Mutations in the vacuolar protein sorting 35 ortholog (VPS35) gene represent a cause of late-onset, autosomal dominant familial Parkinson's disease (PD). A single missense mutation, D620N, is considered pathogenic based upon its segregation with disease in multiple families with PD. At present, the mechanism(s) by which familial VPS35 mutations precipitate neurodegeneration in PD are poorly understood. Here, we employ a germline D620N VPS35 knockin (KI) mouse model of PD to formally establish the age-related pathogenic effects of the D620N mutation at physiological expression levels. Our data demonstrate that a heterozygous or homozygous D620N mutation is sufficient to reproduce key neuropathological hallmarks of PD as indicated by the progressive degeneration of nigrostriatal pathway dopaminergic neurons and widespread axonal pathology. Unexpectedly, endogenous D620N VPS35 expression induces robust tau-positive somatodendritic pathology throughout the brain as indicated by abnormal hyperphosphorylated and conformation-specific tau, which may represent an important and early feature of mutant VPS35-induced neurodegeneration in PD. In contrast, we find no evidence for α-synuclein–positive neuropathology in aged VPS35 KI mice, a hallmark of Lewy body pathology in PD. D620N VPS35 expression also fails to modify the lethal neurodegenerative phenotype of human A53T-α-synuclein transgenic mice. Finally, by crossing VPS35 KI and null mice, our data demonstrate that a single D620N VPS35 allele is sufficient for survival and early maintenance of dopaminergic neurons, indicating that the D620N VPS35 protein is fully functional. Our data raise the tantalizing possibility of a pathogenic interplay between mutant VPS35 and tau for inducing neurodegeneration in PD.

Parkinson's disease-linked D620N VPS35 knockin mice manifest tau neuropathology and dopaminergic neurodegeneration

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Significance

The neuropathological spectrum of Parkinson's disease (PD) brains harboring VPS35 mutations is not yet clear since only a single mutation carrier has been evaluated at autopsy. D620N VPS35 knockin mice developed here represent one of the first models of inherited PD that develop the robust and progressive degeneration of nigral dopaminergic neurons. Furthermore, these mice develop robust and widespread tau-positive pathology and axonal damage but lack signs of Lewy pathology positive for α-synuclein. VPS35 knockin mice indicate that the D620N mutation is sufficient for the development of tau abnormalities but not α-synuclein. The D620N VPS35 knockin mice provide an important new tool for distinguishing the neuropathological consequences and mechanisms of familial VPS35 mutations, including the intersection with other PD genetic risk factors.
6-phosphate receptor (Cl-MPR), sortilin 1 and SorLA, whereas the glucose transporter (GLUT1), ATG9A, and AMPA receptors (GluR1 and GluR2) are alternatively sorted to the plasma membrane (9, 12). While the D620N mutation in VPS35 does not influence the stability, assembly, or subcellular localization of the retromer, its only known molecular defect is an impaired association with the WASH complex via a diminished interaction with the FAM21 subunit, which is suggested to induce the abnormal sorting of select cargo (i.e., ATG9A, Cl-MPR, and GluR1) in cultured cells (15–20). Whether these putative mechanisms extend to the mammalian brain and are important for neuronal degeneration induced by D620N VPS35 in PD is not yet known.

Dominant missense mutations typically exert their pathogenic actions via a gain-of-function mechanism; however, in some cases mutations can manifest in a loss-of-function manner through dominant-negative or haploinsufficient mechanisms. The precise mechanism of action of familial VPS35 mutations is unclear and animal models can often play an informative role in distinguishing such mechanisms. A number of rodent models have been developed to understand both the normal function of VPS35 in the brain and to formally evaluate the pathogenic effects of PD-linked mutations in relevant neuronal circuits and populations. While transgenic mice expressing human VPS35 variants have not yet been reported, the deletion of endogenous VPS35 in KO mice results in embryonic lethality, suggesting a critical role for VPS35 in development (21–23). The lethality of VPS35 KO mice indicates that the heterogeneous D620N mutation is unlikely to manifest disease via a full loss-of-function mechanism. Recent studies demonstrate that heterozygous KO mice or conditional VPS35D620N-Cre KO mice are viable and exhibit the degeneration of dopaminergic neurons in the substantia nigra, a hallmark pathology of PD, indicating that VPS35 function is critical for normal dopaminergic neuronal health (21, 22). While intriguing, these KO mice fail to model the specific mechanisms by which familial VPS35 mutations induce PD and are likely to develop additional neuropathological phenotypes and susceptibilities unrelated to PD etiology that may complicate the identification of disease mechanisms.

To evaluate the effects of VPS35 mutations, we previously developed a viral-mediated gene transfer model in adult rats to formally demonstrate that the overexpression of human VPS35 harboring a D620N mutation in the nigrostriatal pathway is sufficient to induce dopaminergic neurodegeneration and axonal pathology (18). This transgenic rodent model is consistent with a simple loss-of-function effect for the D620N mutation, and neurodegeneration is most likely due to a gain-of-function or partial dominant-negative mechanism. However, certain caveats of this viral-based model include (i) the restricted expression of VPS35 to specific neuronal populations (that preclude the brain-wide assessment of neuropathology), (ii) its relatively acute time course (i.e., 12 wk), (iii) the requirement for non-physiological transgene overexpression, and (iv) the small difference in neuropathology between WT and D620N VPS35 (18). In the present study, we report the development of D620N VPS35 knockin (KI) mice as a model of familial PD. We evaluate the impact of the D620N mutation at physiological expression levels on the development of PD-like neuropathology with chronic aging and its potential relationship with the PD-linked protein α-synuclein, and we address the mechanism of action of this familial mutation. We provide evidence that endogenous D620N VPS35 expression in KI mice is sufficient to recapitulate the progressive degeneration of nigral dopaminergic neurons, a hallmark of PD, and unexpectedly induces widespread tau neuropathology and axonal damage. D620N VPS35 KI mice provide an important tool for understanding the pathophysiological mechanisms underlying PD.

