Cholesterol trafficking and raft-like membrane domain composition mediate scavenger receptor class B type 1-dependent lipid sensing in intestinal epithelial cells
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*These authors contributed equally to this work

Centre de Recherche des Cordeliers, INSERM, UMPC Université Paris 6, Université Paris Descartes Paris 5, CNRS, F-75006 Paris, France

Clinical Innovation Proteomic Platform CLIPP, Université Bourgogne Franche-Comté, F-21000 Dijon, France

Plateforme de Lipidomique, INSERM UMR1231, Université de Bourgogne Franche Comté, F-21000 Dijon, France

Agroécologie, AgroSup Dijon, CNRS, INRA, Université Bourgogne Franche-Comté, F-21000 Dijon, France

EPHE, PSL Research University, F-75006 Paris, France

Center for Pulmonary and Vascular Biology, Department of Pediatrics, University of Texas, Southwestern Medical Center, Dallas, Texas, 75390, USA

Present address: Cell Biology Department of Institut Necker-Enfants Malades (INEM), INSERM U1151-CNRS UMR 8253, Université Paris Descartes Paris 5, F-75993 Paris, France

Present address: Chemosens platform, Centre des Sciences du Goût et de l’Alimentation, INRA, F-21000 Dijon, France

To whom correspondence should be addressed: Dr Véronique Carrière, INSERM U1138 team AL, Centre de Recherche des Cordeliers, 15 rue de l’école de médecine, 75006, Paris, France, Phone: 33144272407, Fax: 33143251615, Email: veronique.carriere@crc.jussieu.fr

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Abbreviations:

AGR2: anterior gradient protein 2 homolog
CALX: calnexin
CTTM: C-terminal transmembrane domain
DRM: detergent-resistant membranes
HDL: high density lipoprotein
HNRH1: heterogeneous nuclear ribonucleoprotein H
HNRPK: heterogeneous nuclear ribonucleoprotein K
ICAM1: intercellular adhesion molecule 1
JAM1: junctional adhesion molecule A
K1C9: keratin, type I cytoskeletal 9
KAP2:engthependent protein kinase type II-alpha regulatory subunit
LDHB: L-lactate dehydrogenase B chain
LPC: lysophosphatidylcholine
LPE: lysophosphatidylethanolamine
MβCD: methyl-β-cyclodextrin
NDUA2: NADH dehydrogenase (ubiquinone) 1 alpha subcomplex subunit 2
PC: phosphatidylcholine
PDE8A: high affinity cAMP-specific and IBMX-insensitive 3',5'-cyclic phosphodiesterase 8A
PE: phosphatidylethanolamine
PI: phosphoinositide
PL: phospholipids
pPE: plasmalogen phosphatidylethanolamine
PPM: postprandial lipid micelles
PS: phosphatidylserine
SM: sphingomyelin
SR-B1: scavenger receptor Class B type 1
STX16: syntaxin-16
VAMP8: vesicle-associated membrane protein 8
VATD: ion transport, such as V-type proton ATPase subunit D
VP9D1: VPS9 domain-containing protein 1
Abstract

Scavenger receptor Class B type 1 (SR-B1) is a lipid transporter and sensor. In intestinal epithelial cells, SR-B1-dependent lipid sensing is associated with SR-B1 recruitment in raft-like/detergent-resistant membrane domains and interaction of its C-terminal transmembrane domain with plasma membrane cholesterol. To clarify the initiating events occurring during lipid sensing by SR-B1, we analyzed cholesterol trafficking and raft-like domain composition in intestinal epithelial cells expressing wild-type SR-B1 or the mutated form SR-B1-Q445A, defective in membrane cholesterol binding and signal initiation. These features of SR-B1 were found to influence both apical cholesterol efflux and intracellular cholesterol trafficking from plasma membrane to lipid droplets, and the lipid composition of raft-like domains. Lipidomic analysis revealed likely participation of d18:0/16:0 sphingomyelin and 16:0/0:0 lysophosphatidylethanolamine in lipid sensing by SR-B1. Proteomic analysis identified proteins, whose abundance changed in raft-like domains during lipid sensing, and these included molecules linked to lipid raft dynamics and signal transduction. These findings provide new insights into the role of SR-B1 in cellular cholesterol homeostasis and suggest molecular links between SR-B1-dependent lipid sensing and cell cholesterol and lipid droplet dynamics.
1. Introduction

Cellular transport and trafficking of cholesterol, as well as the molecular composition of the plasma membrane, are critically involved in the governance of cell fate and cell metabolism. The scavenger receptor Class B type 1 (SR-B1) is primarily known for its function as a cholesterol transporter, particularly in the liver, where it is responsible for the last step in the reverse cholesterol transport pathway, which entails the selective uptake of cholesterol ester from High Density Lipoprotein (HDL). SR-B1 also mediates the bidirectional transport of cholesterol and phospholipids from peripheral tissues or cells to lipoproteins [1]. In the intestine, the functions of SR-B1 regarding cholesterol and lipid metabolism remain unclear. Although SR-B1 is expressed at the apical membrane of enterocytes [2-4], which are the absorptive cells of the intestinal epithelium, its role in intestinal cholesterol uptake has been questioned since NPC1L1 has been demonstrated to be responsible for 70% of cholesterol entry in enterocytes [5-7]. Moreover SR-B1-null mice [8] and mice overexpressing SR-B1 in the proximal intestine [9] do not display alterations in intestinal cholesterol absorption or in trans-intestinal cholesterol efflux [10]. However, other reports showed that SR-B1 is involved in the secretion of triglyceride-rich lipoproteins by enterocytes upon lipid supply [9, 11, 12].

In addition to serving in cholesterol movement, SR-B1 functions as a signal transducer in certain cell types, particularly in endothelial cells and in intestinal epithelium. In endothelial cells, the binding of HDL to SR-B1 triggers the sequential activation of protein kinases, among which the activating phosphorylation of Src kinase represents the earliest signaling event identified [13]. This signaling cascade leads to the production of nitric oxide and it promotes endothelial cell migration, which both contribute to the maintenance of endothelial monolayer integrity by HDL and SR-B1 [13]. In Caco-2/TC7 enterocyte-like cells, SR-B1 triggers intracellular signaling in response to the supply, at the apical pole of the cells, of postprandial lipid micelles (PPM), which mimic the form and composition of the dietary lipids that are present in the gut lumen following a meal. The sensing of PPM by SR-B1 activates MAP kinases and the trafficking of apolipoprotein B, the structural apolipoprotein required for triglyceride-rich lipoprotein assembly and secretion, from the apical region of the cell to basolateral secretory domains [14, 15]. These intracellular events are associated with the
recruitment of SR-B1 into raft-like membrane domains [14, 15], which are known to be signaling platforms [16].

