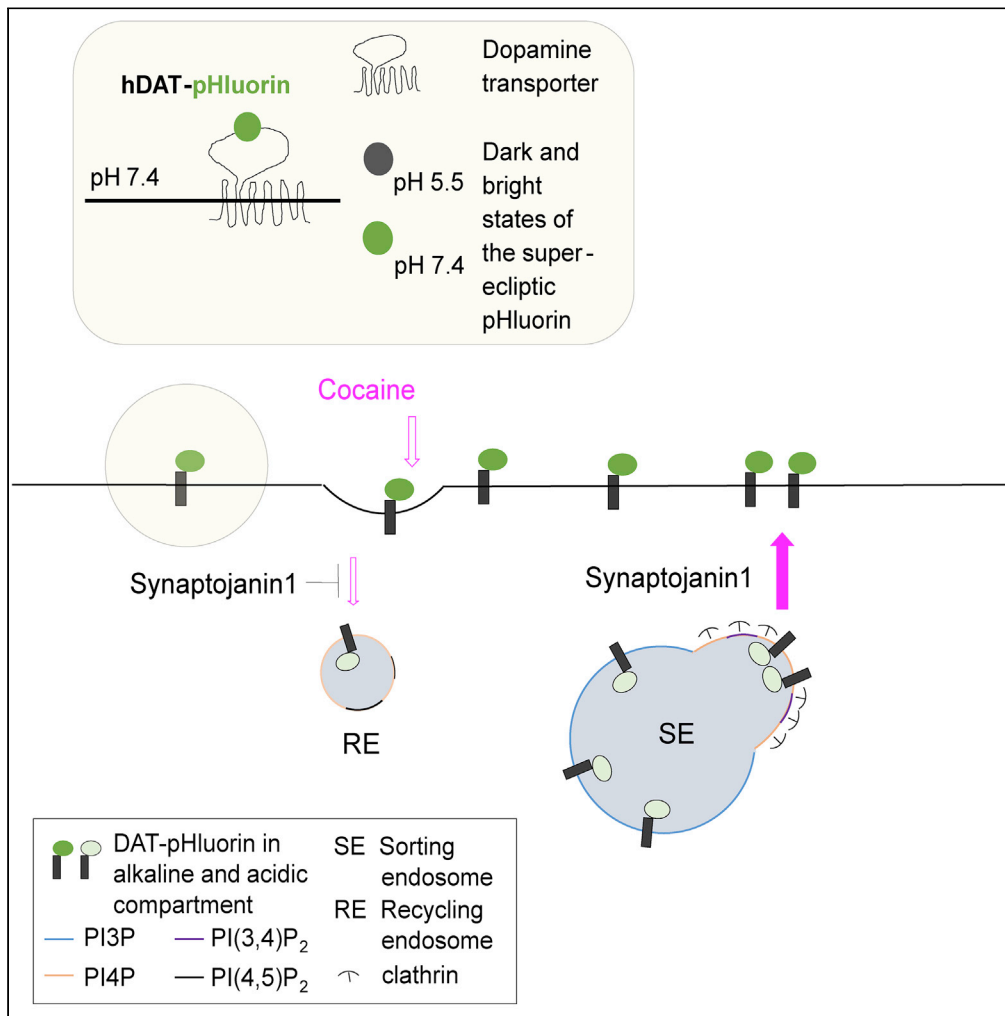


Article

Cocaine-regulated trafficking of dopamine transporters in cultured neurons revealed by a pH sensitive reporter



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Highlights

DAT-pHluorin is engineered for studying dopamine transporter trafficking

DAT is predominantly localized to the plasma membrane in the axons

Cocaine induces a transient internalization of DAT followed by reinsertion

Synaptojanin1 regulates the cocaine-induced DAT trafficking

Saenz et al., iScience 26, 105782
January 20, 2023 © 2022
Rutgers Robert Wood Johnson
Medical School.
<https://doi.org/10.1016/j.isci.2022.105782>



Article

Cocaine-regulated trafficking of dopamine transporters in cultured neurons revealed by a pH sensitive reporter

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SUMMARY

Cocaine acts by inhibiting plasma membrane dopamine transporter (DAT) function and altering its surface expression. The precise manner and mechanism by which cocaine regulates DAT trafficking, especially at neuronal processes, are poorly understood. In this study, we engineered and validated the use of DAT-pHluorin for studying DAT localization and its dynamic trafficking at neuronal processes of cultured mouse midbrain neurons. We demonstrate that unlike neuronal soma and dendrites, which contain a majority of the DATs in weakly acidic intracellular compartments, axonal DATs at both shafts and boutons are primarily (75%) localized to the plasma membrane, whereas large varicosities contain abundant intracellular DAT within acidic intracellular structures. We also demonstrate that cocaine exposure leads to a Synaptojanin1-sensitive DAT internalization process followed by membrane reinsertion that lasts for days. Thus, our study reveals the previously unknown dynamics and molecular regulation for cocaine-regulated DAT trafficking in neuronal processes.

INTRODUCTION

Dopamine (DA) signaling regulates a multitude of brain functions including motor coordination and reward processing. DA reuptake from the extracellular space into the cytosol by the plasma membrane dopamine transporter (DAT) is the primary mechanism for terminating DA signaling in the brain. Lack of DAT expression results in increased extracellular DA and hyperactive locomotion in the knockout mice.¹ Importantly, many psychoactive substances, such as cocaine and amphetamine (AMPH), directly or indirectly target plasma membrane DAT. In addition to acting to increase extracellular levels of DA, these substances are also known to impact DAT trafficking and plasma membrane expression.^{2–6} The potency of these psychoactive substances in altering DA signaling is highly dependent on the amount of surface DAT expression, however, our understanding on the dynamic trafficking and molecular regulation of DAT, especially in neuronal processes, is poorly understood.

The lack of understanding largely reflects a lack of molecular tools to examine the dynamic trafficking of DAT in small neuronal structures like the axons and presynaptic terminals. Current understanding of DAT trafficking is largely based on studies of heterologous cells and isolated synaptosomes. The most commonly used research strategies to examine the surface DAT include (1) biotinylation assays, which has poor temporal resolution; (2) radioactive binding/uptake assays, which entangles DAT localization with function; and (3) imaging studies using fluorescent proteins-tagged DATs or fluorescent DAT ligands,^{7,8} which exhibit limited spatial resolution for neuronal processes. Complex regulatory mechanisms have been proposed for DAT in different cell systems using different methods. For example, the constitutive DAT recycling can be regulated through a protein kinase C (PKC)-dependent⁹ or independent manner.¹⁰ AMPH was shown in most studies to induce DAT internalization via RhoA and TAAR1 signaling^{11–13}; however, AMPH-induced delivery of DAT to the plasma membrane has also been reported.¹⁴ Cocaine was found in various studies to block DAT internalization and to increase its surface expression.^{15–20} Of interest, however, the mechanisms underlying cocaine-regulated DAT trafficking remain elusive, likely because of its inhibitory role on DAT function that hinders the use of many conventional methods. An earlier study by Sorkin et al. further suggested that neuronal DAT trafficking might be regulated by different mechanisms than what we learned from heterologous cells.²¹

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<https://doi.org/10.1016/j.isci.2022.105782>



To better reveal and understand DAT trafficking in neurons, we sought to develop a genetically encoded optical reporter. The pH sensitive “pHluorin” is a variant of green fluorescent proteins (GFP) that fluoresces on deprotonation.²² pHluorin-based assays are best known for analyzing synaptic vesicle recycling when pHluorin is tagged to synaptic vesicle proteins, such as vesicle associated membrane protein 2/VAMP2 (synaptopHluorin),^{22,23} vesicular glutamate transporter 1/VGLut1,^{24,25} and vesicular monoamine transporter 2/vMAT2.^{26,27} In addition, pHluorin has also been successfully engineered into plasma membrane cargo proteins, such as AMPA receptors,^{28,29} transferrin receptors,³⁰ mu-opioid receptors^{31,32} and glucose transporter 4 (GluT4)^{33,34} for analyzing their endosomal trafficking. We have now engineered pHluorin into the second exofacial loop of human DAT. By expressing DAT-pHluorin in cultured midbrain (MB) neurons, we turn DAT trafficking events at nano/micrometer scales to measurable changes in fluorescence using a conventional wide-field microscope. We demonstrate that 75% of axonal DAT is on the plasma membrane, with most intracellular DAT found in the varicosities. Moreover, we reveal the dynamic trafficking of DAT to the axonal surface in response to cocaine, which lasts for up to 4 days. The surface trafficking of DAT is regulated by synaptojanin1, a phosphoinositol phosphatase that was previously known to regulate synaptic vesicle recycling^{35–41} and autophagy.^{41–43} Thus, our study using the newly engineered DAT optical reporter reveals, for the first time, cocaine-regulated DAT trafficking in neuronal processes.

