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Plasma membrane micro domains regulate TACE-dependent TNFR1 shedding in human endothelial cells

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Abstract

Upon stimulation by histamine, human vascular endothelial cells (EC) shed a soluble form of TNFR1 (sTNFR1) that binds up free TNF, dampening the inflammatory response. Shedding occurs through proteolytic cleavage of plasma membrane-expressed TNFR1 catalyzed by TNF-α converting enzyme (TACE). Surface expressed TNFR1 on EC is largely sequestered into specific plasma membrane micro domains, the lipid rafts/caveolae. The purpose of this study was to determine the role of these domains in TACE-mediated TNFR1 shedding in response to histamine. Human Umbilical Vein Endothelial Cells (HUVEC)-derived EA.hy926 cells respond to histamine via H1 receptors to shed TNFR1. Both depletion of cholesterol by methyl-β-cyclodextrin (MBCD) and siRNA knockdown of the scaffolding protein caveolin-1 (cav-1), treatments that disrupt caveolae, reduce histamine-induced shedding of membrane-bound TNFR1. Moreover, immunoblotting of discontinuous sucrose gradient fractions show that TACE, like TNFR1, is present within low density membrane fractions, concentrated within caveolae, in unstimulated EA.hy926 endothelial cells and co-immunoprecipitates with cav-1. Silencing of cav-1 reduces the levels of both TACE and TNFR1 protein and displaces TACE, from low density membrane fractions where TNFR1 remains. In summary, we show that endothelial lipid rafts/caveolae co-localize TACE to surface expressed TNFR1, promoting efficient shedding of sTNFR1 in response to histamine.

Keywords

TACE; Caveolin-1; Lipid rafts; TNF-receptor; Histamine; Inflammation; Endothelium

Introduction

Binding of TNF to a variety of cell types activates signaling pathways that regulate several different processes of medical significance such as apoptosis, inflammation, immunity and metabolism and has been implicated in the pathogenesis of several diseases [1,2]. Although two different TNF receptors (TNFRs) have been identified [3], the majority of endothelial cells (EC) responses appear to be activated through TNFRI (p55-TNFR, CD120a) [4]. TNFRs are ubiquitously expressed in most cultured cell types, but expression is highly