Studies have been conducted to analyze the molecular mechanisms involved in SR-B1-dependent lipid sensing. It has been established that the C-terminal transmembrane domain (CTTM) and the C-terminal cytosolic domain of SR-B1 are required for its property to sense lipid movement and initiate intracellular signaling [17]. The discovery of the effects of a single point mutation in the SR-B1 CTTM domain, which represents the unique domain of SR-B1 able to interact with plasma membrane cholesterol [17, 18], highlighted the importance of plasma membrane cholesterol for SR-B1-dependent lipid sensing and resulting signaling [15]. This mutated form of SR-B1, created by the replacement of the glutamine residue in position 445 of SR-B1 amino acid sequence by an alanine residue (SR-B1-Q445A), provoked a marked decrease in the capacity of SR-B1 to bind plasma membrane cholesterol, whereas the capacities of the receptor to bind HDL and to transport cholesterol were unmodified [15]. In intestinal epithelial cells and endothelial cells expressing SR-B1-Q445A, the addition of ligand (PPM and HDL respectively) failed to activate signaling and subsequent cellular events [15]. Moreover, it was shown that SR-B1-Q445A is not recruited in raft-like membrane domains of intestinal epithelial cells after a PPM supply, contrary to what is observed for wild-type SR-B1 [14, 15]. The initiating cellular events involved in SR-B1-dependent lipid sensing remain unknown. However, plasma membrane cholesterol movement, through cyclodextrin treatment, is sufficient to reproduce, in intestinal epithelial cells and endothelial cells, all the signaling cascades and subcellular events in wild-type SR-B1-expressing cells while SR-B1-Q445A transfected cells are unresponsive [15]. Altogether these results emphasize the critical importance of plasma membrane cholesterol in SR-B1-dependent intracellular signaling, and they suggest that the initiating events in SR-B1-related lipid sensing take place at the plasma membrane and may involve cholesterol trafficking. Although it has been shown that SR-B1 reorganizes the free cholesterol pool in the plasma membrane [19-21], the modifications in the plasma membrane microenvironment that occur upon SR-B1 lipid sensing remain largely unknown.
The purpose of the present study was to analyze the early cellular events involved in intestinal epithelial SR-B1-dependent lipid sensing by analyzing cholesterol trafficking and the remodeling of raft-like membrane domains upon supply of lipid micelles.

2. Materials and Methods

2.1 Cell culture and lipid micelle supply

Caco-2/TC7 cell line is a clonal population of the human colon carcinoma-derived Caco-2 cells, which reproduces to a high degree most of the morphological and functional characteristics of enterocytes [22, 23]. These cells were stably transfected with constructs expressing mouse WT SR-B1 or SR-B1-Q445A as previously described [15]. The presence of SR-B1 at plasma membrane (Suppl. Fig.1) was shown by FACS analysis (BD LSRII FACS system and FlowJo software) performed on unpermeabilized cells as described in [24], using an anti-SR-B1 antibody (NB400-113 Novus Biologicals, Lille, France) and an anti-rabbit antibody conjugated to phycoerythrin (111-116-144 Jackson ImmunoResearch Laboratories Inc. West Grove, PA, USA). As previously described [15], SR-B1-Q445A has a dominant negative effect on signaling triggered by endogenous SR-B1 in both endothelial and Caco-2/TC7 cells. Cells were tested for mycoplasma contamination and found clean. In all experiments, cells were cultured on filter supports for 3 weeks to obtain fully differentiated enterocyte-like cells. Lipid micelles (2 mM sodium taurocholate, 0.6 mM oleic acid, 0.2 mM lysophosphatidylcholine, 0.05 mM cholesterol, and 0.2 mM monoacylglycerol) were prepared in serum-free medium as previously described [14, 15, 25] and added to the upper compartment for the indicated times. When appropriate, lipid micelles were supplemented with 5µM BODIPY-cholesterol (Avanti polar lipids, Alabaster, AL, USA) or with 5µCi/ml [1,2-3H(N)]-cholesterol (57,6Ci/mmol, PerkinElmer, Villebon-sur-Yvette, France). Transfection of Caco-2/TC7 cells with control RNAi and SR-B1 RNAi were performed as previously described [26].

2.2 Cholesterol labelling, trafficking and distribution

The intracellular distribution of cholesterol was analyzed in pulse-chase experiments. Caco-2/TC7 cells were incubated with BODIPY-cholesterol-containing lipid micelles for 1h hour, washed twice
with warm phosphate-buffered saline (PBS), and then incubated with non-fluorescent lipid micelles for one more hour. At the end of the experiment cells were rinsed with cold PBS and prepared for confocal microscopy analysis.

The trafficking of cholesterol from the plasma membrane towards intracellular domains was analyzed as described previously [27]. Briefly, BODIPY-cholesterol was combined with 370 mM methyl-β-cyclodextrin (MβCD, Sigma, Saint-Quentin Fallavier, France) in a 100:1 molar ratio (MβCD/BODIPY-chol). Caco-2/TC7 cells were incubated at their apical pole with MβCD/BODIPY-chol complex in serum-free medium for 1 min at 37°C. The final concentration of MβCD and BODIPY-cholesterol were 0.185 mM and ~1 µM, respectively. Cells were rinsed three times in warm PBS and then incubated with serum-free medium for the indicated times and analyzed by confocal microscopy.

Free cholesterol was visualized by filipin staining. Caco-2/TC7 cells were incubated or not with 3 µg/ml U-18666A (Sigma, Saint-Quentin Fallavier, France) added in both compartments of the filter support for 18h. Cells were then rinsed three times with PBS, fixed with 4% paraformaldehyde (PFA), and incubated with 0.05 mg/ml filipin (Sigma, Saint-Quentin Fallavier, France) in PBS/10% serum for 15 min. After extensive washing with PBS, cells were analyzed by confocal microscopy using a UV filter set.

### 2.3 Cholesterol efflux analysis

Caco-2/TC7 cells were cultured on filter supports and incubated with [1,2-^3^H(N)]-cholesterol containing lipid micelles in the apical compartment for 48h. Apical and basal compartments of the filter were then rinsed three times with serum-free medium. Cells were then collected (T0) or incubated with fresh serum-free medium in the basal compartment and with unlabelled lipid micelles devoid of cholesterol in the apical compartment for 2h. Total radioactivity was measured in cells (T0) and in apical medium (T2h) by liquid-scintillation counting after lipid extraction with chloroform/methanol (2:1, v:v).
2.4 Confocal microscopy analysis

Confocal microscopy analyses were performed as described [26]. Briefly, cells were fixed with 4% PAF, permeabilized by 0.05% saponin in PBS/10% serum and, when appropriate, stained for neutral lipids with LD540 dye [28]. After nuclear staining by DAPI and post-fixation with 4% PFA, images were acquired by laser scanning confocal microscopy (LSM 710 microscope; Carl Zeiss) and analyzed with ZEN software (Carl Zeiss).

2.5 RT-PCR analysis

Total RNAs were extracted from differentiated Caco-2/TC7 cells and used for real-time PCR analysis as previously described [29]. Oligonucleotide sequences were as follows: forward, 5’-GTGAGATGGCAGAGAACGGTGTG-3’ and reverse, 5’-TGCCCTTTGGTCTTTGTCCA-3’ for human PLIN2 mRNA; forward 5’- GCCTTAGCTACAGGAGAGA-3’ and reverse, 5’-TTTCCCTCTGTGACCATTCTC-3’ for human PPIB mRNA (reference gene).

2.6 Western blot analysis

Cell lysates and western blots were performed as previously described [14]. Primary antibodies were used against human SR-B1 (BD 610882, BD Biosciences, Le Pont de Claix, France) and human ANXA2 (BD 610068, BD Biosciences, Le pont de Claix, France).

