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OPEN Motivational, proteostatic and transcriptional deficits precede synapse loss, gliosis and neurodegeneration in the B6.*Htt^{0111/+}* model of Huntington's disease

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We investigated the appearance and progression of disease-relevant signs in the B6.*Htt*^{O111/+} mouse, a genetically precise model of the mutation that causes Huntington's disease (HD). We find that B6.Htt^{0111/+} mice are healthy, show no overt signs of central or peripheral inflammation, and no gross motor impairment as late as 12 months of age. Behaviorally, we find that 4–9 month old B6.*Htt*^{011/+} mice have normal activity levels and show no clear signs of anxiety or depression, but do show clear signs of reduced motivation. The neuronal density, neuronal size, synaptic density and number of glia is normal in B6.*Htt*^{O111/+} striatum, the most vulnerable brain region in HD, up to 12 months of age. Despite this preservation of the synaptic and cellular composition of the striatum, we observe clear progressive, striatal-specific transcriptional dysregulation and accumulation of neuronal intranuclear inclusions (NIIs). Simulation studies suggest these molecular endpoints are sufficiently robust for future preclinical studies, and that B6.*Htt^{Q111/+}* mice are a useful tool for modeling disease-modifying or neuroprotective strategies for disease processes before the onset of overt phenotypes.

Huntington's disease (HD) is an autosomal dominant neurodegenerative disease caused by expansion of a glutamine-coding CAG trinucleotide repeat near the 5' end of the HTT gene¹. The identification of the disease-causing mutation did not point to an obvious set of therapeutic approaches, given that the huntingtin protein is a very large HEAT/HEAT-like repeat solenoid scaffold², which is highly evolutionarily conserved, and vital to normal development³. A key effort in preclinical HD research is understanding the link between polyglutamine expansion in huntingtin and the selective cellular toxicity in corticostriatal circuits which is thought to cause disease symptoms. A range of animal models of the HD mutation have been developed to investigate this question, including knock-in and transgenic mice as well as transgenic rats, sheep, pigs and nonhuman primates⁴. Unique amongst described models, knock-in mice with varying allele sizes provide the ability to quantitatively study the

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Age	Sex	Genotype	n	Measured CAG (SD)	Terminal Body Weight (SD)
3 Months	М	B6.Htt ^{+/+}	9		24.3 (1.7)
3 Months	М	B6.Htt Q111/+	9	113 (3.3)	23.8 (1.4)
3 Months	F	B6.Htt ^{+/+}	9		18.2 (0.7)
3 Months	F	B6.Htt Q111/+	9	114 (2.5)	17.2 (1.4)
9 Months	М	B6.Htt ^{+/+}	9		28.2 (1.8)
9 Months	М	B6.Htt Q111/+	9	113 (5.0)	31.4 (1.8)
9 Months	F	B6.Htt ^{+/+}	9		23.3 (1.2)
9 Months	F	B6.Htt Q111/+	9	113 (3.3)	23.5 (1.6)
12 Months	М	B6.Htt ^{+/+}	9		30.6 (2.6)
12 Months	М	B6.Htt Q111/+	9	N/A	31.4 (1.6)
12 Months	F	B6.Htt ^{+/+}	9		25.2 (3.7)
12 Months	F	B6.Htt ^{Q111/+}	9	N/A	26.1 (2.5)

Table 1. Cohort 2 characteristics.

relationship between CAG size and pathogenesis in an otherwise isogenic system⁵. Recent molecular⁶ and behavioral⁷ analyses of an allelic series of HD knock-in mice reveal very discrete CAG-dependent disease signatures, suggesting these mice recapitulate a cardinal feature of human HD, namely the relationship between CAG tract length and the rapidity of disease progression^{3,8}. Because knock-in models express *Htt* at endogenous levels from the endogenous locus they precisely mimic the genetics of human HD, however their use in preclinical studies has been limited because their overt neurological signs are very subtle compared to transgenic animals^{9,10}. The bulk of preclinical research has therefore been conducted in transgenic models expressing either full-length or short fragments of mutant huntingtin, which display more phenotypes¹¹.

The embracing of transgenic models by the field has largely been driven by the desire to model important signs and symptoms of HD, including progressive striatal atrophy and hyperkinetic motor disturbances. We consider that these phenomena emerge from a substrate of decades of progressive cellular and physiological dysfunction before manifesting in human patients as a set of clinical signs recognizable as "Huntington's disease"¹². Targeting these late features of the disease process, including cell death and emergent motor symptoms, may enable the modification of late HD symptoms. However, this strategy doesn't allow targeting of the earliest disease changes that precede overt clinical disease features, which if clearly identified might ultimately provide paths to prevention and/or early-stage reversal. Finally, no HD mouse model develops chorea-like hyperkinetic movements, and mouse models present with modest striatal volume loss over the lifespan of the mouse^{13,14}, while human disease results in the nearly complete atrophy of this critical structure¹⁵.

We and others acknowledge that knock-in mouse models of HD cannot fully recapitulate clinical symptoms that, in human mutation carriers, require approximately four decades of pathogenic processes to emerge and are followed by a further 15 years to progress to death¹⁶. However, these genetically faithful but subtle mouse models provide us the ability to study the earliest and subtlest changes caused by the CAG expansion of *Htt* - the very processes we wish to understand if we are to develop interventions to prevent the development of clinical HD onset, rather than addressing its symptoms only after significant damage to key processes has been incurred. Towards this end, we have here characterized the earliest observable changes in the murine B6.*Htt*^{Q111/+} (hereafter *Htt*^{Q111/+}) model of the HD mutation¹⁷. Since their creation, these mice have been utilized to test specific hypotheses in a wide range of studies¹⁸, and a single unbiased phenotypic screen¹⁰, but have not been well characterized as a system for conducting preclinical studies. We therefore examined the incidence, progression and statistical power of a range of phenotypes in these mice, and find that they provide a powerful tool for studying hypotheses about the early disease events that stem from the HD CAG expansion mutation. We observe robust transcriptional, proteostatic and motivational phenotypes which progress from 3 to 12 months of age in *Htt*^{Q111/+} mice. Simulation studies suggest interventions that rescue specific signs by 10–25% are readily detectable in appropriately powered preclinical studies, making the *Htt*^{Q111/+} mouse a useful tool for testing disease modifying therapeutics in HD.

