Murine CENPF interacts with syntaxin 4 in the regulation of vesicular transport

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Summary
Syntaxin 4 is a component of the SNARE complex that regulates membrane docking and fusion. Using a yeast two-hybrid screen, we identify a novel interaction between syntaxin 4 and cytoplasmic murine CENPF, a protein previously demonstrated to associate with the microtubule network and SNAP-25. The binding domain for syntaxin 4 in CENPF was defined by yeast two-hybrid assay and co-immunoprecipitation. Confocal analyses in cell culture reveal a high degree of colocalization between endogenously expressed proteins in interphase cells. Additionally, the endogenous SNARE proteins can be isolated as a complex with CENPF in immunoprecipitation experiments.

Introduction
Movement of membrane bound vesicles to and from the cell membrane is dependent on the cytoskeleton. Along with the actin network, microtubules serve as tracks for the efficient and directed movement of organelles and vesicles (Caviston and Holzbaur, 2006; Hennly and Stammes, 2007; Ishiki and Klip, 2005; Soldati and Schliwa, 2006; Vedrenne and Hauri, 2006). A wide range of studies report the importance of the microtubule network in the transport of specific cargos in polarized and non-polarized cells. For example, microtubule-based transport and dynein activity are essential for TrkA signaling to Rap1 and MAPK1/2 as inhibition of dynein activity results in altered vesicular trafficking (Miyamoto et al., 2006; Wu et al., 2007). Similarly, kinesin motors regulate vesicular cargo along microtubules (Schnapp, 2003). Furthermore, a PKA- and calcium-dependent pathway controls non-polarized macrophage secretion of apoE along the microtubule network (Kockx et al., 2007). Additionally, treatment with agents that disrupt microtubule function, such as nocodazole and rotenone, inhibit movement of proteins from the Golgi to the cell surface (Feng, 2006; Zheng et al., 2007). However, identification and characterization of proteins directly linking membrane bound vesicles to the microtubule network remain elusive.

Human centromere protein F (CENPF) (also called mitosin) was independently identified by the Yen and Lee groups utilizing auto-immune antibodies and Rb-binding properties (Rattner et al., 1993; Zhu et al., 1995b). Our group discovered the murine homolog and, although previous publications have used the name LEK1 (Ashe et al., 2004; Goodwin et al., 1999; Pooley et al., 2006; Soukoulis et al., 2005), we now refer to this protein as murine centromere protein F (hereafter referred to as murine CENPF) to more accurately describe the gene product. Murine CENPF, like other family members, is a large protein (2998 aa) and shares significant sequence and domain homology with related proteins in both humans and avians (Ashe et al., 2004; Dees et al., 2000; Goodwin et al., 1999; Liao et al., 1995; Pabon-Pena et al., 2000; Zhu et al., 1995a; Zhu et al., 1995b). The human homolog binds the kinetochore and is an important regulator of mitosis and cell division (Feng et al., 2006; Liang et al., 2004; Liao et al., 1995; Rattner et al., 1993; Yang et al., 2005; Zhu et al., 1995a; Zhu et al., 1997; Zhu et al., 1995b). In addition, both human and murine CENPF bind proteins associated with the microtubule network, including tubulin (Feng et al., 2006) and Nde1 (Soukoulis et al., 2005). The binding of murine CENPF to Nde1 is of particular interest as Nde1 interacts with Lis1 and dynein to modulate the microtubule network in regulation of cell shape and movement (Faulkner et al., 2000; Gibbons, 1996; Rattner et al., 1993; Smith et al., 2000; Zhu et al., 1997). Relevant to the current study, the Lis1 pathway functions with the Golgi network and in membrane trafficking (Kondratova et al., 2005; Liang et al., 2004). Utilizing dominant-negative protein expression and induced suppression of murine CENPF expression, we demonstrate that interference with CENPF function severely alters the microtubule network (Soukoulis et al., 2005). In addition, several studies show that membrane trafficking and positioning of organelles are dependent on interaction of the microtubule network with Nde1 and Lis1 (Banks and Heald, 2001; Faulkner et al., 2000; Gibbons, 1996; Smith et al., 2000; Terada et al., 1996; Xiang et al., 1999).

Further analyses demonstrate that murine CENPF and syntaxin 4 colocalize with components of plasma membrane recycling: SNAP-25 and VAMP2. Depletion of endogenous CENPF disrupts GLUT4 trafficking whereas expression of a dominant-negative form of CENPF inhibits cell coupling. Taken together, these studies demonstrate that CENPF provides a direct link between proteins of the SNARE system and the microtubule network and indicate a diverse role for murine CENPF in vesicular transport.

