Inducible Exoc7/Exo70 knockout reveals a critical role of the exocyst in insulin-regulated GLUT4 exocytosis

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Insulin promotes glucose uptake by triggering the translocation of glucose transporter type 4 (GLUT4) from intracellular vesicles to the plasma membrane through exocytosis. GLUT4 exocytosis is a vesicle fusion event involving fusion of GLUT4-containing vesicles with the plasma membrane. For GLUT4 vesicle fusion to occur, GLUT4 vesicles must first be tethered to the plasma membrane. A key tethering factor in exocytosis is a heterooctameric protein complex called the exocyst. The role of the exocyst in GLUT4 exocytosis, however, remains incompletely understood. Here we first systematically analyzed data from a genome-scale CRISPR screen in HeLa cells that targeted virtually all known genes in the human genome, including 12 exocyst genes. The screen recovered only a subset of the exocyst genes, including exocyst complex component 7 (Exoc7/Exo70). Other exocyst genes, however, were not isolated in the screen, likely because of functional redundancy. Our findings suggest that selection of an appropriate exocyst gene is critical for genetic studies of exocyst functions. Next we developed an inducible adipocyte genome editing system that enabled Exoc7 gene deletion in adipocytes without interfering with adipocyte differentiation. We observed that insulin-stimulated GLUT4 exocytosis was markedly inhibited in Exoc7 KO adipocytes. Insulin signaling, however, remained intact in these KO cells. These results indicate that the exocyst plays a critical role in insulin-stimulated GLUT4 exocytosis in adipocytes. We propose that the strategy outlined in this work could be instrumental in genetically dissecting other membrane-trafficking pathways in adipocytes.

A major function of insulin is to promote glucose uptake into adipocytes and skeletal muscles, a process mediated by the glucose transporter GLUT4 (1–3). Under basal conditions, GLUT4 is sequestered in intracellular storage vesicles (1, 4–6). Insulin stimulation activates a cascade of signaling events that ultimately triggers fusion of GLUT4 vesicles with the plasma membrane (GLUT4 exocytosis), delivering GLUT4 to the cell surface to facilitate glucose uptake (1, 7, 8). Like other intracellular vesicle fusion events, GLUT4 vesicle fusion is driven by a conserved engine composed of soluble NSF attachment proteins (SNAREs)3 and Sec1/Munc18 (SM) proteins (9–17).

To form the SNARE-SM vesicle fusion machinery, exocytic vesicles must first be tethered to the plasma membrane (18–20). An evolutionarily conserved tether factor is the exocyst, a heterooctameric complex comprised of Exoc1/Sec3, Exoc2/Sec5, Exoc3/Sec6, Exoc4/Sec8, Exoc5/Sec10, Exoc6/Sec15, Exoc7/Exo70, and Exoc8/Exo84 (Fig. 1A) (21–26). As a member of the multisubunit tethering complex family, the exocyst interacts with lipids, small GTPases (e.g. Rabs), and SNAREs to promote vesicle tethering in a broad range of eukaryotic exocytic pathways (27–34).

In previous genetic studies, the exocyst was suggested to regulate tethering of GLUT4 vesicles (29, 35–38). In another study, however, manipulations of the exocyst had little effect on GLUT4 exocytosis (39). These genetic studies, however, mainly relied on RNAi and overexpression of dominant-negative mutants, which often suffer insufficient gene silencing as well as off-target effects. Moreover, mutations of the exocyst impair differentiation of adipocytes (40), complicating functional analysis. To definitively establish the role of the exocyst in GLUT4 exocytosis, these limitations needed to be overcome.

In this work, we first analyzed data from a genome-scale CRISPR genetic screen performed in HeLa cells. We found that only a subset of mammalian exocyst genes, including Exoc7, were isolated as significant hits in the screen. By contrast, other exocyst genes did not exhibit a phenotype. Next we developed an inducible adipocyte genome editing system to ablate Exoc7 expression without interfering with adipocyte differentiation.

