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Missense dopamine transporter mutations associate with adult parkinsonism and ADHD

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Parkinsonism and attention deficit hyperactivity disorder (ADHD) are widespread brain disorders that involve disturbances of dopaminergic signaling. The sodium-coupled dopamine transporter (DAT) controls dopamine homeostasis, but its contribution to disease remains poorly understood. Here, we analyzed a cohort of patients with atypical movement disorder and identified 2 DAT coding variants, DAT-Ile312Phe and a presumed de novo mutant DAT-Asp421Asn, in an adult male with early-onset parkinsonism and ADHD. According to DAT single-photon emission computed tomography (DAT-SPECT) scans and a fluoro-deoxy-glucose-PET/MRI scan, the patient suffered from progressive dopaminergic neurodegeneration. In heterologous cells, both DAT variants exhibited markedly reduced dopamine uptake capacity but preserved membrane targeting, consistent with impaired catalytic activity. Computational simulations and uptake experiments suggested that the disrupted function of the DAT-Asp421Asn mutant is the result of compromised sodium binding, in agreement with Asp421 coordinating sodium at the second sodium site. For DAT-Asp421Asn, substrate efflux experiments revealed a constitutive, anomalous efflux of dopamine, and electrophysiological analyses identified a large cation leak that might further perturb dopaminergic neurotransmission. Our results link specific DAT missense mutations to neurodegenerative early-onset parkinsonism. Moreover, the neuropsychiatric comorbidity provides additional support for the idea that DAT missense mutations are an ADHD risk factor and suggests that complex DAT genotype and phenotype correlations contribute to different dopaminergic pathologies.

Introduction

Dopamine is an important neurotransmitter regulating motor activity, cognition, neuroendocrine functions, and reward mechanisms. Moreover, disturbances in dopaminergic signaling are of central importance in several common brain diseases including Parkinsonism, attention deficit hyperactivity disorder (ADHD), autism, schizophrenia, and addiction (1, 2). However, the pathophysiology of these diseases is still poorly understood. Importantly, the spatiotemporal regulation of dopaminergic neurotransmission is critically dependent on the presynaptic dopamine transporter (DAT), which mediates reuptake of released dopamine in the synaptic cleft (3–5). Together with several other neurotransmitter carriers, DAT belongs to the family of neurotransmitter/sodium symporters, also referred to as the solute carrier 6 (SLC6) family of sodium-coupled transporters (3, 5). The significance of DAT in dopamine homeostasis has been supported by genetic deletion of the DAT gene in mice, resulting in an extracellular hyperdopaminergic state concomitantly with a 95% reduction in tissue content of dopamine (6). The DAT-knockout (DAT-KO) mice are characterized by severe behavioral changes including spontaneous hyperactivity, as well as by memory and learning deficits (2, 7). Interestingly, both DAT-KO mice and DAT-knockdown mice with 10% residual DAT expression (8) have been considered animal models of ADHD (9–11). DAT-KO mice might also serve as a model of parkinsonism, although primarily after inhibiting dopamine synthesis with a tyrosine hydroxylase (TH) inhibitor (12, 13). Nevertheless, the significance of altered DAT function in the human diseases remains unclear.

The consequences of DAT deficiency in humans were recently elucidated by the identification of 11 patients, all with homozygous or compound heterozygous missense mutations in the DAT
The conceivable contribution of SLC6A3 coding variants to both movement disorder and psychiatric disorders suggests that the character and severity of functional changes in DAT missense mutations might determine the pathological outcomes. To gain insight into the role of SLC6A3 coding variants in human disease, we sequenced coding exons of SLC6A3 in a cohort of patients with early-onset parkinsonism, dystonia, or related atypical movement disorders. Among these, we identified 1 adult patient who was diagnosed with both early-onset parkinsonism and ADHD and who was compound heterozygous for 2 DAT coding variants. One coding variant was transmitted from his father and changes isoleucine in position 312 to phenylalanine (DAT-I321F), whereas the other variant is the result of a de novo DAT missense mutation, Thr356Met (DAT-T356M), was detected in a patient with autism spectrum disorder (24).

Identification of coding DAT variants in an adult male with parkinsonism and ADHD. DNA from 91 patients with symptoms of early-onset parkinsonism, dystonia, or other unclassified movement disorders were analyzed for mutations in SLC6A3 exons (primer sequences are listed in Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI73778DS1). In 1 patient, we identified 2 unique missense mutations, c.934A>T in exon 7 and c.1261G>A in exon 9, which gave rise to the amino acid substitutions Ile312Phe (DAT-I321F) and Asp421Asn (DAT-D421N), respectively (Figure 1, A and B). The mutations were not registered as polymorphisms in variant databases including the Exome Variant Server (NHGRI GO Exome Sequencing Project [ESP]; http://evs.gs.washington.edu/EVS/; accessed March, 2014) and Ensembl (http://Jun2013.archive.ensembl.org/index.html), which include sequence information for more than 13,000 alleles. Moreover, the mutations were not detected upon analysis of 400 Danish control alleles (data not shown). None of the mutations could be identified in the mother, whereas an unaffected sister carried the DAT-I312F mutation (Figure 1C). Hence, the DAT-I312F allele was presumably transmitted to the siblings from the recently deceased father, while DAT-D421N appears to be a de novo mutation. Sequence alignment revealed both Ile312 and Asp421 as evolutionarily highly conserved (Figure 1D).

To determine the cis-trans allelic position of the mutations, we took advantage of an SNP (c.1215A>G) in exon 9 that cosegregated with the I312F mutation. We designed sequence-specific primers that selectively recognized either the D421N mutation (c.1261G>A), the SNP cosegregating with the I312F mutation (c.1215A>G), or the corresponding WT DNA sequences (Supplemental Figure 1). PCR amplification of genomic DNA from the proband, his sister and mother, showed a pattern consistent with a biallelic position of the D421N and the 1215A>G SNP representing the I312F mutation, as we did not observe any amplification of proband DNA when using the D421N and the SNP selective primers together (Supplemental Figure 1). We also applied the mutation and SNP-specific primers, together with intron-spanning primers, to amplify the proband’s 2 DNA strands for sequencing and consistently observed that the D421N mutation and the 1215A>G SNP were not detected on the same allele (Supplemental Figure 1). Altogether, the data provide strong evidence for a trans-allelic position of the I312F and D421N mutations in the patient. In addition to the DAT-I312F and DAT-D421N mutations, 3 previously reported SNPs were identified in the cohort: c.114A>G (rs6350) with a frequency of 13%, c.1215A>G (rs6347) with a frequency of 40%, and c.1857G>C (rs200712598) with a frequency of less than 1%.

