization signal (nls) (green) in third instar larvae with transgene expression driven by C155. mRFP-HttQ15 is diffusely localized in the cytoplasm of CNS neurons in the ventral nerve cord. (C) Visualization of mRFP-HttQ138 (magenta) and GFP-nls (green) in CNS neurons of third instar larvae with transgene expression driven by C155. Unlike mRFP-HttQ15, mRFP-HttQ138 forms cytoplasmic aggregates throughout the cell bodies of ventral nerve cord neurons. (D and E) Visualization of mRFP-Htt in peripheral multidendritic neurons. While mRFP-HttQ15 exhibits diffuse cytoplasmic localization, mRFP-HttQ138 is found in cytoplasmic aggregates throughout the cell body and neurites. (F–I) Expression of mRFP-Htt (magenta) and GFP-nls (green) driven by the tubP-GAL4 driver in the epidermis (F and G) and salivary gland (H and I). In all cases, mRFP-HttQ15 is diffuse throughout the cytoplasm, while mRFP-HttQ138 forms cytoplasmic aggregates.
distinct cytoplasmic aggregates similar in appearance and localization to those formed by the 588-aa mRFP-HttQ138 protein. GFP-labeled aggregates are found in the cytoplasm of salivary gland cells (Figure 5B) and CNS (Figure 5D) and PNS (Figure 5G) neurons, while no aggregates are observed with expression of a UAS-GFP construct alone (Figure 5, A, C, and F). As observed with the 588-aa fragment, the exon 1 fragment also forms aggregates in axons (Figure 5E) and localizes at nerve terminals. The localization of both 588-aa and 81-aa pathogenic Htt fragments indicates that the neurodegenerative effects induced by these toxic Htt forms are independent of Htt nuclear accumulation in Drosophila. In addition, targeting sequences in exon 1 of Htt are sufficient to localize the protein to neurites in our model.

**Kinetics of HttQ138 aggregate formation**

To examine how Htt aggregates form, we visualized mRFP-HttQ138 dynamics in live larvae. We expressed mRFP-HttQ138 using CCAP-GAL4 that expresses in a single neuron per hemisegment (Park et al. 2003; Vomel and Wegener 2007). This driver allows single-cell resolution for assaying how soluble HttQ138 interacts with larger aggregates. Time-lapse confocal imaging in third instar larval motor neurons expressing mRFP-HttQ138 demonstrates that large aggregates are immobile over a 2-h imaging session. FRAP analysis reveals that HttQ138-positive photobleached aggregates continually add new HttQ138 particles, with a 40% recovery of original fluorescence within 50 min (Figure 6, A–C, Supporting Information, File S1). Two general types of recovery were observed. Larger HttQ138 aggregates tend to recover more quickly by trapping Htt particles (rightmost aggregate in File S1). A second class of aggregates recovers more slowly and displays a gradual and uniform increase in brightness (left three aggregates in File S1). To determine whether the rapid FRAP in axons was due to increased flux of HttQ138 mediated by delivery of new particles by fast axonal transport (FAT), we compared FRAP rates of aggregates in axons to those in neuronal cell bodies aggregates where HttQ138 movement is dictated largely by diffusion. We did not observe a significant difference in aggregation kinetics in these two compartments, indicating that axonal aggregation kinetics are not strictly dependent on FAT (Figure 6B).

We next examined the kinetics of de novo HttQ138 aggregate formation in our model in vivo. During our time-lapse