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3 The abbreviations used are: SNARE, soluble NSF attachment protein; SM, Sec1/Munc18; gRNA, guide RNA; APC, allophycocyanin; dox, doxycycline; GMP-PNP, guanosine 5′-β,γ-imido)triphosphate.
We observed that *Exoc7* KO markedly inhibited insulin-stimulated GLUT4 exocytosis in adipocytes. Insulin signaling, however, remained intact in the KO cells. This study established a critical role of the exocyst in insulin-stimulated GLUT4 exocytosis in adipocytes.

### Results

#### Genome-scale genetic analysis reveals disparate KO phenotypes of exocyst genes

Genome-wide CRISPR screens are a powerful, unbiased approach to interrogate gene functions in mammalian cells (41). The GeCKO V2 CRISPR library contains guide RNAs (gRNAs) targeting virtually all known genes in the human genome, including 12 exocyst genes (42, 43). These 12 exocyst genes included the three paralogues of *EXOC3* (*EXOC3L1, EXOC3L2, and EXOC3L4*) and *EXOC6B*, a paralogue of *EXOC6* (Fig. 1B). Each protein-encoding gene in the library is targeted by six independent gRNAs (42, 43). In a cell population mutagenized by the CRISPR library, each cell usually receives a single gRNA so that only one gene is mutated (43).

Although GLUT4 is not expressed in HeLa cells, exogenously expressed GLUT4 is sorted into insulin-responsive vesicles in HeLa cells and undergoes insulin-stimulated exocytosis (42, 44, 45). HeLa cells expressing a GLUT4 reporter (HA-GLUT4-GFP) were mutagenized using anti-HA antibodies and APC-conjugated secondary antibodies. GFP and APC fluorescence was measured using flow cytometry. To inhibit insulin signaling, 100 nm wortmannin was added. Datasets were normalized to untreated WT cells. Data are presented as mean ± S.D. n = 3. **, p < 0.001; n.s., not significant, p > 0.05.

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ered as significant hits in the HeLa cell screen (Fig. 1B). By contrast, other exocyst genes did not show up as significant hits (Fig. 1B).

We next examined whether exocyst genes are essential to cell viability or growth. KO of an essential gene leads to cell death or growth arrest, which precludes accurate characterization of its function. We cultured the HeLa mutant cell population continuously for 4 weeks without sorting or insulin treatment. The abundance of gRNAs in the mutant cell population was then compared with that in the original CRISPR library. Essential genes were identified based on depletion (dropout) of its corresponding gRNAs in the mutant cell population (46). A total of 1,391 essential genes were identified, encompassing a range of pathways critical to cell physiology, such as protein translation, RNA transcription, and cell cycle regulation (Table S2). However, none of the 12 exocyst genes was among the essential genes (Fig. 1C and Tables S1 and S2), indicating that mutations of exocyst genes do not impair cell viability or growth.

To validate the results of the genome-wide genetic analysis, we used CRISPR genome editing to delete the \textit{EXOC7} gene, a significant hit in the HeLa cell screen (Fig. 1B) (42). We then used a flow cytometry–based assay to quantitatively measure insulin-regulated exocytosis of the HA-GLUT4-GFP reporter in individual cells (Fig. 1D) (42). Insulin stimulation strongly accelerated GLUT4 exocytosis, leading to translocation of the GLUT4 reporter to the cell surface (Fig. 1E and Fig. S1). We observed that insulin-stimulated GLUT4 translocation was abrogated in \textit{EXOC7} KO HeLa cells (Fig. 1E and Fig. S1). KO of \textit{EXOC3}, which was not a significant hit in the screen (Fig. 1B), had no effect on insulin-stimulated GLUT4 translocation (Fig. 1E). These genome editing data confirmed the results of the genetic screen.

These results demonstrated that the majority of mammalian exocyst genes exhibit no phenotype in CRISPR KO studies, likely because of functional redundancy (see “Discussion”). Thus, to genetically determine exocyst functions, it is critical to select an appropriate exocyst gene. In this work, we focused on \textit{EXOC7}, an exocyst gene with the strongest phenotype in the HeLa cell genetic screen (Fig. 1B).

