BRAIN COMMUNICATIONS

Subcellular Localization And Formation Of Huntingtin Aggregates Correlates With Symptom Onset And Progression In A Huntington'S Disease Model

Christian Landles, ¹ Rebecca E. Milton, ¹ Nadira Ali, ¹ Rachel Flomen, ¹
Michael Flower, ¹ Franziska Schindler, ² Casandra Gomez-Paredes, ¹
Marie K. Bondulich, ¹ Georgina F. Osborne, ¹ Daniel Goodwin, ¹ Grace Salsbury, ¹ Caroline L. Benn, ^{1,3} Kirupa Sathasivam, ¹ Edward J. Smith, ¹ Sarah J. Tabrizi, ¹ Erich E. Wanker² and Gillian P. Bates¹

Huntington's disease is caused by the expansion of a CAG repeat within exon 1 of the HTT gene, which is unstable, leading to further expansion, the extent of which is brain region and peripheral tissue specific. The identification of DNA repair genes as genetic modifiers of Huntington's disease, that were known to abrogate somatic instability in Huntington's disease mouse models, demonstrated that somatic CAG expansion is central to disease pathogenesis, and that the CAG repeat threshold for pathogenesis in specific brain cells might not be known. We have previously shown that the HTT gene is incompletely spliced generating a small transcript that encodes the highly pathogenic exon 1 HTT protein. The longer the CAG repeat, the more of this toxic fragment is generated, providing a pathogenic consequence for somatic expansion. Here, we have used the R6/2 mouse model to investigate the molecular and behavioural consequences of expressing exon 1 HTT with 90 CAGs, a mutation that causes juvenile Huntington's disease, compared to R6/2 mice carrying \sim 200 CAGs, a repeat expansion of a size rarely found in Huntington's disease patient's blood, but which has been detected in post-mortem brains as a consequence of somatic CAG repeat expansion. We show that nuclear aggregation occurred earlier in R6/2(CAG)₉₀ mice and that this correlated with the onset of transcriptional dysregulation. Whereas in R6/2(CAG)₂₀₀ mice, cytoplasmic aggregates accumulated rapidly and closely tracked with the progression of behavioural phenotypes and with end-stage disease. We find that aggregate species formed in the R6/2(CAG)₉₀ brains have different properties to those in the R6/2(CAG)₂₀₀ mice. Within the nucleus, they retain a diffuse punctate appearance throughout the course of the disease, can be partially solubilized by detergents and have a greater seeding potential in young mice. In contrast, aggregates from R6/2(CAG)₂₀₀ brains polymerize into larger structures that appear as inclusion bodies. These data emphasize that a subcellular analysis, using multiple complementary approaches, must be undertaken in order to draw any conclusions about the relationship between HTT aggregation and the onset and progression of disease phenotypes.

- 1 Huntington's Disease Centre, Department of Neurodegenerative Disease and UK Dementia Research Institute at UCL, Queen Square Institute of Neurology, UCL, Queen Square, WC1N 3BG, UK
- 2 Neuroproteomics, Max Delbrueck Center for Molecular Medicine, 13125 Berlin, Germany and Berlin Institute of Health (BIH), 10178 Berlin, Germany
- 3 LoQus23 Therapeutics, Babraham Research Campus, Cambridge, CB22 3AT, UK

Correspondence to: Gillian P. Bates, PhD, Huntington's Disease Centre, Department of Neurodegenerative Disease and UK Dementia Research Institute at UCL, Queen Square Institute of

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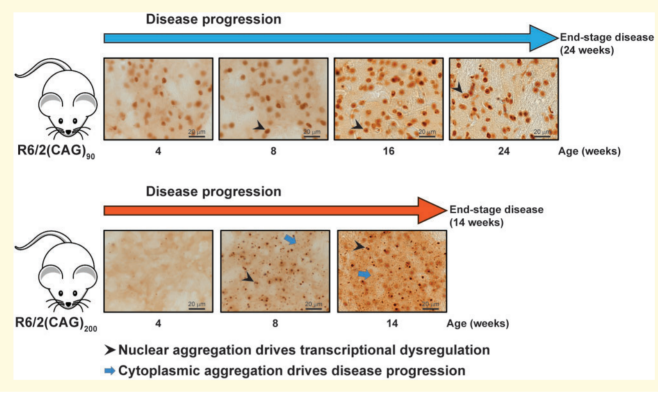
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Neurology, UCL, Queen Square, London WC1N 3BG, UK E-mail: gillian.bates@ucl.ac.uk

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Abbreviations: AGERA = agarose gel electrophoresis for resolving aggregates; PBS = phosphate buffer saline; polyQ = polyglutamine; qPCR = real-time quantitative PCR; SDS = sodium dodecyl sulphate

Graphical Abstract



Introduction

Huntington's disease is an inherited neurodegenerative disorder that manifests with psychiatric, motor and cognitive symptoms (Bates et al., 2015). The mutation is a CAG repeat expansion in exon 1 of the huntingtin gene (HTT) that results in an abnormally long polyglutamine (polyQ)tract in the huntingtin protein (HTT) (Huntington's Disease Collaborative Research Group, 1993). Individuals with less than 36 CAGs remain unaffected, those with 40 or more will develop the disease within a normal lifespan and those with 36-39 CAGs have an increasing chance of developing Huntington's disease (Rubinsztein et al., 1996). CAG repeat expansions of ~ 65 or more are likely to result in symptom onset in childhood or adolescence (Telenius et al., 1993). Huntington's disease causes widespread atrophy of the brain and results in prominent cell loss in the striatum, cortex and other brain regions (Vonsattel and DiFiglia, 1998). HTT aggregation can be detected in the brain by antibody staining and occurs as nuclear inclusions and neuropil aggregates (DiFiglia et al., 1997; Gutekunst et al., 1999).

Although CAG repeat length is the major determinant of age of onset and disease progression, genetic and environmental modifiers also contribute (Wexler et al., 2004). CAG repeats are unstable in Huntington's disease patient brains (Kennedy et al., 2003; Shelbourne et al., 2007; Gonitel et al., 2008) and somatic expansion in brain is correlated with an earlier disease onset (Swami et al., 2009). This somatic instability is recapitulated in mouse models of Huntington's disease (Mangiarini et al., 1997) and is prevented in the absence of the mismatch repair genes Msh2, Msh3, Mlh1 and Mlh3 (Manley et al., 1999; Pinto et al., 2013; Tome et al., 2013). Therefore, the identification of six DNA repair genes as modifiers of the age of onset and/or progression of Huntington's disease through genome-wide association studies: FAN1, MSH3, MLH1, PMS1, PMS2 and LIG1 (GeM-HD Consortium, 2015, 2019; Lee et al., 2017; Moss et al., 2017; Ciosi et al., 2019; Flower et al., 2019) underlined a causative role for somatic CAG instability in

the pathogenesis of Huntington's disease. It also suggested that the pathogenic repeat threshold in brain cells might be larger than that measured in blood and, if so, such a threshold is currently unknown.

The HTT gene contains 67 exons, and we have previously shown that exon 1 does not always splice to exon 2 resulting in the production of a small transcript (Httexon1) that terminates at cryptic polyadenylation sites within intron 1. This occurs in knock-in mouse models (Sathasivam et al., 2013; Franich et al., 2019), where the level of incomplete splicing correlates with the appearance of disease phenotypes, and in Huntington's disease post-mortem brains and cell lines (Neueder et al., 2017). The Httexon1 transcript contains a stop codon at the beginning of intron 1 and is translated to produce the highly pathogenic exon 1 HTT protein (Sathasiyam et al., 2013; Neueder et al., 2018), by the same mechanism as occurs in R6/2 mice (Mangiarini et al., 1996). The extent of this incomplete splicing increases with increasing CAG repeat length, and therefore, the consequence of somatic CAG expansion in the HTT allele is the increased production of the pathogenic exon 1 HTT protein.

The critical role that N-terminal HTT fragments play in the pathogenesis of Huntington's disease is supported by the fact that when CAG repeat size and strain background are comparable, R6/2 and knock-in mice have highly comparable phenotypes at late-stage disease (Kuhn et al., 2007; Woodman et al., 2007; Sathasivam et al., 2010; Labbadia et al., 2011; Mielcarek et al., 2014; Mielcarek et al., 2015). Here, we have established a novel R6/2 line with 90 CAGs $[R6/2(CAG)_{90}]$, a mutation that causes juvenile Huntington's disease, and conducted a molecular, immunohistochemical and behavioural comparison with R6/2 mice carrying ~200 CAGs [R6/2(CAG)₂₀₀]. We find that transcriptional dysregulation occurred earlier in R6/2(CAG)₉₀ mice and correlated with the appearance of aggregated HTT in neuronal nuclei. However, behavioural phenotypes progressed faster in the R6/2(CAG)₂₀₀ mice and this correlated with the accumulation of cytoplasmic aggregates. The length of the polyQ tract, which drives HTT aggregation in Huntington's disease patient brains is unknown; and we show here that the HTT aggregates formed from the expanded polyQ tracts in these two models have different immunohistochemical and molecular properties. Our work demonstrates that correlations between HTT aggregation and disease phenotypes requires a subcellular analysis both at molecular and immunohistochemical levels.