The patient is a male of mixed European descent (aged 40 years at the time of referral) with early-onset parkinsonism and psychiatric symptoms without a family history of movement disorders or psychiatric disease. He suffered from unspecified failure to thrive in infancy and learning problems in school. He was not tested or examined by experts in child psychiatry during childhood or adolescence, but intellectual impairment and behavioral problems with learning difficulties and lack of impulse control were evident to relatives and to school and recreational center staff from early childhood. At age 28, he had insidious onset of right-sided hand tremor. At this time, normal biochemistry excluded disorders such as Wilson’s disease, and a CT brain scan as well as electroencephalography were normal. An amorphophine test resulted in a marked reduction in tremor, but a following therapeutic trial with levodopa treatment was not sufficiently effective to attenuate the tremor and therefore did not support the tentative diagnosis, dopa-responsive dystonia. The patient had a progressive disease course over the following years. Treatment with levodopa was attempted, with minor alleviation of symptoms but concomitant with considerable side effects. At age 35, he had severe coarse tremor in all extremities, truncus, and head during action as well as at rest, accompanied by rigidity and bradykinesia.
Huntington’s disease–like syndrome and parkinsonism were suspected, but our search for extended trinucleotide repeats in HTT, ATXN1-3, CACNA1A, and TBP, mutations in PRKN and GCH1, a hotspot mutation in LRRK2 (Gly2019Ser), and most of the common point mutations and copy number variations in DJ1, PINK, UCHL1, SNCA, and ATP13A2 were all negative. Treatment with the anticholinergic drug orphenadrine (150/150/100/0 mg) was initiated and is hitherto ongoing, with some positive effects in controlling tremor and rigidity. After repeated self-injury behavior, the patient was psychiatrically assessed at age 39 years and diagnosed with ADHD according to DSM-IV (Diagnostic and Statistical Manual of Mental Disorders, fourth edition, American Psychiatric Association, Washington, DC, USA) criteria. Treatment with methylphenidate (Ritalin) produced unacceptable side effects. Symptoms were alleviated according to his caregivers, but not according to the patient.

Presently, the patient has universal tremor (Supplemental Video 1) as well as bradykinesia and rigidity in the upper and lower limbs (Supplemental Videos 2 and 3, respectively). He demonstrates normal-pursuit eye movements, including complete range, and no square wave jerks are seen, but he has saccades with increased latency in initiation, suppressible head movements to initiate, and moderate slowing of movement velocity (Supplemental Video 4). A retropulsion pull test was positive (Supplemental Video 5). Of note, the videos were recorded with the patient on anticholinergic treatment.

To assess possible neurodegeneration in the dopaminergic pathways of the patient, 2 DAT-SPECT scans were performed with an interval of 8 years (2005 and 2013). While both scans showed bilateral reduction of [123I]-FP-CIT binding to DAT in the striatum, DAT availability was further reduced during the 8-year period from approximately 35% to 15% compared with the expected value of age-matched controls (Figure 2A). Moreover, a combined [18F]-Fluorodeoxyglucose ([18F]-FDG) PET/MRI scan was performed in 2013. [18F]-FDG PET showed slight to moderate metabolic reductions cortically in the occipitotemporal junctions bilaterally and intact metabolic activity in the striatum (Figure 2B). MRI volumetric measurements, however, showed a significantly reduced volume of the caudate nuclei after correction for the intracranial volumes (left: 0.195%; right: 0.184%; less than –3 SD). The corrected putaminal volumes were also reduced (left: 0.30%; right: 0.28%), though not significantly (–1.5 SD). Together, these data support a neurodegenerative component of the patient’s disease. Note that despite volume reductions, intact striatal metabolic activity is the common finding, also in late Parkinson’s disease (PD), while cortical reductions may be even more pronounced than those found in this patient (25). Finally, we assessed the level of monoamine metabolites in the cerebrospinal fluid (CSF). The analysis revealed a homovanillic acid (HVA) concentration of 268 nmol/l (reference range, 71–565) and a 5-hydroxyindoleacetic acid (5HIAA) concentration of 104 nmol/l (reference range, 58–220), thus an HVA/5HIAA ratio of 2.6 (reference range, 1.0–3.7).

Figure 1
Identification of missense mutations in SLC6A3. (A) Identified mutations in SLC6A3. Upper panel: nucleotides c.930-938; the c934 A:T mutation, giving rise to DAT-I312F, is denoted as W. Lower panel: nucleotides c.1257-1265; the c1261 G:A mutation, giving rise to DAT-D421N, is denoted as R. (B) Snake diagram of DAT sequence illustrating the location of I312 and D421 in TMs 6 and 8, respectively. (C) Family tree of the patient. The affected patient is indicated by the black square. An unaffected sister also carries the I312F variant (indicated by the gray circle). The mother carries only WT DAT. The diseased father was not available for testing, but was presumably a carrier of the I312F allele (indicated by the gray square). Females are indicated by circles and males by squares. (D) Sequence alignment of 5 vertebrate species showing conservation of I312 as well as D421.
Next, we investigated the cellular processing of the mutants by Western blotting and surface biotinylation. WT DAT expressed in heterologous cells can be found in a mature, fully glycosylated form that elutes at approximately 85 kDa and as an immature ER-glycoform of approximately 55 kDa that is absent from the cell surface (26–28). Consistent with the preserved binding properties of DAT-I312F, Western blot analysis of whole-cell lysates from HEK293 cells showed no reduction compared with WT in the amount of fully glycosylated, mature transporters (Supplemental Figure 3). Likewise, surface biotinylation experiments showed no difference in surface expression of DAT-I312F compared with that in WT (Figure 4, A and B). Interestingly, despite the dramatic reduction we observed in uptake capacity and the absence of measurable [3H]-CFT binding, both Western blot analysis and surface biotinylation showed a significant increase in the levels of mature transporters for DAT-D421N (Supplemental Figure 3 and Figure 4, A and B).

We further assessed the cellular distribution of DAT-I312F and DAT-D421N by immunostainings and confocal microscopy analysis of transfected HEK293 cells (Supplemental Figure 4) and transfected CAD cells (Figure 5), a CN-derived catecholaminergic cell line without endogenous DAT expression (29). Both DAT-I312F and DAT-D421N showed cellular distributions similar to those of WT, with clear plasma membrane staining as well as staining of intracellular structures (Figure 5A and Supplemental Figure 4). To visualize only surface-expressed transporters, we used our recently developed fluorescent cocaine analog JHC 1-64, which binds to correctly folded transporters expressed at the cell surface (30, 31). Following incubation with 10 nM JHC 1-64 and subsequent washing, we observed a strong membrane labeling of cells transfected with WT or DAT-I312F, with no apparent difference in intensity. In contrast, we were unable to detect any membrane staining of DAT-D421N–transfected cells when using 10 nM JHC 1-64 (Figure 5B). However, following incubation with 300 nM JHC 1-64, we observed a strong membrane labeling of DAT-D421N–transfected cells, indicative of a reduced JHC 1-64 affinity for DAT-D421N (Figure 5C). Importantly, labeling at this high JHC 1-64 concentration was blocked by preincubation with nomifensine, a selective DAT blocker (Figure 5D). In summary, both DAT-I312F and DAT-D421N are efficiently targeted to the plasma membrane despite marked functional impairment.

