Corrections

COMMENTARY. For the article “Reprogramming stem cells is a microenvironmental task,” by Mina J. Bissell and Jamie Inman, which appeared in issue 41, October 14, 2008, of Proc Natl Acad Sci USA (105:15637–15638; first published October 8, 2008; 10.1073/pnas.0808457105), the authors note that due to a printer’s error, ref. 4 was omitted and refs. 6–8 appeared out of order. The corrected references appear below.


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BIOCHEMISTRY. For the article “Transport of LDL-derived cholesterol from the NPC1 compartment to the ER involves the trans-Golgi network and the SNARE protein complex,” by Yasuomi Urano, Hiroshi Watanabe, Stephanie R. Murphy, Yohei Shibuya, Yong Geng, Andrew A. Peden, Catherine C. Y. Chang, and Ta Yuan Chang, which appeared in issue 43, October 28, 2008, of Proc Natl Acad Sci USA (105:16513–16518; first published October 22, 2008; 10.1073/pnas.0807450105), the authors note that ref. 43 should instead appear as ref. 33, and refs. 33–42 should be numbered 34–43. The corrected references appear below.


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BIOPHYSICS. For the article “Hydrophobic association of α-helices, steric dewetting, and enthalpic barriers to protein folding,” by Justin L. MacCallum, Maria Sabaye Moghadam, Hue Sun Chan, and D. Peter Tieleman, which appeared in issue 15, April 10, 2007, of Proc Natl Acad Sci USA (104:6206–6210; first published April 2, 2007; 10.1073/pnas.0605859104), the authors note that on page 6209, right column, in Methods, line 3, “antiparallel geometry” should instead appear as “parallel geometry.” This error does not affect the conclusions of the article.

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IMMUNOLOGY. For the article “Inflammation and autoimmunity caused by a SHP1 mutation depend on IL-1, MyD88, and a microbial trigger,” by Ben A. Croker, Brian R. Lawson, Michael Berger, Celine Eidenschenk, Amanda L. Blasius, Eva Marie Y. Moresco, Sosathya Sovath, Louise Cengia, Leonard D. Shultz, Argyrios N. Theofilopoulos, Sven Pettersson, and Bruce Alan Beutler, which appeared in issue 39, September 30, 2008, of Proc Natl Acad Sci USA (105:15028–15033; first published September 19, 2008; 10.1073/pnas.0806619105), the authors request that Sophie Rutschmann, Faculty of Medicine, Imperial College London, London SW7 2AZ, United Kingdom, be added to the author list, between Brian R. Lawson and Michael Berger, and be credited with performing research and analyzing data. The online version has been corrected. The corrected author and affiliation lines, and related footnotes, appear below.


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Correction

**BIOCHEMISTRY.** For the article “Transport of LDL-derived cholesterol from the NPC1 compartment to the ER involves the trans-Golgi network and the SNARE protein complex,” by Yasuomi Urano, Hiroshi Watanabe, Stephanie R. Murphy, Yohei Shibuya, Yong Geng, Andrew A. Peden, Catherine C. Y. Chang, and Ta Yuan Chang, which appeared in issue 43, October 28, 2008, of Proc Natl Acad Sci USA (105:16513–16518; first published October 22, 2008; 10.1073/pnas.0807450105), the authors note that ref. 43 should instead appear as ref. 33, and refs. 33–42 should be numbered 34–43. The corrected references appear below.


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Transport of LDL-derived cholesterol from the NPC1 compartment to the ER involves the trans-Golgi network and the SNARE protein complex