Materials and methods

Mouse breeding and maintenance

All procedures were performed in accordance with the Animals (Scientific Procedures) Act 1986 and were approved by the University Ethical Review Panel. Hemizygous R6/2 mice were bred by backcrossing R6/2 males to (C57BL/6JOlaHsd \times CBA/CaOlaHsd)F1 females (B6CBAF1/OlaHsd, Envigo, the Netherlands). R6/2 and wild type mice were group housed dependent on gender, but genotypes were mixed within cages. The temperature was regulated at 21°C ± 1°C and animals were kept on a 12 h light/dark cycle. All animals had unlimited access to water and breeding chow (Special Diet Services, Witham, UK) and to standard cage environmental enrichment (bedding and play tube). Mouse brains were rapidly dissected, frozen in liquid nitrogen and stored at -80° C.

Genotyping, CAG repeat sizing and somatic instability analysis

Mice were genotyped, and the CAG repeat size was measured by PCR of tail-tip DNA as previously described (Sathasivam et al., 2010). The mean CAG repeat size for all R6/2(CAG)₉₀ mice was 90.55 \pm 1.79, and for all R6/ $2(CAG)_{200}$ mice was 210.24 ± 7.2 (\pm SD). For somatic instability analysis, 50 ng DNA was amplified using HD3 FAM 5'-CCTTCGAGTCCCTCAAGTCCTT-3' and HD5 5'-CGGCTGAGGCAGCAGCGGCTGT-3' primers with Amplitaq Gold and GC enhancer (Thermo Fisher Scientific). Cycling conditions were 95°C for 10 min, 35 cycles of $(95^{\circ}C \text{ for } 30 \text{ s}, 65^{\circ}C \text{ for } 30 \text{ s}, 72^{\circ}C \text{ for } 90 \text{ s})$ and 72°C for 10 min. Samples were run on an ABI 3730xl Genetic Analyser with MapMarker ROX 1000 internal size standards and analysed using GeneMapper v5 software (Thermo Fisher Scientific). The extent of repeat instability was assessed by the somatic instability index (Lee et al., 2010) and expansion index (GeM-HD Consortium, 2019), relative to heart, using a custom R script and confirmed manually. Inclusion threshold for instability and expansion indices was 10% of modal peak height.

Behavioural and phenotype assessment

Body weight

Body weight was monitored weekly, prior to grip strength and rotarod assessment, and recorded to the nearest 0.1 g.

Grip strength

Forelimb grip strength was measured using a Grip Strength Meter (San Diego Instruments, CA, USA) at the following ages: $R6/2(CAG)_{90}$ at 6, 12, 16, 20 and 24 weeks of age, and $R6/2(CAG)_{200}$ at 4, 7, 9, 11 and 13 weeks of age. Mice were held by the base of the tail and allowed to grip the wire, they were gently pulled horizontally away from the apparatus until the wire was released and the maximum tension (g) was recorded. The mice were tested five times and the average performance of the best three trials was used (Hockly *et al.*, 2003).

The apparatus was thoroughly cleaned with 70% industrial methylated spirit between trials.

Rotarod performance

Mice were tested on a standard Ugo Basile 7650 rotarod (Linton Instrumentation, UK), with the modification of a smooth rubber coating over the rotating rod as previously described (Hockly et al., 2003). At the start of each trial, mice were acclimatized to the apparatus by being placed on the rotating rod at 4 rpm for 20 s. Following acclimatization, the rod progressively accelerated from 4 rpm to 40 rpm over a period of 5 min. Latency (s) for mice to fall from the rod was recorded. For the first test session [at 6 weeks of age for R6/2(CAG)₉₀, or 4 weeks of age for R6/2(CAG)200], rotarod performance was tested for three trials per day for four consecutive days. For all later sessions, performance was tested for three trials per day for three consecutive days. In both cases, the data from the last 2 days were used for the analysis. The apparatus was thoroughly cleaned with 70% industrial methylated spirit between testing each mouse.

RNA extraction and real-time quantitative **PCR**

Total RNA was isolated from individual brain regions using the RNeasy mini kit according to the manufacturer's instructions, which includes a DNase I treatment to remove genomic DNA (Qiagen). RNA was quantified using a Nanodrop-1000 spectrophotometer (Thermo Fisher Scientific) and stored at -80°C. cDNA was synthesized using random hexamers with an MMLV RT-ase kit (Invitrogen). Real-time quantitative PCR (qPCR) was then performed using the CFX96 C1000 real-time thermal cycler (Bio-Rad). Housekeeping reference genes were for cortex: Atp5b, Canx, Rpl13a and Ubc, striatum: Atp5b, Eif4a2 and Ubc, cerebellum: Atp5b, Rpl13a and Ubc (Primer Design). Primer sequences and probes used are summarized in Supplementary Table 1. The levels of gene expression were determined by normalizing to housekeeping genes, and the relative levels of the transcripts were determined by using the $2^{-\Delta\Delta Ct}$ method (Livak and Schmittgen, 2001; Benn et al., 2008).

Antibodies

The antibodies used for western blotting, Seprion-ligand ELISA, agarose gel electrophoresis for resolving aggregates (AGERA), filter-trap and immunohistochemistry, together with their respective dilutions, are summarized in Supplementary Table 2. Because of the difference in the length of the polyQ tracts between the two mouse lines, antibodies with a polyQ epitope were avoided, except for the detection of aggregates after formic acid antigen retrieval.

Cytoplasmic and nuclear fractionation

Cytoplasmic and nuclear fractions were prepared as previously described (Davies *et al.*, 1999), their protein concentration was determined using the PierceTM BCA protein assay kit (Thermo Fisher Scientific), and their purity was determined by immunoblotting using anti-histone H3 and anti- α -tubulin antibodies.

Western blotting

Half brains were homogenized (Polytron PT1200E) in ice-cold IGEPAL RIPA buffer [150 mM NaCl, 50 mM Tris-HCl pH 8.0, 1% IGEPAL, 0.5% sodium deoxycholate, 0.1% sodium dodecyl sulphate (SDS)] with 1 mM phenylmethylsulphonyl fluoride, 1 mM dithiothreitol and cOmplete protease inhibitor cocktail tablets (Roche). Protein concentration was determined using the PierceTM BCA protein assay kit (Thermo Fisher Scientific). For immunoblotting, 20-40 µg of total protein in Laemmli loading buffer was denatured at 85°C for 5 min and separated by 10-, 12-, or 15% Criterion TGX Stain-free SDS-PAGE (Bio-Rad) and blotted onto 0.45 µm nitrocellulose membrane (Bio-Rad) by submerged transfer apparatus (Bio-Rad) in [25 mM Tris base, 192 mM glycine, 20% v/v methanol]. For immunoblotting of cytoplasmic and nuclear fractions, either 25 µg of cytoplasm or 12.5 µg of nuclei in Laemmli loading buffer was used. Membranes were blocked for 1 h at room temperature in 3% non-fat dried milk in PBST [phosphate buffer saline (PBS), 0.1% Tween-20], and then incubated overnight with gentle agitation at 4°C with the primary antibody (in PBST with 1% non-fat dried milk). For chemiluminescent detection, blots were washed three times in PBST for 5 min, probed with horse radish peroxidase-linked secondary antibodies (in PBST with 0.5% non-fat dried milk) for 1 h at room temperature and washed three times in PBST. Protein was detected by chemiluminescense (Clarity, Bio-Rad) according to the manufacturer's instructions. The signals were captured on a ChemiDoc Touch Imaging System (Bio-Rad).

Seprion[™]-Pad ELISA

HTT aggregates were captured in Seprion-ligand coated plates (Microsens, PADPCB1 SEP1–01) as previously described (Sathasivam *et al.*, 2010). For individual brain regions, a 2.5% (w/v) total protein lysate was prepared in ice-cold IGEPAL RIPA buffer [150 mM NaCl, 50 mM Tris–HCl pH 8.0, 1% IGEPAL, 0.5% sodium deoxycholate, 0.1% SDS] with 1 mM phenylmethylsulphonyl fluoride, 1 mM dithiothreitol and cOmplete protease inhibitor cocktail tablets (Roche), by homogenizing three times for 30 s in Lysing matrix D tubes (MP Biomedicals) in a Fast-Prep-24TM instrument (MP Biomedicals), and 15 µl was loaded per well. For cytoplasmic and nuclear

fractions, samples were prepared as described, and either $25 \,\mu g$ cytoplasm or $5 \,\mu g$ nuclei diluted in $15 \,\mu l$ ice-cold IGEPAL RIPA buffer was loaded per well.

FRET-based aggregate seeding assay

For cortical brain tissue, a 10% (w/v) total protein lysate was prepared by homogenizing in ice-cold FRET-based aggregate seeding assay buffer [10 mM Tris–HCl pH 7.4, 0.8 M NaCl, 1 mM EDTA, 10% sucrose, 0.25 U/ml benzonase], with cOmplete protease inhibitor tablets (Roche), in Precellys[®] CK14 lysing kit tubes (Precellys[®] 24 Homogenizer). The homogenate was incubated for 1 h at 4 °C on a rotating wheel and protein concentration was determined using the PierceTM BCA protein assay kit (Thermo Fisher Scientific). The FRET-based aggregate seeding assay was performed and evaluated according to the method previously described (Ast *et al.*, 2018) using 2.5 µg of crude lysate per replicate.

Agarose gel electrophoresis for resolving aggregates

For AGERA gels (Weiss et al., 2007), 1.5 g agarose (Bio-Rad) was dissolved in 100 ml 375 mM Tris-HCl, pH 8.8, then once fully dissolved, SDS was added to a final concentration of 0.1%. Gels were gently poured into 10 cm Flowgen BioscienceTM gel trays (8 mm thick), and allowed to set (20-well comb, 2 mm thick). For immunoblotting, either 100 µg cytoplasm or 25 µg nuclei was diluted in non-reducing Laemmli buffer [150 mM Tris-HCl pH 6.8, 33% glycerol, 1.2% SDS and bromophenol bluel, denatured at 95 °C for 5 min, and loaded alongside a Hi-MarkTM HMW pre-stained standard (Invitrogen). Samples were run in ultra-pure Tris/Glycine/SDS [25 mM Tris, 192 mM glycine, 0.1% SDS] running buffer (National Diagnostics) at 100 V until the dye front reached the bottom of the gel, and blotted onto 0.45 µm nitrocellulose membrane by submerged transfer apparatus as described for western blotting above.