DAT-D421N disrupts the second sodium binding site in DAT. According to a homology model of DAT (Figure 6A), based on the structure of the prokaryotic DAT homolog LeuT (32), Ile312 is located in transmembrane (TM) segment 6, without any direct contacts with the substrate or the bound ions (Figure 6B). However, Asp421 aligns with Thr354 in LeuT, a residue that coordinates the sodium ion bound at the second sodium-binding site (Na2). Similarly, Asp421 coordinates the second sodium ion in our DAT model (Figure 6C). The validity of this model is supported by the newly published structure of Drosophila DAT, in which sodium in Na2 is coordinated in a trigonal bipyramidal fashion by residues in 7 TMs 1 and 8, including Asp420, which corresponds to Asp421 in human DAT (hDAT) (33). To address the consequences of the Asp421Asn mutation on sodium binding, we compared the free energy of binding of Na2, calculated for both WT and DAT-D421N bound to dopamine and ions (34, 35). The structural basis for these results encompasses a change in the Na2 binding site, which in WT involves Gly75,
Figure 3

Functional characterization of DAT-I312F and DAT-D421N by [3H]-dopamine uptake and [3H]-CFT–binding experiments. (A) [3H]-dopamine uptake in transiently transfected HEK293 cells. The curves represent average curves from 6 independent experiments, each with triplicate uptake determinations (5 minutes) at the indicated dopamine concentrations. (B) Vmax values of [3H]-dopamine uptake compared with WT. In each experiment, the Vmax values of mutants were normalized to WT. Michaelis-Menten kinetics was applied to fit data. **P < 0.01 and ***P < 0.0001 by 1-sample t test; n = 6. Uptake by DAT-D421N could not be fitted reliably by Michaelis-Menten kinetics and is therefore denoted as not determined (ND). Km values were fitted to 1.7 ± 0.3 μM for WT; 1.8 ± 0.4 μM for DAT-I312F; and 2.1 ± 0.4 for DAT-I312F plus DAT-D421N. (C) Evaluation of [3H]-dopamine uptake at 6.4 μM of dopamine. The uptake was normalized to WT for each experiment. **P < 0.01 and ***P < 0.0001 by 1-sample t test; n = 6. (D) [3H]-CFT binding to transiently transfected COS-7 cells. Curves represent the averages of 4 experiments, each performed in triplicate. Binding data were fitted by nonlinear regression and normalized to WT in each experiment. While DAT-I312F shows preserved binding properties (Kd values [SE intervals] for DAT-I312F and WT were 16.9 [12.1–23.6] nM and 13.7 [12.7–14.8] nM, respectively), no specific binding to DAT-D421N was detected. All data are the means ± SEM. Bmax for WT was 149 ± 39 fmol/105 cells.

Phe76, Val78, Asp421, and Ser422, whereas in DAT-D421N, it is changed to Gly75, Val78, Leu418, and Asn421. In WT, the free energy of sodium binding in Na2 was more favorable (by ~32 kcal/mol) than the solvation energy of sodium in a water box (~102 kcal/mol), showing a robust affinity of DAT for sodium at this site. In contrast, the corresponding energy for sodium in Na2 of DAT-D421N was calculated to be approximately -98 kcal, i.e., similar to the affinity of sodium for the aqueous solution. Thus, binding of sodium in Na2 is weakened considerably in DAT-D421N compared with that in WT. Following the recent publication of the crystal structure of the nonfunctional mutant of Drosophila melanogaster dopamine transporter (dDAT, PDB code 4M48), a second de novo model of hDAT was constructed with the new structure as the template (33). This enabled a comparative investigation of homology model quality and features, which showed that the root mean square deviation between hDAT based on LeuT and dDAT was 1.3Å for the critical residues in TMs 1, 6, and 8 (Supplemental Figure 5). Thus, the critical regions of the substrate dopamine binding site and ion binding sites, which are key to the inferences we describe, are strikingly similar for both models. Moreover, the full comparison of the original LeuT-based homology model with the ones based on dDAT (Supplemental Figure 5) and also with an additional hybrid model constructed with LeuT as a template for TMs 1–10 and dDAT for TMs 11 and 12 (see Supplemental Figure 6) show good structural agreement and support all conclusions drawn from the molecular dynamics (MD) and free-energy perturbation (FEP) results.

In agreement with the predicted loss of sodium affinity in DAT-D421N, determination of [3H]-dopamine uptake in transient HEK293 cells at increasing sodium concentrations (using ChoCl as a substitute) suggested strong impairment of sodium binding to DAT-D421N, with no sign of uptake saturation at a sodium concentration up to 200 mM (Figure 6D). In contrast,DAT-I312F displayed only a small change in sodium dependency in the uptake experiments (EC50 = 70 ± 20 mM for DAT-I312F vs. EC50 = 30 ± 4 mM for WT; n = 3, means ± SEM). The sodium dependency of [3H]-CFT binding was also preserved for DAT-I312F in COS-7 cells (Figure 6E, EC50 = 29 ± 3.6 mM for DAT-I312F vs. EC50 = 46 ± 10 mM for WT; n = 4, means ± SEM) and in HEK293 cells (EC50 = 25 ± 5.3 mM for DAT-I312F vs. EC50 = 30 ± 2.5 mM for WT; n = 5–7, means ± SEM. Data not shown).