**Doxycycline (dox)–induced KO of Exoc7 in adipocytes**

Although HeLa cells are a facile platform for initial analysis of exocyst genes, the role of the exocyst in insulin-stimulated GLUT4 exocytosis needs to be directly examined in adipocytes, a physiologically relevant insulin-responsive cell type. Because the exocyst is required for adipocyte differentiation (40), we developed an inducible adipocyte genome editing system in which a dox-controlled Cas9 expression cassette was stably integrated into mouse preadipocytes (Fig. 2A). Dox was added to transgenic preadipocytes shortly before onset of differentiation so that the gradually diminishing exocyst proteins could support adipocyte differentiation (Fig. 2B). As shown below (Fig. 3), preadipocytes differentiated normally into mature adipocytes when \textit{Exoc7} was deleted using this inducible CRISPR system. Because mutant preadipocytes cannot be cloned out, two distinct gRNAs were used to target the \textit{Exoc7} gene (Fig. 2A). Simultaneous introduction of two distinct gRNAs efficiently abrogates the expression of a candidate gene in a pooled cell population (42, 44). Indeed, we observed that \textit{Exoc7} expression was strongly reduced in adipocytes using this inducible CRISPR system (Fig. 2C).

**Inducible Exoc7 KO permits normal adipocyte differentiation**

A hallmark of adipocyte differentiation is accumulation of lipid droplets, which could be stained with the Nile red dye (Fig. 3A) (2, 47). After extensive testing, we identified a dox induction condition (Fig. 2B) that permitted \textit{Exoc7}–targeted preadipocytes to differentiate normally into mature adipocytes, as indicated by analysis of Nile red–stained lipid droplets using microscopy and flow cytometry (Fig. 3, A and B). Moreover, the overall morphology of \textit{Exoc7} KO adipocytes was indistinguishable from that of WT cells (Fig. 3A). The expression levels of PPARγ, a master regulator of adipocyte differentiation (48), were comparable between WT and \textit{Exoc7} KO cells (Fig. 3C). These data demonstrated that adipocyte differentiation occurred normally when \textit{Exoc7} was deleted using the inducible genome editing system.

**Insulin-stimulated GLUT4 exocytosis is inhibited in Exoc7 KO adipocytes**

We then examined the effect of \textit{Exoc7} KO on the GLUT4 exocytic pathway. We observed that insulin-stimulated GLUT4 translocation was markedly inhibited in \textit{Exoc7} KO adipocytes (Fig. 4A and Fig. S2). Further kinetic analysis showed that the exocytosis rate of GLUT4 was markedly reduced in \textit{Exoc7} KO adipocytes (Fig. 4B), correlating with GLUT4 translocation defects in these cells (Fig. 4A and Fig. S2). Next, we examined the responses of \textit{Exoc7} KO adipocytes to various concentra-
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We found that GLUT4 translocation was diminished in KO cells at all the insulin concentrations we tested (Fig. 4). These results demonstrated that Exoc7 is essential for insulin-stimulated GLUT4 exocytosis in adipocytes.

**Intact insulin signaling in Exoc7 KO adipocytes**

Next we examined whether Exoc7 KO influences insulin signaling. Insulin treatment induced the phosphorylation and activation of AKT/PKB, a downstream serine/threonine kinase in the insulin signaling cascade controlling GLUT4 trafficking (Fig. 5) (49–51). We observed that insulin-induced AKT phosphorylation remained unchanged in Exoc7 KO adipocytes (Fig. 5). Likewise, insulin-induced phosphorylation of AS160, a Rab-binding protein connecting insulin signaling to GLUT4 exocytosis (52), occurred normally in KO cells (Fig. 5). The expression level of sortilin 1 (Sort1), a molecule involved in post-Golgi translocations of insulin. We found that GLUT4 translocation was diminished in KO cells but not preadipocytes. Lipid droplets in the cells were stained with Nile red, and the images were captured using a ×20 objective on an Olympus IX81 microscope. Scale bars = 50 μm. B, quantification of lipid droplet-positive cells (WT or Exoc7 KO). Cells were stained with Nile red and the fluorescence of Nile red was measured using flow cytometry. Data are presented as mean ± S.D. n = 3, n.s., not significant, p > 0.05. C, representative immunoblots showing expression of the indicated proteins in WT and Exoc7 KO adipocytes. Two independent samples were prepared for each cell line. M.W., molecular weight.
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GLUT4 sorting (53, 54), was not altered in Exoc7 KO adipocytes either (Fig. 5). Thus, the exocyst is dispensable for insulin signaling and Sort1 expression, consistent with its role in the downstream vesicle tethering step of GLUT4 trafficking.

