Myrip couples the capture of secretory granules by the actin-rich cell cortex and their attachment to the plasma membrane.

Sébastien Huet, Isabelle Fanget, Ouardane Jouannot, Patricia Meireles, Tim Zeiske, Nathanaël Larochette, François Darchen, Claire Desnos

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Exocytosis of secretory granules (SGs) requires their delivery to the actin-rich cell cortex followed by their attachment to the plasma membrane (PM). How these reactions are executed and coordinated is still unclear. Myrip, which is also known as Slac-2c, binds to the SG-associated GTase Rab27 and is thought to promote the delivery of SGs to the PM by recruiting the molecular motor myosin Va. Myrip also interacts with actin and the exocyst complex, suggesting that it may exert multiple roles in the secretory process. By combining total internal reflection fluorescence microscopy, single-particle tracking, a photoconversion-based assay, and mathematical modeling, we show that, in human enterochromaffin cells, Myrip (1) inhibits a class of SG motion characterized by fast and directed movement, suggesting that it facilitates the dissociation of SGs from microtubules; (2) enhances their motion toward the PM and the probability of SG attachment to the PM; and (3) increases the characteristic time of immobilization at the PM, indicating that it is a component of the molecular machinery that tether SGs to the PM. Remarkably, while the first two effects of Myrip depend on its ability to recruit myosin Va on SGs, the third is myosin Va independent but relies on the C-terminal domain of Myrip. We conclude that Myrip couples the retention of SGs in the cell cortex, their transport to the PM, and their attachment to the PM, and thus promotes secretion. These three steps of the secretory process are thus intimately coordinated.

Introduction

Secretory granules (SGs) store water-soluble hormones and release them by exocytosis. A wealth of mechanistic information has been obtained on membrane fusion, but less is known on the recruitment of SGs at the cell periphery and their attachment to the plasma membrane (PM), which is required for the formation of a pool of ready-to-fuse SGs (Verhage and Sorensen, 2008). SGs are loaded at the trans-Golgi network and transported along microtubules to the cell periphery (Rudolf et al., 2001). They cannot be transferred directly from microtubules to the PM, but diffuse within the actin-rich cortex until they find an attachment site or undergo another microtubule-based run (Huet et al., 2006). Tethering organelles to the actin cortex is thought to promote their dissociation from microtubules and their accumulation at the cell periphery, and thus secretion (Seabra and Coudrier, 2004; Darchen and Desnos, 2012). However, excessive binding to actin restricts the mobility of SGs and their availability for release (Desnos et al., 2003). Actin dynamics (Malacome et al., 2006) as well as molecular motors such as myosin Va (MyoVa) are thought to facilitate the motion of SGs through the actin-rich layer.

MyoVa is associated with SGs and its inhibition impairs SG docking and secretion (Rose et al., 2003; Rudolf et al., 2003, 2011; Varadi et al., 2005; Desnos et al., 2007b). MyoVa is recruited on SGs by the GTPase Rab27a and Myrip (Fukuda and Kuroda, 2002; Desnos et al., 2003; Waselle et al., 2003; Imai et al., 2004). Myrip, which is also known as Slac-2c, is expressed on SGs and retinal melanosomes (Darchen and Desnos, 2012), and like MyoVa, Rab27a, and Rab27b, it controls secretion (Desnos et al., 2003; Waselle et al., 2003; Imai et al., 2004; Ivarsson et al., 2005; Mizuno et al., 2011). Myrip interacts with Rab27-GTP via an N-terminal helix, and with myosin VIIa or MyoVa via a central domain (El-Amraoui et al., 2002; Fukuda and Kuroda, 2002; Desnos et al., 2003; Kuroda and Fukuda, 2005; Klomp et al., 2007; Waselle et al., 2003; Imai et al., 2004). Interestingly, through its C-terminal region, Myrip also interacts with sec6 and sec8, two components of the exocyst complex, and with actin (El-Amraoui et al., 2002; Fukuda and Kuroda, 2002; Desnos et al., 2003; Goehring et al., 2007), suggesting that Myrip can exert MyoVa-independent functions.
Here, we measured the dynamics of SGs beneath the PM in neuroendocrine cells using total internal reflection fluorescence microscopy (TIRFM) combined with single-particle tracking and incorporated the data into a mathematical compartment model of SG trafficking. We found that Myrip plays a dual role in SG dynamics. First, Myrip recruits MyoVa and promotes the retention of SGs in the actin-rich cortex and their transport toward the PM. Second, it allows their docking at the PM. By coupling the delivery of SGs to the PM and the docking reaction, Myrip thus plays a key role in the secretory process.

Materials and Methods

Materials
Anti-chromogranin A/B antibodies were from Abcam; goat anti-Myrip antibodies were from Everest Biotechnology. The anti-myosin Va (Espireafo et al., 1992) was a gift from R. E. Cheney (University of North Carolina, Chapel Hill, NC) or obtained from Sigma (LF-18); anti-actin (AC-74) and anti tubulin (TUB 2.1) antibodies were from Sigma, and anti-Myo Va11 was from Cell Signaling Technology. Cell culture reagents were obtained from PAA Laboratories. DNA purifications were done using kits from Macherey-Nagel. Enzymes for molecular biology were from New England Biolabs.

Constructs
To generate a plasmid encoding Dendra2-tagged neuropeptide-Y (NPY), p-Dendra2C (Evrogen) was digesting using BamHI/EcoRI and sub-cloned into pcDNA3.1. The resultant plasmid was cut with BamHI/NotI, and the fragment was used to replace the BamHI-NotI fragment in pNPY-GFP to generate an in-frame fusion between NPY and Dendra2. The sequence of inserts was confirmed by automated DNA sequencing. pcDNA3-mycRab11b was a gift from M. Cormont (INSERM U895, Centre Méditerranéen de Médecine Moléculaire, Nice, France). Vectors encoding human Myc-tagged-Myrip constructs, human myosin Va, and NPY constructs have been described previously (El-Amraoui et al., 2002; Desnos et al., 2007b). To generate a Myrip rescue construct, three silent mutations corresponding to nucleotides 2, 9, and 11 of Myrip-siRNA1 (sense strand) were introduced by PCR-based mutagenesis in the Myc-tagged Myrip construct. We verified by immuno-fluorescence that the expression of this rescue construct was not reduced by Myrip-siRNA1, in contrast to that of the original Myrip construct.

Cell culture, transfection, and silencing
The BON cell line has been established from a lymph nod metastasis of a human pancreatic carcinoid tumor (Evers et al., 1991) and were provided by C. M. Townsend (University of Texas, Medical Branch, Galveston, TX). In culture, a large majority of the cells extend several processes from the cell body. We used DMEM (Invitrogen) supplemented with 15% FBS. Cells were plated onto collagen-coated glass coverslips before transfection. TIRFM observations were made on collagen-coated glass-bottom dishes (MatTek or World Precision Instruments). SirNA duplexes were provided by MWG Biotech. We used three already validated siRNAs directed against human MyoVa (MyoVa-siRNA1, 5’-GAACAAACUGAGACCUUCUUU; MyoVa-siRNA2, 5’-AA AGUAGGUGGUUCGUAAUU; MyoVa-siRNA3, 5’-AAGGUCAGCUAC CUGAAGAUGA) (Desnos et al., 2007b), Lindsay and McCaffrey, 2011) and two siRNAs targeting Myrip (Myrip-siRNA1, 5’-UAAGAGGAC GUAAACGAAAU; Myrip-siRNA2, 5’-UGGCGAGUACUCAACAUU). The control siRNAs targeting EGFP or luciferase were from Eurofin MWG Operon. Transfection of siRNAs (30 ns) was done with interferin (Polyplus Transfection). Twenty-four hours later, cells were transfected with plasmids using lipofectamine 2000 (Invitrogen). In some cases, BON cells were transfected by electroporation in the presence of siRNA (120 ns) and 24 h later with siRNAs and interferin. In brief, 1.5 × 10^6 cells were collected in 50 μl of PBS containing vector DNAs (2–5 μg), electroporated (600 V/cm, 8 × 3 ms) using a PS10 electropulsator (Jouan), and recovered in warm culture medium before plating. Experiments were done 72 h after initiation of RNA interference. Cell extracts were prepared in parallel for Western blot analysis for protein expression. The blots were scanned and quantified using Image J software (rsweb.nih.gov/ij/). Levels of actin or tubulin were used to normalize the results from different samples.

