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**Objective** — The purpose of this study was to determine whether lysosome trafficking and targeting of acid sphingomyelinase (ASMase) to this organelle contribute to the formation of lipid raft (LR) signaling platforms in the membrane of coronary arterial endothelial cells (CAECs).

**Methods and Results** — By measurement of fluorescent resonance energy transfer (FRET), it was found that in FasL-stimulated CAECs, membrane lamp1 (a lysosome marker protein) or Fas and GM1 (a LR marker) were trafficking together. Cofocal colocalization assay showed that ceramide was enriched in these LR platforms. Further studies demonstrated that these ceramide molecules in LR platforms were colocalized with ASMase, a ceramide producing enzyme. Fluorescence imaging of living CAECs loaded with lysosomal specific dyes demonstrated that lysosomes fused with membrane on FasL stimulation. In the presence of lysosome function inhibitors, bafilomycin (Baf) or glycyl-L-phenylalanine-β-naphthylamide (GPN), these FasL-induced changes were abolished. Moreover, this FasL-induced formation of LR platforms was also blocked in ECs transfected with siRNA of sortilin, an intracellular transporter for targeting of ASMase to lysosomes. Functionally, FasL-induced impairment of vasodilator response was reversed by lysosomal inhibitors or sortilin gene silencing.

**Conclusions** — Lysosomal trafficking and targeting of ASMase are importantly involved in LRs clustering in ECs membrane, leading to the formation of signaling platforms or signalosomes. (Arterioscler Thromb Vasc Biol. 2008;28:2056-2062)

**Key Words:** Fas ligand ■ lysosome ■ coronary circulation ■ vascular endothelium ■ sphingolipid
production of ceramide. It is also unclear where lysosomal ASMase is derived from and how these ASMases are trafficking through lysosomes to LR to form ceramide, ultimately leading to the formation of ceramide-enriched raft signaling platforms. The present study was designed to address these questions.

Materials and Methods

Cell Culture

The primary cultures of bovine CAECs were obtained as we described previously. All studies were performed by using CAECs of 2 to 4 passages.

Fluorescence Resonance Energy Transfer Analysis

CAECs were stained with TRITC-labeled CTXB and fluorescein isothiocyanate (FITC)-labeled antilamp1 (a lysosomal marker protein), anti-CD95 (Fas) antibodies as we described previously and then visualized by confocal microscope. To accurately observe the staining on the cell membrane, these cells were not permeabilized by excluding detergent in the washing and incubation buffer (PBS). Basically, the fusion of lysosomes or secretory vesicles into the plasma membrane results in exposure of lysosomal proteins onto the outer leaflet of the cell membrane when cells were stimulated. It is a general phenomenon that lysosomal proteins can be detected on the outer leaflet of the cell membrane when lysosome-membrane fusion happens. An acceptor bleaching protocol was used to measure the fluorescence resonance energy transfer (FRET) efficiency as described previously. The FRET efficiency was calculated through the following formula: 

\[ E = \frac{(\text{FITCpost} - \text{FITCpre})}{\text{FITCpost} \times 100\%}, \]

as described previously.

Confocal Analysis of LR Clusters and Its Colocalization With Ceramide or ASMase in CAECs

Detection of LR clusters were performed as we described previously. Doses of compounds used in these experiments were based on our previous studies and some preliminary data showing that they could effectively inhibit corresponding cell responses. Colocalization was analyzed by Image Pro Plus software.

RNA Interference of Sortilin

The siRNAs of sortilin gene were purchased from INVIROEN (CAT#HSS109429), which were confirmed to be effective in silencing sortilin gene in different cells by the company. The scrambled small RNA was confirmed as nonsilencing double-stranded RNA and used as control in the present study. Transfection of siRNA was performed using the QIAGEN TransMessenger transfection kit according to the manufacturer’s instructions.

RNA Isolation and Real-Time RT-PCR Analysis

The mRNA levels for Sortilin were analyzed by real-time quantitative PCR. The mRNA levels were presented as Tn as calculated by using 18S mRNA as the reference system as we presented previously.

Western Blot Analysis

The procedures for Western Blot analysis were described in details in our previous publications.

