The norepinephrine transporter (NET) mediates reuptake of synaptically released norepinephrine in central and peripheral noradrenergic neurons. The molecular processes governing availability of NET in the plasma membrane are poorly understood. Here we use the fluorescent cocaine analogue JHC 1-64, as well as several other approaches, to investigate the trafficking itinerary of NET in live noradrenergic neurons. Confocal imaging revealed extensive constitutive internalization of JHC 1-64-labeled NET in the neuronal somata, proximal extensions and presynaptic boutons. Phorbol 12-myristate 13-acetate increased intracellular accumulation of JHC 1-64-labeled NET and caused a parallel reduction in uptake capacity. Internalized NET strongly colocalized with the “long loop” recycling marker Rab11, whereas less overlap was seen with the “short loop” recycling marker Rab4 and the late endosomal marker Rab7. Moreover, mitigating Rab11 function by overexpression of dominant negative Rab11 impaired NET function. Sorting of NET to the Rab11 recycling compartment was further supported by confocal imaging and reversible biotinylation experiments in transfected differentiated CATH.a cells. In contrast to NET, the dopamine transporter displayed markedly less constitutive internalization and limited sorting to the Rab11 recycling compartment in the differentiated CATH.a cells. Exchange of domains between the two homologous transporters revealed that this difference was determined by non-conserved structural elements in the intracellular N terminus. We conclude that NET displays a distinct trafficking itinerary characterized by continuous shuffling between the plasma membrane and the Rab11 recycling compartment and that the functional integrity of the Rab11 compartment is critical for maintaining proper presynaptic NET function.

Reuptake of synaptically released norepinephrine (NE)\(^2\) is the primary mechanism to control duration and strength of noradrenergic neurotransmission. The reuptake is mediated by the NE transporter (NET), which is present in noradrenergic neurons both in the brain and in the peripheral sympathetic nerve system (1–3). NET belongs to the SLC6 family (solute carrier 6) gene family (also referred to as neurotransmitter-sodium symporters) that also includes transporters for the neurotransmitters dopamine, serotonin, glycine, and \(\gamma\)-aminobutyric acid (GABA) (4–6). NET is targeted by many clinically important antidepressant drugs and by several widely abused drugs, such as cocaine and amphetamine. In addition, impaired NET function has been implicated in cardiovascular diseases and in disorders of mood and cognition (7–10).

It is assumed that the amount of NET present in the presynaptic plasma membrane is tightly controlled to sustain NE homeostasis, but the underlying molecular and cellular mechanisms are poorly understood. Several mechanisms have been suggested to regulate NET distribution, including protein kinase C (PKC), whose activation (e.g. by \(G_q\)-coupled muscarinic acetylcholine receptors) can increase NET internalization and reduce NE transport capacity (11–14). Furthermore, amphetamine has been shown to decrease NET surface expression, whereas cocaine was found to enhance NET availability (15–17). An elegant study by Galli and co-workers (16) found evidence that the amphetamine-induced decrease in NET surface levels is Rab11-dependent. The general NET trafficking properties and postendocytic sorting pattern, however, have not been investigated in detail. Of interest, several SLC6 family transporters, such as the dopamine (DAT), serotonin (SERT), and glycine transporters, have been shown to undergo constitutive internalization (18–22). Their fate upon internalization,
however, has not been defined. Whereas some studies suggest that the transporters are recycled back to the plasma membrane
(18, 23), other studies support the notion that DAT and SERT
primarily are sorted to Rab7-positive late endosomes and sub-
sequently to lysosomal degradation upon both constitutive and
stimulated internalization (22, 24, 25).

Here, we employ several different approaches, including use
of the fluorescent cocaine analogue JHC 1-64, to describe NET
surface distribution, internalization, and postendocytic sorting
in live noradrenergic neurons. We show that NET undergoes
extensive constitutive internalization and that internalized NET
mainly is sorted to Rab11-positive recycling endosomes.
Inhibition of Rab11 function by dominant negative Rab11
(Rab11 S25N) impairs NE uptake capacity in noradrenergic
neurons, implying that correct and sufficient postendocytic
recycling is critical for NET function. Finally, we find that NET
displays a markedly higher degree of constitutive internaliza-
tion than DAT and that this difference, together with the dif-
ferential postendocytic sorting of NET compared with DAT,
is determined in part by non-conserved structural elements in the
intracellular N termini of the two transport proteins. Our
results reveal striking differences in the trafficking itineraries of
NET compared with DAT and provide thereby a critical frame-
work for further deciphering the role of NET and DAT in con-
trolling neurotransmitter homeostasis and how this might be
altered in diseased states.

Experimental Procedures

Molecular Biology—The cDNA encoding human NET was
kidnly provided by Dr. Marc G. Caron (Duke University, Dur-
ham, NC) (26). A synthetic gene encoding the human DAT was
a kind gift from Dr. Jonathan A. Javich (Columbia University,
New York). The enhanced green fluorescent protein (EGFP)-
tagged Rab7 and Rab11 constructs (pEGFP-Rab7 and pEGFP-
Rab11) were kind gifts from Dr. Katherine W. Roche (NINDS,
National Institutes of Health, Bethesda, MD) (27). The plasmid
pEGFP-Rab4 was a kind gift from Dr. José A. Esteban (Univers-
sidad Autónoma de Madrid, Madrid, Spain) (28). The NET (1–
128)/DAT construct was a kind gift from Dr. Bruno Giros
(Douglas Mental Health University Institute, Montreal, Can-
ada) (26). The viral vector plasmids pHsSynXW EGFP-Rab4,
Rab7, and Rab11 were generated as described previously (20).
A DAT/NET (590–617) chimeric construct was generated by
introducing a unique restriction site, BsIV, to the coding
sequence of NET by PCR. Subsequently, the C-terminal tail of
NET was ligated into DAT using digestion with BsIV and XbaI
and cloned into pcDNA3.1. DAT (1–59)/NET and NET/
DAT (593–620) chimeras were generated by a two-step PCR
procedure. First, the FLAG-Tac was amplified from pcDNA3 FLAG-Tac
with primers generating an overhang identical to the N-termi-
nal part of NET. Second, NET was amplified from pcDNA3
NET with primers with an overhang identical to the C-terminal
part of FLAG-Tac. The products from the first round of PCR
were used as templates for generating a TacNET fragment that
subsequently was cloned into pcDNA3. All constructs were
verified by dideoxynucleotide sequencing (Eurofins Genomics,
Ebersberg, Germany).

Primary Culture of Superior Cervical Ganglion Neurons—
Postnatally derived rat superior cervical ganglion (SCG) neu-
rons were prepared using a protocol modified from that of
Savchenko et al. (30). Briefly, ganglia from 0–3-day-old rat
pups were dissected and dissociated for 30 min at 37 °C in 3
mg/ml collagenase/0.5 mg/ml trypsin in RPMI medium gently
oxygenated with a mixture of 95% O₂ and 5% CO₂. Digestion
was terminated by adding 10% fetal bovine serum (FBS) in
RPMI medium. Subsequently, cells were briefly centrifuged and
then resuspended in warm RPMI medium supplemented with
3% heat-inactivated FBS and 20 ng/ml nerve growth factor
(NGF). Digested tissue was carefully triturated into single cells
by using increasingly smaller pipette tips. The neurons were
seeded onto coverslips coated with glial cells for immunocyto-
chemistry into similarly coated Lab-Tek chambers for live cell
imaging or into similarly coated 96-well plates for uptake
experiments. On the next day, cultures were treated with 1
μM 5-fluoro-5-deoxyuridine (Sigma) to inhibit further growth.

