

## ORIGINAL ARTICLE

# Mitochondrial targeting of XJB-5-131 attenuates or improves pathophysiology in HdhQ150 animals with well-developed disease phenotypes

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## Abstract

Oxidative damage to mitochondria (MT) is a major mechanism for aging and neurodegeneration. We have developed a novel synthetic antioxidant, XJB-5-131, which directly targets MT, the primary site and primary target of oxidative damage. XJB-5-131 prevents the onset of motor decline in an *HdhQ(150/150)* mouse model for Huntington's disease (HD) if treatment starts early. Here, we report that XJB-5-131 attenuates or reverses disease progression if treatment occurs after disease onset. In animals with well-developed pathology, XJB-5-131 promotes weight gain, prevents neuronal death, reduces oxidative damage in neurons, suppresses the decline of motor performance or improves it, and reduces a graying phenotype in treated *HdhQ(150/150)* animals relative to matched littermate controls. XJB-5-131 holds promise as a clinical candidate for the treatment of HD.

## Introduction

Mitochondrial dysfunction in Alzheimer's (AD) (1), Parkinson's (PD) (2) and Huntington's disease (HD) (3) is implied by the defects in bioenergetics (4,5), mitochondrial fusion/fission (6), mitochondrial movement (7) and transcription (8,9) that are associated with these diseases. Indeed, elevation of oxidative damage and diminished capacity to produce adenosine triphosphate (ATP) characterize neuronal toxicity (10). Treatment with antioxidants and/or supplementation with ATP producing molecules have been logical therapeutic strategies to offset at least some

pathology that develops in patients. However, these attempts have met with marginal success (3,11–13).

Phosphocreatine serves as a spatial and temporal ATP buffer in tissues with high-energy requirements, such as the skeletal muscle and brain. In HD animal models, creatine improves motor performance, reduces brain atrophy, and reduces striatal Htt-aggregate burden (14,15). However, in both HD and PD patients, creatine, although well tolerated in patients, has not been successful in ameliorating disease symptoms. In PD, creatine supplementation improves mood, but without significant effects on the

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Unified Parkinson's Disease Rating Scale (13,16). The beneficial effects of creatine in HD patients were modest even at exceptionally high creatine levels (1–10 g/day), and the CREST-E trial at even higher levels (40 g/day) was stopped early after a futility analysis (<https://nccih.nih.gov/research/extramural/crest-e>).

Dietary supplementation with naturally occurring Vitamin E and Coenzyme Q<sub>10</sub> (3,17) has been a significant focus of antioxidants therapeutics in PD and HD. CoQ<sub>10</sub> (ubiquinone), an essential biological cofactor of the electron transport chain, functions as an important antioxidant in mitochondrial and lipid membranes. However, when administered as a dietary supplement, CoQ<sub>10</sub> tends to be retained in cell membranes, is inefficient in entering the mitochondria (MT) (18,19) and has low permeability to the blood–brain barrier (BBB) (18,19). Nonetheless, CoQ<sub>10</sub> has had some efficacy in both PD and HD mouse models. For example, CoQ<sub>10</sub> treatment improves early motor deficits in several HD mouse models (20). CoQ<sub>10</sub> treatment protects dopaminergic neurons and inhibits  $\alpha$ -synuclein aggregation in a chemically induced 1-methyl-4-phenyl-1,2,3,6-tetrahydropyridine mouse model of PD (19,21). Despite these positive preclinical outcomes, clinical trials for CoQ<sub>10</sub> were stopped for both PD and HD patients due to the lack of efficacy (13,22).

The sources of the failure of HD therapeutics in clinical trials are unclear. The poor outcome of CoQ<sub>10</sub> in humans is thought, at least in part, to occur due to feedback control, i.e. CoQ<sub>10</sub> is the expressed product of an endogenous gene, and the cell compensates for a dietary increase of CoQ<sub>10</sub> by down-regulating CoQ<sub>10</sub> synthesis (11). Even if plasma concentrations rise during treatment, it remains controversial whether dietary supplementation of CoQ<sub>10</sub> will significantly enhance its steady-state level or reach a pharmacological effective *in vivo* concentration in the brain (11,23,24). Indeed, *in vivo*, the effects of CoQ<sub>10</sub> treatment have been marginal, variable or tissue specific (11,23,25). For example, MitoQ, another mitochondrial-targeted CoQ<sub>10</sub> derivative, failed to offset neurodegeneration in PD clinical trials (26), but has unexpected promise in the periphery, as a therapeutic in treating non-alcoholic fatty liver disease (27).

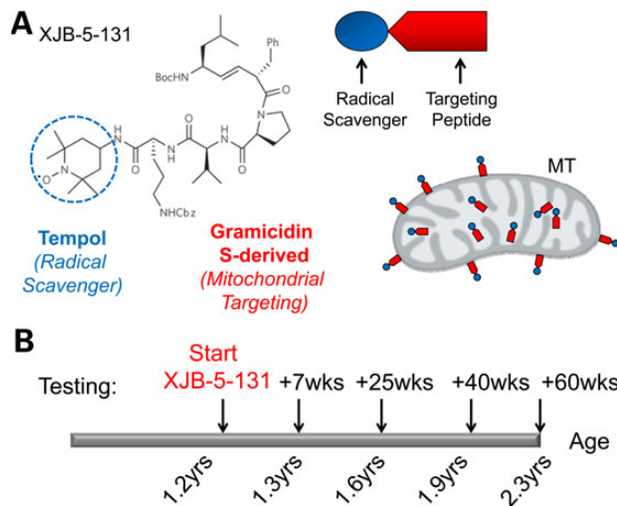
The charge of CoQ<sub>10</sub> and its derivatives can also diminish their efficacy. The loss of the charge gradient is a major feature of MT in dying neurons, yet mitochondrial entry of cationic antioxidants such as MitoQ requires the charge gradient (17). Thus, uptake of these potential-driven antioxidants is self-limiting (28) as there is inevitable depolarization of MT with disease progression. Indeed, MitoQ and MitoVit E protect cultured fibroblasts from Friedrich's ataxia (FRDA) patients; yet, their enhanced potency is abolished in cells pretreated with carbonyl cyanide-4-(trifluoromethoxy)phenylhydrazone, an agent that destroys the charge gradient (29). The diminution of efficacy in FRDA cells raises the issue as to whether MitoQ will have any benefit if treatment starts after disease onset. Many compounds have protective effects in offsetting toxicity in animals if treatment begins early. However, for most compounds, efficacy in counteracting toxicity after disease onset is typically untested, but may be a better predictor of clinical success.

Despite the limitations of antioxidant therapy, the tolerability and/or marginal success in PD and HD patients continues to fuel interest in redox-active therapeutics for neurodegeneration. However, inefficient BBB membrane penetration and the need to sustain pharmacologically effective concentrations in the brain remain significant barriers. To address these issues, we have developed XJB-5-131, a synthetic radical and electron scavenger which directly targets MT, the primary source and the primary target of reactive oxygen species (ROS). XJB's synthetic nature and neutral chemical structure allows it to

permeate membranes. In preliminary characterization, we reported that pretreatment of young animals with XJB-5-131 prevents the decline of Rota-Rod performance in the *HdhQ(150/150)* mouse model for HD. Here, we asked the more difficult, but clinically relevant question: can XJB-5-131 reverse or attenuate toxicity in older animals that have developed disease phenotypes before the start of treatment? Our results demonstrate that XJB-5-131 is a remarkably potent inhibitor of disease progression in animals with well-developed pathology.