DAT-D421N exhibits anomalous dopamine efflux. Amphetamine is a DAT substrate and is capable of promoting the reversal of dopamine transport, which is believed to play a critical role in its psychostimulatory effect (36–39). To further characterize the functional consequences of the Ile312Phe and Asp421Asn mutations, we assessed amphetamine-induced dopamine efflux by amperometry. As expected, amphetamine caused a large transient increase in the amperometric signal for WT, reflecting oxidation of the released dopamine. We observed a similar response for DAT-I312F, while strikingly different properties were seen once more for DAT-D421N, with amphetamine causing a reduction instead of an increase in the amperometric signal (Figure 7, A–C). To further substantiate this finding, we also assessed amphetamine-induced efflux in a [3H]MPP+ superfusion assay. HEK293 cells were preloaded with the metabolically inert DAT substrate [3H]MPP+, and in agreement with the amperometry studies, amphetamine elicited a clear increase in [3H]MPP+ efflux in the WT and DAT-I312F-expressing cells, while no amphetamine-induced efflux was apparent for cells expressing DAT-D421N. Rather, we observed a relative decrease in [3H]MPP+ efflux through DAT-D421N (Figure 7D), similar to the findings in the amperometry experiments. The data most likely reflect a blocking by amphetamine of a constitutive dopamine-MPP+ efflux through DAT-D421N. If this were the case, a DAT inhibitor would be expected to have the same effect on dopamine-MPP+ efflux as that of amphetamine on DAT-D421N, and indeed, this was what we observed. Like amphetamine, cocaine elicited a decrease in the amperometric current for DAT-D421N, but not for that of WT (Figure 7, E and F). Correspondingly,
we found that cocaine significantly reduced [3H]MPP⁺ efflux in HEK293 cells transiently expressing DAT-D421N (Figure 7G).

**DAT-D421N possesses a large cation leak.** We also sought to uncover the functional consequences of the mutations through a basal electrophysiological characterization using the 2-electrode voltage clamp technique in Xenopus oocytes. Previous studies of WT DAT have shown that stimulation with dopamine gives rise to an inward current at negative potentials, believed to represent the stoichiometrically coupled current plus an uncoupled flux of anions (40–42). As might be expected from the uptake properties of the 2 mutants, we observed a dopamine-stimulated inward current for DAT-I312F, but not for DAT-D421N (data not shown). Next, we evaluated whether the mutants showed changes in DAT leak conductances. WT DAT has been shown to possess a small and hardly detectable cocaine-sensitive inward cation leak in sodium that becomes much larger in lithium (40) (Figure 8A). We observed similar properties for DAT-I312F, with a large cocaine-sensitive current in lithium and no detectable cocaine-sensitive current in sodium (Figure 8B). The properties of DAT-D421N, however, were markedly different. We observed a striking cocaine-sensitive leak current in sodium that was similar in size to that of the current in lithium and that had a similar reversal potential (Figure 8C). Furthermore, the leak in lithium showed a leftward shift in the reversal potential and a more pronounced outward current at positive potentials as compared with WT DAT (Figure 8C). Ion substitutions with ChoCl and Na-gluconate demonstrated that the leak current was carried, at least in part, by sodium. That is, the reversal potential changed markedly by substituting sodium with choline (Supplemental Figure 7), while the reversal potential barely changed by shifting from chloride to gluconate, indicating that the current was not carried by anions (Supplemental Figure 7).

**Discussion**

We describe an adult male with early-onset parkinsonism and neuropsychiatric disorder, who is compound heterozygous for 2 SLC6A3 missense mutations: DAT-I312F and DAT-D421N. DAT-I312F was inherited from his deceased father, while the DAT-D421N mutant presumably is a de novo mutation. Importantly, there is increasing awareness that de novo mutations may comprise a considerable segment of the etiology of complex brain disorders (43–45), and, to our knowledge, this patient represents the first case of parkinsonism as well as ADHD, in which a de novo mutation has been identified in DAT. Both mutations, but in particular DAT-D421N, have deleterious consequences for the transporter function, supporting the idea that mutations in SLC6A3 can be a rare monogenic cause of early-onset atypical parkinsonism and possibly ADHD. In general, 5%–10% of parkinsonism cases can be attributed to monogenic causes, giving rise to both dominant and recessive modes of inheritance (46). The most common cause of early-onset parkinsonism is mutation in parkin, which is estimated to account for 1.4% to 8.2% of early-onset cases (46, 47).

DAT deficiency syndrome was recently described as an autosomal recessively inherited disease, manifesting in childhood, typically early infancy, with parkinsonism-dystonia (14, 15, 48). Mutations in SLC6A3 that probably lead to ER retention of the encoded DAT proteins were identified as the cause of disease. According to our cellular analyses, the DAT mutants carried by the present patient showed markedly different properties, with no signs of ER retention. For DAT-D421N, we even observed increased levels of the mature transporter glycoform. Moreover, confocal imaging and biotinylation experiments demonstrated efficient targeting of DAT-I312F and DAT-D421N to the plasma membrane. Nevertheless, both mutants, and in particular DAT-D421N, displayed impaired uptake capacity. Despite a reduced uptake capacity, DAT-I312F bound the cocaine analog [3H]-CFT indistinguishably from that in WT and showed little change in sodium affinity, suggesting that mutation of isoleucine in position 312 to phenylalanine is likely to cause a structural perturbation that indirectly impacts catalytic activity without altering expression and folding of the transporter. In contrast, DAT-D421N showed no detectable [3H]-CFT-binding activity. It is conceivable that this is caused by decreased CFT affinity, preventing the detection of specific binding. Notably, we obtained evidence for decreased affinity of the fluorescent cocaine analog JHC 1-64, i.e., 300 nM was required to visualize DAT-D421N at the plasma membrane of transfected cells. The decreased affinity of these inhibitors, as well as the impaired dopamine uptake, is undoubtedly the result of vastly compromised binding of sodium to DAT-D421N, as supported by our functional studies, computational simulations, and by the recent crystal structure of Drosophila DAT, confirming the participation of the aspartate in sodium coordination (33). DAT is suggested, like LeuT, to possess 2 sodium sites (Na₁ and Na₂) and...
to cotransport 2 sodium ions during each transport cycle (49, 50). Binding of these sodium ions is believed to have a critical role in promoting both substrate binding and binding of cocaine analogs by stabilizing the transporter in an outward-facing conformation (5, 32, 51–53). Moreover, mutation of Thr354 in LeuT, which is the residue corresponding to Asp421 in hDAT, has been shown to contribute to the stabilization of LeuT in an inward-facing conformation (32). It is therefore conceivable that DAT-D421N is predominantly inward facing, leading to a dramatic decrease in apparent dopamine affinity and thereby an impairment of uptake.