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**Figure 3** Salivary gland glue secretion is defective in Drosophila expressing mRFP-HttQ138. (A) Pupae were removed from vials of CS (A), Htt-Q15 (B), or HttQ138 (C) animals expressing the Sgs3 GFP-tagged glue protein. Confocal images of Sgs3-GFP fluorescence are shown in the left panels and mRFP-Htt fluorescence in the right panels. (A) The Sgs3-GFP glue protein can be readily seen in the left panel lining the exterior surface of the pupal case (arrows) following secretion from salivary glands. CS animals do not express Htt-mRFP, with only minor autofluorescence visible in the red channel (right panel). (B) Glue secretion (arrows) is not disrupted in pupae that express mRFP-HttQ15, which localizes diffusely in the salivary gland (arrowhead). (C) Glue secretion is decreased in pupae expressing mRFP-HttQ138, with the Sgs3-GFP protein largely retained in salivary glands (arrows). Aggregation of mRFP-HttQ138 in the pupal salivary glands is noted by the arrowhead in the right panel. (D and E) Sgs3-GFP is abundant in CS third instar larval salivary glands before pupation (D) and depleted from salivary glands following secretion during pupation (E). (F) Sgs3-GFP is retained in pupal salivary glands of mRFP-HttQ138–expressing animals. Bars, 20 μm.
imaging sessions in third instar larvae, we did not observe new aggregates forming. We attempted to visualize aggregate formation in younger animals, but numerous aggregates were already present at the first instar larval stage. It appeared that aggregates that formed early in development grew, but that new aggregates rarely formed in the presence of preexisting ones. To bypass this problem, we used the temperature-sensitive GAL80 repressor to restrict expression of mRFP-HttQ138 through the second instar stage. Expression of mRFP-HttQ138 with the CCAP-GAL4 driver was repressed by GAL80 at 18°C until the larvae reached third instar. We then turned on HttQ138 expression by shifting animals to the GAL80 restrictive temperate of 30°C. Two hours after induction of expression, a low level of diffuse mRFP-HttQ138 was observed in the salivary gland. By 4 hr, small Htt aggregates were observed forming in the salivary gland (Figure 7). After 8 hr of expression, diffuse mRFP-HttQ138 was seen in neuronal cell bodies in the ventral nerve cord, as well as in the proximal regions of axons. By 12 hr, aggregates were seen forming in axons and cell bodies, while salivary gland aggregates increased in size. During the window from 12 to 24 hr after induction, aggregates became larger and more numerous (Figure 7). Beyond 12 hr postinduction, HttQ138 aggregates began growing in size rather than number. The increase in aggregated mRFP-HttQ138 was accompanied by a decrease in the soluble fraction of the protein. We also examined the effectiveness of reducing aggregation by turning off expression of the mRFP-HttQ138 transgene after aggregates had formed, assaying whether cells had the ability to remove aggregates without new HttQ138 expression. After a 24-hr

Figure 4 The 588-aa fragment of mutant human Htt does not undergo cleavage in Drosophila. (A and B) Transient transfection of Drosophila S2 cells with eGFP-HttQ15-mRFP (A) or eGFP-HttQ138-mRFP (B) demonstrates no separation of eGFP (green) and mRFP (magenta) signals, suggesting that the mutant Htt protein does not undergo cleavage in S2 cells. (C–E) Visualization of signal localization in third instar larvae with expression of eGFP-HttQ138-mRFP driven by C155 shows no separation of eGFP (green) and mRFP (magenta) signals in CNS neurons (C), salivary gland cells (D), or epidermal cells (E). (F) Western analysis of brain extracts from control white animals and animals expressing Q138-mRFP or eGFP-Q138-mRFP, showing expression of the Htt protein (asterisks). Immunoblotting with anti-Htt antibodies reveals no differential breakdown product in the double-labeled strain compared to single-labeled lines or control animals that do not express Htt.
pulse of mRFP-HttQ138 expression followed by 72 hr of recovery, we observed a reduction in the size and number of aggregates present (Figure 7). We conclude that new aggregates can form in neurons within 12 hr, while preexisting aggregates rapidly accumulate new HttQ138 protein within minutes. In addition, in the absence of new Htt protein synthesis, neurons are capable of reducing preexisting Htt aggregate size over time.

