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*J Immunol* 2013; 190:4236-4244; Prepublished online 11 March 2013;
doi: 10.4049/jimmunol.1300046
http://www.jimmunol.org/content/190/8/4236

**Supplementary Material**

http://www.jimmunol.org/content/suppl/2013/03/11/jimmunol.1300046.DC1

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Monocyte ADAM17 Promotes Diapedesis during Transendothelial Migration: Identification of Steps and Substrates Targeted by Metalloproteinases

Yoshiaki Tsubota,* Jeremy M. Frey,* Phillip W. L. Tai,†,1 Robert E. Welikson,†,2 and Elaine W. Raines*

Despite expanded definition of the leukocyte adhesion cascade and mechanisms underlying individual steps, very little is known about regulatory mechanisms controlling sequential shifts between steps. We tested the hypothesis that metalloproteinases provide a mechanism to rapidly transition monocytes between different steps. Our study identifies diapedesis as a step targeted by metalloproteinase activity. Time-lapse video microscopy shows that the presence of a metalloproteinase inhibitor results in a doubling of the time required for human monocytes to complete diapedesis on unactivated or inflamed human endothelium, under both static and physiological-flow conditions. Thus, diapedesis is promoted by metalloproteinase activity. In contrast, neither adhesion of monocytes nor their locomotion over the endothelium is altered by metalloproteinase inhibition. We further demonstrate that metalloproteinase inhibition significantly elevates monocyte cell surface levels of integrins CD11b/CD18 (Mac-1), specifically during transendothelial migration. Interestingly, such alterations are not detected for other endothelial- and monocyte-adhesion molecules that are presumed metalloproteinase substrates. Two major transmembrane metalloproteinases, a disintegrin and metalloproteinase (ADAM)17 and ADAM10, are identified as enzymes that control constitutive cleavage of Mac-1. We further establish that knockdown of monocyte ADAM17, but not endothelial ADAM10 or ADAM17 or monocyte ADAM10, reproduces the diapedesis delay observed with metalloproteinase inhibition. Therefore, we conclude that monocyte ADAM17 facilitates the completion of transendothelial migration by accelerating the rate of diapedesis. We propose that the progression of diapedesis may be regulated by spatial and temporal cleavage of Mac-1, which is triggered upon interaction with endothelium. The Journal of Immunology, 2013, 190: 4236–4244.

Inflammation is a key process in disease pathogenesis, and leukocyte transendothelial migration into inflamed tissues controls both initiation and progression of acute and chronic inflammatory diseases. Transendothelial migration is a sequential, multistep process that consists of leukocyte rolling, capture, and firm adhesion to endothelium, followed by locomotion across the endothelium and diapedesis through the intact vessel into the tissue, usually at endothelial junctions (1–3). Engagement of key ligands and their counterreceptors expressed on both leukocytes and endothelial cells have been shown to be responsible for each step of transendothelial migration (1–3). However, very little is known about regulatory mechanisms underlying the progression through each step and transitions between steps.

A molecular mechanism capable of rapidly reducing adhesion molecule interactions is their proteolytic cleavage or “ectodomain shedding” (4). In fact, soluble forms of multiple adhesion molecules involved in each step of leukocyte transendothelial migration are proteolytically shed, and elevated levels are detected in physiological fluids from various inflammatory diseases (4, 5). In addition, recent studies suggest roles for ectodomain shedding in leukocyte trafficking. For example, shedding of Mac-1 (CD11b/CD18), a major integrin dimer on leukocytes, has been reported to accelerate macrophage efflux from an inflammatory site (6) and regulate neutrophil detachment in vitro (7). However, specific steps in leukocyte transendothelial migration modulated by ectodomain shedding have not been determined.