## Results

XJB-5-131 is a bi-functional antioxidant comprising a delivery component conjugated to an antioxidant moiety (Fig. 1A) (30). The delivery portion of the molecule is an alkene peptide isostere modification of the Leu-D-Phe-Pro-Val-Orn segment of the antibiotic gramicidin S (31,32) (Fig. 1A, red). This peptide mimetic directly targets the mitochondrial membrane and delivers the antioxidant nitroxide, 2,2,6,6-tetramethyl piperidine-1-oxyl (TEMPO) (Fig. 1A, blue), to neutralize reactive radical species. We evaluated the efficacy of XJB-5-131 in the *HdhQ(150/150)* HD mouse model, which harbored the disease-length 150 CAG tract knocked into both endogenous alleles (33). These animals are referred to as *HdhQ(150/150)* while the littermate controls are referred to as *HdhQ(wt/wt)*. The *HdhQ(150/150)* knock-in model more closely mimics disease progression relative to fragment models (33), and the slow onset provides a large window to observe phenotypic changes with XJB-5-131 treatment. Homozygous *HdhQ(150/150)* animals were used in the analysis to produce the most robust phenotypes. In the following figures, *HdhQ(150/150)* (HD) and *HdhQ(wt/wt)* [wild-type (WT)] animals were treated either with XJB-5-131 (tr) or the saline vehicle (vh), and the treatment groups are abbreviated in the figures as HD-tr, HD-vh, WT-tr and WT-vh, respectively.



**Figure 1.** The structure of XJB-5-131 and treatment strategy. (A) XJB-5-131 is a bi-functional antioxidant comprising a mitochondrial targeting alkene peptide isostere (blue) attached to an antioxidant component (red). The targeting portion of the molecule is derived from the Leu-D-Phe-Pro-Val-Orn segment of the antibiotic gramicidin S. This peptide mimetic directly targets the mitochondrial membrane and delivers the antioxidant, TEMPO (blue box) that is able to prevent or neutralize oxidative damage. MT targeting is depicted as black lines in the red oblong structure of the MT. (B) A schematic diagram of treatment regimen. Animals were aged to 1.2 years (60 weeks) to develop features of HD pathology. XJB-5-131 treatment was initiated at 1.2 years and continued through age 2.3 years. Animals were tested at 7, 10, 25, 40 and 60 weeks after the start of treatment for changes in CNS and motor activity.

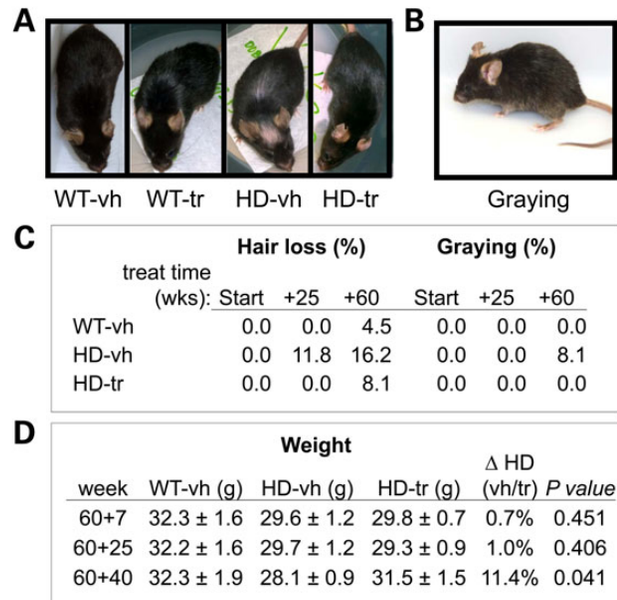
### *HdhQ(150/150)* and littermate controls develop pathology by 60 weeks

HD is characterized by early striatal degeneration, cognitive deficits, involuntary choreiform movements and wasting (34,35), which progress with age. In the experimental paradigm, *HdhQ(150/150)* animals were allowed to age for roughly 1.2 years (60 weeks) to develop features of pathology, at which point treatment began and continued for an additional 1.2 years (60 weeks of treatment) (Fig. 1B). Animals were tested at various times of treatment for physical appearance, central nervous system (CNS) pathology, motor function and the impact of XJB-5-131 on oxidative DNA damage. Collectively, changes in these parameters tested whether the features of disease could be pharmacologically reversed after disease onset (Fig. 1B). Indeed, *Hdh(Q150/Q150)* animals before the start of treatment (60 weeks) had developed measureable pathology (Supplementary Material, Fig. S1). Similar to our previous report, *Hdh(Q150/Q150)* animals weighed less (Supplementary Material, Fig. S1A and B), and performed poorly on the Rota-Rod relative to their *HdhQ(wt/wt)* littermate controls (Supplementary Material, Fig. S1E). At 60 weeks of age, animals of both genotypes had similar neuronal counts (see Fig. 3). However, the level of ubiquitin-containing inclusions were significantly higher in *Hdh(Q150/Q150)* animals (Supplementary Material, Fig. S1C and D). Thus, untreated *Hdh(Q150/150)* animals at 60 weeks were characterized by well-developed phenotypes associated with HD progression.

### Treatment with XJB-5-131 alters physical appearance of aging *HdhQ(150/150)* animals relative to controls

As *HdhQ(150/150)* animals aged, they developed striking physical changes that were blocked by XJB-5-131 treatment. At 1.2 years (60 weeks), animals of both genotypes were well groomed and maintained healthy black coats (Fig. 2A, Panels 1 and 2). However, in *HdhQ(150/150)* animals, 8% developed salt and pepper coats at advanced age (2.3 years), which was prominent around the temples, muzzle and underside (Fig. 2B and C). Furthermore, vehicle-treated animals of both genotypes developed some degree of hair loss at advanced age, but the loss was particularly prominent in older *HdhQ(150/150)* animals (16%) at 2.3 years (Fig. 2A, Panel 3; Fig. 2C). This phenotype was consistent with the excessive grooming activity that has been previously reported in HD animal models (36). XJB-5-131 prevented coat graying (salt and pepper colored coats) (Fig. 2B and C), and significantly reduced hair loss in *HdhQ(150/150)* animals (Fig. 2A and C). The sleek coat appearance of XJB-5-131-treated *HdhQ(150/150)* animals, irrespective of gender, was largely indistinguishable from that of controls at all times tested (Fig. 2A and C).