2.7 Preparation of raft-like/detergent-resistant membranes

Detergent-resistant membranes (DRM) were prepared from a sucrose gradient as previously described [14]. Briefly, 10^6 Caco-2/TC7 cells were homogenized on ice in 2 ml of TRIS-buffered saline (TBS) (10 mM Tris-HCl, pH 8, 150 mM NaCl) containing 1% Triton X-100 and protease inhibitors. Cell homogenates were adjusted to 40% sucrose with 2 ml of 80% sucrose/TBS. The resulting 4 ml were covered with 4 ml of 30% sucrose and 4 ml of 5% sucrose and centrifuged (SW41, L8 Beckman, 18h, 39,000 rpm, 4°C). Sequential 1 ml fractions were collected from the top of the tube and fractions 3 to 5, corresponding to DRM were pooled.
2.8 Lipidomic analysis

Detergent-resistant membranes collected from the sucrose gradient were kept at -80°C until use. Lipid standards di-myristoyl phosphatidylcholine (DMPC), 19:0-lyso phosphatidylcholine (19:0-LPC), di-myristoyl phosphatidylethanolamine (DPME), myristoyl-lysophosphatidylethanolamine (MLPE), di-myristoyl phosphatidylserine (DMPS), d18:1/17:0 sphingomyelin (17:0-SM) and d18:1/17:0 ceramide were used; all of them were purchased from Avanti Polar Lipids (Coger, Paris, France). LC-MS/MS quality grade solvents were purchased from Fischer Scientific (Illkirch, France). Other chemicals of the highest grade available were purchased from Sigma Aldrich (Saint-Quentin Fallavier, France). For quantification of total cholesterol by GS-MS analysis, DRM fractions (20 µl) were spiked with epicoprostanol (2 µg) used as internal standard. Total cholesterol was quantified by GC-MS as previously described [30]. A calibration curve was generated with cholesterol standards processed as DRM fraction samples. For quantification of the different classes of lipids by LC-MS/MS, the DRM fraction (180 µl) was spiked with 20 µl of an internal standard mix containing 1000 ng DMPC, 250 ng 17:0-SM, 500 ng 19:0LPC, 200 ng DMPE, 400 ng DMPS, 100 ng MLPE and 100 ng d18:1/17:0 Cer. Total lipids were further extracted according to the method of Folch et al. [31]. Targeted lipidomic analysis by LC-MS/MS was conducted in Mutiple Reaction Monitoring mode as previously described [32].

2.9 Proteomic analysis

Detergent-resistant membranes collected from sucrose gradient were dialyzed (Flot-A-lyser G2 3.5-5kDa, Spectrumlabs, Breda, The Netherlands) for 3h at 4°C and then concentrated using an amicon Ultra-4 filter (ultracel-3 membrane 3kDa, Millipore, Fontenay-sous-Bois, France) according to the manufacturer’s instructions. DRM were then centrifuged (20 min, 12,000g, 4°C) and kept at -80°C until use. Proteins from DRM samples were processed as previously described [33]. Briefly, proteins were first extracted with a Urea-containing buffer (6M urea, 2.2M thiourea, 5mM EDTA, 0.1%SDS, 5% N-octyl glucoside, 50mM Tris-HCl) and separated shortly (1 cm) on a 1D 4-12 % acrylamide gel electrophoresis (ThermoScientific, Villebon sur Yvette, France). Proteins were excised from gel,
reduced, alkylated and digested by Trypsin Gold Mass Spectrometry Grade, (Promega, Charbonnières-les-Bains, France). Peptides (2 µg) were separated with the nanoRSLC (ThermoScientific) fitted with a C18 trapping column (5 µm average particle diameter; 300 µm inner diameter × 5 mm length; ThermoScientific) and a C18 analytical column (2 µm average particle diameter; 75 µm inner diameter × 150 mm length; ThermoScientific). A 120-minute gradient was performed and peptides were analyzed by nanoLC-MS/MS using an LTQ-Orbitrap Elite mass spectrometer equipped with the Advion TriVersa NanoMate nanospray source. Full-scan spectra from a mass/charge ratio of 400 to one of 1,700 at a resolution of 120,000 full width at half maximum were acquired in the Orbitrap mass spectrometer. From each full-scan spectrum, the 20 ions with the highest intensity were selected for fragmentation in the ion trap.

The acquired data were searched against the International Protein Index/UniProt using Mascot and X!Tandem. Based on tandem mass spectrometry data, peptides and proteins identifications were validated through Peptide- and ProteinProphet software [34]. At the end of the above steps, each injection is described by a list of validated proteins and peptides. Peptides that were not identified in at least 2 of the 3 technical replicates injected per sample were excluded [35]. Retention times were then aligned and peptide intensities extracted using the MASIC software [36]. Using this quantification, a new filter based on the coefficients of variation of each peptide in each sample is applied to discard poorly reproducible peptides [35]. A linear mixed model was used to select differential proteins in DRM from WT SR-B1- and SR-B1-Q445A-expressing cells (Clough et al., 2012). The FDR (false discovery rate) correction was applied to take into account the high dimension (far more tested variables than samples). Only proteins identified with at least 2 peptides fragments and with a fold change higher than 2, with a FDR at 5%, were kept.

2.10 Statistical analysis

For comparing two groups, unpaired Mann-Whitney test was used. For multiple comparisons, one-way ANOVA or two-way ANOVA were performed, followed by Dunnett’s multiple comparison tests
or Bonferroni’s multiple comparison tests, depending on the combinations of comparisons. All analyses were performed using Prism software (GraphPad prism software, La Jolla, CA, USA).

3. Results

3.1 SR-B1-dependent lipid sensing modulates cholesterol distribution in intestinal epithelial cells

We first analyzed the link between SR-B1-dependent lipid sensing and intracellular cholesterol distribution using the incapacity of the SR-B1-Q445A mutant to bind plasma membrane cholesterol and trigger intracellular signaling [15]. The intracellular distribution of cholesterol was studied in WT SR-B1- or SR-B1-Q445A-expressing Caco-2/TC7 cells in a pulse-chase experiment using BODIPY-cholesterol (Fig. 1A and Fig. 1B). Cells were incubated for 1h with BODIPY-cholesterol-containing lipid micelles followed by 1h of chase with unlabelled cholesterol-containing lipid micelles. At the end of the experiment, we observed an increased amount of BODIPY-cholesterol in SR-B1-Q445A-expressing cells as compared to WT SR-B1-expressing cells, at both the apical and intracellular levels. These results revealed impaired cellular cholesterol trafficking in intestinal epithelial cells when SR-B1 cannot trigger lipid sensing.