Coimmunoprecipitation
BON cells were transfected 2 d before the experiment with vectors encoding myc-tagged Myrip and GFP-MyoVa tail. Immunoprecipitation was performed as described previously (Desnos et al., 2003) on cell homogenates prepared in 1% Triton X-100 in the presence of 2 μM latrunculin B (Calbiochem) to recover Myrip that otherwise bound strongly to actin filaments. Extracts were incubated with protein G-Sepharose beads (GE Healthcare) conjugated with required antibodies at 4°C for 2 h. After extensive washes and elution in Laemmli sample buffer, eluates were analyzed by SDS-PAGE and immunoblots.

Immunofluorescence microscopy
Immunocytochemistry was performed as described previously (Desnos et al., 2003). Secondary antibodies were coupled to Cy3-3 (Jackson ImmunoResearch Laboratories), Alexa-488, Alexa-568, or Alexa–350 (Invitrogen). Cells were visualized using a z-motorized Nikon inverted microscope TE2000E equipped with a 100× objective (NA, 1.4) and a CoolSnap ES CCD camera (Roper Scientific). When indicated, Z-series were acquired with Δz = 200 nm, and image stacks were restored using the MetaMorph point spread function-based deconvolution software (Molecular Devices). To evaluate the association of MyoVa with SGs, cells were imaged using the same laser and camera settings. We first identify SGs as NPY-monomeric red fluorescent protein (mRFP)-labeled structures whose brightest pixel had a fluorescence intensity >1.25 times that of the background fluorescence measured in several areas of the cell. Then, we categorized a SG as MyoVa positive if the green fluorescence of this brightest pixel was >1.25 times that of the background green fluorescence of this cell.

High-pressure freezing immunoelectron microscopy
BON cells were cultured on thermanox coverslips (Nunc) and fitted in a 6 mm specimen carrier for rapid freezing in a Leica HPM 100 machine under high pressure (2100 bars). Samples were then rapidly transferred to liquid nitrogen for storage. Cryosubstitution and embedding were performed in a Reichert AFS apparatus (Leica), first in acetone with 0.1% tannic acid at −90°C for 40 h with one change of solution, and then in acetone with 2% osmium during the last 7 h. The cultures were slowly (5°C/h) warmed to −20°C and incubated for additional hours before being warmed (10°C/h) to 4°C. After several rinses with acetone, the samples were warmed to room temperature and incubated in 50% acetone/50% Araldite (Polysciences) for 1 h, followed by 10% acetone/90% Araldite for 2 h. They were then incubated twice in pure Araldite for 2 h. Coverslips were mounted on resin block and cut in parallel to the plan of cells after removing the thermanox coverlip. Sections (80 nm thick) were cut using a Leica Ultracut E, counterstained by incubation with 2% uranyl in water for 10 min and lead citrate for 10 min. The sections were observed in a Philips TENAI 12 (FEI).

TIRFM
TIRFM setup. BON cells were transferred into Locke’s solution containing the following (in mM): 5.6 glucose, 3.6 HCO\textsubscript{3}, 159.6 CI\textsuperscript{–}, 157.6 Na\textsuperscript{+}, 5.6 K\textsuperscript{+}, 5 HEPES-NaOH, 2.5 CaCl\textsubscript{2}, 1.2 MgCl\textsubscript{2}. TIRFM imaging was performed on a custom setup described previously (Huet et al., 2006). The penetration depth δ (the distance along the z-axis over which fluorescence declines e-fold) of the evanescent field used to excite the fluorophore was set to 150 nm. Under the conditions of observation used, one pixel corresponded to 107.5 nm. Stream acquisitions were performed at 9.98 Hz for 40–60 s with an exposure time of 100 ms. Selective excitation was obtained using argon laser lines at 488 and 514 nm, and optical filters (bandpass, 500–540 nm; high-pass, >565 nm) were used for the emission (Melles Griot).

Image analysis.
The density of fluorescent structures was evaluated using Multidimensional Image Analysis (MIA) software (a segmentation algorithm based on wavelets) (Racine et al., 2006) and expressed as the number of NPY-positive structures observed in the evanescent field divided by the size of the cell footprint.
From stacks of images, two-dimensional (x-y) trajectories were obtained by single-particle tracking using MIA software. Mean square displacement (MSD) in the x-y plane was computed using custom routines, and the diffusion coefficient \( D_{xy} \) was then calculated as \( D_{xy} = s/4 \), with \( s \) being the slope of the linear fit to the first 15 points of the MSD curve. To measure \( D_{xy} \) variations along a given trajectory, a rolling analysis window of 2.5 s was used. For each position of the window, \( D_{xy} \) was calculated as described above, and the obtained value was assigned to each point of the trajectory included in this window. Since each point of the trajectory appears in different windows, several \( D_{xy} \) values were assigned to each point and finally averaged. Immobilization periods were defined as portions of trajectories during which \( D_{xy} \) drops below a threshold value \( D_{xy \text{ min}} = 5 \times 10^{-4} \mu m^2 \cdot s^{-1} \), i.e., five times the minimal \( D_{xy} \) measurable with our experimental setup (estimated from the tracking of fluorescent beads immobilized on the coverslip) (Huet et al., 2006). As random walk trajectories may also feature periods of slow motion, we used the duration of immobilization periods as a further criterion of SG attachment. Survival curves for immobilization times (i.e., the cumulative number of immobilization events per trajectory plotted as a function of their minimum duration) were computed for each cell and averaged over the different cells. The best fit for the curves was obtained with the sum of two exponentials, \( N(t) = N_1 \exp(-t/\tau_1) + N_2 \exp(-t/\tau_2) \), using SigmaPlot (Systat Software). \( N_1 \) and \( N_2 \) represent the abundance of each component, and \( \tau_1 \) and \( \tau_2 \) their time constants, respectively. The long-lasting component was taken as an index of SG stalling.

**Minimizing the cross talk between attachment to the PM and diffusive periods.** Based on the two characteristic times observed in survival plots of immobilization times, we calculated a threshold value for the immobilization duration to separate long-lasting immobilization periods from the poorly defined short-lasting immobilization periods. This threshold was calculated to minimize the cross talk between the two populations in the survival curves. The normalized distributions for the short \( (D_{s}(t)) \) and long \( (D_{l}(t)) \) immobilization periods are given by the following equations:

\[
D_{s}(t) = \frac{1}{\tau_1} \exp\left(-\frac{t}{\tau_1}\right),
\]

\[
D_{l}(t) = \frac{1}{\tau_2} \exp\left(-\frac{t}{\tau_2}\right).
\]

For a given threshold \( t_{\text{thresh}} \), the cross talk between the two distributions is equal to the following:

\[
I(t_{\text{thresh}}) = \int_0^{t_{\text{thresh}}} D_{s}(t) \, dt + \int_{t_{\text{thresh}}}^{\infty} D_{l}(t) \, dt.
\]

By calculating the two integrals, we obtain the following:

\[
I(t_{\text{thresh}}) = 1 + \exp\left(-\frac{t_{\text{thresh}}}{\tau_1}\right) - \exp\left(-\frac{t_{\text{thresh}}}{\tau_2}\right).
\]

It is thus possible to obtain the threshold duration which minimizes the cross talk between the two populations:

\[
t_{\text{thresh}} = \tau_1 \times \tau_2 \ln\left(\frac{\tau_2}{\tau_1}\right).
\]

Based on this calculation, we obtained threshold durations between 6 and 8.9 s for the different experiments. Subtrajectories corresponding to directed motion were manually selected; stalled periods include all immobilization periods lasting longer than the threshold value defined above. The remaining subtrajectories were considered as diffusive.