Impaired sodium binding to Na2 might also explain the efflux data showing a reverse effect of amphetamine. Instead of inducing dopamine-MPP+ efflux, as in WT DAT, we observed that amphetamine treatment of cells expressing DAT-D421N produced a decline in the amperometric current as well as a decline in MPP+ efflux, suggesting that amphetamine blocks a constitutive, anomalous leak of substrate through DAT-D421N. The predicted switch in the conformational equilibrium of DAT-D421N to an inward-facing configuration might favor constitutive efflux of dopamine, and under these conditions, amphetamine may act as an inhibitor rather than an inducer of efflux. Indeed, we observed a similar decline in amperometric signal as well as a decline in baseline [3H]-MPP+ efflux following cocaine treatment of DAT-D421N–expressing cells compared with treatment of WT DAT–expressing cells. A similar constitutive anomalous dopamine efflux has been reported for the DAT missense mutation DAT-A559V, identified in a bipolar patient and in 2 brothers with ADHD, and for DAT-T356M, identified in a patient with autism spectrum disorder (24, 54). Importantly, cocaine-sensitive anomalous dopamine efflux has been demonstrated in dopaminergic neurons from DAT-KO mice transfected with DAT-A559V, and a preliminary characterization of DAT-A559V knock-in mice reported elevated

Figure 5
Visualization of DAT mutants by confocal microscopy. (A) Cellular distribution of WT and DAT mutants in transiently transfected CAD cells. DAT was visualized by immunostaining with an N-terminal antibody. No differences were observed in the localization of DAT-I312F or DAT-D421N compared with WT. (B) Confocal live imaging of transfected CAD cells. Cells were stained with the fluorescent cocaine analog JHC 1-64 (10 nM) to obtain specific labeling of DAT present in the plasma membrane. While cells expressing WT, DAT-I312F, or both DAT-I312F and DAT-D421N achieved clear membrane labeling with 10 nM of JHC 1-64, no membrane labeling was observed in cells expressing the DAT-D421N mutant alone. (C) Upon labeling of transfected CAD cells with 300 nM JHC1-64, both WT and DAT-D421N–transfected cells showed clear membrane labeling, which was not observed when cells were preincubated with 1 μM nomifensine, a selective blocker of DAT (D). Images shown are representative of 3 independent experiments. Images of WT and mutants were taken with identical settings.
extracellular dopamine levels (55, 56). Thus, anomalous efflux can even occur in native systems with an intact vesicular dopamine storage system. The identification of anomalous dopamine efflux via DAT-D421N provides additional support to the suggestion that this efflux produced by DAT-coding variants is a potential risk factor in psychiatric disorders (24, 54). The present data might also support the notion that alterations in DAT function or expression levels in general could be a risk factor in ADHD, as indicated by several previous studies (57). It is nonetheless important to note that the patients carrying DAT-A559V and DAT-T356M were heterozygous and that the mutants possessed larger or unaffected dopamine uptake capacities compared with those of DAT-D421N. These patients would therefore be expected to have considerable residual DAT function despite anomalous dopamine efflux, and motonic dysfunctions were not reported for the DAT-A559V and DAT-T356M carriers (21, 24, 54).

The mechanism by which inadequate and/or altered DAT function can lead to parkinsonian symptoms remains unresolved (58, 59). Indeed, the reduced uptake capacity of both DAT-I312F and DAT-D421N is likely of key importance for the dopaminergic dysfunction of the patient, but given the generally much more pronounced phenotype of DAT-D421N, we cannot exclude that the majority of alterations underlying the clinical presentation of the patient can be ascribed to DAT-D421N. It is nonetheless clear that the predicted decreased DAT function will compromise dopamine reuptake and thereby prevent transmitter reuse as well as impair clearance of released dopamine. In addition, impaired dopamine clearance might inhibit dopamine synthesis through overstimulation of presynaptic dopamine D2 autoreceptors and cause downregulation of postsynaptic dopamine receptors (58, 59). In DAT-KO mice, however, presynaptic D2 receptors are downregulated and TH is upregulated (60, 61), which could argue against the involvement of TH inhibition via D2 activation in humans with compromised DAT function. In spite of this, severe dopamine deficiency may develop and inadvertently lead to motoric dysfunction and appearance of parkinsonian symptoms. Interestingly, the majority of DAT-KO mice show a hyperdopaminergic behavioral phenotype and do not display parkinsonian symptoms. The mechanism by which inadequate and/or altered DAT function can lead to parkinsonian symptoms remains unresolved (58, 59).

Figure 6
Molecular modeling of hDAT. (A) Homology model of hDAT based on the outward-open crystal structure of LeuT, a bacterial homolog. (B) I312, indicated in green in TM 6, was seen to have no direct contact with dopamine or ion binding sites. (C) D421, indicated in green in TM 8, was involved in coordinating binding of sodium at the second sodium binding site (Na2). Sodium and chloride ions are indicated in purple and turquoise, respectively. (D) Sodium dependence of [3H]-dopamine uptake. Dopamine uptake was evaluated at increasing concentrations of sodium. The sodium dependence of DAT-I312F–mediated uptake was comparable to that of WT (Kd = 70 ± 20 mM for DAT-I312F vs. Kd = 30 ± 4 mM for WT). DAT-D421N, on the other hand, showed impaired binding of sodium, seen as a linear sodium dependence curve with no tendency to saturate at 200 mM of NaCl. (E) Sodium dependence of [3H]-CFT binding. Sodium dependence of [3H]-CFT binding to DAT-I312F was similar to that of WT (Kd = 29 ± 3.6 mM for DAT-I312F vs. Kd = 46 ± 10 mM for WT), while specific binding of [3H]-CFT to DAT-D421N could not be detected. The curves of D and E are average curves from 3 and 4 experiments, respectively, each performed in triplicate and normalized to WT Bmax. ChoCl was used for equimolar cation substitution of sodium to obtain indicated sodium concentrations. Bmax and Kd values were derived using a one-site–specific binding model.
Figure 7
Amphetamine- and cocaine-induced amperometric currents and MPP⁺ efflux. HEK293 cells were transiently transfected with WT, DAT-I312F, or DAT-D421N. (A) Representative amperometric currents; arrows indicate the application of amphetamine (10 μM). (B) Quantification of the amphetamine-induced peak amperometric currents. (C) Amperometric data are reported as the amphetamine-induced amperometric current recorded 10 minutes after amphetamine application. Data are the means ± SEM; n = 7–9. **P < 0.01 and ***P < 0.001 by Mann-Whitney U test. (D) Quantification of amphetamine-induced [³H]MPP⁺ efflux presented as the AUC of MPP⁺ efflux. [³H]MPP⁺ release was calculated by subtracting the basal release from total release during the first 8 minutes following amphetamine exposure. Data are the means ± SEM; n = 4–5. *P < 0.025 by Mann-Whitney U test. (E) Representative amperometric currents following cocaine exposure (50 μM) of HEK293 cells transiently expressing WT DAT or DAT-D421N. (F) Amperometric data are reported as the cocaine-induced amperometric current 10 minutes after cocaine application. Data are the means ± SEM; n = 3. *P < 0.05 by 1-tailed t test. (G) Quantification of cocaine-induced [³H]MPP⁺ efflux presented as the AUC of MPP⁺ efflux. Cocaine treatments led to a significantly higher reduction in MPP⁺ efflux from cells expressing DAT-D421N compared with WT. *P < 0.05 by Mann-Whitney U test, consistent with a constitutive leak of dopamine.

compared with years in humans. Nevertheless, the level of remaining DAT uptake capacity is likely crucial for whether dopamine deficiency will develop and, accordingly, when parkinsonian symptoms become apparent. It is also possible that the constitutive, anomalous substrate efflux, mediated by DAT-D421N, contributes to dopaminergic dysregulation by further reducing intraneuronal stores of dopamine.