XJB-5-131 treatment remediated other features of the disease. Weight loss is associated with HD pathology in humans. Before treatment started, both male and female mice were lighter relative to their littermate controls (9% lighter and 13.8%, respectively). Untreated male *HdhQ(150/150)* animals lost an average of 6% of their body weight as they aged from 1.2 to 2.3 years (Fig. 2D). Remarkably, in these animals, XJB-5-131 not only prevented further weight loss during the first 25 weeks of treatment (Fig. 2D, 60 + 25 weeks), but also promoted an average 11.4% weight gain by 60 weeks of treatment (Fig. 2D, 60 + 60 weeks). Surprisingly, there was no effect of XJB-5-131 on the weights of female *HdhQ(150/150)* animals (not shown) and they remained smaller than their *HdhQ(wt/wt)* littermates throughout the experiments. For the most part, XJB-5-131 had obvious beneficial effects on the physiology of *HdhQ(150/150)* animals.

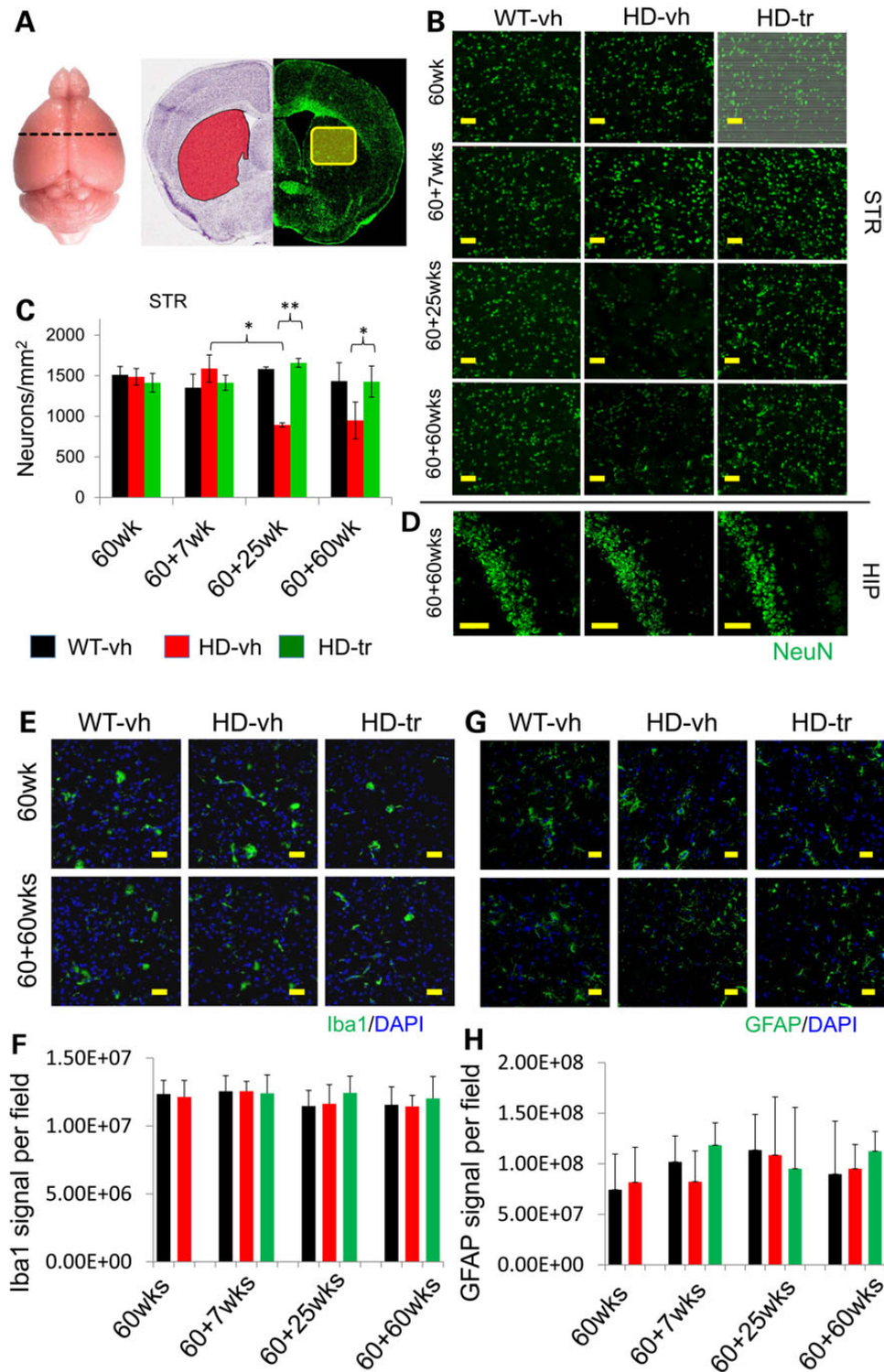


**Figure 2.** XJB-5-131 treatment prevents weight reduction and decreases hair loss and graying phenotypes in *HdhQ(150/150)* and controls with age. (A) A representative example of the hair loss on the forehead and dorsal side (A, Panel 3) that developed in a *HdhQ(150/150)*-vh (HD-vh) mouse at 120 weeks of age. (B) A representative example of 'salt and pepper' coats developed in HD-vh animals at 120 weeks, particularly around the temples, muzzle and underside at old ages (in this case 120 weeks of age). The coat appearance of HD-tr (60 + 60weeks treatment) animals was indistinguishable from that of age-matched WT-vh or WT-tr animals of 120 weeks (A, Panels 1, 2 and 4). (C) Quantification of the hair loss and graying phenotypes (in A and B) among mice at 120 weeks, expressed as a percentage of all animals in their treatment group, WT-vh ( $n = 37$ ), HD-vh ( $n = 46$ ), HD-tr ( $n = 34$ ). (D) The weight of male HD-vh mice with age relative to control littermates. XJB-5-131 promotes weight gain at 40 weeks of treatment ( $P < 0.05$ , Student's *t*-test, one-tailed homoscedastic). WT-vh ( $n = 7$ ), HD-vh ( $n = 12$ ), HD-tr ( $n = 9$ ).

### Treatment with XJB-5-131 reduces neuronal loss and decreases brain lesions in *HdhQ(150/150)* animals

We examined the impact of XJB-5-131 on brain pathology in *HdhQ(150/150)* animals and littermate controls at equivalent ages. The brains of *HdhQ(150/150)* and *HdhQ(wt/wt)* littermates were sectioned and the features compared at the same position in each sample (interaural 3.8 mm, Mouse Brain Atlas) (Fig. 3A). The sensitive striatum (STR) (Fig. 3B) and the more resistant hippocampus (HIP) (Fig. 3D) were stained with anti-NeuN and 4',6-diamidino-2-phenylindole (DAPI) to quantify the neuronal number and density (Supplementary Material, Fig. S2). At the start of treatment (60 weeks), vehicle-treated *HdhQ(150/150)* animals, and their *HdhQ(wt/wt)* littermate controls had similar numbers of neurons in the STR (Fig. 3B and C). However, there was a substantial loss of ~40–50% in aging vehicle-treated *HdhQ(150/150)* animals by 85 weeks (60 + 25 weeks) of treatment that was not observed in the age-matched littermate controls (Fig. 3B and C) (Supplementary Material, Table S1). As judged by histological examination (Fig. 3E–G), there was no measureable gliosis (Fig. 3E and F, anti-Iba1 staining of activated microglia) or changes in the number of glial projections [Fig. 3G and H, anti-glial fibrillary acidic protein (GFAP) staining of astrocytes] in either genotype at any age tested.

