MYADM controls endothelial barrier function through ERM-dependent regulation of ICAM-1 expression

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ABSTRACT The endothelium maintains a barrier between blood and tissue that becomes more permeable during inflammation. Membrane rafts are ordered assemblies of cholesterol, glycolipids, and proteins that modulate proinflammatory cell signaling and barrier function. In epithelial cells, the MAL family members MAL, MAL2, and myeloid-associated differentiation marker (MYADM) regulate the function and dynamics of ordered membrane domains. We analyzed the expression of these three proteins in human endothelial cells and found that only MYADM is expressed. MYADM was confined in ordered domains at the plasma membrane, where it partially colocalized with filamentous actin and cell–cell junctions. Small interfering RNA (siRNA)-mediated MYADM knockdown increased permeability, ICAM-1 expression, and leukocyte adhesion, all of which are features of an inflammatory response. Barrier function decrease in MYADM-silenced cells was dependent on ICAM-1 expression. Membrane domains and the underlying actin cytoskeleton can regulate each other and are connected by ezrin, radixin, and moesin (ERM) proteins. In endothelial cells, MYADM knockdown induced ERM activation. Triple-ERM knockdown partially inhibited ICAM-1 increase induced by MYADM siRNA. Importantly, ERM knockdown also reduced ICAM-1 expression in response to the proinflammatory cytokine tumor necrosis factor-α. MYADM therefore regulates the connection between the plasma membrane and the cortical cytoskeleton and so can control the endothelial inflammatory response.

INTRODUCTION

The endothelium lines the inner side of blood vessels, forming a barrier between the blood and the surrounding tissue that is essential for vascular homeostasis. The endothelium mediates the passage of small molecules and cells from the bloodstream to the tissues, without compromising its integrity, in the presence of continuous osmotic and shear stress. The organization of the endothelial cell surface, the biological fence facing the vessel lumen, is thus essential for integrating signals from different sources that modulate selective permeability, such as mechanical forces, cytokine signaling, and cell–cell interactions (Millán and Ridley, 2005; Simionescu, 2007; Vandenbroucke et al., 2012).

The architecture of the endothelial surface is determined by the links between the plasma membrane, surface proteins, and underlying cytoskeleton. In the current paradigm, lipids and transmembrane proteins are connected to the underlying cytoskeleton like a
picket fence that controls the stiffness of the cell surface and the ability of lipids and proteins to diffuse and signal (Ritchie et al., 2003). The 4.1 family proteins, ezrin, radixin, and moesin (ERM), are the best-known connectors between plasma membrane, transmembrane receptors, and the subcortical actin cytoskeleton (Bretschner et al., 2002; Viola and Gupta, 2007). An additional level of complexity in cell surfaces arises from the fact that the complex mixture of lipids forming the plasma membrane does not distribute homogeneously but can form microdomains with specific subsets of proteins. The model that best explains this heterogeneity, although it is still under debate, posits the existence of liquid-ordered membrane domains enriched in glycolipids and cholesterol, termed membrane rafts. Raft-mediated plasma membrane condensation would regulate signaling, protein trafficking, and compartmentalization (Lingwood and Simons, 2010; Simons and Gerl, 2010).

Some evidence suggests a mutual regulation between ERM proteins and ordered membrane rafts (Viola and Gupta, 2007). On activation, ERMs acquire an open conformation in which the N-terminal Four point one, ezrin, radixin-moesin (FERM) domain interacts directly or indirectly with the cytoplasmic tails of transmembrane receptors and the C-terminal domain binds to actin filaments (Feihon et al., 2010). The FERM domain binds the raft lipid phosphatidylinositol 4,5-bisphosphate (PIP2) and some transmembrane proteins that also partition into ordered membranes (Wojciak-Stothard et al., 1999; Tilghman and Hoover, 2002; Johnson et al., 2008; Kwiatkowska, 2010). ERMs themselves have also been found compartmentalized into ordered membrane domains (Tomas et al., 2002; Gupta et al., 2006; Heyraud et al., 2007). Finally, active ERMs have been shown to regulate membrane raft content (Prag et al., 2007) and prevent membrane raft coalescence (Gupta et al., 2006).

Inflammatory cytokines have the ability to alter the endothelial barrier to cells and solutes by mechanisms that are not completely understood (Vandenbroucke et al., 2008). Disruption of cholesterol-rich membrane rafts has been shown to either promote or ameliorate inflammation in different cell types (Legler et al., 2003; Flemming et al., 2004; Meng et al., 2010). In addition, cytokines such as tumor necrosis factor-α (TNF-α) and interleukin-1β (IL-1β) activate ERM proteins. This activation mediates the increase in endothelial permeability or the transcriptional regulation in response to TNF-α (Kishore et al., 2005; Koss et al., 2006). However, the full role of ERM activation in the endothelial inflammatory response is still not well defined. It is of note that ERM proteins play an important role localizing receptors involved not only in leukocyte adhesion but also in the regulation of endothelial permeability in an inflammatory context, for example, intercellular adhesion molecule (ICAM)-1 or vascular cell adhesion molecule (VCAM)-1 (van Wetering et al., 2003; Clark et al., 2007; Sumagin et al., 2008).

