The Syntaxin 4 N Terminus Regulates Its Basolateral Targeting by Munc18c-dependent and -independent Mechanisms

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To generate and maintain epithelial cell polarity, specific sorting of proteins into vesicles destined for the apical and basolateral domain is required. Syntaxin 3 and 4 are apical and basolateral SNARE proteins important for the specificity of vesicle fusion at the apical and basolateral plasma membrane domains, respectively, but how these proteins are specifically targeted to these domains themselves is unclear. Munc18/SM proteins are potential regulators of this process. Like syntaxins, they are crucial for exocytosis and vesicle fusion. However, how munc18c and syntaxin 4 regulate the function of each other is unclear. Here, we investigated the requirement of syntaxin 4 in the delivery of basolateral membrane and secretory proteins, the basolateral targeting of syntaxin 4, and the role of munc18c in this targeting. Depletion of syntaxin 4 resulted in significant reduction of basolateral targeting, suggesting no compensation by other syntaxin forms. Mutational analysis identified amino acids Leu-25 and to a lesser extent Val-26 as essential for correct localization of syntaxin 4. Recently, it was shown that the N-terminal peptide of syntaxin 4 is involved in binding to munc18c. A mutation in this region that affects munc18c binding shows that munc18c binding is required for stabilization of syntaxin 4 at the plasma membrane but not for its correct targeting. We conclude that the N terminus serves two functions in membrane targeting. First, it harbors the sorting motif, which targets syntaxin 4 basolaterally in a munc18c-independent manner and second, it allows for munc18c binding, which stabilizes the protein in a munc18c-dependent manner.

To generate and maintain membrane polarization in epithelial cells it is essential that proteins are correctly sorted to their target membrane. Different protein machineries are involved in directing this polarized trafficking to the apical and basolateral membranes (1, 2). The final step in membrane targeting is fusion of the transport vesicle to the plasma membrane, which is mediated by members of the SNARE protein family. It has now been well established that epithelial cells express different forms of the SNARE protein syntaxin (3, 4). Syntaxin 3 and 4 specifically localize at the apical and basolateral surfaces, respectively. This polarized distribution is thought to provide a final level of specificity during membrane targeting, allowing for specific recognition of either the apical or basolateral membranes. Indeed, several studies have shown that syntaxin 3 and 4 are involved in, respectively, apical and basolateral membrane targeting in MDCK3 cells (5, 6).

Although these studies showed that syntaxins play a crucial role in polarized membrane targeting, the exact requirements for the different syntaxin forms in the specificity of membrane delivery in epithelial cells is still unclear. We previously showed that redirecting syntaxin 3 to the basolateral membrane resulted in mistargeting of apical proteins (7). Also, mutations that cause basolateral missorting of syntaxin 3 interfered with the formation of tight junctions (8). This suggests that at least syntaxin 3 contributes to the specificity of membrane fusion. There are many differences in the regulation of apical and basolateral transport and whether syntaxin 4 plays an identical role in the specificity of basolateral membrane fusion is currently unknown. Furthermore, it is not known how syntaxin 4 itself is specifically targeted to the basolateral membrane. Syntaxin 4 function has been extensively studied in Glut4 transport and insulin secretion. In these studies, its interaction with the SNARE accessory protein munc18c was shown to be important for its function (9–11).

Munc18c is a member of the S/M (sec1/munc18) protein family, a group of proteins that are essential for exocytosis (12–14). They strongly bind to syntaxins and the different S/M proteins show a high binding specificity for different syntaxins. Recently it has been recognized that they are likely to play an active role in membrane fusion (15, 16). The shared function and strong and specific binding of S/M proteins to syntaxins suggest that they are crucial regulators of syntaxin function, but the exact function of this family of proteins is still unclear. S/M proteins may either stimulate or inhibit the function of syntaxins, which may depend on the way they interact with syntaxins.

In this study we have investigated the requirements for syntaxin 4 and munc18c in basolateral vesicle fusion and cell polarization. We show that syntaxin 4 is essential for basolateral protein delivery but not for the initial establishment of membrane polarity. We identified residues that are required for...
basolateral sorting of syntaxin 4 in MDCK cells. Mutation of these residues resulted in non-polarized targeting of syntaxin 4, but expression of these mutants in syntaxin 4-depleted cells did not result in missorting of other basolateral proteins. We also focused on the role of the munc18c in syntaxin 4 sorting and function. Mutations in syntaxin 4 that abolish munc18c binding or munc18c knockdown did not have any effect on the localization of syntaxin 4. However, introducing mutations in syntaxin 4 that affect both the trafficking and munc18c binding results in intracellular localization of syntaxin 4. We, furthermore, show that a syntaxin 4 mutant that cannot bind munc18c rescues basolateral protein targeting in syntaxin 4-depleted cells as efficient as syntaxin 4. Our data suggests that in MDCK cells munc18c is not directly involved in the polarized localization of syntaxin 4 but is necessary for stabilizing syntaxin 4 at the plasma membrane. Finally, our data indicate that the transport and roles of syntaxin 4 at the basolateral domain are distinctly different from those of syntaxin 3 at the apical surface.

