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Exogenous α-Synuclein Decreases Raft Partitioning of Ca_v2.2 Channels Inducing Dopamine Release

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α-Synuclein is thought to regulate neurotransmitter release through multiple interactions with presynaptic proteins, cytoskeletal elements, ion channels, and synaptic vesicles membrane. α-Synuclein is abundant in the presynaptic compartment, and its release from neurons and glia has been described as responsible for spreading of α-synuclein-derived pathology. α-Synuclein-dependent dysregulation of neurotransmitter release might occur via its action on surface-exposed calcium channels. Here, we provide electrophysiological and biochemical evidence to show that α-synuclein, applied to rat neurons in culture or striatal slices, selectively activates Ca_v2.2 channels, and said activation correlates with increased neurotransmitter release. Furthermore, in vivo perfusion of α-synuclein into the striatum also leads to acute dopamine release. We further demonstrate that α-synuclein reduces the amount of plasma membrane cholesterol and alters the partitioning of Ca_v2.2 channels, which move from raft to cholesterol-poor areas of the plasma membrane. We provide evidence for a novel mechanism through which α-synuclein acts from the extracellular milieu to modulate neurotransmitter release and propose a unifying hypothesis for the mechanism of α-synuclein action on multiple targets: the reorganization of plasma membrane microdomains.

Key words: Ca_v2.2 channels; dopamine release; extracellular alpha-synuclein; lipid rafts

Introduction

α-Synuclein (Syn), an unfolded soluble cytosolic protein, has been reported to accumulate in Lewy bodies, which are intraneuronal aggregates pathognomonic of Parkinson’s disease (PD) (Spillantini et al., 1997). Although the aggregation pathways and the toxicity of aggregated forms of Syn have been studied extensively, little is known about the role of monomeric Syn in the early phases of PD. Syn is highly enriched in presynaptic terminals, but a substantial fraction of this protein is released in extracellular fluids in healthy and PD subjects (Borghi et al., 2000; El-Agnaf et al., 2003). Importantly, healthy dopaminergic grafts implanted in the striatum undergo degeneration accompanied by Syn-containing Lewy bodies (Kordower et al., 2008a,b), suggesting a potential role of secreted extracellular Syn in the onset of the disorder. Although the direct transfer of Syn between neurons has been demonstrated (Hansen et al., 2011; Volpicelli-Daley et al., 2011), little is currently known about the process of release of Syn and its physiopathological function. Duplications and triplications of the SNCA gene encoding α-synuclein are present in familial forms of PD. Recently, the SNCA gene was identified as a risk locus in genome-wide association studies for PD, highlighting the importance of wild-type Syn levels as pathogenic factor also in the sporadic form of PD (Satake et al., 2009; Vendel et al., 2010).

Maintenance of calcium homeostasis through the regulation of presynaptic calcium channels and intracellular calcium stores is fundamental for correct synaptic activity. Calcium dyshomeostasis has been proposed recently as the primary age-related condition driving neurodegeneration in the sporadic form of PD, in which selective subpopulations of neurons undergo degeneration. Dopaminergic neurons expressing higher levels of calcium binding proteins, such as calbindin, calretinin, and parvalbumin, seem to be resistant to degeneration in PD (Damier et al., 1999; Kim et al., 2000; Sóos et al., 2004). Dysfunction in the ion channels and receptors regulating synaptic communication may contribute to the cognitive decline, neurological and motor symptoms associated with chronic disorders (Walsh and Selkoe,
In neurons, presynaptic voltage-gated calcium channels (Ca_{v}) 2.1 and Ca_{v} 2.2 are predominantly responsible for coupling neuronal depolarization to neurotransmitter release. In particular, Ca_{v} 2.2 underlies dopamine (DA) release in the basal ganglia (Bergquist et al., 1998). In SH-SY5Y cells, an increase in calcium influx through Ca_{v} has been suggested to be caused by increased expression of Syn or by extracellular Syn (Hettiarachchi et al., 2009; Melachroinou et al., 2013). In synaptic vesicles, an increase in calcium influx through Ca_{v} 2.2 occurs during incubation with micromolar Syn concentrations; such an effect occurred independently of free radical generation and was suggested to arise through a direct interaction of Syn with Cav2.2 (Adamczyk and Strosznajder, 2006). Mouse models of Syn-induced neurodegeneration exhibit a decreased DA release, similar to that observed in PD patients (Decressac et al., 2012; Lundblad et al., 2012; Nuber et al., 2013). However, increased tonic DA levels attributable to early synaptic dysfunction, preceding the death of dopaminergic neurons and leading to a possible disruption of corticostriatal transmission (Wu et al., 2010), were found in Syn transgenic mice at a young age (Lam et al., 2011), similar to that assessed in other PD models (Goldberg et al., 2003).

Here we evaluate the alteration of synaptic activity attributable to the presence of raised concentrations of Syn in the extracellular space. We analyze in detail the effect of extracellularly applied monomeric Syn on depolarization-evoked calcium influx and Ca_{v} 2.2 calcium current density in neurons and on neurotransmitter release in vitro and in vivo, identifying the molecular mechanism of Syn action as a decrease in raft partitioning of Ca_{v} 2.2 to induce DA release. Such a Syn regulatory mechanism may have important consequences in PD and related synucleopathies.

### Materials and Methods

#### Reagents

All chemicals, unless otherwise stated were purchased from Sigma-Aldrich. The following primary antibodies were used: anti-Cav2.2 α1β (A-11; Santa Cruz Biotechnology catalog sc-271010, RRID: AB_10611357), anti-G-protein β1 (C-16; Santa Cruz Biotechnology catalog sc-379, RRID:AB_2111823), anti-α2δ1 subunit (Santa Cruz Biotechnology catalog sc-133436, RRID:AB_10609764), anti-Hsp90 (Santa Cruz Biotechnology catalog sc-59578, RRID:AB_2295578), anti-flotillin-1 (BD Biosciences catalog #610821, RRID:AB_398140), anti-caveolin-1 (BD Biosciences catalog #610099, RRID:AB_397471), anti-α-synuclein (BD Biosciences catalog #610787, RRID:AB_398108), anti-Na+/K+ ATPase (EMD Millipore catalog #05-369, RRID:AB_309699), anti-Ca_{v} β2 subunit (Sigma-Aldrich catalog #C5738, RRID:AB_258910), anti-synaptophysin (Synaptic Systems catalog #101 011, RRID:AB_887824), anti-synaptin 1A antibody (Santa Cruz Biotechnology catalog sc-58299, RRID:AB_785887), anti-Rab11 (Life Technologies catalog #700184, RRID:AB_10562415), and anti-Ca_{v} 1.2 (Hell et al., 1996). Syn peptides (amino acids 12–23, 34–45, and 61–78) were obtained from Primmibiotech.

#### Rat primary cortical cultures

All experiments were performed in accordance with the Italian Ministry of Health. Primary cultures of rat embryonic cortical neurons were prepared as described previously (Brewer, 1997). Briefly, E18 Sprague Dawley rat embryos (Charles River) were decapitated, and cortices were rapidly dissected and dissociated by incubation in Neurobasal medium (Life Technologies) containing 20 U/ml papain, 1.1 mg/ml cysteine, and 125 μg/ml DNase for 30 min at 37°C. After cell dissociation, cortical neurons were plated in either six-well dishes at a density of 1 × 10^{6} cells/ml on glass coverslips coated with poly-L-lysine (0.1 μg/ml) or 100 mm Petri dishes coated with poly-L-lysine (0.01 μg/ml) at a density of 5 × 10^{8} cells per dish. Cells were cultured in Neurobasal medium supplemented with B-27, GlutaMAX, and penicillin–streptomycin (all from Life Technologies) and maintained at 37°C in 5% CO_{2}. In all experiments, neurons were used between 7 and 14 d in vitro (DIV).