Lentivirus Production and Transduction—Lentivirus was
produced as described previously (20) according to procedures
modified from Naldini et al. (31). The SCG neuronal cultures
were transduced at days 4–5 in vitro, and experiments were
performed 8–11 days after infection.

Cell Cultures and Transfections—Human embryonic kidney
(HEK293) cells (ATCC CRL-1573) were grown in Dulbecco’s
modified Eagle’s medium (DMEM) with HEPES and sodium
bicarbonate (DMEM 1965), and differentiated CATH.a (CAD)
cells (ATCC CRL-11179), originating from mouse brain, were
grown in a 1:1 mixture of DMEM 1965 and Ham’s F-12 medium
(Invitrogen), both supplemented with 10% FBS and 0.01 mg/ml
gentamicin. For transient expression, the cells were transfected
using standard Lipofectamine 2000 protocols (Invitrogen) 2
days prior to experiments.

[^H]Dopamine Uptake—NET uptake experiments were car-
rried out using[^H]DA (30–60 Ci/mmol, PerkinElmer Life Sci-
ences) as substrate. NET transports DA and NE with similar
affinity ($K_m = 0.5–1 μM$), and [^H]DA is commonly used as
NET substrate (26, 32–34). Medium was removed from SCG
neurons cultured on glial cell monolayers in 96-well plates, and
the cells were equilibrated in uptake buffer (25 mM HEPES, pH
7.4, 130 mM NaCl, 5.4 mM KCl, 1.2 mM CaCl₂, 1.2 mM MgSO₄, 1

\[ \text{[^H]DA} \]
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mm l-ascorbic acid, 5 mm d-glucose, and 1 μm catechol-O-methyltransferase inhibitor Ro 41-0960 (Sigma) for 20 min at 37 °C. Subsequently, phorbol 12-myristate 13-acetate (PMA) (1 μm) or vehicle (0.01% DMSO) was added, and neurons were incubated for 30 min at 37 °C. Uptake was initiated by the addition of 100 nM [3H]DA, allowed to continue for 5 min at room temperature, and terminated by rapid aspiration of the medium and two washes with ice-cold uptake buffer. Nonspecific uptake was determined in the presence of 100 nM [3H]DA, allowed to continue for 5 min at room temperature. Nonspecific uptake was determined using increasing concentrations of DA (final concentration 0.05–6.4 μM) with a trace amount of [3H]DA in CAD cells transfected 48 h before the experiment with equal amounts of constructs encoding NET, DAT, NET-DAT chimeras, or TacNET. Uptake was initiated by adding serial dilutions of dopamine/[3H]dopamine and was allowed to continue for 3 min at room temperature. Nonspecific uptake was determined in the presence of 100 μM nomifensine. Subsequently, cells were lysed in 1% SDS and transferred to counting plates (PerkinElmer Life Sciences). Opti-Phase HiSafe3 scintillation fluid (PerkinElmer Life Sciences) was added to the wells, and the samples were counted in a Wallac Tri-Lux β-scintillation counter (PerkinElmer Life Sciences).

Saturation kinetics of DA uptake were determined using increasing concentrations of DA (final concentration 0.05–6.4 μM) with a trace amount of [3H]DA in CAD cells transfected 48 h before the experiment with equal amounts of constructs encoding NET, DAT, NET-DAT chimeras, or TacNET. Uptake was initiated by adding serial dilutions of dopamine/[3H]dopamine and was allowed to continue for 3 min at room temperature. Nonspecific uptake was determined in the presence of 100 μM nomifensine. Subsequently, cells were lysed in 1% SDS and transferred to counting plates (PerkinElmer Life Sciences). Opti-Phase HiSafe3 scintillation fluid was added, and the samples were counted in a Wallac Tri-Lux β-scintillation counter. Uptake data were analyzed with GraphPad Prism version 5.0 (GraphPad Software, La Jolla, CA).

Immunocytochemistry—SCG neurons grown on glial monolayers on coverslips were incubated with 50 nm JHC 1-64 in imaging buffer (25 mM HEPES, pH 7.4, with 130 mM NaCl, 5.4 mM KCl, 1.2 mM CaCl2, 1.2 mM MgSO4, 5 mM d-glucose) for 20 min at room temperature. Subsequently, the specimens were fixed with 4% paraformaldehyde in phosphate-buffered saline (PBS), blocked, and permeabilized in blocking/permeabilization buffer (5% goat serum, 0.2% saponin, PBS), and immunostained with a monoclonal mouse anti-NET antibody (NET-05, mAb Technologies; 1:1000) or a monoclonal mouse anti-tyrosine hydroxylase antibody (Mab318; Millipore, 1:1000) for 30 min at room temperature. Primary antibodies were labeled with a goat polyclonal anti-mouse Alexa Fluor® 488 secondary antibody (Molecular Probes, Inc., Eugene, OR) for 30 min at room temperature. The specimens were mounted on glass slides with Slow-Fade Antifade (Invitrogen) and visualized by confocal microscopy.

Confocal Microscopy—Confocal microscopy was performed using a Zeiss confocal laser-scanning microscope (LSM 510) with an oil immersion ×63 lens with a 1.4 numerical aperture (Carl Zeiss, Oberkochen, Germany) as described (20). Rhodamine-conjugated JHC 1-64, synthesized according to methods described previously (35), and Alexa Fluor® 568 dye were excited using 543-nm laser light from a helium-neon (HeNe) laser source, and emitted fluorescent light was filtered using a 585-nm long pass filter. Alexa Fluor 488 dye and EGFP were excited using a 488-nm laser line from an argon-krypton laser, and the emitted light was filtered using a long pass 505–530-nm barrier filter. Image processing was performed with ImageJ (National Institutes of Health).

JHC 1-64 Live Cell Imaging—NET surface expression was visualized in live SCG neurons grown on glial monolayers in LabTek chambers using the rhodamine-conjugated cocaine analogue JHC 1-64 (50 nm) in imaging buffer for 20 min at 4 °C to label surface NET. Subsequently, the buffer was removed, and the cells were rapidly washed twice with cold imaging buffer and imaged immediately using a Zeiss LSM 510 instrument. To verify the specific binding of JHC 1-64 to NET, some specimens were preincubated with desipramine (100 μM) for 20 min before labeling with JHC 1-64. To visualize NET internalization in live neurons, SCG neurons were first incubated with JHC 1-64 (50 nm) at 4 °C to label surface NET. Subsequently, the buffer was removed and replaced with prewarmed imaging buffer, and internalization was allowed for 1 h at 37 °C. JHC 1-64-labeled NET was then visualized by confocal microscopy carried out at 37 °C. To assess the effect of activation of PKC on NET internalization, SCG neurons were incubated for 1 h at 37 °C in the presence of 1 μM phorbol 12-myristate 13-acetate (PMA) or vehicle (0.01% DMSO) after surface labeling of NET with JHC 1-64 at 4 °C.