**Detection of false positive stalling periods within simulated Brownian trajectories and diffusing fluorescent beads.** Three-dimensional random walk trajectories were simulated as described by Huet et al. (2006). The diffusion coefficient \( (25 \times 10^{-4} \mu m^2 \cdot s^{-1}) \), length of the trajectories (10 to 20 s) and time step between consecutive positions (0.1 s) were chosen to match the values obtained for SGs. Stalling events were detected with the approach used for SGs. The survival curves for immobilization times were based on the simulation of 3000 trajectories.

Time-lapse stacks displaying the diffusion of latex fluorescent beads (500 nm in diameter) within a glycerol/water mix (75–80% of glycerol by volume) were acquired by TIRF microscopy. The beads were tracked using the MIA software. The average diffusion coefficient of these beads \( (D_{xy} \approx 130 \times 10^{-4} \mu m^2 \cdot s^{-1}) \) was approximately five times higher than the value measured for SGs in control conditions. However, to properly estimate the contribution of false positive stalling events within the SG survival curves using the analysis of the bead dynamics, it was necessary to match the ratio between the average diffusion coefficient and the threshold value \( D_{xy \text{ min}} \) for the beads and the SGs. Consequently, stalling periods within bead trajectories were detected with a threshold \( D_{xy \text{ min}} = 25 \times 10^{-4} \mu m^2 \cdot s^{-1} \) and not \( 5 \times 10^{-4} \mu m^2 \cdot s^{-1} \), as used for the SGs. The survival curves for immobilization times were based on 69 bead trajectories whose duration matches that of SGs (16 s).

**Estimation of the diffusion anomaly.** In case of anomalous subdiffusion, the MSD curves can be fitted by the following:

\[
\text{MSD} \propto D \Delta t^\alpha,
\]

with \( \alpha \leq 1.0 \). In the logarithmic representation, Equation 1 becomes

\[
\log(\text{MSD}) \propto \alpha \log(\Delta t).
\]

Thus, the MSD curves calculated from the SG trajectories were plotted in logarithmic representation and fitted using a linear regression, the slope of the fit corresponding to the anomaly parameter \( \alpha \). For the estimation of the diffusion anomaly during stalling and nonstalling periods, the MSD curves were calculated for stretches of trajectories lasting \( >5s \). A median MSD was calculated for each cell, and the fit was performed over the first 25 points of each MSD curve.

**Measuring peripheral SG retention by photoconversion of NPY-Dendra2.** NPY was fused to Dendra2. Dendra2 is a photoactivable fluorescent protein whose emission spectrum shifts from green to red upon brief UV light exposure (Chudakov et al., 2007). A 405 nm diode laser (50 mW; Laser Components) was coupled to our TIRF microscope and aligned with the argon laser using a dichroic mirror. Two days after transfection with NPY-Dendra2, Myrip siRNA, or Myrip constructs, the cells were observed by TIRF with a penetration depth of 100 nm. A 2 s exposure to UV evanescent light was sufficient to photoconvert 64 ± 8% (n = 15 cells) of the green vesicles present in the TIRFIM field. Red signals three times above the background were obtained. Increasing the photoactivation time over 2 s did not significantly increase the red signal but increased the variability. After photoconversion, images were acquired for 1 min at 0.5 Hz using an exposure time of 200 ms. Dendra2 was not significantly photoconverted by such illumination at 514 nm. Images were quantitated using ImageJ. For each cell, the background was measured in areas devoid of SGs and subtracted to all pixel values. Cell boundaries were manually drawn, and the total fluorescence intensity was integrated over the depicted area. Alternatively, images were segmented using MIA software, and the number of vesicles was counted on each frame. Photobleaching was estimated on live cells by measuring the fluorescence decay of immobile SGs, yielding an upper bound estimate of bleaching since immobile SGs are generally attached to the PM and thus close to the interface where the TIRF field is generated. Diffusion of NPY-mRFP within individual SGs may also contribute to the diminution of the fluorescence signal over time. However, this fluorescence loss is likely to be the same under the different conditions since we found, using electron microscopy, that Myrip siRNAs do not change the size of SGs.

**Simulation of the photoconversion experiments.** To simulate the photoconversion experiments, we built up a kinetic model in which SGs can occupy different states: "C" for SGs localized at the cell center, "A" for SGs in the actin cortex, and "D" for SGs attached to the PM. TIRF microscopy does not allow imaging all the SGs present at the cell cortex but only those which are localized in the vicinity of the glass coverslip. To account for this limitation, we included in the model two peripheral SG populations, only one of them being accessible by TIRF imaging.
Due to the short evanescent depth used for photoconversion (−80 nm at 405 nm) most of the photoconverted NPY-Dendra labeled SGs were considered as docked at the PM immediately after the photoconversion. We used this initial condition for starting our simulations. Since the evanescent depth used for imaging is longer than the one used for photoconversion (100 nm instead of 80 nm), we considered that we were imaging all the docked SGs and 25% of the SGs located in the actin cortex. Because this proportion was of SGs located in the actin cortex which can be imaged using a 100 nm evanescent depth could not be estimated precisely, we performed simulations in which the value of \( p_{\text{act}} \) ranged between 0 and 50%. In all cases, we observed a biphasic behavior, the fast phase corresponding to the equilibration between the pool at the PM and the one in the actin, and the slow phase associated with SG movement toward the cell interior. Varying \( p_{\text{act}} \) from 0 and 50% only led to a moderate change in the relative amplitude of the two phases (data not shown).

The exchange rates between the three states were derived from the analysis of SG dynamics by TIRF microscopy (Table 1). To estimate the rates \( k_i \) and \( \lambda_i \) corresponding to the transition \( C \rightarrow A \) and \( C \rightarrow A' \), with \( A' \) being the population of SGs in the actin-rich layer not visible in TIRFM, we had to estimate the relative SG populations in the different compartments at steady state. Based on the images obtained by electron microscopy, we observed that about two-thirds of the SGs display a cortical localization in control cells. Our analysis of SG mobility also showed that, in control conditions, the density of SGs in the actin cortex and at the PM are similar (\( A \approx D \)). Finally, we estimated the proportion of SGs imaged by TIRF as compared to the total number of SGs in the cell. On deconvolved wide-field images of NPY-GFP-expressing BON cells, we counted a total number of labeled vesicles of \( \sim 1500 \) SGs. Knowing that we observed \( \sim 150 \) SGs by TIRF, we considered that \( \sim 10\% \) of the total number of labeled SGs were visible with this approach. Together, these different results allowed estimating the steady SG populations in the different compartments. We obtained \( C \approx 33\% \), \( A \approx 28\% \), and \( A' \approx D' \approx 25.5\% \). Since \( k_i \approx k_{-i} \cdot A/C \) and \( k_i \approx k_{-i} \cdot A'/C \), at steady state, we could estimate that, in the control cells, \( k_i \approx k_{-i} \cdot 0.24 \) and \( k_i \approx k_{-i} \cdot 0.77 \).

Statistical analyses
Values are given as mean ± SE. The significance of differences between two conditions was calculated with Mann–Whitney U test or Student’s t test when data were normally distributed. For multiple comparisons, we used the Kruskal–Wallis test or ANOVA (normal distribution), followed by Dunn’s or Tukey’s post-tests, respectively, using GraphPad Prism version 5.04.