Results

General Animal Health and Inflammation. Consistent with previous reports of $Htt^{Q111/+}$ mice on the CD1 and B6J strains^{9,10}, no body weight changes were observed between 3 and 12 months of age (effect of genotype, $F_{(1,94)} = 3.1$, p = 0.07, mean CAG length = 113, cohort summary in Table 1). Important plasma chemistry parameters, including protein, glucose and ion concentrations, as well as liver enzyme levels, were normal at 4 months of age in $Htt^{Q111/+}$ mice¹⁰. Because we are interested in progressive phenotypes, we examined a suite of clinical chemistry parameters at 12 months of age - no changes were observed in any of these parameters in male $Htt^{Q111/+}$ mice at this age (Table 2). Consistent with findings in younger mice, and coupled with body weight observations, these results suggest that up to 12 months of age $Htt^{Q111/+}$ mice are grossly healthy. Subtle increases in concentrations of pro-inflammatory chemokines and cytokines have been observed in presymptomatic HD mutation carriers¹⁹ and some (e.g. YAC128), but not all (e.g. BACHD), transgenic HD mice by 12 months of age²⁰. To understand whether increased peripherally-detectable inflammation is occurring in $Htt^{Q111/+}$ mice we quantified cytokines, chemokines and acute phase reactants in the plasma of 12-month-old, male $Htt^{Q111/+}$ mice

Analyte	Units	+/+(SD)	Q111/+ (SD)	T-test
EGF	pg/mL	201 (31.7)	211 (24.6)	0.64
Eotaxin	pg/mL	2170 (1044)	1368 (173)	0.18
IL-1 beta	ng/mL	8.975 (2.1)	9.45 (1.3)	0.71
IL-18	ng/mL	20.8 (4)	25.3 (1.5)	0.08
IP-10	pg/mL	60 (42.2)	64.5 (19.8)	0.85
LIF	pg/mL	915.3 (173.6)	722.8 (129.3)	0.13
MCP-1	pg/mL	55.5 (28.9)	43 (17.6)	0.49
MCP-3	pg/mL	293.5 (195.4)	204.8 (30.2)	0.4
MCP-5	pg/mL	17.7 (7.5)	16.5 (3.3)	0.78
M-CSF-1	ng/mL	3.45 (1.1)	4.3 (0.2)	0.19
MDC	pg/mL	2157.5 (1099.5)	1470 (216.9)	0.27
MIP-1 alpha	ng/mL	5.4 (0.8)	5.4 (0.5)	1
MIP-3 beta	ng/mL	3.3 (0.6)	2.5 (0.2)	0.05
Thrombopoietin	ng/mL	35 (7.7)	37.5 (5.2)	0.61
TIMP-1	ng/mL	1.4 (1)	1.15 (0.1)	0.65
VEGF-A	pg/mL	185.5 (28.4)	192.5 (27.8)	0.74
Glucose	mg/dl	253.3 (20.6)	238.2 (85.4)	0.74
BUN	mgld	22.8 (4.2)	23 (4.5)	0.93
Creatine	mg/ld	0.05 (0.05)	0.02 (0)	0.4
Phosphate	mg/dl	6.45 (0.1)	7.68 (1.1)	0.06
Total Protein	g/dl	3.5 (0.1)	3.4 (0.2)	0.48
Albumin	g/dl	2 (0.01)	1.94 (0.1)	0.41
Globulin	g/dl	1.5 (0.05)	1.46 (0.1)	0.82
Albumin/Globulin	Ratio	1.4 (0.06)	1.34 (0.2)	0.91
Total Bilirubin	mg/dl	0.1 (0)	0.18 (0.2)	0.41
ALP	U/L	0.75 (0.5)	0.8 (0.4)	0.88
GGT	U/L	1 (0.8)	1.2 (0.4)	0.65
ALT	U/L	33 (11.4)	40.4 (24.4)	0.6
AST	U/L	60 (17.6)	78.6 (32.4)	0.34
Cholesterol	mg/dl	54.5 (11.2)	56.2 (15)	0.86

Table 2. Clinical chemistry parameters are normal in 12 month old *Htt*Q^{111/+} mice. Top level analytes are from inflammationMAP (Myriad RBM, Austin, TX, USA), lower level analytes are from clinical chemistry panel (Phoenix Central Labs, Mukilteo, WA, USA). Abbreviations: epidermal growth factor (EFG), interleukin (IL), interferon gamma-induced protein (IP), monocyte chemoattractant protein (MCP), macrophage colony-stimulating factor (M-CSF), macrophage-derived chemokine (MDC), macrophage inflammatory proteins (MIP), tissue inhibitor of metalloproteinase (TIMP), vascular endothelial growth factor-A (VEGF-A), alkaline phosphatase (ALP), gamma-glutamyl transferase (GGT), alanine aminotransferase (ALT), aspartate aminotransferase (ASP). Inflammation markers measured below LLoQ (LLoQ listed): Fibroblast growth factor (FGF)–9 (2.4 ng/mL), FGF-basic (9.6 ng/mL), granulocyte-macrophage colony-stimulating factor (15 pg/mL), keratinocyte chemoattractant/human growth-regulated oncogene (0.028 ng/mL), interferon gamma (182 pg/mL), IL-1 alpha (100 pg/mL), IL-2 (70 pg/mL), IL-3 (5.9 pg/mL), IL-4 (60 pg/mL), IL-5 (0.6 ng/mL), IL-6 (4.7 pg/mL), IL-7 (0.19 ng/mL), IL-10 (125 pg/mL), IL-11 (56 pg/mL), IL-12p70 (0.091 ng/mL), IL-17A (0.018 ng/mL), MIP-1 beta (142 pg/mL), MIP-2 (11 pg/mL), oncostatin-M (0.24 ng/mL), stem cell factor (652 pg/mL), tumor necrosis factor-alpha (0.082 ng/mL). N=4 male Htt^{2111/+}, 4 male Htt^{+/+}, subset of 12-month mice from Table 1).

(multi-analyte profiling, Myriad RBM). We successfully quantified 16 of these molecules (Table 2: Eotaxin, EGF, IP-10, IL-1 β , IL-18, LIF, M-CSF-1, MDC, MIP-1 α , MIP-3 β , MCP-1, MCP-3, MCP-5, Thrombopoietin, TIMP-1, VEGF-A) but observed no changes in 12-month-old $Htt^{Q111/+}$ mice. Other important pro-inflammatory molecules (FGF9, FGF-basic, GM-CSF, KC/GRO, IFN- γ , IL-1 α , IL-2, -3, -5, -6, -7, -10, -11, -12p70, -17a, MIP1- β , OSM, SCF, and TNF- α) were also assayed, but fell below the indicated limit of detection in all mice studied (lower limit of quantification, LLoQ, provided in Table 2). These data suggest that peripherally-detectable immune activation is either not occurring in 12-month-old, male $Htt^{Q111/+}$ mice, or is sufficiently subtle to be missed by these analyses.

Striatal Transcriptional Alterations. Altered transcription is an early and prominent feature of HD pathology, and is observed in both animal models and samples from human patients¹². Because we are interested in intervening at the earliest possible point in the pathogenic process, we investigated the time course of transcriptional changes in the central nervous system of $Htt^{Q111/+}$ mice. As an initial, untargeted pilot study,





we examined striatal and cerebellar genome-wide transcriptional changes in 3- and 9-month-old $Htt^{Q111/+}$ mice using mRNA sequencing (RNASeq). At 3 months of age, no robust transcriptional alterations were detected in the striatum of $Htt^{Q111/+}$ mice compared to wild-type mice (Fig. 1a, 5 quantified transcripts - *Lct, Zfp385c, Smpx, Dsp, Crlf1* - have an effect of genotype with a false discovery rate (FDR) of < 5%). By 9 months of age, 726 transcripts were changed in the striatum of $Htt^{Q111/+}$ mice compared to wild-type mice (Fig. 1a, 245 up-regulated, 481 down-regulated at FDR < 5%). The cerebellum, despite robustly expressing mutant Htt^{21-23} , is relatively spared from pathological tissue loss in human HD patients²⁴ and the YAC128 mouse model of HD¹⁴. We therefore compared the rate and scale of transcriptional dysregulation between the cerebellum and striatum of $Htt^{Q111/+}$ mice at 3 and 9 months of age. There were very few significant effects of CAG expansion in Htt in the cerebellum at either age (Fig. 1a, 3 genotype-sensitive transcripts, 2 down-regulated, 1 up-regulated at FDR < 5%). These data confirm that, as in humans with HD and other animal models, progressive striatal transcriptional dysregulation is a feature of aging $Htt^{Q111/+}$ mice.