RESULTS

Engineering and characterization of a pH sensitive DAT reporter

DAT belongs to the family of Na⁺/Cl⁻ dependent neurotransmitter transporters⁴⁴ and contains 12 transmembrane domains. An extensive body of work has shown that DAT substrates, such as DA, AMPH or cocaine, can induce DAT trafficking.^{2,3} To engineer a DAT reporter suited for reporting physiologically relevant DAT endosomal trafficking, our strategy is to insert pHluorin while not, or minimally, disrupting its endogenous functions in substrate binding. Among the six extracellular loops of human DAT cDNA, the second loop was selected due to the optimal length of the loop that provides flexibility for accommodating a tag protein (Figure 1A). Flexible linkers were inserted on both ends of pHluorin to introduce additional flexibility for the proper folding of the reporter. pHluorin was inserted at multiple sites, T173/E174, P194/G195, G199/S201, G209/T210, R219/G220 outside the known functional sites of DAT.^{45–48} Although some insertion sites (T173/E174, G209/T210) resulted in poor expression of the transgene in the initial screen, little difference was observed for the remaining versions of DAT-pHluorin (DAT-pH) in their overall expression patterns (data not shown). We thus proceeded to further validate the DAT-pH at the P194/G195 insertion site, where an HA epitope was inserted in earlier studies.^{49,50}

When expressed in cultured MB neurons, we found that DAT-pH exhibited homogeneous cell body expression similar to endogenous DAT, and it was recognized by an anti-DAT antibody (Figure 1B). DAT-pH was targeted to express in distal axons in both DAergic and non-DAergic neurons in the culture (Figure 1C). In the past, fluorescent proteins such as YFP have been tagged to the N-terminus of DAT to track DAT trafficking in heterologous cells. We next examined the expression of DAT-pH in N2a cells by comparing to other tagged/fluorescent DAT reporters (Figure 1D). Unlike the N-terminally tagged DATs, which often exhibit supraphysiological levels of plasma membrane localization,^{46,48} DAT-pH expression was predominantly intracellular, but exhibited perfect colocalization when coexpressed with DAT-HA (Figure 1D). To determine if a fraction of DAT-pH was indeed targeted to express on the plasma membrane we performed the ³H-DA uptake assay. N2a cells transfected with DAT-pH exhibited a near 40-fold increase in ³H-DA uptake. Although it was only a fraction of the ³H-DA uptake in YFP-DAT expressing cells, DAT-pH in N2a cells performed similarly compared to the endogenous transporter activity measured in SH-SY5Y cells (Figure 1E). A 60-min cocaine exposure inhibited DA uptake for N2a cells expressing DAT-pH by 90%, similar to the cocaine’s effect on endogenous transporters in SH-SY5Y cells (Figure 1F). These data further suggested that pHluorin insertion did not block the essential substrate binding sites on DAT.

As a reporter gene that was designed to retain endogenous functions, an important concern is the potential unregulated expression that leads to altered intracellular signaling. An earlier study by Ryan et al. showed that only 1–2 copies of VGLut1-pHluorin (equivalent to a 10–20% overexpression) expression on the synaptic vesicle was sufficient to serve as a working sensor.⁵¹ Using immunofluorescence labeling of DAT, we found that DA neurons expressing DAT-pH exhibited similar levels of overall DAT expression compared to the untransfected neurons (Figures S1A and S1B). We then analyzed DAT surface expression in transfected and untransfected neurons using a fluorescent cocaine analogue, JHC1-64^{7,8,52} following

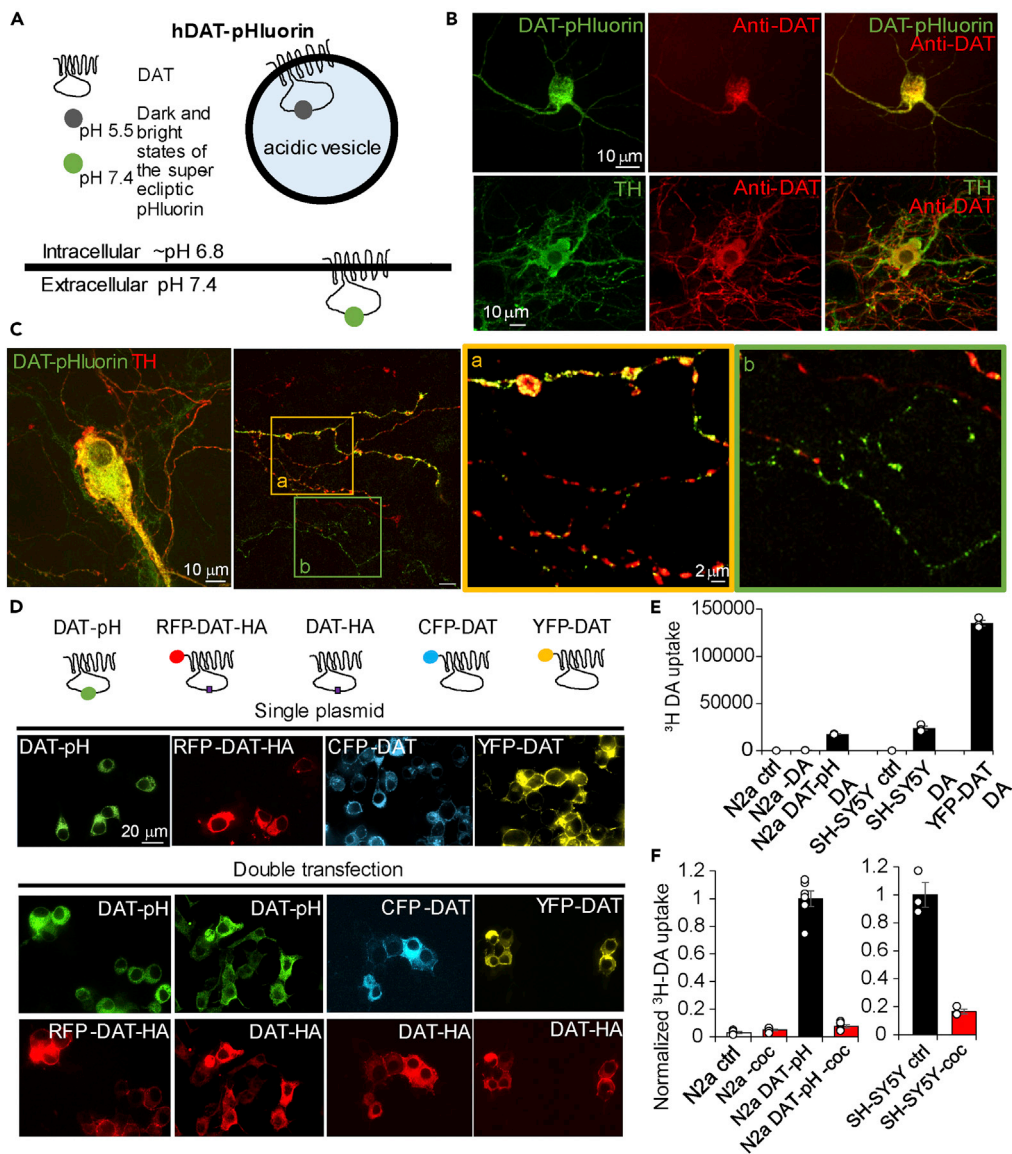


Figure 1. Engineering and validation of a pH sensitive DAT reporter

(A) Working hypothesis of the pH sensitive DAT reporter. The pH sensitive GFP variant, pHluorin, was inserted between the 194 Proline and 195 Glycine of the second exofacial loop of human DAT. This design allows DAT-pHluorin to fluoresce at its brightest when inserted onto the plasma membrane to face a neutral (pH 7.4) extracellular medium; and quenches when endocytosed to face an acidic endocytic compartment.

(B) Top, cultured MB neuron expressing DAT-pH and immunolabeled with anti-GFP (green) and anti-DAT antibodies (red). Bottom, cultured MB neuron immunolabeled with anti-TH (green) and anti-DAT (red) reveals the endogenous protein.

(C) Cultured MB dopamine neurons expressing DAT-pH and immunolabeled with anti-GFP (green) and anti-TH (red) antibodies. From left to right are the cell body, distal axon, and the enlarged view of the two cropped areas of the distal axons containing TH-positive (a) and TH-negative (b) axons.

(D) Comparison of the expression and co-expression patterns of DAT-pHluorin (DAT-pH) and other versions of tagged-DAT in N2a cells. Top, illustrations of the molecular structure and location of the tag for all DAT reporters. Representative images selected from N = 3 repeats.

(E) 3 H-DA uptake for N2a cells, N2a cells expressing DAT-pHluorin, SH-SY5Y cells and N2a cells expressing YFP-DAT. N = 3 replicates.

(F) Bar graphs of normalized 3 H-DA uptake for SH-SY5Y cells, N2a cells and N2a cells expressing DAT-pHluorin in response to cocaine (10 μ M). N = 6 replicates for N2a cells and N = 3 for SH-SY5Y cells. See also Figure S1.