In the current model of ordered membrane domains or rafts, protein machinery associated with these platforms is required to specify their function in the cell (Bauer and Pelkmans, 2006; Lingwood and Simons, 2010). The MAL family consists of integral proteins with several transmembrane domains involved in membrane dynamics (Sanchez-Pulido et al., 2002). Several lines of evidence support a role for these proteins in events related to organization of membrane domains, such as polarized membrane trafficking or plasma membrane reorganization in response to extracellular signaling (Puertollano et al., 1999; Antón et al., 2008; Goldstein Magal et al., 2009; Antón et al., 2011; Aranda et al., 2011). Myeloid-associated differentiation marker (MYADM) is a MAL family protein that regulates membrane order in migrating epithelial cells (Aranda et al., 2011). In this paper, we report that MYADM is a component of endothelial surface rafts. MYADM knockdown induces an inflammatory-like phenotype, altering barrier function through the increase of the adhesion receptor ICAM-1. The rise in ICAM-1 levels is mediated by MYADM-regulated activation of ERM proteins. Collectively our results strongly suggest that the interaction between the plasma membrane and the submembrane actin cytoskeleton at the endothelial surface controls the inflammatory response.

RESULTS

MYADM is confined into ordered domains in the endothelial plasma membrane

Specific types of highly polarized epithelial cells express the MAL family members MAL and MAL2, which are involved in specialized, raft-dependent, apical vesicular trafficking (Puertollano et al., 1999; de Marco et al., 2002). MYADM, in contrast, is expressed ubiquitously and organizes membrane rafts at the epithelial cell surface, thereby affecting cell spreading and motility (Aranda et al., 2011). The endothelium can be considered as a simple squamous epithelium (Palade et al., 1979). Because ordered membrane domains have been shown to be central to endothelial barrier function and signaling (Muppidi et al., 2004; Heyraud et al., 2007; Chidlow and Sessa, 2010; Triantafiliou et al., 2010), we wondered whether these MAL proteins are expressed and play a role in primary human endothelial cells. Detergent-resistant membrane (DRM) fractions were isolated from confluent human umbilical vein endothelial cells (HUVECs) and blotted with antibodies against MAL, MAL2, and MYADM, together with a panel of cell lines used as positive and negative controls. Endogenous MYADM was clearly detected in DRM fractions from all the analyzed cell types, including HUVECs, consistent with the ubiquitous expression pattern previously reported (Aranda et al., 2011). In contrast, MAL and MAL2 were detected only in DRM fractions from epithelial PC3 and Madin-Darby canine kidney (MDCK) cells (Figure 1A). Quantitative PCR (qPCR) assays confirmed that expression of MAL and MAL2 was negligible in HUVECs (Figure 1B). Further analysis of MYADM fractionation in HUVECs showed that this protein is almost exclusively found in DRMs (Figure 1, C and D). The 2B12 monoclonal antibody (mAb), raised against human MYADM is not suitable for immunofluorescence analysis (Aranda et al., 2011). For the study of MYADM distribution, MYADM-green fluorescent protein (GFP) was expressed in human endothelial cells and analyzed by confocal microscopy. MYADM-GFP also partitioned in DRMs, indicating that the GFP tag does not alter MYADM insertion in plasma membrane domains (Figure 1E). MYADM-GFP was localized at the plasma membrane and in cell–cell junctions in confluent endothelial cells, and this distribution overlapped with the staining for filamentous actin and junctional markers, such as vascular endothelial (VE)-cadherin (Figure 1F, insets). A partial colocalization of MYADM-GFP at cell–cell junctions was also found with platelet endothelial cell adhesion molecule (PECAM)-1, 1-p20-catenin, occludin, and nectin-2 (Supplemental Figure S1). In contrast, MYADM-GFP showed little colocalization with caveolin-1, a raft-associated protein essential for vascular homeostasis (Chidlow and Sessa, 2010), or with CD59, a glycosylphosphatidylinositol (GPI)-anchored protein often used as a marker of membrane rafts (Millán et al., 2006), which suggests that MYADM could organize ordered membrane domains differently from caveolae or CD59 in human endothelial cells (Figure 1F).

MYADM knockdown induces an inflammatory-like phenotype by inducing ICAM-1 expression

To gain insight into the role of MYADM in endothelial cells, we investigated the effects of three different small interfering RNAs (siRNAs) that targeted MYADM with low (siMYADM 1), medium
MYADM regulates the endothelial barrier

MYADM silencing caused a clear disruption of the endothelial monolayer and partially disorganized cell junctions, as shown by staining of β-catenin, VE-cadherin, or PECAM-1 (Figure 2, C and D). MYADM reduction induced the appearance of intercellular gaps (Figure 2, C and D, asterisks, and E) and increased both filamentous (F)-actin (Figures 2, C and F, and S2) and permeability in Transwell assays (Figure 2G). Collectively, these results indicate that MYADM knockdown caused endothelial barrier dysfunction.