To investigate the postendocytic fate of internalized NET in live neurons, SCG neurons were transduced with different endosomal markers (EGFP-Rab4, -Rab7, -Rab11, and -Rab11 S25N) on days 4–5 in vitro. Next, transduced (and non-transduced) SCG neurons were incubated with JHC 1-64 (50 nm) to label surface NET before 1 h of internalization at 37 °C. After incubation, the living neurons were imaged by confocal microscopy carried out at 37 °C. To further assess the postendocytic fate of NET, we also visualized lysosomes by incubating non-transduced SCG neurons with LysoTracker Green (50 nm; Invitrogen) during the last 15 min of incubation at 37 °C. Finally, cells were washed twice and imaged at 37 °C.

Antibody Feeding Assay (Internalization Assay)—CAD cells transiently transfected with equal amounts of TacNET and EGFP-Rab7 or EGFP-Rab11 were seeded on coverslips treated with poly-l-ornithine. The cells were incubated with a mouse anti-FLAG antibody (M1 antibody) (1 μg/ml) in DMEM at 4 °C for 30 min to label surface-expressed TacNET. Subsequently, cells were washed three times with DMEM and then incubated in prewarmed medium for 1 h at 37 °C allowing internalization (or at 4 °C as a non-trafficking surface control). Internalization was terminated by washing the cells with ice-cold PBS, followed by fixation in 4% paraformaldehyde at 4 °C for 20 min, permeabilization in blocking/permeabilization buffer, and incubation with anti-mouse Alexa Fluor 568 secondary antibody for 30 min. Finally, the specimens were washed, mounted in ProLong Gold Antifade reagent (Molecular Probes, Invitrogen), and visualized by confocal microscopy.

Colocalization Assay—To evaluate colocalization of the transporters with different endosomal markers, CAD cells were transiently transfected with equal amounts of TacNET and EGFP-Rab7 or EGFP-Rab11. JHC 1-64 internalization assays were performed 48 h post-transfection by labeling surface expressed transporters at 4 °C with JHC 1-64 (50 nm). Subsequently, the cells were incubated for 1 h at 37 °C to allow internalization and imaged at 37 °C. Evaluation of transporter colocalization with EGFP-Rab7 and EGFP-Rab11 was performed
with the investigator kept unaware of the molecular species under investigation.

Quantification of Internalization—Quantification of the fraction of internalized NET in live SCG neurons was performed from single confocal sections. Background subtraction was performed by setting a minimum threshold pixel intensity of 50 in the 8-bit pictures, to ensure that only specific JHC 1-64 signal contributed to the quantification. The entire neuronal somata and proximal extensions (ROI-total) and the cytoplasmic parts of the somata and proximal extensions (ROI-internalized) were manually selected from each image. Internalization was defined as the ratio of the total integrated density of internalized NET (ROI-internalized) and the total integrated density of the entire labeled NET pool (ROI-total). 20–29 cells from at least three independent experiments were used for quantification under each condition. Statistical analyses were performed with GraphPad PrismTM, and significance was determined using one-way ANOVA with Bonferroni’s multiple-comparison test. Differences were considered significant when $p < 0.05$.

Quantification of Colocalization—Quantification of colocalization of NET, DAT, and NET-DAT chimeras with EGFP-Rab7 and EGFP-Rab11 was done using the Van Steensel’s cross-correlation function in the JaCoP Plug-in for ImageJ as described before (36–38). Van Steensel’s cross-correlation function reports the Pearson cross-correlation as a function of the relative movement of the two channels with respect to each other. Masks for the red and green channel were created using permanent settings for each image to avoid background. In a given overlay of the two images, the Pearson coefficient was plotted as a function of a 20-pixel shift in each $x$ direction of one of the images compared with the other. According to the cross-correlation function, complete colocalization results in a cross-correlation function of 1 at $Dx = 0$, and mutually exclusive structures will tend to show no peak or even a dip at $Dx = 0$. Colocalization was quantified for 10–25 cells from three to five independent experiments. Results are means ± S.E.

Quantification of Distances from the Plasma Membrane of JHC 1-64/DAT—Versus JHC 1-64/NET-positive Vesicles—Quantification of the distance of the JHC 1-64 positive vesicles from the plasma membrane in CAD cells transiently expressing NET or DAT was done using ImageJ. The distance from the plasma membrane was measured using the straight tool for 8–12 of the brightest JHC 1-64-positive vesicles per cell in 25–28 cells deriving from at least three independent transfections. Statistical significance was determined using an unpaired $t$ test. Differences were considered significant when $p < 0.05$.

Surface ELISA—To evaluate changes in surface expression of NET upon prolonged incubation with JHC 1-64 or PMA, HEK cells transiently expressing FLAG-tagged TacNET were seeded on 96-well plates. 48 h post-transfection, the TacNET-expressing cells were incubated with 500 nM JHC 1-64, 1 μM PMA, or vehicle (0.01% DMSO) for 1 h at 37 °C. Subsequently, the cells were washed with cold PBS, fixed in 4% paraformaldehyde, and blocked with 5% goat serum in PBS for 1 h at room temperature. The cells were incubated with 1 μg/ml M1 antibody for 1 h in blocking buffer, washed four times with PBS, and incubated with HRP-conjugated goat anti-mouse antibody (1:1000) in blocking buffer for 30 min at room temperature. Subsequently, the cells were washed four times with PBS and incubated with SuperSignal ELISA Femto Maximum Sensitivity Substrate (Pierce) for 30 s with shaking before luminescence was detected on a POLARstar Omega (BMG Labtech, Ortenberg, Germany) instrument. Results are presented as ratios of the surface signal from JHC 1-64- and PMA-treated cells relative to the surface signal from vehicle-treated cells on the same plate. Statistical significance was determined using a one-way ANOVA with Bonferroni’s multiple-comparison test. Differences were considered significant when $p < 0.05$.