**mRFP-HttQ138–induced lethality can be rescued by heterozygous disruption of single loci**

To further define the role of aggregation in HD pathology, we performed forward genetic screens for suppressors of HttQ138-induced lethality or suppressors of HttQ138 aggregation. For the first screen to identify lethality suppressors, we employed a dominant suppressor strategy with the *Drosophila* autosomal deficiency (Df) set to identify chromosomal regions containing loci that could dominantly rescue mRFP-HttQ138 driven by C155 caused pharate adult lethality. We screened for Dfs that rescued this lethality, comparing the number of adult females expressing mRFP-HttQ138 to that of males not expressing the HttQ138 transgene. Deficiencies that dominantly increased the ratio of mRFP-HttQ138 expressing females by 10-fold (female/male ratio = 0.1) were identified as hits. We identified 11 large Dfs, each removing ~100 genes, that suppressed HttQ138-induced lethality, indicating the presence of multiple potential targets that can modify HttQ138 toxicity in a dominant manner (Figure 8A, Table S1). Ten of 11 deletions gave a partial rescue of viability, increasing the number of escapers to ~20–30% of that expected for a full rescue. The viable animals displayed motor defects, as they were unable to climb the walls of the vials or mate, and most died within several days of eclosion, indicating partial rescue. One of the 11 large deletions, *Df(3L)vin7* showed a near complete rescue of viability at eclosion. C155; *Df(3L)vin7*/mRFP-HttQ138 animals showed less severe motor defects.
than other rescuing Dfs, but were unable to climb the vial walls and lived ~10 days posteclosion.

**Reduction of HttQ138 expression and aggregation increases viability in mRFP-HttQ138-expressing Drosophila**

In addition to the viability screen, we conducted a screen for suppressors of aggregation. The same mating scheme was used in both screens. However, the aggregation screen used the lower-expression adult-viable mRFP-HttQ138B insert, instead of the pharate lethal high-expression mRFP-HtrQ138 line. We reasoned the lower-expression line would represent a more sensitized system to identify potentially weak aggregation suppressors that might not be found in strong expression strains. Live wandering third instar larvae expressing mRFP-HttQ138 and heterozygous for each Df were screened under a fluorescent microscope for changes in aggregation. We focused on Htt aggregation within the salivary gland, as these large cells were easily visualized in live animals. We screened for changes in size, density, or brightness of perinuclear mRFP-Htt salivary gland aggregates. Four large deletions, (Df(2R)59AD, Df(2R)AA21, Df(2R)cn9, and Df(3L)vin7) caused a reduction in aggregate density in the salivary gland (Figure 8, B–F). Interestingly, each of these aggregation suppressors was independently identified in the viability screen. Thus, reducing aggregation in our screen was always associated with increased viability. We did not identify any hits that reduced aggregation without increasing viability, suggesting that aggregates may represent a toxic species in this model. While Df(2R)59AD reduced the density and size of salivary gland aggregates, Df(2R)cn9, Df(3L)vin7, and Df(2R)AA21 were most effective in reducing aggregation, with the mRFP-Htt Q138 pattern appearing as diffuse as the nonpathogenic mRFP-HttQ15.

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**Figure 6** FRAP microscopy shows that mRFP-HttQ138 aggregates continue to accumulate HttQ138 in live anesthetized third instar larval axons. (A) Averaged traces comparing FRAP rates of regions of axons with no aggregates, large aggregates that equal or exceed the diameter of the axon, and smaller aggregates less than the diameter of the axon. While regions without aggregates recover quickly due to fast axonal transport, large and small HttQ138 aggregates recover at a slower rate, but to a greater level. The graph shows percentage of initial fluorescence, not total fluorescence. Larger aggregates recover at a greater rate than smaller ones. (B) HttQ138 aggregation kinetics demonstrated by FRAP in different regions of the motor neuron. (C) Time-lapse images of a FRAP experiment showing recovery of an aggregate over 40 min in a live anesthetized third instar larval motor neuron axon.
Figure 7  Acute induction of mRFP-HttQ138 expression by CCAP-GAL4 shows the pattern of aggregate formation in salivary glands, ventral nerve cord, and axons over 24 hr. Expression of UAS-HttQ138 was repressed using tubulin-Gal80<sup>ts</sup> at 19<sup>th</sup> during early development. Animals were moved to 30<sup>th</sup> for the designated time, dissected, and imaged immediately to avoid fixation artifacts.
We attempted finer mapping for each larger Df to define the smallest relevant genetic interval. We were able to map the Df(2R)59AD interval to a region uncovered by Df(2R)59AB, which deletes ~20 genes. The larger Df(2R)cn9 was subdivided to a critical region uncovered by Df(2R)sple-J1, which removes ~39 genes.