Quenching and Dequenching of FM1-43 in BCAECs

As described previously, BCAECs were loaded with 8 μmol/L FM1-43 in 1640 medium with 10% FBS for more than 2 hours at 37°C. After washed with PBS free medium, a low laser power (λ excitation=488 nm) was used to avoid possible fluorescent bleaching. For quenching experiments, after cells were loaded with 8 μmol/L FM1-43 for 2 hours, 1 mmol/L BPB was added in the extracellular medium. For dequenching experiments, cells were loaded simultaneously with 8 μmol/L FM1-43 and 1 mmol/L BPB for 2 hours.

Coronary Arterial Preparation and Microperfusion

Fresh bovine hearts were obtained from a local abattoir. Coronary arteries were prepared for microperfusion to evaluate the endotheli- um-dependent vasodilator response as we described previously. For some vessels, a novel ultrasound microbubble technology was used to transfet the intact endothelium with scrambled small RNA or sortilin-siRNA as we and others described previously.

Statistics

Data are presented as means±SE. Significant differences between and within multiple groups were examined using ANOVA for repeated measures, followed by Duncan multiple-range test. A Student t test was used to detect significant differences between 2 groups. P<0.05 was considered statistically significant.

Results

FasL-Induced FRET Between Lamp1 and LR Component

As shown in Figure 1A, FRET was detected by confocal microscopy between a fluorophore pair, FITC as donor and TRITC as acceptor, which shares the character to allow fluorescence resonance energy transfer. Acceptor (TRITC) bleaching protocol was applied to calculate the FRET efficiency. The left group of images shows a control cell costained with FITC-lamp1 and TRITC-CTXB that underwent an acceptor bleaching protocol. The left group of images from FasL-stimulated (10 ng/mL for 15 minutes) cells. B, Summarized results of detected FRET efficiency between CTXB and lamp1, Fas in CAECs with or without treatment of lysosome function inhibitors. Baf indicates bafilomycin; GPN, glycyl-L-phenylalanine-β-naphthylamide. n=6 batches of cells, *P<0.05 vs control; #P<0.05 vs vehicle+FasL group.

Figure 1. A, Representative images of FRET analysis between FITC-lamp1 and TRITC-CTXB in CAECs. The left group of images obtained from control CAECs and the right group of images from FasL-stimulated (10 ng/mL for 15 minutes) cells. B, Summarized results of detected FRET efficiency between CTXB and lamp1, Fas in CAECs with or without treatment of lysosome function inhibitors. Baf indicates bafilomycin; GPN, glycyl-L-phenylalanine-β-naphthylamide. n=6 batches of cells, *P<0.05 vs vehicle+FasL group.
the postbleaching image of FITC-lamp1 labeling. The overlaid images show the colocalization of both lamp1 and CTXB detected under different protocols. As shown in FRET image (blue in the bottom image), there was very low FRET detected under control condition. The right group of images shows a FasL-stimulated cell that underwent the same FRET protocol. In addition to detected patch formation of green fluorescence and colocalization of both molecules seen in the overlaid images (top panel) in response to FasL, a more intense FRET image (blue one the bottom) was detected in this FasL-treated CAEC, demonstrating that energy transfer occurs between a lysosomal marker-lamp1, and LR component-GM1 ganglioside. This close relationship between lysosome and membrane components may be due to a fusion of lysosome to the cell membrane under this condition.

**FRET Between LR Molecules**

As summarized in Figure 1B, the FRET efficiency between GM1 and lamp1 or Fas were significantly increased on FasL stimulation ($n=6$). Three pairs of molecules including GM1 versus lamp1 or Fas exhibited similar FRET efficiency with a maximum of 23.8% compared to control (5.2%). In the presence of lysosome inhibitors, Baf or GPN, FasL-induced increase in FRET efficiency was significantly attenuated for all three pairs of molecules.