Increased permeability and polymerization of actin are prototypical endothelial responses to several inflammatory stimuli (Pober and Sessa, 2007). We tested whether MYADM knockdown was inducing an inflammatory-like response by following the expression of different receptors previously involved in leukocyte adhesion and permeability and considered typical inflammatory markers (Albelda, 1991; Clark et al., 2007; Sumagin et al., 2008). Western blot analysis indicated that reduction of MYADM levels induced the expression of the adhesion receptor ICAM-1, but not of E-selectin or VCAM-1 (Figure 3, A and B). MYADM knockdown also increased ICAM-1 mRNA, as shown by qPCR experiments (Figure 3C). Immunofluorescence assays revealed the surface accumulation of this receptor in MYADM-depleted cells (Figure 3D). In contrast, no difference was found in the expression of different components of cell–cell junctions involved in modulation of barrier function in an inflammatory context (Fernandez-Martin et al., 2012; Reglero-Real et al., 2012), such as VE-cadherin, β-catenin, p120-catenin, and PECAM-1 (Figure 3A). The Rho GTPases RhoA, Rac1, and Cdc42 have been involved in permeability and F-actin regulation. No statistically significant changes were detected in their activity upon MYADM knockdown (Figure S3, A and B). Also, no significant alteration was detected in the phosphorylation of regulatory myosin

FIGURE 1: MYADM is a transmembrane protein associated with endothelial DRMs. (A) The indicated cell types were extracted with 1% Triton X-100 at 4°C and centrifuged to equilibrium in sucrose density gradients. Aliquots from DRMs were immunoblotted for MYADM, MAL2, human MAL (hMAL), canine MAL (dMAL), and caveolin-1 (Cav-1). (B) RNA from the indicated cells was isolated and the expression of MYADM, MAL2 and MAL transcripts was analyzed by qPCR. Results were normalized to mRNA levels of housekeeping genes (β-actin and GADPH) and represented as percentage of PC3 mRNA levels for MYADM and MAL2 transcripts (left) or Jurkat mRNA levels for the MAL transcript (right). (C) Aliquots from soluble (SOL) and DRM fractions from HUVECs were immunoblotted for MYADM, Cav-1 (as a control of endothelial DRMs), and transferrin receptor (TFR; as a control of transmembrane proteins in the soluble fraction). (D) Quantitation of DRM segregation of the indicated protein from three independent experiments. (E) DRMs were isolated from cells expressing MYADM-GFP for 24 h and immunoblotted for MYADM with anti-MYADM mAb 2B12 and for caveolin-1. (F) MYADM-GFP distributes at the plasma membrane of HUVECs and partially colocalizes with subcortical F-actin and with VE-cadherin. Caveolin-1 and CD59 staining reveals little colocalization of MYADM-GFP with caveolae or with GPI-protein-rich rafts in endothelial cells. Scale bar: 20 μm.
FIGURE 2: MYADM knockdown alters endothelial barrier function. (A) HUVECs were transfected in parallel with control siRNA (siControl) or with three different siRNAs targeting MYADM (siMYADM). At 72 h after transfection, MYADM levels were analyzed by immunoblotting. ERK-1/2 and caveolin-1 (Cav-1) levels are shown as loading controls. (B) Quantitation of MYADM reduction in response to siRNA transfection. The mean ± SEM from three independent experiments is presented. (C and D) HUVECs were transfected with the indicated siRNAs and plated at confluence on fibronectin-coated coverslips. After 72 h, cells were fixed and stained for F-actin and β-catenin (C) or VE-cadherin or PECAM-1 (D) to detect cell–cell junctions. Asterisks in the images indicate intercellular gaps. Scale bar: 20 μm for (C) and 10 μm for (D). (E) Quantitation of intercellular gaps in cells transfected with the indicated siRNAs. (F) Cells were fixed and stained with antibody against total actin or with TRITC–phalloidin. Quantitation of the ratio F-actin/total actin (see Figure S2). (G) HUVECs were transfected with the indicated siRNAs and plated at confluence on Transwells, and permeability assays were performed 72 h after transfection. The mean ± SEM from three independent experiments is shown. *, p < 0.05; **, p < 0.01.
FIGURE 3: MYADM knockdown induces ICAM-1 expression. HUVECs were transfected with the indicated siRNAs for 72 h, and the expression of different proteins involved in endothelial barrier function was analyzed by Western blot (A). Quantitation of ICAM-1 protein (B) and mRNA by qPCR (C). The mean ± SEM from three (B) and two (C) independent experiments is shown. **, p < 0.01. (D) Localization of ICAM-1 in siRNA-transfected cells by immunofluorescence analysis. (E) HUVECs expressing MYADM-GFP for 36 h were stimulated with TNF-α (10 ng/ml) for 6 h to induce detectable levels of ICAM-1. Subsequently the receptor was cross-linked with specific antibodies (X-ICAM-1). Cells were fixed, permeabilized, and stained with TRITC–phalloidin to visualize F-actin. (F) HUVECs expressing MYADM-GFP were starved, stimulated with TNF-α (10 ng/ml) for 6 h to induce expression of adhesion receptors, and then incubated with T-cells for 15 min. Cells were fixed and stained with TRITC–phalloidin to visualize F-actin (top panels) or with TRITC–phalloidin and anti–caveolin-1 antibody (bottom panels). T-cells adhering or transmigrating were morphologically distinguishable by the F-actin staining (red dotted line). Right graphs show intensity profiles across the T-cell–endothelial cell interaction area along the indicated white arrows.
light chain, which indicates that actomyosin-mediated contractility downstream of RhoA is not affected by MYADM depletion (Figure S3C). Although MYADM knockdown induces intercellular gaps, no increase in VE-cadherin tyrosine phosphorylation or decrease in β-catenin association to VE-cadherin was detected upon MYADM knockdown (Figure S4), which suggests the stability of adherens junctions is not regulated by MYADM expression. Collectively these experiments indicate that MYADM plays a role in barrier function and ICAM-1 transcriptional regulation.