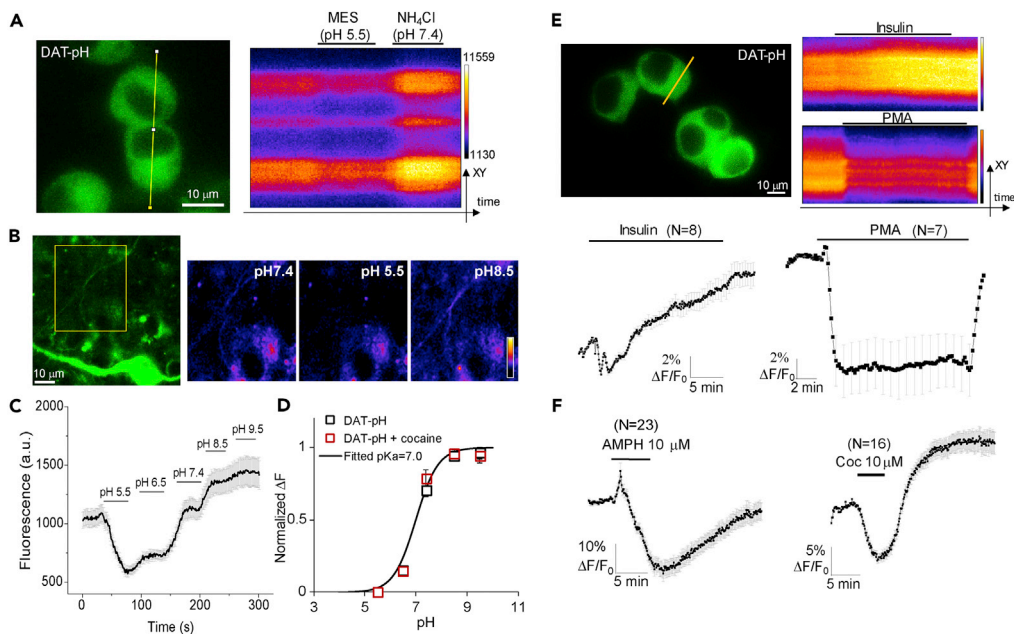


Figure 2. DAT-pHluorin exhibits reliable pH sensitivity and ligand regulated trafficking

(A) N2a cells expressing DAT-pH were perfused with a pH 5.5 MES solution followed by a pH 7.4 NH_4Cl solution. Right, kymographs were generated from the yellow line of the image on the left.

(B) Representative MB neuron expressing DAT-pH and axons (yellow boxed region) were selected for analyzing the pKa. Image intensities were adjusted to visualize the axon. Cell body fluorescence was not saturated at the camera. Right panels are the representative axon with color coding for DAT-pH immunofluorescence taken at different time points during a sequential perfusion of pH-fixed solutions shown in C.

(C) Representative DAT-pH signal when perfused with MES buffered saline at pH 5.5 and 5.6, NH_4Cl solution at 7.4, and Bicine buffered saline at pH 8.5 and 9.5. Error bars are from standard errors of 31 regions of interests along the axon of one neuron.

(D) Summary of normalized changes in DAT-pH responses in (C) from 4 axons at rest (black symbols) and 5 axons after the repeated cocaine treatment (red symbols). Black line represents the fitted change in fluorescence assuming $\text{pK}_a = 7.0$.

(E) N2a cells expressing DAT-pH and DAT-HA were perfused with Tyrode's solution containing insulin ($20 \mu\text{M}$) or PMA ($1 \mu\text{M}$) in separate imaging trials. Kymographs were generated from the yellow lines in the top left panel and the corresponding fluorescence signals of the cells were presented at the bottom. N = cell number, which is indicated next to the trace. Data is represented as mean \pm SEM. Similar responses were verified in three separate experiments.

(F) Averaged responses from N2a cells expressing DAT-pH and DAT-HA perfused with Tyrode's solution containing amphetamine/AMPH ($10 \mu\text{M}$, 5 min) or cocaine/coc ($10 \mu\text{M}$, 5 min). N = cell number. Data is represented as mean \pm SEM from two independent cultures. See also Figure S2.

a validation study conducted in N2a cells (Figures S1C and S1D). We showed that JHC1-64 can bind to DAT-pH in non-DAergic neurons (Figure S1E). Expressing DAT-pH did not lead to an increase in JHC dye binding either at the cell body or the axons (Figures S1F and S1G). The reduced fluorescence at the axons could be because of reduced binding affinity for DAT-pH.

We next examined other functional properties of DAT-pH as a viable reporter. Perfusion of a pH 5.5 MES (2-(N-morpholino)ethanesulfonic acid) solution quenched DAT-pH fluorescence whereas a pH 7.4 NH_4Cl solution revealed its peak fluorescence at the cell body (Figure 2A). To determine if pHluorin insertion in the functional DAT molecule maintains its original pH sensitivity (estimated $\text{pK}_a \sim 7.1$ in synaptopHluorin,²³ we measured the pKa of DAT-pH in the axons of MB neurons by sequentially perfusing a series of pH-fixed solutions following a previously reported protocol²³(Figures 2B and 2C). We found the best fitted pKa to be 7.0 for DAT-pH in MB neurons at rest, and that cocaine ($10 \mu\text{M}$ for 4 days) treatment did not alter the pH sensitivity of the reporter (Figure 2D). Moreover, we examined if DAT-pH exhibits agonist-regulated trafficking. DAT has been shown in earlier studies to internalize on PKC activation,^{46,53} and insulin was able to increase DAT surface expression via activation of the PI3K pathway.⁵⁴ Consistently, DAT-pH exhibited a time-locked insulin-induced increase in fluorescence and a PMA (phorbol myristate acetate)-induced decrease in fluorescence (Figure 2E), suggesting that the new reporter is responsive to PI3K and PKC

regulated trafficking. We then examined psychostimulants including AMPH and cocaine. A brief AMPH (10 μ M) perfusion led to a robust decrease of the fluorescence, which returned to baseline on washout (Figure 2Fleft). Of interest, cocaine (10 μ M) perfusion also induced a transient reduction of the DAT-pH signal, suggesting DAT internalization, which was followed by fast reversal and an increase of fluorescence within a few minutes (Figure 2Fright). Cocaine-induced DAT internalization in such timescale was not reported previously. To verify that the transient reduction of DAT-pH signal was indeed because of loss of surface DAT rather than a change in fluorescence because of conformational changes (such as oligomerization) of the reporter,¹⁹ we performed surface staining in non-permeabilized cells following a 5-min cocaine (10 μ M) incubation (Figures S2A and S2B). Our analysis suggested significant reduction of DAT surface fraction in cocaine treated cells (Figure S2B). Moreover, we did not observe major changes in the relative abundance of monomer, dimer and oligomers of the DAT-pH reporter following a 5-min or 10-min treatment of cocaine (10 μ M) (Figure S2C).

Taken together, our analyses using complementary methods suggest that DAT-pH is a viable reporter for analyzing DAT trafficking in neurons.

Majority of axonal DAT is on the plasma membrane

We next transfected DAT-pH in cultured mouse midbrain neurons. By bath perfusing MES (pH 5.5) and NH₄Cl (pH 7.4) solutions, we were able to extract an MES response that represents the surface pool of DAT and an NH₄Cl response that represents intracellular DAT (Figures 3A, 3B, and S3A). In the cell body and proximal dendrites of the neuron, both surface and intracellular DATs were observed (Figures 3A–3C). Of interest, the intracellular DAT (NH₄Cl response) in majority of the axons (identified by their substantially thinner appearance) was either absent or significantly smaller compared to that from the dendrites and cell body (Figures 3B and 3C). In parallel, the axonal MES response is substantially larger compared to the cell bodies and dendrites (Figures 3B and S3B). To exclude the possibility of quenching intracellular DAT by MES (Figure S3C), we performed surface labeling of pHluorin using an anti-GFP antibody without permeabilizing the cell (Figures S3C and S3D). Our analysis revealed the highest ratio for the surface to intracellular labeling of pHluorin in the axons and the lowest ratio at the cell body (Figure S3E), further suggesting a negligible contribution from the membrane permeability concerns of MES. To determine if DAT-pH yields different responses in DAergic versus non-DAergic neurons from the MB culture, we performed *post hoc* analysis based on tyrosine hydroxylase (TH) immunoreactivity of the neuron analyzed. Across four independent batches of the MB cultures, we did not observe significant differences in either the MES or NH₄Cl responses (Figure S4A).

The dynamic range of DAT-pH response is determined by the acidity of the intracellular structure that it is delivered to. To more accurately determine the surface DAT fraction in each neuronal compartment and the vesicular pH of the DAT-pH in that compartment, we used the Henderson-Hasselbalch equation with the fitted DAT-pH pK_a of 7.0 (Figures 2B–2D), following previously published protocols (Figure 3D, see STAR Methods).^{33,51} We report that neuronal cell body and dendrites contain $23.49 \pm 1.75\%$ (N = 37) and $35.64 \pm 2.68\%$ (N = 35) of surface DAT, respectively, whereas axonal DATs are primarily localized to the plasma membrane ($73.1 \pm 2.38\%$, N = 41) (Figure 3E). On average, DAT-containing vesicles are modestly but significantly more acidic in the axons (pH = 6.85 ± 0.014) compared to those in the cell body (pH = 6.94 ± 0.017) and dendrites (pH = 6.94 ± 0.022) (Figure 3F).