Results
Myrip recruits MyoVa on secretory granules
Myrip forms a complex with MyoVa and Rab27a and is therefore thought to mediate the recruitment of MyoVa onto SGs. To test this possibility, we measured the association of a GFP-tagged construct comprising the globular tail of MyoVa and the exons A, C, E and F (GFP-MyoVa tail) with SGs in an enterochromaffin cell line (BON) derived from a human carcinoid tumor. BON cells store various peptides and serotonin in SGs and secrete them (Kim et al., 2001). They express Rab27a, Myrip, and MyoVa and the three proteins are associated with SGs as in other endocrine cells (Fig. 1) (Desnos et al., 2003, 2007b). Under control conditions, 60% of the NPY- mRFP labeled SGs were decorated with a GFP-MyoVa tail (Fig. 1A–C,H). In contrast, in cells treated with a silencing RNA duplex (siRNA) directed to Myrri mRNA (Fig. 1G), the fraction of SGs labeled by GFP-MyoVa tail was reduced to \( \sim 20\% \) (Fig. 1D–F,H). Moreover, in \( \sim 60\% \) of Myrip-siRNA-treated cells, the GFP-MyoVa tail exhibited a marked soluble cytosolic pattern without any visible enrichment on intracellular structures (Fig. 1J,F). Such increased cytosolic distribution was observed only in 22% of control cells. Overall, the intensity of the cytosolic GFP-MyoVa tail in cell regions devoid of SGs was doubled in Myrip-siRNA-treated cells compared to control cells (mean fluorescence intensity, control cells, 288 ± 32; Myrip knockdown cells, 595 ± 58; \( p < 0.0001 \); \( n = 30 \) cells). The association of MyoVa with SGs was rescued by expressing an siRNA-insensitive Myrip construct, arguing against an off-target effect of Myrip siRNAs (Fig. 1H). These observations indicate that Myrip is needed for the recruitment of MyoVa on SGs.

The intracellular distribution of SGs depends on Myrip
Next, we analyzed the effect of Myrip on the intracellular distribution of SGs. In the majority of control cells, SGs accumulate at the cell periphery and in cell extensions (Figs. 1, 2A, left). In contrast, SGs were evenly distributed in the cytoplasm or concentrated near the nucleus (Fig. 2A, middle, right) in most of the Myrip knockdown cells and in cells in which the Rab-binding domain of Myrip (Myrip-RBD) was expressed to inhibit the binding of endogenous Myrip to Rab27a (Fig. 2B). This effect was first evaluated by visual inspection of cells and manual determination of the fraction of cells exhibiting at least one cell extension with a SG density higher than in the cell center (Fig. 2B). A more quantitative analysis of the mean distance of SGs to the cell center confirmed these results (Fig. 2C). Noteworthy is that the effect of Myrip silencing on SG distribution was rescued by expressing a siRNA-insensitive Myrip construct (Fig. 2B,C). Finally, BON cells were imaged by TIRF to image NPY-labeled SGs present in the subplasmalemmal region (Huet et al., 2006; Desnos et al., 2007b). Most of the SGs observed by TIRFM are localized within the actin-rich cell cortex or at the PM (Oheim and Stuhmer, 2000; Johns et al., 2001; Huet et al., 2006). Myrip silencing and overexpression of Myrip-RBD significantly diminished the number of SGs present in the evanescent field, compared with control cells (Fig. 2D). In contrast, overexpression of full-length Myrip (Myrip-FL) but not of a C-terminally truncated Myrip (Myrip-ΔC, 1–665) had a tendency to increase SG density in the subplasmalemmal region (Fig. 2D), suggesting that the C-terminal region of Myrip contributes to SG retention, although it is not needed for the interaction with MyoVa (Desnos et al., 2003; Kuroda and Fukuda, 2005). We conclude that Myrip promotes the recruitment or the retention (“capture”) of SGs near the PM.

To characterize the effect of Myrip on SG distribution by independent means, we measured the distance of SGs to the PM by electron microscopy. BON cells were treated for 3 d with control siRNAs or Myrip siRNAs, fixed by high-pressure freezing and embedded after cryosubstitution. This technique was shown to preserve the cell architecture and the position of vesicles with respect to the PM (Siksou et al., 2007). Electron-dense SGs were easily identified on EM pictures (Fig. 3A), and their distance to the PM was determined. In agreement with TIRFM data, the number of SGs positioned within 300 nm of the PM was reduced in Myrip knockdown cells, compared with control cells (Fig. 3B). Interestingly, the effect of Myrip silencing on the number of SGs very close to the PM (<50 nm) or morphologically attached to the PM (<15 nm) was even more pronounced. Noteworthy, Myrip silencing had no effect on the size of SGs (not shown) or on the total number of SGs (cells treated with control siRNAs, 2.17 ± 0.27 SG/μm², \( n = 32 \) cells; Myrip knockdown cells, 2.32 ± 0.54 SG/μm², \( n = 30 \) cells).

In agreement with this defect in SG recruitment at the PM, Myrip silencing reduced the secretion of serotonin in BON cells (Fig. 2E), as it does in other neuroendocrine cells (Fukuda and Kuroda, 2002; Desnos et al., 2003; Waselle et al., 2003; Imai et al., 2004; Ivarsson et al., 2005; Mizuno et al., 2011).
Myrip promotes the peripheral retention of SGs

To determine whether the decreased SG density beneath the PM was due to impaired delivery or retention of SGs, we designed an assay to measure the time spent by SGs in the evanescent field. A 405 nm laser diode was coupled to our TIRF microscope and used to photoconvert SG-targeted NPY-Dendra2. A short UV illumination induced a green-to-red shift of the fluorescence emission of Dendra2 in the subset of SGs present in the evanescent field (Fig. 4A, B). In agreement with the effect of Myrip on SG density in the TIRF area (Fig. 2D), Myrip-FL, but not Myrip-H9004C, increased the integrated fluorescence of NPY-Dendra2 measured immediately after photoconversion (data not shown). In contrast, Myrip knockdown reduced this value. Both the fluorescence intensity (Fig. 4C) and the number of visible SGs (Fig. 4D) decreased over time. The decay is likely to reflect SG motion toward the cell interior since under resting conditions secretion is extremely low and cannot contribute to the observed fluorescence decay. Myrip-FL slowed the decay of both the number of labeled SGs and fluorescence intensity, whereas Myrip-siRNA accelerated it, indicating that Myrip promotes the retention of SGs at the cell periphery. Interestingly, Myrip-H9004C slowed the decay of the number of SGs kept in the evanescent field but accelerated the fluorescence loss, suggesting that the overexpression of Myrip-ΔC increased the distance of the SGs to the cell interior.

**Figure 1.** Myrip recruits myosin Va onto SGs. A–F, To evaluate the association of MyoVa to SGs, BON cells were successively transfected with NPY-mRFP (B, E) and with GFP-MyoVa tail (A, D) 3 d and 16 h before cell fixation, respectively. In addition, cells were transfected with luciferase-targeting siRNAs as a control (A–C) or with Myrip siRNA1 (D–F). Z-stacks of epifluorescence images were restored by deconvolution. G, Knockdown of Myrip expression. Three days after transfection of siRNAs, Myrip levels were analyzed by immunoblotting in nontransfected (NT) cells, in cells treated with a control siRNA, with siRNAs directed to Myrip mRNA, and in cells overexpressing Myrip-FL. Myrip levels were lowered by 80 ± 1.8% (Myrip-siRNA1) or by 63 ± 1.8% (Myrip-siRNA2; n = 3). The result of a typical experiment is shown. H, Shown is the percentage of NPY-mRFP-labeled SGs that were also decorated with a GFP-MyoVa tail, a construct comprising the globular tail of MyoVa and the exons A, C, E, and F. The targeting of the GFP-MyoVa tail to SGs was reduced by Myrip silencing and rescued by expressing a siRNA1-insensitive Myrip construct (rescue) (p < 0.0001, ANOVA). Asterisks refer to a Tukey’s post-test; ***p < 0.001; n = 15 cells from two experiments. Cells expressing the rescue construct were identified using an anti-c-Myc antibody and an Alexa-350-labeled secondary antibody. Noteworthy is that the expression of the GFP-MyoVa tail was not reduced by Myrip silencing. I, J, Moreover, in ~60% of Myrip-siRNA1-treated cells, the GFP-MyoVa tail exhibited a marked soluble cytosolic pattern without any visible enrichment on intracellular structures. K, Coimmunoprecipitation of Myrip and myosin Va. Coimmunoprecipitation of the GFP-MyoVa tail with myc-tagged Myrip after expression in BON cells is shown. Immunoprecipitation was done with an anti-c-Myc antibody and immunoblotting with anti-MyoVa antibodies. One-fifteenth of cell extracts and one-fourth of immunoprecipitates were analyzed. The positions of endogenous MyoVa and the MyoVa tail are shown by an arrowhead and an arrow, respectively. Scale bars: 5 μm.
the PM while keeping them at the cell periphery. An increased photobleaching of SGs could also account for the higher fluorescence loss observed upon expression of Myrip-ΔC. SGs would bleach faster if they were closer to the PM and therefore exposed to a greater flux of photons. However, this possibility seems unlikely because the initial fluorescence intensity of individual SGs was not modified by Myrip-ΔC expression, as would be the case if their distance to the PM was lower (data not shown).