Filter-trap retardation assay of SDS insoluble aggregates

Half brains were homogenized in ice-cold IGEPAL RIPA buffer and protein concentration determined exactly as described for western blotting. For filter-trap, two WhatmanTM 3 mm and one 0.2 μ m cellulose acetate membrane filter (Whatman) were equilibrated in (PBS 0.1% SDS) and assembled onto a 96-well MiniFold-1TM dotblot system (Whatman). Next, wells were filled and washed twice with 250 μ l [PBS, 0.1% SDS] and aspirated under vacuum. Then, after filling each well with 200 μ l [PBS, 0.1% SDS], 10 μ g of protein lysate was loaded per well dissolved in 50 μ l [PBS, 0.1% SDS]. Finally, under vacuum, samples were aspirated, and subsequently washed three times in $250 \,\mu$ l [PBS, 0.1% SDS]. Once disassembled, cellulose acetate membranes were blocked for 1 h at room temperature in 3% non-fat dried milk in PBST [PBS, 0.2% Tween-20], and processed exactly as described for western blotting.

Immunohistochemistry

Mice were transcardially perfusion fixed with 4% paraformaldehyde (Pioneer Research Chemical Ltd). Brains were removed and further post-fixed for 6 h at 4 °C before cryoprotection with gradient steps of 20% and 30% sucrose in 0.01 M PBS (Sigma). Once the brains reached equilibrium (had sunk to the bottom of the sucrose solution) they were washed in PBS, embedded in OCT (CellPath Ltd) and stored at -80° C. Coronal brain sections were cut at 30 µm on a cryostat and stored free floating at -20° C in tissue protective solution [30% ethylene glycol, 25% glycerol and 0.05% sodium azide in PBS] until staining.

Sections were washed in PBS before endogenous peroxide activity was quenched using 1% hydrogen peroxide (H_2O_2) in PBS for 30 min followed by three washes in PBS. Prior to immunostaining, non-specific binding was blocked for 1 h using 10% normal goat serum (Sigma) in PBST [0.5% tritonX-100 (Sigma) in PBS]. MW8 primary antibody was applied in fresh blocking solution overnight at 4°C. Sections were washed three times in PBS before application of the biotinylated secondary antibody (1:500, Vector, Peterborough, UK) for 2 h. Sections were washed three times in PBS before signal amplification using Elite ABC reagent. Biotinylated antibody staining was visualized using Sigma FastDAB (0.5 mg/ml) in PBS and activated with H₂O₂. Sections were treated with DAB solution for 20s and rapidly washed with ice-cold PBS to stop the reaction. The sections were washed a further three times with ice-cold PBS to remove the DAB solution completely before mounting on glass slides. To counterstain nuclei, slides were exposed to thionin (Sigma) in [40 mM sodium acetate, 90 mM acetic acid pH 4.3] for 2 min. Slides were then processed through 70%, 90% and 100% alcohols and xylene before coverslipping with DPX mountant (Sigma).

Prior to 4H7H7 primary antibody application, antigen retrieval was used to expose aggregation buried epitopes. Sections were treated three times with 88% formic acid (Alfa Aesar) for 10 min followed by three washes in PBS after each treatment. Then, sections were treated with 0.5 M ethanolamine (Sigma) in PBS (pH 9.5, blocking excess reacting aldehyde sites) for 1 h with the addition of ascorbic acid (EDQM) to a concentration of 5 mM for the final 10 min (reducing Schiff bases). Sections were rinsed in PBS three times and blocked three times with PBST for 10, 30 and 10 min. Biotinylated 4H7H7 (1:3000) was then applied in fresh PBST overnight at 4° C. Sections were washed three times in PBS before signal amplification using Elite ABC reagent (Vector). Sections were washed three times in TI buffer [0.05 M Tris pH 7.4, 0.05 M Imidazole (Sigma) in PBS]. Signals were amplified using the Tyramide Signal Amplification kit (Perkin Elmer) according to the manufacturer's recommendations and then washed three times in PBST prior to a second application of Elite ABC reagent.

Histological images were obtained using a Ziess AxioSkop2 plus microscope fitted with a Ziess AxioCam HRc colour camera. Images were recorded using Ziess AxioVision 4.7.

Statistical analysis

Data were screened for outliers using ROUT or Grubb's Test (GraphPad software), and any outliers were removed before between-group comparisons. Statistical analysis was performed with SPSS (v22 and v26) and R using two-tailed Student's *t*-test, one-way ANOVA, two-way ANOVA or GLM ANOVA, with Bonferroni *post hoc* tests as indicated. Graphs were prepared using GraphPad Prism (v6 and v7) and R. *P*-values less than 0.05 were considered statistically significant.

Data Availability

The authors confirm that all the data supporting the findings of this study are available within the article and its Supplementary material. Raw data will be shared by the corresponding author on request. The custom repeat instability R script can be accessed and run online on request.

Results

The CAG repeat has been maintained at ~ 200 CAGs in our R6/2 colony for more than 15 years. To generate a line with a shorter CAG repeat expansion, R6/2 mice were recovered from cryopreserved sperm (Jackson laboratory; stock initially deposited by James Olson). The three resultant pups included one wild type female, one R6/2 male and one R6/2 female, both of which had CAG repeats of 90 CAGs. The line was established by backcrossing R6/2(CAG)₉₀ males to (CBA/Ca \times C57BL/ 6J)F1 females. We had previously shown that CAG repeats in R6 lines of mice expand when bred through the male line, and contract when bred from females (Mangiarini et al., 1997). Therefore, we transmitted the $(CAG)_{90}$ allele through the female line for six generations with the intention of reducing the CAG repeat length further, but the CAG repeat in this R6/2 allele proved to be very stable, whether bred through the male or female germ lines.

Behavioural phenotypes progress more slowly in R6/2 mice with (CAG)₉₀ than with (CAG)₂₀₀

To determine the course of disease in the $R6/2(CAG)_{90}$ mice, progressive changes in body weight, grip strength and rotarod performance were recorded. The $R6/2(CAG)_{200}$ data presented here were not collected at the same time and are included to allow the reader to draw comparisons in disease progression between the two lines. $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice were sacrificed at 24 and 15 weeks of age respectively at which point, they had reached comparable end-stage disease.

Body weight was measured weekly from 6 to 24 weeks of age for R6/2(CAG)₉₀ mice and 4 to 15 weeks of age for R6/2(CAG)₂₀₀ mice. In both cases, R6/2 mice stopped gaining weight and then lost weight relative to wild type littermates and the decrease in weight was more pronounced in males than in females: R6/2(CAG)₉₀ males (Fig. 1A) [F(age × genotype)_{16,306} = 10.125, P < 0.001], R6/2(CAG)₉₀ females (Fig. 1B) [F(age × genotype)_{16,306} = 3.247, P < 0.001], R6/2(CAG)₂₀₀ males (Fig. 1C) [F(age × genotype)_{11,167} = 6.590, P < 0.001] and R6/ 2(CAG)₂₀₀ females (Fig. 1D) [F(age × genotype)_{11,141} = 2.681, P = 0.004].

Forelimb grip strength was measured on weeks 6, 12, 16, 20 and 24 for R6/2(CAG)₉₀ and on weeks 4, 7, 9, 11 and 13 for R6/2(CAG)₂₀₀ mice. In males, grip strength decreased for R6/2 as compared to wild type mice: $R6/2(CAG)_{90}$ males (Fig. 1E) [F(age × genotype)_{4,85} = 34 810, P < 0.001] and R6/2(CAG)₂₀₀ males (Fig. 1G) [F(age \times genotype)_{4.70} = 9.305, P < 0.001]. In females, the grip strength of wild type and R6/2 mice only changed differently with age for the $R6/2(CAG)_{90}$ females (Fig. 1F) [F(age \times genotype)_{4,88} = 3.781, P = 0.007] but not for R6/2(CAG)₂₀₀ females (Fig. 1H) $[F(age \times genotype)_{4.59} = 1.567, P = 0.193].$ However, grip strength was different between wild type mice and $R6/2(CAG)_{200}$ females (Fig. 1H) [F(genotype)_{1.59} = 29.841, P < 0.001 and the R6/2(CAG)₂₀₀ female grip strength changed with age (Fig. 1H) $[F(age)_{4,59} =$ 37.802, *P* < 0.001].

To assess changes in balance and co-ordination, mice were tested for their ability to walk on an accelerating rotarod during weeks 6, 12, 16, 20 and 24 for R6/ 2(CAG)₉₀ and during weeks 4, 8, 10, 12 and 14 for R6/2(CAG)₂₀₀ mice. There was no difference in the latency to fall between R6/2(CAG)₉₀ and wild type littermates at 6 weeks of age (Fig. 11) [F(genotype)_{1,186} = 1.887, P = 0.178] or R6/2(CAG)₂₀₀ and wild type littermates at 4 weeks (Fig. 1J) [F(genotype)_{1,163} = 1.468 P = 0.234]. In both cases, performance deteriorated with disease progression: R6/2(CAG)₉₀ (Fig. 1I) [F(age × genotype)_{4,186} = 3.447, P = 0.010] and R6/ 2(CAG)₂₀₀ (Fig. 1J) [F(age × genotype)_{4,163} = 4.225, P = 0.003].