**EXPERIMENTAL PROCEDURES**

**Antibodies**—The following antibodies were used. Rabbit anti-syntaxin 4 (Calbiochem, San Diego, CA), rabbit anti-syntaxin 3 (7), rabbit anti-GFP (Rockland, Gilbertville, PA), mouse anti-GFP (Roche Applied Science), rabbit anti-FLAG, mouse anti-FLAG M2, mouse anti-Na,K-ATPase β1 chain, rabbit anti-SNAP23, mouse anti-E-cadherin DECMA (Sigma), mouse anti-E-cadherin, mouse anti-B1-integrin, mouse syntaxin 4 (BD Biosciences), rat anti-B1-integrin clone AII2B (17), rat anti-ZO-1 clone R40.76 (18), rabbit anti-claudin 2 (Invitrogen), mouse anti-p58 clone 6.23.3 (19), mouse anti-gp135 clone 3F21D8 (20), mouse anti-LAMP2 clone AC17 (21), rat anti-occcludin (Zymed Laboratories Inc.), goat anti-Scribble, rabbit anti-β-catenin (Santa Cruz Biotechnology), and rabbit anti-syntaxin 2 and mouse anti-sec6 (StressGen Biotechnologies, Victoria, Canada). The rabbit anti-canine munc18c was raised to a C-terminal peptide of canine munc18c as immunogen (Sigma). Alexa secondary antibodies were obtained from Invitrogen, Li-Cor 800 secondary antibodies were from Li-Cor Biosciences, Lincoln, NE, and HRP-conjugated antibodies from Jackson ImmunoResearch Laboratories, West Grove, PA.

**shRNA, cDNA Constructs, and Mutagenesis**—To generate shRNA constructs, oligos specific for syntaxin 4 and munc18c knockdown were designed using standard procedures. To reduce off-target responses, the sequences were analyzed for possible homology with other sequences searching the dog genome project using the NCBI Blast Server. Oligos designed to knockdown were designed using standard procedures. To determine the efficacy of the shRNA constructs, cells were nucleofected using Amaxa T-buffer and program T-023 and cells were analyzed 2 and 3 days after nucleofection. Stable shRNA-mediated knockdown cell lines were made using the calcium phosphate procedure as previously described (7). Cells were selected with 12.5 μg/ml of blasticidin and screened for knockdown by Western blot analysis. Cell lines were maintained in 10 μg/ml of blasticidin. Prior to experiments, the cells were washed and maintained in normal minimal essential medium with penicillin/streptomycin and 10% FBS. Stable inducible expression cell lines were made as described previously using the T23 MDCK tet-off cell line (7).

**Determination of Integrity of the Monolayer**—The integrity of the epithelial monolayer of cells grown for 5 days on Transwells was determined as described (24). Cells were washed 3 times with PBS with calcium and magnesium (PBS+) PBS-, 0.5 ml of PBS+ with 100 μg/ml of FITC-inulin was added to the apical compartment, whereas 0.5 ml of PBS2- was added to the basolateral compartment. After incubation for 30 min at 37 °C, 200-μl samples were taken from the apical and basolateral compartments. Fluorescence was determined using a fluorescent plate reader. Cells treated with 2 mm EDTA were used as a control for maximal leakage. The results are the mean ± S.D. of three independent experiments.

**Confocal Fluorescence Microscopy**—Immunostaining was done as described previously (7). To stain endogenous syntaxin 4, cells were fixed with methanol at −20 °C for 10 min. Alexa 488 or Alexa 555 were used as secondary antibodies. Alexa 555 phallolidin (Invitrogen) was used to stain filamentous actin. Cells were mounted in Fluorsave supplemented with DAPI. Samples were visualized using a Zeiss LSM 510 with an Axiovert 200M microscope and a C-Apochromat ×63/1.2W Corr lens. Images were analyzed using Imagej for Mac version 1.40g. Composite images with scale bars were made using Adobe Photoshop CS version 9.0 and placed into Illustrator CS2 version 12.0.0.
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For live cell imaging, cells were grown on the bottom of a 6.5-mm Transwell for 5 days as described (25). Cells were infected with adenovirus for VSV-G-EGFP-tsO45 as described (7). We found increased expression levels of VSV-G-EGFP-tsO45 in syn4KD cells, which is likely the result of more exposed adenovirus receptor. We adjusted the amount of virus to obtain similar levels of VSV-G-EGFP-tsO45 (60 infectious units/cell for control cells and 15 infectious units/cell for syn4KD3 cells). Titers were determined using an adenovirus immunoassay kit (Cell Biolabs, San Diego, CA). Infected cells were incubated overnight at 39.5 °C to allow accumulation of VSV-G-EGFP into the ER. Next, the cells were washed in Hanks’ balanced salt solution and incubated for 2 h at 19.5 °C to allow accumulation of VSV-G-EGFP in the Golgi. After this incubation, cells were transferred into minimal essential medium without phenol red containing penicillin/streptomycin, 10% FBS, 20 mM Hapes, pH 7.5, and 10 μg/ml of cytochrome placed on a 35-mm MatTek dish with number 1.5 coverglass (MatTek, Ashland, MA) containing the same medium preincubated at 32 °C in a Perkin-Elmer temperature controlled chamber. The lag time between installing the Transwell and imaging was 2 min. In a typical experiment, Z-stacks with 6 images were obtained with a delay of 30 s for 30–45 min with the Zeiss autofocus macro. Horizontal image drift was corrected using the Stackreg plugin for ImageJ (26). Fluorescence in intracellular areas or areas that overlap the plasma membrane were quantified using ImageJ and represented as a relative increase of fluorescence. This was done for 12 cells for each condition and the average with standard deviations were calculated.