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regulated in tissues. For example, in human kidney and heart, TNFR1 is largely confined to vascular EC. Under specific circumstances, the shedding of membrane TNFR1 (mTNFR1) act by reducing the amount of mTNFR1 and produces a soluble 27-30 kD form of the receptor (sTNFR1) which compete with mTNFR1 to reduce cell sensitivity to TNF [5,6]. Interestingly, both in human umbilical vein (HUV)EC and in the EAhy.926 endothelial cell line, the Golgi apparatus plays an important role in this scenario, functioning as a intracellular reservoir of TNFR1 from which it can be translocated to the plasma membrane and shed in response to stimuli [6-8] such as histamine. The generation of sTNFR1 is triggered by TNF-α-converting enzyme (TACE, also known as ADAM-17 or CD156b) [9-11], a member of the ADAMs (A Disintegrin And Metalloprotease) family of proteases. TACE is widely expressed in a variety of tissues and shows very little sequence similarity to other ADAMs family members [12]. The mechanism of ectodomain shedding is not unique to TNFR1 and it has been reported that a wide range of inflammatory cell surface proteins operate as soluble form [13]. Regarding TACE, it has been recently proposed that the transmembrane domain (TM) is fundamental for efficient cleavage and its amino-acidic sequence can specifically regulate the shedding of a subset of substrate [14]. In other words the mechanism of ectodomain shedding represents an additional post-translational regulatory step which controls the expression of surface molecules. Specifically, the importance of TNFR1 shedding has been demonstrated in patients affected by unexplained episodes of fever and severe localized inflammation who show impaired cleavage of membrane TNFR1 with consequent reduced shedding of the antagonistic soluble receptor [15,16]. Notably, the plasma membrane of mammalian cells contains small lipid microdomains enriched for cholesterol and glycosphingolipids called lipid rafts [17]. In certain cell types, such as EC, fibroblasts and muscle cells, lipid rafts can be clustered into 50-100 nm flask-shaped invaginations called caveolae [18,19]. These organelles may comprise up to 50% of the plasma membrane of ECs in situ [20-22] but are rapidly lost when cells are placed in culture [23], allowing lipid rafts to be dispersed within the plasma membrane. We previously demonstrated that caveolae are retained in the EC-derived cell line EA.hy926 facilitating the study of these organelles in vitro [24]. Caveolae are organized by the caveolin (cav) family of small cholesterol-binding scaffolding proteins, including cav-1, cav-2, and the muscle-specific cav-3 [25-27] that contain a conserved structural motif [28]. Caveolae were initially implicated in endocytosis and transcytosis of macromolecules across the EC lining of blood vessels [29], and it has been appreciated that these organelles contribute to cell signaling by clustering together various cell surface receptors [28], facilitating receptor cross talk [28,30]. Caveolae also play a role in the endocytosis of membrane proteins independently of clathrin-coated pits [31-33]. We previously demonstrated that TNFR1 on the plasma membrane localizes to caveolae in EA.hy926 cells and that pharmacological extraction of cholesterol, which reduces the number of caveolae, blocks ligand-induced TNFR1 internalization [24]. More recently, we found that internalization appears to be independent of cav-1 as it is unaffected by siRNA knockdown of this protein, a treatment that reduces caveolar number but leaves lipid rafts intact [8]. Reduction in cav-1 expression also reduced total TNFR1 protein expression, although the effects on plasma membrane levels were variable. It has been reported that the sheddase activity of TACE is concentrated in lipid rafts and inhibitors of this enzyme increase TNF and TNFRs in lipid rafts [34]. Furthermore, other studies demonstrated that lipid rafts play an important role in the shedding of a variety of different molecules [35,36]. In the present study we extended these findings and propose a key function of cav-1 in both the retaining of TACE within the caveolar network and the shedding of TNFR1 given that the disruption of caveolae inhibits histamine and H1R-mediated sTNFR1 release.
Materials and Methods

Evaluation of TNFR1 surface expression by cytofluorimetric analysis

Replicate cultures of EA.hy926 cells were subjected to specific treatment in six-well plates. For evaluating the involvement of proteing-G to both the shedding and surface expression of TNFR1, cells were pre-incubated in the presence of the indicated doses of Pertussis toxin (Calbiochem-Novabiochem, Nottinghamshire, UK, Ptx for 1h at 37°C. Cells were harvested with 2.5 g/l trypsin 0.2 g/l EDTA in Hanks’ Balanced Salt Solution (HBSS) without phenol red, at 37°C and stained with anti-human CD120a (1 μg/10⁶ cells) or with isotype control antibodies, respectively, for 1 hour at 4°C. Cells were washed once by centrifugation at 1000 RPM in HBSS containing 1% BSA and incubated with streptavidin phycoerythrin for 30 minutes at 4°C. After three additional washes cells were resuspended in HBSS without BSA and acquired using CyAn™ ADP Analyzer 9 color (Beckman Coulter). Fluorescence of 20 × 10³ cells/sample was acquired and analyzed using Summit v4.3 software.