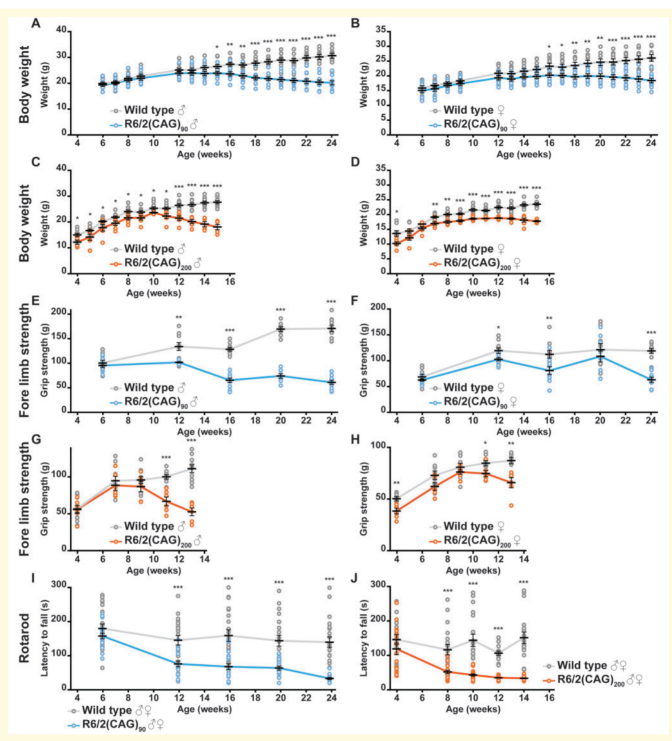


Figure 1 Behavioural and physiological phenotypes progress more slowly in R6/2(CAG)₉₀ **than R6/2(CAG)**₂₀₀ **mice. (A–D)** Failure to gain body weight occurred later in (A) R6/2(CAG)₉₀ males and (**B**) R6/2(CAG)₉₀ females than in (**C**) R6/2(CAG)₂₀₀ males and (**D**) R6/ 2(CAG)₂₀₀ females respectively. (**E–H**) Reduction in fore-limb grip strength progressed more slowly in (**E**) R6/2(CAG)₉₀ males and (**F**) R6/ 2(CAG)₉₀ females than in (**G**) R6/2(CAG)₂₀₀ males and (**H**) R6/2(CAG)₂₀₀ females. A statistically significant reduction in grip strength occurred at 12 weeks of age for R6/2(CAG)₉₀ and 11 weeks of age for R6/2(CAG)₂₀₀ for both genders of each genotype. Discrepancies in the wild type grip strength between the R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ datasets are because the data were collected on different machines by different operators at different times. (**I**, **J**) Rotarod performance, as compared to that of wild type littermates, deteriorated more slowly in (**I**) R6/ 2(CAG)₉₀ mice than in their (**J**) R6/2(CAG)₂₀₀ counterparts. The R6/2(CAG)₂₀₀ data were extracted from the analysis published in Bobrowska et *al.* (2012). Discrepancies in the latency to fall between the wild type data sets are because the data were collected using different machines by different operators at different times. Wild type (CAG)₉₀ (*n* = 20), R6/2(CAG)₉₀ (*n* = 17–20), wild type (CAG)₂₀₀ (*n* = 14–18) and R6/ 2(CAG)₂₀₀ (*n* = 16–19). Statistical analysis was GLM ANOVA with Bonferroni *post hoc* correction *P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001.

Transcriptional dysregulation presents earlier in R6/2(CAG)₉₀ than in R6/2(CAG)₂₀₀ mice

These physiological and behavioural measures indicated that disease onset occurred later and progressed more slowly in the $R6/2(CAG)_{90}$ as compared to the $R6/2(CAG)_{200}$ mice. Transcriptional dysregulation is a well-defined molecular phenotype that has been extensively studied in post-mortem brains (Hodges *et al.*, 2006) and mouse models of Huntington's disease (Langfelder *et al.*, 2016). To determine when transcriptional dysregulation could first be detected, we used qPCR to measure the levels of 'genes of interest', known to be highly dysregulated in Huntington's disease mouse models (Benn *et al.*, 2008), in the striatum, cortex and cerebellum of $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice at 4 and 8 weeks of age (Fig. 2).

There was a dramatic reduction in the levels of the striatal genes: Cnr1, Darpp32, Drd2, Pde10a and Penk1 in $R6/2(CAG)_{90}$ mice at 4 weeks of age, at which point they remained at wild type levels in R6/2(CAG)200 mice (Fig. 2A). By 8 weeks, all five genes were decreased to greater than 50% of wild type levels and dysregulated to a comparable extent in both the R6/2(CAG)₉₀ and R6/ 2(CAG)₂₀₀ lines (Fig. 2B). A similar pattern was evident for the cortical genes, with the expression levels of BdnfIV, Pgam2, Grm2, Hrh3 and Hrt1a decreasing at an earlier age in R6/2(CAG)₉₀ than R6/2(CAG)₂₀₀ mice (Fig. 2C and D). For the cerebellum, changes in Igbp5, Kcnk2 and Pcp4 occurred earlier in R6/2(CAG)90 mice (Fig. 2E and F) but this could not be determined for Nr4a2 and Uchl1 as measurement of expression levels at 4 and 8 weeks of age did not capture the age at which dysregulation first occurred for these two genes (Fig. 2E and F). Therefore, dysregulation of striatal, cortical and cerebellar transcripts occurred earlier in R6/2(CAG)₉₀ than R6/2(CAG)₂₀₀ mice, in contrast to the physiological and behavioural phenotypes which had a later age of onset.

Huntingtin aggregation can be detected earlier in the brains of R6/ 2(CAG)₉₀ than R6/2(CAG)₂₀₀ mice

We next used the Seprion ELISA assay to compare HTT aggregation levels in $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ lysates from striatum, cortex, hippocampus and cerebellum at 2, 4 and 8 weeks of age (Fig. 3A–D). In all brain regions from the $R6/2(CAG)_{90}$ mice, aggregation could be detected as early as 2 weeks of age, and the aggregate load remained higher at 4 weeks than for $R6/2(CAG)_{200}$ mice. However, by 8 weeks, the level of HTT aggregation was comparable between the two R6/2 lines (Fig. 3A–D). We next compared the aggregation levels in brain

between the R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice at end-stage disease, at 24 and 14 weeks of age, respectively. These were relatively comparable for the cortex and cerebellum, but end-stage aggregate levels were higher in the striatum and hippocampus from 14-week-old R6/ 2(CAG)₂₀₀ mice than for the R6/2(CAG)₉₀ line (Fig. 3E– H). Therefore, HTT aggregates formed at an earlier age in the R6/2(CAG)₉₀ lines but accumulated faster in the brains of the R6/2(CAG)₂₀₀ mice.

Next, cortical lysates from 4, 8, 16 and 24 week old R6/2(CAG)90 and 4-, 8-, and 14-week-old R6/ 2(CAG)₂₀₀ mice were separated by SDS-PAGE to assess the soluble and aggregated state of the exon 1 HTT protein (Fig. 3I and J). The migration of exon 1 HTT is retarded by expanded polyQ tracts with the consequence that exon 1 HTT with 90Q migrated at \sim 45 kDa and with 200Q at \sim 90 kDa. In both cases, the level of soluble exon 1 HTT decreased with disease progression as it was recruited into aggregates. At 14 weeks of age, soluble exon 1 HTT could no longer be detected in the cortex of R6/2(CAG)₂₀₀ mice, whereas in R6/2(CAG)₉₀ at 24 weeks, trace amounts were still present (Fig. 3]). In keeping with the Seprion-ligand ELISA data, aggregated HTT could still be detected more readily in the stacking gel at 4 weeks of age in R6/2(CAG)₉₀ lysates than in R6/2(CAG)₂₀₀ lysates (Fig. 3I and J).

CAG repeats undergo somatic instability in R6/2 mice to varying extents that are brain region and tissue specific (Mangiarini et al., 1997). Examination of the western blots (Fig. 3I and I), together with those in Supplementary Fig. 1, suggested that the size of the exon 1 HTT protein might be slightly larger in older $R6/2(CAG)_{90}$ brains as compared to those from younger mice. To investigate whether this might be due to somatic instability, we extracted DNA from available tissues: brainstem, cerebellum, liver, tibialis anterior, gastrocnemius, quadriceps and heart from R6/2(CAG)90 mice at 24 weeks of age and R6/2 (CAG)₂₀₀ mice at 14 weeks of age and prepared GeneMapper capillary electrophoresis traces of the CAG repeat profile (Fig. 4A and B). Consistent with our previous data, we found greater instability in brainstem than cerebellum (Fig. 4 and Supplementary Fig. 2). Similarly, the absence of CAG instability in the three muscles and heart, in both the R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice, was consistent with previous data (Mangiarini et al., 1997). However, unlike R6/2(CAG)₂₀₀ mice in which, as expected, the CAG repeat was very unstable in liver, we only detected a modest level of instability in this tissue in R6/2(CAG)₉₀ mice (Fig. 4 and Supplementary Fig. 2). These data predict that the CAG expansions in specific brain regions in older mice would be larger than the 90 CAGs originally measured in tail DNA prior to weaning.