Results

Generation and Molecular Characterization of D620N VPS35 KI Mice.

We obtained a newly developed conditional D620N VPS35 KI mouse line from The Jackson Laboratory (stock no. 021807) originally developed by the Michael J. Fox Foundation, as recently reported (24). These conditional mice were developed by introducing aloxP-flanked WT minigene (consisting of a splice acceptor, WT VPS35 exons 15–17, and a polyA signal) downstream of intron 14 and replacing endogenous exon 15 (with a D620N mutant version) by homologous recombination in ES cells (Fig. 1A). WT VPS35 is expected to be expressed under normal conditions (via splicing to the minigene), but after Cre-mediated recombination D620N VPS35 is expressed from the endogenous allele (Fig. 1A and B). Upon intercrossing of heterozygous floxed mice (VPS35loxPlox/WT) we were unable to recover homozygotes (VPS35loxPlox/loxPlox) despite the expected frequency of WT and heterozygous mice, suggesting that floxed homozygotes are most likely haploinsufficient due to the D620N mutation.

![Fig. 1. Generation and molecular analyses of D620N VPS35 KI mice. (A) Floxed VPS35 mice were developed by introducing aloxP-flanked WT “minigene” (containing a splice acceptor, WT VPS35 exons 15–17, and a polyA signal) downstream of intron 14 and replacing endogenous exon 15 with a D620N mutant version) by homologous recombination in ES cells. WT VPS35 is expected to be expressed (via splicing to the minigene) but after Cre-mediated recombination D620N VPS35 is expressed from the targeted allele. (B) Germline D620N KI mice were developed by crossing floxed VPS35 mice with CMV-Cre mice to remove the floxed WT minigene. KI mice contain a single loxp site and exon 15 harboring the D620N mutation. (C) PCR genotyping of tail genomic DNA for floxed VPS35 mice (Left) or germline VPS35 KI mice (Right) using PCR primers (arrows) as indicated in A or B. (D) Genomic PCR from liver DNA of germine D620N/WT and WT/WT mice using primers flanking exon 15. PCR products were cloned and sequenced to confirm the presence of the D620N mutation (GAT→AAT) in D620N/WT mice (Lower) compared with WT/WT mice (Upper). (E) Genotype frequencies from crosses of heterozygous (VPS35loxPlox/WT) KI mice. Percentages are from 605 F1 progeny from 70 litters. (F) Kaplan–Meier survivor curves of germline VPS35 KI mice were generated by monitoring cohorts of all genotypes (VPS35loxPlox/loxPlox, n = 139; VPS35loxPlox/loxPlox, n = 302; VPS35loxPlox/loxPlox, n = 125). (G) Western blot analysis of ventral midbrain extracts derived from 3-mo-old D620N VPS35 KI mice using antibodies to endogenous VPS35 (CT, residues 697–797), VP52, VP529, or Ϝ- tubulin. (H) Western blot analysis of hemibrains from floxed VPS35 mice using antibodies to VPS35 (NT; residues 1–311 or CT), VP52, VP529, or Ϝ- tubulin. (I) Northern blot analysis of total brain RNA using a VPS35-specific DNA probe (nucleotides 163–984) confirms normal VPS35 mRNA expression in the D620N/D620N KI mice but reduced VPS35 mRNA in FLOX/FLOX mice relative to their WT/WT littermates.

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likely embryonic-lethal due to compromised VP35 expression from the conditional allele. Notably, homozygous deletion of VP35 in mice is embryonic-lethal (23). As such, germline D620N KI mice were developed by crossing VP35FLOX/WT mice to CMV-Cre transgenic mice to remove the floxed WT minigene (Fig. 1.4 and B).

The resulting heterozygous KI mice (VP35FLOX/D620N/WT) were intercrossed to produce cohorts of VP35FLOX/WT, VP35D620N/D620N, and VP35D620N/WT littermate mice for subsequent age-dependent analysis. KI mice contain a single loxp site and exon 15 harboring the D620N mutation (Fig. 1F), PCR genotyping and genomic sequencing of the targeted exon 15 confirm correct gene targeting and the presence of the D620N mutation in KI mice (Fig. 1 C and D). Notably, heterozygous VP35FLOX/D620N/WT and homozygous VP35D620N/D620N mice are viable, fertile, normal in appearance, produce genotypes at the expected frequency (Fig. 1E) and display normal survival up to 24 mo of age (Fig. 1F). Western blot analysis of ventral midbrain extracts from VP35FLOX/WT, VP35D620N/D620N, and WT mice reveals normal levels of endogenous VP35 as well as the retromer subunits VP26 and VP29 (Fig. 1G), confirming normal expression of VP35 from the mutant allele and no effect of the D620N mutation on retromer stability. Similar experiments on hemibrains from conditional VP35FLOX/WT mice indicate a ~50% reduction of VP35 protein levels, as well as a corresponding reduction of VP26 and VP29, demonstrating that the floxed WT minigene compromises VP35 expression from this conditional allele (Fig. 1H). Normal VP35 mRNA expression levels and splicing in brains of VP35D620N/WT mice do not demonstrate any altered biology by Northern blot analysis of total RNA derived from brain (Fig. 1I). In contrast, we observe ~50% reduced VP35 mRNA levels in the brains of VP35FLOX/WT mice (Fig. I) consistent with Western blot data (Fig. 1H). Therefore, the conditional VP35FLOX/WT mice are equivalent to heterozygous VP35 null mice due to disrupted expression of VP35 from the conditional allele. Collectively, the D620N mutation in VP35 is well-tolerated in mice and does not influence retromer stability or survival, suggesting that it is unlikely to function through a loss-of-function mechanism.