Another potentially important phenotypic property of DAT-D421N is a large cation leak identified by our electrophysiological analyses. This leak is likely linked to the impaired sodium binding to Na₂, thereby supporting a key role of Na₂ in governing cation permeability of the transporter. Interestingly, previous studies have indicated that the uncoupled substrate-induced current in DAT can affect neuronal excitability (42), suggesting that DAT-mediated currents are sufficiently large to impact neuronal function. The leak in DAT-D421N is markedly larger than the substrate-induced current of the WT. Thus, the current might contribute to the dopaminergic dysregulation. Given the large amount of DAT protein believed to be expressed in dopaminergic neurons, one could also speculate that the dopaminergic neurons become more vulnerable as a consequence of the stress enforced by the leak, which could implicate the leak in the neurodegenerative component of the patient’s disease. Note that there is indirect evidence for a potentially harmful effect of cation influx, as blocking sodium influx via sodium channels and the Na⁺/H⁺ exchangers has been shown in several cases to have a neuroprotective effect (62, 63).

Our findings provided strong evidence for dopaminergic neurodegeneration in the patient studied. Comparison of DAT-SPECT scans performed 8 years apart (2005 and 2013) revealed a sizable loss of [¹²³I]-FP-CIT–binding capacity (from 35% to ~15% of normal capacity). It is unlikely that this loss was a consequence of gradually decreased expression of mutant DATs in otherwise intact dopaminergic neurons; rather, it indicates a neurodegenerative component with progressive loss of dopaminergic pathways. Such a neurodegenerative component was supported by striatal volumetric measurements based on a [¹⁸F]-FDG PET/MRI scan, which showed reduced volumes of the caudate nuclei, in accordance with previous findings (64). The neuronal loss might also explain why we do not see an increased ratio between the dopamine metabolite HVA and the serotonin metabolite 5HIAA in CSF, as reported in children with DAT deficiency syndrome (14, 15). Notably, some studies report reduced HVA levels in the CSF of Parkinson’s patients (65, 66), arguing that the normal, not decreased, HVA value of our patient reflects a relative increase in the accumulation and degradation of extraneuronal dopamine as compared with the putative numbers of remaining neurons. Moreover, residual activity of the DAT mutants carried by the proband would also be expected to oppose extracellular dopamine accumulation compared with patients with no DAT expression. Specialized diagnostic laboratories should therefore be aware that for some individuals with DAT mutations, an unremarkable CSF profile might be recorded. Interpretation, in conjunction with clinical details, is therefore essential.

In summary, we describe here what we believe to be the first link between DAT missense mutations and early-onset parkinsonism in adults. Furthermore, the neuropsychiatric comorbidity of the proband strongly supports the notion that complex DAT genotype-phenotype relationship. 

sonian features, except after blockade of dopamine synthesis by inhibition of TH (12). A subpopulation of the mice sporadically develops progressive motor dysfunction with eventual paralysis and death, but this phenotype has been attributed to loss of GABAergic neurons rather than of dopaminergic neurons (13). The discrepancy between humans and mice could indicate that DAT is even more important for sustaining a sufficient pool of releasable dopamine in humans, but could also be related to the time course of the disease, because the maximum time of exposure to the dysfunctional uptake would be 18–24 months in mice.
notype correlations might contribute significantly to different dopaminergic pathologies. Thus, the present findings should provide a critical basis for further exploration of how dopaminergic dysfunction and altered DAT function contribute to brain disorders.

Methods

Subject and procedures. We included 91 unrelated patients from a referral-based hospital cohort. The patients were referred for genetic testing due to early-onset parkinsonism or a related atypical movement disorder. One patient underwent further assessment due to identification of mutation in SLC6A3. In addition, the patient was referred for genetic testing due to a familial history of Parkinson's disease or a related atypical movement disorder. The patients were referred for genetic testing due to early-onset parkinsonism or a related atypical movement disorder. The patients were referred for genetic testing due to early-onset parkinsonism or a related atypical movement disorder.

Saturation kinetics of dopamine uptake in HEK293 cells were determined using a 2-fold dilution row (final concentrations of 0.4–0.05 μM) with constant specific activity. The dilution row was composed of a mixture of [2,5,6] (3H)-dopamine uptake (PerkinElmer) and unlabeled dopamine (Sigma-Aldrich), essentially as described in ref. 72. Briefly, transiently transfected HEK293 cells were seeded onto 24-well plates (175,000 cells per well) coated with polyornithine. On the day of the experiment (48 hours after transfection), cells were washed once in uptake buffer (25 mM HEPES, 120 mM NaCl, 5 mM KCl, 1.2 mM CaCl2, and 1.2 mM MgSO4 supplemented with 5 mM D-glucose, 1 mM ascorbic acid, and 1 μM of the catechol-O-methyltransferase [COMT] inhibitor R-310960, pH 7.4) and equilibrated in uptake buffer for 15 minutes at room temperature.

Figure 8

DAT-D421N possesses a cocaine-sensitive cation leak in sodium. (A–C) I/V diagrams of the cocaine-sensitive steady-state currents (control - cocaine) in sodium chloride (NaCl; squares) and lithium chloride (LiCl; circles) for WT (A), DAT-I312F (B), and DAT-D421N (C). WT and mutant constructs were expressed in Xenopus oocytes and analyzed by the 2-electrode voltage clamp technique as described in Methods. The I/V diagrams were generated in 20-mV steps from –100 to +40 mV. The currents are normalized to the maximum current in lithium at a membrane potential of –100 mV (WT, –402 ± 59 nA; DAT-I312F, –175 ± 27 nA; DAT-D421N 219 ± 31 nA; means ± SEM; n = 4–6).