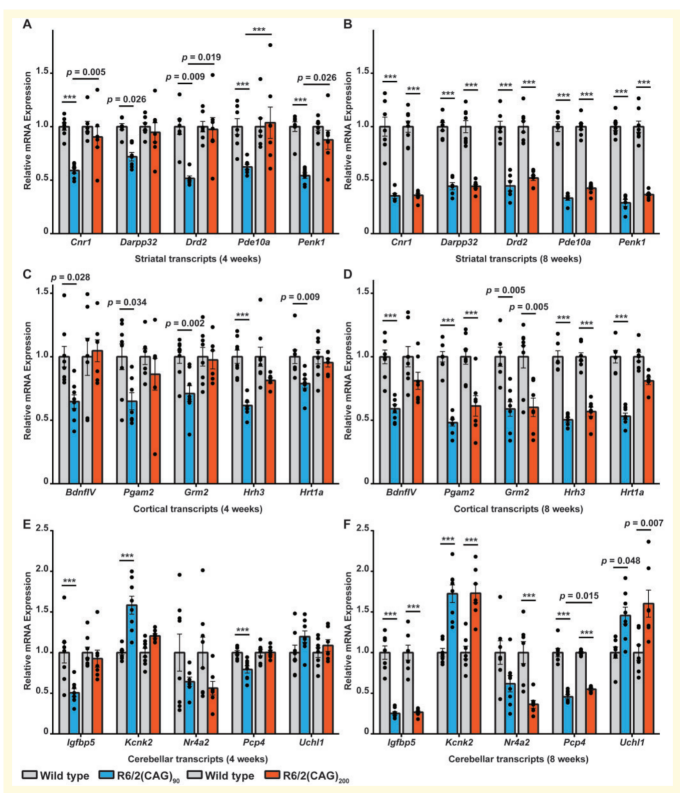


Figure 2 Striatal, cortical and cerebellar genes of interest were dysregulated earlier in $R6/2(CAG)_{90}$ than $R6/2(CAG)_{200}$ mice. (A, B) Comparison of the levels of striatal transcripts, as measured by qPCR, between $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice at (A) 4 weeks and (B) 8 weeks of age. $R6/2(CAG)_{90}$ transcript levels were decreased compared to those of wild type littermates at 4 weeks of age, whereas the $R6/2(CAG)_{200}$ and wild type levels were comparable. At 8 weeks of age, the transcripts were decreased to similar levels in both R6/2 lines. (C, D) Comparison of the levels of cortical transcripts, as measured by qPCR, between $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice at (C) 4 weeks and (D) 8 weeks of age. $R6/2(CAG)_{90}$ transcript levels were decreased compared to those of wild type littermates at 4 weeks of age, whereas the $R6/2(CAG)_{200}$ and wild type levels were comparable. At 8 weeks of age, the Pgam2, Grm2 and Hrh3 transcripts were also decreased in the $R6/2(CAG)_{200}$ mice, whereas BdnflV and Hrt1a, were still expressed at levels similar to wild type. (E, F) Comparison of the levels of cerebellar

The level of expression of the R6/2 transgene is higher in the brains of R6/2(CAG)₉₀ than R6/2(CAG)₂₀₀ mice

To further investigate the mechanism by which aggregation occurs earlier in the $R6/2(CAG)_{90}$ compared to the $R6/2(CAG)_{200}$ line, we measured the level to which the R6/2 transgene was expressed by qPCR. At 2 weeks of age, the R6/2 transgene was expressed at between 2.5and 3-fold greater in the striatum, cortex and cerebellum from $R6/2(CAG)_{90}$ as compared to $R6/2(CAG)_{200}$ mice (Supplementary Fig. 3A). Expression in the cortex decreased from approximately 2.5-fold higher to between 1.5- and 2-fold higher at between 4 and 14 weeks of age (Supplementary Fig. 3B). The levels of endogenous *Htt* were equivalent to that in wild type mice for both the $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice over this period (Supplementary Fig. 3C).

Formation of huntingtin aggregates in neuronal nuclei occurs earlier in R6/2(CAG)₉₀ compared to R6/ 2(CAG)₂₀₀ mice

To investigate the temporal and spatial appearance of HTT aggregates in the brains of R6/2(CAG)₉₀ and R6/ 2(CAG)200 mice, sections were immunoprobed with the MW8 antibody, which detects the C-terminus of the exon 1 HTT protein and readily detects HTT aggregates (Fig. 5). At 4 weeks of age, the level of immunostaining was greater in the R6/2(CAG)₉₀ sections for both cortex (Fig. 5A) and striatum (Fig. 5B). In contrast, there was very little signal in the R6/2(CAG)₂₀₀ cortex (Fig. 5A), and almost none detected in the R6/2(CAG)200 striatum (Fig. 5B). In the R6/2(CAG)₉₀ hippocampus, immunostaining could be detected throughout the pyramidal layer (CA1-CA3) and dentate gyrus, whereas in the R6/ 2(CAG)₂₀₀ animals, immunostaining was only prominent in the CA1 region (Fig. 5C and Supplementary Fig. 7A). In all cases, the immunostaining was predominantly nuclear, appearing as a diffuse signal in R6/2(CAG)₉₀ brains, whereas a more punctate inclusion pattern was present in R6/2(CAG)₂₀₀ sections (Fig. 5A-C). At 8 weeks of age, the nuclear staining pattern in R6/2(CAG)₉₀ sections remained diffuse, whereas in R6/2(CAG)200 sections the inclusions had increased in both number and size

(Fig. 5A–C). To illustrate the location of nuclei, sections from a 8-week-old $R6/2(CAG)_{200}$ mice were immunoprobed with MW8 and counterstained with thionin (Supplementary Fig. 4).

To investigate the nature of the diffuse nuclear immunostain, we then immunoprobed $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ sections from mice at 4 and 8 weeks of age with the 4H7H7 antibody that detects polyQ peptides. No signal was detected since the polyQ epitopes were buried in HTT aggregates (Supplementary Fig. 5). However, upon treatment with formic acid, which exposed the polyQ epitopes (rendering them available for antibody binding), the diffuse nuclear stain was now observed. Therefore, this represented an aggregated form of HTT, and was particularly evident in the brain sections from $R6/2(CAG)_{90}$ mice (Supplementary Fig. 5).

Cytoplasmic aggregates accumulate more rapidly in the brains of R6/ 2(CAG)₂₀₀ as compared to R6/ 2(CAG)₉₀ mice

Cytoplasmic aggregates could be seen in cortical and striatal sections from R6/2(CAG)₂₀₀ mice at 8 weeks of age (Fig. 5A and B), but appeared to be comparatively absent from the equivalent sections from $R6/2(CAG)_{90}$ mice. To investigate the presence of cytoplasmic aggregates in more detail, we focused on the hilus, that lies within the cleft of the dentate gyrus within the hippocampus, an area in which cytoplasmic aggregates are particularly abundant in R6/2(CAG)200 mice. Aggregated HTT accumulated in the neuronal nuclei of the pyramidal cells from the CA1 to the CA3 as well as the granule cells of the dentate gyrus in R6/2(CAG)90 brains at 4 weeks of age, at which time, aggregates were only present in the CA1 region of R6/2(CAG)₂₀₀ brains (Fig. 6A). At this age, cytoplasmic aggregates were not apparent in the hilus in either of the R6/2 lines upon immunoprobing with MW8. At 8 weeks of age, cytoplasmic aggregates were abundant in the hilus of R6/2(CAG)200 brains but were relatively sparse in the comparable regions from R6/ 2(CAG)₉₀ brains (Fig. 6A, Supplementary Fig. 5D). We next compared HTT aggregation in older mice: 16 and 24 weeks for R6/2(CAG)₉₀ and 14 weeks for R6/ 2(CAG)₂₀₀ mice, where 24 weeks and 14 weeks both represent end-stage disease. The extent of cytoplasmic aggregation in the cortex, striatum and hippocampus (both dorsal to the CA1 pyramidal layer and in the hilus) was

Figure 2 Continued

transcripts, as measured by qPCR, between R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice at (**E**) 4 weeks and (**F**) 8 weeks of age. At 4 weeks of age, R6/ 2(CAG)₉₀ *Igfbp5*, *Kcnk2* and *Pcp4* transcript levels were dysregulated compared to those of wild type littermates, whereas the R6/2(CAG)₂₀₀ and wild type levels were comparable. At 8 weeks of age, all transcripts were dysregulated to similar levels in both R6/2 lines. n = 8/genotype (4 male and 4 female). Statistical analysis was one-way ANOVA with Bonferroni *post hoc* correction ***P \leq 0.001. The test statistic, degrees of freedom and *P*-values for the ANOVA are provided in Supplementary Table 3.