MYADM did not appear to associate with ICAM-1, in either ICAM-1 immunoprecipitation (unpublished data) or cross-linking experiments (Figure 3E). In addition, MYADM did not distribute around either adhered or transmigrating leukocytes (Figure 3F), in contrast with caveolin-1, which is enriched in areas where T-cells are transmigrating transcellularly, as previously described (Carman and Springer, 2004; Millán et al., 2006; Keuschnigg et al., 2009). Altogether these data suggest that MYADM depletion induces an increase in ICAM-1 expression but that MYADM is not directly associated with ICAM-1 or involved in ICAM-1 function. To rule out the possibility of an off-target effect of the siRNA oligonucleotides, we followed a rescue-of-function strategy by transfecting siMYADM 2, which targets the 3′ untranslated region and is thus a proper translocation from the cytosol to the plasma membrane and/or cortical actin organization of endothelial cells (Viola and Gupta, 2007). Activation of ERM proteins, connectors between the plasma membrane and the cytoskeleton, increases permeability in microvascular endothelial cells during the inflammatory response (Koss et al., 2006), so we tested the effect of MYADM knockdown on ERM phosphorylation at a specific threonine residue that is required for their conformational change and subsequent activation. Western blotting with an antibody recognizing phosphorylated ezrin (T567), radixin (T564), and moesin (T558) revealed that MYADM silencing increased ERM phosphorylation two- to threefold (Figure 6, A and B). Confocal analysis of MYADM-depleted cells confirmed an increase in phosphorylated ERMs at the plasma membrane (Figure 6C). Both in control and MYADM-depleted cells, pERM appeared particularly enriched in junctional membrane projections, often connecting two adjacent cells. The length of these border protrusions increased in response to MYADM knockdown (Figure 6C, arrowheads, and D). We then examined whether ERM phosphorylation can contribute to endothelial barrier dysfunction. Transfection of a constitutive active form of ERMs (moesin T558D) caused a moderate decrease of cell–cell junction integrity, whereas moesin T558A mutant, which is proposed to be in a closed, inactive conformation, did not significantly alter the distribution of the junctional marker β-catenin (Figure 6, E and F).

ICAM-1 expression increases in response to MYADM reduction or TNF-α requires ERM expression

We then examined whether ERM proteins mediate ICAM-1 protein increase caused by MYADM knockdown. Cells were transfected with control siRNA or siRNA targeting MYADM (siMYADM 2), ERM proteins (siE+R+M), or both (Figures 7A and S5, A and B). MYADM single knockdown increased ICAM-1, whereas simultaneous silencing of MYADM and ERMs reduced ICAM-1 to control levels (Figure 7, A and B). In MYADM-depleted cells, single knockdown of ezrin, radixin, and moesin reduced ICAM-1 expression to a lesser extent than simultaneous triple-ERM silencing (Figure S5C). These findings indicate that the three proteins are playing additive roles. Accordingly, ERM knockdown diminished the increase in leukocyte adhesion caused by MYADM depletion (Figure 7C). Triple-ERM knockdown diminished barrier integrity in control cells, due probably to the essential role of these cross-linker proteins in connecting membrane to subcortical cytoskeleton, so no rescue of function could be performed following this strategy. Together our data indicate that MYADM reduction induces a proinflammatory-like phenotype by regulating ERM protein phosphorylation and ICAM-1 expression. Several protein kinases can potentially regulate ERM phosphorylation in this particular threonine residue. Among them, classical protein kinase C (PKC) family members, serine/threonine kinases that require their conformational change and subsequent activation. Western blotting with an antibody recognizing phosphorylated ezrin (T567), radixin (T564), and moesin (T558) revealed that MYADM silencing increased ERM phosphorylation two- to threefold (Figure 6, A and B). Confocal analysis of MYADM-depleted cells confirmed an increase in phosphorylated ERMs at the plasma membrane (Figure 6C). Both in control and MYADM-depleted cells, pERM appeared particularly enriched in junctional membrane projections, often connecting two adjacent cells. The length of these border protrusions increased in response to MYADM knockdown (Figure 6C, arrowheads, and D). We then examined whether ERM phosphorylation can contribute to endothelial barrier dysfunction. Transfection of a constitutive active form of ERMs (moesin T558D) caused a moderate decrease of cell–cell junction integrity, whereas moesin T558A mutant, which is proposed to be in a closed, inactive conformation, did not significantly alter the distribution of the junctional marker β-catenin (Figure 6, E and F).