1.0 × 1.0 × 1.0 mm³, and an acquisition time of 5 minutes. Anatomical imaging using T2 was performed as follows: turbo spin echo (TSE) sequence, with a TR of 6,000 ms, a TE of 96 ms, a flip angle of 150°, a voxel size of 0.7 × 0.7 mm³, a slice thickness of 4 mm, and an acquisition time of 1.25 minutes. The [18F]-FDG PET scan was attenuation corrected using low-dose CT scanning (69) and compared with a database of healthy subjects (Sciemct 3.0, Syngo-Via; Siemens Healthcare). Volumetric measurements using FreeSurfer version 5.2.0 were performed from T1 MP-RAGE MRI images of the caudate nuclei, putamina, and the intracranial volume and compared with a group of healthy age-matched subjects exposed to identical imaging procedures and analyses.

SNP analysis. Two hundred Danish controls and the patient were subjected to SNP analysis. The probes used were custom-made by Life Technologies to target the variants c.934A>T and c.1261G>A. The analysis was done using the TaqMan SNP Genotyping Assay (Life Technologies) and was performed according to the manufacturer’s instructions.

Site-directed mutagenesis. cDNA encoding the hDAT mutants DAT-I312F and DAT-D421N were made on a pRC/CMV hDAT background using the QuikChange site-directed mutagenesis kit (Stratagene). Constructs were verified by DNA sequencing.

Cell culture and transient transfection. Human embryonic kidney 293 (HEK293) cells and COS-7 cells were maintained as previously described (70, 71). Cath.-a-differentiated (CAD) cells from a catecholaminergic cell line were grown in media composed of 50% DMEM and 50% Ham’s F-12 nutrient mixture (all from Invitrogen) at 37°C in a humidified incubator with 5% CO2. All media were supplemented with 1% penicillin-streptomycin. Transient transfections were performed using Lipofectamine 2000 (Invitrogen), Fugene-6 (Roche Molecular Biochemicals, for amperometry data), or Lipofectamine Plus (Invitrogen, for superfusion assay), according to the manufacturer’s protocols, using a 3:1 transfection reagent/DNA ratio. Assays were conducted 24–48 hours after transfection.
temperature (RT). Uptake was initiated by the addition of the dilution row mixture and terminated after 5 minutes (at RT) by washing twice in ice-cold uptake buffer. Cells were lysed in 1% SDS and transferred onto 24-well counting plates (PerkinElmer). Scintillation fluid was added, and the radioactivity was measured in a Wallac MicroBeta TriLiquid Scintillation Counter (PerkinElmer). Nonspecific binding was determined using cells transfected with a similar amount of empty plasmid. K_{n0} and V_{max} values were derived by fitting Michaelis–Menten kinetics to the background-corrected uptake data using GraphPad Prism 5.0 (GraphPad Software). All determinations were performed in triplicate.

Sodium dose-response curves were obtained by equimolar titration of 200 mM NaCl in anion substitution buffer (250 mM Tris-HEPES, 50 mM K-glucuronate, 12 mM Mg-glucuronate, and 12 mM Ca-glucuronate, pH 7.4) against ChoCl in anion substitution buffer to obtain final Na⁺ concentrations of 200, 130, 80, 40, 20, 10, 5, and 0 mM. These solutions were supplemented with 5 mM D-glucose, 1 mM ascorbic acid, and 1 μM of the COMT inhibitor. Transfected HEK293 cells were washed in wash buffer (200 mM Cho-glucuronate in anion substitution buffer) and equilibrated for 10 minutes in the denuded sodium buffers. Uptake was initiated by addition of a dopamine mixture of unlabeled dopamine and [2,5,6] (H)-dopamine to obtain a final dopamine concentration of 6.4 μM. Uptake was allowed for 5 minutes (WT DAT) or 10 minutes (mutants) and terminated by washing twice in ice-cold wash buffer. Cell lysis, transfer, and counting were carried out as described above. Nonspecific binding was determined using cells transfected with an equivalent amount of plasmid vector without insert. K_{n0} values and maximum uptake (V_{max}) were derived in GraphPad Prism 5.0.

[3H]-CFT binding. Binding assays were carried out as described previously (73) on intact COS-7 cells or HEK293 cells using [3H]β-carbomethoxy-3β-(4-fluorophenyl)tropane (CFT) (76.6 Ci/mmol; PerkinElmer) as radioligand. Cells were washed once in ice-cold binding buffer (25 mM HEPES, 130 mM NaCl, 5.4 mM KCl, 1.2 mM CaCl₂, 1.2 mM MgSO₄, 1 mM L-ascorbic acid, 5 mM D-glucose, and 1 μM Ro 41-0960 [Sigma-Aldrich], pH 7.4). Unlabeled CFT was added at the concentrations indicated in Figure 3D and Supplementary Figure 2, followed by addition of [3H]-CFT (2–5 nM). Cells were then incubated at 5°C for approximately 100 minutes to reach equilibrium, and the assay was terminated by washing twice in ice-cold binding buffer. Transfer and counting were performed as described above. Data were fitted by nonlinear regression using GraphPad Prism 5.0 to calculate maximum binding (B_{max}) and affinity (K_{D}). Sodium dependency curves of [3H]-CFT binding were performed as described for the [3H]-dopamine uptake assay, though cells were incubated for approximately 100 minutes with [3H]-CFT in the sodium concentrations indicated in Figure 6. Nonspecific binding was determined with 1 μM nonfensine (Sigma-Aldrich).

Surface biotinylation. Surface biotinylation experiments were carried out essentially as described previously (28). Briefly, HEK293 cells were seeded in 6-well plates (500,000 cells per well) and transfected with 0.25 μg DNA per well the following day. Twenty-four hours after transfection, cells were washed in ice-cold PBS (pH 7.4) and incubated with biotin for 40 minutes on ice (1 mg/ml in PBS), followed by 2 washes in glycine and PBS (100 mM) and 2 washes in PBS. Cells were lysed in solubilization buffer (25 mM Tris, pH 7.5, containing 150 mM NaCl, 1% Triton X-100, 0.2 mM phenylmethylsulfonyl fluoride, and complete protease inhibitor cocktail; Roche). Lysates were centrifuged for 20 minutes (16,000 ×g). Supernatants were collected, and equal amounts of protein were loaded on avidin beads and left over-night at 4°C. The avidin beads were pelleted by centrifugation and washed 4 times with solubilization buffer. Protein was eluted in 2X loading buffer containing 100 mM DTT and incubated at 37°C for 30 minutes. Total lysates were likewise incubated for 30 minutes with 2X loading buffer containing 100 mM DTT. Avidin beads were removed by filtration. Equal volumes of samples were separated by SDS-PAGE, and DAT was detected as described for Western blotting.