The decline in tissue integrity within the affected brain regions was also obvious during tissue handling (Supplementary



**Figure 3.** XJB-5-131 treatment of *Hdh*(Q150/150) prevents loss of striatal neurons in animals with age. (A) Coronal sections of the forebrain were chosen at interaural 3.8 mm as defined by the Brain Atlas (dotted line). At this position, the dorsal portion (yellow outline) of the caudate putamen of the STR (red region) was assayed for neuronal counts. The left side image is Nissl stained (image credit: Allen Institute for Brain Science) with the STR indicated in red. The right side image is IF stained for neurons using NeuN (green). (B) Images of anti-NeuN IF staining of neurons (green) in the dorsal part of the caudate putamen of the STR of animals at various times of treatment (as indicated), which started at 60 weeks of age. Vehicle-treated control animals is (WT-vh), vehicle-treated *Hdh*Q(150/150) is (HD-vh) and XJB-5-131-treated *Hdh*Q(150/150) animals is (HD-tr). Neuronal counts within the STR indicate HD-vh animals lose neurons by 85 weeks of age, whereas XJB-5-131 treatment from the age of 60 weeks significantly prevents neuronal loss and is similar to WT-vh. [ $n = 3$  mice per sample,  $1 \times 10 \mu\text{m}$  brain slice per animal was tested, 10 imaging fields (>900 cells total) per slice] ( $*P < 0.05$ ,  $**P < 0.005$ , Student's *t*-test, one-tailed homoscedastic) (yellow bar indicates  $50 \mu\text{m}$ ). (C) Quantification of the results from all animals (represented in B). (D) Comparison of striatal neuron loss with the neuron numbers of the HIP [in the pyramidal layer (sp) of Ammon's horn field (CA2)]. At the 60 + 60 weeks of treatment, the control CA3 region of the more resistant HIP displayed no neuronal loss. The color key indicates vehicle-treated control animals is (WT-vh), vehicle-treated *Hdh*Q(150/150) is (HD-vh). The images (E) and quantification (F) of anti-Iba1 staining (green) of the striatum; images (G) and quantification (H) of glial projection stained with an anti-GFAP antibody (green). Images are 3D renderings of  $10 \mu\text{m}$  Z-stacks,  $0.5 \mu\text{m}$  spacing per optical slice (yellow bar indicate  $50 \mu\text{m}$ ). Quantifications were of  $n = 3$  mice per sample, five imaging fields per mouse.

Material, Fig. S3). The STR developed a distinctly spongy consistency with age (Supplementary Material, Fig. S3A and B), and frozen sections became increasingly difficult to slice without tearing the tissue (Supplementary Material, Fig. S3B). The number and volume of the striatal gaps (Supplementary Material, Fig. S3B–D, red) increased with age. Tearing was specific to the affected brain regions, i.e. gaps were present around the STR, but in the same slice were absent in more resistant regions of the brain such as the HIP (Supplementary Material, Fig. S3B and C, blue).

Remarkably, XJB-5-131 treatment attenuated the neuronal loss in the STR of *HdhQ(150/150)* and improved tissue quality in these animals as they aged (Fig. 3B and C). The neuronal number and density (Supplementary Material, Fig. S2) (Fig. 3B and C) in the STR of XJB-5-131-treated *HdhQ(150/150)* animals was similar to control littermates up to 2.3 years (Fig. 3B and C). The decline in neuronal number was region-specific as the neuronal number in the more resistant HIP was indistinguishable among treated groups of both genotypes throughout the experiment (Fig. 3D). The consistency of the STR improved with treatment. Tissue tearing and gaps were infrequent in XJB-5-131-treated *HdhQ(150/150)* animals (Supplementary Material, Fig. S3D), whose brain tissue densities and integrity were similar to those of their control littermates (Supplementary Material, Fig. S3B and D).

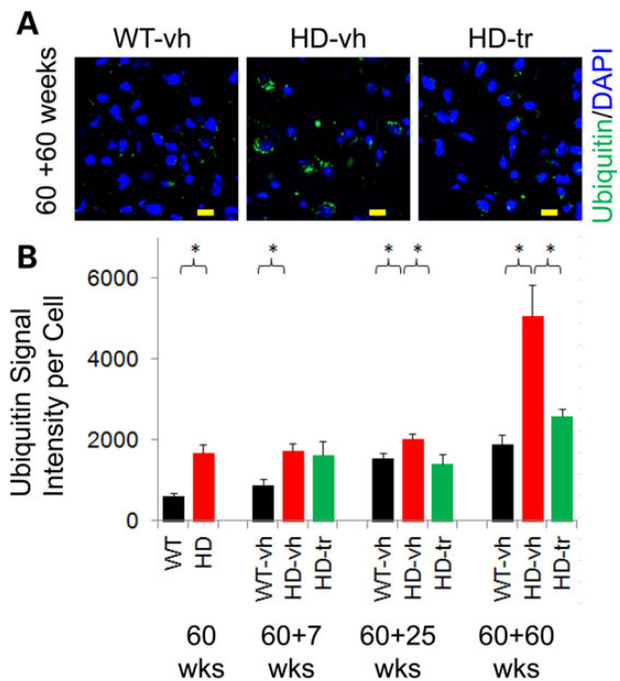
HD pathology is marked by the progressive appearance of inclusions. Indeed, we detected few inclusions in control animals, but aggregates were prominent in age-matched vehicle-treated *HdhQ(150/150)* mice by 60 weeks (Fig. 4A and B). XJB-5-131 treatment suppressed the rise in inclusions compared with vehicle-treated *HdhQ(150/150)* littermates (Fig. 4A and B). Although vehicle-treated *HdhQ(150/150)* animals had developed significant pathology, XJB-5-131 treatment beginning after onset prevented additional loss of neurons and further inclusion formation in animals that had developed disease symptoms before the start of treatment. Taken together, XJB-5-131 had clear benefits in offsetting pathological benchmarks in the CNS of *HdhQ(150/150)* animals.

### Treatment with XJB-5-131 prevents motor decline

Early signs of HD pathology manifest as motor abnormalities which progress with time. To test whether XJB-5-131 was able to improve motor function in aging animals with well-developed pathology, we measured the Rota-Rod performance, grip strength-endurance and open-field activity of 60-week *HdhQ(150/150)* animals at various points following the start of treatment (Fig. 5, from 7 to 60 weeks).

XJB-5-131 administration led to a striking improvement in grip strength-endurance (Fig. 5A and B). Each animal was normalized for its performance at the start of treatment (at 60 weeks) (Fig. 5B) and followed with age. Aging *HdhQ(wt/wt)* littermate maintained good performance within the 25 weeks treatment period (total age was 85 weeks) (Fig. 5C), while vehicle-treated *HdhQ(150/150)* animals substantially declined in the same period (Supplementary Material, Video S1). In contrast, there was no significant decline in grip strength in XJB-5-131-treated *HdhQ(150/150)* animal during the same 25-week treatment period (Fig. 5B, green) (see Supplementary Material, Video S1). XJB-5-131-treated animals were similar to vehicle-treated *HdhQ(wt/wt)* littermates (black), and indistinguishable from XJB-5-131-treated *HdhQ(wt/wt)* (green) controls at advanced age (Fig. 5B).