**Lysosome-Derived Ceramide Production in LR Platforms**

As shown in Figure 2A, CAECs were stained by an Al488-labeled CTXB (as lipid raft marker) and an anticeramide antibody with Cy3 IIO antibody. Under control conditions, both Al488 and Cy3 stainings were diffuse and their colocalization in overlaid images was in tiny yellow spots. When the cells were stimulated by FasL, a number of large patches or spots with colocalization of both components (yellow in overlaid image) on the cell membrane were detected. However, when these CAECs were pretreated with lysosome function inhibitors, Baf or GPN, for 20 minutes, fluorescent patch formation in response to FasL was abolished as shown in the lower panels of images.

**Association of Ceramide With Local ASMase**

As shown in Figure 2B, CAECs stained with Cy3-conjugated ceramide and FITC-labeled ASMase were stimulated with FasL in the presence or absence of lysosome inhibitors, Baf or GPN. The overlay of the 2 images resulted in yellow areas, which indicated clustering or colocalization of ceramide with local ASMase. It was shown that FasL increased colocalization of ASMase and ceramide, which was blocked by both lysosome function inhibitors, Baf and GPN.

**Sortilin siRNA Blocked FasL-Induced Ceramide Production in LR Platforms**

As shown in supplemental Figure IIA through IIC (available online at http://atvb.ahajournals.org), by real-time PCR and Western blot analysis, the silencing effect of sortilin siRNA was verified. Both the mRNA and protein level can be knocked down by more than 80%. Figure 3A presents the results obtained from experiments with sortilin gene silencing, depicting the confocal microscopic analysis of Cy3-conjugated ceramide and Al488-CTXB labeled LR clusters in CAECs stimulated with FasL. Similar to Figure 3, under control condition both CTXB and ceramide stainings were diffuse. When these cells were stimulated by FasL, LR patches were detected with colocalization of ceramide. After these CAECs were transfected with sortilin siRNA, FasL-induced formation of LR platforms with ceramide colocalized was almost completely blocked. However, in scrambled sRNA transfected cells, FasL-induced formation of LR platforms remained unchanged.

**Sortilin siRNA Blocked FasL-Induced FRET in LR Platforms**

As summarized in Figure 3B, FasL significantly increased the FRET efficiency between GM1-gangliosides and lamp1 or CD95, in CAECs without transfection of sortilin siRNA (vehicle and scrambled sRNA-treated cells). In sortilin siRNA transfected CAECs, however, FasL-induced increase in FRET efficiency was significantly decreased to a level close to that obtained in control cells.
Ceramide From ASMase Pathway, but not De Novo Synthesis Pathway Contributed to the LR Platform Formation

As shown in Figure 4A, the ASMase inducer, phosphatidylinositol (PI, 5 μg/mL)\textsuperscript{41} significantly increased the FRET efficiency between FITC-lamp1 and TRITC-CTXB. However, the de novo ceramide synthesis inducer, Arsenic (As\textsubscript{2}O\textsubscript{3}, 1 μmol/L),\textsuperscript{42} had no significant effect on the FRET efficiency.

Caveolin-1 siRNA Failed to Block the Formation of LR Platforms

As shown in Figure 4B and 4C, after BCAECs were stimulated by FasL, some colocalization of caveolin-1 with lamp1 were observed, but not strong compared to staining with CTXB. When BCAECs were knocked down their caveolin-1 gene by a specific siRNA, the FasL-induced colocalization of lamp1 with CTXB on the membrane was still present.

FasL-Induced Lysosome Fusion Into GM1 Ganglioside-Enriched Microdomains

As shown in Figure 5A, representative images of control and FasL-stimulated BCAECs loaded with LysoTracker and Alexa488-labeled CTXB, respectively. Compared to control, FasL caused GM1 clustering on the cell membrane and colocalized with lysosomes.