Using TIRF imaging, we also counted the number of NPY-GFP-labeled SGs that entered or left the evanescent field during time series of TIRF images (Fig. 5A). We observed that knocking down Myrip (Fig. 5B) increased both the rates of SG appearance and disappearance. MyoVa silencing induced similar, but less pronounced, effects (Fig. 5C). The increased rate of SG appearance may result from the higher density of SGs in the cell center observed in Myrip knockdown cells. In contrast, the increased rate of SG disappearance cannot be accounted for by changes in the number of SGs participating in the trafficking, but rather reflects an increase in the rate constant that controls the movement of SGs from the actin-rich layer to the cell center (see the Discussion section for a discussion of these data within the framework of the model depicted in Fig. 10; for the quantification of the different kinetic parameters, see Table 1).

Figure 2. Myrip controls the distribution of SGs and their recruitment at the cell periphery. A, BON cells were transfected with siRNA duplexes or myc-tagged Myrip constructs, and intracellular distribution of SGs was analyzed 3 d later. SGs were stained using an anti-chromogranin A/B antibody or by means of NPY-GFP expression. SGs are generally enriched at the cell periphery and in cell extensions (left). However, upon Myrip silencing or Myrip-RBD expression, SGs are frequently scattered in the cell (middle) or concentrated in the perinuclear region (right). B, Shown are the mean (± SE) percentages of cells with a marked enrichment of SGs at the cell periphery observed in control conditions (NT, nontransfected cells; n = 139 cells from 4 independent experiments; control, n = 255 cells from 6 experiments), upon expression of Myrip-RBD (n = 109 cells from 4 experiments), and upon transfection of Myrip-siRNA1 (n = 190 cells from 6 experiments), Myrip-siRNA2 (n = 125 cells from 3 experiments), and Myrip-siRNA1 plus an myc-tagged siRNA1-insensitive Myrip construct (rescue; n = 81 cells from 2 experiments). Inset, schematic representation of the Myrip constructs used in this study. p < 0.0001 (ANOVA); ***p < 0.001 (Tukey’s post-test); ns, nonsignificant. C, To quantify the cellular distribution of SGs, epifluorescence images of NPY-GFP-expressing cells were segmented using MIA software. For each cell, the mean distance of the thresholded pixels to the centroid of the cell footprint was computed. The graph shows the mean (± SE) of the values obtained in 10 cells for each condition. p = 0.0388 (ANOVA). D, Myrip controls the juxtamembrane SG density. BON cells were transfected with an empty vector (Control, 39 cells from 2 experiments) or a control siRNA duplex (Control-siRNA, 107 cells from 7 experiments), with vectors encoding Myrip-RBD (42 cells from 2 experiments), Myrip-FL (38 cells from 2 experiments), Myrip-ΔC (38 cells from 2 experiments), or with Myrip-siRNA1 (96 cells from 7 experiments) and a Myrip rescue construct (20 cells from 2 experiments); they were also cotransfected with pNPY-mRFP to label SGs. Cells were imaged by TIRFM. Inset, A representative TIRFM image of a BON cell expressing NPY-mRFP. Data are expressed as the mean ± SE of values obtained in the different cells; p < 0.0001 (Kruskal–Wallis). E, Effect of Myrip silencing on secretion. BON cells were treated with control siRNAs or with Myrip-targeting siRNAs. Three days later, they were loaded with [3H]serotonin, washed, and stimulated with Locke’s solution supplemented with 3.7 mM BaCl2 (black and white bars) for 10 min or incubated in nonstimulating bathing medium for the same duration (dashed bars). Released serotonin was measured in the extracellular medium and expressed as a percentage of serotonin contained in the cells (mean ± SE of 6–15 wells). Scale bars: 5 μm. *p < 0.05; ***p < 0.001.
Myrip silencing interferes with SG motion along microtubules

To determine how Myrip controls the retention of SGs at the cell periphery, we characterized SG mobility in the subplasmalemmal region by TIRF imaging and single-particle tracking. The mean lateral diffusion coefficient (\(D_{xy}\)), an index of the mobility of SGs, on entire trajectories was significantly higher in Myrip-silenced cells (\(D_{xy} = 92.3 \pm 7.2 \times 10^{-4} \mu m^2 \cdot s^{-1}\); mean \(\pm SE\); 653 SGs from 10 cells) than in control cells (\(D_{xy} = 24.3 \pm 5.2 \times 10^{-4} \mu m^2 \cdot s^{-1}\); mean \(\pm SE\); 761 SGs from 10 cells; \(p = 0.007\)). The distribution of mean \(D_{xy}\) values computed on entire trajectories is shown in Figure 6A and indicates two major effects of Myrip silencing: (1) an increased proportion of SGs with a very high \(D_{xy}\) value (\(>50 \times 10^{-4} \mu m^2 \cdot s^{-1}\)) and (2) a 50% reduction in the percentage of almost-immobile SGs (\(D_{xy} < 5 \times 10^{-4} \mu m^2 \cdot s^{-1}\)).

To further characterize the effect of Myrip on fast-moving SGs, we manually selected subtrajectories characterized by fast \((D_{xy} > 50 \times 10^{-4} \mu m^2 \cdot s^{-1})\) and directed motion. Such trajectories are severely reduced upon nocodazole treatment, indicating that they correspond to SGs moving along microtubules (Desnos et al., 2007b). The proportion of time spent in a directed motion was increased from 3.8 \(\pm 0.4%\) in control cells to 11.5 \(\pm 1.5%\) in Myrip knockdown cells (mean \(\pm SE\); \(p = 0.002\)). During these periods, SG velocity was 0.39 \(\pm 0.02 \mu m \cdot s^{-1}\) in control cells and 0.63 \(\pm 0.03 \mu m \cdot s^{-1}\) in Myrip-silenced cells (mean \(\pm SE\); \(p = 0.0002\)), and the distribution of velocity values was shifted to higher values by Myrip silencing (Fig. 6B). In addition, the length of microtubule-based runs was increased in Myrip knockdown cells (1.49 \(\pm 0.11 \mu m\) compared to control ones (1.15 \(\pm 0.06 \mu m\); \(p = 0.04\)).