Ca²⁺ imaging

EA.hy926 were cultivated onto 35-mm dishes in medium containing 3.5 μM fura-2-AM (Invitrogen Corporation, CA) for 1 h at 37°C, and then rinsed with Krebs-Henseleit-Hepes (KHH) buffer (140 mM Na⁺, 5.3 mM K⁺, 132.4 mM Cl−, 0.98 mM PO₄²⁻, 1.25 mM Ca²⁺, 0.81 mM Mg²⁺, 5.5 mM glucose and 20 mM Hepes) supplemented with 0.2% fatty acid free BSA or with Hanks’ balanced salt solution (HBSS). Each dish was placed into a culture chamber at 37°C on the stage of an inverted fluorescence microscope (Nikon, TE2000E), connected to a cooled CCD camera (512B Cascade, Roper Scientific, Tucson, AZ). Samples were illuminated alternately at 340 and 380 nm using a random access monochromator (Photon Technology International, NJ) and emission was detected using a 510 nm emission filter. Images were acquired (1 ratio image/s) using Metafluor software (Universal Imaging Corporation, Downingtown, PA). Calibration was obtained at the end of each experiment by maximally increasing intracellular Ca²⁺-dependent fura-2-AM fluorescence with 5 μM ionomycin, followed by recording minimal fluorescence in Ca²⁺-free medium. [Ca²⁺]ᵢ was calculated as previously described [37].

Quantitative RT-PCR analysis

Total RNA was extracted with the TRIZOL Reagent (Invitrogen), according to the manufacturer’s instructions. Briefly, 3 μg of RNA was converted into single-stranded DNA by a standard 20μl RT reaction with the TaqMan RT Reagent (Applied Biosystems, CA). Real time quantitative RT-PCR was performed on a 7500 Real-Time PCR System (Applied Biosystems). Briefly, the cDNA generated from the reverse transcription reactions was amplified by PCR with the SYBR Green JumpStart Taq ReadyMix (Sigma) in a total volume of 25 μl according to the manufacturer’s instructions. The primers used were as follows: TNFR1, 5 ’-AATGCCGAAAGGAATGGGTAGGTCAGG-3’ and 5’- TGCACACGGGTGTCTGTCTCTCTCTCT-3’; TACE, 5’- AGTGCAGTGACAGGAACAGTCCTT-3’ and 5’- GGACACGCCTTTGCAAGTAGCATT-3’; β-Actin, 5’- TGCACACACCTTGTGCTGCTTCT-3’; β-Actin, 5’- TGCACACACCTTGTGCTGCTTCTC-3’ and 5’- CAGCCTTGATAGCATTACAT-3’; β-Actin, 5’- TGCACACACCTTGTGCTGCTTCTC-3’. The level of messenger analyzed was expressed as relative fold change vs. the β-actin messenger RNA and analyzed by means of the 7500 System Software (Applied Biosystems).

Design and transfection of cav-1 siRNA duplexes

Small interfering RNA (siRNA) duplex oligonucleotides against the coding sequence of human cav-1 cDNA (NM_001753) were synthesized and purchased by Integrated DNA
Technologies (Coralville, IA). We selected two target sequences as follows: 5’-
ACCAGAAGGGACACACAGUdTdT-3’ (sense-#538) and 5’-
ACUGUGACGAAAUACUGGUdTdT-3’ (sense #542). Transfection of 80 pmol siRNA in
EA.hy926 was carried out by using Oligofectamine (Invitrogen) (0.3%, v/v), according to
the manufacturer’s instructions. Fresh medium was added 5 h after transfection and
experiments were conducted for 72 h. Non-targeting control siRNA-A (Santa Cruz, CA) was
used as control.

Isolation of caveolae-enriched membranes
Purification of caveolae-enriched membrane fractions was performed as previously
described [24,38] with minor modifications. In brief, cells were washed twice with ice-cold
Dulbecco’s phosphate-buffered saline and scraped with 25 mM MES hydrate buffer pH 6.5
containing 0.15M NaCl, 5 mM EDTA and 0.5% Triton X-100 (MBS) with protease
inhibitors (10μg/ml aprotinin, 10μg/ml leupeptin, 1mM sodium orthovanadate, 10mM NaF,
1mM Pefabloc) and left 20 minutes on ice. Three wells of a 6 well plate were pooled
together and the resultant lysate was subjected to 15 strokes in a Dounce homogenizer and
centrifuged for 10 minutes at 2000 rpm at 4°C to remove nuclei. Clarified post nuclear
supernatants were combined with 90% (w/v) sucrose prepared in MBS, transferred to the
bottom of a Beckman 2 ml ultracentrifuge tubes and overlaid gently with 1 ml of 35% and
0.6 ml of 5% sucrose respectively. The resulting 5-40% discontinuous sucrose gradients
were centrifuged 18 hours at 35.000 rpm at 4°C. After centrifugation, 10 fractions of 200
μl each were harvested from the top to the bottom of the gradients and analyzed either by SDS-
PAGE and immunoblotting.