#### Syn purification

The cDNA encoding the full-length human wild-type Syn inserted in the pET21d plasmid was a kind gift from Dr. B. Lauling (Columbia University, New York, NY). Purification was performed as described previously (Martinez et al., 2003). Bacteria were induced during the exponential phase with 1 mm isopropyl β-D-1-thiogalactopyranoside for 2 h and harvested by centrifugation. The pellet was solubilized in 20 mM HEPES/KOH, pH 7.2, and 100 mM KCI (buffer A) and heated for 5 min at 90°C. The cell lysate was centrifuged at 72,000 × g for 30 min, and the supernatant was loaded on a HiTrap monoQ column (GE Healthcare). Syn was eluted with a liner gradient of KCl from 100 to 500 mM, and the fractions of interest were concentrated using a Centricron centrifugal filter (Millipore) before loading on a Superose 12 column in buffer A (GE Healthcare). The fractions containing Syn were pooled, concentrated, and stored at −80°C. To control Syn purity, 1 mg of purified Syn was loaded on top of a Superdex 75 10/300 column (GE Healthcare) and eluted at 0.5 ml/min on an AKTA Purifier apparatus. Optical density (OD) was continuously measured at 280 nm.

#### Lipid raft isolation

All procedures were performed at 4°C. Lipid raft isolation was performed on cortical neurons at 10 DIV as described previously (Butchbach et al., 2004). Briefly, primary cortical neurons (5 × 10^{6} neurons per dish) were incubated for 1 h at 4°C with 80 μl/dish PBS containing 1% of Triton X-100 and Complete Mini protease inhibitor (Roche Applied Science). The extract was added with sucrose to obtain a 40% final concentration. A total of 900 μl of the extract was placed on the bottom of an ultracentrifugation tube, and a discontinuous 5–30% sucrose gradient was formed on top of the extract. The gradient was then centrifuged at 47,000 rpm for 16 h at 4°C using a MLS 50 rotor (Beckman Coulter). Ten 0.5 ml fractions were collected from the top of each gradient. A total of 400 μl of the sample was added to 100 μl of 5% SDS-PAGE loading buffer (100 μl Tris, pH 7.5, 7.5 μl EDTA, 10% SDS, 10% β-mercaptoethanol, 50% glycerol, and 0.02% bromophenol blue).

#### Biotinylation procedure

The entire biotinylation procedure was performed at 4°C. Labeling of proteins of the plasma membrane was achieved by biotinylation of cortical neurons (10 DIV) and was a slight modification of the protocol reported previously (Butchbach et al., 2004). The cells were washed twice in PBS and once in HBSS and then incubated in HBSS containing 0.6 mg/ml EZ-link Sulfo-NHS-biotin (Thermo Fisher Scientific) for 15 min. The cells were then incubated in quenching buffer (5% BSA in HBSS) for 10 min, scraped, and recovered by centrifugation. Cell pellets were incubated for 1 h with constant shaking in PBS containing 1% Triton X-100 and Complete Mini protease inhibitor (Roche Applied Science). The biotinylated extract was separated in a sucrose gradient (as described above), the fraction was collected, and SDS was added to a final 0.5%. The biotinylated proteins present in each fraction were pulled down by incubation with 100 μl of Neuravidin beads (50% slurry, Thermo Fisher Scientific) for 2 h. Avidin beads were then recovered by centrifugation and washed four times in PBS. After washing, bound proteins were eluted with 1% SDS-PAGE loading buffer (20 μl Tris, pH 7.5, 1.5 μl EDTA, 2% SDS, 2% β-mercaptoethanol, 10% glycerol, and 0.004% bromophenol blue).

#### Immunoblot analysis

SDS-PAGE and electrotransfer of membranes were performed as described previously (Laemmli, 1970). Immunoreactive bands were detected by chemiluminescence, and images were acquired using an LAS AF 4000 apparatus (GE Healthcare). Bands intensities were measured by Image Quant analysis software (GE Healthcare).

#### Filipin quantitative staining

Filipin quantitative staining of fixed cells was performed accordingly to a modification of a commercially available kit (catalog #10009779; Cayman Chemicals). Briefly, cortical neurons (10 DIV) plated on 18 mm
covered sections were fixed with 4% (v/v) PFA, washed in PBS, and incubated for 1 h in PBS containing 1% Triton X-100 (filipin total staining) or PBS (filipin membrane staining). Next, coverslips were incubated for 1 h at room temperature in 10 mM PBS and 10% NGS (Thermo Fisher Scientific), containing 100 µg/ml filipin [freshly prepared from a 10 mg/ml stock solution (in EtOH) stored at −20°C]. Finally, coverslips were incubated for 30 min at room temperature in 500 µl of 1% SDS, and the fluorescence of the solution was measured on a LS55 fluorometer (PerkinElmer Life and Chemical Sciences).

**Fluo-4 imaging**

Rat embryonic cortical neurons were plated onto 25 mm coverslips at a density of 800 neurons/mm² and at 6–10 DIV were incubated with buffer A or 5 µM Syn for 2 h at 37°C. Neurons were then loaded with Fluo-4-AM (2.5 µM; Life Technologies) for 20 min at 37°C. Coverslips were then transferred into a perfusion chamber (Warner Instruments) and superfused with modified Krebs’ solution containing the following (in mM): 135 NaCl, 5 KCl, 1.5 CaCl₂, 0.75 MgCl₂, 0.75 KH₂PO₄, 1.5 MgSO₄, 5.6 glucose, and 20 Na-HEPES, pH 7.4 (adjusted with NaOH). NBQX at 10 µM, 50 µM t-APV, and 100 µM picrotoxin were added to the solution. Nifedipine at 10 µM, 300 nM α-conotoxin GVIA, or 100 nM α-conotoxin N30A (Bachem) was added as described. Fluo-4 fluorescence was recorded at 494 ± 10 nm wavelength at 1 Hz for 3 min. Basal fluorescence was recorded for 1 min, and then Ca²⁺ entry was elicited by a 60 s bath application of a modified Krebs’ solution containing 50 mM KCl with a reciprocal decrease in Na⁺ concentration. A single stimulation was performed per coverslip. Results were expressed as the ratio between the basal fluorescence and the maximum fluorescence of the neurons during 50 mM KCl stimulation. Data were normalized for each experiment to the respective control. Calcium imaging was performed on a Leica AF6000 microscope to acquire images at 1 frame/s.

**GCaMP3 imaging**

Rat embryonic cortical neurons were plated at a density of 400 neurons/mm² and at 10 DIV were transfected with SyGCaMP3 (GCaMP3 with synaptic physiology) (Ronzi et al., 2011) by electroporation. Calcium imaging was performed on a Leica AF6000 microscope to acquire images at 1 frame/s.

**Electrophysiology**

**Superior cervical ganglia neurons for patch-clamp recordings.** Superior cervical ganglia neurons (SCGNs) were acutely dissected from 3- to 6-week-old male Wistar rats according to the guidance of the UK Animals (Scientific Procedures) Act 1986 subject to University of Reading Local Ethics Panel review, as described previously (Bucci et al., 2011). Briefly, SCG were dissected and placed in cold L-15 medium (Sigma Aldrich) containing 5% penicillin–streptomycin (Life Technologies). The ganglia were then acutely dissociated in DMEM containing 0.5 mg/ml trypsin, 1 mg/ml collagenase, and 3.6 mg/ml glucose (all from Sigma Aldrich). After incubation at 37°C for 30–40 min, the digestion reaction was stopped by the addition of 84% Eagle’s MEM, 10% fetal calf serum, 5% horse serum (Lonza), and 1% penicillin–streptomycin (Gibco Invitrogen). After dissociation, cells were plated onto coverslips coated with poly-l-ornithine (Sigma-Aldrich) and incubated at 37°C (5% CO₂) before electrophysiological recordings.