FasL Caused Preloaded FM1-43 Quenched or Dequenched by BPB

As shown in Figure 5B, upper panel, without application of BPB, there are no significant changes in the FM1-43 fluorescence on FasL stimulation, because FM1-43 is not easily out of the cells. In quenching experiments, after BCAECs were loaded with 8 μmol/L FM1-43 for 2 hours, 1 mmol/L BPB were added in the extracellular solution. As shown in middle panel, FasL caused a decrease in FM1-43 fluorescence. In dequenching experiments, BCAECs were loaded simultaneously with 8 μmol/L FM1-43 and 1 mmol/L BPB for 2 hours. As shown in the lower panel, FasL caused an increase in the FM1-43 fluorescence. Supplemental Figure IV showed the colocalization of Lamp1 with AM1-43 after a prolonged incubation (2 hour).
Inhibition of Lysosome Function and Knocking Down Sortilin Gene Reversed FasL-Induced Impairment of Vasodilator Response

As shown in Figure 6, compared to control, FasL significantly attenuated the BK-induced vasodilator responses in small coronary arteries. Preincubation of the arteries with Baf or GPN significantly reversed FasL-induced impairment of the vasodilator response. Pretransfection of sortilin siRNA reversed this FasL-induced action. The transfection efficiency were confirmed by FITC-labeled small RNA (supplemental Figure II).

Discussion

Because FRET can only occur between molecules in a distance within a 10-nm range, increased FRET between lysosomal lamp1 and LR components GM1 should indicate that some lysosomes are indeed fused into cell membrane within LR clusters. This lysosomal trafficking or fusion to the cell membrane of CAECs was dependent on the functional integrity of lysosomes in that inhibition of lysosomal function by Baf or GPN blocked FRET production between lamp1 and GM1. Besides FRET between lamp1 and GM1, we also found that between LR components GM1 and other key molecules in LR platforms, there was strong FRET production in response to Fas activation. It seems that in LR platforms or clusters may constitute a membrane complex and thus transmit and amplify signals. Although there are reports about the lysosomal fusion to the cell plasma membrane for exocytosis of different molecules, to our knowledge the present results provide the first direct evidence indicating that lysosome fusion relates or contributes to the formation of LR signaling platforms, thereby participating in transmembrane signaling.

To address whether trafficking and fusion of lysosomes into the cell membrane are attributed to the formation of LR signaling platforms in CAECs, we determined the functionality of translocated or fused lysosomes in producing ceramide through a lysosome-containing enzyme ASMase in the LR platforms. Given the important role of ceramide in the formation of LR signaling platforms in various cells including CAECs, demonstration of local ceramide production would indicate the formation of LR clusters, because ceramide can spontaneously aggregate to generate large macromdomains or platforms in the cell membrane. By confocal microscopy, we found that FasL-induced LR platforms were indeed abundant of ceramide and that ceramide accumulation there was associated with lysosome function or fusion because inhibition of lysosomal function by Baf or GPN blocked ceramide production or accumulation in LR platforms. In additional groups of experiments, we demonstrated that local ceramide in LR platforms are colocalized with ASMase, indicating that ASMase could be the enzyme producing this sphingolipid for LR clustering. Consequently, in CAECs pretreated with lysosome function inhibitors, Baf or GPN, however, there were no ceramide and ASMase clusters detected even during FasL stimulation. Taken together, all these results indicate that ASMase may be trafficking and fused into the cell membrane along with

![Figure 5 and Figure 6 images](https://example.com/figures)
lysosomes, where ceramide is produced and LR signaling platforms or signalosome are formed in CAECs.

To further demonstrate that ASMase accumulation in the formation of LR signaling platforms is derived from lysosomes, we examined whether targeting of this enzyme to lysosomes is involved. Previous studies have shown that ASMase targeting to lysosomes is via a sortilin-mediated process.45 Sortilin belongs to a family of multiligand type-I receptors with homology to the yeast receptor Vps10p, which is able to transport lysosomal proteins from the Golgi body to lysosomes.45 In the present study, we used validated siRNA to knock down sortilin gene expression in CAECs and found that the ASMase translocation to the cell membrane and the formation of LR platforms were abolished in these CAECs with deficient gene expression of sortilin. By FRET analysis of lamp1 and LRs components, we demonstrated that silencing of sortilin gene almost completely blocked the generation of FRET between various molecular pairs in LR platforms, indicating that sortilin-mediated targeting of ASMase to lysosomes is of importance in LRs clustering. These results also further support the view that ASMase in LR platforms are derived from lysosomes. In addition, this lysosomal ASMase seems to be a driving force to activate LR clustering in CAECs, thereby leading to the formation of signaling platforms. The findings that the ASMase inducer, phosphatidylinositol (PI), but not the ceramide de novo synthesis inducer, As2O3, stimulated the LR platform formation indicate that only the ceramide generated through ASMase activation contributes to FasL-stimulated LR platform formation. As discussed above, lysosomal fusion process with externalization and activation of the acid sphingomyelinase occurs very rapidly after stimulation, which could be within 5 seconds.44,46 This suggests that ASMase activation is almost simultaneously happening with lysosome fusion. If that is the case, we believe two processes are very closely coupled, where any stimuli may trigger each other almost at the same time. In our experiments, it is possible that FasL stimulation leads to lysosome trafficking and at the same time activates ASMase, resulting in lipid raft clustering. At the fusion site, activated ASMase and ceramide were brought into the membrane when fusion occurs. However, when we treated cells with ASMase activator, lysosome trafficking also happened because activation is coupled with such molecular trafficking. This supports the view that two processes are coupled well.