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**DISCUSSION**

Membrane rafts are cholesterol-enriched ordered domains that regulate a plethora of signaling pathways. Cytokines such as TNF-α (Muppidi et al., 2004) and IL-1β (Oakley et al., 2009) or bacterial ligands of Toll-like-receptors (Soong et al., 2004; Triantafilou et al., 2010) induce proinflammatory signals through membrane rafts. Moreover, modulation of cholesterol content in these domains can induce or decrease inflammatory signaling in different cell models (Legler et al., 2003; Flemming et al., 2004; Meng et al., 2010). Collectively these observations suggest that modulation of membrane condensation may itself contribute to the endothelial inflammatory response, probably by mimicking those changes in control endothelial barrier function (Hempel et al., 1997), have been shown to phosphorylate ERM proteins in different cellular contexts (Ng et al., 2001; Ivetic and Ridley, 2004; Adyshev et al., 2011). We addressed whether increased ERM phosphorylation in response to MYADM reduction is dependent on PKC by using a specific inhibitor for this family of kinases. Incubation with Gö6976 for 8 h partially reduced ERM phosphorylation increase in MYADM siRNA–transfected cells, which suggests that MYADM is involved in regulating classical PKC-mediated activation of ERM proteins at the plasma membrane (Figure 7D). Together these results are compatible with a role for MYADM organizing endothelial plasma membrane and thereby regulating the membrane targeting of proteins, such as classical PKCs, that control subcortical protein machinery. Finally, we investigated whether ERM can also regulate ICAM-1 expression in a more physiological context. TNF-α is a proinflammatory cytokine that induces ICAM-1 expression and ERM phosphorylation (Koss et al., 2006; Figure 7, E and F). Triple-ERM knockdown reduced ICAM-1 increment upon TNF-α exposure (Figure 7, G and H). In contrast, MYADM knockdown did not exacerbate ICAM-1 increase in response to this cytokine (Figure 7, I and J), and TNF-α did not significantly alter MYADM levels (Figure S6). Therefore these results indicate that ERM also regulates ICAM-1 expression in response to physiological inflammatory signaling. Collectively our data suggest that a connection between endothelial membrane and cortical actin cytoskeleton modulates the inflammatory response.
clustering or diffusion that different stimuli can induce on protein signaling machinery associated with the plasma membrane. This could be important for chronic inflammatory diseases such as arteriosclerosis, which is triggered by deregulation of homeostasis of raft lipids, as is seen in hypercholesterolemia (Libby, 2002). However, ordered membrane microdomains require protein scaffolds to play specific roles within the cell (Bauer and Pelkmans, 2006). The MAL proteins contain one or two tetra-spanning MARVEL (MAL and related proteins for vesicle trafficking and membrane link) domains that are involved in membrane apposition (Sanchez-Pulido et al., 2002). Different experimental strategies have shown that these proteins have the ability to modulate membrane order or dynamics in a variety of cell types (Puertollano et al., 1997; Goldstein Magal et al., 2009; Aranda et al., 2011). As a consequence, these proteins control the segregation into membrane rafts of signaling proteins, such as the tyrosine kinase Lck (Anton et al., 2008) or the Rho GTPase Rac-1 (Aranda et al., 2011). In this paper, we show that, of the three MAL family proteins with a reported role in raft-mediated function, only MYADM is expressed in endothelial cells. MYADM is also the only member of the MAL family containing two MARVEL domains, rather than one. This feature seems to determine its predominant localization at the plasma membrane, rather than in vesicular compartments (Aranda et al., 2011), although MYADM trafficking has yet to be defined. On the other hand, colocalization analysis clearly shows that MYADM-containing domains are different than those of caveolae or GPI-anchored proteins in endothelial cells, in agreement with our previous studies in other cell types or with other MAL family proteins (Millán et al., 1997; Aranda et al., 2011).