Reversible Biotinylation (Internalization Assay)—Reversible biotinylation experiments were performed using a protocol modified from Schramm and Limbird (39). CAD cells were transiently transfected with equal amounts of NET, DAT, and NET-DAT chimeras 48 h before the experiments. The cells were rapidly cooled by washing with cold PBS and surface-biotinylated with 1.2 mg/ml disulfide-cleavable biotin (sulfo-NHS-SS-biotin; Pierce) for 1 h on ice. Free biotin reagent was washed out by quenching with glycine (100 mM) followed by two washes with PBS. Transporter endocytosis was initiated by incubating the cells with prewarmed medium with no extra treatment, leupeptin (100 μg/ml), or leupeptin (100 μg/ml) and monensin (25 μM) for 1 h at 37 °C. Control plates were kept on ice to prevent internalization. To stop endocytosis and to dissociate the biotin from the cell surface-resident proteins, the cells were washed with cold medium and incubated two times for 20 min with cold MesNa (100 mM) in 50 mM Tris-HCl (pH 8.8), 100 mM NaCl, 1 mM EDTA, 0.2% BSA on ice and subsequently washed twice with cold PBS. To determine total biotinylation, a sample of biotinylated cells was not subjected to reduction with MesNa. Cells were lysed in lysis buffer (25 mM Tris, pH 7.5, 100 mM NaCl, 1 mM EDTA, 1% Triton X-100, 0.2 mM PMSF, 1 protease inhibitor mixture tablet (Roche Applied Science) per 25 ml of buffer and 5 mM N-ethylmaleimide) and centrifuged, and supernatants were collected. Total fractions were collected from all samples and mixed with an equal volume of 2× loading buffer containing 1% SDS, 2.5% β-mercaptoethanol, and 100 mM dithiothreitol (DTT). For pull-down of the biotinylated proteins, the samples were incubated overnight with avidin beads (Pierce) at 4 °C under constant rotation. Beads were washed four times in lysis buffer, and bound proteins were eluted with 2× loading buffer. Proteins were separated by SDS-PAGE and immunoblotted (primary antibodies used: anti-hNET (1:1000) (NET 17–01; Mab-Technologies) to detect wild-type NET; anti-hDAT (1:1000) (Mab369; Millipore) to detect DAT(1–59)/NET; monoclonal anti-DAT (1:500) (Santa Cruz Biotechnology) to detect wild-type DAT and NET(1–128)/DAT; and HRP-conjugated anti-β-actin (1:10 000) (Sigma) in 5% milk powder in PBS plus 0.05% Tween; secondary antibodies used: HRP-conjugated antimouse, HRP-conjugated anti-rat and HRP-conjugated anti-goat (all 1:1000) (Pierce) in 5% milk powder in PBS plus 0.05% Tween). To estimate the relative amounts of transporters present in the blots, the transporter bands were scanned, and the band densities were quantified with ImageJ software.
JHC 1-64 Enables Specific Labeling of Endogenous NET in SCG Neurons—We first assessed whether the fluorescent cocaine analogue, JHC 1-64, could be used as a tool to specifically label surface-resident NET in primary neurons. JHC 1-64 contains a rhodamine moiety, extending from the tropane nitrogen of the parent compound 2β-carbomethoxy-3β-(3,4-dichlorophenyl)tropane with an ethylamino linker (35). We have previously used JHC 1-64 to visualize cell surface expression and internalization of both DAT and SERT (20, 22, 24, 35). Importantly, JHC 1-64 also displays high affinity for NET, as reflected by the potency by which JHC 1-64 inhibits [3H]dopamine uptake in NET-expressing cells (35).

To test whether JHC 1-64 allowed labeling for native NET, we prepared primary cultures of sympathetic SCG neurons known to express high levels of NET (30). These neurons are characterized by large round somata and a very extensive network of neuronal projections with multiple “boutons” probably representing presynaptic NE release sites (40). Labeling of the neurons with 50 nM JHC 1-64, followed by confocal live imaging, revealed a strong fluorescent signal in the rhodamine channel (>585 nm) corresponding to the plasma membrane of the somata (arrow in Fig. 1A) as well as of the neuronal projections and the boutons (Fig. 1A, arrowhead). In general, JHC 1-64 labeling demonstrated the strongest intensity on the plasma membrane of the neuronal projections and the boutons. Consistent with specific visualization of NET, no labeling was observed of the glial cell layer, and preincubation with the NET inhibitor desipramine (100 μM) eliminated essentially all neuronal labeling (a typical SCG neuron is shown in Fig. 1A). The uniform and wide distribution of JHC 1-64-labeled NET in the SCG neurons is further illustrated by the z-scan through live noradrenergic neurons shown in Fig. 1B. The z-scan also highlights the presence of NET in the plasma membrane of the neuronal somata. Notably, NET expression in the plasma membrane of the somata has not been reported in previous studies (16, 40, 41). Taken together, the results suggest that JHC 1-64 efficiently labels endogenous NET in cultured noradrenergic neurons with high signal/noise ratio.

To substantiate that the neurons labeled with JHC 1-64 possessed features of noradrenergic neurons, we co-stained JHC 1-64-labeled SCG neurons with antibodies directed against noradrenergic cell markers. JHC 1-64 labeling was maintained in the fixed and permeabilized neurons, although the staining observed was somewhat weaker and more punctuated compared with live neurons, which most likely can be ascribed to the fixation and permeabilization procedure. JHC 1-64 labeling
was detected in the plasma membrane of the somata as well as the boutons of all neurons positive for NET immunoreactivity (Fig. 1C, top). Of note, the NET antibody also revealed considerable intracellular immunostaining corresponding to both somata and boutons (Fig. 1C, top). Finally, we found strong surface labeling with JHC 1-64 of tyrosine hydroxylase-positive neurons (Fig. 1C, bottom). The tyrosine hydroxylase immunoreactivity did not overlay with signal of JHC 1-64, because tyrosine hydroxylase staining appeared intracellular both in the somata and boutons.

**NET Is Constitutively Internalized in SCG Neurons and Phorbol Esters Increase Internalization**—To determine whether NET undergoes constitutive endocytosis in noradrenergic neurons, surface-expressed NET was labeled with 50 nM JHC 1-64 at 4 °C followed by incubation at 37 °C for 1 h to allow for internalization. Consistent with marked constitutive internalization of NET, confocal live imaging revealed accumulation of multiple JHC 1-64-positive intracellular vesicular structures that were not seen in cells kept at 4 °C (Fig. 2A). Internalized, JHC 1-64-labeled NET was found not only in the somata and proximal extensions but was also clearly visible in the cytoplasm of the boutons (Fig. 2A). Indeed, quantification of confocal sections of live SCG neurons showed significantly higher internalization of JHC 1-64-labeled NET upon incubation at 37 °C (constitutive internalization) compared with the 4 °C control (Fig. 2B). The intracellular accumulation of JHC 1-64-labeled NET was further enhanced upon incubation with the phorbol ester PMA, which activates protein kinase C as well as other kinases (11, 13, 14, 42) (Fig. 2A). Quantification of single sections demonstrated that PMA treatment significantly increased the amount of intracellular JHC 1-64-labeled NET (Fig. 2B).

This finding was supported by [3H]dopamine uptake experiments demonstrating significant decreases in NET uptake capacity in response to PMA, compared with vehicle-treated SCG neurons (Fig. 2C).

**Internalized NET Colocalizes with Rab11-positive Recycling Endosomes and Inhibition of Rab11 Attenuates NET Function**—To investigate the fate of internalized JHC 1-64-labeled NET, we transfected cultured SCG neurons with lentivirus encoding different endosomal markers tagged with EGFP (i.e. EGFP-Rab4, -Rab7, and -Rab11). High transduction efficiency was obtained, and the EGFP-tagged Rab proteins exhibited distinct cellular distribution in agreement with previous reports (43–45). In the transfected neurons, we observed similar clear intracellular accumulation of JHC 1-64-labeled NET after 1 h of incubation at 37 °C as in non-transduced neurons (Fig. 3A and B). Imaging of the neurons revealed strikingly different colocalization patterns between the JHC 1-64-labeled NET and the individual Rab proteins. In the somata and proximal extensions, EGFP-Rab4, a marker of early endosomes and the “short loop” recycling pathway, was present in vesicular structures that showed very little overlap with internalized JHC 1-64-labeled NET (Fig. 3A). Similarly, we observed rather scant overlap between JHC 1-64-labeled NET and the vesicular structures to which the fluorescent signals from EGFP-Rab7, a marker of late endosomes, were localized (Fig. 3A). In contrast, the overlap between JHC 1-64-labeled NET and the signal from EGFP-Rab11, which marks the “long loop” recycling pathway, was strong (Fig. 3A). These observations were supported by quantification using Van Steensel’s cross-correlation analysis, in which the Pearson cross-correlation is calculated as the signal from one channel (i.e. JHC 1-64) is shifted relative to the signal from the other channel (i.e. EGFP) in the x direction pixel per pixel (36–38). The Pearson correlation was plotted as a function of a 20-pixel shift in each x direction of one of the images compared with the other. Consistent with strong colocalization of JHC 1-64-labeled NET with EGFP-Rab11, the cross-correlation function peaked at $\Delta x \approx 0$ with a value of $-0.4$ (Fig. 3A).
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A