Next, we addressed a question whether FasL induced LR clustering is associated with caveolar behavior. We found that after stimulated by FasL, there were indeed a few colocalization of caveolin-1 with lamp1 observed, but not very strong as those with CTXB. When caveolin-1 was silenced in these cells by its specific siRNA, although the colocalization of caveolin-1 and Lamp1 was abolished, a strong CTXB-lamp1 clustering was still detected. It appears that FasL may to some extent stimulate translocation of lysosomal proteins to caveolar area, but this FasL-induced action is much weaker than that occurred in noncaveolar raft areas.

In addition, we performed more experiments to directly confirm the fusion of lysosomes to plasma membrane on FasL stimulation in BCAECs. Compared to control cells, FasL caused CTXB clustering on the cell membrane, which was very close to the lysosomes stained by LysoTracker. This further confirmed the results obtained by using lamp1 staining. We also provided direct evidence that lysosome fusion can be recorded in these cells when they were stimulated by FasL. In these experiments, FM1-43, a proved lysosome specific fluorescence dye when incubated for prolonged period (>1 hour),35 was used to load lysosomes. This dye can be reversibly quenched by bromide phenol blue (BPB),36 which is easier to enter or come out of the lysosomes than FM1-43. In quenching experiments, FasL was found to cause a decrease in the FM1-43 fluorescence. This was attributable to the FasL-stimulated lysosome fusion with the plasma membrane, allowing BPB to enter the lysosomes to quench FM1-43 fluorescence. In dequenching experiments, in contrast, FasL caused an increase in the FM1-43 fluorescence when lysosomes were fused to cell membrane since BPB was moved out of cells. Together, all these direct or indirect evidences strongly suggest that the fusion of lysosomes into the cell plasma membrane occurs in these BCAECs on FasL stimulation.

In our previous studies, it has been demonstrated that FasL, at the dose used in the present study significantly impaired the bradykinin-induced vasodilator response.22,25,29 The present study further found that inhibition of the lysosome function by two inhibitors or blockade of the ASMase targeting to lysosome by silencing sortilin expression with its specific siRNA significantly abolished the damaging effects of FasL. These results provide direct evidence that inhibition of lysosome function or prevention of targeting of ASMase to lysosomes may block the formation of LR platforms and thereby produce beneficial action in protecting the arterial endothelium from detrimental effects of FasL. However, it should be noted that in the present study we found that FasL-stimulated translocation of lysosomal markers is also located where the blebbing phenomenon normally took place during apoptosis (supplemental Figure III). It could be a very early functional change or signaling event during death receptors activation toward cell apoptosis. Therefore, further studies still need to be done to address some questions whether this translocation is specific for transmembrane signaling or whether this translocation is also related to apoptosis, despite that endothelial cells are relatively resistant to apoptotic stimuli.

In summary, the present study demonstrates a critical role of lysosomal trafficking and targeting of ASMase in the formation of LR signaling platforms in CAECs, which was dependent on ceramide production via lysosomal ASMase translocated into the cell membrane via a direct fusion of lysosomes to the plasma membrane. In addition, ASMase targeted to lysosomes through sortilin was the resource of this enzyme for LR signaling, and ASMase, ceramide, and GM1 may constitute a signalosome to carry out transmembrane signal transduction during activation of Fas in CAECs.