ERM proteins are cross-linkers between transmembrane receptors and cortical actin filaments that regulate the structure and function of specific domains at the cell surface, namely microvilli or filopodia, and fundamental signaling processes involved in cell shape, junctional stability, motility, and cytokinesis (Fehon et al., 2010). Cytokines, such as IL-1β and anti–β-catenin antibodies to detect transfected cells and cell–cell junctions, respectively. (F) Quantitation of β-catenin intensity of transfected cells normalized to the intensity of adjacent cells (control). Five hundred thirty regions from 53 transfected cells were analyzed in three different experiments. The mean + SEM is shown. *, p < 0.05.

**FIGURE 6:** MYADM knockdown activates ERM proteins. (A) HUVECs were transfected with the indicated siRNAs for 72 h, and ERM activation was monitored by Western blot with anti-pER(T567)R(T564)M(T558)–specific antibody (pERM) and anti-ERM protein antibody. (B) Quantitation of pERM activation was performed by measuring the ratio of anti-pERM to ERM blotting signal. The mean + SEM from three independent experiments is shown. (C and D) HUVECs were transfected as in (A) and fixed with 10% trichloroacetic acid, and pERM distribution was analyzed by immunofluorescence. Arrowheads indicate the pERM-rich protrusions (C), with their lengths quantitated in (D). (E) The HA-tagged moesin mutants, moesin T558A (TA), in an inactive conformation, or moesin T558D (TD), constitutively active, were transiently expressed in HUVECs for 48 h. Cells were fixed and stained with anti-HA and anti–β-catenin antibodies to detect transfected cells and cell–cell junctions, respectively. (F) Quantitation of β-catenin intensity of transfected cells normalized to the intensity of adjacent cells (control). Five hundred thirty regions from 53 transfected cells were analyzed in three different experiments. The mean + SEM is shown. *, p < 0.05.
FIGURE 7: ERM knockdown prevents ICAM-1 induction mediated by MYADM knockdown or TNF-α. (A) HUVECs were transfected with the indicated siRNAs for 72 h, and MYADM, ERM, and ICAM-1 expression was analyzed by Western blotting. (B) Quantitation of ICAM-1 levels in siRNA-transfected cells. (C) siRNA-transfected HUVECs were plated at confluence, and adhesion assays were performed 72 h after transfection. The mean ± SEM from three independent experiments is shown. (D) HUVECs were transfected with the indicated siRNAs for 72 h. Eight hours before lysis, cells were incubated or not with 100 nm of the PKC inhibitor Gö6976. Subsequently MYADM, ERM, and ICAM-1 expression was analyzed by Western blotting. (E and F) HUVECs were stimulated at different times with TNF-α, lysed, and immunoblotted for the indicated antibodies. (G–J) siRNA-transfected HUVECs were stimulated at different times with TNF-α, lysed, and immunoblotted for the indicated antibodies. Graphs show the mean ± SEM from three independent experiments (H and J). *, p < 0.05; **, p < 0.01.
and TNF-α, or sphingosine 1-phosphate (S1P), all of which are involved in the regulation of the inflammatory response, can also activate ERM proteins (Koss et al., 2006; Adyshev et al., 2011; Huang et al., 2011). This activation, at least for TNF-α and S1P, mediates changes in endothelial barrier function, with different outcomes depending on the stimulus. Ordered domains at the plasma membrane and ERM proteins have thus been independently implicated in inflammation. In this study, we demonstrate that siRNA-mediated modulation of MYADM is sufficient to induce activation of ERM proteins, and this can act as a signal to alter barrier function and to provoke an inflammatory-like response in the absence of inflammatory stimuli. As the immunofluorescence analysis indicates, ERM activation occurs at the plasma membrane and at cell borders. A previous example of cross-regulation between rafts and ERM proteins at the plasma membrane has been reported in B-cells, in which ezrin can modulate the dynamics of ordered domains and their ability to contribute to B-cell receptor (BCR) signaling (Gupta et al., 2006; Viola and Gupta, 2007). Gupta et al. (2006) proposed that, in resting B-cells, active ezrin maintains small raft domains that contain signaling molecules and are linked to cortical F-actin and dispersed on the cell surface. On BCR activation, ezrin becomes transiently inactivated and uncouples actin from the plasma membrane domains. These small rafts then coalesce and promote downstream B-cell signaling. It is therefore plausible that membrane rafts, which are regulated by MYADM, in coordination with the ERM proteins, which connect rafts to the subcortical cytoskeleton, could mediate clustering, diffusion or localization of proteins involved in signaling from inflammatory agonists in endothelial or epithelial cells. In accordance, we have found that, at least for TNF-α, a complete inflammatory response requires ERM protein expression. MYADM was not modulated by TNF-α, and its reduction had no effect on TNF-α-mediated ICAM-1 induction. The identification of the stimuli that act through MYADM and ERM proteins to promote or inhibit inflammation will be important to increase our understanding of the endothelial inflammatory response. ERM proteins are involved in the inflammatory control of protein expression (Kishore et al., 2005) and in connecting adhesion receptors, such as the ICAM family, to subcortical actin (Viola and Gupta, 2007; Reglero-Real et al., 2012). In this paper, we show that ERM expression is required for ICAM-1 expression in response to MYADM suppression or TNF-α. ICAM-1 is a paradigmatic adhesion receptor that regulates leukocyte adhesion. However, its role as a protein essential for controlling endothelial permeability both in vivo and in vitro is emerging (Clark et al., 2007; Sumagin et al., 2008). MYADM knockdown selectively increases ICAM-1 expression, but not many other proteins involved in barrier function, which indicates this protein selectively controls some but not all of the pathways involved in the endothelial inflammatory response. ICAM-1 increase is necessary for MYADM-mediated alteration of barrier properties, which is in agreement with previous reports that link leukocyte adhesion and permeability through dual roles played by ICAM-1 (Clark et al., 2007; Sumagin et al., 2008), VCAM-1 (van Wetering et al., 2003), or PECAM-1 (Graesser et al., 2002; Fernandez-Martín et al., 2012) receptors.