JHC 1-64  EGFP  Merge

EGFP-Rab4

EGFP-Rab7

EGFP-Rab11

B

Somata

Pearson cross correlation

(JHC 1-64)

Rab11  Rab7  Rab4

Somata

Pearson cross correlation

(JHC 1-64)

Rab11  Rab7  Rab4

Boutons

Pearson cross correlation

(JHC 1-64)

Rab11  Rab7  Rab4

Lysotracker

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The peak value for EGFP-Rab7 was lower, and no peak was seen for EGFP-Rab4 (Fig. 3A).

In the boutons, we also observed strong overlap between internalized JHC 1-64-labeled NET and EGFP-Rab11-positive endosomes and little overlap with EGFP-Rab7 (Fig. 3B). Note that expression of EGFP-Rab7 is not unexpected in the axonal compartment, where Rab7 is believed to play a role in retrograde transport (46). For EGFP-Rab4, we interestingly observed an apparent higher degree of colocalization with internalized NET in the boutons compared with the somata and proximal extensions (Fig. 3B). Again, these findings were supported by cross-correlation analysis showing the highest peak value for EGFP-Rab11, hardly any peak for EGFP-Rab7, and a peak value of −0.22 for EGFP-Rab4 (Fig. 3B).

To further substantiate the distinct postendocytic fate of NET, we investigated the colocalization of internalized JHC 1-64-labeled NET with the lysosomal marker LysoTracker that has previously been shown to colocalize with constitutively internalized DAT and SERT (22, 24, 49). Consistent with sorting to a recycling pathway and not to degradation, we observed by live confocal imaging limited colocalization of internalized JHC 1-64-labeled NET with LysoTracker (Fig. 3C).

In light of the robust constitutive internalization, the localization of internalized NET to Rab11-positive endosomes suggests that recycling of NET via the long loop recycling pathway might be critical for maintenance of steady state NET surface levels and consequently NET function. To assess this, we mitigated the function of the Rab11 compartment by overexpressing the dominant negative Rab11 mutant, Rab11 S25N. Rab11 S25N mimics the GDP-bound, inactive form of Rab11 and is commonly used to inhibit Rab11-mediated recycling (16, 50–54). It has been shown that Rab11 S25N can significantly change the morphology of endocytic recycling endosomes and decrease their colocalization with cargo (54). Indeed, we observed a prominent change in the localization of recycling endosomes upon expression of Rab11 S25N (i.e. the disrupted recycling endocytic compartment was localized more peripherally and was dispersed) (compare EGFP-Rab11 with EGFP-Rab11 S25N in Fig. 4A). The overexpression of EGFP-Rab11 S25N resulted moreover in a significant decrease in NET transport capacity compared with neurons only overexpressing EGFP, as illustrated by reduced [3H]DA uptake (Fig. 4B). Overexpression of EGFPRab11, as well as of EGFP-Rab7, did not alter NET transport capacity (Fig. 4B). A role of the Rab11 pathway was further supported by experiments in transfected CAD cells, where co-expression of NET with Rab11 S25N, caused a significant decrease in [3H]DA uptake (Vmax = 72 ± 6% of control (mean ± S.E.), n = 4) compared with cells co-expressing EGFP-Rab11.

FIGURE 3. Internalized NET is primarily sorted to Rab11-positive recycling endosomes. SCG neurons were transduced with lentivirus encoding EGFP-Rab4, -Rab7, or -Rab11 at 4–5 days in vitro. The analysis of postendocytic sorting of NET was performed after 8–11 days in vitro by labeling surface-expressed NET with JHC 1-64 and allowing for internalization for 1 h at 37 °C. Subsequently, the live cells were analyzed at 37 °C by confocal imaging. A, left, confocal live images of internalized JHC 1-64-labeled NET in neuronal somata showing colocalization with EGFP-Rab11-positive endosomes and limited association with EGFP-Rab4 and EGFP-Rab7 positive endosomes. Scale bar, 10 μm. Right, quantification of colocalization between internalized JHC 1-64-labeled NET and EGFP-tagged endosomal markers in neuronal somata using Van Steensel’s cross-correlation function that reports the Pearson cross-correlation as a function of the relative movement of the two channels with respect to each other. The data support strong colocalization of internalized JHC 1-64-labeled NET with EGFP-Rab11, less colocalization with EGFP-Rab7, and no colocalization with EGFP-Rab4 (see “Experimental Procedures” for further details). B, left, confocal live images of internalized JHC 1-64-labeled NET in boutons of SCG neurons showing colocalization with EGFP-Rab11-positive endosomes, some colocalization with EGFP-Rab4-positive endosomes, and limited association with EGFP-Rab7-positive endosomes. Yellow arrowheads, examples of colocalized vesicles. White arrowheads, examples of non-colocalized vesicles. Scale bar, 5 μm. Images are representative of at least three independent experiments. Right, quantification of colocalization using Van Steensel’s cross-correlation function. The data support strong colocalization of internalized JHC 1-64-labeled NET with EGFP-Rab11, some colocalization with EGFP-Rab4, and little colocalization with EGFP-Rab7 (see “Experimental Procedures” for further details). C, constitutively internalized NET demonstrates little colocalization with the lysosomal marker, LysoTracker Green. SCG neurons labeled with JHC 1-64 were incubated with LysoTracker Green during the last 15 min of the internalization period (1 h) at 37 °C, and the live cells were imaged at 37 °C using an LSM 510 confocal microscope. Scale bar, 10 μm. Images are representative of at least three independent experiments.