Immunoblotting
For immunoblotting experiments, scrape-harvested cells were extracted in 150 μl of lysis
buffer (20 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM Na₂EDTA, 1 mM EGTA, 1% Triton,
2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM Na₃VO₄, 1 μg/ml
leupeptin, 2% SDS and 1 mM Pefabloc) for 15 minutes on ice. Unbroken cells and debris
were spun down by centrifugation at 2000 rpm for 10 min at 4°C. 20 μg of cell lysate was
fractionated by SDS-PAGE, transferred to a nitrocellulose membrane (Trans-Blot Transfer
Medium, Bio-Rad, Hercules, CA) and subjected to immunoblotting with primary antibody
followed by incubation with HRP-conjugated secondary antibody. Detection of the bound
antibody was performed by enhanced chemiluminescence (Pierce Biotechnology, Rockford,
IL). For immunoblotting analysis of sTNFR1 cells were treated with indicated chemicals
diluted in 1.6 ml of OPTI-MEM. Culture media were collected and concentrated about 50
times using Vivaspin ultrafiltration spin columns (Sartorius Stedim Biotech GmbH,
Goettingen, Germany), and samples were analyzed by SDS-PAGE electrophoresis and
immunoblotting. Intensities of the bands for specific proteins of interest were quantified and
normalized to the intensity of the band for β-actin with Scion Image application software
(Scion Corporation, Frederick, MD).

Results
Modulation of surface expression of TNFR1 by histamine H1 receptor
In HUVEC histamine causes TACE-mediated shedding of mTNFR1, which in turn
desensitizes TNFR1-mediated responses [6]. Although HUVEC recapitulate many features
of EC in situ, they rapidly lose their caveolae in culture [23], limiting the investigation of
these organelles in vitro. Interestingly, we have recently reported that the EA.hy926
endothelial cell line [39] not only retains caveolae but also expresses higher surface TNFR1
level than HUVEC, which renders this cell line an appropriate model for studying the role of
caveolae in vitro. Moreover, of the four known histamine receptors identified [40],

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EA.hy926 appear to express only H1R and H2R [41]. To evaluate which histamine receptor was involved in TNFR1 cleavage we performed immunoblotting analysis on culture medium collected by both unstimulated- and histamine-stimulated cells in the presence or absence of H1R antagonist mepyramine. Our data suggest that the shedding of TNFR1 was prevented by preincubation of cells with mepyramine, suggesting that H1R was primarily involved in the cleavage of TNFR1. As expected stimulation of cells with 2-3-trifluoromethylphenyl-histamine (TMPH), the specific H1R agonist, did also induce receptor shedding (Fig 1A). FACS analysis confirmed that both histamine and TMPH stimulation reduced the expression of TNFR1 on the plasma membrane (Fig. 1B), and that this effect was prevented by mepyramine (Fig. 1C), further supporting the involvement of only H1R in this mechanism. TACE has been reported as the major ADAM responsible for the cleavage of membrane TNFR1 [9-11]. Therefore, we next employed the TACE inhibitor TAPI-0 to confirm the contribution of TACE to H1R-mediated TNFR1 shedding as reported in (Fig. 1D). We conclude that EA.hy926 cells, like HUVEC, respond to histamine through H1R to induce TACE-mediated shedding of TNFR1 from the plasma membrane generating a soluble form of the receptor (sTNFR1).