We next examined pathway enrichment of dysregulated transcripts using both hypergeometric²⁵ and gene set enrichment analysis (GSEA)²⁶ techniques. The hypergeometric test examines over-enrichment of differentially expressed genes in a specific pathway, while GSEA quantifies whether a set of pathway genes is over-represented at the top or bottom of an ordered list of genes - here, the full list of 9-month striatal transcripts, ordered by the fold-change in $Htt^{Q111/+}$ mice compared to wild-type mice. Hypergeometric testing of 726 genotype-sensitive (FDR < 5%) striatal transcripts (from a gene universe of 19,031 robustly assayed genes) revealed significant enrichment of striatal genotype-sensitive transcripts in 8 pathways with an adjusted p-value < 0.05, notably in neuronal signaling pathways (Fig. 1b). GSEA revealed 15 gene sets significantly enriched (FDR < 0.1) at the high or low end of the list of striatal transcripts, ordered by their genotype expression ratio ($Htt^{Q111/+}/Htt^{+/+}$; Fig. 1c). Nineteen common pathways had *nominal* enrichment p-values < 0.05 in both GSEA and hypergeometric analyses, including Reactome pathways involved in neurotransmission and cellular signaling (e.g. transmission across chemical synapses, neuronal system, signaling by GPCR, and GPCR downstream signaling; Table S1), suggesting synaptic and signaling transcripts are altered at the transcriptional level in the 9-month-old *Htt*^{Q111/+} striatum.

To orthogonally validate the observed striatal transcriptional alterations and examine their relevance as useful preclinical tools in $Htt^{Q111/+}$, we quantified specific transcript levels in the striatum of 3-, 9- and 12month-old $Htt^{Q111/+}$ mice using quantitative real-time polymerase chain reaction (QRT-PCR). As predicted by cross-sectional RNASeq data (Fig. 1), transcript levels of the critical striatal signaling gene *Ppp1r1b*, that encodes the protein dopamine- and cAMP-regulated phosphoprotein, Mr 32 kDa (DARPP32) are normal in $Htt^{Q111/+}$ mice at 3 months of age, but reduced at both 9 and 12 months of age (effect of genotype $F_{(1,47)} = 6.74$, p = 0.01, Figure 2a). Similarly, we find that levels of a suite of additional neuronal signaling and synaptic genes (including: *Cnr1*, *Drd2*, *Homer1*, *Pde10a*, *Penk*, *Scn4b*) are reduced in aged (9- and 12-month-old) but not young (3-month-old) $Htt^{Q111/+}$ mice (Fig. 2a, effect sizes for each target shown in Fig. 2b). Other transcripts increase over time in $Htt^{Q111/+}$ and include genes involved in DNA damage (e.g. *N4bp2*) or immune response (e.g. IsIr2, H60b, Fig. 2a). These results confirm the findings of our RNAseq study and suggest that levels of a range of transcripts are sensitive genotype markers in the aging $Htt^{Q111/+}$ striatum.

Striatal Histology. We also considered a range of histological endpoints in the dorsolateral striatum of 3-, 9- and 12-month-old Htt^{Q111/+} mice. Compared to tissue-level analyses such as RNASeq and QRT-PCR, immunohistochemistry (IHC) enables the identification of specific cell types and analysis of subcellular localization. First, a trivial explanation for the observed transcriptional alterations in the striatum of the $Htt^{Q111/+}$ mice is that the cellular composition or relative cell sizes within the striatum has changed. To examine this possibility, we counted putative neurons, astrocytes and microglia in the dorsolateral striatum (respectively: NeuN-, glial fibrillary acidic protein- and allograft inflammatory factor 1-immunoreactive cells). Between 3-12 months of age, the neuronal density in the dorsolateral striatum of $Htt^{Q111/+}$ mice does not change (Fig. 3a, $F_{(1,100)} = 0.03$, p = 0.87), nor does the cross-sectional area of NeuN-immunoreactive soma in the dorsolateral striatum at 9 months (NeuN-immunoreactive cell size distributions shown in Fig. 3b, two-sample Kolmogorov-Smirnov (K-S) test D = 0.02, p = 0.9). We next quantified astrocytic and microglial density in the dorsolateral striatum of 12 month old $Htt^{Q111/+}$ and $Htt^{+/+}$ mice, finding that neither is increased in 12 month old $Htt^{Q111/+}$ mice (Fig. 3d). Progressive loss of synaptophysin density has been reported in transgenic HD and other neurodegenerative disease mouse models²⁷. We quantified synaptophysin density in the dorsolateral striatum and find that Htt^{Q111/+} and $Htt^{+/+}$ mice have equivalent levels of synaptophysin staining from 3–12 months of age (Fig. 3c, $F_{(1,93)} = 3.71$, p = 0.057). These results suggest that the robust transcriptional alterations we observe in the aging $Htt^{Q111/+}$ striatum (Fig. 1) are not due to altered numbers or size of neurons, loss of synaptic compartments and that overt gliosis is not a component of the phase of HD modeled by the *Htt*^{Q111/+} mice.

We also examined the appearance of aggregated striatal huntingtin using the MW8 antibody, which detects aggregated, but not diffuse, mutant huntingtin²⁸, but to our knowledge has not yet been applied to study Htt^{Q111/+} mice. We restricted our analyses to neuronal cell bodies, immunoreactive for NeuN, a well-described pan-neuronal marker²⁹. In the dorsolateral striatum of 3-month-old *Htt*^{Q111/+} mice there is virtually no neuronal MW8 immunoreactivity, but by 9 months of age we observe robust accumulation of both small punctate and large inclusion staining in neuronal nuclei (Fig. 4a). Between 9 and 12 months of age, increased levels of neuronal intranuclear inclusions (NII) in the $Htt^{Q111/+}$ striatum are accompanied by a reduction in the total nuclear MW8 immunoreactivity. Total neuronal nuclear MW8 immunoreactivity increases from 3 to 9 months of age, before declining at 12-months of age (Kruskal-Wallis: $H_{(5)} = 80.7$, p < 0.0001, Dunn: 3-month genotype p = 0.15, 9-month genotype p < 0.0001, 12-month genotype p < 0.0001; Fig. 4b). This rise and fall in total neuronal nuclear MW8 immunoreactivity is accompanied by a progressive increase in the percentage of cells with large NIIs, from 0% at 3-months to 13% at 9-months and 28% by 12 months of age (Kruskal-Wallis: $H_{(5)} = 87.9$, p < 0.0001, Dunn: 3-month genotype p = 1, 9-month genotype p < 0.0001, 12-month genotype p < 0.0001, Fig. 4c). In parallel, the average size of striatal NIIs in $Htt^{Q111/+}$ mice increases more than 40% from $1.3 \pm 0.7 \mu m$ at 9 months to $1.8 \pm 0.9 \,\mu\text{m}$ at 12 months of age (two-sample K-S test, D = 0.29, p < 0.0001). Impaired autophagy has been proposed to contribute to impaired proteostasis in HD^{29,30}, and in the brain mHTT interacts directly with p62/ Sqstm1, an autophagic receptor protein important for selective macroautophagy³¹. We observed complete co-localization between p62 and MW8 immunoreactivity in striatal NII's in *Htt*^{Q111/+} mice (Fig. 4d) suggesting that, as has been observed in cell culture³¹, p62 is found in NIIs. We finally considered whether quantitative IHC for specific MSN targets is superior to QRT-PCR quantification. As a proof of concept, we quantified neuronal DARPP32 levels in corticostriatal sections. Consistent with mRNA reductions (Fig. 2), we observe reduced neuronal somatic DARPP32 levels in the $Htt^{Q111/+}$ striatum (Fig. 5; Kruskal-Wallis: $H_{(5)} = 19.6$, p = 0.0015, Dunn: 3-month genotype p = 0.12, 9-month genotype p = 0.09, 12-month genotype p = 0.006). We find that, for DARPP32, quantification of IHC data is more variable than QRT-PCR when establishing reductions in the Htt^{Q111/+} striatum.