Co-immunoprecipitation Studies—For co-immunoprecipitation studies, cells were grown on 24-mm Transwells for 5 days. Cells were washed 2 times with PBS+, cut out, and lysed in 800 μl of lysis buffer (25 mM Hapes-NaOH, pH 7.4, 150 mM NaCl, 5 mM EDTA, 1% Triton X-100, 1 mM NaF, 1 mM sodium vanadate and protease inhibitors). Lysates were preclarified with Sepharose CL-4B beads (GE Healthcare) and equivalent amounts of protein were taken for the immunoprecipitation. In most cases 1 μg of antibody or 5 μl of antiserum was added to the lysates and incubated at 4 °C for 30 min. Sepharose-Protein A or G was added and incubated for 2 h at 4 °C while rotating. The beads were washed 4 times with lysis buffer by centrifugation. Proteins were eluted from the beads by boiling for 5 min in sample buffer with 100 mM DTT and the samples were, together with the lysates (5% of total), analyzed by SDS-polyacrylamide gel electrophoresis and Western blotting. For quantitative Western analysis, the PVDF (Immobilon, Millipore) membranes were first blocked with Odyssey Blocking buffer, probed with primary antibodies followed by Li-Cor 800 dye-conjugated antibodies (Li-Cor Biosciences) or Alexa 680-conjugated antibodies in Odyssey Blocking Buffer. Membranes were scanned using an Odyssey Infrared scanner. Protein bands were quantified with Odyssey 2.0 software. In some cases the standard enhanced chemiluminescence technique (ECL) was used to detect proteins. In that case 5% milk powder in PBS/Tween was used as blocking buffer and for incubation with primary and secondary antibodies.

Biotinylation Experiments—Biotinylation experiments for E-cadherin of β1-integrin were done as described (7). Briefly, control and syntaxin 4 knockdown cells were grown for 5 days on Transwells and washed in 3× PBS+. Cells were incubated 2×15 min with PBS+ containing 500 μg/ml of EZ-Link Sulfo-NHS-Biotin (Pierce) at the apical or basolateral side of the Transwell. Reactive biotin was quenched by washing the cells 5 times with 50 mM NH4Cl in PBS+. After washing, cells were lysed in RIPA buffer (20 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.1% SDS, 1% deoxycholate, 1% Triton X-100, 5 mM EDTA and protease inhibitors) and immunoprecipitated as described above. Samples were analyzed by SDS-polyacrylamide gel electrophoresis, transferred to PVDF, and the blots were probed with IRDye 800CW-Streptavidin (LI-Cor, Lincoln, NE). Blots were stripped and reprobed for E-cadherin or β1-integrin.

To analyze p58, cells were biotinylated with EZ-Link Sulfo-NHS-SS-Biotin and biotinylated proteins were pulled down using Neutravidin beads (Pierce). Proteins were eluted from the beads by incubating the beads at 60 °C for 10 min in sample buffer with 100 mM DTT. Samples were analyzed by Western blot and probed with an antibody specific to Na,K-ATPase β1 chain.

For determining the polarity of the different syntaxin 4 constructs, we used different cell lines that express syntaxin 4-WT-EGFP and mutants. Using immunofluorescence, we determined conditions that result in similar expression levels at a cellular level. However, as not every cell shows expression, total expression levels are not the same for each cell line. The different cell lines were biotinylated as described above and the amount of pulled down biotinylated syntaxin 4-EGFP was determined by probing the blots with syntaxin 4 antibody. To determine significance, a paired Student’s t test was performed on all quantitative experiments described above.

RESULTS

Knockdown of Syntaxin 4 Affects Basolateral Polarity—To study the function of syntaxin 4 in epithelial cells, we developed MDCK cell lines that were depleted in syntaxin 4. We designed and tested three shRNA constructs directed to three different sequences in the canine syntaxin 4. Transient expression of two of these constructs (syn4KD2 and syn4KD3) reduced syntaxin 4 levels by 80–90% and were used to make cell lines depleted in syntaxin 4 (Fig. 1A). These hairpins did not affect expression levels of other syntaxins but decreased munc18c expression levels (see supplemental Fig. S1A). This is not the result of nonspecific silencing, but rather reflects the close interdependence of syntaxin 4 and munc18c expression (27, 28). For our next studies we used cell lines syn4KD2 c20 and syn4KD3 c9. As a control we made cell lines transfected with hairpins directed to GFP (GFPKD) or Luciferase (LucKD) (29). Both cell lines were used as controls in all the described experiments.

The syntaxin 4 knockdown (syn4KD) cell lines grew normal on plastic tissue culture plates and did not have an aberrant phenotype. Cell death and proliferation rates were also similar for syn4KD and control cells (not shown). However, when grown on permeable membranes, the cells showed an aberrant morphology with extended bulging apical domains, which
FIGURE 1. Knockdown of syntaxin 4 results in inhibition of basolateral trafficking. A, Western blot analysis of syntaxin 4 expression levels in cells expressing two different hairpins to syntaxin 4 (KD2 and KD3). Two clones for each hairpin are shown. A cell line expressing a hairpin to GFP (GFPKD) was used as a control cell line. B–F, MDCK cell line depleted in syntaxin 4 (syn4KD3) and a control cell line (GFPKD) were grown for 5 days on filter, fixed, and stained. B, gp135/podocalyxin (magenta) and β-catenin (green) (z-scan). C, cells stained for several different basolateral proteins (syntaxin 4, Na,K-ATPase β1, E-cadherin, β1-integrin, and Scribble). D, cells stained for tight junction proteins (occludin, ZO-1, claudin 2, and sec6). Staining for actin is also shown. Images represent projections of several x-y scans. Representative single x-y scans are shown under supplemental Fig. S1B. E, cells stained for laminin (top, projections of x-y scans) and laminin (green) and actin (magenta) (bottom, z-scans). F, cells stained for SNAP23. Scale bars, 20 μm.

stained positive for the apical protein gp135/podocalyxin (Fig. 1B).