**H2R counteracts constitutive shedding of TNFR1**

Since EA.hy926 cells also express H2R which in addition to H1R functions as natural receptor of histamine, we wondered whether this histamine receptor was also involved in the ectodomain shedding of TNFR1. To this purpose, we employed amthamine (AMTHA), the specific H2R agonist. We found that incubation of cells with AMTHA alone did not affect the shedding of TNFR1 but rather appeared to reduce its constitutive release as sTNFR1 (Fig.2A). This results suggested that signaling activated by H2R, possibly through the accumulation of cAMP, may perhaps counteract the release of sTNFR1. In fact, unlike H1R, which has been showed to induce a rise of cytosolic Ca^{2+} concentration [42-44], H2R typically appears to function by increasing adenylate cyclase (AC) activity and accumulation of cAMP [40]. To test this hypothesis we stimulated cells with forskolin, a receptor-independent activator of AC and evaluated the shedding of TNFR1. Forskolin appeared to diminish both constitutive and TMPH-stimulated release of sTNFR1 (Fig. 2B), without significant changes of surface exposed TNFR1 (Fig. 2C). In addition, we investigated the possible involvement of the inhibitory G_i protein by employing pertussis toxin. Our data showed that pre-incubation of cells with Ptx slightly reduced the constitutive release of sTNFR1 (Fig. 2D) while showing a non significant change of TNFR1 surface expression (Fig. 2E), confirming our findings obtained with forskolin. Moreover, histamine-induced shedding of TNFR1 was not affected by Ptx, indicating that histamine H1 and H2 receptors are not G_i-protein linked. These findings suggest that the fate of TNFR1 shedding induced by TACE in EA.hy926 cells may be further regulated by a functional interplay between signaling activated by both histamine H1 and H2 receptor.

**Effects of disrupting caveolae on histamine-induced shedding of TNFR1**

We have previously shown that TNFR1 and cav-1 associate with caveolae in EA.hy926 cells and that this association as well as ligand-induced TNFR1 internalization is disrupted by extraction of cholesterol with MbCD [24]. MbCD functions by extracting cholesterol from the plasma membrane which results fundamental for the correct formation of lipid rafts micro domains. Therefore we wondered whether MbCD could affect TMPH-induced release of sTNFR1. ELISA analysis showed that pre-treatment of cells with MbCD decreased TMPH-induced release of sTNFR1 (Fig. 3A), and FACS analysis confirmed that cholesterol depletion did also reduce the capability of TMPH to decrease the expression of exposed TNFR1 (Fig 3B). Nevertheless, to determine whether the effect of cholesterol extraction was on a proximal signaling event, we examined H1R-dependent calcium responses [40] in the presence of MbCD and found that cholesterol depletion did not interfere with TMPH-
induced calcium mobilization (Fig. 3C), suggesting that action of MbCD may involve downstream events, possibly affecting the process of shedding itself.

**Caveolin-1 down regulation inhibits TMPH-induced TNFR1 shedding**

In certain cell types the expression of the scaffolding protein cav-1 induces the clustering of lipid rafts into vesicle-like organelles called caveolae. To this regard, we investigated the contribution of cav-1 to TNFR1 shedding by evaluating the behavior of cav-1 silenced cells in response to histamine and TMPH. We designed two different siRNA duplex oligonucleotides targeting the coding sequence of human cav-1 cDNA and tested their silencing efficiency by immunoblotting (Fig 4A). Interestingly, we found that reducing cav-1 expression by RNAi did affect both histamine- and TMPH-induced release of sTNFR1 (Fig 4B), suggesting that cav-1 may play an important role in the regulation of TACE activity. In addition, similar to what we had seen with MbCD, cav-1 knock down did not impair TMPH-induced calcium mobilization (Fig 4C). These data support the hypothesis that the mechanism of calcium release induced by TMPH and TNFR1 shedding are functionally uncoupled.