Behavioral Analyses. We cross-sectionally analyzed behavior in several cohorts of 3–9 month old $Htt^{Q111/+}$ mice, including assays of motivation, anhedonia, depression, and anxiety (cohorts described in methods). We first measured reward seeking behavior in $Htt^{Q111/+}$ mice using an operant fixed ratio 1 (FR1) behavioral assay to examine how many sweet rewards mice will perform for in a 5 minute block at 6 months of age. We found that $Htt^{Q111/+}$ mice demonstrate reduced reward attainment relative to $Htt^{+/+}$ mice (Fig. 6a; Genotype: $F_{(1,14)} = 17.0$, p = 0.001: Genotype x Reward Size: $F_{(2,28)} = 8.4$, p = 0.001). In an attempt to discern hedonic from motivational explanations for this reduced reward attainment, we examined preference for either 2% or 4% sucrose in drinking water in a separate cohort of mice. We found no difference between 9 month old $Htt^{+/+}$ and $Htt^{Q111/+}$ mice (2%



Figure 2. Confirmation of progressive transcriptional dysregulation in the striatum of aging $Htt^{Q111/+}$ mice using quantitative real-time polymerase chain reaction. (a) Guided by transcriptional discovery with RNAseq, we quantified a number of transcripts and found that many synaptic and neuronal signalling transcripts were down as mice aged from 3-months to 9- and 12- months. As anticipated, we also found upregulated transcripts related to immune and DNA damage pathways (N = 60, subset of 5 mice per row from Table 1). Data are presented as boxplots. (b) Corresponding longitudinal effect sizes for each transcript (highlighted in colored bands above) are displayed in the same color below, with whiskers representing the 95% confidence interval range. Along the x-axis, transcripts are ordered by increasing effect size at 12-months, though robust effects are seen at both 9- and 12-months. Our results demonstrate that these transcripts are sensitive genotype markers that make suitable targets for assessing rescue in interventional trials using the $Htt^{Q111/+}$ mouse model.



Figure 3. No evidence of changes in striatal cell distribution, neuronal cell size or a synaptic marker in $Htt^{QIII/+}$ mice through 12 months of age. (a) NeuN+ cell density is not changed between $Htt^{QIII/+}$ and $Htt^{+/+}$ mice at 3, 9, or 12 months of age (three image frames analyzed per mouse). ANOVA: Genotype $F_{(1, 100)} = 0.03$, p = 0.87, Age $F_{(2, 100)} = 98.9$, p < 0.0001, Interaction $F_{(2, 100)} = 2.79$, p = 0.07. Data presented as boxplots. (b) Distribution of striatal NeuN+ cell sizes does not differ between $Htt^{QIII/+}$ and $Htt^{+/+}$ striata at 9 months of age. Two-sample K-S test D = 0.02, p = 0.98, N = 2038 cells, one image frame analyzed per mouse. (c) Synaptophysin staining intensity was examined at 3, 9, and 12 months of age, showing no differences between $Htt^{QIII/+}$ and $Htt^{+/+}$ mice (one image frame analyzed per mouse). ANOVA: Genotype $F_{(1, 93)} = 3.7$, p = 0.057, Age $F_{(2, 93)} = 16.7$, p < 0.0001, Interaction $F_{(2, 93)} = 1.5$, p = 0.22. Data presented as boxplots. (d) Glia counts at 12 months show no difference in numbers of GFAP+ astrocytes (green; t(32) = 1.2, p = 0.25.) and Aif1+ microglia (red; t(31) = 0.2, p = 0.9) in $Htt^{QIII/+}$ versus $Htt^{+/+}$ striata. Representative $Htt^{+/+}$ section shown. Data presented as boxplots.

sucrose Genotype x Sucrose concentration: $F_{(1,38)} = 0.03$, p = 0.87; 4% sucrose Genotype x Sucrose concentration: $F_{(1,35)} = 0.41$, p = 0.53, Figure S1). However, we did find $Htt^{Q111/+}$ mice demonstrate a stronger preference for sucrose pellets (Genotype: $F_{(1,14)} = 15.9$, p = 0.001, Fig. 6b). Taken together, these tests demonstrate no inherent lack of sucrose preference in $Htt^{Q111/+}$ mice that is present in $Htt^{+/+}$ mice.

To explore the potential contributions of anxiety or depression to the observed reward attainment changes we conducted several tests of these phenotypes in a separate cohort of 9-month old $Htt^{+/+}$ and $Htt^{Q111/+}$ mice. First we conducted the Porsolt swim test, a measure of behavioral despair³⁰, where we observe no differences between 9-month old $Htt^{+/+}$ and $Htt^{Q111/+}$ mice in the duration of time spent inactive during the task (Fig. 6c; t(38) = 0.8, p = 0.9). Similarly, using the elevated plus maze task as a measure of anxiety³², we found that 9-month-old $Htt^{+/+}$ and $Htt^{Q111/+}$ mice do not differ in the amount of time spent in the open arms (Fig. 6d; t(37) = 0.7, p = 0.5), or in the percentage of total arm entries into the open arms, (t(37) = -1.3, p = 0.2). As a final measure of anxiety levels in $Htt^{Q111/+}$ mice, we also employed the light/dark exploration task ^{32,33}, finding that $Htt^{+/+}$ and $Htt^{Q111/+}$ mice spend the same amount of time in the light and dark halves of the apparatus (Fig. 6e; t(38) = 0.04, p = 0.9) and enter the light compartment of the maze a similar number of times (t(38) = -0.6, p = 0.5). Altogether, these data suggest that up to 9-months of age $Htt^{Q111/+}$ mice do not display anxiety-related or depressive-like symptoms, and declines observed in reward seeking behavior may reflect presently undefined cognitive or motivational issues worthy of additional study.