Key words: CENPF, Syntaxin 4, GLUT4, Rab11a, VAMP2, SNAP-25
such fusion events (Chen and Scheller, 2001; Ishiki and Klip, 2005). Apposing SNARE proteins, which consist of vesicle-associated membrane proteins (VAMPs), plasma membrane-associated syntaxins, and cytoplasmic synaptosomal-associated proteins (SNAPs), form coiled-coil aggregates that are important in regulating membrane fusion events (Aikawa et al., 2006; Chen and Scheller, 2001; Ishiki et al., 2005; Martin et al., 1998; McMahon et al., 1993; Rowe et al., 1999). Plasma membrane trafficking of proteins between subcellular domains and translocation to the cell surface is mediated, in part, by SNARE proteins (Chen and Scheller, 2001; Jahn and Sudhof, 1999; Mallard et al., 2002; Sollner et al., 1993; Wicke et al., 2000). The SNARE protein syntaxin 4 is an integral membrane protein that localizes to the plasma membrane and is essential in vesicular docking and fusion (Aikawa et al., 2006; Bajohrs et al., 2005; Band et al., 2002; Pooley et al., 2006). Specifically, syntaxins are critical in vesicular transport of GLUT4-containing vesicles in skeletal muscle, cardiomyocytes and adipose tissue after insulin stimulation (Bryant et al., 2002; Cain et al., 1992; Martin et al., 1996; Pessin et al., 1999). Determining the physical linkage of SNAREs to the microtubule network is essential for understanding the role of this cytoskeletal component in the myriad of events that control vesicular transport.

In a recent study, we reported that murine CENPF physically associates with SNAP-25 (synaptosomal-associated protein 25), and together these proteins form a complex with Rab11a, myosin Vb, and VAMP2 in the recycling endosome pathway (Pooley et al., 2006). Furthermore, disruption of endogenous murine CENPF function by dominant-negative expression or protein knockdown severely retarded the recycling endosome network and transferrin trafficking (Chen and Scheller, 2001). Although this is the only report of CENPF regulating vesicular transport, the family has been shown to function with the cytoskeleton (Goodwin et al., 1999; Feng et al., 2006), and from this, we postulate that CENPF may have an extensive role in controlling SNARE-mediated vesicular transport by the microtubule network. Further data is crucial to establish the roles for CENPF in the diverse processes of vesicular transport.

In the current study, using yeast two-hybrid (Y2H) and biochemical analyses, we demonstrate that syntaxin 4 and murine CENPF physically interact. These data are consistent with the hypothesis that murine CENPF is a critical component in the dynamic regulation of plasma membrane trafficking with the microtubule network through its interaction with SNARE proteins and Nde1 (Soukoulis et al., 2005). Using genetic, immunolocalization, and immunoprecipitation studies, we demonstrate that both transiently expressed and endogenous cytoplasmic CENPF directly associate with syntaxin 4 at the Golgi complex and this complex also contains VAMP2 and SNAP-25. Additionally, disruption of CENPF interferes with cell coupling in NIH3T3 fibroblasts, demonstrating inhibition of gap junction function at the cell membrane (Francis and Lo, 2006) and indicating the essential role of CENPF in membrane trafficking. Finally, we show that disruption of CENPF function inhibits GLUT4 trafficking, a system used to model syntaxin 4 function in membrane trafficking in 3T3 adipocytes. Thus, the present study establishes a physical link between murine CENPF and the SNARE complex and suggests a role for CENPF in the regulation for vesicular transport by the microtubule network.