Tight regulation of adipocyte genome editing using the inducible CRISPR system

Next we further characterized the inducible adipocyte genome editing system. We observed that Cas9 was expressed only in dox-treated adipocytes, correlating with loss of Exoc7 expression in these cells (Fig. 6A). In contrast, Exoc7 was expressed at WT levels in adipocytes that contained the CRISPR cassette but were not induced with dox (Fig. 6A). Thus, Cas9 expression and genome editing were strictly dependent on dox induction.

Finally, we sought to rule out an off-target effect of the Exoc7 KO phenotype observed in adipocytes. To this end, we expressed the human EXOC7 gene in Exoc7 KO preadipocytes (Fig. 6B). We observed that the human EXOC7 gene, which was not targeted by the mouse Exoc7 gRNAs, fully restored insulin-stimulated GLUT4 exocytosis in Exoc7 KO adipocytes (Fig. 6C and Fig. S2). Thus, the defect of GLUT4 exocytosis in KO cells was caused by loss of Exoc7 expression. These results also suggest that the inducible CRISPR system permits specific gene ablation in adipocytes.

Discussion

In this work, we provide direct evidence of a critical role of the exocyst in insulin-stimulated GLUT4 exocytosis in adipocytes. This genetic study was made possible by an inducible adipocyte genome editing system that overcame the limitations of previous genetic assays. A major advantage of this system is that the timing of genome editing can be coordinated with adipocyte differentiation. Upon dox induction, the Exoc7 gene was deleted, terminating new protein production. Existing Exoc7 proteins were gradually degraded, but the remaining proteins were able to support differentiation of adipocytes. Exoc7 proteins were eventually eliminated upon completion of adipocyte differentiation. In this system, Cas9 expression and adipocyte genome editing are tightly regulated by dox induction. No leaky genome editing was observed, even after prolonged culture of uninduced cells.

The exocyst promotes vesicle tethering by interacting with molecules on exocytic vesicles (27, 28). The small GTPase Rab10 is anchored to GLUT4 vesicles and is known to bind to the Exoc6 subunit of the exocyst complex (36). It is conceivable that the Rab10–exocyst interaction facilitates tethering of GLUT4 vesicles to the plasma membrane. RalA, another vesicle-associated GTPase, also binds to the exocyst and is expected to contribute to GLUT4 vesicle tethering (29, 34, 55, 56). Although vesicle tethering is clearly a key function of the exocyst, in yeasts, the exocyst also directly interacts with SNAREs and SM proteins (30, 31, 57, 58), suggesting a more active role of the exocyst in the vesicle fusion reaction. Further studies will be needed to determine whether the exocyst directly regulates the vesicle fusion machinery and whether it is involved in other steps of GLUT4 trafficking, such as movement of GLUT4 vesicles on cytoskeletons.

Insulin-stimulated GLUT4 exocytosis was markedly inhibited but not abolished in Exoc7 KO adipocytes. KO of Exoc7 using two independent gRNAs was highly efficient so that the partial inhibition of GLUT4 exocytosis was not due to incomplete gene ablation. Instead, the partial inhibition phenotype was likely due to the existence of a parallel tethering mechanism or the unique features of adipocytes. The narrow cytoplasm of adipocytes may enable GLUT4 vesicles to partially bypass the requirement of the exocyst in vesicle tethering. Another notable observation from this study is that GLUT4 exocytosis increased with higher concentrations of insulin in both WT and Exoc7 KO cells. This finding suggests that the exocyst is not a direct target of insulin stimulation in GLUT4 trafficking.