Power Analysis for preclinical studies in Htt^{Q111/+} **mice.** To establish the utility of the natural history data collected here (QRT-PCR and IHC), we conducted several power analysis studies to understand whether these results would be useful in a preclinical trial setting. We considered a hypothetical 2×2 factorial experiment, with two genotypes ($Htt^{+/+}$ vs. $Htt^{Q111/+}$) and two treatment groups (baseline vs. a hypothetical treatment). We simulated data for $Htt^{+/}$ and $Htt^{Q111/+}$ mice in the 'treatment' condition by drawing values from a random normal distribution with the same mean, variance, and covariance as our real natural history data. Assuming a 50% rescue from a hypothetical intervention, we established the power of single molecular endpoints in a preclinical



Figure 4. Progressive accumulation of p62 - and huntingtin-immunoreactive neuronal intranuclear inclusions (NIIs) in the striatum of aging $Htt^{Q111/+}$ mice. (a) Images of the dorsolateral striatum taken in 5- μ m sections of 12-month mice triple labelled for neurons (Red; NeuN), nuclei (Blue; DAPI) and aggregated huntingtin (Green; MW8). Accumulation of both large neuronal-nuclear HTT aggregates, as well as small huntingtin nuclear speckles is present in $Htt^{Q111/+}$, but absent in $Htt^{+/+}$ mice. (b) Total neuronal nuclear MW8 immunoreactivity increases from 3 to 9 months of age, before declining at 12 months of age (Kruskal-Wallis: $H_{(5)} = 80.7$, p < 0.0001, Dunn: 3-month genotype p = 0.15, 9-month genotype p < 0.0001, 12-month genotype p < 0.0001; N = 108, breakdown in Table 1. (c) Using a neuronal (NeuN) mask with particle inclusion from 0.5-5 μ m² shows a median of 10% of neurons contain large nuclear aggregates by 9-months, which increases to 27% at 12 months of age. The presence of NIIs provides a robust measure for huntingtin accumulation with an extremely robust effect size for genotype comparisons at 9- and 12-months (d = 1.6 and 6.7, respectively; error bars = 95% confidence interval). Kruskal-Wallis: $H_{(5)} = 87.9$, p < 0.0001, Dunn: 3-month genotype p = 1, 9-month genotype p < 0.0001; N = 108, breakdown Table 1. (d) Images colubelled for aggregated huntingtin (Green; MW8), and autophagy adaptor protein p62 (Red), demonstrate p62 is colocalized with aggregated huntingtin in 12-month $Htt^{Q111/+}$ mice.

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study with 10 animals per arm (4 arms - $Htt^{+/+}$ and $Htt^{Q111/+}$ mice in a treatment or control arm). By 12 months of age, several individual endpoints provide reasonable power to detect this 50% rescue, including reductions in striatal *Scn4b* mRNA levels and an increase in MW8-immunoreactive aggregate counts (Fig. 7a). We next considered whether combining information from multiple molecular endpoints would improve the power to detect a partial rescue. We trained an elastic net logistic regression model to classify $Htt^{Q111/+}$ versus $Htt^{+/+}$ mice using a weighted combination of the QRT-PCR (*Scn4b, Drd1a, Cnr1, Darpp32, Homer1*) and IHC (mHTT aggregates, mHTT area, DARPP32, cell counts, neuronal ratio) endpoints in the striatum. This model distinguished $Htt^{Q111/+}$ vs. $Htt^{+/+}$ mice with > 90% accuracy in training data from 9- and 12-month-old mice. The model assigns non-zero weights to 0 endpoints in 3-month-old mice, 9 in 9-month-old mice, and 9 in 12-month-old mice (regression coefficients for individual endpoints provided in Fig. 7d). With a sample size of 10, this multivariate model would have 80% power to detect a 35% rescue in 12-month-old mice or a 60% rescue in 9-month-old mice (Fig. 7b). With n = 20, the model had 80% power to detect a 25% rescue in 12-month-old mice or a 40% rescue in 9-month-old mice (Fig. 7c). These results suggest that combining information from multiple molecular endpoints can improve power to detect subtle effects of treatments.



Figure 5. Histological analysis confirms protein-level reductions in DARPP32 in the aging $Htt^{Q111/+}$ striatum. (a) DARPP32 staining with NeuN positive cell traces (Yellow) to demonstrate the inclusion areas for DARPP32 quantification (Top). Representative images of DARPP32, NeuN, and DAPI triple-labelled dorsolateral striatum highlights interneurons (Red), medium spiny neurons (MSN; Yellow), and non-neuronal cells (Blue). B. DARPP32 immunofluorescence per neuronal cell body is reduced in $Htt^{Q111/+}$ mice by 12-months (Fig. 5; Kruskal-Wallis: $H_{(5)} = 19.6$, p = 0.0015, Dunn: 3-month genotype p = 0.12, 9-month genotype p = 0.09, 12-month genotype p = 0.006, N = 108, breakdown in Table 1). Data are presented as boxplots, effect size error bars = 95% confidence interval.

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Discussion

We have here characterized the health and progression of phenotypes in the $Htt^{Q111/+}$ mouse, which accurately model the zygosity and Htt expression level of human HD patients. We found that, in contrast to transgenic N-terminal HTT fragment models of the disease, which have rapidly progressing and terminal disease, the $Htt^{Q111/+}$ mice were grossly healthy through 12 months of age in terms of body weight, plasma chemistry and both central and peripheral inflammation, extending previous observations of these mice¹⁰. Despite their generally healthy state in their first year of age, during this interval they presented striatal-specific, progressive, molecular changes consistent with those observed in other animal models of HD and indeed human mutation carriers, as well as specific behavioral phenotypes consistent with early HD. Using simulation studies, we demonstrated that these animals provide a powerful tool for experimental therapeutics targeting these early molecular changes.

In a recent review, Menalled and Brunner¹¹ demonstrated that less than 5% of preclinical studies of reviewed therapeutic agents in HD were tested in knock-in models of the disease. The R6 and N171 lines of mice, which transgenically express short fragments of mHtt, were much more commonly used, together comprising 72% of all preclinical studies examined. Translation from animal models of neurodegeneration to human clinical trials has been disappointing, with a large number of agents predicted to be useful in transgenic animals failing to provide clinical benefit in human HD, for example¹¹. A major benefit of screening therapeutics in short fragment mouse models of HD is that they present robust behavioral phenotypes, including motor alterations, affective impairments and cognitive alterations⁴. Similar, though less pronounced, phenotypes have been observed in full-length YAC128 and BACHD mice⁹. Distinguishing central effects from peripheral ones is difficult in these animals because they either progressively lose (e.g. R6/2³⁴ and N171^{34,35}) or gain (YAC128³⁶ and BACHD⁹) significant amounts of body weight in parallel with the development of progressive molecular and behavioral phenotypes. Large observational studies have suggested that HD patients may, on average, lose weight^{9,37}, as observed in short fragment mouse models of HD. However, more recently, careful and well-powered studies of presymptomatic HD mutation carriers and early stage patients reveals they have normal body composition and plasma chemistry³⁸, consistent with our observations in aging *Htt*^{Q111/+} mice (Table 1).