Myrip promotes SG attachment to the plasma membrane

Next, we focused on stalled SGs. While diffusing with an average \(D_{xy}\) equal to \(25 \times 10^{-4} \mu m^2 \cdot s^{-1}\) in control conditions, SGs frequently display periods of highly restricted mobility with \(D_{xy}\) values as low as \(10^{-4} \mu m^2 \cdot s^{-1}\). Several authors have suggested that these stalling periods correspond to the attachment of SGs to the PM (Huet et al., 2006; Desnos et al., 2007b; Nofal et al., 2007; Karatekin et al., 2008; Yizhar and Ashery, 2008; Ostrowski et al., 2010). Before analyzing the consequences of impairing Myrip on this class of SG motion, we first investigated further whether transient stalling could be used as a criterion of attachment to the PM. The fluctuations in SG mobility were monitored by measuring the variations of \(D_{xy}\) along the SG trajectories, using a rolling analysis window, and immobilization periods were defined as portions of trajectories during which the \(D_{xy}\) value dropped below a threshold of \(D_{xy} = 5 \times 10^{-4} \mu m^2 \cdot s^{-1}\) (Fig. 7A, B). The choice of this threshold was driven by the observation that, within the 2 s preceding exocytosis, most of the SGs, which are probably attached to the PM, display a \(D_{xy}\) value of \(<5 \times 10^{-4} \mu m^2 \cdot s^{-1}\) (Desnos et al., 2007b). However, due to the stochastic nature of Brownian motion, even purely diffusive trajectories characterized by an average diffusion coefficient of \(10^{-3} \mu m^2 \cdot s^{-1}\) may exhibit short periods during which \(D_{xy}\) is lower than \(D_{min}\). To distinguish such false positive stalling events from real periods of SG attachment, it was thus necessary to set a minimum crossing time, \(t_{min}\). Therefore, we compared the distribution of immobilization times between SG tracks and simulated Brownian trajectories. The survival curves displayed on Figure 7C show that with \(t_{min}\) set to 5 s, false positive stalling events were contributing by \(<1%\) to the events detected for SGs. Stalling periods were also detected on trajectories of latex beads diffusing in a glycerol/water mix, and we confirmed that, with this minimum crossing time, the probability of detecting stalling events was at least 100 times higher for SGs than for beads (Fig. 7C). To further substantiate the fact that stalling periods reflect a constrained behavior, we compared the anomaly of the diffusion between stalling periods longer than 5 s and the rest of SG trajectories. The anomaly parameter \(\alpha\) quantifies the restriction of the diffusion: the lower the value of \(\alpha\), the more restricted the diffusion, with \(\alpha\) ranging between 0 and 1. The stalling periods were characterized by a strong restriction of SG motion, with \(\alpha \approx 0.4\), in agreement with
an attachment to the PM. In contrast, outside these stalling periods, SG diffusion showed only moderate restriction with $\theta_{H9251}/H_{11015}0.8$ (Fig. 7D). Finally, using the fact that, by TIRF microscopy, changes in SG fluorescence intensity report changes in the position along the vertical axis, we measured the difference in the altitude of SGs between two successive periods of immobilization. We found that, in most of the cases, the difference was $H_{11021}/H_{20}$ nm (i.e., the resolution of our microscope in the $z$-axis) (Fig. 7E).

The data thus suggest that immobile SGs are at the same altitude, i.e., at the PM. Survival curves for SG immobilization times were best fitted with two exponentials (Fig. 7C). It is noteworthy that both the short-lasting (characteristic time, $\sim 4$ s) and the long-lasting (characteristic time, $\sim 15$ s) components were observed in single cells and thus cannot be attributed to the heterogeneity of the population of cells (Fig. 7C). If slow diffusion can contribute a significant fraction of the fast component of the survival curves, it is not the case of the slow component, which we therefore used as a refined criterion of SG attachment to the PM.

We reported previously that MyoVa promotes SG attachment to the PM (Desnos et al., 2007b), an effect that could be due to an increase in either the occurrence or the duration of attachment events. To resolve this issue, we analyzed the effect of MyoVa silencing on the distribution of immobilization times. MyoVa depletion reduced the occurrence of the long-lasting component but not its characteristic time ($H_{9270}2$) (Fig. 7G, I, J). This reduction in the occurrence of SG immobilization was observed with three different MyoVa siRNAs, indicating that it was not due to off-target effects. We conclude that MyoVa promotes the recruitment of SGs to the PM rather than the stability of the immobilized state.

Similarly, Myrip silencing reduced the occurrence of the long-lasting component (Fig. 7H, K). In addition, it reduced the characteristic time, $\tau_{H9270}$, of this immobilization process (Fig. 7H, L). It is noteworthy that survival curves were rescued by expressing a siRNA-insensitive Myrip construct arguing against an off-target effect of the siRNAs. Since Myrip silencing also reduced the number of SGs present in the TIRF area (Fig. 2D), its overall effect was to diminish the number of attached SGs per surface area by $H_{11011}/H_{65}$%. This conclusion is consistent with the data obtained by EM showing a severe reduction in the number of SGs morphologically attached to the PM in Myrip knockdown cells and accounts for the secretory defects observed in these cells.
Myrip restricts the diffusion of SGs within the actin-rich cortex

Next, we characterized the effect of Myrip on the mobility of SGs when they were neither stalled nor moving along microtubules (see Materials and Methods). During such periods, SGs display Brownian motion and are therefore diffusing with or within the actin-rich cortex (Fig. 7D) (Huet et al., 2006). Both Myrip and MyoVa silencing increased the proportion of time spent by SGs in the diffusive state (Fig. 8A) and the mobility of SGs in the diffusive state (Myrip knockdown cells, $D_{xy} = 22.6 \pm 1.3 \times 10^{-4} \mu m^2 \cdot s^{-1}$ vs control, $14.9 \pm 1.9 \times 10^{-4} \mu m^2 \cdot s^{-1}$; 10 cells, $p = 0.01$; MyoVa-silenced cells, $D_{xy} = 24.5 \pm 2.9 \times 10^{-4} \mu m^2 \cdot s^{-1}$ vs control, $14.0 \pm 1.5 \times 10^{-4} \mu m^2 \cdot s^{-1}$; 16 cells, $p = 0.003$). These results suggest that the Rab27/Myrip/myosin Va complex restricts the mobility of SGs within the actin-rich cell cortex, most likely by tethering SGs to actin filaments, to maintain SGs in the vicinity of release sites.

The C-terminal domain of Myrip participates in SG attachment to the plasma membrane

The above-mentioned difference between the effects of Myrip and MyoVa silencing on the duration of long-lasting SG immobilization suggested that Myrip may exert MyoVa-independent actions. To investigate this possibility, we overexpressed Myrip, Myrip-ΔC (which can interact with MyoVa), or Myrip-RBD (which does not bind to MyoVa) into BON cells and measured its effect on SG mobility. Overexpression of Myrip-RBD or Myrip-ΔC increased the mean diffusion coefficient, $D_{xy}$, computed on entire trajectories (Fig. 9A) and reduced the percentage of SGs with very low $D_{xy}$ values ($<5 \times 10^{-4} \mu m^2 \cdot s^{-1}$) by ~35% compared to control cells or to cells expressing Myrip-FL ($p < 0.0001$, ANOVA). We then measured the duration of immobilization events along trajectories and found that Myrip-RBD and Myrip-ΔC reduced the occurrence of long-lasting immobilization periods by 53% and 27%, respectively. Myrip-ΔC also decreased the characteristic time of long-lasting immobilization compared to Myrip-FL ($p = 0.0004$), suggesting that the C-terminal region of Myrip contributes to the stability of SG attachment at the plasma membrane.

Discussion

In this study, we identified Myrip as a factor that couples the capture of SGs in the actin-rich cell cortex and their attachment to the PM, a process mandatory for exocytosis. Myrip thus plays a major role in the accumulation of SGs at the cell periphery and in the replenishment of the releasable pool of SGs.

Inferring the functional state of SGs from their mobility

To understand SG recruitment at release sites, our approach was to track single SGs, to analyze their motion along trajectories and to infer their functional state from their behavior. Previous studies have shown the existence of three classes of SG motion and
suggested that (1) rapid and asymmetric trajectories correspond to SGs moving along microtubules, (2) stalled SGs are attached to the PM, and (3) the remaining trajectories correspond to random walks within the actin-rich cell cortex (Huet et al., 2006; Desnos et al., 2007b).