Results
Identification of syntaxin 4 as a murine CENPF binding partner
Despite the significant amount of literature characterizing CENPF family members, most work has focused on the C terminus of the protein (Clark et al., 1997; Feng et al., 2006; Konstantinidou et al., 2003; Liao et al., 1995; Zhou et al., 2005; Zhu et al., 1995a; Zhu et al., 1997; Zhu et al., 1995b). Relatively little was known concerning potential molecular interactions and functions in the major N-terminal regions of the molecule until recently. Feng et al. identified a microtubule binding domain in the initial 385 amino acids of human CENPF (Feng et al., 2006), and our group demonstrated the interaction of human CENPF with Nde1 and SNAP-25 (Pooley et al., 2006; Soukoulis et al., 2005). To identify binding partners in this domain, and thus, ascribe molecular function, we conducted an extensive Y2H screen using the N-terminal coiled-coil domain (amino acids 1-689; coding sequences for exons 1-6) of murine CENP-F (Feng et al., 2006), and our group identified the interaction of murine CENPF with Nde1 and SNAP-25 (Pooley et al., 2006; Soukoulis et al., 2005). To identify binding partners in this domain, and thus, ascribe molecular function, we conducted an extensive Y2H screen using the N-terminal coiled-coil domain (amino acids 1-689; coding sequences for exons 1-6) of murine CENPF as bait. From this screen, we identified specific proteins known to regulate organellar positioning and membrane trafficking, one of which was the cytoplasmic SNARE protein SNAP-25 (Pooley et al., 2006). The screen yielded CENPF interaction with a second SNARE protein: syntaxin 4, the binding partner of SNAP-25 proteins, which helps mediate vesicular docking and fusion. Y2H analysis was used to further define binding domains within these proteins. From this assay, the first 474 amino acids of murine CENPF (NTmCENPF) were determined to be required for syntaxin 4 interaction (Fig. 1), and further reduction of this sequence eliminated syntaxin 4 binding altogether. Interestingly, our group identified this region of CENPF for its interaction with SNAP-25, whereas Feng et al. recognized the same area for the interaction of the human homolog with tubulin (Feng et al., 2006; Pooley et al., 2006). Y2H analysis determined
that the C-terminal 144 amino acids of syntaxin 4 were indispensable for CENPF binding (Fig. 1).

Transient protein expression within mammalian cells further confirmed the interaction of CENPF and syntaxin 4. The minimal syntaxin 4 binding domain of CENPF (myc-tagged N-terminal 474 amino acids of CENPF, termed NTmCENPF), and GFP-tagged syntaxin 4 were expressed in COS-7 cells. Lysates were prepared and co-immunoprecipitations (co-IP) were conducted with the appropriate antibodies. As seen in Fig. 2F, we were able to co-precipitate GFP-syntaxin 4 utilizing NTmCENPF, whereas all control experiments demonstrated no spurious GFP-syntaxin 4 association with CENPF. Syntaxin 4 did not react with any other sequences in CENPF except NTmCENPF (our unpublished data). Taken together, these results demonstrate that the N-terminal 474 amino acids of murine CENPF are necessary and sufficient for syntaxin 4 association.

Transiently expressed N-terminal murine CENPF and syntaxin 4 colocalize in cells

Knowing that these proteins interacted in Y2H and co-IP analyses, we investigated the subcellular localization of these proteins. Transient expression of NTmCENPF in COS-7 cells results in strong protein localization to the perinuclear region of cells and more diffusely to the cell periphery (Fig. 2A). As previously reported for syntaxins in COS cells (Banfield et al., 1994; Quinones et al., 1999), transient expression of GFP-syntaxin 4 leads to protein localization in multiple foci located throughout the cell (Fig. 2B). This punctate perinuclear distribution is very similar to previous reports of exogenously-expressed syntaxin in COS cells (Banfield et al., 1994; Quinones et al., 1999). However, in cells co-expressing both proteins, NTmCENPF redistributed to GFP-syntaxin 4-positive foci with a high degree of colocalization (Fig. 2C-E). Taken together, these data support our Y2H and co-IP data and demonstrate an interaction between murine CENPF and syntaxin 4.

Endogenous CENPF and syntaxin 4 associate in mammalian cells

We next examined the endogenous localization of CENPF and syntaxin 4 in murine cell lines. Two cell lines previously shown to express endogenous CENPF and syntaxin 4 were used: C2C12 myoblasts and 3T3 L1 adipocytes (Pooley et al., 2006; Soukoulis et al., 2005; Tortorella and Pilch, 2002). As seen in the myoblast line in Fig. 3A, confocal analysis demonstrated significant colocalization in the perinuclear region of the cell extending into the cell periphery. Band et al. (Band et al., 2002) have previously observed this perinuclear to peripheral distribution of endogenous syntaxin 4 in cultured NRK cells. In the present study, overlap was not absolute, as the staining pattern of CENPF extended further in the cell periphery than that of syntaxin 4. This is to be expected given that both proteins have been shown to bind other proteins and function in multiple pathways.