Proteases responsible for shedding of a large number of cell-surface proteins involved in leukocyte transendothelial migration on both leukocytes and endothelial cells are zinc-dependent endopeptidases composed of a family of matrix metalloproteinases (MMPs), which include secreted and membrane types (MT-MMPs), and the ADAM (a disintegrin and metalloproteinase) family of transmembrane metalloproteinases (4, 5). Analyses of specific enzymes have identified some key proteolytic cleavage events, but most studies have focused on regulation of neutrophil and lymphocyte transendothelial migration. For example, knockdown of ADAM10 on T cells, HUVECs, or both impairs T cell transendothelial migration in vitro (8). Although VE-cadherin was identified as a pos-
possible endothelial target of ADAM10, the relative role of VE-cadherin cleavage was not determined (8). Neutrophil transendothelial migration is inhibited by soluble forms of junctional adhesion molecule (JAM)-A, which is shed from endothelium by ADAM10 and ADAM17 (9). Recently, ADAM17 shedding of L-selectin (CD62L) was shown to limit neutrophil recruitment, but monocyte emigration is independent of CD62L shedding (10). A more limited number of studies have examined monocyte transendothelial migration. An Ab to MT1-MMP impairs transendothelial migration of human monocytes but only when HUVECs are preactivated and MCP-1 is present (11). Although ICAM-1 has been proposed as a possible substrate (12), the step(s) targeted by MT1-MMP is still unclear. Thus, there is a need to evaluate particular steps targeted by regulated proteolysis during transendothelial migration and to better define the enzymes and substrates involved.

In this report, we identify the metalloproteinase-regulated mechanisms underlying transendothelial migration used by monocytes, cells of central importance in the outcome of acute, and chronic inflammatory disease pathogenesis. In the presence of GM6001, which blocks metalloproteinases expressed on both monocytes and endothelial cells, we show that diapedesis is significantly delayed without any marked effect on adhesion or locomotion across the endothelial surface. To interrogate possible substrates whose impaired cleavage contributes to the delay of diapedesis, we separately screened monocytes and endothelial cells for adhesion molecules whose cell surface expression was elevated by GM6001 following their coinubcation, which mimics transendothelial migration. Despite the presence of multiple substrates on both cells, only monocyte surface levels of Mac-1 increase upon coinubcation with GM6001. We further show that ADAM10 and ADAM17 are responsible for metalloproteinase-dependent shedding of Mac-1 and that absence of monocyte ADAM17 leads to a significant prolongation of diapedesis. In contrast, depletion of monocyte ADAM10, or endothelial ADAM10 or ADAM17, does not affect diapedesis. Therefore, we demonstrate that monocyte ADAM17 promotes diapedesis and suggest that ADAM17 cleavage of Mac-1 may serve as a regulatory mechanism.

Materials and Methods

Human endothelial cell culture and monocyte isolation

HUVECs (Cascade Biologics–Invitrogen) were cultured in gelatin-coated flasks using M199 supplemented with 20% FCS, EGM-2 SingleQuots (Lonza), and antibiotics and used up to passage 3. Human PBMCs were freshly isolated from citrate anticoagulated whole blood of healthy donors by Ficoll–Paque Plus (Amersham Biosciences) separation and subjected to enrichment for monocytes by negative selection using the Monocyte Isolation Kit II (Miltenyi Biotec) plus biotinylated Abs against CD42b (Genetex) in some experiments to remove platelets (13). Enriched monocytes (>90% estimated by FACS analysis) were resuspended in assay medium (phenol red-free M199 with 20% FCS and 20 mM HEPES [pH 8]) and kept on ice unless specified. Metalloproteinase activity was blocked by a 30-min preincubation with 50 μM GM6001 (Elastin Products) and 0.1% DMSO as a control, unless specified otherwise. Because GM6001 blockade is reversible, all subsequent experiments also included inhibitor or control. The University of Washington Human Subjects Review Committee has approved all protocols.

Transwell assays

HUVECs were seeded at 3.0 × 104 cells/well onto Costar Transwell filters (3-μm pore, 6.5-mm diameter) previously coated with 5 μg/ml human fibronectin and grown to confluence. PBMCs (3.0 × 105 cells/100 μl) were labeled with calcein-AM, and migrated leukocytes in the lower well and bottom filter were counted using ImageJ software (National Institutes of Health).