Modest Motor Deficits in D620N VP35 KI Mice. To evaluate the potentially deleterious impact of D620N VP35 expression on motor behavior, we conducted open-field, rotarod, and gait analyses on VP35FLOX/WT and VP35D620N/D620N mice compared with WT littermate control mice at different time points as the mice aged. VP35FLOX/WT and VP35D620N/D620N mice do not exhibit altered body weight compared with VP35FLOX/WT control mice (SI Appendix, Fig. S1). In the open-field paradigm, VP35FLOX/WT and VP35D620N/D620N mice exhibit novelty-induced locomotor activity similar to that of WT mice at 3 and 13 mo of age, including measures of distance and speed (SI Appendix, Fig. S1). Rotarod analysis using an accelerating paradigm reveals similar motor balance, coordination, and learning between genotypes at 13 mo of age (SI Appendix, Fig. S1). Gait analysis assessed using a digital Catwalk system in D620N KI mice reveals a nonsignificant trend of gait impairment (i.e., average speed, limb swing speed, and cadence) in VP35D620N/D620N mice at 6.9, and 12 mo (SI Appendix, Fig. S1), whereas base of support (distance between front or hind paws) and paw print position (distance between right or left paws) are normal (SI Appendix, Fig. S1). Taken together, the expression of D620N VP35 has a rather modest impact on motor function at advanced age.

D620N VP35 Expression Induces Progressive Dopaminergic Neurodegeneration. To determine whether the expression of D620N VP35 in mice influences the integrity of the nigrostriatal dopaminergic pathway with age, cohorts of KI mice were aged up to 13 mo. The number of tyrosine hydroxylase (TH)-positive dopaminergic and total Nissl-positive neurons in the substantia nigra pars compacta (SNpc) of VP35D620N/WT, VP35D620N/D620N, and WT littermate mice were assessed using unbiased stereological counting methodology. The morphology of dopaminergic neuronal soma and their neurite projections appear grossly normal in VP35D620N/WT and VP35D620N/D620N mice (Fig. 2A). At ~3 mo of age, VP35D620N/WT mice display a modest nonsignificant reduction of nigral dopaminergic neuronal number compared with their VP35FLOX/WT or WT littermates (Fig. 2B). Remarkably, at 13 mo of age, VP35D620N/D620N mice exhibit a significant loss (38 ± 4.0%) of dopaminergic neurons whereas VP35D620N/WT mice also exhibit significant dopaminergic neuronal loss (31 ± 3.4%) relative to WT mice (Fig. 2C). The loss of TH-positive neurons is due to neuronal degeneration rather than to a loss of TH phenotype, as indicated by a corresponding significant loss of total Nissl-positive SNpc neurons in VP35D620N/WT (31 ± 3.5%) and VP35D620N/WT (22 ± 2.6%) mice compared with WT mice (Fig. 2C). Unexpectedly, the density of TH-positive dopaminergic nerve terminals in the striatum of VP35D620N/WT and VP35D620N/D620N mice at 13 mo is normal compared with their WT littermates (SI Appendix, Fig. S2), suggesting a compensatory resprouting of the remaining dopaminergic axonal processes. To further examine the impact of D620N VP35 expression on nigrostriatal pathway function, we monitored the levels of biogenic amines in the striatum of VP35FLOX/WT and WT mice at 14 to 15 mo of age by HPLC with electrochemical detection. Consistent with the normal density of striatal dopaminergic nerve terminals, D620N VP35 KI mice display normal levels of striatal dopamine and its metabolites [3,4-dihydroxyphenylacetic acid (DOPAC) and homovanillic acid (HVA)] as well as other biogenic amines including serotonin and norepinephrine (Fig. 2D). Dopamine and other biogenic amines are also normal in the prefrontal cortex and olfactory bulb (SI Appendix, Fig. S2), which receive projections from dopaminergic neurons in the ventral tegmental area. Dopamine and serotonin turnover in the striatum, prefrontal cortex, and olfactory bulb are not altered in the KI mice (SI Appendix, Fig. S2). Collectively, heterozygous and homozygous D620N KI mice develop robust and progressive substantia nigra dopaminergic neuronal degeneration with age, thereby recapitulating one of the major pathological hallmarks of PD.
Axonal Damage in D620N VPS35 KI Mice. Since the neuropathological spectrum associated with familial VPS35 mutations in PD is not yet clear, we next evaluated the development of PD-related neuropathology using a number of classical markers of neuronal damage and degeneration. Reactive gliosis consistently accompanies neuronal degeneration in PD and other neurodegenerative diseases. However, evidence for astroglisis (GFAP) or microgliosis (Iba1) is not observed throughout the brains of VPS35D620N/WT and VPS35D620N/D620N mice at 13 mo (SI Appendix, Fig. S3). Axonal pathology was next evaluated using either immunostaining for amyloid precursor protein (APP), a sensitive marker of axonal damage, or staining with Gallyas silver, which specifically labels degenerating neuronal soma and processes. VPS35D620N/WT and VPS35D620N/D620N mice at 13 mo display the accumulation of spherical axonal inclusions containing APP as well as increased cytoplasmic APP within neurons of the striatum and substantia nigra (Fig. 3A). VPS35WT/WT mice do not develop these abnormalities. The accumulation of APP-positive axonal inclusions suggests that the expression of D620N VPS35 may also promote axonal degeneration. Gallyas silver staining of striatum and substantia nigra sections from 13-mo-old VPS35D620N/WT and VPS35D620N/D620N mice reveals neurite degeneration, as indicated by the silver/black staining of neurites (Fig. 3B). Quantification of silver-positive neurites in the striatum confirms a significant increase in neurite degeneration in VPS35D620N/WT and VPS35D620N/D620N mice compared with WT mice (Fig. 3C). No evidence of neurite degeneration is observed in D620N KI mice at 3 mo of age (see Fig. 9E). Neurite degeneration is more prominently detected outside of the nigrostriatal pathway, with extensive silver-positive axonal and neuronal degeneration present in the hippocampal CA1-3 region, cerebellum, and brainstem (but not the cortex) of VPS35D620N/WT and VPS35D620N/D620N mice at 13 mo (SI Appendix, Fig. S4). VPS35WT/WT/mice display minimal silver-positive neurites or neurons (Fig. 3 B and C and SI Appendix, Fig. S4). Collectively, our data reveal marked and widespread axonal pathology in the brains of D620N VPS35 KI mice.