Syntaxin 4 function has been studied extensively in 3T3-L1 adipocytes (Cain et al., 1992; Pessin et al., 1999; Volchuk et al., 1996). Therefore, in prelude to our functional studies of this novel interaction, we examined the subcellular distribution of CENPF and syntaxin 4 in this cell line (Fig. 3C). The intense perinuclear staining of both CENPF and syntaxin 4 mirrored the cytoplasmic expression observed in C2C12 cells, but there was more significant staining of both proteins at the cell periphery than in the myoblast cell line (Fig. 3B). This pattern of syntaxin 4 localization reflects results previously reported for this cell type (Band et al., 2002). Furthermore, differentiated 3T3-L1 adipocytes demonstrated a high degree of endogenous CENPF and syntaxin 4 colocalization (Fig. 3C). Protein localization was broader in the differentiated cells and extended to the cell periphery.

To corroborate these results, we probed for the presence of endogenous complexes containing both CENPF and syntaxin 4 using immunoprecipitation of C2C12 lysates. As seen in Fig. 3D, CENPF was readily coprecipitated with syntaxin 4 (lane 4). By contrast, precipitations with beads alone (lane 2) and non-immune IgG (lane 5) were negative. As a positive control, we coprecipitated CENPF from the same lysate with an antiserum against the CENPF binding partner SNAP-25 (Fig. 3D, lane 3). Taken together, these data, identifying endogenous colocalization and biochemical interaction through antibody co-immunoprecipitation, support the hypothesis that CENPF associates with the endogenous SNARE complex within eukaryotic cells. 
Transient expression of NTmCENPF redistributes endogenous SNAP-25 and syntaxin 4

In an effort to define the role of murine CENPF in the regulation of vesicular function, we examined whether expression of its syntaxin-4-binding domain might influence subcellular localization of the trafficking apparatus in C2C12 cells. As seen in Fig. 4, forced expression of NTmCENPF resulted in the concentration of syntaxin 4 in perinuclear foci (compare Fig. 3 with Fig. 4A). SNAP-25, a known binding partner of both syntaxin 4 and CENPF also accumulated in this position but not completely, as other regions of the cytoplasm remained weakly positive for the anti-SNAP-25 antibody (Fig. 4B). Utilizing the trans-Golgi network (TGN) marker golgin-97, we determined that these NTmCENPF–syntaxin 4–SNAP-25 foci colocalized within TGN (Fig. 4C) and not with markers of the early or recycling endosome (data not shown).

VAMP2, but not VAMP3, also accumulated in the TGN of cells expressing NTmCENPF (Fig. 4D). We would also note that expression of GFP-syntaxin 4 alone also results in accumulation of endogenous CENPF in the TGN in C2C12 cells (data not shown).

Cotransfection was used to determine whether redistribution of components was a consistent feature of transient expression of NTmCENPF and syntaxin 4. As seen in Fig. 5A, expression of exogenous proteins resulted in the same accumulation of product in the Golgi and/or TGN. Consistent with expression of NTmCENPF alone, VAMP2 (Fig. 5B) and SNAP-25 (data not shown) were readily seen in this compartment, whereas VAMP3, Rab11a and EEA1 were excluded from the TGN. The dramatic redistribution of syntaxin 4, VAMP2 and SNAP-25 to the TGN after expression of NT-mCENPF suggests a role for this protein in vesicular transport.

Disruption of CENPF function interferes with cell coupling

Coupling of cells is dependent, in part, on the efficient trafficking of connexin molecules to and from the cell membrane (Wei et al.,...
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2004). Transfer of molecular dyes in cultured cells is an efficient and sensitive measure of connexin-based cell coupling (Francis and Lo, 2006). Dye coupling has the advantage of allowing the direct visualization of gap junction communication and function in a community of cells. For our purposes, 3T3 fibroblasts were transfected with GFP vector, GFP-syntaxin and NTmCENPF, or GFP-SNAP-25 and NTmCENPF. Living transfected cells were identified by GFP expression and loaded with sulforhodamine101 dye (dark red cell with yellow spot, Fig. 6A). Dye transfer from the injected cell to first tier cells (those in direct contact with the injected cell) and second tier cells (those in contact with first tier cells) were quantified using established methods (Francis and Lo, 2006). As seen in Fig. 6, dye is efficiently transferred to first and second tier cells after injection into control GFP transfected cells. In addition, this figure demonstrates that control transfection of cells in the first and second tier cells adjacent to the injected cells have no apparent affect on cell coupling. By contrast, expression of NTmCENPF with either GFP-labeled SNAP-25 or syntaxin 4 in the injected cell alone significantly inhibits the transfer of dye to neighboring cells, especially second tier cells (Fig. 6A,B). Inhibition of coupling is observed with co-expression of NTmCENPF with either syntaxin 4 or SNAP-25 even though NTmCENPF–syntaxin 4 complexes accumulate in the TGN (Fig. 5A) whereas SNAP-25–NTmCENPF concentrates in the recycling endosome (Pooley et al., 2006). Expression of NTmCENPF alone results in the same disruption of dye transfer (data not shown). These data provide direct evidence that CENPF function is important for proper cell coupling.