To determine whether suppression of aggregation was associated with a change in mRFP-HttQ138 expression, we quantified Western blots of flies expressing mRFP-HttQ138 with C155 in the Df background (Figure 9, A and B). As a control for UAS-GAL4 transgene regulation, we also quantified expression of UAS-CD8-GFP in the Df backgrounds (Figure 9C). All four deletions that reduced aggregation resulted in reduced mRFP-HttQ138 and CD8-GFP protein expression. However, two single gene suppressors (lab14, Pbc02324; see below) had no effect on salivary gland aggregation or transgenic protein expression by Western analysis. To further characterize the suppressors, we measured HttQ138 mRNA levels using semiquantitative RT-PCR. Despite showing distinct effects on protein level, all Dfs decreased HttQ138 mRNA levels. These results are consistent with the observation that Htt aggregation is strongly influenced by expression levels of the protein and that toxicity in our model correlates with HttQ138 expression level (Figure 1C).

We next determined whether the reduction in salivary gland HttQ138 aggregates and suppression of lethality by these Dfs resulted in alteration in the subcellular distribution or density of Htt aggregates in peripheral nerves. Htt aggregates in axons have been suggested to cause axonal transport defects and contribute to HD pathogenesis (Li et al. 1999, 2001; Gunawardena et al. 2003; Lee et al. 2004). We found a significant (P < 0.05) decrease in the number of mRFP-HttQ138 aggregates >1 μm diameter in axons in the Df(3L)vin7 background, but no change in the other Dfs (Figure 10). Interestingly, Df(3L)vin7 had the strongest effect on suppressing lethality (Figure S1) and

Figure 8  Suppression of lethality with a Df haploinsufficiency screen. (A) Male/female viability ratio of mRFP-HttQ138–expressing Df lines. Homozygous mRFP-HttQ138 females were crossed to C155A; Df/Bal males for 81 Df lines on chromosome II and 60 Df lines on chromosome III. Few female escapers expressing mRFP-HttQ138 are ever seen in control crosses (male/female ratio = 0.01). All Dfs with a viability ratio >0.1 were identified as hits. (B–F) Confocal images of mRFP-HttQ138 aggregates in the salivary glands of third instar larvae from controls (B) and animals heterozygous for Df(3L)vin7 (C), Df(2R)sple-J1 (D), Df(2R)AA21 (E), and Df(2R)59AB (F). Bars, 100 μm.

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improving motor performance in HttQ138-expressing animals. We also analyzed Syt 1 distribution along axons, which we have previously observed to coaggregate with Htt in axonal aggregations (Lee et al. 2004), while remaining diffuse in axons from controls or HttQ15-expressing animals. Abnormal aggregation of Syt 1 was still observed in the rescued animals (Figure 10, A–G), indicating that although the Dfs reduced aggregation in salivary glands and suppressed lethality, they did not prevent axonal aggregation of HttQ138. The identification of dosage-sensitive suppressors of mutant Htt toxicity that reduce lethality without disrupting aggregation suggests that pathways downstream of aggregate formation can be targeted for neuroprotection in HD. The differences in HttQ138 mRNA levels vs. HttQ138 aggregates also suggest that subtle changes in transcript level can have a dramatic effect on the concentration reached within a cell that is required to trigger aggregation.