XJB-5-131 improvement on Rota-Rod performance of *HdhQ(150/150)* animals was equally striking (Fig. 5C). The vehicle-treated *HdhQ(150/150)* animals were characterized by reduced motor

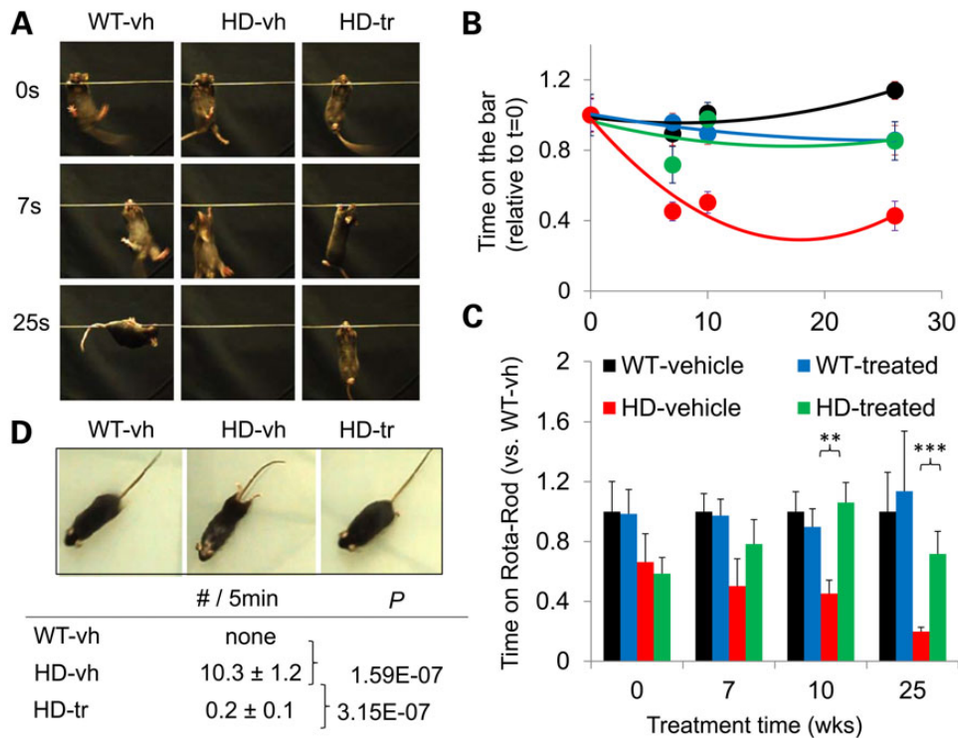


**Figure 4.** XJB-5-131 treatment of *HdhQ(150/150)* animals reduces inclusions in the brain. (A) The signal staining intensity for anti-ubiquitin antibody (green) after 60 weeks of XJB-5-131 treatment (60+60 weeks). Inclusions are elevated in the striatal cells of HD-vh mice relative to striatal cells WT-vh, and are suppressed in HD-tr by XJB-5-131 treatment. Scale bars indicate 10  $\mu$ m. Blue is DAPI staining of DNA in the nucleus. (B) Quantified results of (A) for WT-vh (black) HD-vh (red), and HD-tr (green), as indicated. The start of treatment is 60 weeks; the time of treatment is n, as in (60+n). There are significantly fewer inclusion bodies (measured by IF signal of ubiquitin) after XJB-5-131 treatment in the striatal cells of HD-tr relative to WT-vh and HD-vh and at all measured time points (n = 3 animals per genotype, >250 cells each) (\*P < 0.05, Student's t-test, one-tailed homoscedastic).

performance on the Rota-Rod, which progressed with age. XJB-5-131 treatment of *HdhQ(150/150)* animals significantly improved performance that was well developed at the beginning of treatment (Fig. 5C). Even at advanced age, the Rota-Rod performance of XJB-treated *HdhQ(150/150)* animals was better than that at the beginning of treatment (Fig. 5C, compare 0 and 25 weeks), and was similar to their age-matched *HdhQ(wt/wt)* littermates at all ages tested.

We also tested the open-field activity in vehicle- and XJB-5-131-treated *HdhQ(150/150)* animals using video imaging. Quantified for each animal was the speed of motion, acceleration speed, body area, body axes ratio and the orientation of movement from individual animals, measured at the same time each day (see the 'Materials and Methods' section) (Supplementary Material, Fig. S4A). The activity of animals in the open field was highly variable and sensitive to handling conditions (examples in Supplementary Material, Video S2). Some *HdhQ(150/150)* animals did not move (Supplementary Material, Fig. S4B), but others were active and did not appear distinct from control littermates. Similarly, most control animals had strong exploratory activity (Supplementary Material, Fig. S4C), but others had poor motility. Consequently, the average activity among treated and untreated animals for these parameters was not statistically different.

In contrast, quantitative video analysis revealed a significant frequency of 'foot dragging' in vehicle-treated *HdhQ(150/150)* animals that was not observed in control or in XJB-5-131-treated *HdhQ(150/150)* littermates (Fig. 5D and Supplementary Material,



**Figure 5.** XJB-5-131 treatment of *HdhQ(150/150)* improves motor function with age. (A) Representative examples of the grip test for three treatment groups at three ages, as indicated. The *HdhQ(150/150)*-vh animals lose the ability to grip the bar with age, while treatment with XJB-5-131 prevents this deterioration, as HD-tr animals appear equivalent to vehicle (vh) or treated (tr) WT animals (see Supplementary Material, Video S1). (B) Quantification of (A). The results for the change in performance of each animal were normalized to their performance relative to the start of treatment, which was denoted as one for each animal. Treated diseased animals performed significantly better with age over their untreated counterparts ( $P < 0.05$ ) ( $P =$  Student's t-test, one-tailed homoscedastic). (C) The quantified results of the Rota-Rod assay also indicate that XJB-5-131 improved performance of *HdhQ(150/150)* relative to untreated animals (vehicle, vh). Vh and treated (tr) *HdhQ(150/150)* animals were normalized to the performance of their WT littermates for each time point. The color key identifies the genotypes and treatment groups. (D) XJB-5-131 treatment in *HdhQ(150/150)* animals reduces rear feet dragging phenotypes. A sample video is provided (Supplementary Material, Video S3). The *HdhQ(150/150)* untreated or vehicle-treated mice occasionally dragged their back feet during forward motion at 20–40 weeks of treatment (80–100 weeks of age). Treated *HdhQ(150/150)* had a significantly reduced occurrence of this phenotype ( $P =$  Student's t-test, one-tailed homoscedastic) ( $n = 3$  mice per sample type, five videos of 5 min each were scored per mouse).