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**Supplementary Figure Legends**

**SFig. 1.** A. Intracellular lamp-1 staining.

**SFig. 2.** Efficiency of sortilin gene silencing in CAECs by its siRNA. A: Real time RT-PCR detection of mRNA; B: A gel document of Western blot analysis; C: Densitometric analysis of gel documents via Western blotting. Values are means ± SE; n=4, * P<0.01 vs. control; # P<0.01 vs. scrambled sRNA transfected cells. D. Transfection of the intact endothelium of small bovine coronary arteries with scrambled small RNA labeled with fluorescein (FITC-siRNA). Control: the endothelium transfected with non-fluorescence scrambled small RNA. EC(-): the endothelium was carefully removed after FITC-siRNA transfection.

**SFig. 3.** A. Dynamic DIC images of a BCAEC stimulated by 10 ng/ml Fas L. Note that the location of arrowheads indicates blebs formation on the cell membrane. B. Fluorescence resonance energy transfer (FRET) between FITC-labeled lamp1 antibody and a lysosome-unrelated external protein, TRITC-conjugated anti-IgG secondary antibody in the absence (A) or presence (B) of lysosome function inhibitor, GPN. Note that only a portion of the cell was bleached and the increase in the donor fluorescence intensity only occurred in bleached area, where the intensity of donor fluorescence in non-bleached area remained almost unchanged. When lysosomal function inhibitor, 200 µM GPN was applied to pretreat the cells, this FRET can not be blocked.
Supplemental Materials About Selection of Baf and GPN

It has been reported that Baf is a macrolide antibiotic that specifically inhibits vacuolar H\(^+\)-ATPase (V-H\(^+\)-ATPase), resulting in a failure to pump protons into lysosomal lumen for acidification. Decreased acidification of lysosomal lumen has been shown to lead to loss of many lysosome functions. GPN is another mechanistically different inhibitor of lysosome function. GPN as a substrate of lysosomal cathepsin C may be hydrolysed, and its hydrolysis products could be accumulated in the vesicle, resulting in a reversible osmotic swelling and in this way interfering with lysosomal functions. It has been reported that the V-ATPase inhibitors concanamycin A and bafilomycin A lead to Golgi swelling in tobacco BY-2 cells (Protoplasma. 2004 Dec; 224(3-4):255-260). Since ASMase functions in an acidic environment, it is presumed that Baf will interfere with ASMase function. There is no report demonstrating that GPN could interact with Golgi trafficking or ASM function. This possible non-specific action on the Golgi activity is the rationale for us to use two mechanistically different tool compounds for inhibition of lysosome function.

Supplemental Protocol for Caveolin-1 Small RNA Interference

Based upon an established protocol of Caveolin-1 small RNA interference in bovine endothelial cells(Gonzalez E, Nagiel A, Lin AJ, Golan DE, Michel T. Small interfering RNA-mediated down-regulation of caveolin-1 differentially modulates signaling
pathways in endothelial cells. J Biol Chem. 2004 Sep 24;279(39):40659-69.), we used a caveolin-1 siRNA duplex corresponding to bases 223–241 from the open reading frame of the bovine caveolin-1 mRNA: 5′-CCA GAA GGA ACA CAC AGU U-dTdT-3′. Small interfering RNA duplex oligonucleotides were purchased from QIAGEN, Inc. (Valencia, CA). Transfecting BAEC at 50–70% confluence maintained in RPMI-1640/10% fetal bovine serum; transfections with siRNA (20 nM) used siLentFectTM-Lipid (2:1, Bio-Rad laboratories, Hercules CA), following protocols provided by the manufacturer. Experiments were conducted 48 h after transfection. After labeling of FITC-Lamp-1, the labeling of caveolin-1 was followed. 0.3% Triton X-100 was added to each solution to permeabilize cells. Rabbit anti-caveolin-1 Antibody (1/200 dilution; BD Pharmingen) were detected with Texas red-conjugated second Antibody (1/500 dilution).