Dephosphorylation of human dopamine transporter at threonine 48 by protein phosphatase PP1/2A up-regulates transport velocity

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Several protein kinases, including protein kinase C, Ca²⁺/calmodulin-dependent protein kinase II, and extracellular signal–regulated kinase, play key roles in the regulation of dopamine transporter (DAT) functions. These functions include surface expression, internalization, and forward and reverse transport, with phosphorylation sites for these kinases being linked to distinct regions of the DAT N terminus. Protein phosphatases (PPs) also regulate DAT activity, but the specific residues associated with their activities have not yet been elucidated. In this study, using co-immunoprecipitation followed by MS and immunoblotting analyses, we demonstrate the association of DAT with PP1 and PP2A in the mouse brain and heterologous cell systems. By applying MS in conjunction with a metabolic labeling method, we defined a PP1/2A-sensitive phosphorylation site at Thr-48 in human DAT, a residue that has not been previously reported to be involved in DAT phosphorylation. Site-directed mutagenesis of Thr-48 to Ala (T48A) to prevent phosphorylation enhanced dopamine transport kinetics, supporting a role for this residue in regulating DAT activity. Moreover, T48A-DAT displayed increased palmitoylation, suggesting that phosphorylation/dephosphorylation at this site has an additional regulatory role and reinforcing a previously reported reciprocal relationship between C-terminal palmitoylation and N-terminal phosphorylation.

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suggestive of transporter modulation by other interacting proteins or other PKC-responsive residues in DAT.

There are many other serine and threonine residues in DAT intracellular domains that represent potential phosphorylation sites that may contribute to kinase-mediated functions. To search for additional phosphorylation sites, we have utilized MS previously to identify phosphorylation of rat DAT (rDAT) at PKC site Ser-7 and extracellular signal–regulated kinase site Thr-53 (16, 17). In a series of studies, we demonstrated that these residues mediate complex kinase- and phosphorylation-dependent functions, including regulation of cocaine analog affinity and amphetamine-induced DA efflux (16, 17). A further picture emerging from these studies is that regulation of DAT function occurs through integration of multiple post-translational inputs. For example, transport kinetic capacity is established by reciprocal phosphorylation of Ser-7 and palmitoylation of Cys-580 (18). Interaction of these effects could occur at the level of the enzyme pathways for phosphorylation (kinases and phosphatases) and palmitoylation (protein acyltransferases and acyl protein thioesterases) (19), which could be regulated by distinct signaling cascades. Alternatively, modification status could be regulated through accessibility of the sites, which may be affected by transporter activity, interactome alterations, or N-terminal conformation. Identification of enzymes that catalyze modification of these sites and determining how they are regulated or have access to DAT domains is thus crucial for elucidating DAT regulatory mechanisms.

Reversible protein phosphorylation is mediated by coordinated control of protein kinases and protein phosphatases (PPs). Most studies on DAT regulation by phosphorylation have focused on kinase inputs, with less known regarding dephosphorylation mechanisms. It has previously been shown that okadaic acid (OA), an inhibitor of PP1/2A, down-regulates DAT activity and increases DAT phosphorylation level by reducing dephosphorylation (8, 10, 12, 20, 21). Studies with specific PP1/2A peptide and small molecule inhibitors support dephosphorylation of DAT in rat striatum by PP1 and to a lesser extent PP2A (21), and the catalytic subunit of PP2A (PP2Ac) has been identified in co-immunoprecipitation complexes with DAT in mouse striatal synaptosomes (22). Interactions between DAT and PP2Ac in the rat caudate putamen were increased by cocaine self-administration followed by 3 weeks of abstinence (23), supporting a role for this enzyme in regulating DAT responses to drug exposure. However, the phosphorylation sites regulated by these enzymes remain unknown.

To investigate these issues, we aimed at identifying DAT-associated protein phosphatases and characterize their respective phosphorylation sites. Here, we demonstrate that PP1 and PP2A were complexed with both mouse striatal DAT (mDAT) and heterologously expressed human DAT (hDAT) and that pharmacological inhibition of PP1/2A increased phosphorylation of hDAT at Thr-48, a previously unknown DAT phosphorylation site. Functional analysis of this site via site-directed mutagenesis revealed a role for its involvement in DA uptake and reciprocal palmitoylation events in hDAT.

Results

PP1 and PP2A associate with DAT

To screen for DAT association with PPs, we applied co-immunoprecipitation (co-IP) followed by LC-coupled tandem MS (LC-MS/MS) to confirm the association between DAT and PP2Ac, as well as to screen for association of any other protein phosphatases with DAT. DAT protein complexes were isolated from striatal synaptosomes prepared from WT and DAT−/− knockout (KO) mice by immunoprecipitation with anti-DAT antibody. Proteins were separated by SDS-PAGE with colloidal blue staining. Each lane was excised and divided into 12 pieces each for in-gel tryptic digestion, and the resulting tryptic peptides were subjected to LC-MS/MS. Specific associations were determined by comparison of proteins from the WT and KO samples (Fig. 1A).