Large varicosities are distinct axonal sites that contain DAT in acidic intracellular organelles

The relative distribution of intracellular versus surface DAT at rest in different subcellular compartments of the neuron is reflective of constitutive DAT recycling.^{9,48} In all axonal structures examined, a fraction of these exhibited a relatively large NH₄Cl response (Figure S3B), suggesting that parts of the axons might be subjected to active DAT internalization. However, it remains unclear whether DAT is internalized at random positions along the axon, or if there are designated structures that support DAT turnover. To address this question, we examined three different structures in the axon: (1) Synaptic boutons, which are active sites for synaptic vesicle recycling and exhibit punctate staining for synapsin (Figure 4A); (2) axonal shafts, which exhibit diffuse synapsin staining; and (3) large varicosities that often exhibit DAT-pH outlining at rest (Figure 4B) but weak or diffuse synapsin staining. These varicosities, although not observed in all axons, are not uncommon for MB neurons. They have been previously reported to be potential axonal branching points⁵⁵ and are presently viewed to be highly relevant in neural injury and neurodegeneration.^{56–58} We found that only varicosities, but not axonal shafts or boutons exhibited a robust NH₄Cl

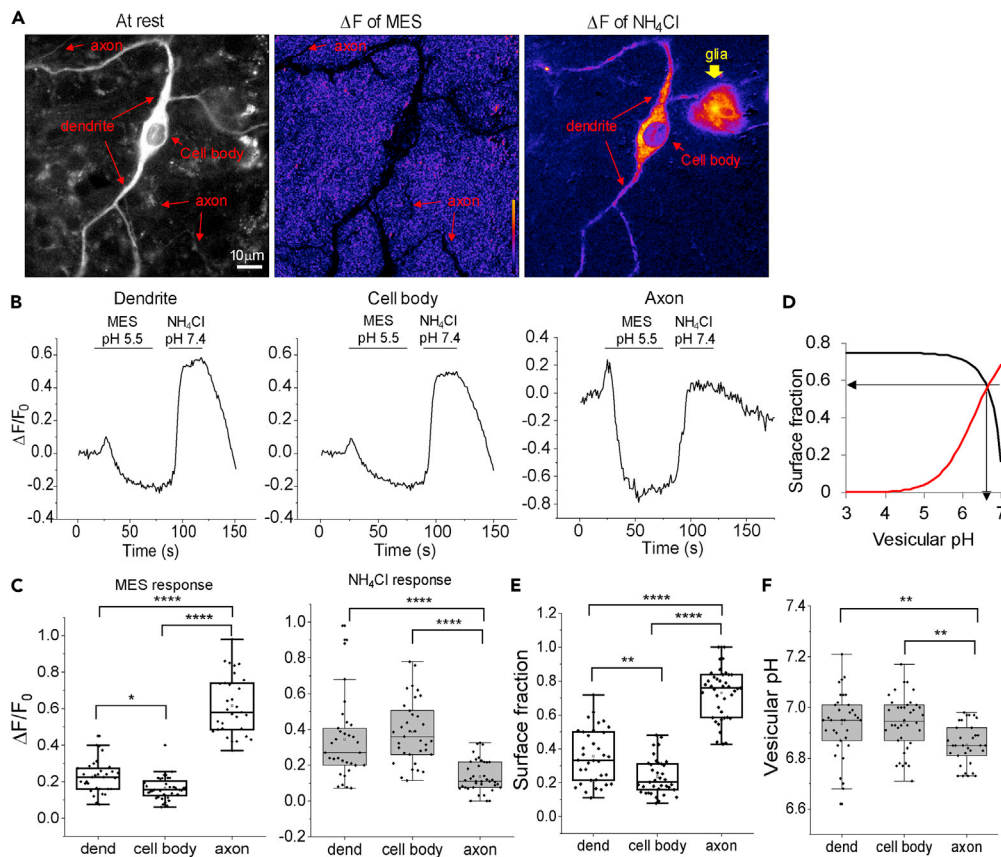


Figure 3. Majorities of axonal DAT is on the plasma membrane

(A) Representative imaging field containing a cultured midbrain neuron and a neighboring glia expressing DAT-pHluorin. From left to right: fluorescent image at rest in gray scale, ΔF image of the negative response to the MES solution perfusion and ΔF image of the positive response to the NH_4Cl solution perfusion. Arrows point to the cell body, neurites, and the glial cell, which were visible at different stages.

(B) Representative DAT-pHluorin signals measured from the dendrite, cell body and axon during a sequential MES and NH_4Cl perfusion experiment.

(C) Box and whisker charts for the $\Delta F/F_0$ of the MES and NH_4Cl responses measured at different compartments of the midbrain neuron. Each data point represents an averaged response from one cell across multiple trials. $N = 33$ for dendrites, $N = 36$ for cell bodies and $N = 31$ for axons. $F = 159.67$, $p = 2.06 \times 10^{-31}$, one-way ANOVA for MES responses. **** $p = 0.0000$ comparing axons versus dendrites and axons versus cell bodies; * $p = 0.044$ comparing dendrites versus cell bodies, Bonferroni *post hoc* following one-way ANOVA. $F = 22.56$, $p = 7.29 \times 10^{-9}$, one-way ANOVA for NH_4Cl responses. **** $p < 1.00 \times 10^{-5}$ comparing axons versus dendrites; **** $p < 1.00 \times 10^{-7}$ axons versus cell bodies; $p = 0.90$ comparing dendrites versus cell bodies, Bonferroni *post hoc* following one-way ANOVA. Data from four batches of cultures.

(D) Illustration for the calculation of surface fraction and vesicular pH based on the $\Delta F/F_0$ measurements obtained in (B) using the Henderson-Hasselbalch equation (see STAR Methods).

(E and F) Box and whisker charts summarizing the surface fraction (E) and vesicular pH (F) of DAT-pH in different cellular compartments. $F = 131.72$, $p = 6.37 \times 10^{-30}$, one-way ANOVA for surface fraction. **** $p = 0.0000$ comparing axons versus dendrites and axons versus cell bodies; ** $p = 0.0012$ comparing dendrites versus cell bodies, Bonferroni *post hoc* following one-way ANOVA. $F = 7.14$, $p = 0.0013$, one-way ANOVA for vesicular pH. ** $p = 0.0063$ comparing axons versus dendrites; ** $p = 0.0026$ comparing axons versus cell bodies; $p = 1.00$ comparing dendrites versus cell bodies, Bonferroni *post hoc* following one-way ANOVA. See also Figures S3 and S4.

response (Figures 4B and 4C). No significant difference was observed for the MES responses (Figure 4C). We further calculated the surface DAT fraction using the Henderson-Hasselbalch equation and found that axonal shafts and boutons exhibited a similar $\sim 75\%$ surface fraction of DAT and similar acidity for DAT-containing vesicles (Figures 4D and 4E). No significant difference was observed between TH+ and TH- neurons (Figure S4B). Varicosities, however, exhibited only $\sim 20\%$ surface DAT (Figure 4D) and DAT-containing

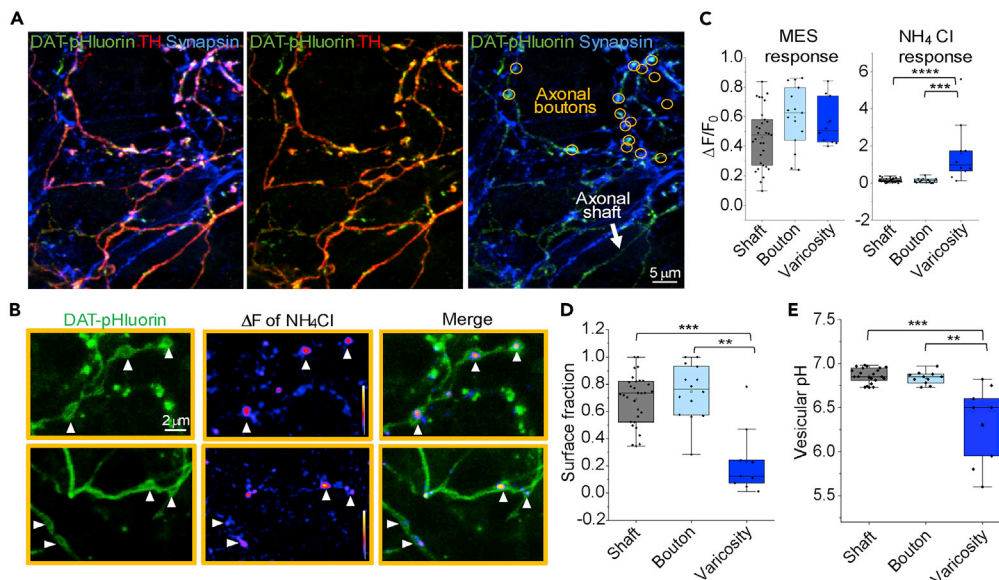


Figure 4. Varicosities contain a large portion of acidic intracellular DAT

(A) Representative immunofluorescence images of the axons expressing DAT-pH labeled with anti-GFP, anti-TH and anti-synapsin1/2. Orange circles indicate synapsin-enriched axonal boutons and the white arrow points to an axonal shaft with light synapsin staining.

(B) Representative axonal segments expressing DAT-pH and responded robustly to NH_4Cl perfusion. From left to right: DAT-pH axon at rest (left), the ΔF image of the NH_4Cl response of the axon (middle) and the overlay of the two images (right). The arrowheads point to varicosities with DAT-pH contours and NH_4Cl -responding centers.

(C) Box and whisker charts summarizing the MES and NH_4Cl responses for axonal shaft (N = 31), synapsin boutons (N = 15) and large varicosities (N = 10). $F = 2.90$, $p = 0.064$, one-way ANOVA for MES responses. Chi-Square = 19.62, $***p = 5.50\text{E-}5$, Kruskal-Wallis test for NH_4Cl responses. $***p = 7.3\text{E-}5$ comparing varicosity versus shaft, $***p = 8.41\text{E-}4$ comparing varicosity versus bouton, $p = 1.00$ comparing bouton versus shaft, Mann-Whitney test with Bonferroni correction.