**Sub cellular localization of H1R and TACE in EA.hy926 cells**

Despite the limitations in the use of cav-1 silencing to quantify shedding, it is still useful as an approach to study receptor localization since it disrupts caveolae while dispersing rather than causing loss of low density lipid rafts. Some G protein coupled receptors, the category to which H1R belongs, localize to lipid rafts [45,46] and TACE has also been localized to lipid rafts in some cell types [34]. Disruption of lipid rafts and caveolae by pharmacological depletion of cholesterol has been also reported to affect the release of ADAMs substrates [47-49]. Therefore, we next evaluated the distribution of both H1R and TACE by means of sucrose gradient fractionation in EC. Under our experimental procedures, H1R was mainly excluded from low density membranes in either case (Fig. 5A). By constrast, TACE was extensively detected along the sucrose gradient obtained from cells transfected with the control dsRNA duplex (Fig. 5B, upper panel). In contrast, cav-1 silencing reduced TACE expression in high density membranes and its localization within the low density membranes (Fig 5B, lower panels). Moreover, we found that immunoprecipitation of cav-1 followed by immunoblotting with anti TACE antibody revealed the interaction between these two proteins that was also unchanged after histamine treatment (Fig. 5C). Although we did not investigate the origin of this association this result suggest that cav-1 expression may be significant for sequestrating TACE within the caveolar network, placing the enzyme in proximity to its substrate. Interestingly, we showed that cav-1 silencing reduced TNFR1 protein but not mRNA expression in EA.hy926 cells, although it did not cause TNFR1 to dissociate from (now dispersed) lipid rafts [8]. We therefore evaluated if cav-1 silencing affected TACE protein expression. Besides confirming the overall reduction of TNFR1 induced by cav-1 down regulation [8] (Fig. 6A), we found that TACE protein expression, but not its mRNA level, was also decreased in cav-1 knock down cells (Fig. 6B), indicating that the level of TNFR1 shedding can be regulated by the reduction of both TACE and TNFR1.

**Discussion**

ADAM family of metalloproteases has been linked to diverse biological processes leading different groups to explore novel potential regulators which can activate or inhibit these proteins both under physiological and pathological conditions [12,50,51]. Since TNF and TNF receptors are targets of TACE/ADAM17, understanding the mechanism which regulate TACE activity may represents an important target for modulating the inflammatory response. In the present study we extended our previous works on the relationship between
TNFR1 and caveolae [8,24] and reported the contribution of cav-1 to TACE-mediated shedding of the membrane-bound TNFR1. Histamine evokes a variety of effects in different systems such as nervous, endocrine and cardiovascular by binding to four different receptors. However, the availability of specific agonists and antagonists [52] represents an important tool for studying the contribution of histamine receptors in different cell types. Notably, it has been reported that primary human ECs, such as HUVEC, rapidly loose caveolae after few cell doubling in culture [23], limiting the investigation of caveolae in primary EC. Therefore, we employed the permanent cell line EA.hy926 established by fusing primary HUVEC with a thioguanine-resistant clone of A549 which shows functions characteristic of human vascular endothelium [39]. Unlike cultured HUVEC, EA.hy926 retain an extensive caveolar network and express higher level of surface TNFR1 compared to primary cells, which make them an appropriate model to explore the contribution of caveolae to TNFR1 signaling in vitro. To this regard, we found that similarly to what had been reported in HUVEC [6], histamine also stimulated TACE-mediated TNFR1 shedding in EA.hy926. Moreover, by employing the specific H1R antagonist mepyramine we show that this mechanism was specifically mediated through the recruitment of H1 receptor and the contribution of TACE was confirmed by pretreatment of cells with TACE inhibitor TAPI-0. Importantly, H1 and H2 receptors usually coexist in blood vessels [53,54] and are also co-expressed in EA.hy926 [41], allowing us to investigate the contribution of H2R to the shedding of TNFR1. We found that stimulation of histamine H2 receptor with AMTHA not only failed to induce TNFR1 shedding, but also appeared to diminish the constitutive release of the cleaved TNFR1. Given that H2R recruitment was reported to induce cAMP accumulation via Gs-protein activation [55,56], we hypothesized that the second messenger may function by counteracting H1R-induced shedding of membrane TNFR1. Our data, in fact, demonstrate that forskolin-mediated activation of AC reduced TMPH-induced sTNFR1 accumulation, which suggests a contribution of cAMP on TACE-mediated receptor shedding. Forskolin is generally used to induce intracellular cAMP accumulation by directly activating AC bypassing H2R recruitment. Moreover, additional experiments performed in the presence of Ptx, which catalyzes ADP-ribosylation at cysteine residue of G_i, blocking the AC inhibitory pathway, indicate that histamine H1 and H2 receptors are not G_i-protein linked in EA.hy926 cells. Therefore, our data indicate the involvement of cAMP in the inhibition of TNFR1 shedding, although other signaling might affect TACE activity in response to AMTHA. In fact, it has been reported that H2R stimulation also affect other second messengers, such as arachidonic acid, and phospholipase A2 that may in turn disturb TACE-mediated shedding [56]. Although the mechanism of H2R-mediated TACE inhibition remains unknown it is possible that the mechanism of TNFR1 shedding might be affected by opposing responses activated by H1R and H2R stimulation.