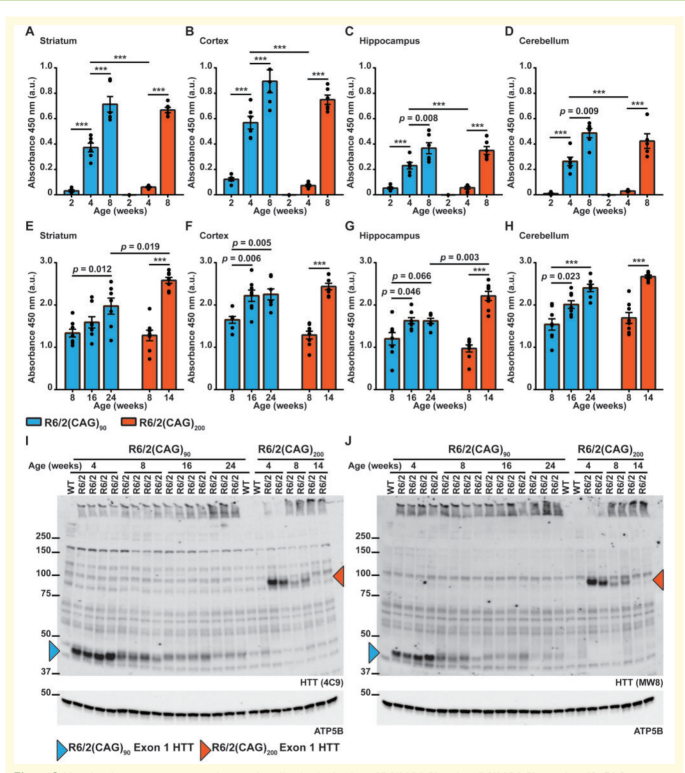


Figure 3 Huntingtin aggregates were detected earlier in the brains of R6/2(CAG)₉₀ **than R6/2(CAG)**₂₀₀ **mice. (A–D)** Seprionligand ELISA analysis of aggregated HTT in the (A) striatum, (B) cortex, (C) hippocampus and (D) cerebellum of R6/2(CAG)₉₀ and R6/ 2(CAG)₂₀₀ mice at 2, 4 and 8 weeks of age. Aggregated HTT was detected at 2 weeks of age in all brain regions from R6/2(CAG)₉₀ mice but not until 4 weeks in R6/2(CAG)₂₀₀ mice. In all brain regions, the level of HTT aggregation was higher in R6/2(CAG)₉₀ mice than in R6/2(CAG)₂₀₀ mice at 4 weeks of age. By 8 weeks of age, the level of HTT aggregation was comparable between R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice for all brain regions. (E–H) Seprion-ligand analysis of aggregated HTT in the (E) striatum, (F) cortex, (G) hippocampus and (H) cerebellum of R6/ 2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice from 8 weeks of age until end-stage disease at 24 and 14 weeks of age respectively. (A–D) n = 6/genotype. (E–H) n = 8/genotype. The absorbance values cannot be compared between (A–D) and (E–H) as these experiments were not performed at the same time. Statistical analysis was two-way ANOVA with Bonferroni *post hoc* correction ****P \leq 0.001. The test statistic, degrees of freedom and *P*-values for the ANOVA are provided in Supplementary Table 4. (I, J) Western blot analysis of soluble and aggregated exon 1 HTT protein in

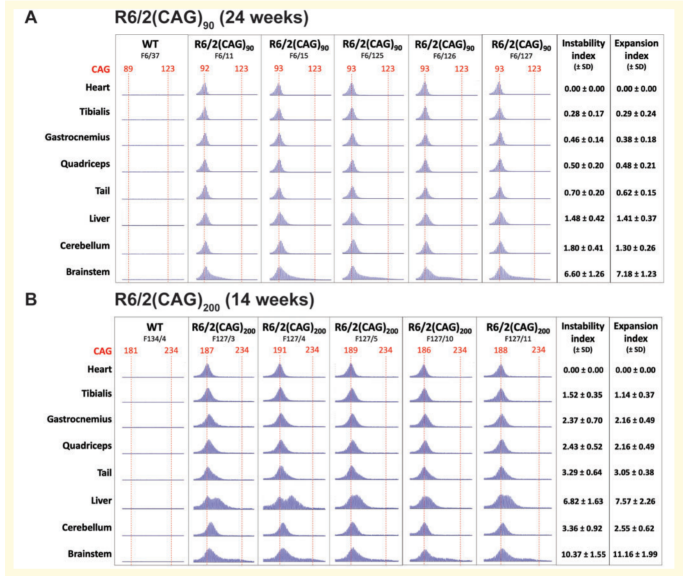


Figure 4 Comparison of somatic CAG instability in tissues from R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice. Capillary electrophoresis traces for (**A**) R6/2(CAG)₉₀ mice at 24 weeks of age and (**B**) R6/2(CAG)₂₀₀ mice at 14 weeks of age. The mode CAG repeat length is indicated at the top of the trace in red and the red dashed line helps to visualize the extent to which the CAG repeat has been somatically unstable during the lifespan of each mouse. Each vertical column shows the traces obtained from the tissues of a single mouse. The mean, standard deviation and data points for the instability and expansion indices of each tissue are illustrated in Supplementary Fig. 2, and the numerical values for these measures are listed on the right hand side of the tables (±SD) in (**A**) and (**B**). Somatic CAG instability was detected in the brain stem from both models, but was only prominent in the liver from the R6/2(CAG)₂₀₀ mice. *n* = 5/genotype. The mean CAG repeat size for R6/2(CAG)₉₀ mice was 93.11 ± 0.86 (SD) and for R6/2(CAG)₂₀₀ mice was 190.53 ± 3.17 (SD). Statistical analysis was one-way ANOVA with Bonferroni *post hoc* correction, ****P ≤ 0.001. The test statistic, degrees of freedom and P-values for the ANOVA are provided in Supplementary Table 5. WT = wild type.

Figure 3 Continued

brain lysates from R6/2(CAG)₉₀ mice at 4, 8, 16 and 24 weeks of age and from R6/2(CAG)₂₀₀ mice at 4, 8 and 14 weeks of age, immunoprobed with either (I) 4C9 or (J) MW8 antibodies. ATP5B was used as a loading control. Aggregated HTT was prominent in the stacking gel for R6/ $2(CAG)_{90}$ mice at 4 weeks, but not from R6/2(CAG)₂₀₀ mice until 8 weeks of age. The migration of soluble exon 1 HTT was retarded by the expanded polyQ tracts and the levels of this soluble protein diminished with age in both lines as they were recruited into aggregates. The full-sized blots for the loading controls are shown in Supplementary Fig. 12. WT = wild type.

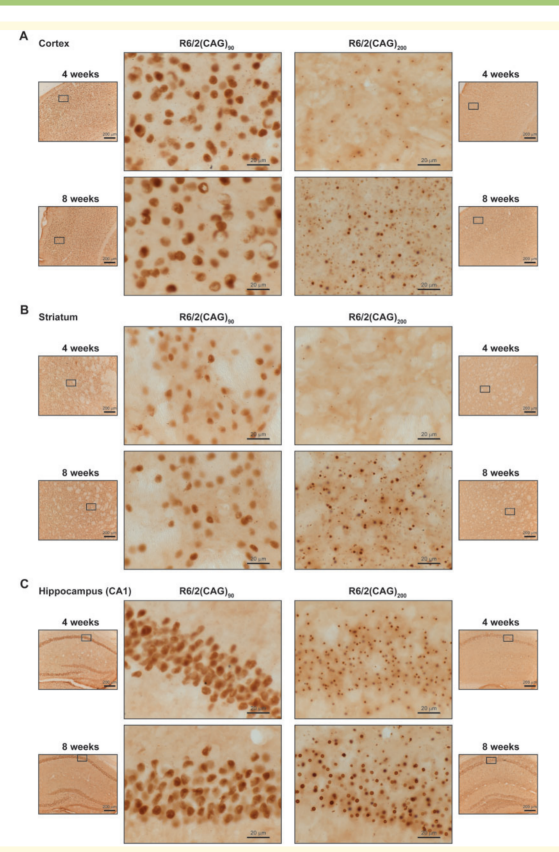
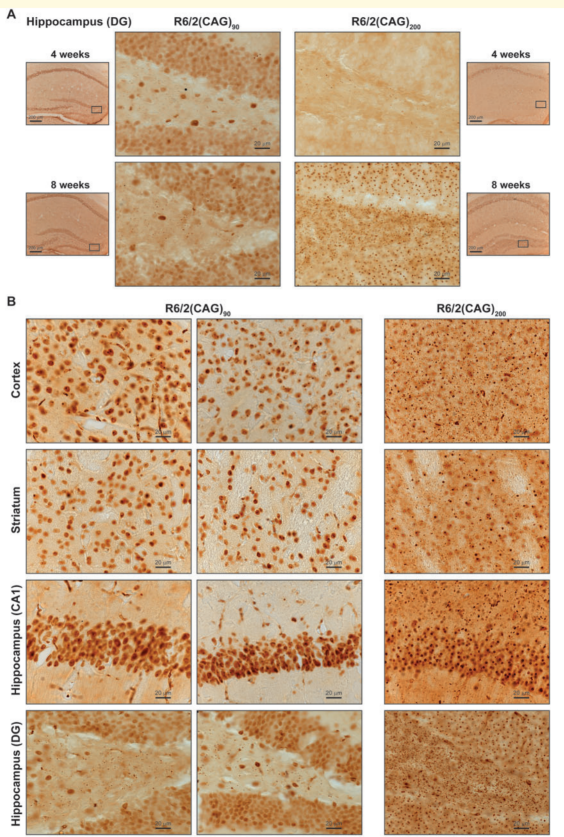


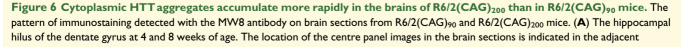
Figure 5 Aggregated HTT accumulated in neuronal nuclei in the brains of R6/2(CAG)₉₀ at a younger age than in R6/2(CAG)₂₀₀ mice. The pattern of immunostaining detected with the MW8 antibody on brain sections from R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice at 4 and 8 weeks of age is shown for the (**A**) cortex, (**B**) striatum and (**C**) hippocampus. The location of the centre panel images in the brain sections is indicated in the adjacent thumbnails, and the position of the thumbnails within the tissue section is illustrated in Supplementary Figs 6 and 7. The wild type control sections are shown in Supplementary Figs 8 and 9. n = 3/R6/2 genotype and n = 1/wild type control. Centre panels, scale bar = 20 µm; thumbnails, scale bar = 200 µm.