(D and E) Calculated surface fraction (D) and DAT-containing vesicular pH (E) based on the Henderson-Hasselbalch equation using the measurements in C. Data from four batches of cultures. Chi-Square = 16.45, $***p = 2.69\text{E-}4$ Kruskal-Wallis test for surface fraction. $***p = 5.85\text{E-}4$ comparing varicosity versus shaft, $**p = 0.0018$ comparing varicosity versus bouton, $p = 0.96$ comparing bouton versus shaft, Mann-Whitney test with Bonferroni correction. Chi-Square = 16.95, $***p = 2.09\text{E-}4$ Kruskal-Wallis test for NH_4Cl responses. $***p = 3.45\text{E-}4$ comparing varicosity versus shaft, $**p = 0.0032$ comparing varicosity versus bouton, $p = 1.00$ comparing bouton versus shaft, Mann-Whitney test with Bonferroni correction. See also [Figure S4](#).

vesicles were substantially more acidic with an average pH of 6.31 ± 0.14 (Figure 4E). Thus, our data suggests that the constitutive recycling of DAT is similar between axonal shafts and synaptic boutons, whereas large varicosities stand out as unique structures potentially involved in DAT endocytosis and degradative processing.

Cocaine exposure elicits a redistribution of DAT to the axonal plasma membrane

Cocaine has been shown to not only inhibit DAT's reuptake activity but also to increase DAT surface expression,^{15–17,19} or to prevent DAT from internalization.^{18–20} The precise manner by which cocaine changes DAT surface expression/availability, especially in neurons, is not clear. An increase in DAT binding sites were reported in the striatal/synaptic regions of primates after chronic cocaine use.^{59,60} In rodent models of substance use disorder, voltammetry studies have suggested reduced DA uptake, hence, less surface DAT in the nucleus accumbens (NAc) of mice given extended cocaine access,^{61,62} whereas in the intermittent access model, an enhanced cocaine potency suggested increased DAT surface expression.⁶³

To better understand cocaine-induced DAT trafficking in neurons, and to specifically pinpoint the changes of DAT surface expression in various subcellular compartments of the neuron, we expressed DAT-pH in cultured midbrain neurons and subjected the cell culture to either a 1-day or a 4-day cocaine (10 μM) treatment. We then compared surface DAT fraction and DAT-containing vesicular pH from cocaine treated cells

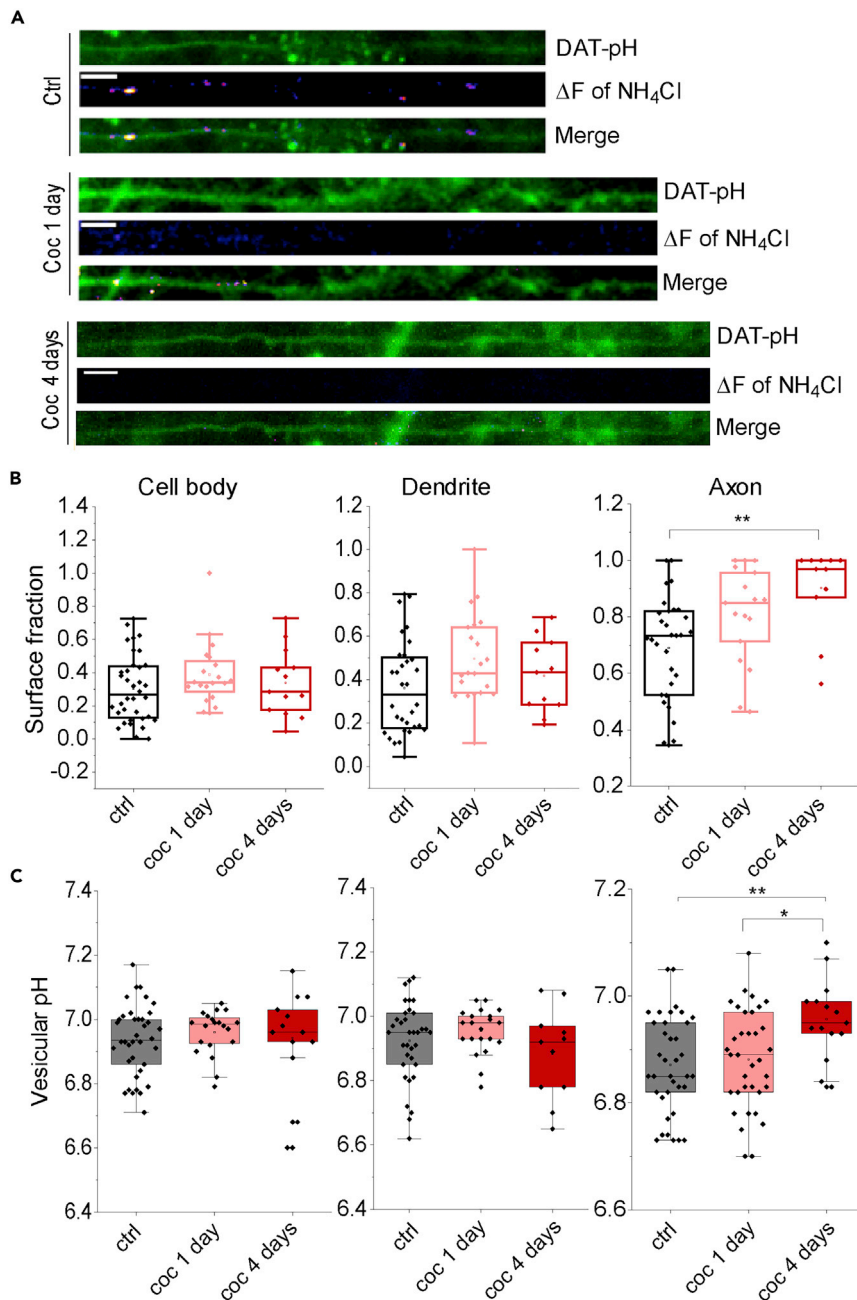


Figure 5. Cocaine exposure results in an increase in surface DAT fraction in the axon

(A) Representative NH_4Cl responses (ΔF of NH_4Cl) at the axon in cells without exposure to cocaine (control/ctrl), those that have been exposed to cocaine for 24 h (coc 1 day) and those that have been exposed to cocaine for 4 days (coc 4 days). The baseline DAT-pH images were shown above the NH_4Cl ΔF images and the overlay images (merge) were shown at the bottom. Scale: 5 μm .

(B and C) Box and whisker plots for surface fraction (B) and vesicular pH (C) at the cell body, dendrites and axons in cocaine treated and non-treated neurons. Cell body: ctrl N = 32, coc 1 day N = 20, coc 4 days N = 13, $F = 1.17$, $p = 0.32$ for surface fraction and $F = 0.34$, $p = 0.71$ for vesicular pH, one-way ANOVA. Dendrite: ctrl N = 32, coc 1 day N = 21, coc 4 days N = 11, $F = 2.78$, $p = 0.070$ for surface fraction and $F = 1.50$, $p = 0.23$ for vesicular pH, one-way ANOVA. Axon: ctrl N = 31, coc 1 day N = 17, coc 4 days N = 11, Chi-Square = 12.57, $**p = 0.0019$ for surface fraction, Kruskal-Wallis ANOVA. $**p = 0.0044$ comparing ctrl versus coc 4 days, $p = 0.098$ comparing ctrl versus coc 1 day, $p = 0.22$ comparing coc 1 day versus coc 4 days, Mann-Whitney test with Bonferroni correction. $F = 5.19$, $p = 0.0075$ for vesicular pH, one-day ANOVA. $**p = 0.0071$

Figure 5. Continued

comparing ctrl versus coc 4 days, * $p = 0.023$ comparing ctrl versus coc 1 day, $p = 1.00$ comparing coc 1 day versus coc 4 days, Bonferroni post hoc following one-way ANOVA. Data from six batches of cultures for control, four batches of cultures for coc 1 day and three batches of cultures for coc 4 days.

to cells without cocaine treatment from the same batch. Of interest, DAT surface fraction remained relatively steady at the cell bodies and dendrites (Figure 5B). The axons exhibited a significant increase in surface DAT by day 4 (Figure 5B), which is consistent with the gradual reduction of the NH_4Cl response (Figure 5A) and a significant alkalization of the DAT-containing vesicles (Figure 5C).

The cocaine-induced DAT trafficking is regulated by synaptojanin1

To further investigate the mechanisms underlying cocaine-regulated DAT trafficking, we examined DAT-pH signals in response to a shorter cocaine treatment. Similar to the N2a cells (Figure 2F), neuronal cell bodies and axons exhibited a cocaine-induced reduction of DAT-pH fluorescence, which recovered and reversed to an increase in DAT-pH fluorescence within 5 min following cocaine removal (Figures 6A and 6B). Recent studies have suggested molecules important for clathrin coat removal, such as synaptojanin1 (Synj1) or auxilin1, play a crucial role in DAT expression and/or trafficking.^{40,57,58} We therefore decided to investigate if Synj1 deficiency influences cocaine-regulated DAT trafficking using this new reporter. Of interest, in neurons carrying a heterozygous mutation of *Synj1* (*Synj1*^{+/-}), a larger deflection was observed (Figures 6A and 6B), which could suggest increased endocytosis of DAT or delivery of DAT to more acidic intracellular structures. Nonetheless, the slightly altered internalization phase did not impact the steady-state surface delivery of DAT-pH at the cell body (Figure 6A) but completely abolished this process in the axons of *Synj1*^{+/-} neurons (Figure 6B). *Synj1* encodes a phosphoinositide phosphatase, synaptojanin1 (*synj1*) which has been well-characterized for its role in clathrin-mediated synaptic vesicle (SV) recycling.^{37,38,40,41,64–66} *Synj1* deletion results in an accumulation of clathrin-coated vesicles within the synaptic terminal.^{36,37,67} Our data demonstrates the first evidence that Synj1 also regulates cocaine-induced DAT trafficking in the axons.