MS identified 39 representative DAT peptides in the immunopurified protein complex from the WT mouse striatum, which covered 34.89% of the mouse DAT sequence (Fig. 1B and Table S1). In the same protein complex from the WT striatum, MS identified three peptides for PP1 α, β, and/ or γ catalytic subunits (Table 1). In addition, PP2A regulatory subunits A and B as well as PP2A catalytic subunit α/β were identified as co-immunoprecipitated proteins with DAT exclusively in the WT striatum (Table 1 and Fig. 1 (C–E)). The association between DAT and PP2A regulatory subunit α/β was confirmed by co-IP coupled with immunoblotting for mouse striata and HEK293 cells stably expressing YFP-tagged hDAT (YhDAT-HEK cells) (Fig. 2). Immunoreactivities of PP2A regulatory subunit α were observed in immunopurified DAT complexes from WT striata and YhDAT-HEK cells but not KO mouse striata and parent HEK293 cells, which do not express DAT (Fig. 2). PP1 α catalytic subunit was also co-purified with DAT in YhDAT-HEK cells (Fig. 2B). MS identified co-purified PP2B with 24 peptides in the mouse WT striata, but parallel MS analysis also detected five peptides in the KO mouse striata (Table S2). Moreover, this association was not verified in a heterologous cell system examined in parallel (Fig. S1), and thus, we excluded this protein for further studies.

Identification of phosphorylation in DAT

We detected phosphorylation of DAT residue Thr-53 in the mouse striatum by LC-MS/MS (Fig. S2), confirming previous findings obtained by immunoblotting with a Thr-53–phosphospecific antibody (16). However, because this identification could not provide any evidence for PP1/2A-responsive phosphorylation sites, we stably expressed hDAT in LLC-PK1 cells in the presence or absence of a PP1/2A inhibitor OA to determine which phosphorylation sites were affected.

After immunopurification of hDAT from cell lysates, we repeated the MS analyses as applied for mouse DAT proteins. MS identified DAT from three different gel regions (Fig. 3, A and B), which corresponded to three immunoreactive YFP-DAT bands (Fig. 2B, left). MS identified a few phosphopeptides in hDAT under basal conditions, but treatment with various concentrations of OA increased the MS identification rate for hDAT phosphopeptides, especially the sequence
36EQNGVQLTSSTLTNPR51 (Table 2). We manually confirmed these phosphopeptides with a peak representing neutral loss of phosphoric acid, a signature of phosphopeptides, in MS/MS spectra. This tryptic peptide contains two serine (Ser-44 and -45) and three threonine (Thr-43, -46, and -48) residues that could represent potential phosphoacceptor

Figure 1. Identification of protein phosphatases in the mDAT complex by LC-MS/MS. A, flow diagram of co-IP and MS for the identification of DAT interacting PPs from WT and DAT−/− KO mouse striata. Immunoblot images show DAT protein expression exclusively in the WT striatal lysate. B (top), the double-charged (2+) peptide obtained at m/z 638.85 was fragmented to produce an MS/MS spectrum with y- and b-ion series that identified the sequence LAYAITPEKDR (amino acids 591–601) from mouse DAT. B (bottom), sequence coverage of mouse DAT with identified peptides (boldface type) by MS/MS in mouse striata. C (left), the MS/MS spectrum, obtained at m/z 897.41 (2+), for SPDTNYLFMGDYVDR (amino acids 75–89) of PP2A catalytic subunit (or isoforms). C (right), sequence coverage of mouse PP2Aα (Uniprot ID: PP2AA_mouse) with peptides identified by MS/MS in the DAT complex. E, the MS/MS spectrum representing VLMASGPYNLHR (amino acids 486–498) of co-purified PP2A 65-kDa regulatory subunit A isoform with DAT.
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**Table 1**

| Protein name | Peptide sequence | Residues | Mascot ions score | KO | WT |
|--------------|-----------------|----------|-------------------|----|----|
| PP1A_MOUSE | EIFQPFLLELPILIEEL | 111-121 | 44-60 | 44 | 44 |
| PP1B_MOUSE | IKYPENFFLLR | 332-342 | 44-60 | 32 | 44 |
| PP2AA_MOUSE | SPDTNYLFMGDYVDR | 75-89 | 35-51 | 35 | 35 |
| PP2AB_MOUSE | LQEVPHEGPM ox | 186-206 | 50-75 | 50 | 50 |

The ions score for an MS/MS match is based on the calculated probability, \( p \), that the observed match between the experimental data and the database sequence is a random event. The reported score is \( 10^{\log(p)} \).