One difficulty is to distinguish random walks from stalling periods. Here, we used a method that combined a selection of immobilization periods characterized by a low $D_{xy}$ value with the analysis of their durations to provide an index of SG confinement. Analyzing the distribution of immobilization times helped to distinguish attachment (the long-lasting component of the survival curves) and slow random walk. Subtrajectories selected by $D_{xy}$ thresholding and longer than 5 s were characterized by a nonlinear relationship between MSD and time, indicating a subdiffusive behavior. This is in agreement with our previous finding that these periods feature a negative autocorrelation of the motion along the z-axis, suggesting that during these periods, SGs do not move freely but are attached (Desnos et al., 2007b). SG immobilization could result from binding to other structures than the PM, especially to actin filaments. However, actin structures are not immobile (Desnos et al., 2007b) (our unpublished observations), and thus cannot support stable SG immobilization. Furthermore, we found that SGs that un-

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**Figure 7.** Myrip promotes SG attachment to the plasma membrane. **A, B.** Example of a single SG trajectory and detection of immobilization periods. The $x$–$y$ displacement of a single SG is shown in A. The initial point of the trajectory is depicted by a red dot. The $D_{xy}$ values were computed along the trajectory using a rolling analysis window and plotted against time in B. Below a threshold value $D_{xy} = 5 \times 10^{-4}$ $\mu$m$^2$ s$^{-1}$ (red line), SGs were considered immobile. In this example, the SG displayed a constrained behavior during two subtrajectories. Period 2 was categorized as an immobilization period. **C.** The cumulative distribution of the number of immobilization events identified in control trajectories (filled circles, 32 cells) is plotted as a function of the minimum duration of immobilization. The best fit was obtained with the sum of two exponentials, indicating the existence of two different processes (black line, slow component; blue line, fast component). A similar distribution is found in single cells; an example is shown in gray. Ninety-nine percent of the immobilization periods detected in simulated Brownian trajectories (red) and fluores-cent bead trajectories (green) were shorter than 5 s, and their distribution can be fitted by a single exponential.

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**Table 1.** Summary of the results of the statistical analysis. p values were calculated using a Dunn’s multiple comparison test. $\dagger$ indicates a significant difference from control, $\dagger\dagger$ from siRNA1, $\dagger\dagger\dagger$ from siRNA2, $\dagger\dagger\dagger\dagger$ from siRNA3. $p < 0.05$ is considered significant.

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**Figure 8.** Immunoblot analysis of Myrip and MyoVa expression in Myrip and MyoVa knockdown cells. **A.** Western blot analysis of Myrip and MyoVa expression in control (C), Myrip-siRNA1 (M), Myrip-siRNA2 (M2), and Myrip-siRNA3 (M3) cells. **B.** Quantification of Myrip and MyoVa expression in control (C), Myrip-siRNA1 (M), Myrip-siRNA2 (M2), and Myrip-siRNA3 (M3) cells. **C.** Western blot analysis of MyoVa and Myrip expression in rescue cells (R). **D.** Quantification of MyoVa and Myrip expression in rescue cells (R). **E.** Western blot analysis of MyoVa and Myrip expression in MyoVa knockdown cells (M). **F.** Quantification of MyoVa and Myrip expression in MyoVa knockdown cells (M). **G.** Western blot analysis of MyoVa and Myrip expression in MyoVa rescue cells (R). **H.** Quantification of MyoVa and Myrip expression in MyoVa rescue cells (R).
A robust criterion of SG attachment to the PM. We think that this approach provides a simple compartment model to describe SG delivery to the cortex. Different parameters controlling SG dynamics at the cell cortex, such as actin filaments (density ρA) and microtubules (density ρD), were identified. The remaining parts of the trajectories were considered diffusive. A, B, Proportion of time spent in the diffusive (A) or immobile (B) state for SGs from control, Myrip knockdown cells, and MyoVa knockdown cells (10–20 cells per condition). The mean length of trajectories was ~17 μm in control cells, 14 μm in MyoVa knockdown cells, and 10 μm in Myrip knockdown cells. *p < 0.05; **p < 0.01; ***p < 0.001.

dergo multiple immobilization periods during their trajectory generally immobilize at the same altitude. As actin filaments are present at different altitudes, this observation also suggests that SGs immobilize at the PM. We think that this approach provides a robust criterion of SG attachment to the PM.

SG mobility is high in BON cells compared to primary neuromedin cells such as adrenal chromaffin cells and pancreatic β cells (Degtyar et al., 2007; Michael et al., 2007). As we can collect many examples of the different possible behaviors and transitions between different classes of motion, this particular feature of BON cells facilitates the analysis of SG recruitment at exocytotic sites.

A compartment model to analyze Myrip-dependent SG trafficking

Our data indicate that Myrip has profound effects on SG distribution, mobility, and membrane attachment. To evaluate the different parameters controlling SG dynamics at the cell cortex, we designed a simple compartment model to describe SG delivery to the PM (Fig. 10 A). The model posits that SGs are transported from the cell center (C) to the cell periphery (most likely along microtubules) and then enter a thin (~1 μm diameter) actin-rich layer (A) into which they diffuse until they find an attachment site (D). As suggested by Huet et al. (2006), we considered that the transition C→D is not allowed, i.e., that SGs coming from the cell center cannot reach the PM directly. The values of A (0.22 ± 0.025 SGs/μm²) and D (0.22 ± 0.025 SGs/μm²) were derived from the density of SGs observed by TIRFM (density = A + D) and from the proportion of the time spent in the stalled or diffusive states, as deduced from the analysis of Dxy along trajectories (Figs. 7, 8). At steady state, the number of SGs arriving in the TIRF area should be similar to the number of SGs exiting the TIRF field; hence, k-1A = k+1C. Consistently, SG appearance and disappearance rates were found to be similar (Fig. 5). From the rate of SG disappearance (k-1A) and the measured value of A, we obtained k-1A = 0.016 ± 0.002 s⁻¹. Similarly, at steady state, the number of SGs diffusing in the cell cortex that immobilize at the PM is compensated by a similar number of SGs that come apart. Thus, k+1A = k-1D. The detachment rate constant k̂-2 determines the characteristic lifetime, τ, of the immobilized state (τ = 1/k̂-2), which we inferred from the survival curves shown in Figure 7 (number of stalled SGs still immobilized after a time t). The exponential fit to the slow component of these curves gave a value for k̂-2 (k̂-2 = 0.056 ± 0.007 s⁻¹), and hence for k̂1, the rate constant of SG attachment to the PM [k̂1 = k̂-2D/A] = 0.082 ± 0.012 s⁻¹.

Using this compartment model, we simulated the photoconversion experiment shown in Figure 4 (Fig. 10 B). Both the experimental and simulated curves display a biphasic behavior which can be well fitted with the sum of two exponential components. The fast one (characteristic time, 6.5 s, compared with 5.9 s for the experimental curve) represents the equilibration of A and D.
The consequences of Myrip and MyoVa knockdown on SG dynamics at the cell cortex were analyzed using the same compartment model (Table 1). Myrip silencing induced a marked increase in $k_{-2}$ and $k_{-1}$ values and a decrease in $k_{2}$ values, indicating decreased retention in the actin-rich cell cortex, decreased transport toward attachment sites at the PM, and lowered stability of SG attachment. Lowering MyoVa levels induced a slight increase in $k_{-1}$ and a decrease in $k_{2}$, but no significant change in the rate of SG detachment from the PM. Therefore, we conclude that the effects of Myrip can be partly accounted for by the recruitment of MyoVa and that Myrip also exerts a MyoVa-independent role in SG attachment to the PM. In addition, the effect of Myrip and MyoVa on $k_{2}$ suggest that the motor activity of MyoVa promotes the delivery of SGs at the PM, and therefore that MyoVa does not merely act as a tether.