Accumulation of Abnormal Tau Protein in Brains of D620N VPS35 KI Mice. To further determine the pathological consequences of VPS35 mutations in vivo, and given the prominent axonal pathology observed in KI mice (Fig. 3), we next assessed alterations in the microtubule-associated protein tau (MAPT). Tau normally localizes to axonal processes via its interaction with microtubules but can dissociate from axons to form intraneuronal neurofibrillary tangles (NFTs) consisting of paired helical filament tau, the hallmark pathology of neurodegenerative tauopathies such as Alzheimer’s disease (25, 26). Notably, some PD brains harboring LRRK2 mutations as well as a number of LRRK2 rodent models can manifest tau-positive neuropathology (27–29), whereas common variation in the MAPT gene is associated with increased PD risk (30). Given that NFTs contain tau species that are abnormally hyperphosphorylated at multiple sites and adopt abnormal conformations (25, 26, 31), we used antibodies that detect specific disease-associated phosphorylation (AT8, PHF1, or CP13) or conformational (MC1) epitopes of tau. Intriguingly, we find evidence for prominent tau-positive neuropathology in the D620N VPS35 KI mice. Tau abnormalities are prominent by 15 mo of age in VPS35D620N/WT and VPS35D620N/D620N mice compared with WT mice (SI Appendix, Fig. S5). Tau abnormalities (AT8 and MC1) are detected most prominently in the cortex, hippocampus, and cerebellum of aged VPS35D620N/WT and VPS35D620N/D620N mice, followed by marked tauopathic changes in brainstem and ventral midbrain, and only modest tau pathology in the striatum (SI Appendix, Fig. S6). Somatodendritic tau pathology is generally more prominent in VPS35D620N/WT mice relative to VPS35D620N/D620N mice compared with general background staining observed in VPS35WT/WT mice (Fig. 4 and SI Appendix, Figs. S5 and S6).

Fluorescent colocalization studies in the hippocampus further reveal that abnormal tau accumulation (AT8 and MC1) is largely confined to the dendritic arbor of pyramidal neurons (i.e., stratum oriens, radiatum, and molecular layers) and to occasional NeuN-positive neuronal soma in the stratum oriens and pyramidal layers of KI mice at 13 mo (Fig. S4). Quantitation of AT8 and MC1 fluorescence intensity levels in the hippocampal CA1 and CA3 subregions confirms a significant increase in tau pathology in the VPS35WT/WT and VPS35D620N/WT mice compared with WT mice (Fig. SB). In the substantia nigra, the significant accumulation of total (Tau25) and abnormal (AT8 and MC1) tau is observed specifically in TH-positive dopaminergic neurons of D620N KI mice at 13 mo of age (Fig. 6) and is detectable as early as 3 mo of age (SI Appendix, Fig. S5). Throughout the brains of D620N KI mice, we do not generally find evidence for tau-positive NFTs, inclusions, or neuritic pathology containing fibrillar forms of tau (Figs. 4–6 and SI Appendix, Figs. S5 and S6), suggesting instead that abnormal tau most likely accumulates rather than aggregates in somatodendritic compartments most consistent with earlier forms of “pre-tangle” pathology (25, 31).

To further explore the aggregation state of tau in D620N KI brains, we conducted biochemical studies to determine the steady-state levels, solubility, or abnormal phosphorylation of tau. At 13 mo of age, 1% Triton-soluble extracts derived from D620N KI mice fail to reveal consistent alterations in the steady-state levels (Tau5), phosphorylation (AT8), or abnormal conformational (MC1) of tau protein in the ventral midbrain and hippocampus of VPS35D620N/WT and VPS35D620N/D620N mice (SI Appendix, Fig. S4). VPS35WT/WT mice display minimal silver-positive neurites or neurons (Fig. 3 B and C and SI Appendix, Fig. S4). Collectively, our data reveal marked and widespread axonal pathology in the brains of D620N VPS35 KI mice.
Appendix, Fig. S7). A central feature of human tauopathies is the loss of tau solubility and the accumulation of hyperphosphorylated tau that can be isolated from brain tissue with sarkosyl extraction (25). Accordingly, sequential detergent extraction (low-salt-soluble, 1% Triton X-100-soluble, and sarkosyl-insoluble fractions) was conducted on brain regions with prominent tau pathology (hippocampus and cerebellum) from D620N KI mice at 13 mo. In general, D620N VPS35 expression does not obviously influence overall retromer stability. To evaluate the interaction of VPS35 with retromer complex subunits in the brain of D620N KI mice, we conducted coimmunoprecipitation (co-IP) assays with a VPS35-specific antibody using soluble, 1% Triton X-100, and sarkosyl-insoluble fractions. Using similar co-IP assays, we can detect the robust interaction of V5-tagged WT VPS35 with endogenous WASH1 and FAM21 in HEK-293T cells and the marked impairment of both interactions with D620N VPS35 (Fig. 7C), as previously reported (16, 19). Our data suggest that endogenous VPS35 in the mouse brain does not appreciably interact with the WASH complex. Next, we conducted confocal colocalization studies to understand the pathogenic effects of the D620N mutation in dopaminergic neurons of the substantia nigra. Intriguingly, we find a marked significant reduction of VPS35 and WASH1 levels selectively in intact TH-positive dopaminergic neurons of both VPS35D620N/D620N and VPS35D620N/WT mice compared with their WT littermates at 3 mo (Fig. 7D and E). These data suggest a selective retromer deficit (VPS35 and WASH complex) in nigral dopaminergic neurons due to the D620N mutation, whereas retromer levels and assembly are generally normal in the brains of KI mice (Figs. 1G and 7A and B).