We then analyzed more precisely the intracellular localization of BODIPY-cholesterol in both WT and SR-B1-Q445A-expressing cells. In enterocytes as well as in Caco-2/TC7 cells, the supply of PPM is known to induce the formation of lipid droplets [26, 37], which store triglycerides and cholesterol esters. We analyzed the localization of BODIPY-cholesterol and of lipid droplets that were stained with the neutral lipid dye LD540. We observed that BODIPY-cholesterol co-localizes with lipid droplet staining (Fig 1C) in both WT and SR-B1-Q445A-expressing cells. However, SR-B1-Q445A cells display a more intense co-labeling (Fig. 1C, merge), which could be related to an increased accumulation of lipid droplets (Fig 1C, LD540 staining) in these cells as compared to WT SR-B1-expressing cells. Interestingly, such an accumulation of lipid droplets is also observed in Caco-2/TC7 cells transfected with RNAi for SR-B1 (Suppl. Fig. 2). We thus analyzed, in WT- and SR-B1-Q445A-expressing cells, the expression of PLIN-2 (Fig. 1D), which is known to be associated with lipid droplets and to modulate their formation and/or their stabilization [37, 38]. Before the addition of lipid micelles, we observed that the two cell lines express similar PLIN-2 mRNA levels. After a 24
hour incubation with PPM, an increase of PLIN-2 mRNA abundance is observed in both cell lines; however, the upregulation occurs to a greater extent in SR-B1-Q445A cells as compared to WT SR-B1 cells. Altogether these results indicate that SR-B1-dependent lipid sensing participates to the dynamics of lipid droplets.

We next studied whether the observed perturbation of cholesterol distribution in SR-B1-Q445A-expressing cells is associated with a modification of cholesterol efflux. Cells were incubated for 48h with [3H]-cholesterol-containing lipid micelles and then chased for 2h with lipid micelles devoid of cholesterol in the apical compartment (Fig.1E). We found that the apical efflux of [3H]-cholesterol is lower in SR-B1-Q445A-expressing cells as compared to WT SR-B1-expressing cells, whereas basal efflux of [3H]-cholesterol is similar in both cell types. Thus, the modification of cholesterol distribution in SR-B1-Q445A cells is associated with attenuated apical cholesterol efflux.

3.2 SR-B1 is involved in cholesterol trafficking from plasma membrane

Having previously observed that plasma membrane cholesterol movement is required for SR-B1-dependent lipid sensing [15], we analyzed the cholesterol trafficking between plasma membrane and intracellular compartments in WT SR-B1- and SR-B1-Q445A-expressing cells (Fig. 2). Cells were pulse-labelled by the addition of BODIPY-cholesterol/methyl-β-cyclodextrin complexes (Bdp-Chol/CD) to the apical compartment for 1 min at 37°C, as previously described [27], and then chased with a medium without Bdp-Chol/CD for 5min or 60min. This short pulse allows limited incorporation of BODIPY-cholesterol into the cells, thus avoiding a saturation of the labelling and a better visualization of cholesterol trafficking from the plasma membrane [27]. After a 1min pulse, the intensity of sub-apical BODIPY-cholesterol labelling is similar in both cells (1’ pulse, Fig. 2A and quantification in Fig. 2B). During the pulse-chase, the intensity of sub-apical BODIPY-cholesterol labelling rapidly decreased in WT SR-B1-expressing cells, probably due to a rapid metabolism or efflux of BODIPY-cholesterol. In contrast, in SR-B1-Q445A cells BODIPY-cholesterol continued to label the plasma membrane after 5 min of chase, and accumulated in intracellular compartments after 60 min of chase.
We next analyzed the capacity of plasma membrane cholesterol in WT SR-B1- and SR-B1-Q445A-expressing cells to be distributed in intracellular compartments. We visualized free cholesterol using filipin staining and studied free cholesterol movement using the properties of compound U-18666A to inhibit cholesterol transport leading to an intracellular accumulation of cholesterol [39]. Before U-18666A treatment, a similar localization of free cholesterol at the plasma membrane is observed in both WT and SR-B1-Q445A cell lines (Fig. 2C, ctrl panels). After U-18666A treatment, intracellular filipin signal is observed in WT SR-B1-expressing cells, indicating that free cholesterol is accumulating in vesicular structures (Fig. 2C, square showing an enlargement of the XY plan). By contrast, in SR-B1-Q445A-expressing cells treated with U-18666A, filipin staining is mainly observed at the plasma membrane, indicating that free cholesterol remains trapped at this level. The quantification of filipin fluorescence intensity showed a higher signal at plasma membrane (Fig. 2D) in SR-B1-Q445A cells treated with U-18666A (as compared to U-18666A-treated WT SR-B1 cells) that is associated with a decreased intracellular filipin dot numbers (Fig. 2E). Altogether these results reveal an impairment of cholesterol trafficking in cells expressing the SR-B1-Q445A mutant.

3.3 The expression of SR-B1-Q445A modifies the lipid composition of raft-like domain

We showed previously, in intestinal epithelial cells, that SR-B1-dependent lipid sensing is associated with the recruitment of SR-B1 in raft-like membrane domains [14, 15]. Raft domains of the plasma membrane are signaling platforms characterized by a large amount of cholesterol and sphingolipids that can be isolated by their properties to be resistant to detergent action [40]. We thus determined whether the observed perturbations of cholesterol trafficking in SR-B1-Q445A cells are associated with alterations in raft lipid composition. We isolated detergent-resistant membranes (DRM) from WT SR-B1 and SR-B1-Q445A cells treated or not with PPM for 15min, which is a time compatible with the activation of lipid signaling [14]. We conducted LC-MS/MS studies to determine lipid composition of DRM and analyzed the total concentration of 10 lipid subclasses, cholesterol, ceramides, phosphatidylcholine (PC), lysopPC (LPC), phosphatidylethanolamine (PE), lysoPE (LPE), plasmalogen PE (pPE), phosphoinositiide (PI), phosphatidyserine (PS) and sphingomyelin (SM), as well as several individual species in each subclass. In the absence of lipid micelle treatment (Fig. 3),
we observed that DRM from SR-B1-Q445A-expressing cells display higher concentrations of cholesterol (+32%, Fig. 3A), SM (+94%, Fig. 3B), PE (+24%, Fig. 3C), pPE (+72%, Fig. 3D), PI (+59%, Fig. 3E) and PS (+51%, Fig. 3F), as compared to DRM from WT SR-B1-expressing cells. By contrast, no change in the total concentration of PC, LPC, LPE and ceramides is observed between the two cell lines (Fig. 3G, H, I, J). Among the individual lipid species analyzed in the different lipid subclasses, d18:0/16:0 SM (Fig. 3K), one of the most represented SM, and d18:0/C20:0 ceramide (Fig. 3L), a minor species in this subclass, displayed the greatest increase of concentration (approximately a 3 fold change) in DRM from SR-B1-Q445A cells as compared to WT cells. Overall, the lipidomic analysis revealed that DRM from SR-B1-Q445A cells display higher concentrations of lipids (Fig. 4A), in particular of phospholipids (PL) (Fig. 4B), than those from WT SR-B1 cells. The qualitative analysis of lipid composition of DRM, achieved by the calculation of SM to total PL ratio (Fig. 4C) revealed that DRM from SR-B1-Q445A cells are enriched in SM as compared to WT SR-B1 cells. Altogether, these results show that the presence of the mutated form of SR-B1 modifies the lipid composition of DRM.