While presymptomatic HD mutation carriers do not show clinically obvious motor impairments, by definition, they do show clear sub-clinical cognitive impairments³⁹ and progressive affective alterations. Amongst affective changes in HD mutation carriers, apathy has emerged as the most widely observed⁴⁰, and uniquely amongst neuropsychiatric symptoms, progresses in a continuous fashion with disease state^{40,41}. Like presymptomatic HD mutation carriers, previous studies of $Htt^{Q111/+}$ mice revealed normal gross motor behavior and grip strength through 12 months of age^{9,10,42}. Recent work with q175 HD knock-in mice suggests they present subtle motivational alterations detected using mixed fixed-/progressive-ratio operant tasks⁴³. Similarly, reduced executive function has been observed in $Htt^{Q111/+}$ mice using delayed non-match to sample tasks⁴². The data presented here (Fig. 6) suggest that 9-month old $Htt^{Q111/+}$ mice do not have enhanced anxiety or depression-like behaviors, but do show motivational deficits, most notably a reduced performance on a fixed-ratio 1 task, despite normal hedonic drive for sweet stimuli at this time point. These behavioral studies, in conjunction with the body weight and



Figure 6. Presence of Htt^{QIII} allele results in reduced reward seeking behavior independent of anxiety-like or depressive behavior. (a) $Htt^{QIII/+}$ mice display reduced reward attainment compared to $Htt^{+/+}$ mice across three reinforcement durations (Rft = 40 ms, 60 ms, & 100 ms with 100 ms being 5 µL) during FR1 operant testing (Genotype: $F_{(1,14)} = 17.0$, p = 0.001, Genotype x Reward: $F_{(2,28)} = 8.4$, p = 0.001). (b) $Htt^{QIII/+}$ mice exhibit a stronger preference for sucrose pellets during the hours sucrose consumption task compared to $Htt^{+/+}$ mice (Genotype: $F_{(1,14)} = 15.9$, p = 0.001). (c) $Htt^{QIII/+}$ and $Htt^{+/+}$ mice do not differ in the amount of time spent inactive during the Porsolt swim task. (d) $Htt^{QIII/+}$ and $Htt^{+/+}$ mice do not differ in the amount of time in the open arms of the elevated plus maze. (e) $Htt^{QIII/+}$ and $Htt^{+/+}$ mice do not differ in the amount of time spent in the light compartment of the apparatus during the light/dark exploration task (a) and (b) are presented as mean +/- SEM. (c), (d) and (e) are presented as boxplots.

general health data presented here, suggest that $Htt^{Q111/+}$ mice more closely resemble pre-symptomatic mutation carriers than do short fragment models of HD. Importantly, this suggests interventions targeting apathy, the single most prevalent and progressive psychiatric manifestation of HD^{41,40}, can be tested in knock-in models of HD.

Transcriptional dysregulation in HD has long been noted in both human samples and samples taken from experimental models. Here, using mRNA sequencing techniques we provide quantitative evidence about the degree to which striatal transcriptional changes exceed transcriptional changes in a relatively spared tissue in HD, the cerebellum. Indeed, at 9 months of age, the striatum has more than 250-fold more differentially expressed transcripts than does the cerebellum at this age (Fig. 1A). This is consistent with the fact that the cerebellum is grossly spared from pathological volume change in both human patients²⁴ and full-length mouse models of HD¹⁴. A concern with transcriptional changes in pathologically affected tissue is that tissue architecture changes (in either cellular composition or relative cell size changes) may explain alterations in mRNA abundance in tissue level analyses. The evidence presented here suggests that robust transcriptional dysregulation in the striatum of $Htt^{Q111/+}$ mice precedes any alterations in the density of striatal neurons, astrocytes, microglia or synapses. In fact, of the presented phenotypes, altered transcript levels are amongst the most powerful features for distinguishing $Htt^{+/+}$ from $Htt^{Q111/+}$ mice at 9–12 months of age (Fig. 7). These data confirm that striatal transcriptional dysregulation occurs in the absence of changes in cell number in early HD, occurring the absence of neurodegeneration, gliosis or overt loss of synapses. Also consistent with previous observations of $Htt^{Q111/+}$ mice at younger ages¹⁷, we





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observe progressive striatal-specific accumulation of neuronal intranuclear inclusions (NIIs) in $Htt^{Q111/+}$ mice. Similar to aggregates found in human HD, these large inclusions are primarily restricted to neurons (we do not observe any glial NIIs), and co-stain with autophagic cargo marker p62 (Fig. 4). Our power analyses suggest that treatments that alter either mHTT levels, or improve proteostasis and thereby alter aggregate formation, can be robustly tested *in vivo* using the $Htt^{Q111/+}$ mice with adequate power.

While we find that the *Htt*^{Q111/+} mice provide a number of useful reflections of human HD, there are also significant limitations to their truly modeling Huntington's disease. First, even healthy captive mice live only approximately 2 years⁴⁴, while human HD generally has onset only after 30–50 years of relatively unaffected life⁴⁵. This discrepancy in lifespan may explain why very long CAG repeats (here, approximately 113 - Table 1) are required to lead to robust molecular or behavioral changes. Repeats of this length would be predicted to cause juvenile onset HD in the human context, with fulminant behavioral and neuropathological consequences that are not observed in any mouse models described⁴⁶. Alternatively, or in addition, the different neural circuity⁴⁷, transcriptional and proteomics networks^{6,48} of the human brain could explain the selective vulnerability of human neurons to mutant-huntingtin induced cell dysfunction and death.

In summary, we present data suggesting that the *Htt*^{Q111/+} mice are grossly healthy during their first year of life, yet present a range of molecular and behavioral alterations consistent with presymptomatic HD mutation carriers. Our power analyses revealed that these changes can provide sufficiently powered endpoints for preclinical studies of neuroprotective therapies for HD with a reasonable duration and number of subjects. In conjunction with other recently described cognitive and affective^{10,42,43} behavioral phenotypes in these animals, we propose that knock-in models provide greater face validity, and thereby potential translatability, than transgenic models of HD for future preclinical studies.

Methods

Mice. We used 4 cohorts (detailed below) of mixed-gender B6.*Htt*^{Q111/+} mice for the described studies (Research Resource Identifier:IMSR_JAX:003456). The creation of the *Htt*^{Q111/+} line has been described⁵. Experiments for cohort 1 were conducted in accordance to NIH Guide for the Care and Use of Laboratory Animals and was approved by the Massachusetts General Hospital (MGH) Subcommittee of Research Animal Care (SRAC). Experiments for cohort 2 and 3 were conducted in accordance to NIH Guide for the Care and Use of Laboratory Animals and approved by the Western Washington University (WWU) animal care and use committee. Experiments for cohort 4 were conducted in accordance to the UK Animals (Scientific) Act 1986, European Union Directive 2010/63/EU and approved by the Cardiff University local ethical review committee.

Cohort 1 (RNA sequencing). The cohort used to generate the RNASeq data presented in Fig. 1 was generated at Massachusetts General Hospital (MGH). Cohorts of 58 total mixed-gender littermate mice (12×3 -month and 16×9 -month $Htt^{Q111/+}$; 12×3 -month and 16×9 -month $Htt^{+/+}$) mice were generated by crossing B6. $Htt^{Q111/+}$ males with C57Bl/6J females. The size of the Htt CAG tract ranged from 126 to 135 (mean 131), with no significant differences observed across the groups of this cohort. The mice were part of a larger study of the impact of dietary fat on metabolism⁴⁹ - the mice described in Fig. 1 were fed chow from weaning until 9 months of age with either 60% kcal/fat (Open Source Diet D12492) or 45% kcal/fat (Open Source Diet D1245). Striatal and cerebellar transcriptional effects of these diets were collapsed for the analysis presented in Fig. 1.