Cell lysate preparation and Western blot analysis. Transiently transfected HEK293 cells were washed twice in ice-cold PBS and lysed in solubilization buffer (25 mM Tris, pH 7.5, containing 150 mM NaCl, 1% Triton X-100, 0.2 mM PMSF, and complete protease inhibitor cocktail; Roche). The lysates were incubated at 4°C for 20 minutes with end-over-end rotation, followed by a 20-minute centrifugation at 16,000 ×g. Protein concentration in the supernatant was determined using a BCA Protein Assay Kit (Pierce Biotechnology). The supernatant was mixed with 4× SDS loading buffer containing 100 mM DTT and incubated for 30 minutes at 37°C. An equal amount of protein was separated by SDS-PAGE (using Any kD Gels from Bio-Rad) and transferred to Immobilon-P membranes (Millipore). Membranes were blocked for 1 hour in blocking buffer (PBS containing 0.05% Tween-20 and 5% dry milk) and incubated with primary rat anti-DAT antibody (1:1,000, MAB369; Millipore) in blocking buffer for 1 hour at RT or overnight at 4°C. Membranes were then washed 3 times in PBS containing 0.05% Tween-20 and incubated for 1 hour with secondary HRP-coupled goat anti-rat antibody (1:4,000 in blocking buffer). After washing, the blots were visualized by chemiluminescence (ECL kit; Amersham Biosciences) using AlphaEase (Alpha Innotech) and quantified with ImageJ software (NIH). To verify equal protein loading, membranes were reprobed for β-actin (Sigma-Aldrich).

Immunocytochemistry and JHC 1-64 live imaging of DAT. Immunocytochemical staining for DAT in transiently transfected CAD cells or HEK293 cells was performed using an N-terminal primary antibody (rat-anti DAT 1:1,000, MAB 369; Millipore) and Alexa Fluor 488 as a secondary antibody (goat anti-rat, 1:500), as described previously (72). For live imaging of DAT with 10 nM or 300 nM JHC 1-64, cells were grown in polyornithine-coated 2-well Lab-Tek chambers (Nunc). The procedure was carried out as described previously (72). As a control for DAT specificity, cells were preincubated for 20 minutes with 1 mM nomifensine. Confocal microscopy analysis was performed using a Zeiss LSM 510 confocal laser-scanning microscope (Carl Zeiss) with an oil-immersion 63 × 1.4 numerical aperture objective (Carl Zeiss). Alexa Fluor 488 dye was excited with a 488-nm laser line from an argon-krypton laser, and detection of the emitted light was done using a 505- to 530-nm bandpass filter. JHC 1-64 was excited with a 543-nm helium-neon laser, and fluorescence was recorded using a 560-nm long-pass filter. Images were combined using ImageJ software.

Molecular modeling and simulations. The molecular models for hDAT were constructed with MODELLER 9v10 (74) program and based on either the recent outward-open crystal structure of the bacterial member of the NSS family LeuT (PDB code: 3TTT and the sequence alignment detailed previously (75), or on the recently published dDAT structure (PDB code: 4M48) (76). The substrate, dopamine, was positioned in the binding site by aligning the amine group and the hydrophobic portion with those of the substrate in the LeuT structure (77). The final model also included 2 Na⁺ ions positioned equivalently to those in LeuT and a Cl⁻ ion coordinated by residues Asn82, Tyr102, Ser321, and Asn353 of DAT, based on the chloride binding site described previously (78, 79). For the MD simulation of DAT models, we prepared a lipid bilayer model composed of the following mixture of neutral lipids: 60% palmitoyl-oleoyl-phosphatidylcholine (POPC) (shared equally on the upper and lower leaflets), 25% 1-palmitoyl-2-oleoyl-phosphatidylethanolamine (POPE) (on the lower leaflet), 5% PIP₂, and 10% sphingomyelin (SGM), shown to be essential for DAT binding to the membrane (80). Thus, the simulated system, composed of the DAT model immersed in an explicit representation of the water/lipid bilayer/water environment, was constructed with the VMD software and equilibrated using NAMD software version 2.7 (81) and the all-
atom CHARMM27 force field. The simulated system, including the transporters embedded in a membrane patch and water layers on each side containing Na+ and Cl− ions (corresponding to a concentration of 150 mM NaCl), was composed of approximately 340,000 atoms in a box with final dimensions of 168 × 165 × 153 Å (at the end of 100 ns of free equilibration).

During the equilibration of the DAT model in its environment, the backbone was initially fixed and then harmonically constrained, and water was restrained by small forces from penetrating the protein-lipid interface. The constraints were released in 3 steps of 300 ps each, changing the force constants gradually from 1 to 0.5 kcal/mol Å2 and 0.1 kcal/mol Å2, respectively. A constant temperature (310 K) was maintained with Langevin dynamics, and 1 atm constant pressure was achieved by using the hybrid Nosé-Hoover Langevin piston method on a flexible periodic cell (35).

To model the mutant DAT construct with D421N, the mutation was introduced with an FEP carried out with NAMD in the system described above and as described in detail previously (34, 35). According to this protocol, the coupling parameter α was varied from 0 to 1 such that each window did not exceed 3 kcal/mol. In general, the hysteresis tests did not exceed 1 kcal/mol between annihilation and creation.

Amperometry. Transiently transfected HEK293 cells plated on 35-mm dishes were washed with Krebs-Ringer HEPS (KRH) buffer (130 mM NaCl, 4.8 mM KCl, 2.2 mM CaCl₂, 1.1 mM MgSO₄, 1.2 mM KH₂PO₄, and 25 mM HEPS, pH 7.4) containing 10 mM D-glucose, 100 μM parabanic acid, and were incubated with 1 μM ascorbic acid and NaCl, 10 mM HEPES, 34 mM D-glucose, 1.5 mM CaCl₂, 0.5 mM MgSO₄, and 1.3 mM KH₂PO₄. Dopamine was measured by amperometric (oxidation) currents using a carbon fiber electrode (5 mm; Dagan Corporation). The electrode was positioned in apposition to the plasma membrane and held at +700 mV with respect to the bath ground (a potential further investigations).

Statistics. Unless otherwise stated, we have applied a 1-sample t test. For all datasets in which WT DAT was compared with more than 1 mutant, the significance level α was adjusted to account for multiple tests. We applied the Bonferroni’s correction, αcorr = α/n, with n being the number of individual tests and α = 0.05. Thus, the significance level was set at P < 0.025 when performing 2 individual comparisons and P < 0.0166 when performing 3 individual tests.

Study approval. Patients were included from a referral-based hospital cohort, and sequencing of SLC6A3 in patients was approved by the Danish Regional Committee on Biomedical Research Ethics (Copenhagen, Denmark). Informed consent was obtained from the proband prior to any further investigations.

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