After the addition of lipid micelles, changes in DRM lipid composition were observed. Lipid micelle treatment promotes a marked increase in LPC concentration, but this modification occurs to the same extent in both cell lines (Suppl Fig. 3). This dramatic increase in LPC concentration probably results from the incorporation of exogenous LPC, provided by PPM, within cell membranes. More interestingly, after the supply of lipid micelles, the alterations in the concentrations of certain lipids differed in DRM of WT SR-B1 as compared to SR-B1-Q445A cells (Table 1). In DRM from WT SR-B1-expressing cells, the concentration of d18:0/16:0 SM increased by 17% upon lipid micelle provision, and the concentration of 18:0/0:0 LPE increased by 43%. In contrast, no alterations were observed in these lipids in DRM from SR-B1-Q445A-expressing cells after PPM treatment. These results indicate that d18:0/16:0 SM and 18:0/0:0 LPE may be involved in SR-B1-dependent signaling in response to lipid micelles. On the other hand, several lipid concentrations in DRM were decreased after PPM supply in SR-B1-Q445A-expressing cells whereas they remained unchanged after such treatment in WT SR-B1 cells (Table 1). This included a decrease of total ceramide (-28%) and of total PE (-10%) as well as several individual lipid species in different subclasses (Table 1).
Altogether these results show that lipid composition of raft-like membrane domains is modified during SR-B1-dependent lipid sensing.

3.4 SR-B1-dependent lipid signaling modifies the protein composition of raft-like domains.

In order to identify proteins in raft domains that could be involved in the SR-B1-dependent lipid sensing and resulting signaling in intestinal epithelial cells, we analyzed and compared changes in the amounts of DRM proteins after PPM supply in WT SR-B1- versus SR-B1-Q445A-expressing cells (Table 2). We isolated detergent-resistant membranes from both cell lines before and 15 min after PPM supply. The quantity of total protein was similar in all DRM (data not shown). Proteins were identified by mass spectrometry, and changes in protein amounts induced by PPM treatment for each cell line were determined.

We first analyzed modifications of protein amounts in DRM from WT SR-B1-expressing cells after PPM supply. We found that PPM supply promotes significant changes in the abundance of 41 proteins in DRM from WT SR-B1 cells, 25 of them being more represented and 16 of them less represented as compared to the condition without PPM treatment (Table 2). These proteins have been classified in several function categories according to UniProt Consortium annotations.

We then determined whether the amounts of these 41 identified proteins were also modified after PPM supply in DRM from SR-B1-Q445A cells. We observed that 5 of them (Table 2, green lines) displayed similar directional change in abundance in DRM from SR-B1-Q445A as compared to WT SR-B1 cells after PPM supply, suggesting that these proteins are not involved in SR-B1-dependent lipid sensing. More importantly, we found that 6 proteins, namely anterior gradient protein 2 homolog (AGR2), keratin, type 1 cytoskeletal 9 (K1C9), VPS9 domain-containing protein 1 (VP9D1), L-lactate dehydrogenase B chain (LDHB), Syntaxin-16 (STX16) and heterogeneous nuclear ribonucleoprotein H (HNRH1) displayed opposite variations in DRM from SR-B1-Q445A as compared to WT SR-B1 cells after PPM supply (Table 2, red lines). AGR2 is involved in protein processing, K1C9 belongs to an intermediate filament family, VP9D1 participates in ion transport, LDHB is involved in oxidation-reduction processes, STX16 plays a role in vesicular docking and fusion, and HNRH1 participates in the maturation of mRNA. Displaying disparate recruitment to
DRM in WT versus SR-B1-Q445A cells, these six proteins are candidates for molecules involved in SR-B1-dependent signaling in response to lipid movement. Additional candidates are the 30 proteins whose abundance changed after PPM supply in DRM from WT SR-B1 but not in SR-B1-Q445A cells (Table 2, uncoloured lines). These proteins belong to functional categories related to 1) cell adhesion/junction and cytoskeleton, such as the junctional adhesion molecule A (JAM1) and the intercellular adhesion molecule 1 (ICAM1), which are involved in cell-cell contact; 2) protein interaction/processing, such as peroxisomal membrane protein PEX14 involved in protein-protein docking, and calnexin (CALX) involved in protein folding; 3) oxidation/reduction, such as the NADH dehydrogenase (ubiquinone) 1 alpha subcomplex subunit 2 (NDUA2); 4) ion transport, such as V-type proton ATPase subunit D (VATD); 5) pre-mRNA processing, such as Heterogeneous nuclear ribonucleoprotein K (HNRPK); and 6) signal transduction, such as the cAMP-dependent protein kinase type II-alpha regulatory subunit (KAP2), protein FA83B and the High affinity cAMP-specific and IBMX-insensitive 3',5'-cyclic phosphodiesterase 8A (PDE8A). The vesicle-associated membrane protein 8 (VAMP8), which is involved in vesicular trafficking, is also present among the 30 proteins that remain unchanged after PPM supply in DRM from SR-B1-Q445A cells as compared to WT SR-B1 cells.

Altogether these results showed that SR-B1-dependent lipid sensing is associated with changes in the amount of several proteins in detergent-resistant membranes.

4. Discussion

SR-B1-dependent lipid sensing has been previously characterized in endothelial cells and in intestinal epithelial cells [14, 15, 41]. The discovery that the glutamine at the position 445 in the C-terminal transmembrane domain of SR-B1 was crucial in this process revealed that the lipid sensing is dependent on the interaction of SR-B1 with plasma membrane cholesterol [15]. The plasma membrane cholesterol sensing is also required for the recruitment of SR-B1 in raft-like membrane domains upon lipid micelle stimulation in intestinal epithelial cells. In the present study, we analyzed changes in the lipid and protein composition of raft-like domains occurring during SR-B1-dependent lipid sensing in intestinal epithelial cells, and also the impact of this sensing on cholesterol trafficking. We discovered
that a defect in SR-B1-mediated lipid sensing leads to an alteration of cholesterol distribution and to a decreased apical cholesterol efflux. Associated with these events, we observed, using SR-B1-Q445A or RNA interference, an exaggerated accumulation of lipid droplets in the absence of efficient SR-B1-dependent lipid sensing. Such a link between SR-B1 and lipid droplet dynamics could be suggested from previous reports. Indeed, SR-B1 is found associated with lipid droplets during fat absorption in enterocytes [42, 43]. SR-B1 is also present in mature adipocytes, where it could be involved in lipid storage in adipose tissue [44, 45]. Interestingly, our proteomic analysis of DRM from WT SR-B1 cells revealed increased DHB2 protein abundance in DRM after PPM supply, and this did not occur in DRM from SR-B1-Q445A-expressing cells. Although DHB2, which belongs to the family of 17b-hydroxysteroid dehydrogenases, is mainly known for its role in steroid metabolism, it has been shown to be associated with lipid droplets in intestinal epithelial cells. Its depletion increased triglyceride secretion, suggesting a role for this enzyme in the control of the balance between lipid storage and lipid secretion in these cells [29].