**Enhanced DA uptake in the dephosphomimetic T48A mutant**

To investigate the functional effect of Thr-48 dephosphorylation, we examined DA uptake mediated by the dephosphomimetic T48A mutant. The construct was transiently expressed in LLC-PK1 cells and showed plasma membrane expression comparable with that of the WT DAT by surface biotinylation analysis \( (p > 0.05; \text{Fig. 5A}) \). To validate that our cell surface analysis was within the linear range of detection and not saturated, we determined a NeutrAvidin binding curve using increasing amounts of cell lysate (25–200 \( \mu \)g of protein) from our cell surface biotinylation assay (Fig. S4). We found that our assessment of DAT surface levels was within the linear range of our assay. In saturation analyses performed in parallel and normalized to DAT surface levels, there was no difference in \( K_m \) (WT, sites, and the Mascot database search also indicated probabilities for phosphorylation at Ser-45, Thr-46, or Thr-48.

To determine the identity of the phosphorylation site, we mutated Thr-48 to Ala to prevent phosphorylation and expressed the T48A hDAT mutant in heterologous cells under OA treatment. MS analysis showed no phosphorylation of this peptide in either basal or OA-stimulated conditions (Tables S3 and S4). Therefore, we concluded that Thr-48 is a PP1/2A-responsive phosphorylation site in hDAT (Fig. 3C). As an additional control, we analyzed the phosphorylation status of this peptide derived from rDAT, which contains isoleucine rather than threonine at position 48 (Fig. S3), although several other Ser and Thr residues are present in this sequence. The resulting tryptic peptide also showed no phosphorylation, providing further support for the specific regulation of Thr-48 phosphorylation status by PP1/2A inhibition.

**Inhibition of PP1/2A increases phosphorylation of DAT at Thr-48**

OA-inhibited hDAT dephosphorylation was indicated by the increased number of phosphopeptides isolated (Table 2). To determine the relative increase in Thr-48 phosphorylation mediated by OA, we applied “stable isotope labeling with amino acids in cell culture” (SILAC), which distinguishes peptides derived from different conditions by the stable isotope-introduced mass tag \( (24, 25) \). The LLC-PK1 cells stably expressing hDAT were cultured in media containing either Lys-0/Arg-0 (“Light”) or Lys-6/Arg-10 (“Heavy”) in the presence or absence of OA (0.5 \( \mu \)M for 1 h). Because the doubly charged (2\( + \))-peptide \( ^{36}\text{EQNGVQLTSTLTNPR}^{52} \) contains one Arg and Lys-6/Arg-10 labeling would increase its mass by 5 Da \( (m/z = 10 \text{ Da}/2^+) \), we could compare the identified phosphopeptide peak with the peak that is located at +5 or −5 Da according to the labeling of the identified phosphopeptides. SILAC certainly indicated increased levels of phosphopeptide for Thr-48 in the presence of OA (Fig. 4A, left and middle). When treated with OA in both Light and Heavy culture media, MS identified comparable phosphopeptides for Thr-48 in the Light- and Heavy-labeled DAT at a different \( m/z \) of 5 Da (Fig. 4A, right). However, the increase of tryptic peptide peaks by OA was not observed in the corresponding unphosphorylated peptide \( ^{36}\text{EQNGVQLTSTLTNPR}^{51} \) (Fig. 4B) and the hDAT \(^{591}\text{LAYIAPKDR}^{601} \) peptide (Fig. 4C).
1.7 ± 0.6 μM; T48A, 1.7 ± 0.5 μM; p > 0.05), but $V_{\text{max}}$ was significantly increased in T48A-DAT (WT, 558 ± 72 pmol/min/mg; T48A, 847 ± 142 pmol/min/mg; p < 0.01, n = 5) (Fig. 5B). Enhanced activity in the dephosphomimetic mutant suggests that uptake is reduced by phosphorylation of the site and returns to basal levels by PP1/2A-mediated dephosphorylation. 

The lower $V_{\text{max}}$ in the WT protein than in the T48A form is presumably driven by basal phosphorylation of this site. 

Other phosphorylation sites, including Ser-7, are regulated by PKC and PP1/2A (17) and could contribute to outcomes in WT and T48A-DATs. Hence, in order to determine whether Thr-48 mediates transport regulation by PP1/2A or PKC, we performed DA uptake analyses in the absence or presence of OA, the PKC activator phorbol 12-myristate 13-acetate (PMA), or both compounds together. DA uptake activity in WT and T48A-DATs was significantly decreased in all three conditions (Fig. 5C). In the WT protein, levels of down-regulation induced by OA and PMA (17 ± 11% and 29 ± 8%, respectively) were similar to that seen in previous studies (12, 18, 20), and there was no additivity with the combined PMA/OA treatment (Fig. 5C, 30 ± 10%), suggesting a shared mechanism of action. In addition, there was no difference in down-regulation responses between WT and T48A-DAT for each treatment (p > 0.05; Fig. 5C), suggesting that the inability of Thr-48 to undergo phosphorylation had no influence on either PKC or PP1/2A responses. 

We further examined amphetamine-stimulated DA efflux by the dephosphomimetic T48A mutant. LLC-PK1 cells transiently expressing WT or T48A-DAT were preloaded with [3H]DA and then treated with amphetamine (3 μM) to stimulate DA efflux. In superfusion experiments, no significant differences of efflux were observed at each fraction between WT and T48A-DAT (p > 0.05) (Fig. 5D).