**Whole-cell voltage-clamp recordings.** Whole-cell recordings from isolated SCGNs were performed as described previously (Bucci et al., 2011) using borosilicate glass electrodes with resistances between 5 and 7 MΩ when filled with an intracellular solution [in mM: 140 CsCl, 10 HEPES, 0.1 CaCl₂, 1.0 MgCl₂, 4.5 ATP, 0.2 Na-GTP, and 1.0 EGTA, pH 7.3 (adjusted with CsOH)] and an extracellular solution [in mM: 160 tetraethylammonium bromide, 10 HEPES, 1 KCl, 1 MgCl₂, and 4 glucose, with 10 mM Ba²⁺ as charge carrier, pH 7.4 (adjusted with Sigma 7–9 base)]. Membrane currents were acquired at a holding potential (V_H) of −70 mV using an Axopatch 200B patch-clamp amplifier and WinWCP version 4.0.7 software (John Dempster, University of Strathclyde, Glasgow, UK). Linear components of capacitive and leak currents were subtracted using the standard P/4 protocol. Series resistance compensation of >70% was typically used. Data were analyzed offline using WinWCP version 4.0.7, OriginPro version 7.0 (Microcal), and MATLAB (MathWorks) software. I–V data were fitted with the modified Boltzmann equation:

\[ \text{current density} = \frac{G_{\text{max}}(V - V_{\text{rev}})}{1 + \exp(V - V_{\text{rev}}/k)} \]

where \( G_{\text{max}} \) is the maximum conductance, \( V_{\text{rev}} \) is the null potential, \( V_{0.5} \) is the voltage at which 50% of the current is activated, and \( k \) is the slope factor. To determine \( V_{0.5} \) and \( k \) values accurately, data from normalized tail current amplitude was fitted with a standard Boltzmann function. Statistical significance (taken as \( p < 0.05 \)) was determined using one-way ANOVA, Student’s t test, or Mann–Whitney U test, as indicated.

**Synaptic transmission between SCGNs.** Cells were prepared as described previously (Mochida et al., 1996). Briefly, males Wistar rats were decapitated on postnatal day 7 under diethylthreanesthesia according to the guidelines of the Physiological Society of Japan. SCGNs were isolated and maintained in culture for 6–7 weeks in a growth medium of 84% Eagle’s MEM, 10% fetal calf serum, 5% horse serum, 1% penicillin–streptomycin (all from Life Technologies), and 25 mg/ml nerve growth factor (2.5 S, grade II; Alomone Labs).

Paired intracellular recordings were made from two proximal neurons using microelectrodes filled with 1 M potassium acetate (70–90 MΩ) as described previously (Mochida et al., 2003). Briefly, SCGNs were superfused with modified Krebs’ solution containing the following (in mM): 136 NaCl, 5.9 KCl, 2.5 CaCl₂, 1.2 MgCl₂, 11 glucose, and 3 Na-HEPES, pH 7.4 (adjusted with NaOH). Action potentials (APs) were generated in a neuron by current injection (at 0.1 Hz) through the intracellular recording electrode and EPSPs recorded from the synchronically coupled neuron. Data were analyzed using software written by the late Ladislav Tauc (National Center of Scientific Research, Paris, France). Through-out, data were obtained at 32–35°C. Collected data were analyzed with Origin software (Origin Lab) and are reported as means ± SEMs. Statistical significance (taken as \( p < 0.05 \)) was determined using Student’s t test.

**Fast-scan cyclic voltammetry**

Male Wistar rats were anesthetized with halothane and decapitated. The brain was removed, and coronal slices containing the dorsal striatum were cut at a thickness of 300 µm in cold carboxygenated artificial CSF (aCSF; in mM: 125 NaCl, 2.5 KCl, 0.3 KH₂PO₄, 26 NaHCO₃, 2.4 CaCl₂, 1.3 MgSO₄, and 10 d-glucose) using a VT1000S vibrating microtome (Leica Microsystems). Dorsal striatal brain slices were allowed to recover for at least 1 h at room temperature before carboxygenated aCSF. For recordings, slices were superfused with 32°C carboxygenated aCSF at a flow rate of 1 ml/min. Experimental recordings started a minimum of 20 min after transfer to the slice chamber. Carbon fiber electrodes (5 µm; Goodfellow) were made as described previously (Kuhr and Wightman, 1986). Electrodes were inserted −100 µm into the slice. The potential of the working electrode was held at −0.4 V versus Ag/AgCl between scans and was ramped to +1.3 V at 300 Vs and back at −0.4 V every 100 ms via an A-M Systems isolated pulse stimulator. The triangular waveform was computer controlled using a HEKA EVA8 potentiostat and an ESA Bioscience fast-scan cyclic voltammetry interface analog-to-digital converter via TH-1 software (ESA Biosciences). Axonal DA release in the dorsolateral striatum was evoked using a twisted bipolar stimulating electrode (Plastics One). Stimulations were delivered every 2 min by a single electrical pulse (1 ms, 400 µA). Syn at 5 µM (dissolved in aCSF) was superfused 10 min after the first measure. For α-conotoxin experiments, drug was superfused (at 1 ml/min) starting from 4 min before the first recording until the end of the experiment. Background-subtracted cyclic voltammograms were obtained by subtracting the current obtained before stimulation from all recordings. The peak oxidation current for DA in each voltammogram was converted into a measure of DA concentration by post-calibration of the electrode using 1 µM DA (Sigma-Aldrich).
Data were normalized to the first five recordings (10 min) of their respective control period and plotted against time (means ± SEMs). Statistical significance (taken as p < 0.05) was determined by two-way ANOVA (time × treatment as variables).

Microdialysis

In vitro Syn microdialysis was performed to verify Syn diffusion through the microdialysis membrane. A 4 mm, 100 kDa cutoff membrane (CMA12 guide; CMA Microdialysis) was connected to a microdialysis syringe pump (CMA-402 syringe pump; CMA Microdialysis). aCSF (in mM: 147 NaCl, 2.7 KCl, 1.2 CaCl2, and 0.8 MgCl2) plus 10 μM Syn was used as a perfusion medium at a flow rate of 0.1 μl/min. The microdialysis cannula was immersed in 50 μl of aCSF, and the appearance of Syn in the solution was measured by SDS-PAGE and Western blot performed on 2 μl fractions sampled at the indicated times.

In vivo Syn microdialysis was performed in awake and freely moving rats (males Wistar). A guide cannula (CMA12 guide; CMA Microdialysis) was stereotaxically implanted in the right striatum under isoflurane anesthesia and cemented. Stereotaxic coordinates for probe positioning were chosen according to a rat brain atlas: anteroposterior, +1.0 mm; lateral, 3.0 mm; and dorsoventral, −6.5 mm relative to bregma. Approximately 24 h after surgery, dialysis probes were inserted into the guide cannula. The dialysis probes were perfused during implantation into the brain and for 1 h afterward with aCSF. At 1 h after the operation, animals were returned to their home cages. On the next day, the dialysis probes were connected to a syringe pump and perfused with aCSF at 1.0 μl/min for 30 min (washout period). After this stage, the perfusion rate was changed to 0.25 μl/min, and aCSF was perfused for 2 h (equilibration period). The solution was then changed to aCSF containing buffer A, and perfusion was continued for 3 h at 0.25 μl/min. Samples were collected every 60 min into collection tubes containing 2 μl of perchloric acid (1M). After the experiment, animals were disconnected from the system and were left in their home cage overnight. On the second day, the dialysis probes were connected again to a syringe pump and perfused according to the protocol described above with addition of aCSF containing Syn (10 μM). The samples collected during hour 3 of Syn or buffer A application were used for analysis. DA content in samples collected with and without Syn application was determined by HPLC.