The impact of SR-B1 on cellular cholesterol metabolism has been mainly understood in the context of selective cholesteryl ester uptake from HDL and in the bidirectional flux of free cholesterol between cells and HDL (for review [21]). SR-B1 is known to be localized to caveolae/lipid rafts and to interact with several phospholipids [46-48]. A number of these studies indicated that SR-B1-dependent cholesterol flux may be mediated by changes in the lipid organization of the plasma membrane and particularly that of the raft membrane domains [49-52]. Our lipidomic analysis revealed that the presence of SR-B1-Q445A per se modifies the lipid composition of DRM as compared to cells expressing the wild type form of SR-B1. These modifications, which are characterized by increases in cholesterol, sphingomyelins, and several phospholipids, result in a relative enrichment of sphingomyelins versus total phospholipids in DRM. It has been observed that sphingomyelins are critical for cholesterol sequestration in the plasma membrane [53] and for the regulation of cholesterol efflux [54]. Moreover, modulation of SM amount in HDL or in cell membrane modified SR-B1-dependent cholesterol flux [55, 56]. Interestingly, two point mutations in the extracellular loop of human SR-B1 (S112F and T175A) result in a reduced efflux of free cholesterol to HDL and in the impairment of the redistribution of free cholesterol pools at the plasma...
membrane [57]. Thus, our findings provide evidence that, through its capacity to bind plasma membrane cholesterol, SR-B1 governs the lipid composition of raft-like membrane domains, thereby preparing a favourable environment for lipid sensing processes.

We analyzed additionally whether SR-B1-dependent lipid sensing is associated with modifications in the lipid composition of DRM. Lipidomic analysis showed that PPM supply provokes an enrichment of d18:0/16:0 SM and 18:0/0:0 lysophosphatidylethanolamines in WT SR-B1 cells. This enrichment is not observed in DRM from SR-B1-Q445A after PPM supply, suggesting that these lipids may be involved in SR-B1-dependent lipid sensing and in the resulting intracellular signaling. Although these lipids belong to classes of lipids recognized to be involved in cell signaling, their specific roles in such processes are largely unknown. Sphingomyelins are important structural component of lipid rafts that can be metabolized into ceramides, which induce changes in membrane fluidity [58] and impairment of receptor-dependent signaling [59, 60]. Lysophosphatidylethanolamines are minor components of the cell membrane which are able to induce calcium-related and MAPK signaling in several cell lines [61-64].

In addition to the changes in the lipid composition of DRM that occurred during SR-B1-dependent lipid signaling, we compared alterations in protein abundance in DRM from WT SR-B1- and SR-B1-Q445A-expressing cells after PPM supply. Proteins present in DRM are generally hydrophobic and their identification by proteomic analysis is dependent on their intrinsic properties and on the detergents used to solubilize and extract them [65, 66]. Moreover, we selected only proteins that were covered by more than 2 peptides. SR-B1 itself was not reproducibly identified, in reproducible manner, by our proteomic analysis procedure, probably due to its low solubility. Nevertheless, our study revealed modifications in the abundance of a variety of proteins involved in several cellular processes (Table 2). Most of them are related to the cytoskeleton and oxidation/reduction metabolism. It is well known that the dynamics of plasma membrane organization and lipid raft assembly are dependent on the actin cytoskeleton and that they consume large amounts of ATP (for review [67]). Additionally, many proteins involved in oxidation/reduction process are located in DRM [68] and they were identified to be altered by PPM supply (such as COX2, VATD, ATPK, VATF). Moreover, this dynamic organization of the plasma membrane is linked to the
endosomal/lysosomal system, where an ATP-dependent acidification of vesicles occurs [69, 70]. In our study, we also observed changes in the abundance of syntaxin-16 (STX16) and vesicle-associated membrane protein 8 (VAMP8). These proteins belong to the SNARE (soluble N-ethylmaleimide-sensitive factor attachment protein receptor) family, which is involved in membrane fusion and vesicular trafficking [71]. SNARE proteins participate also in cholesterol trafficking [72], and particularly in its movement from lipid droplets to mitochondria during steroidogenesis [73, 74]. SNARE proteins have been implicated additionally in autophagy [75], which is a cellular process that controls lipid droplet dynamics in intestinal epithelial cells [26]. Moreover, our proteomic analysis showed changes in the abundance of proteins involved directly in signal transduction. These included KAP2 (cAMP-dependent protein kinase type-II), and PDE8A (phosphodiesterase 8A), which are linked to cAMP-dependent signaling [76, 77], and FA83B (Protein FAM83B), which is involved in epidermal growth factor receptor activation and downstream RAS/MAPK and PI3K/AKT/TOR signaling cascades [78, 79]. Finally, we observed an increased abundance of PEX14 (Peroxisomal membrane protein PEX14) in DRM from WT SR-B1 cells after PPM supply that did not occur in DRM from SR-B1-Q445A cells. PEX14 is an important component of peroxisomal import machinery and it interacts directly with other peroxin proteins such as PEX5 [80, 81], which binds peroxisomal targeting signal type 1 (PTS1)–containing proteins such as SR-B1 [82]. Interestingly, PEX14 is also able to interact with LC3 protein involved in autophagy in a process dependent on nutrient availability [83]. Although the discrete functions of these proteins that are pertinent to their recruitment to DRM in response to PPM are yet to be elucidated, it is apparent that the dynamics of DRM composition are greatly influenced by the capacity of SR-B1 to bind plasma membrane-associated cholesterol and to initiate signaling in response to PPM.

In summary, our studies demonstrated that a point mutation in SR-B1 abolishing the property of the receptor to sense lipids induces modifications in lipid composition of raft-like membrane domains and cell cholesterol distribution in intestinal epithelial cells. The SR-B1-dependent lipid sensing influences both apical cholesterol efflux and intracellular cholesterol trafficking from the plasma membrane to lipid droplets. Two saturated lipids (d18:0/16:0 sphingomyelin and 16:0/0:0 lysophosphatidylethanolamine) and several proteins involved in lipid raft dynamics and signal
transduction play an important role in lipid sensing by SR-B1 (Fig. 5). Our findings, which provide new insights into the role of SR-B1 in cellular cholesterol homeostasis, suggest molecular links between SR-B1-dependent lipid sensing and cell cholesterol and lipid droplet dynamics.

5. Acknowledgements

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6. Conflict of interest

No conflict of interest declared.

7. Authors contributions

E.M., S.G. G.L., C.T., J-P.B., S.D., M.R. and V.C. designed and conducted the experiments and analyzed the data. M.R. and V.C. wrote the paper. F.S-P. coordinated lipidomic and proteomic analyses. C.M. and P.W.S. created SR-B1-Q445A mutant. A.L., M.R., F.S-P, C.M. and P.W.S. revised the manuscript. All authors reviewed the results and approved the final version of the manuscript.