The basolateral staining for β-catenin was reduced in syn4KD cells (Fig. 1B), indicating that basolateral protein delivery is affected. To investigate this further, we stained for several other basolateral proteins (Fig. 1C, images show projections of Z-scans supplemental Fig. 1B shows single x-y scans of these images). As expected, syntaxin 4 expression was strongly reduced as the syn4KD cells are characterized by the absence of the typical honeycombed staining of syntaxin 4 (Fig. 1C, residual fluorescence is most likely background staining). Antibodies to the basolateral proteins E-cadherin, β1-integrin, and in particular p58 (which recognizes β1-chain of the Na,K-ATPase (30)) showed substantial increase of intracellular staining (Fig. 1C). Most of this staining was diffuse but we also observed localization in intracellular compartments for all antibodies. Similar results were obtained for knockdown cells lines made with different hairpins (supplemental Fig. 1C). Interestingly, the basolateral localization of E-cadherin was less sensitive to syntaxin 4 ablation as compared with other basolateral proteins. However, when cells were grown for longer periods (7–9 days on filter), a more significant reduction of basolateral E-cadherin staining was observed (supplemental Fig. S1D). We also analyzed the localization of Scribble, a polarity protein that directs the formation of the basolateral domain (31) and is localized basolateral (32). Scribble localized mostly basolateral in syn4KD cells, although some diffuse intracellular Scribble was observed (see also supplemental Fig. S1B).

We next examined the localization of several tight junction proteins and the exocyst protein sec6 (Fig. 1D). Interestingly, syntaxin 4 depletion affected some, but not all tight junction proteins. Staining for ZO-1 and the exocytotic protein sec6 were not affected by syntaxin 4 knockdown. However, occludin and claudin 2 showed aberrant localization in the syntaxin 4KD cells, as some tight junction proteins are not correctly localized we would expect significant effects on the integrity of the epithelial layer. However, determining the epithelial integrity using leakage of FITC-inulin from the apical compartment showed no significant difference between control and syn4KD3 cells (supplemental Fig. 1E). This suggests that syn4KD cells partly maintain their tight junctions and form a monolayer impermeable for small molecules.

We next tested if secretion of basolateral proteins was affected in syn4KD cells. For this, we stained cells with an antibody for the secreted protein laminin-111 (which recognizes laminin β1 and γ1 chains) and found less deposited laminin and increased intracellular staining (Fig. 1E). This indicates that syntaxin 4 is also required for basolateral secretion. Finally, we found that localization of endogenous SNAP23, another plasma membrane SNARE protein that interacts with syntaxin 4, was not changed when syntaxin 4 was depleted (Fig. 1F).

Together, the data show that depletion of syntaxin 4 resulted in inhibition of basolateral and tight junction protein delivery. Apical targeting was not affected but the apical domain was greatly expanded.

Syntaxin 4 Knockdown Differentially Affects Basolateral Protein Delivery—Because our data indicated that basolateral localization of E-cadherin and β1-integrin was differentially affected by syntaxin 4 knockdown, we analyzed membrane transport and correct polarized delivery in more detail for several different basolateral proteins. For this, we quantified the localization of E-cadherin, β1-integrin, and Na,K-ATPase β1 using surface biotinylation. Surface biotinylation of E-cadherin showed that a small but significant portion of this protein was missorted to the apical surface (Fig. 2A, top blot, bottom, bar graph; open bars, apical E-cadherin). However, the larger effect was seen in total delivery to the plasma membrane, as we found that the total surface biotinylation was reduced in syn4KD cells (Fig. 2B, open bars: surface-biotinylated protein as fraction of total immunoprecipitated protein, 73 ± 10% for syn4KD3).
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When we repeated these experiments for $\beta_1$-integrin, we also found a small but significant increase in apical localization (not shown). Total surface biotinylation of $\beta_1$-integrin was substantially reduced (Fig. 2C, open bars: surface biotinylated protein as fraction of total immunoprecipitated protein, 68 ± 17%). Interestingly, we also found differences between the two proteins, as $\beta_1$-integrin but not E-cadherin expression was significantly reduced in syn4KD3 cells compared with control cells (Fig. 2, B, closed bars, E-cadherin (98 ± 14%) versus C, closed bars, $\beta_1$-integrin (66 ± 10%). Expression levels are indicated as a percentage of control). Finally, we analyzed basalateral localization of Na,K-ATPase $\beta_1$ by surface biotinyation. In the syn4KD cells surface biotinylation was strongly reduced (Fig. 2D, open bar, 53 ± 13%) with little effect on protein expression level (Fig. 2D, closed bar, 90 ± 7%), suggesting that a considerable amount of the protein was localized intracellular.

Thus, when comparing the results of the different basolateral proteins we found that syntaxin 4 knockdown affects basolateral targeting of protein differently. Basolateral localization of E-cadherin was affected to a lesser extent compared with $\beta_1$-integrin and Na,K-ATPase $\beta_1$, in which syntaxin 4 knockdown results in intracellular protein accumulation with little apical mistargeting. These results are in agreement with the immunofluorescence data shown in Fig. 1C. Together, our data suggests that the loss of function of syntaxin 4 mainly results in failure to deliver proteins to the plasma membrane, rather than cause their missorting to the apical surface.