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Mammalian cells acquire cholesterol mainly from LDL. LDL enter the endosomes, allowing cholesteryl esters to be hydrolyzed by acid lipase. The hydrolyzed cholesterol (LDL-CHOL) enters the Niemann–Pick type C1 (NPC1)-containing endosomal compartment en route to various destinations. Whether the Golgi is involved in LDL-CHOL transport downstream of the NPC1 compartment has not been demonstrated. Using subcellular fractionation and immunoadsorption to enrich for specific membrane fractions, here we show that, when parental Chinese hamster ovary (CHO) cells are briefly exposed to 3H-cholesteryl linoleate (3H-CL-LDL), newly liberated 3H-LDL-CHOL appears in membranes rich in trans-Golgi network (TGN) long before it becomes available for re-esterification at the endoplasmic reticulum (ER) or for efflux at the plasma membrane. In mutant cells lacking NPC1, the appearance of newly liberated 3H-LDL-CHOL in the TGN-rich fraction is much reduced. We next report a reconstituted transport system that recapitulates the transport of LDL-CHOL to the TGN and to the ER. The transport system requires ATP and cytosolic factors and depends on functionality of NPC1. We demonstrate that knockdown by RNAi of 3 TGN-specific SNAREs (VAMP4, syntaxin 6, and syntaxin 16) reduces ≥50% of the LDL-CHOL transport in intact cells and in vitro. These results show that vesicular trafficking is involved in transporting a significant portion of LDL-CHOL from the NPC1-containing endosomal compartment to the TGN before its arrival at the ER.

Various bodily cells receive cholesterol mainly via LDL, the major cholesterol carrier in the blood cholesteryl ester (CE). In human LDL, cholesteryl linoleate (CL) is the predominant CE. LDL enters the cells mainly via LDL receptor mediated endocytosis (1). The endocytosed CE undergoes hydrolysis in an early endosomal compartment enriched in acid lipase (2); the cholesterol released then emerges in endosomes containing the protein Niemann–Pick type C1 (NPC1); from the NPC1 compartment(s), the LDL-derived cholesterol (LDL-CHOL) moves to various destinations including the plasma membrane (PM) and the endoplasmic reticulum (ER), etc. (2–4). At the ER, the LDL-CHOL can be re-esterified to CE by acyl-CoA:cholesterol acyltransferase (ACAT). There are 2 ACAT isoenzymes, ACAT1 and ACAT2. In healthy humans, ACAT1 is the main isoenzyme in most cell types, whereas ACAT2 is the main isoenzyme in intestinal enterocytes (5).

The NPC1 and NPC2 proteins play important roles in LDL-CHOL transport. NPC1 is a multispan membrane protein localized to a unique set of endosomes (6, 7); the NPC1-GFP fusion protein is also present in vesicles budding off from the endosomes (8). NPC2 is a soluble glycoprotein in the lumen of the late endosomes/lysosomes (LE/LYS) (9). Both NPC1 and NPC2 bind to cholesterol (10–15) and may cooperate in endosomal cholesterol transport (16). Whether the Golgi is involved in LDL-CHOL transport downstream of the NPC1 compartment has not yet been demonstrated. To address these issues, we performed pulse–chase experiments in CHO cells with LDL that contained 3H-cholesteryl linoleate (3H-CL-LDL) to monitor the fate of 3H-LDL-CHOL liberated in a time-dependent manner. We also developed an in vitro system that efficiently recapitulates the NPC1-dependent re-esterification of 3H-LDL-CHOL and used this system to study the mechanism of LDL-CHOL transport.