Increased palmitoylation in the dephosphomimetic T48A-DAT

Previously, we reported a reciprocal mechanism between DAT phosphorylation at Ser-7 and palmitoylation at Cys-580 that acted in concert to set levels of DA uptake capacity (18). Therefore, we examined whether DAT palmitoylation could also be affected by Thr-48 phosphorylation status. We transiently expressed WT and T48A-DAT in LLC-PK1 cells in the absence and presence of OA and assessed transporter palmitoylation levels by an acyl-biotinyl exchange (ABE) assay. In this procedure, free thiols in the protein are blocked using methyl
Figure 3. Identification of PP1/2A-responsive DAT phosphorylation at Thr-48 by LC-MS/MS. A, representative colloidal blue–stained SDS-polyacrylamide gel image with proteins immunopurified by an anti-DAT antibody (Ab) from LLC-PK₁ cells heterologously expressing hDAT in the absence (−) and presence (+) of OA. Arrowheads and square bracket, bands and region containing DAT protein identified by LC-MS/MS. The numbers refer to the positions of molecular weight markers. B, sequence coverage of human DAT with identified peptides (boldface type) by MS/MS from transfected LLC-PK₁ cells. The putative transmembrane segments (S₁–S₁₂) of hDAT are underlined. C, MS/MS-identified hDAT phosphorylation site at Thr-48 (pT₄₈) from transfected LLC-PK₁ cells upon OA treatment. Shown is the MS/MS spectrum from the precursor ion at m/z 913.40 (2⁺) assigned to hDAT EQNGVQLTSSTLpTNPR (amino acids 36–51) with phosphorylation at Thr-48, which was verified by β-eliminated y-ions and neutral loss of H₃PO₄ from the precursor. The phosphosite localization probabilities determined by the Mascot delta score (38) are shown in parentheses. pT, phosphorylated threonine; deN, deamidated asparagine.
methanethiosulfonate (MMTS), endogenous palmitate groups are cleaved with hydroxylamine (NH2OH), and the liberated free SH groups are tagged using sulfhydryl-reactive (N-(6-(bi-

Table 2

| Peptide sequence | No treatment | Okadaic acid |
|------------------|--------------|--------------|
| hDAT WT (LLC-PK1) | ^{28}EVELILKEEQNGVQLTSSLTLPNPR | 0:19 \(^{a}\) | 2:11 (0.25 \(\mu\)M, 2 h) |
|                  | ^{28}EQNGVQLTSSLTLPNPR | 0:28 | 11:29 (0.5 \(\mu\)M, 2 h) |

\(^{a}\) The values in boldface type denote the sum of the two different recovered phosphopeptide sequences displayed in the second column versus the sum of the two corresponding unphosphorylated peptides recovered in three individual experiments using three different concentrations of OA.

Figure 4. Relative quantification of OA-induced DAT phosphorylation at Thr-48 by SILAC. A, the doubly charged (2\(^{+}\)) phosphopeptide (EQNGVQLTSSLTLPNPR) presenting phosphorylated hDAT at Thr-48 (pT48) was abundant in the samples from hDAT-LLCPK1 cells upon OA treatment (0.5 \(\mu\)M for 1 h) but limited in the untreated samples. hDAT-LLCPK1 cells were cultured in metabolically different culture media with either Light medium containing normal lysine (12C6, 14N2) and arginine (12C6, 14N4) or Heavy medium containing isotopic lysine (13C6, 14N2) and arginine (13C6, 15N4) in the absence and presence of OA, which provides a difference of 5 Da between unlabeled and labeled doubly charged EQNGVQLTSSLTLPNR peptides. B, the corresponding unphosphorylated peptide (EQNGVQLTSSLTLP) was not affected by OA treatment in both Light and Heavy culture conditions. C, a representative hDAT peptide (LAYAIAPEKDR) demonstrates the overall ratio of hDAT and no effect of OA in hDAT expression between two different culture media. SILAC labeling at one Lys and one Arg provided the 8-Da (\(m/z\) 6 + 10 Da/2\(^{+}\)) difference of the labeling.

Discussion

The results show that in basal conditions, palmitoylation of T48A-DAT was significantly enhanced relative to the WT protein (54 \(\pm\) 14%, \(p < 0.05\), versus WT, \(n = 9\)) (Fig. 6), supporting a role for Thr-48 phosphorylation in regulating this modification. However, in contrast to our findings that palmitoylation was reduced by PMA (18), we found no effect of OA on this modification (Fig. 6), indicating mechanistic differences between functions regulated by PKC versus PP1/2A.
KO mice that eliminated nonspecific and false-positive proteins. PP2A is characterized as having a heterodimeric core enzyme and a heterotrimeric holoenzyme, where the former is composed of scaffold and catalytic subunits and interacts with one of four regulatory subunits to form a holoenzyme (26). Our MS analysis not only confirmed the association with the previously immunoblotting-detected PP2A catalytic subunit (22) but also identified a scaffold subunit, PP2A PR65α/H9251, and a regulatory subunit, B55, in the immunopurified DAT protein complex from mouse striata (Table 1). MS also presented PP1 association with DAT, which we previously showed is a major contributor to rat striatal DAT dephosphorylation (8, 21).