Inhibition of CENPF function inhibits glucose transport

Syntaxin 4 has a critical role in GLUT4 trafficking in insulin-responsive tissues (Martin et al., 1998; Cain et al., 1992; Pessin et al., 1999; Volchuk et al., 1996). Insulin stimulates the translocation of intracellular GLUT4 vesicle pools to the plasma membrane in target tissues, which include cardiac myocytes, skeletal muscle and adipose tissue. Activation of insulin receptors triggers a large increase of GLUT4 vesicle trafficking and exocytosis as compared to basal conditions. Additionally, although studies have concentrated on the function of SNAP-23 in GLUT4 trafficking, SNAP-25 has been isolated from GLUT4-positive cells (Jagadish et al., 1996). Thus, confident that this model could directly measure murine CENPF function in vesicular transport, we analyzed whether inhibition of CENPF function would inhibit GLUT4 trafficking. Briefly, 3T3-L1 cells were differentiated and CENPF function was inhibited using MO knockdown, a method previously described by our group (Ashe et al., 2004; Soukoulis et al., 2005; Pooley et al., 2006) in conjunction with the 2-deoxy-D-glucose transport assay (Kawanishi et al., 2000). The level of GLUT 4 trafficking was compared between differentiated 3T3-L1 adipocyte cultures treated with control (SC) or those treated with CENPF MO. To determine whether CENPF MO treatment was specific for CENPF depletion, replicate SC control and experimental MO-treated cultures were processed for western blot analysis. Fig. 7A demonstrates that CENPF protein levels drop significantly with experimental but not control MO application. (A) NTmCENPF and GFP-syntaxin 4 colocalize with the TGN marker golgin-97. (B) VAMP2 also colocalizes to a high degree with NTmCENPF and GFP-syntaxin 4. (C) VAMP3 does not demonstrate any significant redistribution to NTmCENPF–GFP-syntaxin 4 foci. (D) Rab11a, a marker of recycling endosomes, does not demonstrate noticeable colocalization. (E) Early endosomes, stained with EEA1, also show no significant redistribution to NTmCENPF–GFP-syntaxin 4 foci. Therefore, the NTmCENPF–GFP-syntaxin 4 complex is specific for localization at the TGN. NTmCENPF staining is indicated in blue, GFP-syntaxin 4 in green and the third marker, as indicated, is in red. Bar, 10 μm.

Fig. 5. Transiently expressed NTmCENPF and syntaxin 4 colocalize at the TGN in COS-7 cells. COS-7 cells were cotransfected with NTmCENPF and GFP-syntaxin 4, and only those cells expressing both transient proteins were analyzed. (A) NTmCENPF and GFP-syntaxin 4 colocalize with the TGN marker golgin-97. (B) VAMP2 also colocalizes to a high degree with NTmCENPF and GFP-syntaxin 4. (C) VAMP3 does not demonstrate any significant redistribution to NTmCENPF–GFP-syntaxin 4 foci. (D) Rab11a, a marker of recycling endosomes, does not demonstrate noticeable colocalization. (E) Early endosomes, stained with EEA1, also show no significant redistribution to NTmCENPF–GFP-syntaxin 4 foci. Therefore, the NTmCENPF–GFP-syntaxin 4 complex is specific for localization at the TGN. NTmCENPF staining is indicated in blue, GFP-syntaxin 4 in green and the third marker, as indicated, is in red. Bar, 10 μm.
observed with insulin treatment in SC MO-treated cultures and this response was statistically significant compared with no insulin treatment (P < 0.01). By contrast, glucose uptake was not observed in CENPF MO-treated cultures with or without insulin treatment. These data indicate that MO treatment alone is neither stimulatory nor inhibitory to glucose uptake.