**Accumulation of Intracellular Vesicles in syn4KD Cells**—Our findings that syntaxin 4 knockdown inhibited the steady state localization of basolateral proteins at a different extent may be explained by differential stability of these proteins at the basolateral surface, rather than by differences in basolateral targeting. We therefore analyzed how syntaxin 4 knockdown affects trafficking of newly synthesized protein to the basolateral membrane. For this, we used live cell imaging of the temperature-sensitive mutant of the VSV-G protein (tsO45) tagged to EGFP. This mutation causes accumulation of the protein in the ER, whereas a subsequent incubation at 19.5 °C further accumulates the protein in the trans-Golgi network. We followed the synchronous transport of VSV-G-EGFP to the plasma membrane after removing the temperature block (33). In control cells an increase of lateral fluorescence was observed due to arrival of VSV-G-EGFP at the lateral membrane (Fig. 3A). This was accompanied by a decrease in intracellular fluorescence as
VSV-G-EGFP is transported out of the trans-Golgi network. The delivery of VSV-G-EGFP to the plasma membrane was clearly inhibited in the syn4KD cells (Fig. 3B). Semiquantitative analysis of fluorescence of 12 individual control and syn4KD cells showed that maximum fluorescence at the plasma membrane was reached after 20 min in control cells (Fig. 3C, open circles). In the syn4KD cells, maximum fluorescence levels are reached after 40 min (open squares) and this fraction of fluorescence is ~30% compared with the control cells. This number is likely an overestimation as our system does not distinguish between actual plasma membrane localization and localization of transport vesicles close to the plasma membrane. Total intracellular fluorescence did not decrease in the syn4KD cells to the same level as the control cells (Fig. 3C, closed symbols, and D). These data suggest that depletion of syntaxin 4 does not inhibit the exit of newly synthesized VSV-G-EGFP from the Golgi area, but does cause an accumulation of intracellular transport vesicles containing VSV-G-EGFP as a result of inhibition of membrane fusion.

**Munc18c Knockdown Mimics the Syntaxin 4 Phenotype**—Several studies have shown that expression levels of syntaxin 4 and munc18c are tightly linked (27, 28). To determine how expression of munc18c affects syntaxin 4 expression and localization in MDCK cells, cell lines depleted in munc18c were
made. Two of the 6 shRNA hairpins we tested gave a knockdown of 80% compared with a control vector upon transient transfection. We made cell lines using munc18cKD2 in which knockdown was about 80–90% (Fig. 4A) and found that munc18c knockdown reduced expression of syntaxin 4 (Fig. 4A). We were not able to stain for endogenous munc18c, using several different antibodies and fixation protocols. However, as Western blotting showed such significant reduction, it is likely that most of the cells were depleted in munc18c. Similar to what we observed in syn4KD cells, munc18cKD cells showed extended apical domains (Fig. 4B, munc18cKD2 c20, similar results were obtained with other clones). The localization of E-cadherin was not grossly affected (Fig. 4B), but both Na,K-ATPase β1 (Fig. 4C) and β1-integrin (not shown) showed increased intracellular staining and reduced staining at the lateral membrane. Using biotinylation studies we found that munc18c depletion resulted in reduced basolateral surface labeling of β1-integrin (Fig. 4D). Thus, the phenotype of these cell lines resembled syntaxin 4 knockdown cells. However, the phenotype of munc18c knockdown was less severe than observed for the syn4KD cells. When we stained for endogenous syntaxin 4 in munc18cKD cells we found, although difficult to detect, some residual basolateral staining (Fig. 4E).

FIGURE 4. Munc18c knockdown has a similar phenotype as syntaxin 4 depletion. A, expression levels were determined by Western blotting of canine munc18c and syntaxin 4 in control and different clones of munc18c knockdown cells (munc18cKD2). GAPDH was used as a loading control. B, immunofluorescence of munc18c KD2 c20 and control cells (LucKD) grown for 5 days on filter. Cells were stained for gp135/podocalyxin (top, with x-z scans of gp135) and E-cadherin (bottom, with x-z scans of E-cadherin). Scale bars are 20 μm. C, immunofluorescence of munc18c KD2 c20 and control cells (LucKD) grown for 5 days on filter. Cells were stained for Na,K-ATPase β1. Scale bar, 20 μm. D, the relative biotinylation of total immunoprecipitated β1-integrin was quantified compared for control cells (LucKD) and munc18cKD c20 cells and the results are normalized to control cells (open bars). Relative expression levels of β1-integrin in lysates of control and munc18c KD c20 cells are shown as closed bars. The results are shown as a mean ± S.D. of three independent experiments (*, p < 0.001; **, p < 0.001). E, control (LucKD), syn4KD3 c9, and munc18c KD c20 stained for syntaxin 4. Scale bar, 10 μm.
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was not affected by FLAG-munc18c expression. Expression levels of syntaxin 4 levels remained low when FLAG-munc18c expression was induced (Fig. 5A, FLAG-munc18c syn4KD3). These results show that syntaxin 4 expression is essential for basolateral delivery of proteins and that the reduced levels of munc18c do not contribute to the observed phenotype of the syn4KD cells.

N-terminal Region of Syntaxin 4 Contains Basolateral Sorting Signal—We recently showed that exchanging the first N-terminal 40 amino acids of syntaxin 3 with those of syntaxin 4 results in complete basolateral localization of syntaxin 3 (7). As this suggests that the N-terminal sequence of syntaxin 4 contains basolateral sorting signals, we further characterized these signals by making several deletion mutants of rat syntaxin 4. To visualize the mutants we ecto-tagged syntaxin 4 with EGFP at the C terminus (supplemental Fig. S2A). Ecto-tagging SNAP proteins at the C terminus does not effect the localization and function of these proteins (34, 35). Indeed, wild type (WT) EGFP-tagged syntaxin 4 showed a similar lateral localization as untagged syntaxin 4. We and others have previously shown the importance of the first 8 amino acids of the N-terminal sequence for munc18c binding (7, 36, 37). However, transient expression of syntaxin 4 with N-terminal deletions up to 15 amino acids had no major effects on the localization of syntaxin 4 (Fig. 6A), although removal of the first 10 amino acids completely blocked binding to munc18c (see below, Fig. 7A). This suggests that basolateral targeting of syntaxin 4 is independent of munc18c binding.