Previous MS-based proteomics studies have identified DAT-interacting proteins including PKCs and CaMKIIs, but not PP2A, in mouse and rat brains (7, 27, 28). In other SLC sodium carrier transporters, MS approaches have shown the interaction of PP2A with norepinephrine transporter and organic cation/carnitine transporter OCTN2 (29, 30), but PP2A-responsive dephosphorylation sites were not determined in these transporter proteins. Although MS is a very powerful and accurate tool to identify phosphorylation sites (31), some additional procedures are required for MS sample preparation to identify protein kinase/phosphatase-specific sites by MS. For this purpose, a target protein kinase/protein phosphatase needs to be identified, and experimental methods are determined accordingly. Because we detected a few phosphopeptides for DAT purified from LLC-PK1 cells under basal culture conditions, we presumed that the association of PP1/2A maintains the dephosphorylated state of DAT. Therefore, we pharmacologically inhibited PP1/2A using OA in LLC-PK1 cells expressing hDAT to prevent dephosphorylation. Then MS with stable isotope labeling manifested the increased phosphorylation at Thr-48 in hDAT, indicating PP1/2A-directed dephosphorylation at this site in hDAT (Fig. 4).

We observed a significant effect of the dephosphomimetic T48A mutant on DA uptake (Fig. 5B) but not on amphetamine-stimulated DA efflux (Fig. 5D), indicating the specific role of dephosphorylation at Thr-48 in the forward transport mechanism. The enhanced DAT-mediated DA uptake rate in the T48A mutant strongly supports a distinctive role of this PP1/
cells in the presence and absence of OA by chymotryptic in-gel digestion (48TNPRQSPVEAQDRETW63; data not shown). However, we could not determine its phosphorylation status upon OA treatment by SILAC because MS did not identify phosphorylation at Ser-53 in DAT by means of the OA-induced increase of $^{32}$PO$_4$ incorporation in rDAT (12). Therefore, OA-induced dephosphorylation at Thr-48 may also be related to Thr-48 phosphorylation.

In our previous studies, the OA treatment of rat striatal slices caused increased phosphorylation primarily on distal N-terminal serines (21, 32) and Thr-53 (equivalent to hDAT Ser-53) (16). In addition, our previous MS study pinpointed phosphorylation at hDAT Ser-7 in HEK293 cells upon OA treatment (17). In the present study, however, we could not identify OA-responsive phosphorylation at Ser-7 in LLC-PK$_1$ cells, probably due to different profiles of protein kinases and protein phosphatases between experimental cell lines. This also may be related to different PP response to OA and against OA-induced cell toxicity between the two cell lines (data not shown). We also identified phosphorylation at hDAT Ser-53 in LLC-PK$_1$.
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17), this study demonstrated the significance of the N terminus in DAT regulation by not only phosphorylation but also dephosphorylation. As we have found up- or down-regulation of DA activity in DAT dephosphomimetic mutants of potential PP1/2A-directed sites, a comprehensive study needs to be performed for phosphorylation/dephosphorylation effects on DAT functions, and for this purpose, identification of specific protein kinases/phosphatases as well as their specific sites of action is a prerequisite. This is the first time a protein phosphatase–directed site in SLC6 neurotransmitter transporters with functional involvement has been identified, representing progress toward establishing a mechanism for the modulation of SLC6 transporters by coordinated phosphorylation/dephosphorylation and, in the case of DAT, additionally palmitoylation.

Experimental procedures

Animals

Functional DAT knockout mice were generated by crossing heterozygous mice expressing the CRE recombinase under the DAT promoter (34). The mice have a C57BL/6J background, and the genotype was confirmed twice for each mouse. The mice were bred at the Core Unit of Biomedical Research, Division of Laboratory Animal Science and Genetics, Medical University of Vienna (Vienna, Austria). All animal care procedures were in accordance with the ARRIVE guidelines, the Austrian animal protection law, and the Austrian animal experiment by-laws that implement European law (directive 2010/63/EU) into Austrian law; they were approved by the local animal ethics committee at the Medical University of Vienna and authorized by the Austrian Federal Ministry of Science and Research (GZ: BMWFVW-66.009/0016-WF/V/3b/2015).