8. References

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Figure legends

**Figure 1:** SR-B1-dependent lipid sensing modulates cholesterol processing and lipid droplet dynamics. (A) Pulse-chase experiments were conducted on Caco-2/TC7 cells stably transfected with WT SR-B1 or SR-B1-Q445A. Cells were cultured on semi-permeable filters in the presence of BODIPY-cholesterol-containing lipid micelles in the apical compartment of the filter for 1h (pulse), and then with unlabelled cholesterol-containing lipid micelles for one more hour (chase). At the end of the experiment, BODIPY-cholesterol (Bdp-Chol; purple) distribution was analyzed by confocal microscopy. Diamidino-2-phenylindole (DAPI; blue) is used for visualization of nuclei. Bar = 10µm. (B) Summary data (mean ± SEM) for Bdp-Chol fluorescence quantified at both apical and intracellular XY planes (5 fields per condition). ***p<0.001 versus WT SR-B1 expressing cells. (C) Transfected Caco-2/TC7 cells described in (A) were cultured for 1h with Bdp-Chol-containing lipid micelles. Neutral lipids of lipid droplets were stained with LD540 dye (purple). Co-localization of lipid droplets and Bdp-Chol (green) was analyzed by confocal microscopy. Enlarged views of the fields designated by white rectangles are displayed on the right. DAPI (blue) is used for nucleus staining. Bar = 10µm. (D) Cells expressing WT SR-B1 (WT) and SR-B1-Q445A (Q445A) were cultured in the presence (+) or absence (-) of PPM for 24h. The mRNA levels of PLIN2 were quantified by RT-PCR. Results are expressed as the ratio (mean ± SEM, n=6) of PLIN2 mRNA levels to cyclophilin mRNA level used as control. *p<0.05 and ***p<0.01 as compared to the same cell line in absence of PPM; #p<0.05, n=5. (E) Cells were cultured in the presence of [3H]-cholesterol-containing lipid micelles in the apical compartment for 48h. At the end of treatment (T0), cells were washed and then incubated for 2h with unlabelled lipid micelles devoid of cholesterol in the apical compartment. Radioactivity was then quantified in the apical (left panel) and basal (right panel) medium. Results are expressed as percentage of radioactivity counted in the corresponding medium to the radioactivity counted in cells at T0, mean ± SEM of 6 independent cultures. * p<0.05, ns: non-significant.

**Figure 2:** SR-B1 is involved in cholesterol trafficking from plasma membrane. (A) Pulse-chase experiments were conducted on Caco-2/TC7 cells stably transfected with WT SR-B1 or SR-B1-Q445A. Cells were cultured on semi-permeable filters and incubated in presence of BODIPY-cholesterol combined with methyl-β-cyclodextrin (Bdp-Chol/CD) in the apical compartment for 1
min (pulse), and then with culture medium without cholesterol for 5 min and 1 hour (chase). BODIPY-Cholesterol (purple) distribution was analyzed by confocal microscopy. DAPI (blue) is used for visualization of nucleus. Sub-apical XY planes are shown. Bar= 10µm. (B) Summary data (mean ± SEM) for Bdp-Chol fluorescence intensity quantified at sub-apical XY planes (5 fields per condition) are expressed as arbitrary units (a.u.). **p<0.01 and ***p<0.001 versus WT SR-B1 expressing cells. (C) Cells were incubated or not (ctrl) with U-18666A for 18h, then free cholesterol was visualized by filipin staining (purple). Rectangles and squares on the right of the XY planes represent enlargement of the selected area. Bar= 10µm (D) Filipin fluorescence intensity at plasma membrane reported to surface area was quantified in both cell lines with or without U18666A treatment. Results are expressed as arbitrary units (a.u.), mean ± SEM, (5 fields per condition). ####p<0.0001 as compared to WT SR-B1 in absence of U-18666A, ****p<0.0001 as compared to U-18666A-treated WT SR-B1 cells, ns: non-significant (E) Intracellular filipin dot numbers were quantified in both cell lines after U18666A treatment. Results are expressed as mean ± SEM (5 fields per condition). ****p<0.0001 as compared to WT SR-B1.

Figure 3: SR-B1-Q445A-expressing cells display changes in detergent-resistant membrane lipid composition as compared to WT SR-B1 cells. Detergent-resistant membranes were isolated from WT SR-B1- and SR-B1-Q44A-expressing cells. Concentrations of six lipid subclasses (A-J), of d18:0/16:0 SM (K) and d18:0/C20:0 ceramide (L) were determined by LC-MS/MS: cholesterol (chol), SM (sphingomyelin), PE (phosphatidylethanolamine), pPE (plasmalogen PE), PI (phosphoinositide) and PS (phosphoserine), PC (phosphatidylcholin), LPC (lysopPC), LPE (LysoPE) and ceramides. Results are expressed in µM as mean ± SEM from three independent experiments. *p<0.05,**p<0.01 and ***p<0.001 as compared to WT SR-B1-expressing cells.

Figure 4: Changes of total lipid and total phospholipid concentrations, as well as sphingomyelin to phospholipid ratio in detergent-resistant membranes of SR-B1-Q445A cells as compared to WT SR-B1 cells. Concentration in DRM fractions of (A) total lipid calculated by summing all lipid subclasses and (B) total phospholipid calculated by summing the PC, PE, PS, PI, LPC, pPE, LPE obtained from LC-MS/MS experiments. Results are expressed in µM as mean ± SEM from three independent experiments. *p<0.05 as compared to WT SR-B1-expressing cells. (C) The SM/PL ratio was calculated from concentrations of total SM and total PL for each condition. Results are expressed as arbitrary units (a.u.) ± SEM from three independent experiments, *p<0.05 as compared to WT SR-B1-expressing cells.

Figure 5: SR-B1-dependent lipid sensing is linked to cholesterol trafficking and modifications of lipid and protein composition of detergent-resistant membranes in intestinal epithelial cells. In
intestinal epithelial cells, SR-B1-dependent lipid sensing in response to lipid micelles requires plasma membrane cholesterol (Chol) interaction with the glutamine at position 445 (Q445) in the C-terminal transmembrane domain of SR-B1 [14, 15]. In this study, we identified that SR-B1-dependent lipid sensing is linked to intracellular cholesterol trafficking from plasma membrane and towards lipid droplet (LD) as well as cholesterol efflux. Lipidomic analysis of detergent-resistant membranes (DRM) revealed that the enrichment of d18:0/16:0 SM and 18:0/0:0 PE is associated with SR-B1-dependent lipid sensing. Moreover, we observed an increased abundance of 22 proteins and a decreased abundance of 14 proteins in DRM after lipid micelles supply, highlighting their potential role in SR-B1-dependent lipid signaling. SM: spingomyelin; PE: phosphatidylethanolamine; N: N-terminal region.

Supplemental Figure 1: Presence of SR-B1 at plasma membrane on unpermeabilized Caco-2/TC7 cells expressing WT SR-B1 and SR-B1-Q445A. Caco-2/TC7 cells transfected or not with WT SR-B1 or SR-B1-Q445A were detached with accutase and suspended in binding buffer (D-PBS 0.15% BSA). Unpermeabilized cell suspensions were incubated for 1h with anti-SR-B1 antibody (NB400-113) and then with a secondary antibody coupled to phycoerythrin. Cells were washed three times with fresh D-PBS and the fluorescence of each sample was evaluated. Histograms show phycoerythrin fluorescence intensity (PE-A) for negative sample (secondary antibody alone, red histogram), endogenous SR-B1 in untransfected cells (green histogram), WT SR-B1 expressing cells (yellow histogram) and SR-B1-Q445A expressing cells (blue histogram).

Supplemental Figure 2: Loss of SR-B1 expression results in lipid droplet accumulation. Caco-2/TC7 cells were transfected with control RNAi (siCTRL) or SR-B1 RNAi (siSR-B1). Differentiated cells were then incubated for 24h with PPM containing a fluorescent C12-fatty acid (FA568). (A) Lipid droplets were visualized with the fluorescent fatty acid (red) and nucleus with DAPI (white). Bar= 10µm. (B) After PPM treatment cells were lysed and western blot analyses were performed to determine the efficiency of SR-B1 RNAi. An antibody against annexin A2 (ANXA2) was used as control.