FIGURE 4. Prolonged inhibition of Rab11 attenuates NET function. A, postendocytic sorting of NET in SCG neurons transduced with lentivirus encoding EGFP-tagged wild-type Rab11 or dominant negative Rab11 (Rab11 S25N). The analysis was done by labeling surface-expressed NET with JHC 1-64 followed by internalization for 1 h at 37 °C. Subsequently, the live cells were imaged at 37 °C using an LSM 510 confocal microscope. Both panels show representative somata. Overexpression of the EGFP-Rab11 S25N compared with EGFP-Rab11 changed the morphology of the endocytic recycling compartment. Scale bar, 10 μm. B, lentivirus-mediated transduction of Rab11 S25N resulted in a significant decrease in the [3H]DA uptake rate compared with neurons overexpressing EGFP. Data are means ± S.E. (error bars), n = 3–5, p < 0.05, one-way ANOVA, Dunn’s multiple-comparison test.
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**FIGURE 5. NET and DAT have distinct postendocytic fate in transfected CAD cells.** A, confocal live cell images and corresponding transmission images of untransfected CAD cells (left), cells transiently expressing NET (middle), or cells transiently expressing DAT (right). Cells were incubated with JHC 1-64 (50 nM) for 20 min at 4 °C before imaging. Scale bar, 10 μm. A, confocal live microscopy images of internalized JHC 1-64-labeled NET exhibiting strong colocalization with EGFP-Rab11-positive endosomes and limited association with EGFP-Rab7-positive endosomes. B, quantification of colocalization using Van Steensel’s cross-correlation function. The data support strong colocalization of internalized JHC 1-64-labeled NET with EGFP-Rab11 and less colocalization with EGFP-Rab7 (see "Experimental Procedures" for further details). C, confocal live microscopy images of internalized JHC 1-64-labeled DAT exhibiting limited colocalization with EGFP-Rab11-positive endosomes and stronger association with EGFP-Rab7-positive endosomes. Scale bar, 10 μm. D, quantification of colocalization using Van Steensel’s cross-correlation function. The data support the strongest colocalization of internalized JHC 1-64-labeled NET with EGFP-Rab7 and less colocalization with EGFP-Rab7 (see “Experimental Procedures” for further details). Error bars, S.E.

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ing NET and wild-type Rab11. Co-expression of NET with dominant negative Rab4 (Rab4 S25N), however, did not cause any change in NET uptake rate compared with the cells expressing wild-type Rab 4 (V_max = 100 ± 9% of control (mean ± S.E.), n = 3). Altogether, our results suggest that to a major extent, constitutively internalized NET is sorted to the Rab11 recycling compartment in SCG neurons and that impairment of the Rab11 pathway results in decreased NET function.

**NET and DAT Have Distinct Trafficking Properties**—We have recently characterized in detail the postendocytic fate of constitutively internalized DAT and SERT, which both appeared to be sorted largely to late endosomes and lysosomal degradation (21, 22). Thus, we did not expect that constitutively internalized NET would be sorted primarily to the Rab11 recycling compartment in primary neurons. To further characterize the fate of NET, we wanted to directly compare NET with DAT; therefore, we turned to a heterologous expression system. To distinguish between the Rab11 recycling compartment and the Rab7 late endosomal compartment, NET and DAT were transiently transfected into catecholaminergic CAD cells (55, 56). Importantly, we observed no background labeling of these cells with JHC 1-64 (Fig. 5A). In contrast, we observed strong plasma membrane labeling of CAD cells transiently expressing either NET or DAT (Fig. 5A). Next, we co-expressed NET or DAT with either EGFP-Rab7 or EGFP-Rab11. The cells were labeled with JHC 1-64, followed by incubation at 37 °C for 1 h to allow constitutive internalization. Similarly to our findings in noradrenergic neurons, we observed strong JHC 1-64 labeling of the plasma membrane in NET expressing CAD cells. Moreover, we observed accumulation of multiple intracellular JHC 1-64-positive vesicles during the 37 °C internalization period, consistent with constitutive internalization. Such vesicular structures were not observed in cells kept at 4 °C, and no labeling or accumulation of JHC 1-64 were seen in the presence of the NET inhibitor desipramine (data not shown). Importantly, internalized, JHC 1-64-labeled NET colocalized prominently with EGFP-Rab11 but only to a limited extent with EGFP-Rab7 (Fig. 5A). This was supported by cross-correlation analysis indicating marked colocalization with EGFP-Rab11 and less with EGFP-Rab7 (Fig. 5B). In agreement with our previous findings for DAT (24), we also observed strong JHC 1-64 labeling of DAT-expressing CAD cells and intracellular accumulation of JHC 1-64-labeled DAT upon incubation at 37 °C (Fig. 5C). Internalized JHC 1-64-labeled DAT, however, showed substan-
In independent experiments, JHC 1-64 vesicles that appeared more perinuclear (2.4 ± 0.1 µm) versus 4.2 ± 1.2 µm (NET) (means ± S.E.) (Fig. 6B). Moreover, our images indicated that the constitutive NET internalization was more intense compared the DAT internalization. We quantified accordingly the fraction of JHC 1-64-labeled NET and DAT, which confirmed that a significantly higher fraction of NET was internalized compared with DAT in CAD cells transiently expressing NET and DAT (Fig. 6C).

**Internalized TacNET Colocalizes with Rab11 Endosomes**

We showed previously that JHC 1-64 has a very slow off-rate from DAT, making it a reliable marker of DAT trafficking (20). To assess whether the off-rate from NET likewise was slow, we used a procedure similar to the one that we used previously for DAT (20). We assessed the capability of JHC 1-64 to inhibit [3H]DA uptake in NET-expressing CAD cells upon prolonged incubation and wash-outs to permit dissociation and preclude reassociation of JHC 1-64. The cells were incubated with 500 nM JHC 1-64 or vehicle for 30 min, followed by washing and incubation for 20, 40, 60, or 90 min at 37°C. After two additional washes, [3H]DA uptake was measured, and we found that uptake was still inhibited to almost the same extent (~90%) as it was when measuring uptake with no washout immediately after the addition of 500 nM JHC 1-64 (data now shown). This strongly suggests that the off-rate of JHC 1-64 is very slow also from NET and that JHC 1-64 therefore should be a reliable tool for tracking both NET and DAT in live cells even over extended time periods.

We wanted subsequently to exclude the possibility that binding of JHC 1-64 to the transporter affects the internalization and postendocytic sorting properties of NET described above. We therefore generated a fusion construct where the intracellular N terminus of NET was fused to the intracellular N terminus of NET (Fig. 7A). Because Tac contains an N-terminal M1 FLAG recognition epitope is available. Previously, we have made similar constructs for DAT and SERT (TacDAT and TacSERT) that have proven very useful for studying the trafficking properties of these transporters (21, 22). Similar to TacDAT and TacSERT, TacNET exhibited clear functional activity, although its maximum uptake capacity was somewhat reduced as compared with wild-type NET (Km = 0.30 ± 0.06 µM for TacNET versus 0.35 ± 0.08 µM for wild-type NET, Vmax = 23 ± 5% of wild-type NET (means ± S.E.), n = 3). For the antibody-based internalization assay, CAD cells expressing TacNET were first incubated with M1 antibody at 4°C to label surface TacNET. Confocal imaging of the cells upon fixation (no permeabilization) and immunostaining with Alexa Fluor 568-conjugated secondary antibody revealed a fluorescent signal corresponding to the plasma membrane upon fixation (no permeabilization) and immunostaining with Alexa Fluor 568-conjugated secondary antibody revealed a fluorescent signal corresponding to the plasma membrane.
membrane, suggesting that TacNET was expressed at the cell surface (Fig. 7B). We then analyzed cells after 1 h at 37 °C and observed marked intracellular accumulation of TacNET immunosignal in vesicular structures, consistent with constitutive internalization (Fig. 7B) as was observed for the JHC 1-64-bound transporter. This accumulation appeared, as would be expected, generally stronger upon treatment of the cells with PMA during the 1-h internalization period (Fig. 7B). We also performed experiments where we visualized surface TacNET using both M1 immunostaining with Alexa Fluor 488-conjugated secondary antibody and JHC 1-64 at the same time by incubating the cells with JHC 1-64 for 20 min at 4 °C prior to
incubation with the M1 antibody at 4 °C. In addition, we visualized TacNET internalization by labeling the surface-expressed TacNET with antibodies (M1 and Alexa Fluor 488-conjugated secondary antibody) and JHC 1-64 following internalization for 1 h at 37 °C. Importantly, the fluorescent signals from the two different channels almost completely overlapped (Fig. 7C).