An unexpected observation of this study is that the majority of mammalian exocyst genes exhibit no phenotype in CRISPR KO studies. A lack of phenotype is likely due to compensation by (a) redundant gene(s). Indeed, both Exoc3 and Exoc6 have paralogous genes that are expected to be functionally redundant (Fig. 1B). Exoc2, Exoc4, and Exoc5 may also have functionally redundant paralogues that remain to be uncovered. Alternatively, these three exocyst genes might not be fully targeted in the CRISPR screen. In the CRISPR library, six independent
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To identify essential genes, the unsorted passage control mutant population of HeLa cells was cultured continuously for 4 weeks. The abundance of gRNAs in the passage control population was analyzed by deep sequencing as described above. The depletion of gRNAs was calculated by comparing the passage control population with the initial GeCKO V2 CRISPR DNA library. A CRISPR score was calculated for each gene based on fold changes in the abundance of its corresponding gRNAs using an established algorithm (46). A CRISPR score of \(-0.25\) was set as the cutoff, corresponding to a total of 1,391 essential genes in these cells. This number is consistent with the essential genes identified in previous genome-wide genetic studies (46, 63, 64).

**Genome editing in HeLa cells**

Two independent gRNA sequences were selected to target early constitutive exons of a candidate gene. One of the gRNAs was subcloned into the pLenti-CRISPR-V2 vector (Addgene, 52961). The other gRNA was subcloned into a modified version of the pLentiGuide-Puro vector (Addgene, 52963), in which the puromycin selection marker was replaced with a hygromycin selection marker. CRISPR plasmids were transfected into 293T cells along with pAdVantage (Promega, E1711), pCMV-VSVG (Addgene, 8454), and psPax2 (Addgene, 12260) as described previously (42). The 293T culture media containing lentiviral particles were harvested daily for 4 days and centrifuged in a Beckman SW28 rotor at 25,000 rpm for 1.5 h. Viral pellets were resuspended in PBS and used to infect HeLa cells. Infected HeLa cells were consecutively selected using 1 \(\mu\)g/ml puromycin (Sigma, 3101118) and 500 \(\mu\)g/ml hygromycin B (Thermo, 10687010).

The sequence of gRNAs targeting the human EXOC3 gene was CGCCCCGACGCTGAGACA (guide 1), and CAAAGACTCCACGACTGCG (guide 2). The sequence of gRNA targeting the human EXOC7 gene was TTCTCGTCTGCTTGTGAC (guide 1), and AATCTTGCCATGCTTCCC (guide 1).

**Measurement of insulin-stimulated GLUT4 exocytosis**

Insulin-stimulated GLUT4 exocytosis was measured using flow cytometry as described previously (42, 44, 65). Briefly, cells were washed three times with KRH buffer (121 \(\mathrm{mM}\) NaCl, 4.9 \(\mathrm{mM}\) KCl, 1.2 \(\mathrm{mM}\) MgSO\(_4\), 0.33 \(\mathrm{mM}\) CaCl\(_2\), and 12 \(\mathrm{mM}\) HEPES (pH 7.0)). After incubation in KRH buffer for 2 h, the cells were treated with 100 \(\mathrm{nM}\) insulin for 30 min. When applicable, 100 \(\mathrm{nm}\) wortmannin (Sigma, W1628) was added to the adipocytes 10 min before insulin treatment. After insulin stimulation, the cells were rapidly chilled, and surface GLUT4 reporters were stained using anti-HA antibodies (BioLegend, 901501) and APC-conjugated secondary antibodies (eBioscience, 17-4014). Subsequently, the cells were dislodged using Accutase (Innovative Cell Technologies, AT 104), and GFP and APC fluorescence was measured on a CyAN ADP analyzer (Beckman Coulter). To calculate normalized surface levels of GLUT4 reporters, the mean APC fluorescence (surface staining) was divided by mean GFP fluorescence (total reporters), and the obtained values were normalized to those of untreated WT samples. Data from populations of \(~5,000\) cells were analyzed.