Supplemental Figure 3: Modification of lysophosphatidylcholine concentration in detergent-resistant membranes after PPM supply. WT SR-B1- and SR-B1-Q445A-expressing cells were cultured in absence (-) or in presence (+) of lipid micelles (PPM) for 15 min. Detergent-resistant membranes were isolated through a sucrose gradient and lysophosphatidylcholine (LPC) concentrations were determined by LC-MS/MC. Results are expressed in µM as mean ± SEM from three independent experiments. ***p<0.001 as compared to the condition without lipid micelles for each cell line.
**Fig. 2**

**A**

1' Bdp-Chol/CD pulse

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<th>5' chase</th>
<th>60' chase</th>
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<tr>
<td>SR-B1-Q445A</td>
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Bdp-Chol/DAPI

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**C**

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<tr>
<td>U-18666A</td>
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FILIPIN

**D**

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<td>1.5</td>
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<tr>
<td>+</td>
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FILIPIN fluorescence intensity at plasma membrane (a.u.)

**E**

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<tr>
<td>SR-B1-Q445A</td>
<td>ns</td>
</tr>
</tbody>
</table>
Fig. 3
Fig. 4

(A) Lipids

(B) PL

(C) SM/PL ratio

WT SR-B1 vs. SR-B1-Q445A
lipid micelles

In DRM:
- d18:0/16:0 SM
- 18:0/0:0 PE
- 22 proteins
- 14 proteins

SR-B1
Q445

DRM

N

efflux

Chol

plasma membrane

cytoplasm

Signaling

LD

Fig. 5
Suppl Fig. 1
Suppl Fig. 3
WT SR-B1- and SR-B1-Q445A-expressing cells were cultured in absence (-) or in presence of lipid micelles (PPM) for 15 min. Detergent-resistant membranes (DRM) were isolated through a sucrose gradient and lipid concentrations were determined by LC-MS/MC analysis. Results (mM) are expressed as mean ± SEM from three independent experiments. The species representing more than 10% of the total amount of species analyzed in their lipid subclass are underlined. Statistical changes (variation and p value) are in bold.
Table 2: List of proteins whose abundance in DRM changes after PPM supply in WT SR-B1- versus SR-B1-Q445A-expressing cells.

<table>
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<td>212.49 2.34E-05</td>
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<td>Q9UJU6</td>
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<td>cell adhesion/junction and cytoskeleton</td>
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<td>9.42 0.012</td>
<td>P40221</td>
<td>TCPZ, T-complex protein 1 subunit zeta</td>
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<td>LAP2B, Lamina-associated polypeptide 2 isoforms b/g</td>
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<td>Q9YSK8</td>
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<td>P37059</td>
<td>DHB2, Estradiol 17-beta-dehydrogenase</td>
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<td>FXR1, Fragile X mental retardation syndrome-related protein 1</td>
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<tr>
<td>2.54 0.01</td>
<td>P40879</td>
<td>S26A3, Chloride-anion exchanger</td>
<td>ion transport</td>
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<td>0.956</td>
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<tr>
<td>2.46 0.003</td>
<td>Q9H6A9</td>
<td>PCX3, Pecanex-like protein 3</td>
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<td>2.28 0.017</td>
<td>P36578</td>
<td>RL460S, ribosomal protein L4</td>
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<tr>
<td>2.17 0.015</td>
<td>Q9Y624</td>
<td>JAM1, Junctional adhesion molecule A</td>
<td>cell adhesion/junction and cytoskeleton</td>
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<td>0.545</td>
</tr>
<tr>
<td>2.13 3.22E-06</td>
<td>P35527</td>
<td>K1C9, Keratin, type I cytoskeletal 9</td>
<td>cell adhesion/junction and cytoskeleton</td>
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<td>0.0019</td>
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<tr>
<td>2.12 1.41E-06</td>
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<td>CALX, Calnexin</td>
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<tr>
<td>2.06 5.95E-05</td>
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<td>VATFV-type proton ATPase subunit F</td>
<td>ion transport</td>
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<td>0.0061</td>
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<tr>
<td>0.50 0.031</td>
<td>P13861</td>
<td>KAP2, cAMP-dependent protein kinase type II-alpha regulatory subunit</td>
<td>signal transduction</td>
<td>0.81</td>
<td>0.547</td>
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<tr>
<td>0.49 0.002</td>
<td>Q15363</td>
<td>TMED2, Transmembrane emp24 domain-containing protein 2</td>
<td>vesicular trafficking/autophagy</td>
<td>0.66</td>
<td>0.0197</td>
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<tr>
<td>0.47 0.0036</td>
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<td>VP9D1, VP9 domain-containing protein 1</td>
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<td>0.46 0.0067</td>
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<td>LDHB, L-lactate dehydrogenase B chain</td>
<td>oxidation-reduction</td>
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<td>IMB1, Importin subunit beta-1</td>
<td>protein interaction/processing</td>
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<tr>
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<td>Q9BV40</td>
<td>VAMP8, Vesicle-associated membrane protein 8</td>
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<td>Q14662</td>
<td>STX16, Syntaxin-16</td>
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<td>VAT1, Sycnatic vesicle amine transport</td>
<td>oxidation-reduction</td>
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<td>PHB2, Prohibitin-2</td>
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<td>0.32 0.001</td>
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<td>ICAM1, Intercellular adhesion molecule 1</td>
<td>cell adhesion/junction and cytoskeleton</td>
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<td>0.29 9.3E-05</td>
<td>Q5T0Z8</td>
<td>CF132, Uncharacterized protein C6orf132</td>
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<tr>
<td>0.25 0.002</td>
<td>P08574</td>
<td>CY1, Ubiquinol-Cytochrome-C Reductase Complex</td>
<td>oxidation-reduction</td>
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<tr>
<td>0.17 1.92E-06</td>
<td>P31943</td>
<td>HNRH1, Heterogeneous nuclear ribonucleoprotein H</td>
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<tr>
<td>0.14 0.024</td>
<td>Q60658</td>
<td>PDE8A, High affinity cAMP-specific and IBMX-insensitive 3',5'-cyclic phosphodiesterase 8A</td>
<td>signal transduction</td>
<td>0.92</td>
<td>0.950</td>
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WT SR-B1- and SR-B1-Q445A-expressing cells were cultured in absence (-) or in presence of lipid micelles (PPM) for 15 min. Detergent-resistant membranes (DRM) were isolated through sucrose gradient and protein enrichment was determined by tandem mass spectrometry analysis. Proteins with a fold change higher than 2 in DRM from WT SR-B1-expressing cells after PPM supply are listed. Fold change variation of these proteins in DRM from SR-B1-Q445A-expressing cells are reported. P values <0.05 are in bold. Proteins with similar direction change in their abundance in DRM after PPM in WT SR-B1- and SR-B1-Q445A-expressing cells are highlighted in green. Proteins with opposite variations induced by PPM in DRM in WT SR-B1- versus SR-B1-Q445A expressing cells are highlighted in red. Proteins unchanged in abundance in DRM after PPM supply in SR-B1-Q445A-expressing cells are indicated by a lack of highlighting.