Discussion
The CENPF family regulates diverse cellular processes
The CENPF family has diverse roles in cell cycle, division and differentiation. Although sequences in the C terminus have even been shown to regulate gene expression (Ma et al., 2006; Zhou et al., 2005), many important functions associated with CENPF are thought to be mediated through the microtubule network (Ashe et al., 2004; Dees et al., 2000; Pooley et al., 2006; Rattner et al., 1993; Soukoulis et al., 2005; Zhu et al., 1995b). For example, human CENPF binds the kinetochore and is highly expressed at the G1-S boundary (Zhu et al., 1997; Zhu et al., 1995b). The kinetochore is located at the centromere of the chromosome and serves as the site for microtubule spindle attachment during mitosis (Cleveland et al., 2003; Rieder and Salmon, 1998). Silencing of CENPF results in misalignment of chromosomes during mitosis and premature cell death (Yang et al., 2005), demonstrating a central role for this protein in microtubule regulation of mitosis and cell division. CENPF is also critical in the modulation of cell shape and motility. This protein directly interacts with Nde1, a regulator of dynein and microtubules (Soukoulis et al., 2005; Vergnolle and Taylor, 2007). Dynein, a microtubule-based motor, is also important for chromosome positioning and segregation through this interaction (Cleveland et al., 2003; Heald and Walczak, 1999; Sharp et al., 2000). In turn, Nde1 interacts with Lis1 in organelle positioning and movement, especially relating to developing neurons in the CNS (Feng et al., 2000; Feng and Walsh, 2004). Taken together, CENPF family proteins demonstrate diverse roles in organelle dynamics associated with microtubule-based processes. Although it is clear from this current study and previously reported data that CENPF family proteins are involved in organelle positioning, to date, the interacting proteins and regulators of such functions are largely unknown. Through a series of experiments to identify interacting proteins, we have now determined that murine CENPF provides a link between the MT network and vesicular transport.

Interaction with syntaxin 4 reveals a potentially broad role for murine CENPF in vesicular transport
Using genetic, biochemical and immunochemical analyses, we demonstrate that murine CENPF has a direct interaction with
syntaxin 4. This interaction is mediated through sequences in the N-terminal portion of CENPF that are highly conserved in various vertebrate genomes (NCBI search) and predicted to form the same tertiary coiled coil structure. The highly conserved nature of these sequences, their broad distribution in vertebrate genomes, the ubiquitous nature of CENPF expression in developing organisms, and the diversity of putative cellular function suggests a general role for this protein in cell function.

Both transiently expressed NTmCENPF and endogenous CENPF associate with syntaxin 4 in co-IP analyses. Furthermore, NTmCENPF and syntaxin 4 colocalize specifically to the TGN. Expression of the truncated form of NTmCENPF results in accumulation of this protein and syntaxin 4 in the TGN. As seen in previous studies (Rowe et al., 1999; Salaun et al., 2004; Takuma et al., 2002; Washbourne et al., 2001), exogenous expression of syntaxin 4 leads to its accumulation in the TGN (Fig. 5). Our data demonstrate that NTmCENPF also localizes to the TGN with the SNARE proteins SNAP-25 and VAMP2, both of which are associated with vesicular movement to and from the cell membrane. This is expected as both SNAP-25 and VAMP2 interact with syntaxin 4 (Pevsner et al., 1994). This result suggests a function for CENPF in vesicular transport and provides a method to analyze its potential role in this basic cell activity.

Murine CENPF is critical in regulation of vesicular transport

Previous studies on CENPF have demonstrated its roles in mitosis and cell division (Feng et al., 2006; Hussein and Taylor, 2002; Konstantinidou et al., 2003; Liao et al., 1995; Zhou et al., 2005; Zhu et al., 1995a; Zhu et al., 1997; Zhu et al., 1995b) and even gene transcription (Ma et al., 2006; Zhou et al., 2005). Other, in vivo and in vitro, analyses of CENPF and its binding partners suggest additional roles in cell movement and organelle positioning and translocation (Sasaki et al., 2000; Soukoulis et al., 2005; Vergnolle and Taylor, 2007). Identification of mCENPF interaction with syntaxin 4 and the dramatic redistribution of these proteins with exogenous expression of these binding partners strongly suggests a potential role in regulation of vesicular transport. To test the significance of CENPF-syntaxin 4 interaction, we employed two different experimental interventions. First, GLUT4 assay was used as a model of plasma membrane trafficking, as syntaxin 4 is critical in GLUT4 vesicle trafficking (Foster and Klip, 2000; Thurmond et al., 1998; Watson and Pessin, 2001). In depleting cells of CENPF, there was a significant decrease in the amount of labeled glucose recovered from 3T3-L1 adipocyte plasma membranes as compared with control groups. This is completely consistent with its role in movement of receptors to the cell surface by vesicular transport (Foster and Klip, 2000; Thurmond et al., 1998; Watson and Pessin, 2001). In addition, the inhibition of dye transfer between cells is a model to test the integrity of cell coupling after disruption of CENPF function by directly assaying for gap junction function at the cell surface (Francis and Lo, 2006; Lauf et al., 2002; Moskalewski et al., 1994). Our analyses show that disruption of murine CENPF function significantly diminished dye transfer between treated cells and clearly demonstrates inhibition of gap junction activity at the cell surface, resulting from mis-regulated protein trafficking to the cell membrane. Taken together with our previous work, the current data predict a broad role for CENPF in vesicular transport.