Preparation of mouse striatal synaptosomes

Mice were dispatched by cervical dislocation, and subsequently the brain was removed immediately to dissect the dorsal striata. The right and left striata of one mouse were homogenized in ice-cold buffer containing 0.32 M sucrose, 5 mM sodium phosphate (pH 7.4), 0.1 mM sodium fluoride, and protease inhibitors (Complete®, Roche Applied Science). The suspension was centrifuged at 800 × g for 10 min, and subsequently the supernatant was centrifuged at 38,000 × g for 90 min. The resulting crude synaptosomal membrane pellet was suspended in the homogenization buffer.

cDNA constructs

The hDAT pcDNA3.0 vector was created by excising the hDAT coding region from a His6-HA-hDAT vector (35) using endonucleases Acc65I and EcoRI and ligating the excised fragment into pcDNA3.0 using these same restriction endonuclease sites. The hDAT pEYFP vector was created by excising the hDAT coding region from the hDAT pcDNA3.0 vector using endonucleases Acc65I and Xbal and ligating the excised fragment into pEYFP-C1 using these same restriction endonuclease sites. Mutagenesis to create DAT T48A was performed using hDAT pcDNA 3.0 as the template with the Stratagene QuickChange® kit, and codon substitution was verified by sequencing (Eurofins MWG, Louisville, KY). For production of stable transformants, LLC-PK1 cells (ATCC, Manassas, VA) were transfected (X-tremeGENE HP, Roche Applied Science) and after 48 h maintained under selection with 600 µg/ml G418 (8). HEK293 and tsA201 cells were transiently transfected with the indicated vector using the Turbofect (Thermo Scientific) according to the manufacturer’s protocol.

Cell culture and transient transfection

Parental LLC-PK1 cells and LLC-PK1 cells stably expressing hDAT (hDAT-LLCPK1), T48A hDAT (T48A-LLCPK1), and rat DAT were maintained in α-minimum essential medium supplemented with 5% FBS, 2 mM L-glutamine, 200 µg/ml G418, 25 IU/ml penicillin, and 25 µg/ml streptomycin. Parental HEK293 cells and HEK293 cells stably expressing YFP-tagged human DAT (YhDAT-HEK293) as well as tsA201 cells were cultured in Dulbecco's modified Eagle's medium with 10% FBS, 4.5 mg/ml glucose, 25 IU/ml penicillin, and 25 µg/ml streptomycin. Additionally, 250 µg/ml G418 was added to the medium for stable cell lines to maintain DAT protein expression. Cells were maintained in a humidified atmosphere of 5% CO2 at 37 °C.

IP and immunoblotting (IB)

Mouse striata membranes and cell lysates from heterologous cells expressing DAT proteins were prepared as described previously (16). For IP, tissue and cell lysates were solubilized in 0.5 and 1% Triton X-100 lysis buffer, respectively, and incubated overnight with a goat anti-DAT (sc-1433, Santa Cruz Biotechnology, Inc.) or rabbit anti-GFP polyclonal antibodies (A6455, Life Technologies, Inc.). Antigen–antibody complexes were then incubated with protein G beads (GE Healthcare) for 4 h at 4 °C. After six washes, bound proteins were eluted in SDS sample buffer at 95 °C for 3 min. Eluted proteins were size-fractionated on SDS-polyacrylamide gels and visualized by a colloidal blue staining (Invitrogen).

For IB, proteins were separated on SDS-polyacrylamide gels and transferred to polyvinylidene fluoride membranes (Waters, Milford, MA), which were then blocked and immunostained with anti-DAT (sc-74580, Santa Cruz) or PP2B-A (sc-17808, Santa Cruz) antibodies. For IB, proteins were separated on SDS-polyacrylamide gels and transferred to polyvinylidene fluoride membranes (Waters, Milford, MA), which were then blocked and immunostained with anti-DAT (sc-74580, Santa Cruz) or PP2B-A (sc-17808, Santa Cruz) antibodies.

In-gel digestion

The protein bands were directly excised from SDS-polyacrylamide gels, destained with 50% acetonitrile in 50 mM ammonium bicarbonate, and dried in a speed-vacuum concentrator. After reduction and alkylation of Cys, gel pieces were washed and dehydrated. Dried gel pieces were swollen with 25 mM ammonium bicarbonate (pH 8.0) containing 10 ng/µl trypsin (Promega) or chymotrypsin (Promega) and incubated at 37 °C for 2–18 h. Digested peptides were extracted with 50% acetonitrile in 5% formic acid (FA) and concentrated in a speed-vacuum concentrator (16).

LC-MS/MS

An electrospray ionization (ESI)- ion trap (IT) mass spectrometer (HCT and Amazon speed ETD, Bruker) and an ESI-quadrupole-TOF (QTOF; Compact, Bruker) coupled with an Ultimate 3000 nano-HPLC system (Dionex) were used for LC-
MS/MS data acquisition. A PepMap100 C-18 trap column (300 μm × 5 mm) and PepMap100 C-18 analytic column (75 μm × 150 mm) were used for reverse-phase chromatographic separation with a flow rate of 300 nl/min. The two buffers used for the reverse-phase chromatography were 0.1% FA and 80% acetonitrile in 0.08% FA with variable gradient conditions. Eluted peptides were then directly sprayed into the mass spectrometer, and the MS/MS spectra were interpreted with the Mascot search engine (version 2.4.1, Matrix Science, London, UK) against Swissprot database (556,196 sequences, released in November 2017), and the taxonomy was restricted to Homo sapiens (human; 20,244 sequences) for human DAT from cell lines as well as to Rodentia (rodents; 26,656 sequences) for mouse proteins and rat DAT purified from the mouse striata and heterologous cells, respectively.