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using the FlowJo software (FlowJo, LLC, v10) based on experiments run in biological triplicates.

To visualize GLUT4 exocytosis using confocal microscopy, Hela cells and preadipocytes were cultured as described above. After incubation in KRH buffer for 2 h, the cells were stimulated with 100 nM insulin for 30 min. The cells were then fixed using 4% paraformaldehyde (Sigma, P6148) and blocked using 2% BSA (Thermo, 064985). Surface GLUT4 reporters were stained using anti-HA antibodies and Alexa Fluor 568–conjugated secondary antibodies (Thermo, A-11004). Nuclei were stained with Hoechst 33342 (Sigma, D9642). Images were captured using a ×100 oil immersion objective on a Nikon A1 laser-scanning confocal microscope.

**Inducible adipocyte genome editing**

To engineer an inducible genome editing system, we first created a TLCV2-DeGFP construct (deposited to Addgene, 133302) by deleting the GFP sequence from TLCV2, a lentiviral vector (Addgene, 87360) (66), using site-directed mutagenesis. Next, one gRNA targeting the mouse Exoc7 gene was subcloned into the BsmBI site of the TLCV2-DeGFP vector. The second gRNA targeting mouse Exoc7 was subcloned into the BbsI site of the pmU6-gRNA vector (Addgene, 53187). Next, the expression cassette of the second gRNA, including the mU6 promoter, the gRNA, and the gRNA scaffold, was amplified by PCR with KpnI sites introduced at both ends. The PCR product was subcloned into the KpnI site of the TLCV2-DeGFP vector containing the first gRNA. The resulting construct, TLCV2-Exoc7-Ex5&10 (deposited into Addgene, 133303), was transfected into 293T cells to produce lentiviruses as described above for HeLa cells. Lentiviral particles were used to infect preadipocytes, which were derived from inguinal white adipose tissues. These preadipocytes were a spontaneously arising immortal cell line (a gift from Dr. Shingo Kajimura) that could be passaged for at least 13 generations without losing differentiation capacity. The transduced preadipocytes were selected using 3.5 μg/ml puromycin.

The primers for site-directed mutagenesis of the TLCV2 vector were CGAGGAGAATCTTGCCCCAGCTACCTAATCCGGCGAGCGCCATC (forward) and GATGTGCGCTAGGCTAGCGCTACCGGACCCCCAGCATCCTGCAGAACG (reverse). The sequence of gRNA targeting the mouse Exoc7 gene was AGAAGCTGCTGTTTGAGCGA (guide 1), and TTCTAGAGCTTTGGCCCCAA (guide 2). The primers for amplification of the expression cassette of the second gRNA were CGCGGTACCACCGGTATAGATCCGACGCCGCA (forward) and GCCGCGTGTTACCCGCCGCTCAGTACAA (reverse).

Preadipocytes were cultured in DMEM supplemented with 10% FBE and penicillin/streptomycin to ~95% confluence before a differentiation mixture was added at the following concentrations: 5 μg/ml insulin (Sigma, I0516), 1 mM tri-iodo-L-thyronine (T3, Sigma, T2877), 125 μM indomethacin (Sigma, I-7378), 5 mM dexamethasone (Sigma, D1756), and 0.5 mM 3-isobutyl-1-methylxanthine (Sigma, I5879). After 2 days, the cells were switched to DMEM supplemented with 10% FBE, 5 μg/ml insulin, and 1 mM T3. After another 2 days, fresh DMEM containing 10% FBE and 1 mM T3 was added. Adipocytes were usually analyzed 6 days after addition of the differentiation mixture. To induce Cas9 expression, dox (Sigma, D3447) was added 24 h before differentiation at a final concentration of 2 μg/ml. Dox was included in the cell culture media throughout the differentiation process.