**Mapping of suppressors of HttQ138-induced lethality**

To begin identifying loci that underlie suppression of HttQ138-induced phenotypes, we attempted finer mapping of hits from the viability and aggregation screen by testing smaller and overlapping Dfs within these regions using stocks available from the Bloomington Stock Center (Table S2). In many cases, coverage across the original Df region was not sufficient to map suppression to individual loci. It is also possible that the ability of some Dfs to suppress toxicity may have resulted from additive effects of haplo-insufficiency for several genes within the deleted region. However, we were able to successfully refine 3 of the original 11 large deletions to two individual loci. The large Df suppressors Df(3L)st-f13 and Df(3L)brm11 overlapped a 43-gene region from 72C1 to 72D5. Df(3L)ED220 further refined the area to 72C1–72D4, a 20-gene interval with mutant stocks available for 12 predicted loci (Figure S1). We tested the 12 lines and

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**Figure 9** Quantification of transgenic Htt expression. (A) Western blot with C155-driven HttQ138 expression in control (white) and the indicated genotypes from pupal head extracts probed with anti-Htt antibodies and control anti-actin antibodies. (B) Quantification of C155-driven HttQ138 protein expression in control (white) and the indicated genotypes from pupal head extracts. The control containing C155; UAS-mRFP-HttQ138 was normalized to one for the genotypic comparisons. (C) Quantification of C155-driven GFP-CD8 expression in control (white) and the indicated genotypes from adult head extracts by Western blot analysis with anti-GFP antibodies. (D) HttQ138 mRNA levels were measured in pupal head extracts by quantitative RT-PCR and normalized to control (white) expression. Error bars indicate SEM. *P < 0.05 by Student’s t-test.
observed that stock 10887 rescued HttQ138-expressing animals at greater than the 10% expected ratio. Line 10887 (PBc02324) is a piggyBac transposable element insertion into the 5' region between two genes, CG5830 and mRps31. RT-PCR of line 10887 revealed that CG5830 is downregulated twofold, while mRps31 is not significantly affected by

Figure 10 HttQ138 and Synaptotagmin aggregation in peripheral axons. (A–G) Confocal images of third instar larval peripheral nerves expressing mRFP-HttQ138 (magenta, left panel) and immunostained with anti-synaptotagmin I antibodies (green, center panel) from controls (A) and animals heterozygous for PBc02324 (B), lab14 (C), Df(3L)win7 (D), Df(2R)sple-J1 (E), Df(2R)AA21 (F), and Df(2R)59AB (G). Bars, 10 μm. (H) Quantification of HttQ138 aggregate number for 100-μm axon segments for 25 segments (n = 5 larvae) of the indicated genotypes. Aggregates >0.5 μm were counted using the “find 2D nucleus” function of the Velocity version 5.4 software (Perkin-Elmer). Error bars indicate SEM; *P < 0.05 using Student’s t-test.
The remaining interval we were able to refine to a single mutation was \(\text{Df(3R)Tpl10}\), which deletes the 83C1–84B2 region and uncovers 173 genes. Screening other Dfs in this region revealed that \(\text{Df(3R)MAP117}\) and \(\text{Df(3R)MAP2}\) significantly rescued viability of HttQ138-expressing animals (Figure S1). These Dfs overlap the 84A1–84A5 interval, which contains 47 genes with 18 stocks available that disrupt loci within. We tested these strains and found that stock 2092 rescued mRFP-HttQ138-induced lethality, increasing the expected adult viability ratio to 24% vs. 0% in strains expressing HttQ138 alone. Stock 2092 is an X-ray-induced amorphic mutant (\(\text{lab}^{14}\)) of the \(\text{labial}\) gene. \(\text{Labial}\) is one of 8 homeobox genes in the Antennapedia cluster that play critical developmental roles in anterior–posterior body axis specification (Brody 1999). To determine whether other members of the Antennapedia HOX complex could also suppress HD pathology in our model, we tested mutations in additional members of the Antennapedia cluster for their ability to rescue HttQ138-induced lethality. Haplo-insufficiency for mutations in \(\text{Sex combs reduced (Scf)}^{\text{CP1}}, \text{22% viability ratio}; \text{Scf}^{\text{E}}, \text{24% viability ratio}; \text{proboscopedia (pr)}^{\text{l}}, \text{22% viability ratio}, \text{Deformed (Dfd)}^{\text{E}}, \text{15% viability ratio}\) and \(\text{Ultrabithorax (Ubx)}^{\text{G1}}, \text{12% viability ratio}\) also resulted in a significant rescue of HttQ138-induced lethality.