HPLC analytical procedure

Measurements of DA in microdialysis samples were performed using HPLC with electrochemical detection (ALEXYS LC-EC system) equipped with a reverse-phase column (3 μm particles, ALB-215 C18, 1 × 150 mm; Antec) at a flow rate of 200 μl/min and electrochemically detected by a 0.7 mm glass carbon electrode (VT-03; Antec). The mobile phase contained 50 mM H3PO4, 50 mM citric acid, 8 mM KCl, 0.1 mM EDTA, 400 mg/l octanesulfonic acid sodium salt, and 10% (v/v) methanol, pH 3.9. The sensitivity of the method allowed detection of ~3 fmol of DA. Dialysate samples (11 μl) were injected into HPLC without any additional purification.

Results

Extracellular Syn induces an increase in evoked calcium entry through Ca2,2

Syn has been shown to be released in physiological and pathological conditions in vitro and in vivo. First, we confirmed the release of Syn from cultured rat neurons over a period of 48 h; Syn was found in the medium in picomolar concentration. Neurons showed no sign of damage or leakage of cytosolic proteins as shown by the absence of GAPDH in the media (Fig. 1A). To determine whether extracellular Syn can affect intracellular calcium levels, we applied highly purified Syn to rat cortical neurons (10 DIV). The monomeric state of purified Syn was assessed by gel filtration and SDS-PAGE (Fig. 1B). During incubation of the purified protein with cultured neurons, the formation of oligomers was found at 48 h (Fig. 1C). The absence of oligomeric or cleaved products of the protein after 2 h incubation was confirmed by gel filtration. Fractions collected from the column showed that the bulk of Syn eluted with proteins of 14–20 kDa, as assessed by a calibration column, and indicate that Syn was monomeric (Fig. 1D). Neurons were incubated with or without recombinant Syn (5 μM) for 2 h at 37°C and subsequently loaded with the calcium dye Fluo-4 (Fig. 2A). In the presence of AMPA and NMDA ionotropic glutamate receptor blockers, we did not detect any significant effect on basal calcium level during Syn application, whereas we measured an ~12% increase in calcium influx after depolarization of neurons with 50 mM KCl (Fig. 2A, middle). We evaluated the time course of effects of Syn preincubation and found a significant increase in calcium influx after 30 min, which was higher and more reproducible after 2 h preincubation (Fig. 2A, right). We applied nifedipine (10 μM), a selective blocker of Ca2,1, to analyze their possible involvement in Syn action. Under these conditions, we found a significantly larger (~22%) increase in calcium signal during Syn exposure, suggesting that, in the absence of nifedipine, we could have underestimated the change in fluorescence because of partial dye saturation (Fig. 2B, left). To determine which calcium channels are targeted by Syn, we used, in addition to nifedipine, the specific calcium channel inhibitors ω-agatoxin IVA and ω-conotoxin GVIA to block Ca2,1 and Ca2,2, respectively (Fig. 2B, middle and right). ω-Agatoxin had no effect on Syn-induced increase in calcium entry, whereas ω-conotoxin completely abolished the effect of Syn, indicating that Ca2,2 are selectively affected by Syn in cortical neurons. Because of the power relationship between calcium influx and exocytosis (Dodge and Rahamimoff, 1967), small changes in influx might have profound effects on transmitter release; however, to exclude the possibility that the effect of Syn could be unspecific, we used three distinct Syn peptides: (1) P34 (amino acids 34–45), i.e., the glycosphingolipid-binding domain of Syn; (2) P61 (amino acids 61–78), i.e., the cholesterol...
Extracellular Syn increases whole-cell Ca\textsubscript{2.2} current density
and synaptic transmission in isolated SCGNs

Rat SCGNs were used to measure the Syn effect on Ca\textsubscript{2.2} activity because, in these neurons, 85% of calcium current is carried by \omega-conotoxin-sensitive Ca\textsubscript{2.2} (Stoehr et al., 1997). Whole-cell calcium currents (measured as \(I_{\text{Ca}}\)) in SCGNs cultured 1–2 d were assessed after 2 h preincubation with purified recombinant Syn at various concentrations (Fig. 3A, representative traces in central panel). The modulatory effect of Syn was measured against separate pools of internal control recordings from SCGNs within the same preparation. Pretreatment of neurons with 5 \(\mu\text{M}\) Syn in the culture medium caused an ~70% increase in calcium current density at 0 mV and a corresponding increase in \(G_{\text{max}}\) (Fig. 3A; Table 1). In these experiments, control Syn had no significant effect on \(V_{\text{rev}}\) (Table 1). At a 10 times lower concentration (0.5 \(\mu\text{M}\)), Syn also produced a significant increase of calcium current density and \(G_{\text{max}}\) with no significant difference between the two concentrations tested, whereas 0.1 \(\mu\text{M}\) Syn, the lowest concentration applied, failed to produce any significant effect compared with controls (Fig. 3A, right; Table 1). To investigate the effect of Syn on \(V_{\text{rev}}\) and slope factor \(k\), we fitted Boltzmann functions to tail current amplitude data measured at ~120 mV; these protocols use short activation pulses to avoid any contaminating voltage-dependent current inactivation and so give the most accurate measures of transition between closed and open states. Concentrations of Syn that affected current density and \(G_{\text{max}}\) were fit to the total protein, binding domain of Syn that has been shown to induce membrane leakage and to aggregate (Crowet et al., 2007; Fantini et al., 2011), and (3) P12 (amino acids 12–23), a peptide with no recognized function. None of these peptides determined measurable effects on calcium entry (Fig. 2C), indicating that the whole protein, albeit unfolded, is required for its activity on Cav2.2.

Ca\textsubscript{2.2} are localized presynaptically and are major regulators of neurotransmitter release (Westerbroek et al., 1992; Catterall, 2000; Garner et al., 2000). To investigate the action of extracellular Syn on calcium influx specifically in the presynaptic compartment, the calcium indicator GCaMP3 was targeted to synaptic vesicles as a chimera with the synaptic vesicle associated protein synaptophysin (SyGCaMP3) (Li et al., 2011). In neurons expressing SyGCaMP3, the fluorescent signal in presynaptic boutons was increased during KCl perfusion (Fig. 2D). In the selected ROI, fluorescence reached a peak lasting ~7 s (Fig. 2E). Incubation of neurons with Syn induced an ~15% increase in calcium entry at the presynapse that was also completely abrogated by \omega-conotoxin (Fig. 2F), suggesting a possible activity of extracellular Syn in the regulation of neurotransmitter release by Ca\textsubscript{2.2}.