Retromer Subunit Assembly and Levels in Brains of D620N VPS35 KI Mice. Western blot analyses of ventral midbrain extracts from VPS35D620N/D620N, VPS35D620N/WT, and WT mice at 3 mo reveal normal levels of the cargo-selective complex subunits VPS35, VPS26, and VPS29 (Fig. 1G). Therefore, the D620N mutation does not obviously influence overall retromer stability. To evaluate the interaction of VPS35 with retromer complex subunits in the brain of D620N KI mice, we conducted coimmunoprecipitation (co-IP) assays with a VPS35-specific antibody using soluble hemibrain extracts from VPS35D620N/D620N and WT mice. WT and D620N VPS35 interact robustly and equivalently with VPS26 in brain (Fig. 7A). However, we are unable to detect an interaction of WT or D620N VPS35 with the WASH complex subunits FAM21 and WASH1 (Fig. 7A). We could further confirm the equivalent interaction with retromer subunits VPS26A, VPS26B, and VPS29 by mass spectrometric analysis of VPS35 IPs from WT or D620N KI brains, but we are unable to detect any WASH complex subunits (Fig. 7B). Using similar co-IP assays, we can detect the robust interaction of V5-tagged WT VPS35 with endogenous WASH1 and FAM21 in HEK-293T cells and the marked impairment of both interactions with D620N VPS35 (Fig. 7C), as previously reported (16, 19). Our data suggest that endogenous VPS35 in the mouse brain does not appreciably interact with the WASH complex. Next, we conducted confocal colocalization studies to understand the pathogenic effects of the D620N mutation in dopaminergic neurons of the substantia nigra. Intriguingly, we find a marked significant reduction of VPS35 and WASH1 levels selectively in intact TH-positive dopaminergic neurons of both VPS35D620N/D620N and VPS35D620N/WT mice compared with their WT littermates at 3 mo (Fig. 7D and E). These data suggest a selective retromer deficit (VPS35 and WASH complex) in nigral dopaminergic neurons due to the D620N mutation, whereas retromer levels and assembly are generally normal in the brains of KI mice (Figs. 1G and 7A and B).
specific to Lewy pathology in PD (36). Biochemical analyses of detergent-soluble (1% Triton X-100) extracts from the ventral midbrain of 13-mo-old D620N KI mice indicate normal steady-state levels and phosphorylation (Ser129) of α-synuclein (SI Appendix, Fig. S9), consistent with immunohistochemical data. Therefore, D620N KI mice lack evidence of α-synuclein accumulation or neuropathology, in contrast to VPS35 KO mice (21, 22).

To further explore a potential pathological interaction of VPS35 with α-synuclein, we crossed the D620N VPS35 KI mice with human A53T-αSyn transgenic mice (line G2-3), a robust and well-characterized model of α-synuclein–induced neurodegeneration (37). VPS35D620N/D620N mice were first crossed with hemizygous A53T-αSyn mice to generate F1 progeny (i.e., VPS35D620N/WT and VPS35D620N/D620N (αSyn+)) that were subsequently intercrossed to generate F2 cohorts for direct comparison (Fig. 8). A53T-αSyn transgenic mice display shortened lifespan largely owing to the robust degeneration of brainstem and spinal cord neurons leading to limb paralysis, autonomic dysfunction, and premature death (37). Accordingly, we assessed the impact of physiological D620N VPS35 expression on the survival of α-Syn mice to generate F1 progeny (i.e., VPS35D620N/WT and VPS35D620N/D620N (αSyn+) that were subsequently intercrossed to generate F2 cohorts for direct comparison (Fig. 8). A53T-αSyn transgenic mice display shortened lifespan largely owing to the robust degeneration of brainstem and spinal cord neurons leading to limb paralysis, autonomic dysfunction, and premature death (37). Accordingly, we assessed the impact of physiological D620N VPS35 expression on the survival of "

Fig. 6. Tau abnormalities in nigral dopaminergic neurons of D620N VPS35 KI mice. Immunofluorescent confocal colocalization analysis of the 5Npc from 13-mo-old VPS35D620N/WT, VPS35D620N/D620N, and control VPS35WT/WT mice coabeled with (A) Tau5 (total tau), (C) AT8, or (E) MC1 antibodies (green) and the dopaminergic neuronal marker TH (red). Merged images indicate the accumulation of Tau5-, AT8-, and MC1-positive immunofluorescent signals within the somatodendritic compartment of nigral dopaminergic neurons of D620N/WT and D620N/D620N mice. (Scale bars: 15 μm.) Bar graphs indicating CTCF values (mean ± SEM, n = 4 animals per group) of (B) Tau5, (D) AT8, or (F) MC1 fluorescence intensity in TH-positive dopaminergic neurons. **p < 0.001 by one-way ANOVA with Bonferroni’s post hoc test compared with WT/WT mice.

Lack of α-Synuclein Pathology or Functional Interaction in D620N VPS35 KI Mice. α-Synuclein plays a central role in the pathogenesis of PD, where it comprises a key component of Lewy pathology (30, 32–34). Prior studies have revealed a functional interaction of VPS35 and α-synuclein, whereby VPS35 depletion in yeast and worm models can exacerbate human α-synuclein–induced toxicity (35), whereas α-synuclein accumulates in the brains of mice with VPS35 depletion (21, 22). As such, we evaluated brains from aged D620N VPS35 KI mice for alterations in α-synuclein levels or aggregation. VPS35D620N/WT and VPS35D620N/D620N mice do not reveal consistent alterations in the levels, distribution, or morphology of phosphorylated Ser129-α-synuclein above background levels throughout the brain (SI Appendix, Fig. S9), a robust marker of pathological α-synuclein.
A53T-α-Syn mice (Fig. 8). A53T-α-Syn mice exhibit premature lethality occurring over a period of 8 to 18 mo of age. However, the survival of VPS35D620N/WT/WT and VPS35D620N/WT/αSynTr mice does not significantly differ from that of VPS35WT/WT/αSynTr mice. VPS35D620N/WT and VPS35D620N/D620N mice exhibit normal survival relative to VPS35WT/WT mice over 24 mo (Fig. 8). Our data indicate that the lethal neurodegenerative phenotype of A53T-α-Syn transgenic mice is not influenced by endogenous D620N VPS35 expression, in contrast to simpler models with VPS35 deletion (35, 38). We next conducted preliminary experiments by unilaterally delivering mouse α-synuclein preformed fibrils (PFFs) to the striatum of 6-mo-old D620N mice; Fig. 9, insets) to explore the impact of the D620N mutation on the development of α-synuclein pathology (SI Appendix, Fig. S10). At 30 d postinjection, we observe pSer129-α-synuclein–positive pathology localized to dopaminergic neurons of the ipsilateral substantia nigra of D620N mice; however, qualitatively similar α-synuclein pathology is also observed in VPS35D620N/WT mice (SI Appendix, Fig. S10). These data suggest that the D620N mutation does not regulate the initial progression and propagation of α-synuclein pathology in the α-Syn-PFF model. Taken together, these data fail to provide robust evidence for a pathological interaction of α-synuclein in the D620N VPS35 KO mouse model.