Regulation of the interaction between the plasma membrane and the underlying cytoskeleton is also essential in mechanotransduction (Chien, 2007). Cholesterol-rich membrane domains are required for signaling propagated by mechanic forces induced by shear stress (Yamamoto et al., 2003) or stretch (Zeidan et al., 2003). Laminar shear stress elicits proinflammatory signals that promote ICAM-1 up-regulation in vitro (Tzima et al., 2002). Pathological alteration of these forces in vivo, such as disturbed blood flow, are proinflammatory and proatherogenic (Chien, 2007). It is therefore possible that membrane and actin reorganization induced by MYADM knockdown regulates pathways related to cell stiffness and mechanotransduction. Finally, further understanding of the structure of membrane ordered domains, their protein machinery, and their involvement in inflammatory signaling could lead to the design of new therapies for chronic inflammatory diseases.

**MATERIALS AND METHODS**

**Materials**

Monoclonal antibody to human MYADM (mAb 2B12) was generated as previously described (Aranda et al., 2011). Rabbit anti–ICAM-1, anti–caveolin-1 (N-20), and anti–ERK1/2 antibodies were obtained from Santa Cruz Biotechnology (Santa Cruz, CA). Mouse anti–ICAM-1 antibody was from R&D Systems (Abingdon, UK). Mouse monoclonal anti–VE-cadherin antibody was from BD Biosciences (San Jose, CA). Rabbit anti–β-catenin and TRITC/FITC–phalloidin was from Sigma- Aldrich (St. Louis, MO). Anti–transferrin receptor antibodies were purchased from Zymed (San Francisco, CA). Anti–ERM, anti–phospho–ERM, and specific anti–ezrin, anti–radixin, and anti–moesin antibodies were from Cell Signaling Technology (Boston, MA). Hemagglutinin (HA)-tagged moesin-TD (T558D) and moesin-TA (T558A) were from J. Délon (Université René Descartes, Paris, France).

**Cell culture and transfection**

HUVECs were obtained and cultured as previously described (Millán et al., 2010) and were always plated at confluence on fibronectin-coated dishes for 24–72 h. HeLa, MDCK, and PC3 cells were obtained from the American Type Culture Collection (Manassas, VA) and grown in DMEM supplemented with 5% fetal bovine serum. Isolation of HeLa clones stably expressing MYADM-GFP and human memory T-cells for adhesion assays has been described elsewhere (Millán et al., 2006; Aranda et al., 2011). HUVECs were transiently transfected with 1–5 μg plasmid DNA/10^6 cells with calcium phosphate and used for experiments 24–72 h after transfection. For siRNA transfection, we devised a protocol for delivery of siRNA with high efficiency into primary endothelial cells that is a modification of our previous method (Millán et al., 2006). HUVECs were plated at subconfluence (10^6 cells on each well of a six-well dish) in EBM-2 medium (Lonza, Walkersville, MD) with no antibiotics. The following day, cells were transfected by mixing 4 μl of oligofectamine (Invitrogen, Carlsbad, CA) with siRNA to a final concentration of 100 nM. At 24 h after transfection, cells were trypsinized and plated at confluence onto different dishes for parallel assays, such as permeability, adhesion, immunofluorescence, or Western blotting. Assays were performed 72 h after transfection. Sequences of siRNA duplexes targeting MYADM were 5′-CGAGATCACTGGCTATATG-3′ (siMYADM 1), 5′-GATGTAAGCCTGCAGCCGCA-3′ (siMYADM 2), and 5′-GGTCTAAGACTCTCCCAAG-3′ (siMYADM 3). Ezrin was targeted with the siRNA oligonucleotide 5′-CAAGAGGGCACCUCACUU-3′, moesin with 5′-AUAGAAGUGCAUAAGUC-3′, and radixin with 5′-GAACUGGCAUGAAGAACAU-3′ and 5′-GAACUGGCAUGAAGAACAU-3′. Sequences of siRNA controls were obtained from Dharmaco (Lafayette, CO).