The search parameters were used with a mass tolerance of 20 ppm (QTOF) or 0.5 Da (IT) and an MS/MS tolerance of 0.1 Da (QTOF) or 0.5 Da (IT). Carbamidomethylation on Cys, oxidation on Met, phosphorylation on Ser/Thr, and deamidation on Asn/Gln were allowed with two missing cleavage sites. The Mascot cut-off score was set to 10, and each filtered MS/MS spectrum exhibiting possible phosphorylation was carefully examined based on the existence of a neutral loss of H3PO4 (97.977 Da) (16, 36).

**SILAC**

hDAT-LLCPK₁ cells were grown for at least five cell divisions in either normal lysine (13C6, 14N2; “Lys-0”) and arginine (13C6, 14N2; “Arg-0”) media or in isotopic Lys (13C6, 14N2; “Lys-6”) and Arg (13C6, 15N2; “Arg-10”) labeling media in the absence and presence of okadaic acid. After labeling, cells were solubilized in lysis buffer containing 1% Triton X-100, 20 mM Tris-Cl (pH 8.0), 150 mM NaCl, 1 mM EDTA, 1 mM sodium orthovanadate, 5 mM NaF, 5 mM sodium pyrophosphate, and a protease inhibitor mixture. The cell lysates were mixed at a 1:1 ratio based on protein quantification for IP with an anti-GFP antibody. After size fractionation by SDS-PAGE, hDAT proteins were in-gel trypsin-digested and analyzed by ESI-QTOF as described above. After Mascot database searches, relative spectral intensities were examined using Compass DataAnalysis (version 4.2, Bruker) software.

**DA uptake and cell surface biotinylation assays**

Cells were grown in 24-well plates to ~80% confluence and then transected (XtremeGene, Sigma) with WT hDAT or T48A hDAT cDNA (pcDNA3.0). After 36 h post-transfection, cells were rinsed twice with 0.5 ml of 37 °C Krebs-Ringer/HEPES (KRH) buffer (25 mM HEPES, 125 mM NaCl, 4.8 mM KCl, 1.2 mM KH₂PO₄, 1.3 mM CaCl₂, 1.2 mM MgSO₄, 5.6 mM glucose, pH 7.4) followed by incubation at 37 °C for 30 min with vehicle (MeSO₄, 1 μM OA, 400 nM PMA, or both OA and PMA followed by a [3H]DA transport assay. Uptake was performed in triplicate for 8 min and initiated by the addition of 10 nM [3H]DA plus 3 μM unlabeled DA. For saturation analysis, unlabeled DA concentration varied from 0.3 to 20 μM, with nonspecific uptake determined in the presence of 100 μM (−)-cocaine. Cells were rapidly washed two times with ice-cold KRH buffer and solubilized in 1% Triton X-100, and radioactivity contained in lysates was assessed by liquid scintillation counting.

For determination of cell surface expression, cells were plated, transfected, and treated in parallel with those analyzed for DA transport. These cells were washed three times with ice-cold PBS/Ca-Mg (138 mM NaCl, 2.7 mM KCl, 1.5 mM KH₂PO₄, 9.6 mM Na₂HPO₄, 1 mM MgCl₂, 0.1 mM CaCl₂, pH 7.4) and incubated twice with a 0.5 mg/ml concentration of the membrane-impermeable reagent sulfo-NHS-SS-biotin for 25 min at 4 °C in PBS/Ca-Mg. The biotinylating reagent was removed, and the reaction was quenched by two sequential incubations with 100 mM glycine in PBS/Ca-Mg for 20 min at 4 °C. Cells were washed with PBS/Ca-Mg and lysed at 4 °C with 250 μl of radioimmunoprecipitation assay buffer (10 mM sodium phosphate, pH 7.3, 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, 1% sodium deoxycholate, 0.1% SDS) containing protease inhibitor. Total cell lysates (50 μg of protein) were incubated with NeutrAvidin beads overnight at 4 °C. The beads were washed three times with radioimmunoprecipitation assay buffer, and bound proteins were eluted with 40 μl of Laemmli buffer (62.5 mM Tris-HCl, pH 6.8, 20% glycerol, 2% SDS, 5% β-mercaptoethanol, and 0.01% bromophenol blue), followed by immunoblotting with a DAT-specific polyclonal antibody (C-20, Santa Cruz, Inc., sc-1433).

DA uptake values from the saturation analysis were normalized to DAT surface expression by dividing WT and T48A uptake values by relative surface abundance for each, where the WT surface abundance was set to 1 and the T48A was either less than or greater than 1, as determined by cell surface biotinylation with equal amounts of total protein for WT and T48A samples loaded on the high-capacity NeutraAvidin resin. V₅₀, Kₘ, and V₅₀/Kₘ values were determined by nonlinear regression analysis of the normalized saturation analysis uptake values.