The D620N Mutation Supports Survival and Is Inconsistent with a Full Loss-of-Function Mechanism. Our data indicate that homozygous D620N VPS35 KI mice display a normal lifespan (Fig. 1F), in contrast to homozygous KO mice (and VPS35D620NFLOX mice; Fig. 1A) which are embryonic-lethal (23), suggesting that the D620N mutation is not likely to act via a loss-of-function mechanism. To further address this mechanism, we crossed VPS35D620N/WT mice with VPS35D620NFLOX mice (containing a single VPS35 null allele) (Fig. 1, A, H, and J) to create compound heterozygous VPS35D620NFLOX progeny. These mice harbor a single D620N allele and a VPS35 null allele and exhibit ~50% VPS35 protein levels in the brain similar to VPS35D620NFLOX mice, as expected (Fig. 9A). Importantly, we find that the VPS35D620N/VPS35D620NFLOX mice are viable, grossly normal, produced at the expected genotypic frequency (Fig. 9B), and exhibit normal survival over 24 mo (Fig. 9C). At ~3 mo, the VPS35D620N/VPS35D620NFLOX mice display a normal number of substantia nigra dopaminergic neurons (Fig. 9D) and lack axonal pathology (Fig. 9E) relative to WT littermate mice. These data are in contrast to conditional KO mice with homozygous VPS35 deletion in DAT-positive neurons that develop robust dopaminergic neuronal loss at 2 to 3 mo of age (22). We further evaluated tau pathology in the VPS35D620N/FLOX mice at 24 mo of age. Both VPS35D620N/WT and VPS35D620N/FLOX mice markedly accumulate abnormal tau (AT8 and M1C) in hippocampal subregions that is completely absent from VPS35D620NFLOX mice (Fig. 9F). The burden of tau pathology in the CA3 subregion is similar between VPS35D620N/WT and VPS35D620N/FLOX mice, whereas in the CA1 subregion of VPS35D620N/FLOX mice the accumulation of abnormal tau shifts from dendrites to the soma of pyramidal neurons (Fig. 9F). As compound heterozygous VPS35D620NFLOX mice alone are sufficient for the development of tau pathology, these data suggest that D620N VPS35 can induce pathology irrespective of the presence of WT VPS35. Furthermore, VPS35 depletion in the VPS35D620N/FLOX mice is not sufficient to induce tau pathology (Fig. 9F and SI Appendix, Fig. S8), implying that the D620N mutation impacts tau protein via a gain-of-function mechanism. Since VPS35 and VPS35D620N/FLOX mice are embryonic-lethal (23), our data collectively demonstrate that a single copy of the D620N allele...
(in compound VPS35<sup>D620N/FLOY</sup> mice) is sufficient for maintaining normal survival and early dopaminergic neuronal health in mice, indicating that D620N VPS35 is fully functional and highly unlikely to manifest disease via a loss-of-function mechanism (i.e., via a haploinsufficient or dominant-negative effect).

**Discussion**

Here, we report the development and phenotypic assessment of a germline D620N VPS35 KI mouse model of PD with advancing age. We demonstrate that the expression of D620N VPS35 at endogenous levels is sufficient to induce the progressive degeneration of nigrostriatal pathway dopaminergic neurons, thereby recapitulating the major neuropathological hallmark of PD. KI mice develop rather modest motor deficits with age consistent with the normal density of striatal dopaminergic nerve terminals and dopamine levels. In addition, D620N VPS35 expression induces widespread axonal damage and tau-positive pathology throughout the brain, including within the substantia nigra. Tau pathology is characterized by the accumulation of abnormal hyperphosphorylated and conformational-specific epitopes within somatodendritic compartments and is most consistent with an early "pre-tangle" form of pathology. Heterozygous VPS35 null mice do not similarly develop tau pathology at advanced age. Surprisingly, we find no evidence for α-synuclein neuropathology in the D620N KI mice, a second pathological hallmark of PD. Furthermore, the D620N mutation is unable to modify the normal survival and dopaminergic neuronal health in VPS35 null mice, indicating that the D620N VPS35 protein is fully functional and is unlikely to act via a loss-of-function mechanism. Taken together, our data demonstrate that the PD-linked dominant D620N mutation in VPS35 is able to induce neuronal degeneration, axonal damage, and tau pathology, most likely via a gain-of-function mechanism. Our findings formally establish an important contribution of familial VPS35 mutations to the development of PD.

D620N VPS35 KI mice are one of the first models of inherited monogenic PD that develop the robust and progressive degeneration of nigral dopaminergic neurons. Importantly, a heterozygous or homozgyous D620N mutation produces similar neurodegenerative phenotypes in the KI mice. For tau pathology, the heterozygous KI mice even exhibit modestly enhanced pathology compared with homozygous mice. These data are consistent with a threshold effect of the D620N mutation for the development of neurodegeneration, rather than a gene dosage effect, which would lend support to a gain-of-function mechanism for this dominantly inherited mutation. This idea is supported by the normal viability and survival of homozygous D620N KI mice [in contrast to embryonic lethality in homozygous KO mice (23)] and by the ability of a single D620N allele to support normal survival in mice. Therefore, the D620N mutation does not compromise the overall function of VPS35, a finding supported by studies in yeast (18), and is not consistent with a simple loss-of-function mechanism. The observation that heterozygosity for the D620N mutation is sufficient for the development of fulminating pathology in the KI mice, whereas the viral-mediated overexpression of human D620N VPS35 can also produce similar phenotypes in rodents (18, 39), would tend to argue against a partial dominant-negative effect of this pathogenic variant.