The presence of the FLAG epitope in TacNET furthermore enabled the employment of an ELISA-based assay to quantify surface expression of the transporter (24). According to this assay, incubation of the cells for 1 h at 37 °C with PMA caused a significant decrease in TacNET surface expression (Fig. 7D), confirming our observations by confocal imaging that PMA indeed causes redistribution of TacNET away from the plasma membrane. In contrast, incubation of the cells with JHC 1-64 at a saturating concentration (500 nM) for the same period of time did not cause any change in TacNET surface expression (Fig. 7D). This indicates that JHC 1-64 does not by itself affect transporter trafficking.

To obtain evidence that labeling of the transporter with JHC 1-64 also did not affect the postendocytic fate of NET, we performed antibody-based internalization assays in CAD cells expressing TacNET together with either EGFP-Rab7 or EGFP-Rab11. Consistent with the live imaging of native NET, constitutively internalized M1-labeled TacNET exhibited extensive colocalization with EGFP-Rab11 and showed little colocalization with EGFP-Rab7 (Fig. 7E). Thus, our TacNET experiments suggest that JHC 1-64 labeling does not affect the basic trafficking properties of the transporter.

Inhibition of Recycling Increases NET, but Not DAT, Intracellular Accumulation—Next, we employed a biochemical internalization assay (reversible biotinylation) to further investigate the trafficking properties of NET and DAT independent of fluorescent confocal imaging. CAD cells expressing non-tagged NET were biotinylated, using the reducible biotinylation reagent sulfo-NHS-SS-biotin, whereupon internalization was permitted by incubating the cells at 37 °C for 1 h. By treating the cells with the reducing agent MesNa, biotin residing on the cell surface was removed, and subsequently, intracellular biotinylated proteins were isolated and analyzed by Western blotting. The results strongly supported extensive internalization of NET by revealing significantly stronger NET immunoreactivity for cells incubated at 37 °C for 1 h as compared with cells kept at 4 °C (Fig. 7A, lanes 2 and 3). Interestingly, inhibition of lysosomal degradation by the lysosomal protease inhibitor leupeptin during the internalization period did not cause any further increase in the amount of NET immunoreactivity and thus intracellular NET levels. However, additional inhibition of recycling by monensin, a cation ionophore blocking recycling (57), caused a significant increase in NET levels (Fig. 8A), supporting our results from the imaging experiments indicating that NET is primarily recycled and not targeted to degradation upon internalization.

Our confocal fluorescence imaging results were further supported by reversible biotinylation experiments performed on DAT-expressing CAD cells. The experiments showed that DAT undergoes constitutive internalization as well (i.e. we found stronger DAT immunoreactivity for cells incubated at 37 °C for 1 h as compared with cells kept at 4 °C (Fig. 8B, lanes 2 and 3). Importantly, we observed different effects of leupeptin and monensin on DAT as compared with NET; inhibition of
lyosomal degradation (leupeptin) showed a significantly increased intracellular accumulation of DAT, and additional inhibition of recycling (leupeptin and monensin) did not further increase the DAT immunosignal (Fig. 8B), in agreement with DAT being sorted to a larger degree to degradation than to recycling. It is also interesting to note that the apparent degree of internalization was lower for DAT compared with NET as determined with recycling and degradation blocked by leupeptin and monensin (12 ± 1 versus 41 ± 2%, respectively, n = 3–4; Fig. 8, compare A and B).

The NET and DAT N Termini Are Critical for Internalization and Postendocytic Sorting—Our findings suggest strikingly different trafficking itineraries for NET compared with DAT. To test whether specific NET or DAT domains encoded information sufficient to direct internalized transporter to specific endosomal compartments, we assessed the endocytic properties of chimeras between NET and DAT where the N and C termini were exchanged between the two proteins. All chimeras into NET (DAT(1–59)/NET), we observed internalization properties more similar to those seen for wild-type DAT. The apparent rate of internalization was lower at control conditions and, although there was still an increase in the DAT(1–59)/NET immunosignal when the cells were incubated with both leupeptin and monensin, the increase was more modest (Fig. 8C). As a result, the degree of internalization, as determined with recycling and degradation blocked by leupeptin and monensin, was higher for wild-type NET compared with DAT(1–59)/NET (41 ± 2 versus 16 ± 2%, respectively, n = 3–4) (Fig. 8, A and C). Similarly, by introducing the NET N terminus into DAT (NET(1–128)/DAT), we observed internalization properties more similar to those seen for wild-type NET. Blocking lysosomal degradation with leupeptin modestly increased NET(1–128)/DAT immunosignal, whereas additional blocking of recycling with monensin had a significant effect on NET(1–128)/DAT immunosignal compared with control conditions (37 °C, no treatment) (Fig. 8D). As a result, the degree of internalization, as determined with recycling and degradation blocked by leupeptin and monensin, was higher for NET(1–128)/DAT compared with wild-type DAT (32 ± 8 versus 12 ± 1%, respectively (mean ± S.E.), n = 3–4) (Fig. 8, B and D).

Discussion

Despite the physiological and pharmacological significance of NET, the cellular processes that control the trafficking of the transporter to and from the plasma membrane are still rather poorly understood. Here, we provide novel insight into the trafficking itinerary of NET by employing several different approaches. In particular, we take advantage of the fluorescent cocaine analogue JHC 1-64, which provides a powerful tool enabling direct visualization of surface distribution, internalization, and postendocytic sorting of endogenous NET even in single boutons of live neurons. Importantly, JHC 1-64 itself is unlikely to have any significant effect on NET trafficking. By using the TacNET construct, which adds an extracellular FLAG antibody epitope to the transporter, we observed that JHC 1-64 had no detectable effect on NET surface levels. An antibody feeding assay results also showed that TacNET was internalized constitutively to the same Rab11-positive pathway as JHC 1-64-bound NET.

Labeling of live SCG neurons with JHC 1-64 revealed a uniform distribution of NET in the plasma membranes of the extensions and boutons. We also observed a clear labeling of the plasma membrane of the somata, which, to our knowledge, has not been described before. The role of NET in the somata remains to be determined; however, it is attractive to consider that the wide and uniform distribution of NET in SCG neurons reflects the importance of volume transmission for norepinephrine action (58). We should note that a similar uniform distribution in the plasma membrane of SCG neurons and other neuronal preparation was not observed in previous studies based on immunostaining with NET antibodies. Rather, these studies revealed more punctate expression (30, 40, 41, 59).