A key finding of this report is that both cholesterol depletion with MbCD and knock down of cav-1 by RNAi markedly reduced H1R-dependent TNFR1 shedding, and neither treatments reduced calcium mobilization caused by H1R signaling. This suggest that the mechanism of shedding itself may be independent of calcium mobilization induced by H1R. Notably, cav-1 silencing reduces the expression level of TNFR1 [8], highlighting the importance of cav-1 for receptor stability. On the other hands, cholesterol depletion which pharmacologically disrupt caveolea/lipid rafts would only induce surface redistribution of TNFR1 which is also sufficient to affect the shedding mechanism itself, possibly by destabilizing specific protein interactions. To this regard we demonstrated that short MbCD treatment prevented co-localization of cav-1 and TNFR1 [24]. We have also showed that intrinsic signaling by TNFR1 is not inhibited by these treatments [8], suggesting that localization of the receptor to caveolea may be important for interactions with other caveolar proteins. Since TNFR1 shedding is mediated by TACE which has been reported to localize to lipid rafts [34], structures that are concentrated within caveolea, we hypothesized that caveolea/lipid rafts may be important for regulating the compartmentalization of TACE and
its substrates. By employing a sucrose gradient fractionation technique we found that TACE localized to cholesterol- (e.g. lipid rafts micro domains) and caveolin-enriched domains, which in EA.hy926 EC are largely confined to caveolae. In contrast, H1R appeared to be restricted to high density membranes that does not contains lipid raft domains. These findings allow to exclude a direct interaction between H1R and TACE. Interestingly, sucrose gradient fractionation performed on of cav-1 silenced cells showed that knock down of cav-1 displaced TACE from the light-density fractions and apparently reduced its expression within high-density membrane fractions. However, while TNFR1 similar to TACE can be co-immunoprecipitated with cav-1 from EA.hy926 cell lysates, cav-1 knock down does not displace TNFR1 from (now dispersed) lipid rafts [8]. Data generated in HUVEC at early passages demonstrated similar effects of cav-1 knock-down on histamine-induced shedding (data not shown) even though at lower extent. This phenomenon is not unexpected since primary HUVEC rapidly lose their caveolae in culture and for unknown reason show a smaller amount of exposed TNFR1 compared to EAhy.926. In conclusion, data reported here suggest that cav-1 may play several crucial roles in the shedding of TNFR1 from EC, including maintenance of caveolae network, co-localizing TNFR1 and TACE within caveolae and sustaining higher levels of both proteins. The reduction in the expression of both TNFR1 and TACE followed by cav-1 silencing may be thus the cause of the reduced shedding of TNFR1 receptor, which suggest the importance of cav-1 for maintaining this mechanism controlled. In other words, caveolae/lipid rafts would play a crucial structural role preserving receptor cross-talk rather than regulating receptor signaling. In conclusion, our data suggest that specific targeting of caveolae signaling platform may represent an innovative approach to control endothelial cell functions and might have prospective for management of inflammatory disorders.