Materials and Methods

Yeast two-hybrid screen

This screen was previously described by Pooley et al. (Pooley et al., 2006). Briefly, a large N-terminal region of cytoplasmic murine CENPF (amino acids 1-689, termed ‘LCR’) was utilized in the Matchmaker Y2H System 3 (BD Biosciences Clontech). Library plasmids were isolated from yeast colonies that survived on quadruple dropout medium (QDO; SD-, Ade-, His-, Leu-, Trp-, X-a-Gal) and exhibited lacZ expression. The inserts were then sequenced by the Vanderbilt Sequencing Core Facility and identified using NCBI Blast (Altschul et al., 1990). For each identified protein product, false positive tests with empty vector and random protein matings were conducted to eliminate spurious interactions, according to manufacturer’s recommendations.

Cell culture, transfection and constructs

COS-7, NIH3T3-L1 and C2C12 cells (ATCC) were maintained in Dulbecco’s modified Eagle’s medium supplemented with 10, 10 and 20% FBS, respectively, 100 μg/ml penicillin-streptomycin and L-glutamine, in a 5% CO2 atmosphere at 37°C. 3T3-L1 pre-adipocytes were differentiated by treatment with insulin, dexamethasone and isobutyl methylxanthine as previously described (Front and Lane, 1985), and cells were used for experimentation 9-12 days after initiation of differentiation. For transfection, cells were grown to 50-75% confluency and transfected with DNA using FuGENE 6 (Roche) according to manufacturer’s recommendations. Murine N-terminal CENPF (NTmCENPF) was constructed by placing the N-terminal 474 amino acids of CENPF into the CMV-myc vector and full-length syntaxin 4 was placed into the EGFP-C3 vector (BD Biosciences Clontech).

Immunostaining and microscopy

For transient and endogenous studies, cells were gently washed with 1× PBS and fixed with either 4% paraformaldehyde to visualize endogenous proteins or with methanol to visualize transient protein for 20 minutes. Subsequently, cells were washed with 1× PBS, permeabilized with 0.25% Triton X-100 in 1× PBS for 10 minutes, and blocked for at least 1 hour in 2% BSA in 1× PBS at room temperature. Primary antibodies were incubated overnight at 4°C. Cells were then washed three times in 1× PBS and secondary antibodies were added for 1 hour at room temperature. Cells were again washed three times with 1× PBS and coverslips mounted with AquaPoly/Mount (PolySciences). Cells were visualized by fluorescence microscopy with an AX70 (Olympus), or for confocal analysis, with a LSM510 (Zeiss) microscope. Images were captured using Magnafire (Olympus). All images of control and experimental cells were processed identically.

Co-immunoprecipitation using transient transfections

COS-7 cells were grown on 10 cm plates; lysis were harvested 48 hours post transfection. The ProFound Mammalian Myc Tag Co-IP Kit (Pierce) was utilized according to manufacturer’s protocol. Briefly, cells were washed once with ice-cold TBS, incubated with M-Per extraction reagent (Pierce) containing protease inhibitor (Sigma), and centrifuged at 16,000 g for 20 minutes at 4°C. Lysate protein concentration of the supernatant was determined using a bichinchoninic acid solution assay (Pierce). For 2 hours, 100 μg total lysate was incubated with gentle shaking at 4°C with 10 μl anti-myc agarose slurry. Columns were washed three times with 1× TBS-Tween. Protein was eluted with 2× non-reducing sample buffer (Pierce) at 95°C for 5 minutes. To reduce proteins for SDS-PAGE analysis and western blot analysis, 2 μl 2-mercaptoethanol was added. Total lysate supernatant (10 μl) was used to confirm protein expression. Blots were developed using NBT-BCIP (Roche) and scanned (Hewlett-Packard) to produce digital images.