VPS35 KO (heterozygous or conditional) mice also develop dopaminergic neurodegeneration (21, 22), although it is unclear whether this is a selective effect given that VPS35 is broadly expressed throughout the CNS and is critically required for normal development (23). The molecular mechanism underlying the D620N mutation is not yet clear. Although our data support a gain-of-function effect as the most plausible mechanism for the D620N mutation in vivo, several studies have shown that this mutation impairs the interaction with and endosomal recruitment of the WASH complex by VPS35 in cell culture models (16, 19), a finding replicated in the present study. We demonstrate that the WASH complex is neither a major nor robust interacting protein of VPS35 in the brain, suggesting that it either forms a labile complex with the retromer or that it binds at levels undetectable by our current methods. Despite the lack of WASH complex binding, we do find evidence for a deficiency of VPS35 and WASH complex components in nigral dopaminergic neurons of D620N KI mice. Future studies will be important to determine how the D620N mutation influences VPS35 protein turnover in these neurons and whether or how retromer deficiency is sufficient to induce dopaminergic neurodegeneration.

A recent study using a version of this germline D620N KI mouse reports an increase in evoked striatal dopamine release (24), whereas an independent D620N KI model oppositely exhibits impaired dopamine release in young mice (40). These alterations in dopaminergic neurotransmission in young D620N KI mice may indicate an early dysfunction of dopaminergic neurons before their subsequent degeneration at advanced age. Another study demonstrates the dominance of the D620N mutation in VPS35 KI mice or in patient-derived human neuroblasts and monocytes robustly increases LRRK2-mediated Rab10 phosphorylation (41). These findings suggest that PD-linked VPS35 mutations may exert their pathogenic actions by increasing LRRK2 kinase activity via an unknown mechanism, which would further lend support to a gain-of-function mechanism for the dominant D620N mutation.

The neuropathological spectrum of PD brains harboring VPS35 mutations is not yet clear since only a single mutation carrier has so far been evaluated at autopsy (4, 7). Unfortunately, the postmortem examination of D620N KI mice also fails to reveal similar abnormalities in nigral dopaminergic neurons. The development of tau pathology in KI mice is reminiscent of familial LRRK2 mutations. LRRK2 mutation carriers with PD often develop Lewy pathology but some cases instead exhibit tau pathology in neurons or glial cells (28, 29, 42), whereas numerous mutant LRRK2 rodent models report abnormalities in tau metabolism or phosphorylation as the most consistent pathological feature (27, 43–45). Whether the same is true for VPS35 mutation carriers is not yet clear, but VPS35 KI mice indicate that the D620N mutation is sufficient for the development of tau abnormalities but not α-synuclein. Notably, the absence of similar tau pathology in heterozygous VPS35 null mice with age indicates that the D620N mutation exerts this effect via a gain-of-function mechanism. Additional aging up to 24 mo may be required for the development of α-synuclein pathology in the KI mice, or they could potentially develop soluble oligomeric forms of α-synuclein that may not be detectable using the methods applied here. The observation that D620N KI mice are unable to enhance α-synuclein–induced phenotypes that develop in transgenic mice or preformed fibril-based models would tend to argue against meaningful pathological changes in α-synuclein.

Tau pathology can occur in familial and sporadic PD brains albeit much less prominently than Lewy pathology (28, 29, 46). Importantly, common variation in the MAPT gene is consistently associated with an increased risk of developing PD and most likely arises through increases in tau transcript expression (30). In the D620N KI mice, tau pathology develops before axonal damage. This could potentially be due to early microtubule instability within axons that would initially lead to tau disassembly from microtubules and its redistribution and accumulation.
within somatodendritic compartments of neurons. Inhibition of axonal transport and subsequent axonal damage would likely result from microtubule instability. Our data therefore potentially link abnormal retromer function to microtubule instability but the mechanistic underpinnings remain unclear. These effects could result from the inappropriate recycling of specific axonal cargo by the retromer; however, little is presently known about retromer function and specialized cargo within neurons (47). The recent observation of LRRK2 hyperactivation in D620N VPS35 KI mice could potentially link the microtubule network to tau abnormalities (41), as familial mutations that enhance LRRK2 kinase activity (48, 49) are reported to impair microtubule stability (50, 51), increase tau phosphorylation (52, 53), and promote the neuron-to-neuron transmission of WT tau (54). Alternatively, VPS35 mutations may act upon tau metabolism independent of a primary effect on microtubule stability. For example, the D620N mutation has been suggested to impair the delivery of cathepsin D to the lysosome via the abnormal retromer-dependent endosomal sorting of its receptor, CI-MPR (15, 38), whereas reduced lysosomal cathepsin D is linked to tau-induced neurotoxicity (55). VPS35 mutations could also potentially influence tau-directed kinases, as the hyperphosphorylation of tau is known to reduce its affinity for microtubules and promote their destabilization (26, 31, 56). Future studies should focus on the relationship between VPS35 mutations and tau pathology, including the mechanism for inducing tau abnormalities and whether tau itself is important for dopaminergic neurodegeneration in the D620N VPS35 KI mice.

Collectively, our data demonstrate that the physiological expression of PD-linked D620N VPS35 in mice is sufficient to recreate the progressive degeneration of nigral dopaminergic neurons that is the major pathological hallmark of familial and sporadic PD. VPS35 KI mice surprisingly reveal prominent tau pathology, which may represent an important and early feature of mutant VPS35-induced neuronal degeneration in PD. The D620N VPS35 KI mice provide an important tool for elucidating the complex pathophysiological mechanisms underlying neuritic degeneration in PD, including the interaction with other important PD genetic risk factors (i.e., MAPT/tau and LRRK2).