Static assay of monocyte transendothelial migration

The in vitro system previously reported to model monocyte transendothelial migration under static conditions using confluent HUVEC monolayers grown on type I collagen gels (14) was modified to enable simultaneous recording of multiple conditions. Briefly, flat collagen gels (∼0.12-mm thickness) were prepared in 4-well silicon chamber (9 mm diameter, ∼3-mm depth), which is made in a 40-mm glass-bottom dish (Wilco Wells) in combination with Secure-Seal Adhesive Spacer and Press-to-Seal Silicon Isolator (Molecular Probes). Detailed methodology is available upon request. The collagen gels were coated with human plasma fibronectin, and HUVECs were plated and grown to confluence. After 30 min of equilibration at 37°C and 30 min of pretreatment with DMSO or GM6001, both in assay medium, experiments were started by addition of a monocyte suspension (50 μl droplet/well, 3–4 × 105 cells) to the HUVECs (100 μl/well). Multiple fields were randomly chosen so that at least 25 diapedesis events/condition could be monitored. The recording was started after 20 min for unactivated HUVECs and 10 min for TNF-activated HUVECs. Non-migrating leukocytes were identified by Ficoll–Paque Plus (Amersham Biosciences) separation and subjected to flow analysis or splenic leukocytes that ultimately underwent diapedesis, and monocytes undergoing transmigration at impaired endothelial junctions were excluded.

Flow assay of monocyte transendothelial migration

Analysis of transendothelial migration under shear flow was performed as described previously (15). Briefly, HUVECs were grown to confluence on 25-mm-diameter fibronectin-coated glass coverslips and preactivated for 4 h with 10 ng/ml TNF-α, followed by preincubation with DMSO (0.1%) or 50 μM GM6001 for 30 min at 37°C. The prepared coverslip was inserted into the in vitro flow device as described previously. Platelet-free monocytes were kept at 8°C and used within 6 h of preparation. Preincubation of monocytes with DMSO or GM6001 at 37°C for 5 min to minimize the loss of CD62L, which is essential for adhesion under flow conditions. The monocytes were then introduced into the flow chamber as a bolus (105 cells/100 μl), followed by drawing them across the HUVEC monolayer with a flow rate estimated to be 0.5 dyn/cm2 for 30 min in flow medium (PBS with 0.75 mM Ca2+ and Mg2+ and 125 mM NaCl) at 37°C. The degree of shear stress used induces maximal adhesion of monocytes at the beginning of the assay. Acquisition of time-lapse ×20 differential interface contrast images was started when ~10 monocytes were adherent in the monitored field, and images were collected every 10 s thereafter.

Analysis or monocyte migration captured by time-lapse video microscopy

Monocyte migration on endothelium was analyzed in a blinded manner by two separate investigators using ImageJ with MTtrackJ plug-in software (E. Meijering, Biomedical Imaging Group Rotterdam and the Eramus MC–University Medical Center, Rotterdam, The Netherlands). Monocyte cell bodies were traced on the time-lapse images by moving the MTtrackJ pointer when they displaced more than their radii. Analyses were done only on monocytes that ultimately underwent diapedesis, and monocytes undergoing transmigration at impaired endothelial junctions were excluded.

Flow cytometry-based screening of metalloproteinase substrates on coincubated endothelial cells and monocytes

HUVECs were grown to confluence in human plasma fibronectin (5 μg/ml)-coated 10-cm dishes and stained for 10 min with 1 μM CellTracker Green CMFDA (Molecular Probes) 16 h prior to use. Monolayers were stimulated for 4 h with TNF-α at 10 ng/ml and subjected to preincubation with DMSO or GM6001 for 20 min. Freshly prepared PBMCs (107/ml) were similarly preincubated and then coincubated with the HUVEC monolayers in the presence of DMSO or GM6001 for 2 h. The cocultures were washed three times to remove nonadherent cells and collected with ice-cold PBS supplemented with 0.1% BSA and 10 mM EDTA (1 ml/dish) using cell lifters. Following incubation with FcR blocking reagent (Miltenyi Biotec), cell suspensions were stained at 4°C for 30 min with anti-CD14 and saturating amounts of PE-labeled Abs for candidate molecules (Supplemental Table I). Simultaneously, cells incubated separately for 2 h were collected and stained as described above as well as with the same antibody combinations. All stained cells were analyzed on FACSscan (BD Biosciences), and flow data were analyzed using FlowJo 8.4 software (Tree Star).