16 weeks

24 weeks

14 weeks



To further investigate the distribution of soluble and aggregated HTT in the nucleus and cytoplasm, we prepared nuclear and cytoplasmic fractions from R6/ 2(CAG)₉₀ hemispheres at 4, 8, 16 and 24 weeks of age and from R6/2(CAG)₂₀₀ hemispheres at 4 and 8 weeks of age. The nuclear and cytoplasmic fractions were western blotted and immunoprobed with the S830 antibody (Fig. 7A). Soluble exon 1 HTT was detected in the cytoplasmic fractions and the levels decreased with age for both R6/2 lines, as the soluble protein was recruited into aggregates (Fig. 7A). Consistent with our previous data, soluble exon 1 HTT could not be detected in the nuclear fraction from R6/2(CAG)200 mice. However, we were able to detect a weak signal corresponding to soluble exon 1 HTT in the R6/2(CAG)₉₀ nuclear fraction (Fig. 7A). Examination of the stacking gel showed that, consistent with the immunohistochemical analysis (Fig. 5), nuclear aggregation could be detected at a younger age in R6/2(CAG)₉₀ than in R6/2(CAG)₂₀₀ mice (Fig. 7A). This was further confirmed by Seprion ELISA, showing that the level of aggregated HTT was greater in the R6/2(CAG)₉₀ nuclei at 4 weeks of age but then accumulated rapidly so that by 8 weeks of age the levels in the R6/2(CAG)₂₀₀ nuclei, as measured by the Seprion ELISA were greater (Fig. 7B). The cytoplasmic fractions cannot be used to study aggregation, as a considerable proportion of the cytoplasmic aggregates are lost during the preparation of the fractions, as indicated by the lack of aggregate signal in the cytoplasmic stacking gel (Fig. 7A).

Aggregated HTT in R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mouse brains have different molecular properties

In order to further investigate differences in the HTT aggregates formed in the brains of the $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ brains, we separated the nuclear fractions from 4- and 8-week-old mice by AGERA (Fig. 8A). At 4 weeks of age, the degree of aggregation was not only greater in $R6/2(CAG)_{90}$ mice but the migration was more compact than in $R6/2(CAG)_{200}$ mice. At 8 weeks of age, the level of aggregation was more comparable, but the migration of the aggregates was still visually more compact in $R6/2(CAG)_{90}$ samples.

We then used cortical samples to investigate HTT aggregate properties over the entire disease course of both R6/2 lines. The Seprion ELISA (Fig. 3B and F) and immunohistochemical analyses (Fig. 5A and Supplementary Fig. 5A) had been consistent in demonstrating that the aggregate load was greater in the R6/ 2(CAG)₉₀ cortex at 4 weeks of age. At 8 weeks, although the Seprion ELISA assay suggested that aggregate levels were relatively comparable (Fig. 3A-H), the immunohistochemical staining pattern with MW8 was very different: appearing as a diffuse signal in the R6/2(CAG)90 nuclei and as inclusion bodies in the R6/2(CAG)₂₀₀ nuclei. The filter retardation assay traps larger aggregates on filters, whilst allowing smaller aggregates to wash through (Scherzinger et al., 1999). Filtration of cortical lysates through cellulose acetate filters showed that aggregates from the R6/2(CAG)₂₀₀ brains were trapped on the filters at both 8 and 14 weeks of age, whereas this was not the case from R6/2(CAG)₉₀ brains (Fig. 8B). At all ages, up to 24 weeks of age, aggregates from the $R6/2(CAG)_{90}$ cortices were never trapped on the cellulose acetate membrane (Fig. 8B). Finally, we used the FRET-based aggregate seeding assay (Ast et al., 2018) to assess the seeding competence of aggregates from the same cortices as had been used for the filter-trap assay (Fig. 8C-E). Aggregates from the R6/2(CAG)₉₀ cortices were already seeding competent by 2 weeks of age (Fig. 8C and D), this had increased at 4 weeks but then did not increase any further up to late-stage disease at 24 weeks of age (Fig. 8D). The seeding competence of the R6/2(CAG)₂₀₀ cortical lysates was similar to wild type at 2 weeks of age (Fig. 8C and E) and then increased considerably between 4 and 8 weeks (Fig. 8C and E). Therefore, the cortical aggregates from 2- and 4-week-old R6/2(CAG)₉₀ mice were smaller and had a greater seeding competence than those from $R6/2(CAG)_{200}$ mice.

Discussion

Here, we have used R6/2 transgenic mice expressing an exon 1 *HTT* transgene with $(CAG)_{90}$ and $(CAG)_{200}$ repeats to investigate the effects of polyQ expansions on the molecular and immunohistochemical properties of HTT aggregates and their phenotypic consequences. We showed that nuclear aggregation occurred earlier in R6/ $2(CAG)_{90}$ mice and that this correlated with the onset of transcriptional dysregulation. However, cytoplasmic

Figure 6 Continued

thumbnails, and the position of the thumbnails within the tissue section is illustrated in Supplementary Figs 6 and 7. The wild type control sections are shown in Supplementary Figs 8 and 9. (**B**) The cortex, striatum, CA1 region of the hippocampus and hilus at 16 and 24 weeks of age for the R6/2(CAG)₉₀ and 14 weeks of age for the R6/2(CAG)₂₀₀ mice. The location of the images from within the tissue sections is illustrated in Supplementary Fig. 10. The wild type control sections are shown in Supplementary Fig. 11. n = 3/R6/2 genotype and n = 1/wild type control. (**A**) Centre panels, scale bar = 20 µm; thumbnails, scale bar = 20 µm. (**B**) scale bar = 20 µm.

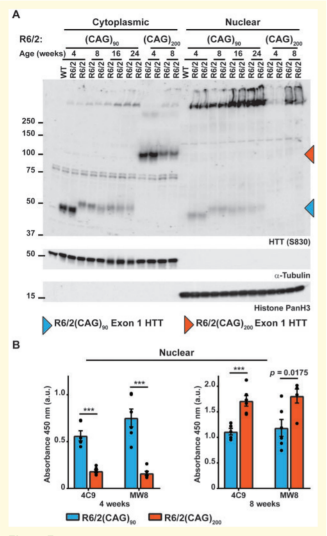


Figure 7 Soluble exon | HTT was detected in the nuclear fraction from R6/2(CAG)₉₀ but not from R6/2(CAG)₂₀₀ brains. (A) Western blot of cytoplasmic and nuclear fractions from R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ hemispheres at the ages indicated immunoprobed with the S830 antibody. A fragment that migrated as soluble exon 1 HTT was detected in the nuclear fraction of R6/ 2(CAG)₉₀ but not R6/2(CAG)₂₀₀ mice. S830 was used for the western blot presented here, as MW8 and 4C9 both detected cross-hybridising bands, which co-migrated with the R6/2(CAG)₉₀ protein, making the presence of exon I HTT in the nuclear fraction harder to interpret. The purity of the nuclear and cytoplasmic fractions was indicated by immunoblotting with antibodies to α tubulin and histone panH3 respectively. (B) Seprion ELISA analysis of HTT aggregation in nuclear fractions from R6/2(CAG)₉₀ and R6/ 2(CAG)₂₀₀ hemispheres at 4 and 8 weeks of age immunodetected with 4C9 or MW8 antibodies (n = 6/genotype). Statistical analysis was two-way ANOVA with Bonferroni post hoc correction. ***P < 0.001. The test statistic, degrees of freedom and P-values for the ANOVA are provided in Supplementary Table 6. WT = wild type.

aggregates accumulated more rapidly in the R6/ $2(CAG)_{200}$ mice, and more closely tracked with the progression of behavioural phenotypes and with end-stage disease. We found that aggregate species formed in the

R6/2(CAG)₉₀ brains have different properties to those in the $R6/2(CAG)_{200}$ mice. Within the nucleus, they retained a diffuse punctate appearance throughout the course of the disease, and could be partially solubilized by detergents. There were a greater number of aggregate seeds in the brains of younger R6/2(CAG)₉₀ mice than those from R6/2(CAG)₂₀₀. Aggregates from R6/2(CAG)₉₀ brains were more compact on AGERA gels and, unlike aggregates from R6/2(CAG)200 brains, did not become trapped after filtration through cellulose acetate membranes as the disease progressed. These data emphasize that understanding the relationship between HTT aggregation and the onset and progression of disease phenotypes in Huntington's disease mouse models, requires a subcellular analysis using multiple complimentary approaches. Soluble HTT and its N-terminal fragments can only be detected in the cytoplasm on western blots in R6/2, BACHD and HdhQ150 mice (Benn et al., 2005; Landles et al., 2010; Gu et al., 2015), and therefore, we had not expected to find that an exon 1 HTT fragment migrated as a soluble protein in the nuclear fraction from the R6/ 2(CAG)₉₀ brains. We have interpreted this as having been released from HTT aggregates for the following reason. The first 17 amino acids of HTT contain a very potent nuclear export signal (Cornett et al., 2005; Zheng et al., 2013), which effectively expels soluble exon 1 HTT from the nucleus in transgenic mice (Benn et al., 2005). This endogenous nuclear export signal is potent and was shown to override an exogenous nuclear localization sequences (NLS), both in cell culture (Cornett et al., 2005) and in vivo (Benn et al., 2005), with the consequence that, despite the addition of an nuclear localization sequence, an exon 1 HTT protein with 20 glutamines was cytoplasmic in transgenic mice (Benn et al., 2005). In contrast, HTT proteins with polyQ tracts in the mutant range accumulate in the nucleus because they form detergent insoluble aggregates. In keeping with this, the addition of an nuclear localization sequence to a HTT protein containing an expanded polyQ tract accelerates the appearance of aggregates in the nucleus, and drives all of the mutant HTT into these nuclear aggregate structures (Schilling et al., 2004; Benn et al., 2005; Gu et al., 2015), which can be solubilized with formic acid (Landles et al., 2010; Gu et al., 2015). Therefore, we infer that, unlike R6/2(CAG)200, the R6/2(CAG)90 brains contain an aggregate species in the nucleus, from which soluble HTT can be released upon exposure to mild detergents in the sample preparation.