Additional deletion of amino acids caused intracellular localization of syntaxin 4 (Δ20, Δ30, and Δ40), although a significant amount was still directed to the lateral membrane. Less cells express syntaxin 4-Δ20, -Δ30, and -Δ40 expression, which is likely caused by degradation. Indeed, intracellular syntaxin 4-Δ40 co-localized with the endosomal/lysosomal protein Lamp-2 (Fig. 6B) (21).

Together our results indicate that a sorting signal is present in the region just after the first 15 amino acids. We determined that amino acids Leu-25 and Val-26 are conserved among different species (supplemental Fig. 2C). Introduction of a V26A, and to a lesser extent L25A mutation, resulted in significant apical localization, but no significant intracellular staining (Fig. 6C). Other mutations of the acidic amino acids located in the 15–20-amino acid region of syntaxin 4 had little effect on its localization (Fig. 6C and supplemental Fig. S2B). This is unexpected as deletion of this region resulted in intracellular localization (Fig. 6A).

To exclude any effect of the EGFP tag, we also analyzed cells stably expressing C-terminal myc-tagged syntaxin 4 WT and
mutants and found a similar apical localization (supplemental Fig. S3A) for syntaxin 4-L25A,V26A. Mutations in EGFP that prevent dimerization (A206K) (38) or oligomerization (C49S) (39) did not change the localization of syntaxin 4-WT or syntaxin 4-L25A,V26A (supplemental Fig. S3B). Thus, Leu-25 and Val-26 are important for correct localization of syntaxin 4. We confirmed this finding in several other epithelial cells, including Fisher rat thyroid cells and LLC-PK1 cells (supplemental Fig. S3C).

Interestingly, the deletion mutants of syntaxin 4 localized partly intracellular, whereas the L25A,V26A mutant was missorted to the apical surface, but did not accumulate intracellularly. This suggests that an additional sorting determinant is present in the N-terminal region. Because the L25A,V26A mutant, but not deletion mutants binds munc18c, we investigated if munc18c is involved in the localization of syntaxin 4.

For this, we introduced the L8K mutation, which decreases munc18c binding (10). This mutant showed a normal lateral localization (Fig. 6C). However, mutation of both the sorting sequence and the munc18c binding peptide (L8K, L25A, and V26A) showed a non-polarized localization and high expressing cells showed intracellular staining (Fig. 6C and supplemental Fig. S3D). The localization of this mutant resembled deletion mutants Δ20-Δ40 and the syn3/4 chimera, suggesting that N-terminal peptide binding of munc18c contributes to correct localization of syntaxin 4.

To analyze the apical and basolateral steady state localization by cell surface biotinylation, we generated syntaxin 4 mutants as described above with an extracellular EGFP tag. We expressed these under control of the tet-inducible system (40), which allowed us to obtain similar expression levels of the different constructs (Fig. 6D). Expression of EGFP-tagged syn-
taxin 4-WT, syntaxin 4-L8K, syntaxin 4-L25A,V26A, syntaxin 4-L8K,L25A,V26A, and syntaxin 4-Δ30 did not affect polarized cell morphology and the EGFP-tagged proteins localized similarly to the transiently expressed constructs (not shown). Consistent with our immunofluorescence data, there was a significant increase of apical syntaxin 4-L25A,V26A, syntaxin 4-L8K,L25A,V26A, and syntaxin 4-Δ30 as compared with syntaxin 4-WT, whereas no significant changes were observed for syntaxin 4-L8K (Fig. 6D).

Together, our results show that residues Leu-25 and Val-26 within the N-terminal region are required for correct basolateral localization and act by a munc18c-independent mechanism. Our data, furthermore, indicates that munc18c plays a separate role in the polarized localization of syntaxin 4.

L25A,V26A Mutation in Syntaxin 4 Does Not Affect Munc18c Binding—Recent published crystal structure data showed that only the first 12 amino acids of the N-terminal peptide of syntaxin 4 bind to munc18c (36). To confirm this, and to exclude that the L25A,V26A mutation inhibited binding, we transiently expressed different syntaxin 4-EGFP constructs or the pEGFP-N1 control vector in MDCK cells that express FLAG-tagged munc18c in a tet-regulatory manner (7). FLAG-munc18c co-immunoprecipitated with syntaxin 4-WT but not with syntaxin 4-L8K, syntaxin 4-Δ10, or the control EGFP (Fig. 7A), thus confirming previous reported results (10). Even with longer exposure times we did not observe any co-immunoprecipitation between syntaxin 4-L8K or syntaxin 4-Δ10 with munc18c. Despite the fact that syntaxin 4-L8K and FLAG-munc18c do not interact, co-expression of FLAG-munc18c increased expression of both syntaxin 4 WT and 4-L8K (Fig. 7B).