We next sought to determine the domain function of Synj1 involved in this regulation by expressing mutant human SYNJ1 (hSJ1) in N2a cells. In our previous work, we have shown that the Parkinson disease (PD)-linked R258Q mutation in the SAC1 phosphatase domain abolishes the PI(3)P and PI(4)P hydrolysis activities of the enzyme without affecting the primary 5'-phosphatase (5'-PPase) activity; however, the PD-linked R839C mutation reduces the PI(4,5)P₂ hydrolysis (by 60%) and abolishes PI(4)P hydrolysis (Figure 6C).⁴¹ Mice carrying the Synj1 R258Q mutation exhibit DAT clusters specifically in the dorsal striatum,⁴⁰ suggesting dysregulation of DAT trafficking via the SAC1 enzyme. Another mutation in the 5'-PPase domain, hD769A/mD730A, which was previously reported to abolish the PI(4,5)P₂ hydrolysis³⁸ was also included in this analysis to perform a comprehensive dissection of the lipid signaling pathways. We found that wild-type (WT) SJ1 expression did not influence the surface delivery of DAT despite a smaller internalization response (Figure 6D). Both the RC and DA mutations interfered with DAT surface delivery, however, the RQ mutation in the SAC1 domain exhibited the most statistically significant impairment compared to the WT SJ1 (Figures 6D and 6E). Thus, our work using a newly engineered DAT optical reporter suggested an essential role of Synj1's phosphatase activities in cocaine-regulated DAT trafficking. Specifically, although the 5'-PPase is more important for gating DAT internalization, the SAC1 enzyme is perhaps essential, especially for neuronal axons, to support DAT surface delivery following internalization.

DISCUSSION

DAT trafficking is an integral part of DAT function and bears tremendous importance in DA-related disorders. However, our understanding on neuronal DAT trafficking is limited primarily because of the lack of research tools that are capable of revealing DAT trafficking events in small structures such as the axons. Here, we report the engineering of a DAT reporter, DAT-pH. We show that DAT-pH senses pH with a stable pKa of 7.0, while exhibiting DA uptake, cocaine sensitivity and regulated trafficking. We show that in cultured MB neurons, DAT-pH exhibit 25% of surface expression at cell bodies but 75% in axons. We further identified large varicosities as potential hotspots for axonal DAT internalization and degradation. Finally, we report a novel role of Synj1 in assisting cocaine-induced DAT trafficking to the axonal surface.

Compared to existing tools that measure DAT trafficking, DAT-pH offers multiple advantages: (1) DAT-pH is capable of revealing DAT localization or constitutive recycling in small neuronal structures that otherwise

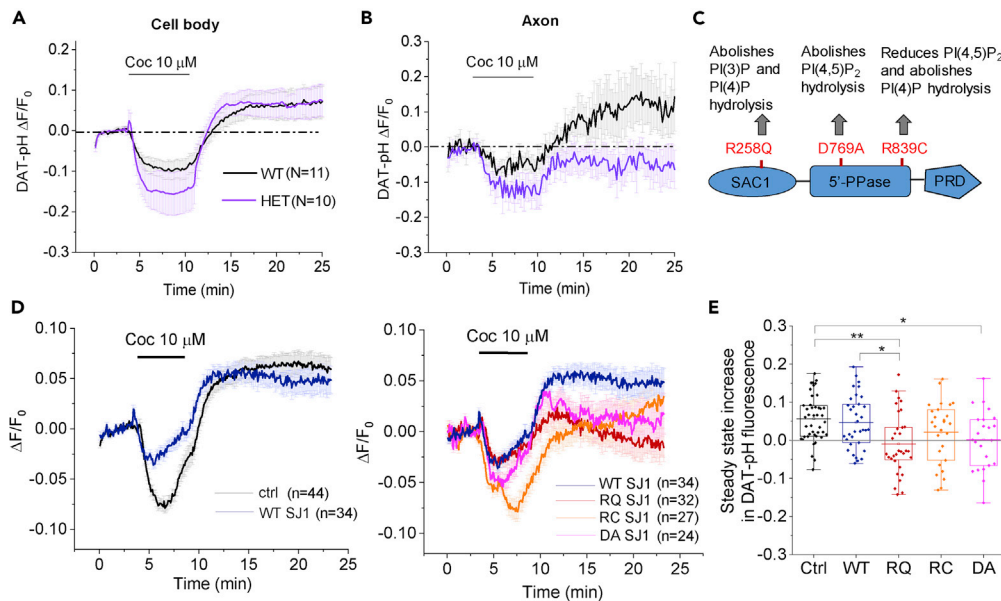


Figure 6. The cocaine-induced DAT trafficking is regulated by Synj1

(A and B) Averaged DAT-pH signals in response to an acute perfusion of cocaine measured from the cell body (A) or axon shafts (B) of the littermate MB neurons. Data is represented as mean \pm SEM N = cell number from three independent batches of cultures.

(C) Synj1 domain structure, mutants used in the study and the previously reported impact of these mutations on phosphoinositide metabolism.

(D) Averaged DAT-pH signals in response to an acute perfusion of cocaine measured from control (ctrl) N2a cells or those transfected with WT hSJ1 or hSJ1 mutants. Left, overlay of the responses from control cells (black) and cells expressing WT SJ1 (navy). Right, overlay of the responses from WT and mutant SJ1 transfected cells. Data is represented as mean \pm SEM from two independent batches of cultures. N = cell number.

(E) Box-whisker charts for the average of the final 3 min of the DAT-pH responses, which was denoted as steady state increase across all test groups. $F = 5.21$, $p = 5.78E-4$, one-way ANOVA. ** $p = 0.0014$ comparing ctrl versus RQ, * $p = 0.018$ comparing WT versus RQ, * $p = 0.041$ comparing ctrl versus DA, $p = 1.00$ comparing WT vs. RC, $p = 0.22$ comparing WT versus DA, Bonferroni post hoc analysis following one-way ANOVA.

requires electron microscopy (EM) or super resolution imaging analysis. We report a compartmentalized expression pattern of DAT in neurons, which resembles previous EM analyses in rodent brains.^{21,68,69} These results lend credibility to visualizing actual DAT trafficking events in neurons. (2) DAT-pH is a superior tool for revealing the dynamic changes of DAT in real-time for cell lines and neurons. Fluorescent DAT ligands, which have also been reported to label and track plasma membrane DAT,^{7,8} often exhibit competitive binding with cocaine and native DA. Using DAT-pH we report for the first time the dynamic internalization and surface delivery of DAT on cocaine exposure, which is regulated by a phosphoinositide phosphatase, Synj1. It is also worth noting that although N2a cells initiated the reverse trafficking of DAT within 5 min of cocaine exposure, neurons did so after cocaine removal. These kinetic differences between a neuron and a heterologous cell, although not yet understood, would not be possible with conventional tools. (3) DAT-pH, being a genetically encoded reporter, has the potential to be used *in vivo* or in combination with another optical sensors, such as RdLight1⁷⁰ that measures DA. The DAT is an essential molecule for shaping DA transients in the brain, however, little is known about how trafficking of DAT is regulated by DA or other psychostimulants; and whether/how the translocation of DAT in turn modifies DA transients in the brain. For example, the somatodendritic DAT has been shown to not only mediate DA uptake but also participates in reverse transport of DA.⁷¹⁻⁷⁴ Thus, real-time measurement of DAT surface expression and DA transients may advance our understanding of DA signaling in an unprecedented way.

The DAT-pH reporter and its future derivatives are timely tools for investigators across the fields to understand DAT dysregulation in substance use disorders, affective disorders, and Parkinsonian neurodegeneration. Using DAT-pH we revealed a previously uncharacterized mode of axonal DAT trafficking especially in response to cocaine exposure. Interestingly, WT SJ1 overexpression resulted in a less robust internalization

but did not affect the steady state DAT surface expression. The RQ SJ1 induced a similar internalization as the WT SJ1 but failed to deliver DAT to the surface. Thus, the internalization phase that we observed appeared to be independent from DAT surface delivery. We are yet to understand the biological significance and the molecular mechanisms underlying the cocaine-induced DAT internalization. However, we are the first to demonstrate that the cocaine-induced surface expression of DAT is regulated by Synj1. Mutations in both phosphatases of *SYNJ1* have been implicated in early onset Parkinsonism.^{75–78} Remarkably, genetic variants of *SYNJ1* are also associated with bipolar disorder.^{79–81} The lack of ligand-induced DAT trafficking in the axons of *Synj1*^{+/-} neurons may contribute to the altered DA signaling in the pathogenesis of these disorders, which we will investigate in the future. We envision that similar analyses can be carried out to assess other disease risk genes involved in PD, substance use disorders and psychosis.

Taken together, using a pH sensitive DAT reporter our study demonstrated previously uncharacterized biology of DAT trafficking in different parts of the neuron. We suggest immense potential for this reporter in revealing greater molecular details of DAT trafficking and DA signaling in physiological states as well as in disease pathogenesis.

Limitation of the study

There are a few limitations to be noted while interpreting data obtained using DAT-pH: First, DAT-pH measures pH rather than direct surface expression, which brings richness to the data (such as the DAT-containing vesicular pH) but also caveats when interpreting a singular measurement in a time-lapse study. Second, the DAT-pH analysis used in this study was unable to distinguish surface internalization from membrane lateral diffusion, another important form of DAT trafficking in the axon.^{7,82} Indeed, we showed that the cocaine-induced increase in surface DAT at the cell body (Figure 6A) was not measurable after 24 h (Figure 5B), suggesting potential lateral diffusion. Future endeavors in developing ratiometric reporters for DAT would circumvent such concerns. Third, to apply DAT-pH analysis on small molecules known to induce DAT oligomerization,^{20,83} we have yet to test in a more rigorous way whether DAT oligomerization would alter the pKa of DAT-pH. Lastly, the expression of the current DAT-pH in neuronal axons is relatively weak, which limits signal/noise ratio for axonal DAT trafficking and *in vivo* imaging at the striatum or nucleus accumbens. Further optimization of the reporter design is desired to expand the types of applications for DAT-pH.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at <https://doi.org/10.1016/j.isci.2022.105782>.