Results

Nocodazole or Brefeldin A (BFA) Affects the Re-Esterification and Efflux of 3H-LDL-CHOL For most of the studies reported here, we used a parental CHO cell line, 25RA (17) and a mutant NPC1 cell line, CT43 cells. 25RA cells contain a mutation in the sterol regulatory element-binding protein (SREBP) cleavage-activating protein (SCAP), which constitutively activates SREBP processing in a sterol-independent manner (18). CT43 cells are NPC1-null cells derived from 25RA cells (19). Findings made in this cell system are generally applicable to other cell types (8, 20–22). To monitor the hydrolysis and transport of 3H-LDL-CHOL liberated from 3H-CL, we pulse-labeled cells with 3H-CL-LDL for 30 min, then chased the cells without 3H-CL-LDL for up to 6 h [supporting information (SI) Fig. S1A]. The percentage of 3H-LDL-CHOL re-esterified by ACAT1 was used to estimate 3H-LDL-CHOL arrival at the ER. We also used the availability of labeled sterol to cyclodextrin (CD) in medium for 10 min to examine the movement of 3H-LDL-CHOL to the PM (2). The results indicate that in the 25RA cells, the re-esterification of 3H-LDL-CHOL occurs and increases linearly with time for up to 6 h, whereas in CT43 cells, re-esterification remains uniformly deficient (Fig. S1B). As the chase time increases from 20 min to 2 h, both the percentage re-esterification and the CD-mediated efflux of 3H-LDL-CHOL increased significantly in 25RA cells but not in CT43 cells (Fig. S1C Middle and Bottom) (2). The percentage hydrolysis of 3H-CL in CT43 cells was similar to that of 25RA cells (Fig. S1C Top). These results confirm previous studies (2, 3) and indicate that in cells lacking functional NPC1, the movement of LDL-CHOL from the NPC1 compartment to the ER and to the PM is defective. Using the pulse–chase protocol, we tested the effects of several compounds (cytochalasin D, wortmannin, latrunculin A, nocodazole, BFA, etc.) in 25RA cells for their ability to interfere with re-esterification or with efflux of 3H-LDL-CHOL. Because these compounds can also affect the endocytosis/internalization of the LDL (23), we treated cells with the test...
compound only during the chase but not during the pulse. The results (Fig. S1D) showed that nocodazole, an agent that disrupts the microtubules and disintegrates the Golgi within minutes (24), inhibited the re-esterification and efflux of \(^{3}H\)-LDL-CHOL by \(\approx 50\%\) (Fig. S1D Middle and Bottom). BFA, which causes fusion of the Golgi with the ER (25), inhibited efflux to the PM by \(\approx 40\%\) and increased re-esterification of \(^{3}H\)-LDL-CHOL by \(\approx 20\%\). These results suggest the involvement of the Golgi in the transport of LDL-ChOL and are consistent with a previous result using BFA in human fibroblast cells (26). Other drugs tested did not show significant effects on re-esterification (data not shown).

**Fate of \(^{3}H\)-LDL-CHOL Downstream of the NPC1 Compartment.** It is possible that the effects of nocodazole or BFA on re-esterification and on efflux of \(^{3}H\)-LDL-CHOL described above may be unrelated to their ability to disrupt the Golgi function. To further test the possibility that \(^{3}H\)-LDL-CHOL may move from the NPC1-containing compartment to the Golgi, we monitored the fate of \(^{3}H\)-LDL-CHOL by performing subcellular fractionation of labeled cell extracts. We first showed that the cellular organelles are separated into 6 major fractions (Figs. 1A and S2). NPC1 and NPC2 are principally located in fraction 6 (Fig. S2) but not in fraction 2, where ER/LYS are located. The same NPC1/NPC2 distribution patterns occurred in 25RA and WT CHO cell extracts (data not shown). These results suggest that in CHO cells, NPC1, and NPC2 are located in fractions distinct from the LE/LYS. In our current work, we designate the endocytic compartment(s) that contain NPC1 as “the NPC1 compartment.”

We next performed pulse–chase experiments and tracked the labeled cholesterol in various subcellular organelles. The results (Fig. 1B Left) show that at 0 time, in both the 25RA (blue) and the CT43 cell extracts (red), \(^{3}H\)-LDL-CHOL was enriched in fractions 4 and 5. At 20-min chase, in 25RA cells, fraction 5, rich in the Golgi and the early endosome (EE), was enriched in \(^{3}H\)-LDL-CHOL. In the CT43 cells, \(^{3}H\)-LDL-CHOL was enriched in fraction 2, rich in LE/LYS, consistent with the previous finding that in mutant NPC1 cells, cellular cholesterol accumulates abnormally within the LE/LYS (27). At later time point (2 h), in 25RA cell extracts, the \(^{3}H\)-LDL-CHOL appeared in fractions with higher densities (i.e., PM, ER, and mitochondria), whereas in CT43 cell extracts, the \(^{3}H\)-LDL-CHOL remained in fraction 2. The distribution of \(^{3}H\)-CL are very similar in 25RA and CT43 cells (Fig. 1B Right). For \(^{3}H\)-CO, it appeared as lipid droplets (fractions 1) in 25RA cells after 2 h, whereas no \(^{3}H\)-CO was produced in CT43 cells (data not shown). The drug U-18666A causes treated cells to exhibit a phenotype similar to that of NPC mutants (28). We performed additional experiments and found that the distribution patterns of \(^{3}H\)-LDL-CHOL from 25RA cells treated with U-18666A for 24 h were essentially the same as those of the CT43 cells (Fig. S2A). Additional experiments show that when 25RA cells were pulsed with \(^{3}H\)-CL-LDL and chased for 2 h, \(^{3}H\)-LDL-CHOL accumulated in the lower-density endosomal fractions (fractions 2 and 4) in nocodazole-treated cells during the chase, suggesting that rapid Golgi disintegration and microtubule disruption may create a transient NPC1-null phenotype, causing a significant portion of the LDL-ChOL to accumulate in the endosomes.