**Superfusion experiments**

Substrate efflux assays were performed as described previously (16) with the following modification. The LLC-PK₁ cells transiently expressing WT or T48A-DAT were preincubated with 0.4 μM [3H]DA (55 Ci/mmol; American Radiolabeled Chemicals, St. Louis, MO) for 60 min at 37 °C in a final volume of 0.1 ml of KRH buffer/well and subsequently transferred into superfusion chambers. Immediately, superfusion was initiated with KRH buffer at 25 °C at a perfusion rate of 0.7 ml/min. After 45 min, a stable efflux of radioactivity was achieved, and the experiment was started with the collection of 2-min fractions. After three fractions, amphetamine (3 μM) was added to stimulate DA efflux. Finally, the cells were lysed with 1% SDS. Afterward, the amount of tritiated substrate present in each vial was determined by β-scintillation counting (PerkinElmer Life Sciences). Efflux of tritium was expressed as a fractional rate (i.e. the radioactivity released during a fraction was expressed as the percentage of the total radioactivity present in the cells at the beginning of that fraction).

**Analysis of palmitoylation by ABE**

Analysis of palmitoylation by ABE was performed as described previously (37). Briefly, cells were treated with vehicle or 1 μM OA for 30 min, followed by cell membrane prepa-
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Membrane pellets were solubilized in lysis buffer (50 mM HEPES, pH 7.0, 2% SDS, 1 mM EDTA) containing protease inhibitors and 25 mM N-ethylmaleimide to block free thiols. Lysates were incubated at room temperature for 1 h with mixing followed by acetone precipitation and resuspension in lysis buffer containing MMTS and incubation at room temperature overnight with end-over-end rotation. Excess MMTS was removed by three sequential acetone precipitations followed by resuspension of the final pellet in 75 mM Tris-HCl, pH 7.4, as control or 0.7 M NH2OH, pH 7.4, to cleave the thioester bonds. NH2OH was removed by three sequential acetone precipitations followed by resuspension of the precipitated proteins in 240 μl of 4SB buffer. Samples were diluted with 900 μl of 50 mM Tris-HCl, pH 7.4, containing 0.1% SDS, and biotinylated proteins were extracted using NeutrAvidin resin. Proteins bound to the resin were eluted with sample buffer (60 mM Tris, 4% SDS, 50 mM Tris, 5 mM EDTA, pH 7.4). Each sample was divided into two equal portions that were treated for 2 h at room temperature with 50 mM Tris-HCl, pH 7.4, as control or 0.4 mM HPDP biotin and incubated at room temperature for 1 h with mixing followed by acetone precipitation as described previously (18). Membrane pellets were solubilized in lysis buffer (50 mM HEPES, pH 7.0, 2% SDS, 1 mM EDTA) containing protease inhibitors and 25 mM N-ethylmaleimide to block free thiols. Lysates were incubated at room temperature for 1 h with mixing followed by acetone precipitation and resuspension in lysis buffer containing MMTS and incubation at room temperature overnight with end-over-end rotation. Excess MMTS was removed by three sequential acetone precipitations, followed by resuspension of the precipitated proteins in 300 μl of a buffer containing 4% (w/v) SDS (4SB: 4% SDS, 50 mM Tris, 5 mM EDTA, pH 7.4). Each sample was divided into two equal portions that were treated for 2 h at room temperature with 50 mM Tris-HCl, pH 7.4, as control or 0.7 M NH2OH, pH 7.4, to cleave the thioester bonds. NH2OH was removed by three sequential acetone precipitations followed by resuspension of the precipitated proteins in 240 μl of 4SB buffer. Samples were diluted with 900 μl of 50 mM Tris-HCl, pH 7.4, containing 0.4 mM HPDP biotin and incubated at room temperature for 1 h with end-over-end mixing. Unreacted HPDP biotin was removed by three sequential acetone precipitations followed by resuspension of the final pellet in 75 μl of lysis buffer without MMTS. Samples were diluted with 50 mM Tris-HCl, pH 7.4, to contain 0.1% SDS, and biotinylated proteins were extracted using NeutrAvidin resin. Proteins bound to the resin were eluted with sample buffer (60 mM Tris, pH 6.8, 2% SDS, 10% glycerol) containing 100 mM DTT plus 3% β-mercaptoethanol and subjected to SDS-PAGE. DAT was identified by immunoblotting using polyclonal antibody (C-20, sc-1433, Santa Cruz Biotechnology).

Statistics

Palmitoylated and cell surface DAT band intensities were quantified by densitometry using Quantity One (Bio-Rad) software. The palmitoylated DAT signal was normalized to the cytosol and promotes reverse transport. J. Neurosci. 15, 4102–4108 CrossRef Medline

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**Regulation of dopamine transporter by protein phosphatases**