**Expression of the EXOC7 rescue gene**

To express the EXOC7 rescue gene, we first created a SHC003BSD construct (deposited into Addgene, 133300) by replacing the puromycin selection marker of the SHC003 vector (Sigma, SHC003) with a blastocidin selection marker. We then created a SHC003BSD-DeGFP construct (deposited into Addgene, 133301) by deleting the GFP sequence from SHC003BSD using site-directed mutagenesis, introducing a Sall site at the same time. The human EXOC7 gene was amplified using PCR with a Sall site and three tandem copies of FLAG tags introduced at the 5′ end. A Nhel site was introduced to the 3′ end of the PCR product. The EXOC7 PCR product was then subcloned into the Sall and Nhel sites of the SHC003BSD-DeGFP vector. The resulting SHC003BSD-EXOC7 construct (deposited into Addgene, 133298) was transfected into 293T cells to produce lentiviral particles, which were used to infect preadipocytes. The transduced preadipocytes were selected using 10 μg/ml of blastocidin (Thermo Fisher Scientific, BP2647).

The primers for site-directed mutagenesis of the SHC003BSD vector were CAGATCCGCTAGCCGCTACCGTGCGCACCCGACATCTGGCGACGAACG (forward) and CGTGCTCTGCAGATGCTGGGGGCTGACGTCGACGGTAGCGCTAGCGATCTCG (reverse). The Primers for amplification of the human EXOC7 gene were CGGCTAGTCCTGGGCTGCTAGCGATCTCG (reverse). The Primers for site-directed mutagenesis of the SHC003BSD vector were CAGATCCGCTAGCCGCTACCGTGCGCACCCGACATCTGGCGACGAACG (forward) and CGTGCTCTGCAGATGCTGGGGGCTGACGTCGACGGTAGCGCTAGCGATCTCG (reverse). The Primers for amplification of the human EXOC7 gene were CGGCTAGTCCTGGGCTGCTAGCGATCTCG (reverse).

**Immunoblotting**

Cells grown in 24-well plates were lysed in a SDS protein sample buffer. Cell lysates were resolved on 8% BisTris SDS-PAGE and probed using primary antibodies and horseradish peroxidase–conjugated secondary antibodies. Primary antibodies used in immunoblotting included mouse monoclonal anti-Exoc7 antibodies (Santa Cruz Biotechnology, sc-365825), mouse monoclonal anti-Sort1 antibodies (Santa Cruz Biotechnology, sc-76576), mouse monoclonal anti-PPARγ antibodies (Santa Cruz Biotechnology, sc-7273), mouse monoclonal anti-α-tubulin (DSHB, clone 12G10), rabbit monoclonal anti-Rab10 antibodies (Cell Signaling Technology, 8127), rabbit polyclonal anti-phospho-AKT (Ser-473) antibodies (Cell Signaling Technology, 9271), rabbit polyclonal anti-AKT antibodies (Cell Signaling Technology, 9272), rabbit monoclonal anti-AS160 antibodies (Cell Signaling Technology, 2670), rabbit monoclonal anti-phospho-AS160 (Thr-642) antibodies (Cell Signaling Technology, 1340), and mouse monoclonal anti-FLAG antibodies (Sigma, F1804).

**Lipid droplet staining and analysis**

To analyze lipid droplets using microscopy, preadipocytes and adipocytes grown in clear-bottom 96-well plates were fixed using 4% paraformaldehyde. To stain lipid droplets, Nile red

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(Sigma, N3013) was added to the cells at a final concentration of 5 μg/ml in PBS. After 15 min, the cells were washed three times with PBS and visualized on an Olympus IX81 microscope using a ×20 objective. Nile red fluorescence was detected using the FITC channel. For flow cytometry analysis of lipid droplets, cells were stained with Nile red in the presence of 10 μg/ml Hoechst 33342. Subsequently, the cells were detached using Accutase, and the fluorescence of Nile red and Hoechst 33342 was measured on a CyAN ADP analyzer. Data from populations of ~5,000 cells were analyzed using FlowJo software based on experiments run in biological triplicates. Fluorescence of unstained WT preadipocytes was used as a control blank.

Statistical analysis

Statistical significance was calculated using datasets from at least three independent experiments. Student’s t test was used to compare two experimental groups, whereas one-way analysis of variance was used to compare more than two groups.

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