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AD designed, performed the research and wrote the paper, AF designed the research and wrote the paper, JSP designed the research, BE and CG performed the research, EZ critically read the paper.

References


Figure 1. Effect of histamine receptors on TNFR1
(A) Cells were cultured in the presence or absence of mepyramine as described in “Materials and Methods” prior to histamine treatment for 30 minutes. Concentrated medium from treated cells were analysed by SDS-PAGE and immunoblotted with TNFR1 antibody. TMPH was used as specific agonist of H1R. NT represent cells stimulated with vehicle alone. Data are representative of three different experiments. (B) Effect of histamine and TMPH on the expression of surface TNFR1. (C) Effect of mepyramine on histamine-induced downregulation of membrane TNFR1. (D) Inhibition of TMPH-induced down regulation of membrane bound TNFR1 in the presence of TACE inhibitor, TAPI-0.
Figure 2. Forskolin reduced both constitutive and TMPH-stimulated release of TNFR1
(A) Effect of 500 μM AMTHA and 100 μM TMPH on sTNFR1 release. (B) Cells were preincubated with 10μM forskolin or with vehicle alone prior to TMPH treatment and soluble TNFR1 was measured by ELISA assay. (C) Values indicate the mean fluorescence intensity measured by FACS analysis of membrane TNFR1 in both forskolin- and vehicle-treated cells. Effect of Ptx treatment on both sTNFR1 release (D) and membrane TNFR1 expression. Data are representative of two independent experiments. (*, P value <0.05).
Figure 3. Evaluation of MbCD effect on TNFR1 shedding and Ca\(^{2+}\) mobilization
(A) ELISA analysis of sTNFR1 from culture media of cells pre-incubated with 5mM MbCD for 30 minutes and then stimulated with TMPH. (B) FACS analysis showing the effect of MbCD on expression of surface TNFR1 in response to TMPH. (C) Effect of cholesterol depletion by MbCD on TMPH-induced calcium mobilization. (*, P value <0.05).
Figure 4. Involvement of cav-1 in the release of TMPH- and histamine-induced release of sTNFR1

(A) Immunoblotting of cav-1 expression showing the efficiency of the two cav-1 dsRNA sequences used for RNAi experiments in EA.hy926 cell line. (B) EA.hy926 cells were first subjected to cav-1 knock down as described in “Materials and Methods” and then treated with either histamine or TMPH for 30 minutes. The amount of sTNFR1 was measured by ELISA assay from cell culture supernatants. (C) Cav-1 knock down did not affect TMPH-induced calcium mobilization. Bars show the increase of calcium mobilization after stimulation with TMPH. Data are representative of results of three independent experiments. (*, P value <0.05).
Figure 5. Effect of cav-1 knock down on proteins distribution and transcripts level

(A) Sucrose gradient fractionation was performed in order to isolate low buoyant fractions enriched in caveolin proteins. Immunoblotting of fractions harvested from the top (low sucrose density) to the bottom (high sucrose density) show that H1R was excluded from caveolin-enriched fractions. (B) In resting cells TACE appeared to localize both with low and high sucrose density membranes. By contrast, cav-1 down regulation by RNAi induced displacement of TACE mainly from the caveolin-enriched membranes (lower panels). (C) Cav-1 immunoprecipitates were immunoblotted for TACE in order to evaluate protein:protein interaction. IgG served as negative control.
Figure 6. Effect of cav-1 RNAi on both TACE and TNFR1 protein expression
(A) Immunoblotting analysis showing the effect of cav-1 knock down on TACE, TNFR1 and cav-1 protein expression. β-Actin (indicated as β-Act) was used as equal loading control (left panel). Densitometric analysis of replicate immunoblotting experiment showing the effect of cav-1 RNAi on TACE, TNFR1 and cav-1 protein expression (right panel). (B) Effect of cav-1 silencing on both cav-1 and TACE mRNA transcripts. (*, P value <0.05)