**DRM isolation**

Different cell lines and HUVECs grown to confluence in 100-mm dishes were lysed for 20 min in 1 ml of 25 mM Tris-HCl (pH 7.5), 150 mM NaCl, 5 mM EDTA, 1% Triton X-100, 4°C, as indicated (Millán et al., 2002). DRM fraction was isolated by centrifugation to equilibrate in a sucrose discontinuous density gradient in a swinging bucket SW40 rotor (Beckman Coulter, Brea, CA) at 39,000 rpm (188,000 × g) for 20 h (Brown and Rose, 1992; Millán et al., 2006). Fractions were harvested from the bottom sucrose layer (40%; soluble) and from the above interphase between 5 and 30% sucrose.
layers (DRMs). Equivalent aliquots were subjected to immunoblot analysis with the appropriate antibodies.

**ICAM-1 cross-linking and immunofluorescence**

Anti–ICAM-1 antibody (1 μg/ml) was added to TNF-α–stimulated HUVECs for 45 min. Cells were rinsed and incubated with 1 μg/ml of fluorophore-coupled secondary antibody for 30 min to induce receptor clustering. Cells were rinsed and fixed with 4% paraformaldehyde for 20 min, blocked with TBS (25 mM Tris, pH 7.4, 150 mM NaCl) for 10 min, permeabilized for 5 min with phosphate-buffered saline (PBS) containing 0.2% Triton X-100 at 4°C, and incubated at 37°C with tetramethylrhodamine isothiocyanate (TRITC)-labeled phallolidin to visualize F-actin. Specimens were mounted in DAKO fluorescent mounting medium (DAKO, Ely, UK). For immunofluorescence analysis the adhered T-cells, unlabeled T-cells were added to transfected HUVECs stimulated with TNF-α for 6 h. Cells were fixed and stained with the indicated antibodies and with TRITC-phallolidin to visualize T-cells. For F-actin quantification, confocal images from cells stained for TRITC-labeled phallolidin were exported in formats compatible with ImageJ software (http://rsb.info.nih.gov/ij). ImageJ was used to obtain the mean fluorescence intensities from different images of confluent cells.

**Adhesion assays**

Starved memory T-cells were labeled with 2′,7′-bis-(2-carboxyethyl)-5-(and-6)-carboxyfluorescein, acetoxymethyl ester (Molecular Probes, Eugene, OR) in serum-free medium for 20 min, washed with medium containing 1% BSA, and incubated for 15 min at 37°C with confluent HUVECs previously transfected with different siRNAs. Nonadhered cells were washed off, and adhesion was determined by fluorescence measurement in a Fusion α-FS fluorimeter (Perkin Elmer-Cetus, Waltham, MA). Data were processed and statistical significance was determined using Student’s t test (Microsoft Excel; Redmond, WA). For immunofluorescence analysis of adhered T-cells, unlabeled T-cells were added to transfected HUVECs stimulated with TNF-α for 6 h. Cells were fixed and stained with TRITC-phallolidin (to visualize T-cells) and with the indicated antibodies.

**Permeability assays**

Permeability assays were carried out as previously described (McKenzie and Ridley, 2007). siRNA-transfected HUVECs were plated at confluence on fibronectin-coated Transwells of 0.4-μm diameter. FITC-dextran (Mr 42,000; Sigma–Aldrich) was applied apically at 0.1 mg/ml and allowed to equilibrate for 60 min before a transcellular assay in which cells were incubated in PBS for 20 min and the calcium levels were subsequently restored by replenishment with normal EBM-2 medium. Data were normalized to the mean resistance detected before the calcium switch.

**qPCR analysis**

One microgram of RNA from HUVECs, PC3, Jurkat, and HepG2 cell lines was subjected to reverse transcription with the High Capacity RNA-cDNA kit (Applied Biosystems, Bedford, MA). qPCR was performed from the resulting cDNA in a thermocycler CFX 384 (Bio-Rad, Hercules, CA) using the SsoFast EvaGreen Supermix (Bio-Rad) and the following forward and reverse primers, previously designed with ProbeFinder software (Roche Diagnostics, Indianapolis, IN): MAL2, 5′-GACAATCTCGCCGACCTACTC-3′; 5′-AGACAGACCCCCACCAAC-3′; MYADM, 5′-GGCCTGCTCTCAGCTACG-3′; 5′-TAGCCACGTGTCATGAC-3′; MAL, 5′-AGACTCCCTGGTCACCTTG-3′; 5′-CTCGTTCGATCGTATG-3′. Parallel qPCR from β-actin and GAPDH was performed to normalize data from each cell type with the following primers: β-actin, 5′-CAGGCGACAGGGCGCTG-3′; 5′-GTCTGAGGCCTCTCTGCTCT-3′. GAPDH, 5′-AGCCCATCGTCCAGACAC-3′; 5′-CCGCCCCATACGACAAAT-3′.  

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**REFERENCES**