**Figure 2.** Extracellular Syn increases calcium influx through Ca\textsubscript{2.2}. A. Left, Representative trace of a calcium transient of a cortical neuron obtained after perfusion for 60 s with 50 mM KCl (starting point at 60 s). Neurons were incubated with 5 \(\mu\text{M}\) Syn (red dots) for 2 h at 37°C or without (black dots); neurons were loaded with Fluo-4 and imaged at 1 frame/s. Data are expressed as \(F/F_0\) (ratio between basal fluorescence in the first 20 s and fluorescence at each data point). Middle, Bar chart showing the increase in fluorescence calculated at the point of maximum value in Syn-treated neurons (red; \(n = 150\)). Data were pooled from different experiments and normalized to the respective control (black; \(n = 150\)) in each experiment. Right, Bar chart showing the increase in fluorescence, calculated as above, in neurons treated with 5 \(\mu\text{M}\) Syn for 20 min, 2 h, and 4 h (red; \(n = 83, n = 63, n = 67\)). Data were normalized to the control (black; \(n = 120\)) in each experiment. Statistical significance determined by one-way ANOVA. * \(p < 0.05\). B, Quantification of Fluo-4 fluorescence during KCl depolarization in Syn-treated neurons (red), calculated as above, in the presence of 10 \(\mu\text{M}\) nifedipine (nife; \(n = 600\)), 100 \(\mu\text{M}\) \omega-agatoxin (aga; \(n = 450\)), or 300 \(\mu\text{M}\) \omega-conotoxin (cono; \(n = 250\)), each bath applied 3 min before KCl perfusion. Data are expressed as mean ± SEM of \(F/F_0\) normalized to the respective control (black; same \(n\) as Syn-treated samples). C, Bar chart showing the increase in fluorescence calculated as above, in neurons treated with 5 \(\mu\text{M}\) P12, P34, and P61 Syn peptides for 2 h (red; \(n = 150\), \(n = 375\), \(n = 167\)). Data were pooled from different experiments and normalized to the respective control (black; \(n = 150\), \(n = 376\), \(n = 100\)) in each experiment. D, Series of microphotographs showing a group of positive synapses (black arrow) in neurons transfected with SyGCaMP3 and depolarized with 50 mM KCl in the presence of nifedipine and in the absence (top) or presence (bottom) of Syn. KCl was perfused starting at frame 12 and caused synaptic boutons to emit fluorescence immediately. ctrl, Control. E, Representative trace of the increase in calcium in neurons transfected with SyGCaMP3 and depolarized as above. Neurons were incubated with 5 \(\mu\text{M}\) Syn (red dots) or without (black dots) as in A. Images were acquired at 1 frame/s. F, Left, Quantification of the fluorescence in SyGCaMP3-positive boutons (red; \(n = 100\)) in Syn-treated neurons normalized to the respective control (black; \(n = 100\)). Right, Quantification as in the left panel of neurons perfused with 300 \(\mu\text{M}\) \omega-conotoxin (cono) 3 min before KCl. Data were expressed as the mean ± SEM of \(F/F_0\) normalized to the respective control. Statistical significance in A–C and F determined by nonparametric t test. * \(p < 0.05\).
Figure 3. Extracellular Syn increases \( I_{\text{BA}} \) currents and synaptic transmission in SCGNs. A, Left, Mean calcium current density–voltage relationships for control (black circles; \( n = 8 \)) and 5 \( \mu \text{M} \) Syn-treated (red circles; \( n = 10 \)) SCGNs. Continuous lines are modified Boltzmann fits to each current density–voltage curve. Currents were evoked using 50 ms depolarizing test pulses in 10 mV intervals from a \( V_{\text{CH}} = -70 \text{ mV} \). Data are shown as mean ± SEM. Middle, Superimposed representative traces showing currents at 0 mV in control (black) and after treatment with 5 \( \mu \text{M} \) Syn for 2 h (red). Right, Bar graph summarizing the effect of Syn on \( I_{\text{BA}} \) and showing data for 0.1 \( \mu \text{M} \) Syn (light pink; \( n = 5 \)), 0.5 \( \mu \text{M} \) (dark pink; \( n = 6 \)), 5 \( \mu \text{M} \) (red; \( n = 10 \)), and the respective controls (black; \( n = 6 \), \( n = 6 \), \( n = 8 \), respectively). Statistical significance determined by nonparametric \( t \) test. * \( p < 0.05 \). B, Plot of capacitance versus current at 0 mV recorded in SCGNs incubated with buffer (black circles; \( n = 14 \)) or Syn (0.5 and 5 \( \mu \text{M} \), red circles; \( n = 16 \)). Dotted lines represent regression fits for buffer-treated (black) and Syn-treated (red) neurons. C, Left, Superimposed current density–voltage curves showing \( I_{\text{BA}} \) (P1; black circles) and effect of a depolarizing prepulse to +120 mV (P2; black squares) in control SCGNs used in A. Right, Superimposed current density–voltage curves showing \( I_{\text{BA}} \) (P1; red circles) and effect of a depolarizing prepulse to +120 mV (P2; red squares) in 5 \( \mu \text{M} \) Syn-treated SCGNs used in A. Middle, Bar chart summarizing the facilitation ratio (P2/P1) expressed as the mean ± SEM in control (black) and 5 \( \mu \text{M} \) Syn-treated (red) SCGNs. Ctrl, Control; PP, prepulse. D, Left, EPSP amplitude evoked at 0.1 Hz in SCGN synapses. At \( t = 0 \) min, buffer (black circles; \( n = 4 \)) or Syn (red circles; \( n = 6 \)) were drop applied to the 2 ml bath (5 \( \mu \text{M} \) final concentration). EPSP amplitude was normalized to that before protein bath application for each neuron. The pink and gray lines represent smoothed values (10-point moving-average algorithm) of the EPSP amplitudes. Right, Bar chart showing the increase of the normalized EPSP amplitude expressed as the mean ± SEM at \( t = 40 \) min in control (black) and Syn-treated (red) SCGNs. Right, Bottom, Bar chart showing the increase of the normalized EPSP amplitude expressed as the mean ± SEM at \( t = 40 \) min in control (black) and Syn-treated (red) SCGNs. Statistical significance determined by nonparametric \( t \) test. * \( p < 0.05 \).

Table 3. Effects of Syn on \( I_{\text{BA}} \) current density  

<table>
<thead>
<tr>
<th>( I-V )</th>
<th>Current density at 0 mV (pA/pF)</th>
<th>( G_{\text{max}} ) (nS/pF)</th>
<th>( V_{\text{rev}} ) (mV)</th>
<th>( n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ctrl</td>
<td>-19.6 ± 2.0</td>
<td>0.33 ± 0.03</td>
<td>54.4 ± 3.4</td>
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</tr>
<tr>
<td>Syn 5 ( \mu \text{M} )</td>
<td>-33.5 ± 2.6*</td>
<td>0.56 ± 0.04*</td>
<td>61.5 ± 2.3</td>
<td>10</td>
</tr>
<tr>
<td>Ctrl</td>
<td>-18.4 ± 2.6</td>
<td>0.32 ± 0.12</td>
<td>51.4 ± 1.3</td>
<td>6</td>
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<tr>
<td>Syn 0.5 ( \mu \text{M} )</td>
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<td>0.63 ± 0.07*</td>
<td>52.3 ± 1.5</td>
<td>6</td>
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<tr>
<td>Ctrl</td>
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<td>0.32 ± 0.14</td>
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<td>5</td>
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<tr>
<td>Syn 0.1 ( \mu \text{M} )</td>
<td>-19.1 ± 8.5</td>
<td>0.32 ± 0.14</td>
<td>53.6 ± 1.0</td>
<td>5</td>
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</table>

Current densities from control and Syn-treated SCG neurons were plotted against voltage and fitted with a single Boltzmann function. * \( p < 0.05 \) versus control (Ctrl). Statistical significance for \( I-V \) data, expressed as mean ± SEM, was determined by Student’s \( t \) test.