Cohort 2 (Molecular natural history). The cohort used to generate the molecular natural history data, presented in Figs 2–5 and 7, was bred and maintained at the Jackson Labs (Bar Harbor, Maine) and shipped to the WWU vivarium approximately 2 weeks before the endpoints under investigation. Three groups of mice were sacrificed at 3 (90 \pm 4 days), 9 (295 \pm 2 days) or 12 (369 \pm 5 days) months of age (Table 1).

Cohort 3 (Behavioral cohort, WWU). The cohort used to collect the behavior data at WWU (Porsolt swim test, elevated plus maze, light/dark box, sucrose solution preference) were bred and maintained at Jackson Labs (Bar Harbor, Maine) and consisted 40 female mice ($20 Htt^{+/+}$; $20 Htt^{Q111/+}$). Htt CAG tract ranged from 107 to 119 (mean 114). Mice were shipped to the WWU vivarium at 8 months of age and acclimated to reversed light cycle conditions (lights on from 12 am to 12 pm) for four weeks prior to testing at 9 months of age. Behavioral assays were conducted between 1:00 pm and 5:00 pm during the active phase.

Cohort 4: (Behavioral cohort, Cardiff). The Cardiff cohort consisted of 16 mice (8 $Htt^{+/+}$; 8 $Htt^{Q111/+}$), bred in-house from founders originally obtained from Jackson Labs (Bar Harbor, Maine), was used for the FR1 and sucrose pellet preference experiments. The mice were housed under 12-hour light/dark cycle with free access to food and water outside of experimental periods. FR1 operant testing began when the mice were 4 months of age, followed by the sucrose pellet preference experiment at 6 months of age. All experimental procedures were initiated between 8 am and 9 am.

Tissue isolation and sample processing. After a three hour fast, mice in the molecular natural history cohort were euthanized via IP sodium pentobarbital injection (Fatal Plus, Henry Schein). Whole blood was collected by cardiac puncture with EDTA $(25 \,\mu\text{M})$ and plasma was extracted. Mice were then transcardially perfused with phosphate buffered saline (PBS) to clear tissues of blood. Whole brains were extracted and placed in a Brain Slicer Matrix (Zivic). A midline longitudinal cut was made to separate hemispheres. Coronal cuts were made in the left hemisphere 1- and 4-mm posterior to the junction between the olfactory bulb and cortex, resulting in a 3-mm thick corticostriatal block of tissue (Figure S2) that was formalin fixed (6–8 hours), paraffin embedded, sectioned (5- μ m), and mounted onto glass slides for immunohistochemistry. The contralateral hemisphere was dissected into striatum, cortex, and cerebellum, and flash frozen for molecular analyses.

Plasma chemistry and immune profiling. Plasma clinical chemistry was measured using an AU2700 Chemistry Analyzer (Beckman Coulter) at Phoenix Central Laboratories (Mukilteo, WA, USA). A panel of 37 cytokines, chemokines and acute phase reactants was assessed using a multiplexed cytometric bead array immunoassay (Mouse InflammationMAP 1.0, Myriad RBM, Inc., Austin, TX, USA).

Library construction, RNA Sequencing and RNASeq analysis. For the RNA Sequencing studies in Fig. 1, RNA was extracted using the RNeasy Lipid Tissue Mini Kit (Qiagen). Quality control for RNA was conducted using the Agilent 2100 Bioanalyzer with Agilent RNA 6000 Nano kit. Libraries for sequencing were constructed using the Illumina TruSeq RNA Sample Prep Kit and sequenced on a HiSeq 2000 (2×50 bp) to a read depth of $2.4 \times 10^7 \pm 6.2 \times 10^6$ per sample. The fastq files were aligned using the default parameters of SNAPR⁵⁰ (https://github.com/PriceLab/snapr) against the GRCh38 genome assembly, along with the transcriptome assembly gtf file from Ensembl, GRCh38.75. SNAPR generates read counts for both genes and transcripts simultaneously with alignment. All alignments were performed on Amazon EC2 c3.8xlarge instance using a Ubuntu14.04 base AMI. Differential gene expression was conducted using the edgeR package⁵¹, and pathway enrichment using HTSanalyzeR⁵² in Bioconductor⁵³.

mRNA quantification – *QRT-PCR.* Total RNA was extracted from the right hemi-striatum using the RNeasy Lipid Tissue Mini Kit (Qiagen). Tissue was homogenized in 500 μ L of Qiazol Lysis Reagent for one minute at 4,000 rpm using a homogenizer (BeadBug; Benchmark Scientific). Aqueous and organic phases were separated by the addition of 100 μ L of chloroform. All subsequent steps in were performed according to the manufacturer's protocol. Reverse transcription was performed using the Superscript III First Strand Synthesis System (Life Technologies) following the manufacturer's protocol. Quantitative real-time polymerase chain reaction (QRT-PCR) was conducted with the resulting cDNA using the Real-Time StepOne System (Applied Biosystems). Transcripts were analyzed by the relative quantitation standard curve method and normalized to a transcript unregulated by mutant Htt expression (*Actb*). All taqman probes were purchased from Life Technologies and were as follows: Drd1a: Mm02620146_s1, Drd2: Mm00438545_m1, Cnr1: Mm01212171_s1, Darpp32: Mm00454892_m1, Penk: Mm01212875_m1, N4bp2: Mm01208882_m1, Islr2: Mm00623260_s1, Pde10a: Mm00449329_m1, H60b: Mm04243254_m1, and Scn4b: Mm01175562_m1.

Immunohistochemistry. Deparaffinization washes were as follows: $2 \times 3 \text{ min}$ xylenes, 3 min xylenes:ethanol (1:1), 3 min 95% ethanol, 3 min 80% ethanol, 3 min 70% ethanol, 3 min 50% ethanol, H_2O rinse. Heat-mediated antigen retrieval: 20 min at 95 °C in citrate buffer (pH 6) or Tris-EDTA buffer (pH 9). Sections were blocked in 20% goat serum in PBS for 1 hour, followed by overnight incubation in primary antibody at 4 °C; GFAP-stained slides were blocked with a mouse-on-mouse kit (Vector). Following wash, secondary antibody incubation was 1 hour at room temperature. Slides were mounted and sealed with DAPI fluormount-G (Southern Biotech). AIF1 slides were prepared from free-floating 40-µm coronal sections. Primary antibodies: mouse anti-aggregated HTT (1:750; DSHB; MW8), mouse anti-SQSTM1/p62 (1:300; Abcam; AB56416), rabbit anti-NeuN (1:750; Millipore; ABN78), mouse anti-DARPP32 (1:250; SCBT; H-3; sc-271111), rabbit anti-AIF1 (1:500; Wako), mouse anti-GFAP (1:500; Millipore; MAB3402). Secondary antibodies: Alexa 488 anti-mouse (1:1000; Life Technologies), Alexa 568 anti-rabbit (1:1000; Life Technologies).