ACKNOWLEDGMENTS

We would like to thank Dr. Amy Newman for providing the cocaine analogue, JHC1-64 dye; Drs. Zhiping Pang, Timothy Ryan, and Jeffrey Diamond for providing valuable insight for the manuscript; and Dr. Loren Runnels for sharing the scintillation counter. The work is supported by an NINDS R01 grant (NS112390) awarded to P-Y.P., a Rutgers Brain Health Institute Pilot grant awarded to P-Y.P. and D.J.B., and a NINDS diversity supplement for J.S. (3R01NS112390-02S1).

AUTHOR CONTRIBUTIONS

J.S., D.J.B., and P-Y.P. conceptualized the project, J.S., O.Y., E.K., M.A., and P-Y.P. contributed to data collection and analysis. X.Z. and E.D-B. participated in the radioactive DA uptake study. D.J.B. and P-Y.P. obtained the funding. J.S. and P-Y.P. wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

One or more of the authors of this paper self-identifies as an underrepresented ethnic minority in their field of research or within their geographical location. One or more of the authors of this paper received support from a program designed to increase minority representation in their field of research.

Received: August 18, 2022

Revised: November 7, 2022

Accepted: December 7, 2022

Published: January 20, 2023

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Chicken anti-GFP	Thermo Fisher	Cat# A-10262; RRID: AB_2534023
Rabbit anti-GFP	Thermo Fisher	Cat# A-11122; RRID: AB_221569
Guinea pig anti-synapsin1/2	Synaptic System	Cat#106004; RRID: AB_1106784
Mouse anti-TH	Sigma	Cat# T2928; RRID: AB_477569
Rabbit anti-TH	Novus Biologicals	Cat# NB300-109; RRID: AB_10077691
Rat anti-DAT	Millipore	Cat# MAB369; RRID: AB_2190413
Bacterial and virus strains		
Bacteria: DH5 α	Thermo Fisher	Cat# 18265017
Bacteria: XL-10 Gold	Agilent Technology	Cat# 200314
Chemicals, peptides, and recombinant proteins		
Cytosine β -D-arabinofuranoside hydrochloride (ARA-C)	Sigma-Aldrich	Cat# C6645
Cocaine hydrochloride	Sigma-Aldrich	Cat# C5776
³ H-DA (Dihydroxyphenylethylamine (Dopamine), 3,4-[Ring-2,5,6-3H]-, 250 μ Ci (9.25MBq)	PerkinElmer	Cat# NET673250UC
D-Amphetamine Hemisulfate Salt	Sigma-Aldrich	Cat# A5880
Experimental models: Cell lines		
Neuro-2a (N2a) cells	ATCC	Cat# CCL-131; RRID: CVCL_0470
SH-SY5Y cells	ATCC	Cat# CRL-2266; RRID: CVCL_0019
Experimental models: Organisms/strains		
C57BL/6J mice	IMSR_JAX:	Cat# 000664; RRID: IMSR_JAX:000,664
Synj1 +/- mice	Pietro DeCamilli laboratory at Yale University	N/A
Oligonucleotides		
DAT-N Forward: cgagctcaagcttccaattctgcagtcgacgg	This paper	N/A
DAT-N ₁₉₄ Reverse: gctagtgcctaccgtaggtagggcatc	This paper	N/A
DAT-C ₁₉₅ Forward: tagtggcaccgggtgactccagtgg	This paper	N/A
DAT-C Reverse: cggacggctagagccggtggatcccg	This paper	N/A
CFP Forward: gctagcgtaccggtcaagaattgccacc	This paper	N/A
CFP Reverse: aattcactctcaggcgaattctgcagtcg	This paper	N/A
Recombinant DNA		
Plasmid: pEYFP-C1-DAT	Addgene	Cat# 90228; RRID: Addgene_90228

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Continued

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Plasmid: tagRFP-C1-DAT-HA	Addgene	Cat# 90265; RRID: Addgene_90265
Plasmid: pCAG-ECFP	Addgene	Cat# 32597; RRID: Addgene_32597
Plasmid: vGlut1-pHluorin	Timothy A. Ryan laboratory at Weill Cornell Medical College	N/A
Plasmid: Eevee-iAkt-PM	Kazuhiro Aoki at National Institute for Basic Biology, Japan	N/A
Software and algorithms		
ImageJ	Schneider et al. ⁸⁴	https://imagej.nih.gov/ij/
OriginPro 2018/2023	Origin Lab	https://www.originlab.com/
Other		
JHC1-64 dye	Amy Newman Laboratory at NIH	N/A

RESOURCE AVAILABILITY**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Ping-Yue Pan (pingyue.pan@rutgers.edu).

Materials availability

DAT-pHluorin generated in this study will be shared by the lead contact upon request.

Data and code availability

- All data reported in this paper will be shared by the [lead contact](#) upon request.
- This paper does not report original code.
- Any additional information required to reanalyze the data reported in this paper is available from the [lead contact](#) upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS**Animals**

C57BL/6J mice (RRID: IMSR_JAX:000664) were originally purchased from the Jax lab. The *Synj1*^{+/-} mice³⁶ were gifted by the Pietro DeCamilli laboratory at Yale University. *Synj1*^{+/-} mice were crossed to the C57BL/6J mice to generate littermate pups for MB cultures. Mice were housed in the pathogen-free barrier facility at the Rutgers Robert Wood Johnson Medical School SPH vivarium. Handling procedures were in accordance with the National Institutes of Health guidelines approved by the Institutional Animal Care and Use Committee (IACUC # PROTO201800183). Both male and female mice were used for preparing primary neuron cultures.

Cell lines

N2a cells (RRID: CVCL_0470) from male origin were cultured and passaged following an ATCC suggested protocol using culture media containing DMEM (Thermo Fisher, 11965118), 10% fetal bovine serum (FBS) (Atlantic Biologicals, S11550), and 5% 10 U/mL Penicillin-Streptomycin (Thermo Fisher, 15140122). Cells were trypsinized using 0.05% Trypsin-EDTA (Gibco, 25300-054) and plated at 30% confluency.

SH-SY5Y cells (RRID: CVCL_0019) of female origin were cultured and passaged according to the ATCC suggested protocol using DMEM/F-12(1:1) culture media (Gibco, 11320-033) containing 10% FBS (Atlantic Biologicals, S11550), and 5% 10 U/mL Penicillin-Streptomycin (Thermo Fisher, 15140122). For experiments, cells were trypsinized using 0.05% Trypsin-EDTA and plated at 50% confluency.

Primary cell culture

Midbrain cultures^{26,27} were prepared as described previously. Ventral midbrains (containing both VTA and SN) were dissected from P0 male and female mouse pups and digested using papain (Worthington, LK003178) in a 35 °C water bath with gentle stirring and constant oxygenation. Midbrain neurons were then seeded within a cloning cylinder on the #1.5 cover glasses pretreated with Poly-L-ornithine (Sigma, P3655). Cells were plated at a density of 199,000 cells/cm² and grown in the Neurobasal-A based medium supplemented with GDNF (10 ng/mL, EMD Millipore, GF030).

METHOD DETAILS

Constructs and cloning

Human DAT cDNA was amplified from YFP-hDAT (Addgene, 90228) and pHluorin was amplified from vGlut1-pHluorin (a gift from Timothy A. Ryan and pHluorin is an original asset of James E. Rothman). The hDAT-pHluorin cDNA was inserted in the modified pCAGGS backbone (cut out from the Eevee-iAkt-pm sensor,⁸⁵ a gift from Kazuhiro Aoki) using the ECoR-I and Xba-I restriction sites. The plasmid contains a CAG (combination of chicken beta-actin promoter with CMV enhancer) promoter. An Age-I restriction site was inserted first by two-step PCR amplification using primers at the desired position of hDAT, followed by enzyme digestion and pHluorin insertion. The same linkers in vGlut1-pHluorin were used (TGSTGGSGGTGG and SGGTGGSGGTGGSGGTG) on both ends of pHluorin in DAT-pH. The correct orientation of the pHluorin was confirmed by sequencing. DAT-HA was amplified from RFP-HA-DAT (Addgene, 90265) using ECoR-I and Xba-I sites and subcloned into the same backbone as the DAT-pH. All primers were synthesized by IDT custom DNA oligo services and all subcloning enzymes and buffers were purchased from New England Biolabs (NEB). DH5 α (Thermo Fisher, 18265017) or XL-10 Gold (Agilent Technology, 200314) competent cells were used for plasmid amplification.

The following constructs were purchased directly from Addgene: pEYFP-C1-DAT (90228), tagRFP-C1-DAT-HA (90265) and pCAG-ECFP (32597). CFP-DAT was subcloned using the following primers: forward-5'-gctagcgcctaccggcaagaattcgccacc-3' and reverse-5'-aattcactcctcaggcgaattctgcagctcg-3' to replace EYFP using the Age-I and EcoR-I enzymatic sites.