**LDL-ChOL Moves from the NPC1 Compartment to the TGN.** Subcellular fractionation can provide only partially purified membranes. To further explore the possibility that Golgi membranes may participate in LDL-ChOL transport, we used beads coated with antibodies against syntaxin 16, a TGN-protein marker, and performed selective immunoadsorption (IA) on post-nuclear supernatants (PNSs) prepared from 25RA cells. Beads coated with nonspecific rabbit antibodies were used as control. Immunoblot analysis revealed that the anti-syntaxin 16 adsorbed most of the syntaxin 16 and syntaxin 6-positive membranes (Fig. 2A); these membranes also contained a significant amount of the protein called vesicle-associated membrane protein 4 (VAMP4) and a small amount of caveolin-1 (cav1). These results are consistent with previous findings that syntaxin 6, syntaxin 16, and VAMP4 form the soluble N-ethylmaleimide (NEM)-sensitive factor (NSF) attachment protein receptor (SNARE) complex at the TGN (29), and that cav1 and syntaxin 6 can form a tight complex in the TGN membranes (30). The syntaxin 16-positive membranes did not contain any detectable amount of the early endosomal marker EEA1. We next performed pulse–chase and monitored the presence of \(^{3}H\)-LDL-CHOL in the syntaxin 16-rich membranes by IA. The results indicate (Fig. 2C) that in the 25RA cell extract, \(^{3}H\)-LDL-CHOL was associated with the syntaxin 16-rich membranes at 0 time; the association was increased after the 20-min chase. In the CT43 cell extracts, however, such association was less at 0 time, and no increase in association was observed after the 20-min chase. To expand on this observation, we carried out similar IA experiments by using beads coated with anti-syntaxin 6-specific antibodies or with nonspecific rabbit IgGs. We used a pool of OptiPrep fraction 5 membranes isolated from the 25RA cells as the starting materials. The immunoblot analysis revealed that anti-syntaxin 6 immunosorbent essentially all of the syntaxin 6-positive membranes; these membranes also contained a small amount of cav1 but did not contain EEA1 or NPC1 (Fig. 2B). We next performed a pulse followed by a 20-min chase in both the 25RA cells and the CT43 cells, isolated OptiPrep fraction 5 from the cell extracts, and carried out the IAs from these fractions. The results show that in the 25RA cell extracts, a significant amount of \(^{3}H\)-LDL-CHOL was associated with the syntaxin 6-rich membranes; whereas in the CT43 cell extracts, such association was much lower (Fig. 2D). Together, these
results are evidence that a significant portion of the LDL-CHOL is present in syntaxin 6- and syntaxin 16-rich TGN membranes, and that the movement of LDL-CHOL to the TGN depends on functional NPC1.