Both syntaxin 4-WT and syntaxin 4-L25A,V26A, but not syntaxin 4-L8K or syntaxin 4-L8K,L25A,V26A also interacted with endogenous canine munc18c (munc18c, Fig. 7, C and D). This was confirmed by reciprocal co-immunoprecipitations using transiently transfected syntaxin mutants (supplemental Fig. S4). These residues (Leu-25 and Val-26) are not required for munc18c binding but replacing Leu-8 with lysine or deletion of 10 amino acids of the N-terminal region resulted in complete loss of munc18c binding.

Munc18c Stabilizes Syntaxin 4 at Plasma Membrane—To examine if munc18c knockdown affected syntaxin 4 localization we nucloefected control or munc18c knockdown cells (munc18cKD2 c20) with syntaxin 4-WT and syntaxin 4-L25A,V26A. Consistent with the basolateral localization of the syntaxin 4-L8K mutant, munc18c knockdown did not affect syntaxin 4-WT localization (Fig. 8A). However, depletion of munc18c caused syntaxin 4-L25A,V26A to localize exclusively lateral as compared with the non-polarized localization in the control cells (Fig. 8B). A possible explanation is that munc18c binding to syntaxin 4-L25A,V26A may mask a potential apical endocytosis signal and depletion of munc18c allows apical syntaxin 4-L25A,V26A to be endocytosed. It is expected that syntaxin 4-L25A,V26A expressed in munc18cKD cells localize also intracellularly as observed for the syntaxin 4-L8K,L25A,V26A mutant (Fig. 6C). This is, however, not the case and this is likely the result of insufficient expression levels to detect intracellular syntaxin 4-L25A,V26A.

We next examined if munc18c contributes to the apical localization of syntaxin 4-L25A,V26A. For this we used the cell line that expresses FLAG-munc18c under a tet-inducible promoter. Overexpression of munc18c results in increased levels of ectopically expressed syntaxin 4 (see Fig. 7B) (7). As expected, transiently expressed syntaxin 4-L25A,V26A in cells that do not express FLAG-munc18c (+dox) localized partially apically (Fig. 8C, +dox). Induction of FLAG-munc18c expression (~dox) increased the expression levels, but did not alter the localization of syntaxin 4-L25A,V26A (Fig. 8C, ~dox). In contrast, expression of syntaxin 4-L8K,L25A,V26A or syntaxin 4-Δ30 in FLAG-munc18c expressing cells resulted in significant apical localization and an increased expression of this mutant as compared with cells that do not express FLAG-munc18c (Fig. 8, D and E, compare +dox to ~dox). This was unexpected because we were not able to find any interaction between FLAG-munc18c and syntaxin 4-L8K,L25A,V26A or syntaxin 4-Δ30 by immunoprecipitation studies (Fig. 7 and supplemental Fig. S4 and data not shown). Without FLAG-munc18c expression (+dox), syntaxin 4-Δ30 and Δ40 localized partly intracellularly. In FLAG-munc18c expressing cells (~dox) no intracellular staining was observed (supplemental Fig. S5, C and D, +dox). Nevertheless, these data show that munc18c overexpression increases syntaxin 4-L25A,V26A localization at the apical membrane.

Binding of munc18c Is Not Essential for Rescue of Basolateral Transport in Syntaxin 4 Knockdown Cells—We show that specific mutations of syntaxin 4 in the N-terminal region resulted in mislocalization of syntaxin 4. We next examined if these syntaxin 4 mutants were still able to function in protein deliv-
residues are present in the N terminal that are essential for the function of syntaxin 4 as syntaxin 4-Δ30 and Δ40 were not able to rescue Na,K-ATPase transport. In the case of apical localized syntaxin 4-L25A,V26A, we did not observe any mislocalization of Na,K-ATPase β1 to the apical membrane (supplemental Fig. S6C). These results show that munc18c binding to syntaxin 4 per se is not required for the function of syntaxin 4 in basolateral trafficking in MDCK cells yet the N terminal contains residues that are important for its function.

**DISCUSSION**

We show that depletion of syntaxin 4 results in intracellular accumulation of basolateral and tight junction proteins. All basolateral proteins we tested so far show decreased basolateral targeting in the syn4KD cells, although E-cadherin was less sensitive to the loss of syntaxin 4 compared with the other basolateral proteins. This would be in agreement with the function of E-cadherin as an initiator of basolateral polarity and acting upstream in the biogenesis of basolateral polarity (41). Also, E-cadherin at the plasma membrane will likely be stabilized by trans-interactions. Intracellular E-cadherin accumulation increased in cells grown for longer periods on filter, indicating that recycling of E-cadherin after establishment of polarity is affected by depletion of syntaxin 4. Delivery of claudin 2 and occludin is also inhibited by syntaxin 4 depletion, indicating that syntaxin 4 is also involved in the delivery of these tight junction proteins. However, we find no effect on the permeability for small molecules in syn4KD cells compared with control cells, suggesting that not all tight junction components are affected by syntaxin 4 knockdown. This is also apparent from the unchanged localization of ZO-1. Despite reduced basolateral delivery, cells maintain their polarity and only a small fraction of the tested proteins were found to localize at the apical domain.