Table 3. Effects of PPF on Syn modulation of \( I_{\text{BA}} \) current density  

<table>
<thead>
<tr>
<th>( I-V )</th>
<th>Current density at 0 mV (pA/pF)</th>
<th>( V_{\text{rev}} ) (mV)</th>
<th>( n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ctrl</td>
<td>-19.6 ± 2.0</td>
<td>54.4 ± 3.4</td>
<td>8</td>
</tr>
<tr>
<td>Ctrl + PP</td>
<td>-29.6 ± 3.5*</td>
<td>51.8 ± 2.4</td>
<td>10</td>
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<tr>
<td>Syn 5 ( \mu \text{M} )</td>
<td>-33.5 ± 2.6*</td>
<td>61.5 ± 2.3</td>
<td>10</td>
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<tr>
<td>Syn 5 ( \mu \text{M} ) + PP</td>
<td>-43.0 ± 3.1***</td>
<td>58.5 ± 1.0</td>
<td>6</td>
</tr>
<tr>
<td>Ctrl</td>
<td>-18.4 ± 2.6</td>
<td>54.6 ± 2.3</td>
<td>6</td>
</tr>
<tr>
<td>Ctrl + PP</td>
<td>-21.0 ± 2.4*</td>
<td>54.7 ± 2.7</td>
<td>6</td>
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<tr>
<td>Syn 0.5 ( \mu \text{M} )</td>
<td>-37.6 ± 3.9*</td>
<td>52.3 ± 1.5</td>
<td>6</td>
</tr>
<tr>
<td>Syn 0.5 ( \mu \text{M} ) + PP</td>
<td>-43.1 ± 5.6***</td>
<td>52.2 ± 1.1</td>
<td>6</td>
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<tr>
<td>Ctrl</td>
<td>-19.0 ± 7.1</td>
<td>51.4 ± 1.3</td>
<td>6</td>
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<tr>
<td>Ctrl + PP</td>
<td>-22.1 ± 7.9*</td>
<td>52.5 ± 1.2</td>
<td>6</td>
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<tr>
<td>Syn 0.1 ( \mu \text{M} )</td>
<td>-19.1 ± 8.5</td>
<td>53.6 ± 1.0</td>
<td>5</td>
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<td>Syn 0.1 ( \mu \text{M} ) + PP</td>
<td>-23.0 ± 9.1***</td>
<td>53.7 ± 1.1</td>
<td>5</td>
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</table>

Current densities from control and SCGNs were plotted against voltage and fitted with a single Boltzmann function. * \( p < 0.05 \) versus control (Ctrl); ** \( p < 0.05 \) versus control + PP; *** \( p < 0.05 \) versus Syn. Data were expressed as mean ± SEM. Statistical significance determined by Student’s \( t \) test.

Calcium channels in SCGNs are physiologically modulated by voltage-dependent G-proteins (Dolphin, 1996), and G-protein modulation is characterized by an inhibition of current amplitude (Stephens et al., 1998). Therefore, we examined the effect of Syn on endogenous G-protein modulation. Transient dissociation of G-proteins from the channel induces facilitation of calcium currents that can be measured in response to a brief, strongly depolarizing voltage step [named prepulse facilitation (PPF)]. A prepulse of +120 mV was applied to control neurons to measure endogenous G-protein modulation (Fig. 3C; Table 3). In the control group (Fig. 3C, left; Table 3), the ratio between the currents recorded before (P1) and after (P2) the PPF step was 1.43 ± 0.10 (\( n = 10 \)) at 0 mV. After pretreatment with 5 \( \mu \text{M} \) Syn (Fig. 3C, right; Table 3), the P2/P1 ratio was 1.31 ± 0.11 (\( n = 10 \))
at 0 mV, showing that PPF of \( I_{\mathrm{K}} \) caused a similar level of current relief from basal G-protein inhibition compared with controls (Fig. 3C, middle inset). Thus, together with the above, acutely isolated SCGNs exhibit enhanced calcium currents when pre-treated with Syn, but Syn effects cannot be explained by a change in channel sensitivity to G-protein modulation.

We next investigated the modulatory role of Syn in synaptic transmission, taking advantage of the synapses formed among SCGNs in long-term culture (Fig. 3D). This is a well characterized model for studying fast cholinergic transmission and modulation of Ca\(_{2.2}\) (Mochida et al., 2003; Stephens and Mochida, 2005).

Incubation of neurons with 5 \( \mu \text{M} \) Syn caused an increase in EPSP amplitude (Fig. 3D, left), with the maximum increase reached \( \sim 40 \text{ min} \) after Syn application. The increase in synaptic transmission was reversible; washing out Syn from the bath returned synaptic transmission to control levels after an additional 40 min. These results indicate that Syn plays a modulatory role in evoked synchronous transmitter release, leading to an \( \sim 33\% \) increase in EPSP amplitude compared with control (Fig. 3D, right). The observed effect of Syn on synaptic transmission is consistent with the reported increase in calcium current through presynaptic Ca\(_{2.2}\).

Syn-dependent activation of Ca\(_{2.2}\) leads to increased DA release in the striatum

Little is known about synaptic dysfunctions at the premanifest stage of PD. In mice overexpressing Syn, tonic DA release is increased at early stages of the pathology (Lam et al., 2011), and we show that Syn-dependent activation of Ca\(_{2.2}\) at the presynapse leads to altered neurotransmitter release in SCGNs. Increased extracellular DA may be a common mechanism contributing to substantia nigra dopaminergic neuron stress and eventual demise, subsequently leading to downregulation of synaptic activity in the striatum. To assess the effect of extracellular Syn on DA release in the striatum, we performed fast scan cyclic voltammetry experiments on striatal slices perfused with 5 \( \mu \text{M} \) purified Syn. Axonal DA release in the dorsolateral striatum was evoked using a twisted bipolar stimulating electrode. Stimulations were delivered every 2 min by a single electrical pulse (1 ms, 400 \( \mu \text{A} \)). Syn dissolved in aCSF was superfused 10 min after the first measure (control period), and, after 20 min of treatment, DA release increased by \( \sim 30\% \) (Fig. 4A), in agreement with the observed increase in EPSP amplitude (Fig. 3D). Similarly to that observed for calcium entry, superfusion with \( \omega \)-conotoxin primarily ablated DA release (Fig. 4A). \( \omega \)-Conotoxin also reduced basal DA levels by \( \sim 60\% \), indicating that, in dorsolateral striatum preparations, Ca\(_{2.2}\) are the predominant calcium channels supporting neurotransmission (Phillips and Stamford, 2000). The large increase in DA release induced by excess Syn suggested a physiological role for Syn in the regulation of neurotransmitter release in the striatum. We wondered whether sequestration of constitutively released Syn from neurons would modify DA release in the opposite way. Perfusion of slices with anti-Syn antibody induced a significant \( \sim 10\% \) decrease in DA release (Fig. 4B), indicating that Syn, in physiological conditions, acts as a positive regulator of neurotransmission.

To investigate the effect of in vivo acute administration of Syn on the release of DA in the striatum, we performed microdialysis to perfuse Syn in the striatum and measured DA release from the same area (Fig. 4C). The ability of Syn to diffuse through a microdialysis membrane with 100 \( \mu \text{kDa} \) cutoff was determined under in vitro microdialysis conditions, and the passage of Syn into the solution outside the membrane was verified by immunoblot (Fig. 4C, top). Then, a 4-mm-long microdialysis cannula passing through the whole length of striatum was placed in the dorsal striatum of the rat. Freely moving rats were perfused at a rate of 0.25 \( \mu \text{l/min} \) with aCSF.
solution with or without 10 \mu M Syn. Samples were collected for 60 min during hour 3 after addition of Syn into the perfusion solution and measured by HPLC. These experiments revealed a significant ~50% increase in DA release in the Syn-treated group compared with respective control values obtained during perfusion of solution without Syn (Fig. 4C, bottom).