Materials and Methods

Animals. Mice were maintained in a pathogen-free barrier facility and provided with food and water for ad libitum consumption and exposed to a 12-h light/dark cycle. Animals were treated in strict accordance with the NIH Guide for the Care and Use of Laboratory Animals (57). All animal experiments were approved by the Van Andel Institute Animal Care and Use Committee. Conditional (“floxed”) D620N VPS35 KI mice (strain 021807) and CMV-Cre transgenic mice (strain 006054) were obtained from The Jackson Laboratory. Human A53T α-synuclein transgenic mice (line G2-3) have been described (37). Mice were identified by genomic PCR (24, 37).

RNA Extraction and Northern Blot Analysis. Total RNAs were extracted from hemispheres and subjected to Northern blot analysis using a 32P-labeled DNA probe (nucleotides 163–984 of mVPS35). Signal was detected using a Typhoon phosphor-imaging system (Amersham). Expression Plasmids and Antibodies. Human V5-tagged VPS35 plasmids were obtained from Addgene (no. 21691; ref. 58) and previously described (18). Mouse VPS35 cDNA was obtained from Dr. Macon. Primary antibodies used were VPS35 (ab57632 or ab154647; Abcam), VPS26 (ab23892; Abcam), VPS29 (ab10166; Abcam), VPS18 (Sigma), FAM21a-D (S;13; Santa Cruz), V5 and V8-HRP (Life Technologies), TH (NB300-109; Novus Biologicals or clone TH-2; Sigma), α-synuclein (clone 42; BD Biosciences), p53R129-α-synuclein (clone EP1536Y; Abcam), APP (clone 22C11; Millipore), tau (clone Tau5; Millipore), phospho-tau (clones AT8, PHF1, CP13, and MC1), provided by Peter Davies, Feinstein Institute.

Cell Culture and Transient Transfection. HEK-293T cells were maintained and transfected using X-tremeGENE HP Reagent (Roche) as described (59). For co-IP assays, hemibrains from 3-mo-old mice or transfected HEK-293T cells were prepared in IP buffer (20 mM Hepes-KOH, pH 7.2, 50 mM potassium acetate, 200 mM sorbitol, 2 mM EDTA, 0.1% Triton X-100, and 1 Complete Mini protease inhibitor mixture (Roche)) and subjected to IP with antibodies to VPS35 or V5 (5 µg) as described (59). IPs and input lysates were subjected to Western blotting, detection, and quantitation as described (59). For M5, anti-VPS35 IPs from WT or VPS35 KI mouse brain were subjected to analysis by nanoliquid chromatography/tandem MS (LC-MS/MS) (ThermoFisher Q Exactive) at M5 Bioseworks as previously described (59).

Sequential Fractionation of Brain Tissues. Brain tissues from KI mice were harvested and subjected to sequential fractionation in low-salt, Triton X-100, and sarkosyl buffers as described previously (54). For standard biochemical analysis, brain tissues were homogenized in lysis buffers containing 1% Triton X-100 or 2% SDS (Triton-insoluble), as described (18, 60).

Measurement of Biogenic Amines. Homogenized tissues were separated on a 150–× 4.6-mm Microsorb MV C18 100-5 column (Agnilent Technologies) and analyzed by HPLC-electrochemical detection (ECD) to detect dopamine, HVA, DOPAC, serotonin (5-HT), 5-hydroxyindoleacetic acid (5-HIAA), noradrenaline, and 3-methoxy-4-hydroxyphenylglycol (MHPG), as previously described (61, 62).

Immunohistochemistry and Immunofluorescence. Mice were perfused with 4% paraformaldehyde (pH 7.4), cryoprotected in 30% sucrose, and dissected into 35-µm-thick coronal sections. Immunostaining by incubation with primary and biotinylated secondary antibodies (Vector Labs), ABC reagent (Vector Labs), and DAB (Vector Labs) has been described (18, 60). Images were captured using an Axio Imager M2 (Zeiss) with color CCD camera (AxioCam; Zeiss). Silver staining was conducted on serial sagittal sections with the PD NeuroSilver kit II (PD Neurotechnologies) according to the manufacturer’s instructions. Images were captured using a ScanScope XT slide scanner (Aperio) at a resolution of 0.5 µm per pixel. Images were quantified using a custom CellProfiler v3.1.5 pipeline to isolate black neurites and particles. The percent of total pixels associated with black neurites was calculated for each image, sampled across three to six images per animal. Fluorescence immunostaining of brain sections was conducted using primary and secondary antibodies conjugated to Alexafluor-488 or -594 (Life Technologies) according to the manufacturer’s instructions. Images were captured using a ScanScope XT slide scanner (Aperio) at a resolution of 0.5 µm per pixel. Immunofluorescence and quantitation of SNpc dopaminergic neurons was performed using NIH ImageJ (v1.3). The mean optical density across six to eight sections per animal was determined.

Stereotactic Delivery of Preformed Fibrils. Mouse α-synuclein was expressed in Escherichia coli and purified as previously described (63). To generate fibrils, 5 mg/mL of purified α-synuclein in 1× PBS, pH 7.4, was shaken at 1,000 rpm for 7 d, sonicated with 60 pulses at 10% power (30 s total time, 0.5 s on, 0.5 s off), aliquoted, and stored at −80 °C. Sedimentation assays and thioflavin T assays were performed as described (63, 64) to ensure fibril quality. Male adult littermate mice (n = 3 mice per genotype) were subjected to stereotactic surgery for the unilateral delivery of PFF–dyn into the dorsal striatum. The following coordinates were used relative to bregma: anterior–posterior, +0.2 mm; mediolateral–ventral, +2.0 mm; and dorsal–ventral, −2.6 mm. Mice received PFF–dyn (5 µg) in 1 µL at a flow rate of 0.1 µL/min. Animals were euthanized at 30 d postinjection.

Behavioral Assessments. Behavioral testing was performed on male and female mice at ages indicated for each test, as described (60). Open-field testing using an Ethovision XT video tracking system (Noldus Information Technology), accelerating rotorod (5 to 50 rpm over 5 min), and digital Gait analysis using the automated CatWalk XT system (Noldus) were conducted.

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