### Myrip and MyoVa mediate the retention of SGs at the cell periphery

Our observations demonstrate that Myrip and MyoVa mediate the retention of SGs at the cell periphery. Three mechanisms are likely involved. First, Myrip mediates the association of SGs with the actin-rich layer. This conclusion is supported by the increase in the rate constant $k_{-1}$ and the increased mobility of the SGs that were diffusing within the actin-rich layer observed in Myrip knockdown cells. Both MyoVa (Desnos et al., 2007b) and Myrip may mediate the association of SGs with actin filaments. The C-terminal region of Myrip, which directly interacts with F-actin (Fukuda and Kuroda, 2002; Desnos et al., 2003), contributes to the restriction in SG mobility during diffusive periods (see Results) (Fig. 9), suggesting that Myrip directly tethers SGs to actin.

The second mechanism that contributes to SG retention at the cell periphery is suggested by the effect of Myrip on SG motion along microtubules. SGs move faster along MTs in the absence of Myrip (present study) or MyoVa (Desnos et al., 2007b), suggesting that powering SG motion along microtubules is made more difficult in the actin-rich environment by the interaction of Myrip or MyoVa with F-actin. This interaction may therefore promote the detachment of the newly arrived SGs from microtubules. This capture mechanism involving dissociation from microtubules and retention in the actin-rich layer is likely to have broad implications in the polarized distribution of various organelles, such as melanosomes, calcium stores, or recycling endosomes (Desnos et al., 2007a; Wagner et al., 2011).

The third mechanism underlying SG retention in the subplasmalemmal region is the attachment of SGs to the PM. The probability of SG attachment to the PM (the A-to-D transition in Figure 10) is decreased in both Myrip and MyoVa knockdown cells, indicating that the two proteins promote SG delivery to release sites. This effect is probably due to the motor activity of MyoVa directed toward the PM, where F-actin barbed ends are enriched. In addition, we observed a specific effect of Myrip de-

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**Table 1. The dynamics of SGs quantified according to the model depicted in Figure 10**

<table>
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<tr>
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<th>Control SiRNA</th>
<th>MyoVa SiRNA1</th>
<th>Myrip SiRNA</th>
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<td><strong>Density (SG/μm²)</strong></td>
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<td>0.23 ± 0.023**</td>
<td>0.35 ± 0.032</td>
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<td><strong>Docked (%)</strong></td>
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<td>35.6 ± 3.7**</td>
<td>24.4 ± 2.4***</td>
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<td><strong>A (SG/μm²)</strong></td>
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<td>0.08 ± 0.013***</td>
<td>0.08 ± 0.011**</td>
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<tr>
<td><strong>k₂ (s⁻¹)</strong></td>
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<td>0.045 ± 0.012*</td>
<td>0.040 ± 0.006*</td>
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<tr>
<td><strong>k⁻₁ (s⁻¹)</strong></td>
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<td>0.064 ± 0.001</td>
<td>0.088 ± 0.009***</td>
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<td><strong>k₋₁ (s⁻¹)</strong></td>
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<td>0.022 ± 0.002**</td>
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<tr>
<td><strong>k₅C (SG/μm²/s²)</strong></td>
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<td>0.0027 ± 0.0002</td>
<td>0.0087 ± 0.001**</td>
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<tr>
<td><strong>N/cells</strong></td>
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<td>10</td>
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SG density refers to the number of SGs detected by TIRFM. The density of SGs attached to the PM (D) or diffusing in the actin-rich layer (A) or in the cell center (C). Straight yellow lines represent microtubules along which SGs travel to reach or to leave the cell periphery. Thin lines represent the actin filament network. Transitions between the three states C, A, and D are governed by the constants $k₁$, $k₋₁$, $k₂$, and $k₋₂$. In agreement with previous findings (Huet et al., 2006), the transition between A and D is forbidden: SGs must dissociate from MTs and diffuse for a while within the actin cortex before they can attach at the PM. See Table 1 for a quantification of the rate constant values. Since the characteristic residency time at the cell periphery (A + D, ~60 s) is longer than the sum of the residency time in A (~15 s) and in D (~15 s), we conclude that SGs can undergo several local searches and successful membrane attachment periods before going back to C. B. Simulation of the photoconversion experiments. Shown is the comparison of the photoconversion experiment (control condition as in Fig. 5D, fitted with 2 exponentials; solid black line) and a simulation (gray) based on the model shown in A (for details, see Materials and Methods). The fast component reflects the equilibration between SGs attached to the PM and SGs diffusing in the actin-rich layer. The slow component represents the exit of SGs from the actin-rich layer.
pletion on the stability of SG immobilization at the PM, indicating that Myrip, but not MyoVa, is a component of the attachment machinery. This effect is mediated by the C-terminal region of Myrip since overexpression of Myrip-ΔC but not of full-length Myrip reduced the stability of SG immobilization.

Mizuno et al. (2011) previously characterized the effect of Myrip on the trafficking and exocytosis of SGs in insulin-secreting cells. In agreement with previous studies (Waselle et al., 2003; Ivarsson et al., 2005) and this report, they found that Myrip silencing reduced insulin secretion and SG recruitment at the cell periphery. Based on the effect of Myrip overexpression on the accumulation of SGs in the actin-rich layer, they suggested that Myrip only acts by tethering SGs to the actin-rich layer, via MyoVa. However, this observation supports the conclusion that Myrip tethers SGs to actin filaments (Desnos et al., 2003; Kuroda and Fukuda, 2005) (present study), but does not contradict the proposed additional role of Myrip in SG docking at the PM.

A direct role for Myrip in SG attachment to the PM

The fact that Myrip silencing and Myrip-ΔC, but not MyoVa, modify the duration of immobilization events indicates that the C-terminal region of Myrip is involved in the attachment of SGs to the PM. This region of Myrip interacts with components of the exocytosis (Goehring et al., 2007), a multimeric protein complex involved in tethering reactions between membrane compartments. The exocytosis targets several transport vesicles to the PM (Lipschutz and Mostov, 2002; He and Guo, 2009), and preliminary evidence supports its role in SG docking (Tsuboi et al., 2005). The reported effect of the C-terminal region of Myrip on SG secretion and melanosome distribution (Waselle et al., 2003; Imai et al., 2004; Ramalho et al., 2009) may therefore be related to the Myrip/exocyt interaction. Future studies will have to test the possibility that the exocyt complex mediates the effect of Myrip on SG docking. The SNARE proteins SNAP25 and syntaxin-1, as well as the syntaxin-1 partner Munc18 and the vesicular protein synaptotagmin-1, have also been implicated in SG docking at the PM (Toonen et al., 2006; de Wit et al., 2009). A key issue is to determine the relationship among Myrip, the exocytosis, and this SNARE-based docking complex. One possibility is that Myrip mediates the first interaction with the PM, in concert with the large exocyt complex, and that SNAREs and synaptotagmins then assemble to mediate short-range interactions between SGs and the PM and prime SGs for exocytosis. The fact that previously immobilized SGs undergo a 20 nm step toward the PM (Karatekin et al., 2008) or a small lateral movement (Degtyar et al., 2008) may therefore be related to the Myrip/exocyt interaction. Future studies will have to test the possibility that the exocyt complex mediates the effect of Myrip on SG docking. The SNARE proteins SNAP25 and syntaxin-1, as well as the syntaxin-1 partner Munc18 and the vesicular protein synaptotagmin-1, have also been implicated in SG docking at the PM (Toonen et al., 2006; de Wit et al., 2009). A key issue is to determine the relationship among Myrip, the exocytosis, and this SNARE-based docking complex. One possibility is that Myrip mediates the first interaction with the PM, in concert with the large exocyt complex, and that SNAREs and synaptotagmins then assemble to mediate short-range interactions between SGs and the PM and prime SGs for exocytosis. The fact that previously immobilized SGs undergo a 20 nm step toward the PM (Karatekin et al., 2008) or a small lateral movement (Degtyar et al., 2007) shortly before fusion supports this possibility and suggests that different-sized tethering complexes are sequentially formed during the secretory process (Karatekin et al., 2008).

References