**Transport of LDL-CHOL from Endosomes to the ER in Vitro.** We next used the re-esterification of 3H-LDL-CHOL as a reporter for its arrival at the ER, and developed an in vitro assay that recapitulates the NPC1-dependent transport of LDL-CHOL in intact cells. We found that cells permeabilized with 0.001–0.003% digitonin at 4 °C for 5–15 min constituted a robust and reproducible in vitro system for LDL-CHOL transport in 25RA cells (outlined in Fig. 3A). We loaded the 3H-CL-LDL in intact 25RA cells by performing a pulse, then prepared semi-intact cells by permeabilizing the cell monolayers. We then added various factors, including rat liver cytosol, ATP/GTP, and oleoyl-CoA (as a substrate by ACAT1 in vitro), individually or in combination, and monitored the re-esterification of 3H-LDL-CHOL with time. We also conducted parallel experiments in AC29 cells. AC29 cells are mutant cells derived from 25RA cells, but AC29 cells lack ACAT1 (31). As shown in Fig. 3B Top, after incubation at 37 °C for 3 h, the hydrolysis of 3H-CL-LDL significantly increased in both 25RA and AC29 cells; however, significant re-esterification of 3H-LDL-CHOL occurred in 25RA cells but not in AC29 cells (Fig. 3B Bottom), demonstrating the dependency of re-esterification on ACAT1. Omitting ATP/GTP or cytosolic factors from the incubation mixture severely diminished re-esterification activity; omitting oleoyl-CoA partially reduced the activity (Fig. 3B Bottom). A separate experiment demonstrated that adding GTPyS (a nonhydrolyzable analog of GTP) also diminished the re-esterification activity in reconstituted 25RA cell system (data not shown). Additional experiments (Fig. 3C) show that after incubating at 37 °C for >1 h, 3H-LDL-CHOL re-esterification in reconstituted 25RA cell system increased with time for up to 3 h.

By contrast, in reconstituted CT43 cell system, re-esterification activity remained undetectable. This demonstrates that the re-esterification of 3H-LDL-CHOL in the reconstituted system depends on functional NPC1. In reconstituted 25RA cell system, 3% re-esterification occurred in 2 h and 5% in 3 h (Fig. 3C); in intact 25RA cells, 12% re-esterification occurred in 2 h, and 20% in 3 h (Fig. S1B). Thus, the efficiency of the reconstituted system is ~20–25% of that observed in intact cells. Vesicular trafficking requires soluble ATPas present in the cell cytosol. NEM inhibits ATPas and disrupts vesicular trafficking events in vitro. We tested the possible involvement of soluble ATPas in our in vitro system. The results show that NEM abolished the activity of the cytosol, whereas simultaneous inclusion of DTT partially prevented the inhibitory action of NEM (data not shown).

The results described in Fig. 3B and C suggest that the movement of LDL-CHOL from the NPC1 compartment to the ER in vitro involves vesicular transport. Numerous protein cargos undergo retrograde transport from the EE/recycling endosomes to the TGN by vesicular trafficking (32). To test this hypothesis that the transport of LDL-CHOL from the NPC1 compartment to the TGN may involve a similar mechanism, we performed an LDL-CHOL transport assay in reconstituted 25RA and CT43 cell systems for 2 h, then used the labeled PNSs to perform 1A experiments, using anti-syntaxin 6 or nonspecific antibodies as the immunoprobes. The results (Fig. 3D) show that a significant increase in association between 3H-LDL-CHOL and the syntaxin 6-rich membranes occurred in the reconstituted 25RA cells but not in the reconstituted CT43 cells. An additional experiment (Fig. 3E) showed that in reconstituted 25RA cells, 3H-LDL-CHOL was associated with the TGN-rich membranes, both ATP and cytosolic factors are

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**Fig. 2.** Association of 3H-LDL-CHOL with syntaxin 6-rich and syntaxin 16-rich membranes in intact 25RA cells. The 25RA and CT43 cells were pulsed with 3H-CL-LDL and chased in Medium A for 20 min. (A and C) The PNSs were immunoadsorbed with rabbit anti-syntaxin 16 (antiS16) antibodies or nonspecific rabbit antibodies (CNTL). (B and D) The PNSs were fractionated by Optiprep density gradient ultracentrifugation; the syntaxin 6-rich fractions (fraction 5) were isolated and immunoadsorbed with rabbit anti-syntaxin 6 (antiS6) antibodies or nonspecific rabbit antibodies (CNTL). (A and B) Associated proteins were analyzed by immunoblot using appropriate antibodies as indicated. The 25RA cells were used. (C) 3H-LDL-CHOL counts in the eluates were analyzed after lipid extraction and TLC. Values are presented as the percentage of total labeled sterol. Data are means ± SD from 2 experiments. *P < 0.05 by Student’s t test.