**Both aggregated and soluble HttQ138 are likely to represent toxic species**

By comparing the relative contributions of decreased aggregation and decreased soluble HttQ138 measured experimentally to increased life span, we observed a general trend whereby decreasing Htt expression causes a decrease in aggregation and an increase in viability (Figure 11). However, two suppressors uncovered in the screen displayed increased viability without a significant change in aggregation in peripheral motor axons \(\text{Df(2R)59AB}\) and \(\text{Df(2R)sple-j1}\). mRFP-HttQ138 expression in these backgrounds is decreased by Western analysis and RT-PCR, but there is no change in the number of axonal aggregates (Figure 8, C and E). Additionally, \(\text{Pbc020324}\) and \(\text{lab}^{14}\) show increased life span without a significant change in aggregation in any cell type. These suppressor backgrounds indicate some increase in viability can be achieved independent of decreasing aggregated HttQ138. In these lines, the increase in life span is correlated with a decrease in soluble HttQ138, arguing that this form of HttQ138 may also contribute to toxicity.

**Discussion**

Many neurodegenerative diseases associated with protein misfolding have been modeled in \textit{Drosophila}, including Parkinson’s disease (Feany and Bender 2000), Alzheimer’s disease (Wittmann et al. 2001), spinocerebellar ataxia type 1 (Fernandez-Funez et al. 2000) and type 3 (Warrick et al. 1999), and Huntington’s disease (Jackson et al. 1998; Steffan et al. 2001; Gunawardena et al. 2003; Lee et al. 2004). These models replicate many neuropathological features characteristic of the diseases, such as late onset, progressive neurodegeneration, and formation of inclusions containing the mutant protein. Here we describe the generation of a new \textit{Drosophila} HD model in which expression of a 588-aa N-terminal fragment of human Htt containing a 138 poly(Q) tract results in pharate adult lethality. By engineering a fluorescent tag on the mutant Htt fragment, we were able to visualize the location, trafficking, and aggregation of Htt in both neuronal and non-neuronal cells in live \textit{Drosophila}. As such, we were able to screen independently for suppressors of Htt aggregation by following HttQ138-mRFP localization in live animals. The screens resulted in the identification of seven large Dfs...
uncovering genomic regions that are capable of suppressing Htt-induced lethality without altering HttQ138 aggregate formation and four additional Dfs that suppressed both Htt-induced lethality and HttQ138 aggregation. Our findings indicate the presence of gene products downstream, or independent, of aggregation that can dominantly reduce HD toxicity. Our results also indicate that expression levels of mutant Htt are critical for disease pathology, as all Dfs we identified that reduced Htt expression levels by ~50% increased viability. As such, targeted approaches that reduce mutant Htt expression by relatively modest amounts may have profound effects on toxicity in HD patients.