Image acquisition and analysis. For all immunolabeled sections, 12-bit images were acquired with an IX-81 laser-scanning confocal microscope with Fluoview 1000 software (Olympus) using a 60x/1.42 NA oil objective, with the exception of AIF/GFAP which used a 40x/1.30 NA oil objective. For each secondary antibody, acquisition parameters were set such that a brain section with no primary antibody emitted no fluorescent signal and settings were maintained for the entire set of sections. Z-stack numbers and thickness were kept consistent for each set of sections. Maximum z-projections were compiled using ImageJ⁵⁴. NeuN and/or DAPI masks were automatically created using ImageJ default thresholds, and used to measure neuronal and/or nuclear expression of MW8, p62, and DARPP32 (Fig. 5B). AIF1 and GFAP were manually counted. Synaptophysin was quantified as percent area of image with synaptophysin immunoreactivity. Experimenters were blind to genotype for antibody application, image acquisition, and analysis.

Behavior. *Elevated Plus Maze.* The plus maze was made of white acrylic with two open arms ($25x5 \times 0.5$; room lux ~475) and two closed arms ($25x5 \times 16$ cm) extending from a central platform elevated 50 cm above the floor. Each session began by placing a mouse in the intersection of the arms with its head directed toward the closed arm located opposite of the experimenter and ended after 10 minutes of free exploration. Activity was video recorded and analyzed using EthoVision XT 8 (Noldus).

Light/Dark Exploration. The light/dark exploration box used in this experiment was constructed of acrylic plastic divided into a clear, brightly illuminated side $(27 \times 27 \times 30 \text{ cm})$; room lux ~475) and a black, fully enclosed side $(18 \times 37 \times 30 \text{ cm})$ separated by a 5.5 × 5.5 cm opening that permitted the animals to move freely between each compartment. Each trial began by placing a mouse in the illuminated section of the apparatus facing the entrance to the dark section, and ended after 10 minutes of free exploration. Activity was video recorded and analyzed using EthoVision XT 8 (Noldus).

Porsolt Swim Test. Mice were placed into a clear 8-quart bucket (28 cm tall, 22 cm in diameter) filled ³/₄ full with room temperature water. Each mouse was placed into the bucket for a single 6-minute session, and swimming activity was recorded using a camera mounted on a tripod oriented toward the side of the apparatus. Inactivity

during the last 4 minutes of each session was assessed by two separate experimenters blind to genotype (interrater reliability = 0.89), and a single composite score was calculated for subsequent analyses.

Sucrose Consumption Test. Mice were given a sucrose eating challenge to determine whether free access to sucrose would produce different levels of consumption between the genotypes. For these sessions the mice were given 95% sucrose pellets (AIN-76A, TestDiet, Richmond, IN) for 10 hours per day for 4 days, and normal lab chow for 14 hours. The amount of sucrose and chow consumed was measured daily by weight of 5 mg sucrose pellets/chow consumed and expressed as g/kg of bodyweight.

Sucrose Preference Test. Two-bottle sucrose preference testing occurred in the WWU vivarium over eight days. On day 1, animals were single housed and provided access to two identical 50 mL dual bearing sipper tubes containing tap water. On day 2, one of the tap water tubes was replaced with a 2% sucrose solution, and mice were provided free access to both sipper tubes for 72 hours. On day 5, the sipper tube containing the 2% sucrose solution was replaced with a 4% sucrose solution, and mice were provided free access for another 72 hours. The location of the sipper tube containing sucrose solution relative to tap water was counterbalanced between experimental animals and rotated each day to avoid place preference. Every 24 hours, experimenters weighed both sipper tubes to determine consumption from each. The amount consumed from each tube was expressed as a percentage total consumption per day; average consumption for each sipper tube during each 72-hour period was compared between genotypes

Operant Testing. Operant testing was conducted in 16 aluminium/steel 9-hole box ($14 \text{ cm} \times 13.5 \text{ cm} \times 13.5 \text{ cm}$) operant chambers (Campden Instruments, UK). On the rear wall of each chamber situated 15 mm from the grid floor, a horizontal 9-hole light array was fixed that had 9 response holes (11 mm diameter and 2 mm apart) that contained lights at the rear and infra-red (IR) sensors at the front, such that nose poke responses to light stimuli could be detected with an IR beam break. For the present experiments only the central response hole (hole 5 of 9) was used with the other holes blocked. On the inner of the front wall a food magazine was placed that allowed the mouse to recover sweet liquid rewards (Yazoo strawberry milk, Campina Ltd, UK), delivered by peristaltic pump after a successful response to the light stimuli. Initial training consisted of non-contingent reward presentations to the magazine that were signalled by the illumination of a light in the magazine. Removal of the head from the magazine was detected by an IR beam across the magazine entrance which then reset the trail. This was followed by training the mice to respond with a nose poke to illuminated hole 5 on the light array. A response in lit hole 5 extinguished the light and illuminated the magazine light to signal reward delivery. On removal of the head from the magazine, the magazine light was extinguished and after a 2s intertrial interval, a new trial began with the illumination of the stimulus light in hole 5 thereby continuing the fixed ratio 1 (FR1) schedule of reinforcement. Once all mice could successfully complete 30 trials in a 30 minute session, the reward seeking probes began. Nine 45 minute FR1 sessions were run using, 3 (x3) counterbalanced reward pump durations (40 ms, 60 ms, 100 ms), to determine sensitivity to different reward magnitudes with the key outcome measure being number of rewards obtained.

Statistical Analysis. Statistics were processed in R 3.2.3⁵⁵. Data were tested for normality (Anderson-Darling test)⁵⁶ and homoscedasticity (Levene's test)⁵⁷. If data met parametric assumptions, we fit linear models analysed by ANOVA. If data violated these parametric assumptions, Kruskal-Wallis test was used as an omnibus test and followed up with Dunn's test to determine whether pairwise differences between genotypes were significant with Bonferroni corrections for multiple post-hoc comparisons. Data presented in Figs 2–6 used boxplots - horizontal lines indicate 25th, 50th and 75th percentile, while the vertical whiskers indicate the range of data. Data falling outside 1.5 times the interquartile range are graphed as isolated points, but were not excluded from statistical analysis⁵⁸. Simulated distributions for power analysis were constructed with the mvrnorm function in the MASS package^{59,60}. The elastic net classifier was constructed using glmnet⁵⁹. Endpoint QRT-PCR and IHC data and code for power analysis are available at https://github.com/seth-ament/hd_endpoints. Graphics were produced using ggplot2⁶¹ and Illustrator (Adobe).

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Author Contributions

R.M.B. and S.R.C. completed a majority of the work (tissue collection, tissue processing, data analysis, manuscript preparation) and should be considered co-first authors. J.P.C. assisted with histological analyses. R.M.W. with M.E.M. prepared the RNAseq tissue. C.F., R.M.W., S.A.A. and N.D.P. designed analyses, aligned and analyzed the RNAseq data. S.P.B. and S.B.D. designed, ran and analysed the operant experiments and sucrose consumption task. D.S., E.W., B.S., L.J., A.G., J.A., M.A. conducted behavioral experiments including elevated plus maze, forced swim test, sucrose preference and light/dark box under the supervision and direction of J.P.C. and S.M., S.A.A. designed and ran the multi-endpoint power analysis with input from N.D.P. and J.B.C., V.C.W assisted with the conception and design of studies and analytic approaches J.B.C. conceived the experiments and wrote the manuscript. All authors have seen and approved the manuscript in its final form.

Additional Information

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