**TABLE 1**

| Transporter | $K_m$ | $V_{max}$ | % of NET |
|-------------|------|---------|----------|
| DAT         | 1.90 ± 0.15 | 400 ± 80 |          |
| NET         | 0.46 ± 0.12 | 100     |          |
| NET(1–128)/DAT | 0.64 ± 0.05 | 230 ± 48 |          |
| DAT/NET(590–617) | 1.90 ± 0.40 | 310 ± 44 |          |
| DAT(1–59)/NET | 1.18 ± 0.58 | 156 ± 19 |          |
| NET/DAT(591–620) | 0.48 ± 0.13 | 96 ± 9  |          |

The transport kinetics of dopamine were assessed in CAD cells transiently expressing wild-type and chimeric transporters. The $K_m$ and $V_{max}$ values were determined by nonlinear regression fits determined from 3–6 independent experiments performed in triplicates. $K_m$ values are reported as the mean ± S.E., and $V_{max}$ values are expressed relative to NET (mean ± S.D.).
Most likely, this punctate distribution is an artifact of the fixation and permeabilization procedure used for the immunostainings. Our immunostainings also revealed more punctate NET distribution compared with JHC 1-64 labeling of live cells, and fixation and permeabilization also caused the JHC 1-64 labeling to become more punctate (Fig. 1B). Of interest, both previous immunostainings and our own showed extensive intracellular NET immunosignals in the somata and in the boutons (16, 30, 40, 41). The presence of such large intracellular pools of NET indirectly suggests that the transporter is dynamically shuffled between intracellular compartments and the plasma membrane. Indeed, we obtained support for such extensive trafficking of NET by demonstrating fast constitutive internalization of NET not only in the cytoplasm of the somata and proximal extensions but also in the cytoplasm of single boutons that conceivably represent presynaptic release and
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The internalization was moreover increased in SCG neurons upon stimulation with PMA, demonstrating that NET expressed in neurons can be subject to regulation by protein kinase C and other kinases activated by phorbol esters. It was shown before that NET natively expressed in rodent trophoblasts undergoes endocytosis in response to PMA (14), and earlier studies have supported PMA-induced internalization of NET in transfected cells (11, 13, 14, 42). However, PMA-induced trafficking of endogenous NET in neurons has not been demonstrated before.

Upon sequestration from the plasma membrane of SCG neuronal somata, proximal extensions, and boutons, NET showed considerable colocalization with EGFP-tagged Rab11. Rab11 mediates cargo recycling back to the plasma membrane from the pericentriolar recycling endosomes and marks the long loop recycling pathway (52, 54). Altogether, our results suggest that constitutively internalized NET is mainly sorted to the long loop Rab11-dependent recycling pathway and only to a limited extent to degradation. Moreover, we found that the integrity of the Rab11-dependent pathway was important for NET function, because overexpression of a dominant negative mutant (Rab11 S25N) (54, 62) significantly reduced uptake capacity in the SCG neurons. Additional support for sorting of NET to recycling was obtained by expressing NET in CAD cells where confocal imaging and biotinylation experiments showed evidence for the same sorting pattern as in the SCG neurons. Of further interest, Matthies et al. (16) showed by use of a NET endodomain antibody that upon amphetamine-induced internalization, NET displays colocalization with both Rab4 and Rab11 in SCG neurons, indicating that amphetamine might increase NET internalization and sorting to both fast and slow recycling pathways (16). Indeed, we also observed some colocalization of internalized JHC 1-64-labeled NET with Rab4 in the boutons of the SGN neurons, indicating that NET also to some degree might recycle via the Rab4 pathway.

We and others have shown that the closely related SLC6 transporters DAT and SERT, like NET, undergo considerable constitutive internalization (18, 20, 22, 23), although a recent study has indicated that DAT only might undergo modest constitutive internalization (18, 20, 22, 23). However, recent findings and in contrast to NET, DAT and SERT are mainly sorted to late endosomes and lysosomes, and, if recycled, they are more likely to follow the Rab4 short loop recycling pathway (22, 24). In contrast, the neuronal glycine transporter GlyT2 was shown to colocalize with Rab11 and to be efficiently recycled (19). The postendocytic fate of DAT, however, has been debated (18, 23, 24), but our direct comparison of NET and DAT in CAD cells revealed clear differences between the two proteins. In contrast to NET, JHC 1-64-labeled, internalized DAT colocalized mostly with EGFP-Rab7 but somewhat less with EGFP-Rab11 and showed a markedly lower degree of internalization. Importantly, this difference in internalization kinetics is not likely to explain the differential colocalization pattern observed by confocal imaging. Reversible biotinylation experiments indicated that NET is indeed more likely to recycle to a larger degree than DAT and that DAT function, in contrast to NET function, seems to be unaffected by dominant negative Rab11 mutants (24, 64).

To identify structural elements governing postendocytic sorting and/or internalization rates, we undertook a chimeric approach as previously done to identify domains critical for substrate selectivity and ion dependence in NET and DAT (26, 65). The results suggest that the N termini of NET play critical roles in determining the rate and postendocytic fate of the transporter. Only little is known about intrinsic sorting signals in NET and DAT as well as in other SLC6 family transporters. NET was found to sort to the basolateral membrane, and DAT was found to sort to the apical plasma membrane in polarized epithelial cells. In addition, it was observed that this differential sorting pattern was determined by the N termini (66). An apical localization signal for GAT1 (GABA transporter 1) also appeared to reside in the N terminus, although a basolateral localization signal for GAT2 was located to the C terminus (67–69). The involvement of the N termini in differential targeting in epithelial cells is interesting, but to what degree it directly relates to our observations remains to be investigated. Our insight into structural elements driving constitutive and regulated internalization of SLC6 family transporters is also limited. A short sequence in the DAT C terminus (FREKLAYAIA) was shown to be important for constitutive and PKC-mediated internalization of DAT, but the motif is only partially conserved in NET (70). Interestingly, a short N-terminal segment, which is proximal to TM1 and involved in regulation of transporter conformation (71), was demonstrated to affect constitutive DAT internalization (72), but because the segment is conserved between NET and DAT, it cannot determine the differences between the two transporters observed in this study. It is also unlikely that differences in N-terminal ubiquitination would determine the difference between DAT and NET. N-terminal ubiquitination can regulate PMA-induced DAT internalization (73) but appeared not to affect constitutive internalization and sorting (24).

Striking differences in trafficking itineraries have been observed for other homologous membrane proteins. The G protein-coupled δ- and μ-opioid receptors are differentially sorted between degradative and recycling pathways following agonist-induced endocytosis (74, 75). In a similar way, dopamine D2 receptors are sorted to degradation, whereas D1 receptors are recycled after agonist-induced endocytosis (76, 77). Notably, sorting of D2 receptors to degradation is not determined by the absence of a recycling signal but by binding of the receptor to GASP (G protein-coupled receptor-associated sorting protein). It would be interesting to see whether similar protein-protein interactions control DAT/NET internalization and sorting. It will also be interesting to clarify the precise biological significance of the distinct trafficking itineraries for NET and DAT, and whether or not these can be related to neuropsychiatric disease or chronic drug use. For example, differential up- and down-regulation of brain NET and DAT, respectively, was reported upon chronic cocaine or methamphetamine exposure in humans (47, 78). For homologous

3 J. Eriksen and U. Gether, unpublished observation.
receptors, distinct sorting mechanisms are believed to enable differential physiological adaptation to agonist exposure (48, 74, 75). For homologous transport proteins like NET and DAT, it is tempting to suggest that the rapid cycling of NET enables faster adjustment of transporter availability to the immediate cellular needs. It is possible that such tighter control may be required in, for example, peripheral noradrenergic neurons to ensure proper physiological responsiveness of the sympathetic nervous system.

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