**Extracellular Syn does not affect Ca\textsubscript{v2.2} interaction with major regulatory proteins**

Ca\textsubscript{v2.2} function is widely reported to be dictated by their interaction with a number of binding partners (Jarvis and Zamponi, 2007). Therefore, we investigated whether Syn directly binds Ca\textsubscript{v2.2} to affect their kinetics or indirectly affects other regulatory proteins. Neurons, incubated with 5 \mu M Syn for 2 h, were lysed and immunoprecipitated with anti-Ca\textsubscript{v2.2} antibody. Syn band was present also in the absence of anti-Ca\textsubscript{v2.2} antibody and in the absence of Syn addition because of unspecific binding of endogenous Syn to agarose beads, but it was not significantly increased in the anti-Ca\textsubscript{v2.2} + Syn sample (1.07 ± 0.19-fold vs control in the absence of Syn, n = 3), excluding a direct binding between the two proteins (Fig. 5A). To further evaluate the possibility that Ca\textsubscript{v2.2} activation was attributable to impaired binding of G-protein \textbeta{1} subunit to the channel (Dolphin, 2003), we coimmunoprecipitated with anti-Ca\textsubscript{v2.2} antibody. The difference in the amount of G-protein \textbeta{1} subunit coimmunoprecipitated with the channel in the presence of extracellular Syn was not significant (1.05 ± 0.11-fold vs control in the absence of Syn; n = 3; Fig. 5A), a result that is consistent with the observed lack of change in PPF in electrophysiological experiments. Furthermore, Ca\textsubscript{v2.2} is activated by protein kinase C (PKC; Stea et al., 1995), but in the presence of the PKC inhibitor GF109203X (2-[1-(3-dimethylaminopropyl)-1H-indol-3-yl]-3-(1H-indol-3-yl)maleimide) (Fig. 5B), there was no change in Syn-dependent increase in \omega-conotoxin-sensitive calcium entry, excluding PKC as a mediator of Syn effect. To evaluate the possibility that Syn could induce a change in the binding of syntaxin 1A to Ca\textsubscript{v2.2} and remove the tonic inhibition of the channel (Davies et al., 2011), we analyzed the interaction between the two proteins in the absence or presence of Syn. No differences were observed between control and Syn-treated samples (1.03 ± 0.26-fold vs control in the absence of Syn; n = 3; Fig. 5C). Syntaxin 1A modulation of Ca\textsubscript{v2.2} has been reported to be associated with a stabilization of channel inactivation (Bezprozvanny et al., 1995). Consistent with a lack of effect on syntaxin 1A binding, Syn had no effect on Ca\textsubscript{v2.2} voltage dependence of inactivation (Fig. 5D).

Together, these data suggest that Syn neither binds directly to Ca\textsubscript{v2.2} nor acts by disrupting binding of effectors of major pathways incident on Ca\textsubscript{v2.2}.

**Extracellular Syn decreases plasma membrane cholesterol content**

Ion channel activity is also widely reported to be regulated via the cholesterol content of the plasma membrane (Levitan et al., 2010), with calcium current shown to be decreased by cholesterol loading (Jennings et al., 1999; Toselli et al., 2005). Therefore, we measured total and plasma membrane cholesterol by staining with filipin, a cholesterol binding molecule, in permeabilized and nonpermeabilized neurons, respectively (Fig. 6A). Although methyl-\beta-cyclodextrin (M\textbeta{CD}), as expected, depleted both total...
and membrane cholesterol, Syn induced a selective \( \sim 30\% \) decrease in plasma membrane cholesterol (Fig. 6A, right), whereas total cholesterol was unchanged. To assess the role of cholesterol on calcium influx in isolated cells, we used M\(_{\beta}CD\) and cholesterol-loaded M\(_{\beta}CD\) (cholesterol-M\(_{\beta}CD\)) to decrease and increase cholesterol content, respectively. We found that KC1-evoked calcium entry in the absence of calcium channel inhibitors was increased by M\(_{\beta}CD\), an effect mimicked by Syn (Fig. 6B), whereas in neurons loaded with cholesterol, evoked calcium entry was less than half of that in controls (Fig. 6D). We next investigated whether cholesterol loading of neurons could prevent Syn-induced depletion of plasma membrane cholesterol and calcium increase; indeed, both effects were blocked when neurons were incubated with cholest-M\(_{\beta}CD\) and Syn together (Fig. 6C,D). Together, these results indicate that a decrease in plasma membrane cholesterol content induced by extracellular Syn is causal to the increase in intracellular calcium. We next wondered whether cholesterol loading would act in the opposite manner to Syn and act to inhibit neurotransmitter release. SCGNs treated with 0.13 and 1.3 mg/ml chol-M\(_{\beta}CD\) showed a dramatic reduction in EPSP amplitude compared with control (Fig. 6E). To monitor in real time the possible effect of cholesterol loading on Syn-driven increase in EPSP (Fig. 3D), we recorded EPSP from SCGNs loaded with cholest-M\(_{\beta}CD\) in the presence of Syn (Fig. 6F). The initial increase in amplitude was almost completely reversed after 20 min, with the effect of Syn decreasing to 40% of the initial value after 30 min. During washout, the reduction of EPSP amplitude persisted for the duration of the recording time (100 min; Fig. 6F), indicating that cholesterol loading strongly attenuated Syn effect on neurotransmission. Together, these results suggest a causal role for cholesterol in mediating Syn action.

**Ca\(_{\text{2,2}}\)** at the plasma membrane relocate into cholesterol-poor domains during Syn action

An increased activity of Ca\(_{\text{2,2}}\) could be attributable to an increase in their surface expression. We characterized the amount of Ca\(_{\text{2,2}}\) by cell-surface biotinylation and found that Syn does not affect Ca\(_{\text{2,2}}\) expression at the neuronal plasma membrane (1.17 \pm 0.10-fold vs control in the absence of Syn; \( n = 4; \) Fig. 7A). The reduced cholesterol level in the plasma membrane of neurons exposed to Syn was suggestive of a change in microdomain composition and partitioning of proteins. To address this hypothesis, we analyzed the potential changes in raft partitioning of different proteins in response to Syn treatment. Neuronal extracts were loaded at the bottom of a nonlinear sucrose gradient (Fig. 7B) and were shown to contain the same protein and cholesterol content in each fraction between control and Syn-treated neurons (Fig. 7C). It is well established that proteins, such as flotillin-1 and caveolin, represent markers for lipid rafts and partition into Triton X-100-insoluble cholesterol-rich microdomains (Bickel et al., 1997). In our gradient, flotillin-1 and caveolin were recovered in fractions 5 and 6 (Fig. 7D), we recorded EPSP from SCGNs loaded with cholest-M\(_{\beta}CD\) in the presence of Syn (Fig. 6F). The initial increase in amplitude was almost completely reversed after 20 min, with the effect of Syn decreasing to 40% of the initial value after 30 min. During washout, the reduction of EPSP amplitude persisted for the duration of the recording time (100 min; Fig. 6F), indicating that cholesterol loading strongly attenuated Syn effect on neurotransmission. Together, these results suggest a causal role for cholesterol in mediating Syn action.

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fraction containing solubilized proteins (Fig. 7E). As a control for cholesterol depletion, we treated neurons with MβCD and then loaded extracts on a sucrose gradient (Fig. 7F). However, MβCD primarily depletes cholesterol both inside the cell and at the membrane (Fig. 6A) and, as expected, induced an almost complete solubilization of Cav2.2 and also of proteins tightly associated with lipid rafts, such as flotillin-1.

Finally, we examined the behavior of surface exposed Cav2.2, whose activity accounted for the increased calcium currents observed in the presence of Syn above. We found that ~40% of the biotinylated channels that were in rafts in control conditions shifted to cholesterol-poor domains of the plasma membrane during Syn incubation (Fig. 8A, left); in contrast, α2β-1 subunits, which also partition in lipid rafts and are widely reported to modulate trafficking and to direct Cav2.2 into lipid rafts (Davies et al., 2006; Robinson et al., 2010), remained in cholesterol-rich domains (Fig. 8A, right). These data indicate that the presence of extracellular Syn cause a relocation of cell-surface Cav2.2 into cholesterol-poor domains of the plasma membrane (Fig. 8B). This change in the environment of the channel is consistent with increased channel activation, calcium entry, and release of neurotransmitters.