Fig. S5) (see Supplementary Material, Video S3). As they pushed themselves forward to initiate movement, these animals were often slow to fully engage their hind legs (Supplementary Material, Fig. S5) (see Supplementary Material, Video S3). These instances were manually counted and corroborated by video analysis as arcs when centroid speed is plotted versus body area (see Supplementary Material, Video S3). An average of 10 occurrences of 'foot dragging' were detected in *HdhQ(150/150)* animals during a 5 min period, while no instances were observed in XJB-5-131-treated littermates or in age-matched *HdhQ(wt/wt)* littermates (Fig. 5D). Collectively, XJB-5-131 significantly reduced the motor deficits in *HdhQ(150/150)* mice.

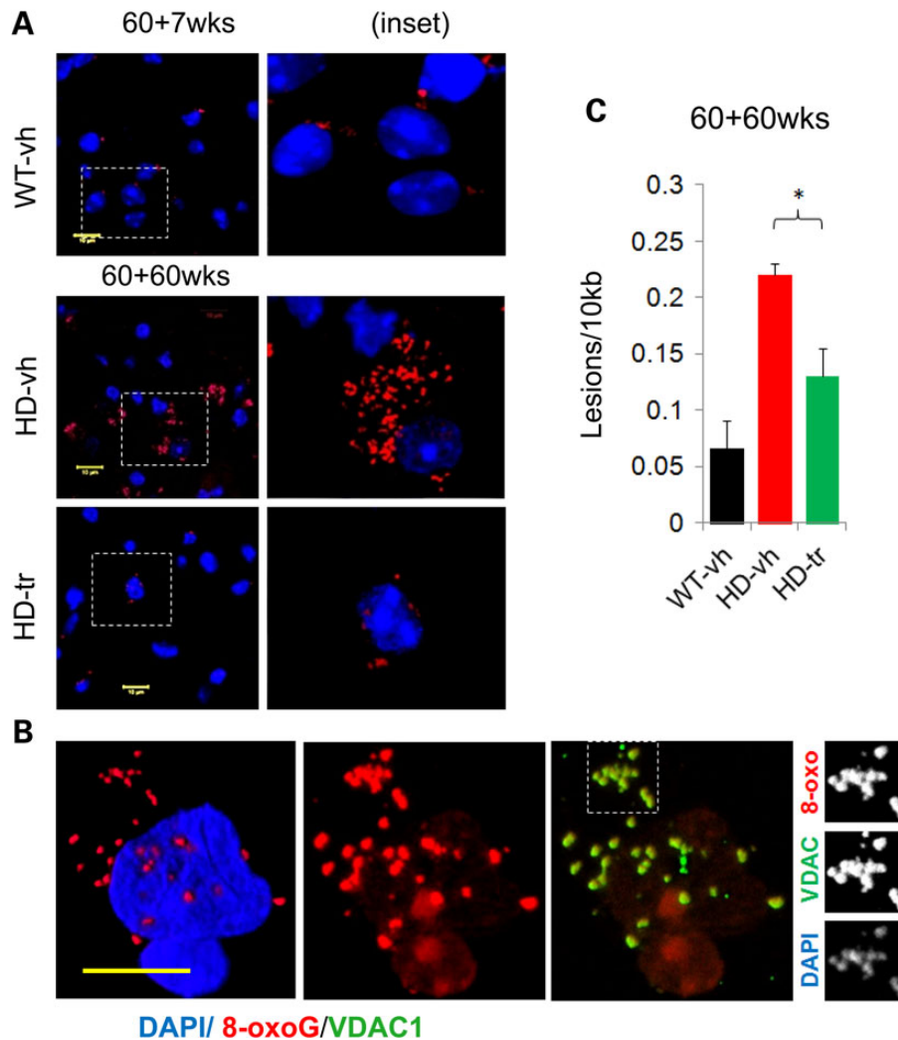
#### Treatment with XJB-5-131 reduces oxidative damage to MT and nuclear DNA in neurons

XJB-5-131 is designed to target MT and reduce oxidative damage. Indeed, we previously published that boron-dipyrromethene (BODIPY)-fluorescent (FL)-XJB-5-131, a FL-BODIPY-labeled derivative of XJB-5-131, crosses the plasma membrane of MT in primary cultured striatal neurons, and co-stained with MitoTracker Deep Red (37). If the beneficial effects of XJB-5-131 occurred by its expected mechanisms, then XJB-5-131 should reduce the level of oxidative damage in brain mtDNA. In the first experiments, we sliced brains sections, used an antibody

to 8-oxo-7,8-dihydroguanine (8-oxo-G) on each slice to determine whether XJB-5-131 reduced staining intensity (Fig. 6).

Oxidation occurs continually throughout life. As expected, the level of 8-oxo-G staining was obvious in the histology section of both genotypes, even in the brain sections of *HdhQ(wt/wt)* animals at early treatment times (Fig. 6B, WT-vh, 60+7 weeks, shown is the STR). However, the level of 8-oxo-G increases significantly in the aging brain, and some of the 8-oxo-G converts to other forms of oxidative damage, which are no longer detected by the 8-oxo-G antibodies (38,39). Thus, antibody staining intensity poorly discriminated between the lesion level in brain tissue from aging *HdhQ(wt/wt)* and *HdhQ(150/150)* animals (Supplementary Material, Table S2), but served as a robust probe of signal reduction by XJB-5-131. Indeed, XJB-5-131 treatment dramatically reduced the level of 8-oxo-G staining in the brains of both *HdhQ(150/150)* and *HdhQ(wt/wt)* littermates (Fig. 6B, compare Panels 2 and 3) (Supplementary Material, Table S2).

In the histology sections, oxidation was present in the nuclear DNA, as judged by the diffuse staining intensity throughout the neuronal nucleus (Fig. 6A, overlay the blue DAPI in Panel 1 to the co-incident diffuse, red 8-oxoG antibody staining in Panels 2 and 3). However, the majority of 8-oxo-G staining was confined to organelles (Fig. 6A, Panels 1 and 2), which were identified as MT by co-staining with the mitochondrial voltage-dependent anion channel (VDAC) (Fig. 6A, compare the red 8-oxo-G staining in



**Figure 6.** XJB-5-131 reduces oxidative damage in MT and somatic DNA. (A) At the start of treatment there are low levels of oxidative DNA lesions in untreated *HdhQ(wt/wt)* (WT-vh) animals as visualized by anti-8-oxo-G IF staining. There is little discernible difference between WT and *HdhQ(150/150)* (HD) littermates at these early ages (up to 85 weeks of age) (image not shown). At late ages (120 weeks), there is a multi-fold increase in DNA lesions in HD animals, which is suppressed by XJB-5-131 treatment, as seen both by IF staining for 8-oxo-G (A,B) and by (C) q-PCR analysis ( $n = 3$  mice per group) ( $*P < 0.05$ ) ( $P =$  Student's *t*-test, one-tailed homoscedastic). (B) 8-oxo-dG (red) is present both in MT and in the nucleus of striatal cells of old HD mice [in this case an *HdhQ(150/150)*-vh mouse 120 weeks of age]. (B, Panel 2) The localization of the oxidative damage is primarily perinuclear foci, along with a fainter and diffuse nuclear pattern. (B, Panel 3 and insets) The foci co-localize with the MT [anti-VDAC (green)] and also with DAPI (viewable when DAPI imaging exposure is increased). Yellow bars represent 10  $\mu\text{m}$  (images  $\sim 21 \mu\text{m}$  confocal Z-stack 3D reconstruction, frame: 84.9  $\mu\text{m}^2$ ).