Cell culture transfection

For N2a cells and SH-SY5Y cells, transfection was carried out following a company suggested protocol the day after plating using LipofectamineTM3000 (Thermo Fisher, L3000015). The transfection mixture was left in the medium until the day of imaging (typically within 24-48). For primary neurons, LipofectamineTM 2000 (Thermo Fisher, 11668019) was used for transfection at DIV (days *in vitro*) 7 following a company suggested protocol. The DNA-lipofectamine mixture was washed out after 45 min incubation at 37 °C and the growth medium was replaced with a fresh medium supplemented with an antimetabolic agent, ARA-C (Sigma-Aldrich, C6645).

Radioactive DA uptake assay

Radioactive DA uptake assay was performed following previous published protocols.⁸⁶ N2a cells and SH-SY5Y cells were seeded in triplicates in the 24-well plate two days before the uptake assay. Cells typically reach 80% confluency on day of testing. For cocaine inhibition, cells were treated with 10 μ M cocaine (Sigma, C5776) for 50 min in the cell culture incubator prior to the uptake assay based on published results to achieve maximum inhibition.⁸⁷ ³H-DA (Dihydroxyphenylethylamine (Dopamine), 3,4-[Ring-2,5,6-3H]-, 250 μ Ci (9.25MBq)) was purchased from PerkinElmer (NET673250UC). 50 nM ³H-DA was diluted in the uptake buffer containing 130 mM NaCl, 1.3 mM KCl, 10 mM HEPES, 1.2 mM MgSO₄, 1.2 mM KH₂PO₄, 2.2 mM CaCl₂, 10 mM glucose as well as 50 mM L(+) ascorbic acid (Millipore, AX1775-3) and 50 mM pargyline (Sigma Aldrich, P8013.) from freshly prepared stock before the experiment. 10 μ M cocaine was added when testing cocaine inhibition. The pH of the buffer was adjusted to 7.4. Cells were incubated with the ³H-DA at room temperature for 10 min before washing with an additional uptake buffer without ³H-DA. Scintillation fluid was added to the wells immediately and transferred to the tubes. The radioactive signal was read by the scintillation counter (Beckman, LS 6500). For cocaine inhibition analysis, duplicate cultures were made for Western blot analysis and all uptake counts were normalized to the total DAT proteins in the culture.

Live imaging and data processing

For live cell imaging, cells on cover glass were mounted on a custom-made laminar-flow chamber with constant gravity perfusion (at a rate of ~0.2-0.3 mL/min) of a Tyrode's salt solution containing 119 mM NaCl, 2.5 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 25 mM HEPES, 30 mM Glucose, 10 μM 6-cyano-7-nitroquinoxaline-2,3-dione (CNQX), and 50 μM D, L-2-amino-5-phosphonovaleric acid (AP-5) and buffered to pH 7.40. The NH₄Cl medium contains: 50 mM NH₄Cl, 70 mM NaCl, 2.5 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 25 mM HEPES, 30 mM Glucose, 10 μM CNQX, and 50 μM AP-5, buffered to pH 7.40. The MES medium contains: 25 mM MES, 70 mM NaCl, 2.5 mM KCl, 2 mM CaCl₂, 2 mM MgCl₂, 30 mM Glucose, 10 μM CNQX, and 50 μM AP-5, buffered to pH 5.50. Perfusion of Tyrode's solution is regulated by Valvelink 8.2 and the NH₄Cl or MES solutions were perfused by pipettes. All chemicals were purchased from Sigma-Aldrich. Temperature was clamped at ~31.0 °C on the objective throughout the experiment. Images were acquired using a highly sensitive, back-illuminated EM-CCD camera (iXon+ Model DU-897E-BV, Andor Corp., CT, USA). Nikon Ti-2 wide-field microscope is modified with Spectra-X (Lumencor) as the light source for shuttered illumination. pHluorin fluorescence excitation and collection were using a Nikon Plan APO 60X 1.40 NA objective using 525/50m emission filter and 495LP dichroic filters (Chroma, 49002).

Images were sampled at 1 Hz using the Elements software and analyzed in ImageJ. Cell body responses were measured by regions of interests (ROIs) drawn by the freehand tool based on the ΔF image of the NH₄Cl response. Dendritic responses were measured using the 8x8 pixels (2 μM) circular ROIs based on the ΔF image of the NH₄Cl response. Axonal responses were measured using the 6x6 pixels (1.5 μM) circular ROIs based on the basal fluorescence image and the ΔF image of the MES response. For varicosity of a large size, several circular ROIs might be placed to cover the area. The average response of all ROIs from one of the three above compartments (cell body, dendrite or axon) was extracted using the Time Series Analyzer plugin. All responses were background subtracted followed by baseline normalization in OriginLabs. Typically, two or three trials of sequential perfusions were carried out with 5-10 min spacing for each imaging field and the averaged peak response was measured as the MES or the NH₄Cl response of the cell.

DAT-pH pKa measurements

Measurements were made at midbrain neuron axons due to the highest surface fraction and least interference with endogenous buffers. For the pH 5.50 and pH 5.60 solutions we used 25 mM MES (pKa 6.1) (Millipore-Sigma, M3671) to replace HEPES in Tyrode's. For the pH 8.50 and pH 9.50 solutions we used 25 mM BICINE (pKa 8.3) (Millipore-Sigma, B3876) to replace HEPES in Tyrode's. All solutions were perfused by hand pipetting during a period of 5 min imaging. Typically, data from each axon is averaged from two sequential perfusion experiments. DAT-pH fluorescence changes were normalized to the peak ΔF.

Calculation for surface fraction and vesicular pH

The calculation of surface fraction and vesicular pH was performed following previous published methods.⁸⁸ At any given pH (pH_i), the deprotonated DAT-pHluorin molecules determines the fluorescence. Based on the Henderson-Hasselbalch equation, the fluorescence change during perfusion of a pH 7.4 membrane permeable NH₄Cl solution is determined by the following function assuming pKa = 7.0:

$$dx = \frac{(1/(1 + 10^{7.0-7.4}) - 1/(1 + 10^{7.0-pH_i}))}{1/(1 + 10^{7.0-pH_i})}$$

The surface fraction as a function of pH_i is then plotted by

$$Fr = \frac{dx - \frac{dF}{F_0} \text{ of NH}_4\text{Cl response}}{dx * \frac{dF}{F_0} \text{ of NH}_4\text{Cl response} + dx}$$

During perfusion of a pH 5.5 MES solution, the ratio of surface to internal DAT-pH can be calculated as the following

$$\Phi = \frac{1/(1 + 10^{7.0-5.5})}{1/(1 + 10^{7.0-pH_i})}$$

The surface fraction as a function of pH_i is then plotted by

$$Fr = \frac{\frac{dF}{F_0} \text{ of MES response}}{1 - dx + \Phi * (1 - \frac{dF}{F_0} \text{ of MES response})}$$

The surface fraction and vesicular pH are determined at the point where two functions intersect.

JHC1-64 dye staining

Staining was performed following previously published protocols.^{7,8} Briefly, midbrain cultures were washed three times in ice cold JHC-buffer containing: 25 mM HEPES, pH 7.4, with 130 mM NaCl, 5.4 mM KCl, 1.2 mM CaCl₂, 1.2 mM MgSO₄, 1 mM L-ascorbic acid, 5 mM D-glucose. Cell culture was then incubated with 50 nM JHC1-64 dye diluted in JHC buffer at 4°C for 30 min. After incubation the dye was washed away three times with ice-cold JHC- buffer followed by immediate fixation with 4% PFA at room temperature.

Immunofluorescence and data analysis

Immunocytochemistry was performed following previously published procedures.^{26,27} The following antibodies were used for immunofluorescence: chicken anti-GFP (Thermo Fisher, A-10262, 1:1000), rabbit anti-GFP (Thermo Fisher, A-11122, 1:1000). Guinea pig anti-synapsin 1/2 (Synaptic System, 106004, 1:500), mouse anti-TH (Sigma, T2928, 1:1000), rabbit anti-TH (Novus Biologicals, NB300-109, 1:1000), rat anti-DAT (Millipore MAB369 1:1000). Immunofluorescence was analyzed using a Nikon Ti-2 wide-field microscope or a Nikon CREST spinning disk confocal microscope. The same imaging parameters were set for each batch of culture. Image stacks were taken at different focal planes at 0.9 μm interval to include the whole cell and a maximum projection image was generated for each stack via ImageJ for analysis. All analyses were done manually. Synapsin 1/2-positive boutons were determined based on the bright punctate staining pattern.

Controlled substances

Cocaine hydrochloride (Sigma C5776) and D-Amphetamine Hemisulfate Salt (Sigma A5880) were purchased from Sigma through authorized institutional distributors. For cocaine experiments, cells were treated with 10 μM cocaine for 24 (1 day) or for 4 consecutive days. Treatment typically starts on DIV (days *in vitro*) 12-13 and imaging experiments were performed between DIV 13 and DIV 17.

QUANTIFICATION AND STATISTICAL ANALYSIS

All data are replicated within the same batch and across multiple batches of cells. For datasets that follow normal distribution and exhibit equal variance, Student's *t* test or ANOVA test was used, followed by Bonferroni correction for *post hoc* analysis. For datasets that reject normal distribution and exhibit non-homogenous variance, Mann-Whitney test or Kruskal Wallance test was used, followed by Bonferroni correction for *post hoc* analysis. Separate sensitivity tests were performed for non-parametric datasets after log transformation and results were similar to those from non-parametric tests prior todata transformation.