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**Fig. 3.** Cholesterol transport assay in semi-intact cells. (A) Procedure used to conduct the LDL-CHOL transport assay in vitro. (B) Hydrolysis and re-esterification in the presence or absence of ATP/GTP, or rat liver cytosol, or oleoyl-CoA as indicated, for 3 h at 37 °C. (C) Re-esterification in semi-intact 25RA or CT43 cells during the 3-h incubation time. (D and E) 3H-LDL-CHOL associated with syntaxin 6-rich membranes in semi-intact 25RA or CT43 cells with or without incubation for 2 h (D) or in semi-intact 25RA cells with or without ATP or rat liver cytosol during incubation as indicated (E). Values are presented as the percentage of total labeled sterol. Data are means ± SD from 2 experiments. **P < 0.01 by Student’s t test.
Knocking Down VAMP4/Syntaxin 6/Syntaxin 16 Diminishes the Re-esterification of ³H-LDL-CHOL in CHO Cells. The SNARE complex VAMP4/syntaxin 6/syntaxin 16 is required, suggesting that transport of LDL-CHOL from the NPC1 compartment to the TGN involves a vesicular mechanism.

The results show that VAMP4 KD (with iv4; or iv4; or iv4; or iv4) also efficiently reduced the VAMP4 protein expression in treated WT cells, without affecting the expression of other proteins examined (Fig. S5A). The re-esterification of ³H-LDL-CHOL in iv4 or iv4 treated WT cells was reduced by 11% or 28% (Fig. 4B Left). In WT cells treated with U-18666A, the re-esterification of ³H-LDL-CHOL remained very low with or without iv4 treatment (Fig. 4B Right). We next asked whether the role of VAMP4 in LDL-CHOL transport could be demonstrated in our in vitro system. We performed VAMP4 KDs in 25RA cells, and then conducted the in vitro assay for transport of LDL-CHOL to the ER. The results show that VAMP4 KD (with iv4; or iv4; or iv4) reduced the re-esterification of ³H-LDL-CHOL by 42% or 58%. To test whether VAMP4 is involved in the trafficking of LDL-CHOL to the TGN, we treated 25RA cells with an iv4 or with a CNTL siRNA for 48 h, performed the pulse, then chased for 30 min in intact cells (Fig. 4E) or chased in the reconstituted cell system for 2 h (Fig. 4F), then used the PNSs to perform IA experiments. The results show that a significant increase in association of ³H-LDL-CHOL to the syntaxin-6 rich membranes occurred in CNTL treated intact cells or reconstituted cell system, but not in iv4 treated intact cells or reconstituted cell system (Fig. 4 E and F). These results suggest that VAMP4 is involved in transporting LDL-CHOL to the TGN.

We next performed similar siRNA experiments on syntaxin 6. In the 25RA, CT43, and WT cells, after 72-h incubation, the 2 syntaxin 6 siRNAs (iS61 and iS62) reduced the syntaxin 6 expression by ≈50% without significantly affecting the expression of several proteins examined (Fig. S5B). The results of the LDL-CHOL trafficking assay in intact cells showed that iS60 or iS62 reduced the re-esterification of ³H-LDL-CHOL by 31% or 17% in 25RA cells (Fig. S5A Left) and by 57% or 29% in WT cells, respectively (Fig. S5B Left). KD of syntaxin 6 did not affect the transport of ³H-LDL-CHOL to the PM in 25RA cells (Fig. S5C). In the reconstituted 25RA cell system, syntaxin 6 KD (with iS60 or iS62) reduced the re-esterification of ³H-LDL-CHOL by 48% or 32% (Fig. S5D).