Panel 2 with green VDAC antibody staining in Panel 3). Indeed, the individual fluorophore patterns (Fig 6A insets) confirmed that 8-oxo-G co-stained with mtDNA and VDAC in the same organelles. The oxidative damage in MT occurred in both neurons and in glial cells, but the lesion level was roughly 3-fold higher in neurons (67% neurons versus 22% of glia) (Supplementary Material, Table S2).

To more accurately compare the overall damage, we isolated the DNA in the MT of the striatum, and used quantitative polymerase chain reaction (qPCR) to measure the level of lesions directly in *HdhQ(wt/wt)* and *HdhQ(150/150)* animals. qPCR intensity increases linearly with the lesion load, and detects a wide range of DNA oxidative lesion based on their ability to slow or to block polymerase progression in DNA (38). Importantly, 8-oxo-G has little impact in polymerase progression, and the detection of this common lesion does not dominate the PCR products. Thus, qPCR better reflected differences in oxidative damage between the *HdhQ(wt/wt)* and *HdhQ(150/150)* animals by quantifying lesions that are typically present at the lower level.

Indeed, qPCR provided evidence that the level of oxidative DNA damage in mtDNA was higher in the brains of vehicle-treated *HdhQ(150/150)* animals relative to their *HdhQ(wt/wt)* littermates (Fig. 6C), and confirmed that XJB-5-131 reduced the level of oxidative damage in *HdhQ(150/150)* animals. Thus, XJB-5-131 operated by its expected mechanism in brain MT. Collectively, the reduced load of oxidative damage after XJB-5-131 treatment was accompanied by rescue of neurons in XJB-5-131-treated *HdhQ(150/150)* animals and improvement in their motor performance.

## Discussion

There are no effective therapeutics available that offset the devastating progressive effects of HD or other neurodegenerative diseases. Previously, we have reported that the synthetic radical and electron scavenger, XJB-5-131, attenuates the decline in Rota-Rod performance if treatment begins at a young age before phenotypes develop (37). Here, we report that XJB-5-131 treatment attenuates or reverses the effects of disease if treatment begins

after disease onset. In aging animals with well-developed pathology, XJB-5-131 treatment promotes weight gain, attenuates neuronal loss, reduces inclusion formation, prevents the performance decline in multiple motor tests and decreases oxidative damage in the brains of aging *HdhQ(150/150)* animals relative to age-matched vehicle-treated *HdhQ(150/150)* animals. The treated *HdhQ(150/150)* animals were often indistinguishable from age-matched *HdhQ(wt/wt)* control animals. XJB-5-131 also reduces the somatic expansion of the CAG repeat tracts in these animals (40), and this feature has been confirmed in R6/2 animals (A. Polyzos and J. Morton, Univ. of Cambridge, submitted for publication). To our knowledge, this is the first demonstration of a sustained reversal/attenuation of pathophysiology by an antioxidant therapy.

The key to its efficacy is the mitochondrial targeting function of XJB-5-131. A peptide mimetic moiety in this compound targets the antioxidant directly to the mitochondrial membrane (Fig. 1A) where the compound is enriched ~600-fold relative to the cytosol. Neither untargeted TEMPO (antioxidant component) nor the peptide mimetic (targeting component) alone are effective in offsetting toxicity (30). We show here that XJB-5-131 effectively reduces ROS in brain MT, but also reduces 8-oxo-G staining in the mtDNA, consistent with its functions in mitochondrial targeting and antioxidant efficacy. Thus, the peptide mimetic portion of XJB-5-131 delivers the antioxidant moiety directly to the site where reactive species are formed (Fig. 1A), and there the drug exerts a sustained homeostatic effect.

TEMPO, the active nitroxide moiety in XJB-5-131, can undergo one- or two-electron transfer processes to hydroxylamines or oxoammonium cations (41,42) thereby supporting redox homeostasis. Due to these redox cycling abilities, XJB-5-131 prevents the leakage of electrons during the reduction of O<sub>2</sub> to H<sub>2</sub>O at cytochrome c-oxidase and avoids the formation of superoxide radical anion (31). The nitroxide moiety is able to return the electrons to the electron transport chain, and therefore XJB-5-131 can exert antioxidant properties without significantly diminishing ATP generation.

XJB-5-131 overcomes some potential sources of past clinical failures of antioxidant therapies. Since XJB-5-131 is a synthetic antioxidant it is not modulated by gene expression, and is capable of achieving and maintaining pharmacologically effective concentrations in cells. XJB-5-131 crosses the BBB and is readily visualized in the brains of rats by *in vivo* EPR imaging (43). Thus, the synthetic compound has proven direct access to the neurons. Furthermore, XJB-5-131 is charge neutral and does not influence the mitochondrial membrane potential nor does it depend on the electrostatic gradient to enter MT. This property of XJB-5-131 provides a mechanism for improving mitochondrial function even after membrane depolarization.

Taken together, direct targeting of a synthetic antioxidant to MT is poised to provide a level of specificity and efficacy that enables a viable therapeutic strategy. Because mitochondrial decline is a common and central feature of toxicity in the brain, targeted nitroxides such as XJB-5-131 are expected to be useful in a wide spectrum of neurodegenerative diseases, or in other deleterious ROS-induced conditions and mitochondrial malfunctions. Indeed, we have previously shown that XJB-5-131 and the related JP4-039 have efficacy in a rat model of hemorrhagic shock (44), a mouse model of radiation mitigation (45), a rat model of traumatic brain injury (43) and on cardiac resistance to ischemia-reperfusion-induced oxidative stress in aged rats (46).

A therapeutic compound is far less useful if it cannot reverse the disease phenotypes that have developed in a patient before treatment. The lack of such efficacy in animal testing may account, at least in part, for the current disconnect between preclinical and clinical outcomes. Delivery of XJB-5-131 to neuronal

protects MT against damage before or after onset in models of advanced HD. XJB-5-131 has not yet been tested in humans for efficacy, however, based on the preclinical results reported here, a compound with these or similar properties is desirable. Individuals with HD are typically aware that the disease gene is passed down in their family line, but not all individuals in a family will choose to be genetically tested. Thus, subjects who have a confirmed diagnosis may opt to start treatment with an effective therapeutic at a young age. At the same time, a compound that is effective after the development of phenotypes would bring enormous benefits to patients who choose not to be tested or are unaware of the disease but develop symptoms later in life.

The need for new therapeutics for neurodegeneration is extraordinary. Due to rapid increases in the elderly populations, age-related diseases are expected to grow exponentially in the coming years (47). Among these, neurodegenerative diseases will be the most devastating in terms of their emotional and economic impacts. Because the number of affected individuals will grow significantly, the gap between the size of the problem and our capabilities for treatment will widen. Recently, a reduction in the level of expanded mhtt protein by anti-sense RNA to human Huntington has led to reversible efficacy in an animal model (48). XJB-5-131 alone is able to produce the same outcome based on its distinct mechanism of action. These features support XJB-5-131's role as a promising clinical candidate, and warrant further investigation of this class of compounds for their efficacy either as a stand-alone therapeutic or in combination with other treatments.