The ability to follow Htt dynamics in live animals using our fluorescently tagged HttQ15 and HttQ138 transgenes revealed several important aspects of our *Drosophila* HD model. We find that both pathogenic and nonpathogenic versions of Htt are localized to the cytoplasm of all cell types examined. In humans, cleavage of Htt is thought to be important in generation of toxic Htt fragments (Qin and Gu 2004), with several studies indicating that small cleaved N-terminal fragments enter the nucleus to form intranuclear inclusions that contribute to pathogenesis (Difiglia et al. 1997; Sieradzan et al. 1999). Although evidence suggests that intranuclear aggregates can contribute to HD pathology (Davies et al. 1997; Becher et al. 1998), several studies indicate that pathogenic Htt aggregates in the cytoplasm and neurites play a causative role (Li et al. 1999; Sapp et al. 1999). As such, where toxic Htt fragments responsible for HD pathology reside in neurons remains to be conclusively identified. Using Htt fragments that were tagged at the N terminus with GFP and at the C terminus with mRFP, we demonstrate that cleavage of Htt is not apparent in *Drosophila*. In addition, expression of a smaller Htt exon 1 fragment still localizes to the cytoplasm. It is possible that addition of a GFP moiety alters Htt localization, although we observed the same pattern with an HA-tagged Htt transgenic protein as well (Lee et al. 2004). Thus, pathology in this HD model occurs secondary to cytoplasmic poly(Q) Htt localization.

To characterize the kinetics of Htt aggregation, we used live imaging of fluorescently tagged mRFP-HttQ138. Several studies indicate mutant Htt can disrupt FAT (Gunawardena et al. 2003; Szelenyi et al. 2003; Lee et al. 2004), but the mechanism by which defects occur is unclear. Since Htt is selectively toxic to neurons, an attractive model is that disrupted FAT may confer toxicity due to transport defects. Our FRAP data suggest that large immobile aggregates acquire new HttQ138 puncta. We determined the rate of FRAP recovery within aggregates was the same in both axons and the cell body, suggesting that aggregation kinetics are not solely dependent on the higher local concentrations of Htt induced by FAT. We took advantage of the Gal80* repressor to view early events in aggregate formation and visualize the aggregation progression. We observed that soluble HttQ138 forms aggregates that grow in size over a 12-hr window, decreasing the amount of soluble Htt present in the cell. Incorporation of HttQ138 into preexisting aggregates was much faster, suggesting that once aggregates are formed, they represent an active sink for accumulating new Htt protein on a minute timescale.

A key question in the HD field is whether Htt aggregates are toxic, neuroprotective, or simply by-products of the disease process. This issue is of critical importance for therapeutic considerations, as many current efforts are aimed at reducing Htt aggregation, assuming that this will decrease toxicity. Our previous observations that Htt aggregates accumulate in axons and impede axonal transport (Lee et al. 2004) suggested a model in which axonally localized aggregates can disrupt neuronal function. Whether aggregates in other cellular compartments cause toxicity is still an open question. The ability of drugs like Congo Red (Heiser et al. 2000; Sanchez et al. 2003), minocycline (Chen et al. 2000; Smith et al. 2003), and the transglutaminase inhibitor cystamine (Dedeoglu et al. 2002) to block Htt aggregation and reduce behavioral phenotypes in R6/2 Htt poly(Q) mice is suggestive of aggregate toxicity as well. Molecular (Warrick et al. 1999; Fernandez-Funez et al. 2000; Jana et al. 2000; Cummings et al. 2001; Vacher et al. 2005) and chemical (Yoshida et al. 2002) chaperones that reduce aggregate formation have been shown to reduce cytotoxicity in several poly(Q) disease models. Likewise, intracellular antibodies (Lecerf et al. 2001; Khoshnan et al. 2002; Colby et al. 2004; Wolfgang et al. 2005) that bind mutant Htt epitopes and suppress aggregate formation provide some neuroprotection in animal models. Our genetic screens support a model where aggregate formation and soluble Htt (monomers or oligomers) may both contribute to toxicity, as every Df that reduced aggregation or soluble Htt levels showed an increase in viability. The observation that we did not identify any haplo-insufficient loci capable of reducing aggregation without altering Htt expression levels suggests that aggregation inhibition alone may require more potent pharmacological effects than can be achieved with only a 50% reduction in activity of a single protein. In contrast, the finding that *Df(2R)spleJ1* and *Df(2R)59AB* increase viability without changes in aggregation suggests that HttQ138 toxicity can be reduced without altering Htt aggregation.