Transcriptional dysregulation is a well characterised molecular phenotype for Huntington's disease that has been extensively studied in mice (Luthi-Carter *et al.*, 2000; Langfelder *et al.*, 2016) and post-mortem brains (Hodges *et al.*, 2006), demonstrating that mice usefully replicate this phenotype (Kuhn *et al.*, 2007). Here, we have monitored the expression levels of specific transcripts, reproducibly altered in the brains of Huntington's disease mouse models (Kuhn *et al.*, 2007; Langfelder

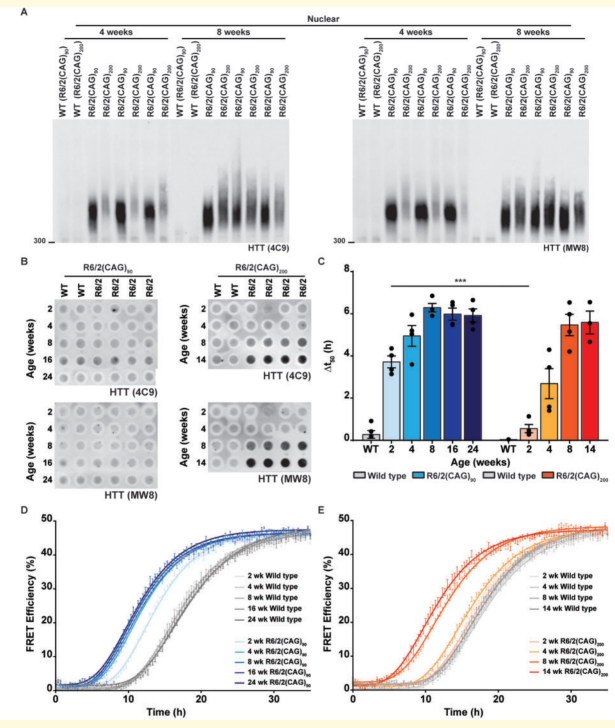


Figure 8 HTT aggregates formed in R6/2(CAG)₉₀ and **R6/2(CAG)**₂₀₀ brains have different molecular properties. (**A**) Nuclear fractions from R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ hemispheres at 4 and 8 weeks of age were fractionated by AGERA and immunoprobed with either 4C9 or MW8 antibodies. (**B**) Cortical lysates from R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ mice were filtered through cellulose acetate filters and immunoprobed with either 4C9 or MW8 antibodies (wild type, n = 2; R6/2, n = 4/genotype/age). (**C**) Quantification of mutant huntingtin seeding activity (HSA) in brain extracts from the R6/2(CAG)₉₀, R6/2 (CAG)₂₀₀ and wild-type control mice (n = 4/genotype/age). Results for wild type mice are shown as an average Δt_{50} value. Cortical samples were from the same mice as had been used for the filter retardation assay in (**B**). (**D**) Effect of cortical homogenates from R6/2(CAG)₉₀ and wild type mice at 2, 4, 8, 16 and 24 weeks of age on the Ex1Q48-CyPet and –YPet (1:1 mixture, 1.2 µM) co-aggregation. (**E**) Effect of cortical homogenates from R6/2(CAG)₂₀₀ and wild type mice at 2, 4, 8 and 14 weeks of age on the Ex1Q48-CyPet and –YPet (1:1 mixture, 1.2 µM) co-aggregation. Statistical analysis was two-way ANOVA with Bonferroni *post hoc* correction. The test statistic, degrees of freedom and *P*-values for the ANOVA are provided in Supplementary Table 7. ***P < 0.001. WT = wild type.

et al., 2016), as markers of transcriptional dysregulation. At 4 weeks of age, all striatal, cortical and some cerebellar transcripts were dysregulated in $R6/2(CAG)_{90}$, but not in $R6/2(CAG)_{200}$ mice. Therefore, the onset of transcriptional dysregulation correlated with the appearance of HTT aggregation and not with the presence of soluble exon 1 HTT which would have been shuttling between the nucleus and cytoplasm in both R6/2 lines. It was striking that the level of transcriptional dysregulation in both the striatum and cortex was comparable between $R6/2(CAG)_{90}$ and $R6/2(CAG)_{200}$ mice at 8 weeks of age, although the immunoprobing with MW8 revealed quite different nuclear structures: a diffuse signal for $R6/2(CAG)_{200}$.

The accumulation of cytoplasmic aggregates was much faster in R6/2(CAG)200 brains and correlated with disease progression. In fact, cytoplasmic aggregates remained relatively sparse in R6/2(CAG)₉₀ brain regions, even at end-stage disease. In a previous study, end-stage disease occurred much earlier, and the range of neurological phenotypes was more extensive in R6/2 mice with aggregate pathology in both the nucleus and cytoplasm, as compared to those with comparable aggregation occurring in the nucleus only (Benn et al., 2005). In keeping with this, heterozygosity for HDAC4, a cytoplasmic protein that interacts with HTT (Mielcarek et al., 2013; Federspiel et al., 2019), delayed aggregation in the cytoplasm, improved behavioural phenotypes, and delayed end-stage disease in R6/2 mice, whilst the levels of nuclear aggregation and transcriptional dysregulation were unchanged (Mielcarek et al., 2013).

The work presented in this study contributes to a larger data set showing that the relationship between CAG repeat length and disease progression in R6/2 mice is complex, with phenotypes generally worsening with increasing CAG repeat length up to ~150 CAGs, after which phenotypes improve as the CAG repeat length increases further (Dragatsis et al., 2009; Morton et al., 2009; Cummings et al., 2012; Morton et al., 2019). The level of huntingtin expression at both RNA and protein levels was lower in R6/2 mice with expansions in 400 CAG range as opposed to 150 CAGs (Dragatsis et al., 2009), consistent with the prediction that unusual DNA structures formed by super-long CAG repeats might impair transcription (Duzdevich et al., 2011). As CAG repeats expanded beyond 150 CAGs, the proportion of cytoplasmic HTT aggregates increased dramatically, presumably because exon 1 HTT with super-long CAG expansions can no longer diffuse into the nucleus. Therefore, the onset and progression of the phenotype is determined by the consequences of a given CAG repeat length on transgene expression, protein aggregation and the subcellular localization of aggregate pathology.

A number of factors could influence the difference in the subcellular location of the aggregates in the R6/ $2(CAG)_{90}$ and R6/ $2(CAG)_{200}$ lines. The kinetics of HTT aggregation are concentration and polyQ length dependent (Scherzinger et al., 1999), and given that we found that the transgene is more highly expressed in the R6/2(CAG)90 mice, both of these factors will influence the differential rate of aggregate formation between the R6/2(CAG)₉₀ and R6/2(CAG)₂₀₀ lines. (i) The probability of aggregates forming in a specific subcellular location will depend upon local exon 1 HTT concentrations, and possibly, how these are influenced by interactions with other proteins and the local proteostasis network. (ii) The concentration of exon 1 HTT in the nucleus will be determined by whether it is small enough to diffuse across the nuclear membrane, a size determined by the polyQ repeat length, which at the cellular level might have increased through the somatic expansion of the CAG repeat. Somatic expansions are more likely to result in the exclusion of exon 1 HTT from the nucleus in the $R6/2(CAG)_{200}$ than the $R6/2(CAG)_{90}$ mice. (iii) Once aggregates have formed in one subcellular location, the recruitment of soluble exon 1 HTT into these aggregates will deplete the levels available for aggregation elsewhere.

HTT aggregates formed in the nuclei of R6/2(CAG)₉₀ mice could be distinguished from those in $R6/2(CAG)_{200}$ mice by immunohistochemistry with MW8, appearing as a diffuse nuclear signal, as compared to a discrete inclusion body, throughout the course of disease in these two lines. This diffuse pattern of nuclear aggregation is similar to that previously published for knock-in mice with 92 glutamines (Wheeler et al., 2000), although at the time of publication, it was not understood that this represented an aggregated form of HTT. Our work demonstrates that the simple correlation of HTT aggregate load in the brain with disease phenotypes is likely to be misleading, and that this must be performed at a subcellular level. Here, we have identified distinct contributions from nuclear and cytoplasmic aggregates to molecular and behavioural phenotypes, and this should be extended to distinguish the consequences of aggregates in other subcellular regions e.g. the synapse. We have shown that aggregates formed from HTT proteins with different polyQ expansions also have different molecular properties as defined by filter retardation, gel electrophoresis and aggregate seeding potential. More sophisticated approaches will now be required to understand what structures these represent, and to distinguish the effects they might have on cellular function. Whilst we do not understand the impact of somatic CAG repeat expansion in generating the pathogenic length of polyQ tracts in Huntington's disease patient brains, it is important that we understand the properties of HTT proteins with a range of polyQ expansions.

Supplementary material

Supplementary material is available at *Brain Communications* online.

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Competing Interests

C.L.B. is an employee of LoQus23 Therapeutics. The authors report no other competing interests.

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