We examined the basolateral targeting of syntaxin 4 and identified residue Leu-25 and to a lesser extent Val-26 as essential for basolateral sorting of syntaxin 4. These residues resemble a dileucine motif, preceded by a region of acidic residues. Dileucine motifs act as endocytosis signals, but can also act as basolateral target signals (2, 42). The motif in rat and mouse syntaxin 4 differs from the consensus dileucine motif, as the first acidic amino acid is at the −5 and not −4 position relative to the first leucine (see supplemental Fig. S2C). Mutations of these acidic residues in syntaxin 4 do not affect basolateral sorting. However, a truncation including the acidic residues but not the leucine/valine motif caused missorting (Fig. 7A, syn4-Δ20). This is similar to what was previously reported for the sorting of CD147, in which removal of acidic amino acids resulted in apical missorting but substitution of these residues with alanine did affect trafficking (43). Dileucine motifs can bind the μ, β, or γ/σ1 subunits of AP1, the adaptor involved in the trans-Golgi network to plasma membrane transport (42, 44–46). MDCK cells express the epithelial specific adaptor subunit μ1B and it was shown that this subunit is essential for the correct sorting of some basolateral proteins (47, 48). However, in LLC-PK1 cells, a cell line that does not express μ1B (48), syntaxin 4 localizes mostly basolateral. Also, the L25A,V26A mutant shows increased apical localization in this cell line.
suggesting that μ1B is not directly involved in sorting of syntaxin 4 to the basolateral membrane.

It is still unclear if munc18 proteins contribute to syntaxin localization. One study showed that syntaxin 1A was targeted correctly in munc18-1-deficient mice (49). However, other studies found that knockdown of munc18-1/-2 in PC12 cells resulted in decreased expression and intracellular mislocalization of syntaxin 1A (50, 51).

In our system, knockdown of munc18c does not interfere with targeting of syntaxin 4 to the plasma membrane. Syntaxin 4 localization is also not affected by the introduction of deletions Δ10 and Δ15 or a mutation (L8K) that affects binding to munc18c. On the other hand, although the L25A,V26A mutation is missorted, it does not affect munc18c binding. From these results we conclude that munc18c is not required for syntaxin 4 targeting to the plasma membrane. However, removal of the first 20 amino acids of the N terminus of syntaxin 4 or introducing mutations that affect both munc18c binding and plasma membrane sorting (L8K,L25A,V26A) results in intracellular localization of syntaxin 4. This suggests that munc18c stabilizes syntaxin 4 at the plasma membrane.

Indeed, depletion of munc18c results in reduced expression of syntaxin 4 or vice versa. Also the L8K mutation in syntaxin 4 affects its stability, as the expression of this mutant was lower after transient nucleofection compared with syntaxin 4-WT cells. In addition, overexpression of munc18c results in a decrease of endogenous munc18c. This suggests that munc18c levels are tightly controlled. Such a close interdependence of munc18c- and syntaxin 4-expression has been reported before (27, 28) and was also shown for other plasma membrane SNAREs (49, 52).

Membrane stabilization of syntaxin 4 by munc18c may also explain our localization studies of the syntaxin 4-L25A,V26A mutant. Syntaxin 4-L25A,V26A expression in munc18c-depleted cells results in reduced apical localization. Similarly, the munc18c binding mutant L8K,L25A,V26A is not localized at the apical membrane. However, munc18c overexpression results in stabilization of the L8K,L25A,V26A mutant and Δ30 and Δ40 deletion mutants at the apical membrane. This is an unexpected result because co-immunoprecipitation studies showed no interaction between munc18c and any of the L8K and deletion mutants, in both control and munc18c overexpressing systems (Fig. 8). Several explanations for this discrepancy are possible. The interaction of the L8K mutant with munc18c may be weak and lost during co-immunoprecipitation. Indeed, residual interaction between syntaxin 4 mutated in the N-peptide has been reported, as Latham and co-workers (10) found some interaction between syntaxin 4-L8K pulled down with munc18c-GST munc18c. Using FRET analysis it has been shown that another syntaxin 4 N-peptide mutant deficient in munc18c binding interacts in vivo with munc18c (37). Also, we have previously shown that syntaxin 4 without the N-terminal peptide binds munc18c (7) in vitro. Another possible explanation is that syntaxin 4 N-peptide mutants are part of a larger complex of syntaxins. It was recently shown that both syntaxin 1A and syntaxin 4 form large clusters of syntaxins, via interactions of their core regions (54, 55). The syntaxin 4-L8K could be localized together with endogenous syntaxins in clusters and these clusters may be stabilized by overexpression of munc18c that is bound to endogenous syntaxin.

Absence of munc18c interaction does not affect the function of syntaxin 4 in our system. Expression of syntaxin 4-L8K in syntaxin 4-depleted cells rescues Na,K-ATPase β1 localization to a similar extent as syntaxin 4-WT. Also, the effect of munc18c knockdown on delivery of basolateral proteins is not as pronounced as for syntaxin 4 and may rather be the effect of reduced syntaxin 4 levels.

Interestingly, syntaxin 4 mutant with a deletion of the first 30 residues did not rescue Na,K-ATPase β1 localization in the syntaxin 4 knockdown cells. This suggests that other residues in this region may be important for the function of syntaxin 4.

Taken together our data suggest that in epithelial cells, munc18c binding to the N-terminal peptide may not be essential for docking and fusion. Nevertheless, these results show the importance of stabilization of syntaxin 4 at the plasma membrane.

Depletion of syntaxin 4 causes intracellular accumulation of basolateral proteins but no substantial missorting to the apical membrane. Expression of the apical sorted syntaxin 4 mutant L25A,V26A in syntaxin 4-depleted cells does not result in mislocalization of Na,K-ATPase β1. This suggests that apical localization of syntaxin 4 is not sufficient to redirect trafficking of basolateral proteins. It is likely that specific combinations of SNAREs, tethering proteins, and other proteins (53, 56) will provide specificity in membrane fusion. How the different proteins of the tethering/fusion machinery interact and how this is regulated is still an open question. Further studies will be required to identify such combinations of protein complexes for the different trafficking pathways.

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