Although neuronal toxicity associated with Htt poly(Q) expansion has been the emphasis of most HD studies, the Htt protein is expressed in many nonneuronal tissues, including testes, liver, heart, lungs, and pancreatic islets (Ferrante et al. 1997; Cattaneo et al. 2005). HD patients have been shown to have a higher risk of diabetes (Lalic et al. 2008), and mouse HD models show pancreatic pathology (Martin et al. 2008), indicating that Htt poly(Q) expansion may also cause defects in nonneuronal cells. We found that HttQ138 expression in *Drosophila* salivary glands results in defective glue secretion (Figure 3), suggesting that nonneuronal defects from HttQ138 expression exist in our model as well. The ease with which the Sgs3-GFP glue secretion assay can be performed opens up the possibility of additional screens for second-site suppressors that alleviate salivary gland dysfunction in HttQ138-expressing animals. It
that ANTP genes are upregulated in HD and contribute to neuropathology. Several Hox genes, including labial, proboscipedia, and Sex combs reduced, were increased in Htt-poly(Q) brains, suggesting that haplo-insufficiency for these loci might function to reduce Htt expression to a less toxic level. Further studies will be needed to dissect this link between the homeotic genes and HD pathology.

Examining the pattern by which suppressor mutations interact with HttQ138 provides insight into the role of aggregates in toxicity. The weakest suppressors had no alteration in aggregation or HttQ138 expression, suggesting some minor rescue is possible without disrupting aggregation. A larger increase in viability occurs with decreased expression of HttQ138, but little to no change in aggregated Htt, suggesting that soluble HttQ138 is a toxic species in these cases. The final group of suppressors increased viability even more, while decreasing both soluble and aggregated Htt. In summary, we have generated a new HD model in Drosophila that allows for in vivo analysis of pathogenic Htt localization, aggregation, and dynamics. Using this model, we identified several genetic suppressors that can reduce HttQ138-mediated toxicity. The most robust suppressors reduced both soluble and aggregated Htt levels, suggesting that toxicity is likely to be associated with both forms of the mutant protein.

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Huntingtin Aggregation Kinetics and Their Pathological Role in a *Drosophila* Huntington’s Disease Model

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**Figure S1**  Fine mapping for two genetic intervals uncovered in the initial suppressor screen for HttQ138 pupal lethality. (A) Suppressors *Df(3L)st-f13* (Bloomington stock 2993) and *Df(3L)bbrm11* (Bloomington stock 3640) overlap the 72C1-72D5 interval. Mapping with additional aberrations contained within the Bloomington Stock Center demonstrated that stock 10887 (PBr02324) rescued HttQ138 expressing animals at 15% of expected viability. (B) Suppressor *Df(3R)Tpi10* (Bloomington stock 1990) was sub-mapped to a single genetic aberration within the region defined by stock 2092 (*lab<sup>L5</sup>*) that increased the expected adult viability ratio to 24%.
Figure S2  Quantitative RT-PCR was performed on the Pbc02324 p-element insertion mutant. The transcript levels of both genes surrounding the insertion, CG5830 and mRp531, were measured in homozygous mutant heads compared to white flies. Both genes appear slightly downregulated in the mutant, but only CG5830 was statistically significant. Error bars indicate SEM, with * denoting p<0.05 using Student’s t-test.
File S1

Time-lapse confocal images of CCAP-GAL4; mRFP-HttQ138 showing FRAP of four aggregates of various sizes.

Frame rate is 1 per 60 seconds for a total time of 60 minutes.

File S1 is available for download at as a movie file.
Table S1  Table showing the 166 Bloomington deficiency kit strains that were tested in the initial screen, along with the number of viable flies and changes in salivary gland aggregation. The viability data is represented graphically in figure 8A.

Table S2  Table showing stocks used for mapping of screen hits. Control data is listed on the first page. Each additional page heading represents mapping from the original large Df identified in the screen. The cells within each page list most of the smaller deficiencies tested in each region and their results for viability and/or aggregation where applicable.

Tables S1 and S2 are available for download as excel files.