Co-immunoprecipitation of endogenous protein complexes containing murine CENPF

C2C12 cells were lysed with Nonidet P-40 buffer with gentien sonication. Whole cell lysates were recovered and samples containing 2-3 mg total protein were preclariﬁed with Gammabind Plus Sepharose (Amersham Biosciences) for 20 minutes with gentle rotation at 4°C. Cell lysates were collected and incubated overnight with 3 μg polyclonal syntaxin 4 antibody (Sigma). Gammabind Plus Sepharose was added to bind the antibody-protein complex. Beads were washed 3 times with cold 1× PBS and proteins were eluted with Laemmli sample buffer at a boiling temperature for 5 minutes. Proteins were resolved on a 6% SDS-PAGE gel and analyzed by western blotting. Protein lysate (20 μg) was loaded to visualize CENPF in whole cell lysate.

MO antisense oligomer treatment

Production of and methods utilizing morpholino oligonucleotides (MO) to speciﬁcally knock down endogenous CENPF have been previously reported (Ashe et al., 2004; Pooley et al., 2006; Soukoulis et al., 2005).

2-Deoxy-Glutamine glucose transport assay

Forty-eight hours after MO addition, 3T3-L1 adipocytes were serum-starved for 1 hour. The cells were then incubated with 100 nM insulin in KRH buffer for 20 minutes. Glucose transport was initiated by addition of 0.5 mM 2-deoxy-[1,2-3H]glucose (0.25 μCi). After 10 minutes, transport was terminated by washing the cells three times with cold KRH buffer. Cells were then solubilized with 0.5% SDS, and the incorporated radioactivity was measured by liquid scintillation counting. All quantitative data are representative of three separate experiments conducted over 3 days, each with n=6-8. As a control to demonstrate the insulin-dependent nature of glucose transport with morpholino inhibition, standard control and CENPF MO-treated cultures were assayed for glucose uptake with and without insulin stimulation. A
one-sample Student’s t-test was used after normalization to standard control (SC) cell populations.

Cell coupling
Cultured 3T3 fibroblasts were transfected with GFP alone as a control, both NtMnCENPF and GFP-SNAP-25, or both NtMnCENPF and GFP-syntaxin 4. Living transfected cells were identified and to quantitatively assess dye coupling, intracellular impedance was carried out with microelectrode filled with with sulfonamide101. The fluorescent dye was injected into the impaled cells iopthetrophonically using a current pulse of 1-3 nA of 0.5 second duration once per second for a total duration of 2 minutes. After an additional 3 minutes, the total extent of dye spread was recorded as the number of surrounding cells containing the injected dye. Transfer of dye was quantified and outlined according to published methods in first and second tier cells (Francis and Lo, 2006).

Antibodies
Rab11a (a gift from James Goldenring, Vanderbilt University) and murine CENPF antibodies were previously described (Poole et al., 2006; Soukkalits et al., 2005). SNAP-25, syntaxin 4, and β-tubulin antibodies were obtained from Sigma; Golgin-97 was obtained from Molecular Probes; VAMP2 and VAMP3 antibodies were purchased from StressGen; syntaxin 4, α- and γ-GFP antibodies were obtained from BD Biosciences; Alexa-Fluor-488- and Alexa-Fluor-568-conjugated secondary antibodies were also utilized (Molecular Probes). For triple labeled immunofluorescence studies, polyclonal anti-my (NISU) was directly labeled with the Xenon Alexa-647 labeling kit (Molecular Probes). Alkaline phosphatase-conjugated secondary antibodies for western blot were also purchased from Sigma.

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References
Alkawa, Y., Xia, X. and Martin, T. F. (2006). SNAP25, but not syntaxin 1A, recycles Rab11a (a gift from James Goldenring, Vanderbilt University) and murine CENPF antibodies were previously described (Poole et al., 2006; Soukkalits et al., 2005). SNAP-25, syntaxin 4, and β-tubulin antibodies were obtained from Sigma; Golgin-97 was obtained from Molecular Probes; VAMP2 and VAMP3 antibodies were purchased from StressGen; syntaxin 4, α- and γ-GFP antibodies were obtained from BD Biosciences; Alexa-Fluor-488- and Alexa-Fluor-568-conjugated secondary antibodies were also utilized (Molecular Probes). For triple labeled immunofluorescence studies, polyclonal anti-my (NISU) was directly labeled with the Xenon Alexa-647 labeling kit (Molecular Probes). Alkaline phosphatase-conjugated secondary antibodies for western blot were also purchased from Sigma.