We next performed siRNA KD of syntaxin 16. In 25RA, CT43, or WT cells, after 72-h incubation, the 3 different syntaxin 16 siRNAs reduced syntaxin 16 expression by 50% (iS161), 80% (iS162), or 70% (iS163) of the control value (Fig. S5C). Syntaxin 6 expression was also modestly reduced by the syntaxin 16 KD, in agreement with a previous report (33). In 25RA cells, the re-esterification of ³H-LDL-CHOL was reduced by 42% (iS161), 73% (iS162), or 58% (iS163) (Fig. 5A Left). In WT cells, the re-esterification of ³H-LDL-CHOL was reduced by 48% (iS161), 56% (iS162), or 58% (iS163) (Fig. 5B Left). Syntaxin 16 KD in 25RA cells also did not significantly affect the transport of ³H-LDL-CHOL to the PM (Fig. 5C). In the reconstituted 25RA cell system, syntaxin 16 KD (with iS162 or iS163) reduced the re-esterification of ³H-LDL-CHOL by 50% or 45% (Fig. 5D). Together, these results implicate VAMP4/syntaxin 6/syntaxin 16 in the LDL-CHOL transport pathway in both the 25RA and WT CHO cell lines.

Discussion

Previous studies, using filipin as a probe for cholesterol, Blanchette-Mackie and colleagues (34) performed cytochemical freeze-fracture electron microscopy and showed that cholesterol content increased in the Golgi when human fibroblasts were incubated with LDL for 24 h, suggesting the involvement of the Golgi in LDL-CHOL transport. However, because of the low sensitivity of the filipin-staining method, the LDL incubation time was necessarily prolonged; this and other technical difficulties prevented the investigators from determining the role of the Golgi in LDL-CHOL transport (discussed in refs. 35–38). In our current work, we
Our results show that neither VAMP4, syntaxin 6, nor syntaxin 16 KDs reduce 3H-LDL-CHOL re-esterification in intact 25RA, WT cells, or reconstituted 25RA cell system. The 25RA, CT43, or WT cells were transfected with control (CNTL) or syntaxin 16 (iS16, iS16) siRNA and incubated for 72 h, then pulsed with 3H-CL-LDL. (A) Re-esterification in the 25RA or CT43 cells; chase time, 3 h. Data are means ± SD from 3 experiments. (B) Re-esterification in the WT cells treated with or without 0.3 μM U-18666A; chase time, 6 h. Data are means ± SD from 2 experiments. (C) Efflux to CD in the 25RA cells; chase time, 3 h. Data are means ± SD from 3 experiments. (D) Re-esterification in reconstituted 25RA cell system. Incubation time, 3 h. Data are means ± SD from 2 experiments. **P < 0.01 by Student’s t test.

We propose a working model to summarize a pathway for LDL-CHOL transport (Fig. S7): the endocytosed CE in LDL undergoes hydrolytic cleavage in a specialized EE; the CHOL liberated enters the NPC1-containing endosomes. By mechanisms(s) yet to be revealed, cholesterol-rich vesicles form; these vesicles move to the TGN by vesicular trafficking that involves the SNARE proteins VAMP4, syntaxin 6, and syntaxin 16. From the TGN, the LDL-CHOL moves to the ER for re-esterification by ACAT1, or moves to the PM. Additional trafficking mechanisms(s) also exist to transport LDL-CHOL. For example, in mutant NPC cells, overexpressing Rab9, a small GTPase enriched in the LE/LYS, efficiently rescues the LE/LYS cholesterol accumulation phenotype (45, 46). The following scenario may occur: In NPC1-null cells, LDL-CHOL enters the endosomes but cannot exit normally. It rapidly moves to the LE/LYS where it accumulates. A Rab9-dependent NPC1-independent cholesterol translocation pathway occurs between the LE/LYS and the TGN but operates at low efficiency. Overexpressing the Rab9 protein increases the efficiency of this pathway and bypasses the need for NPC1. It will be important to further dissect the molecular machineries involved in transporting LDL-CHOL. The availability of the reconstituted cell system described in this work should facilitate these studies.