## Materials and Methods

### Animals, breeding and weights

The Institutional Animal Care and Use Committee approved all procedures. Animals were treated under guidelines for the ethical treatment of animals, and approved by IACUC protocol #274005 at Lawrence Berkeley Laboratory. All animal work was conducted according to national and international guidelines. The following mouse models were used: homozygous *HdhQ(150/150)*, which have been previously described and characterized (33). The original *HdhQ(150/wt)* line (33) was used in these experiments, and not a derived line that is commercially available, but was generated later (strain B6.129P2-Htt<sup>tm2Det1</sup>/150) (Jackson Labs, Bar Harbor, ME, USA). The *HdhQ(150/wt)* were generated on the C57Bl6 background. The homozygous mice are referred to as *HdhQ(150/150)* and their WT littermates are referred to as *HdhQ(wt/wt)*.

### Treatments and groups

Mice were divided into three treatment groups: untreated, vehicle treated (vh) and XJB-5-131 treated (tr) (at 2 mg/kg) for each genotype [*HdhQ(150/150)* and their WT littermates]. There are 30 mice in each group. XJB-5-131 was administered by intra-peritoneal injection three times per week for the duration of the study. Vehicle treatments were identical except that XJB-5-131 was replaced by filtered phosphate-buffered saline (PBS). Mice were weighed, and tested for behavior beginning at 60 weeks, and after 7, 10, 25 and 40 or 60 weeks of treatment. Treatment began ~60 weeks, followed by treatment period of up to 60 weeks. The animal age at the end of the study was ~120 weeks.

### XJB-5-131

XJB-5-131 was synthesized as previously reported (3). Lyophilized, powdered XJB-5-131 was reconstituted in dimethyl sulfoxide at a

concentration of 1 mg/μl, as previously described (49). These samples were aliquoted and kept at –80°C. On the day of injection, the XJB-5-131 solution was mixed with 0.2 μm filtered and pre-warmed PBS (100°C) and heated for 10 s to reach a final concentration of 2 mg/kg mouse body weight in 200 μl solution. The solution (200 μl) was injected within 30 min of preparation.

### Rota-Rod

Motor activity was tested with a Rota-Rod (UgoBasile, USA/Italy), as previously described (7,37,40). Weights of each animal were recorded once per week. Animals in each group were evaluated for Rota-Rod performance and grip strength at the indicated ages. Mice were lowered on to the already spinning Rota-Rod at the required speed (20 rpm was used in this study). The amount of time the animals stayed on the Rota-Rod was determined by a built-in magnetic trip-switch, which was stopped when the animal fell-off. Mice were measured for Rota-Rod activity for a maximum of 120 s, with three attempts given for each mouse. Animals were trained for one session each day for 5 consecutive trials, and tested for 3 consecutive days at each time point (0, 7, 10 and 25 weeks of treatment). The best (longest) performance was recorded and used for the comparative analysis.

### Grip test

For the grip strength-endurance test, mice were lowered onto a parallel rod ( $D < 0.25$  cm) placed 50 cm above a padded surface. The mice were allowed to grab the rod with their forelimbs, after which they were released and scored for the length of time they could hold onto the bar (maximum 30 s). Mice were tested consecutively three times for each time point (at 0, 7, 10 and 25 weeks of treatment). The maximum length of time they were able to hold on was recorded for analysis.

### Antibodies, immunofluorescence and quantification

Primary antibodies used were mouse anti-NeuN Alexa488 conjugate (Millipore #MAB377X) (used at 1:400), mouse anti-GFAP Cy3 conjugate (Abcam #ab49874) (used at 1:400), rabbit anti-Iba1 (Wako #019-19741) (used at 1:400), rabbit anti-Ubiquitin (DAKO #Z0458) (used at 1:400), mouse anti-8-oxo-dG (Trevigen #4354-mc-050) (used at 1:400) and rabbit anti-VDAC (Thermo Fisher #PA1-954A) (used at 1:400). Secondary antibodies used were goat anti-rabbit Alexa-555 conjugated (Invitrogen #A31630) (used at 1:400), goat anti-mouse Alexa-488 conjugated (Invitrogen #A31620) (used at 1:400). Brain sections cryoembedded in optimal cutting temperature (OCT) were sliced (10 μm thick using a Leica Cryostat set at: –14°C for the sample and –12°C for the blade) and placed onto Histobond microscope slides (VWR). They were immediately fixed and OCT removed in 100% methanol (10 min). Samples were rehydrated in 75, 50, 25 and 0% ethanol in PBS (2 min each). Tissue was treated with Image-iT FX signal enhancer (Thermo Fisher #I36933) to reduce autofluorescence (30 min) and blocked (2–18 h) in blocking solution (PBS, 3% bovine serum albumin, 5% goat serum, 0.7% donkey serum, 0.03% triton X-100). Antibody staining was performed overnight, followed by three washes with PBS (5 min ea.). Secondary antibody was applied later as required along with 0.5 μm DAPI (1–2 h) followed by three washes in PBS. Slides were coated with Vectashield + DAPI, sealed with a coverslip and stored (–20°C) until they were imaged. Slides were imaged using a Zeiss 710 confocal microscope using either of 20 × (0.8N/A)/air, 40 × (1.2N/A)/water or 100 × (1.4N/A)/oil lenses. Image analysis was done using ImageJ: Fiji (50).

### Video imaging and foot dragging quantification

Mouse behavior in the open-field arena was tracked at high-resolution from raw videos (30 Hz) processed with Matlab scripts (The MathWorks) as a custom-made adaptation of freely available animal tracking software (51). Quantitative analyses of low-level kinematic quantities were based on the animal centroid coordinates, its instantaneous speed and acceleration, and morphological features such as the animal's body area, and its orientation via an ellipsoidal fit on its posture. Foot dragging effects were suggested via phase plots combining centroid speed as a function of body area.

### Lesion counts by quantitative PCR

The relative level of mtDNA lesions in the mouse brain was performed as previously described (52,53). The determination of mtDNA lesion number consisted of amplifying a 10 kbp mtDNA fragment by performing an initial denaturation for 45 s at 94°C, followed by 22 cycles of denaturation for 15 s at 94°C, annealing/extension at 64°C for 12 min, and a final extension for 10 min at 72°C. We used the following primer nucleotide sequences: 5'-CCAGTCCATGCAGGAGCATC-3' (forward) and 5'-CGAGAAGA GGGGCATTGGTG-3' (reverse). Levels of damage were calculated as the relative amplification of the *HdhQ(150/150)* STR compared with the WT *Hdh(wt/wt)* controls. The data were corrected for possible changes in mtDNA replication rates. The results were derived from three qPCR assays in triplicate on each animal.  $n = 7$  WT-vh mice;  $n = 9$  HD-vh mice and  $n = 5$  HD-tr mice.

### Supplementary Material

Supplementary Material is available at HMG online.

Conflict of Interest statement. None declared.

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