**Discussion**

Our study focuses on the pathological role of increased concentrations of extracellular Syn, a condition that might arise by the release of Syn from neurons in which Syn is overexpressed, because of the presence of multiple SNCA gene copies (Singleton et al., 2003; Farrer et al., 2004) or polymorphism in regions of the gene that regulate its expression (Chiba-Falek et al., 2006). Our results provide evidence that Syn alters the organization of the plasma membrane of neurons, acting on the external leaflet of the membrane to lower the content of plasma membrane cholesterol and thereby altering the partitioning of surface-associated proteins. Syn action leads to increased presynaptic calcium influx and neurotransmitter release mediated by Cav2.2. There is much evidence that Syn function is related to its association with plasma and vesicular membranes at synaptic terminals (Sharon et al., 2001) and in particular with lipid rafts (Fortin et al., 2004). In vitro studies suggest an effect of Syn on the integrity of lipid bilayers inducing lipid clustering (Madine et al., 2006). It was shown recently in SH-SYSY cells that membrane fluidity was affected by exogenous Syn (Melachroinou et al., 2013). The observed effect of Syn on the composition of plasma membrane may involve altered endocytosis or recycling of lipid and proteins, leading to incorrect positioning of proteins in microdomains with different lipid composition, rather than causing differences in the amount of proteins at the cell surface. In our study, an effect of Syn on membrane retrieval to the intracellular compartment was ruled out by the measurements of capacitance, which appeared unchanged. However, it is possible that membrane retrieval of raft domains is accompanied by recycling of cholesterol-poor vesicles at the membrane, leaving the surface area intact. In particular, we found that Cav2.2 repositioned in cholesterol-poor domains with important effects on channel electrophysiological properties. The reported reduction in slope factor from calcium tail current amplitude measurements suggests an improvement of channel voltage sensitivity, such that the gating charges move more easily and, thus, that channel activation is improved. The large increase in current conductance with an accompanying improved voltage sensitivity of the channel is likely a result of the compartmentalization of the channel in less stiff membrane domains in which the membrane deformation energy required for the transition from closed to open state is lower (Lundbaek et al., 1996; Levitan et al., 2000). It has been shown that lipid rafts composition affects calcium channels currents (Davies et al., 2006; Levitan et al., 2010). In neuroblastoma × glioma NG-108 hybrid cells, cholesterol loading selectively reduces Cav2.2 currents by decreasing the channel open probability (Toselli et al., 2005). Conversely, cholesterol depletion by MβCD, which disrupts lipid rafts, has been shown to induce increase Cav2.1 current in recombinant cells (Davies et al., 2006) and to cause a trend to increase current in caveolin-1 transfected NG-108 cells (Toselli et al., 2005). These reports suggest that association of Cav2.2 in a lipid-rich environment results in reduced calcium current, whereas in a lower cholesterol environment, current is increased. Here we show that, in both central and peripheral neurons, cholesterol depletion mimics Syn action and that cholesterol loading has opposite effects to Syn on synaptic transmission. The improved transition between the closed and open states of the channels in a cholesterol-poor environment seems independent of the association of the channel with α2δ-1 subunit on the surface. Thus, the localization of surface α2δ-1 subunit, highly enriched in rafts domains (Robinson et al., 2010), was unaffected by the presence of Syn. α2δ subunits control the functional expression of the α1 subunit, modulating its trafficking from the cell soma and leading to a significant increase in synaptic abundance of α1 (Dolphin, 2009). The comparable amount of surface-exposed Cav2,2 in control and Syn-treated neurons indicated that α2δ-1-dependent trafficking of channels to the membrane was unchanged. Overall, we demonstrate that the Syn-mediated shift of Cav2,2 into cholesterol-poor domains is sufficient to induce enhancement of calcium current independently of α2δ association with the channel.

Our data suggest that, in the presence of Syn, channels are shifted into a low-cholesterol environment, and the ensuing increase in calcium influx has a profound effect on release. It has
been shown that cholesterol depletion does not influence the confinement of Ca$_{\text{v}}$1 in proximity to release sites at bipolar cells ribbon synapses (Thoreson et al., 2013), suggesting that lipid rafts are not required to cluster Ca$_{\text{v}}$1, but rather clusters of channels are maintained by interaction with presynaptic proteins and cytoskeletal elements. Effects may be synapse specific, in that the same group reports that cholesterol depletion does affect Ca$_{\text{v}}$1 channel mobility at photoreceptor synapses in the retina (Mercer et al., 2011, 2012). Importantly, in these studies, Ca$_{\text{v}}$1 channel mobility was assessed via the use of antibodies against the major retinal a2$eta$-4 subunit. It may be that a2$eta$ subunits are important for trafficking of Ca$_{\text{v}}$2.2 from the endoplasmic reticulum to the plasma membrane (Dolphin, 2012), but once at the membrane, association is more labile, consistent with reported low-affinity association of a2$eta$ with Ca$_{\text{v}}$2.2 in plasma membrane proteomic studies (Muller et al., 2010). As above, our data strongly suggest that Syn effects on release occur in the absence of movement of a2$eta$-1, which we show to be retained in cholesterol-rich domains. Therefore, it is possible that, during exposure to Syn, Ca$_{\text{v}}$2.2, independently from a2$eta$-1 and lipid rafts, remain positioned at lower cholesterol regions within active zones in which they sustain neurotransmitter release. It is also reported that cholesterol depletion decreases synaptic vesicles release by altering membrane curvature (Rituper et al., 2013). However, extracellular Syn did not affect internal cholesterol levels here, arguing against a major contribution to effects on release in this study. Another possible explanation for the observed increase in Ca$_{\text{v}}$2.2 activity is that changes in membrane organization may move the channels into proximity of a second messenger, such as phosphatidylinositol 4,5-bisphosphate, which in turn modulates channel activity (Roberts-Crowley et al., 2009). The importance of Ca$_{\text{v}}$2.2 in neurotransmission at the presynapse is well documented, and these channels are the main players in the control of striatal DA release (Phillips and Stamford, 2000). We provide evidence that DA release is significantly increased in a conotoxin-sensitive manner in striatal slices superfused with Syn, and this result correlates with the increase in calcium entry at the presynapse, as assessed by the use of SyGCaMP3. Together, these data identify a mechanism that would explain the increase in DA release found at premanifest stages of PD in Syn transgenic mice (Lam et al., 2011) and in virus-injected mice at early stages of the pathology (Chung et al., 2009). The increase in DA is reproduced in vivo by microdialysis of Syn directly into the striatum, suggesting a tonic increase in DA levels attributable to Syn accumulation in the synaptic cleft. Early alterations at the synaptic terminals level might be responsible for the “dying back” of dopaminergic neurons starting at the axon terminal, sensitizing DA neurons in the initial, presymptomatic, stages of disease. The underlying regulatory mechanism we propose—Syn-mediated disorganization of lipid rafts (Fig. 8B)—may also occur in other areas prone to neurodegeneration in PD and affect the function of other plasma membrane-associated proteins, possibly explaining the selective vulnerability of particular subset of neurons depending on their lipid and protein composition. As an example, the Ca$_{\text{v}}$2.2 knock-out mice model displays reduction in pain sensitivity (Saegusa et al., 2002), and pain is one of the non-motor symptoms of PD; therefore, we can speculate a causal relationship between hyperactivation of CaV2.2 and pain sensitivity in PD.

References