**Materials and Methods**

**LDL-Derived Cholesterol Trafficking Assays.** The 25RA and CT43 cells were cultured in 6-well dishes in Medium A for 2–3 days, pulsed with 30 μg/ml 3H-CL-LDL in Medium A for 30 min at 37 °C, and chased in Medium A for 0–6 h (Fig. S1A). The labeled lipids were extracted and analyzed on silver nitrate impregnated TLC plates (47); the percentage hydrolysis of 3H-CL and the percentage re-esterification of 3H-CL-LDL were calculated as described (2). To conduct the cholesterol efflux experiments, cells were incubated with 4% 2-hydroxypropyl-β-CD in Medium D at 37 °C during the last 10 min of the chase period (2). Previously, we employed a “release of a temperature block” method to monitor the fate of 3H-CL-LDL (2). Control experiments show that the pulse–chase protocol described here reproduces all of the findings described in ref. 2. The WT cells were cultured in Medium A for 2 days, switched to Medium D with or without 0.3 μM U-18666A for 24 h, and chased in Medium A with or without U-18666A.

**Immunoadsorption.** Four hundred microliters of PNS material mixing with 200 μl of 60% OptiPrep solution, cleared by centrifuging at 15,000×g for 10 min, was used as the starting material. Alternatively, PNS was subject to 5–25% OptiPrep gradient ultracentrifugation for 3 h; fraction 5 was used as the starting material. 150 μl of the starting material were incubated for 1.5 h on ice with 150 mM NaCl and 2 μl of anti-syntaxin 16 antiserum or 2 μg of anti-syntaxin 6 IgG, or the same amount of nonspecific rabbit antiserum or IgGs in final vol. of 200 μl; the mixtures were incubated for 30 min on ice with 40 μl of μMac Protein A MicroBeads (Miltenyi Biotec), loaded onto the μColumn, the flowthrough (FT) was collected. After washing the beads with 800 μl of HB containing 150 mM NaCl, the proteins bound to the beads were eluted with 100 μl of 1 × SDS/PAGE sample buffer. Equal aliquots of the total input, eluate and FT samples were analyzed by immunoblots using appropriate antibodies. The 3H-labeled lipids bound to beads were collected by eluting beads with 500 μl of chloroform/methanol (2:1) and analyzed by TLC (2).

**LDL-CHOL Transport Assay in Digitonin-Permeabilized Semi-Intact Cells.** The assay was based on the assay developed to monitor protein transport between organelles in vitro (29). Cells were grown in Medium A in 6-well dishes, labeled with 30 μg/ml 3H-CL-LDL in Medium A for 30 min at 37 °C, then placed on ice,
washed twice with ice-cold transport buffer (TB, 25 mM Hepes-KOH, pH 7.2, 115 mM potassium acetate, 2.5 mM MgCl₂), and incubated for 10 min with 20 μg/ml (0.002%) digitonin in TB at 4 °C. The permeabilized cells were washed with ice-cold TB and incubated for 20 min at 4 °C to remove the residual cytosol. Over 99% of cells become permeabilized, as assessed by trypan blue staining. The semi-intact cells were incubated at 37 °C in 500 μL of TB with the following reagents: 200 μg of rat liver cytosol (Xenotech), the ATP regeneration system (1 mM ATP, 10 mM creatine phosphate (Alfa Aesar), 10 units/ml creatine phosphokine, 0.1 mg/ml BSA, 10 mM Hepes-KOH, pH 7.2), 250 μM GTP, 50 μM oleoyl-CoA, and 0.8 mM magnesium acetate, for various times. After incubation, the reaction was stopped by adding 500 μL of 0.4 M NaOH, and the cells were incubated for 30 min at room temperature. Labeled lipids were extracted and analyzed by TLC. To monitor the transport of LDL-CHOL to syntaxin 6-rich region in vitro, the semi-intact cells with or without incubation for 2 h in transport reaction mix, as indicated, were scraped and homogenized by using a glass homogenizer (with 20 strokes), followed by centrifugation at 16,000 × g for 10 min in 20% OptiPrep. The resultant PNSs were incubated with appropriate antibodies as indicated, then incubated with μMacs Protein A MicroBeads, and processed in the same manner as the immunoadsorption procedure described above.

Additional materials and methods are described in SI Materials and Methods.

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