Knockdown of ADAM10 and ADAM17 in primary monocytes

Two reports small interfering RNA (siRNA) sequences for knockdown of ADAM10 and ADAM17 were synthesized by Ambion, and those with the
best downregulation of target ADAMs were used for the experiments shown: ADAM10-construct, 5′-AGA CAU UAU GAA GGA UUA U3′ (16), and ADAM17-construct, 5′-GCU UGC UUC UUU GUU CUC A3′ (17). A nontargeting siRNA (Silencer Negative Control number 1; Ambion) was used as control.

Platelet-free primary monocytes (5 × 10⁶) were transfected with siRNA (300 nM siRNA/reaction) using a Nucleofector II Kit and a Human Monocyte Nucleofector Kit (Amaxa). To achieve maximal knockdown, monocytes were allowed to recover at 37°C for 16 h in IMDM (Lonza) containing 20% human plasma–derived serum (or FCS to evaluate soluble CD18 dimers). Monocytes to be used for transmigration were incubated in a 50-mL conical tube on an orbital shaker to give an estimated shear of 7.5 dyn/cm² to maintain monocyte migratory properties as determined in pilot experiments. Monocytes were subjected to an Annexin V–based Dead Cell assay and then grown to confluence on fibronectin-coated porous Transwell inserts to assess migration in the presence of 50 μM GM6001 or 0.1% DMSO (vehicle control). Before coincubation, the HUVEC monolayers and leukocytes were separately preincubated for 30 min with vehicle or GM6001. HUVECs were unactivated (A) or preactivated with 0.1 ng/mL TNF-α for 4 h (B). Representative data from at least three different experiments are shown. (C) Monocyte migration across preactivated HUVEC monolayers (10 ng/mL TNF-α) was determined by FACS analysis for mononuclear cell transendothelial migration as shown in (B). Monocyte numbers in the presence of GM6001 are expressed as ratios relative to those of DMSO.

The effects of GM6001 on monocyte transendothelial migration were also tested under flow conditions that more closely model physiological conditions. Monocytes were drawn across a TNF-α–activated endothelial layer for 30 min in medium including vehicle or GM6001 at an estimated flow rate of 0.5 dyn/cm², conditions that induce maximal initial adhesion to allow evaluation of post-adhesion events (15). In vehicle control (Supplemental Video 1), monocytes rapidly adhere, spreading on the activated endothelium followed by locomotion for a relatively short distance to reach endothelial junctions. Monocytes then undergo diapedesis, thereby finishing their paracellular transendothelial migration, which is particularly easy to distinguish under flow conditions. It takes monocytes ~10 min to complete all of these steps (data not shown). However, in the presence of GM6001 (Supplemental Video 2), monocyte tails remain on the apical surface of endothelial monolayers for longer durations until they complete transendothelial migration (Fig. 2B), and GM6001 delays the completion of transendothelial migration under flow conditions as observed under static conditions (Fig. 2A). Taken together, our data indicate that metalloproteinase activity is rate limiting and is required for optimal monocyte transendothelial migration.

Statistical analysis

For analysis of two groups within a single experiment of time-lapse microscopy using monocytes from a single donor, unpaired *t* tests were performed after data were confirmed to fulfill the criteria. Otherwise, Mann–Whitney *U* tests were applied. In comparison of means or medians obtained from three or four experiments, paired *t* tests were performed. To validate significance for fold differences, paired *t* test was performed after log transformation. All statistical analyses were performed two-sided using InStat (GraphPad), and *p < 0.05* was considered significant.

Results

A broad-spectrum metalloproteinase inhibitor impairs transendothelial migration of monocytes

We put forward the hypothesis that proteolytic shedding of cell surface proteins provides a mechanism to aid in the rapid transition of cells between different steps and to coordinate the complex, multistep process of leukocyte recruitment to inflammatory sites (4). To test this hypothesis, we examined the impact of a broad-spectrum, zinc-dependent metalloproteinase inhibitor, GM6001, on net migration of human PBMCs across HUVEC monolayers using a Transwell assay. As shown in Fig. 1A, GM6001 impairs net migration of leukocytes across unactivated endothelium to 40% of vehicle control. When endothelial cells are preactivated by the inflammatory stimulant TNF-α (0.1 ng/mL for 4–5 h), GM6001 reduces net leukocyte migration to a similar extent (Fig. 1B). We confirmed that monocytes are the major cell type that migrates across endothelial monolayers under all conditions tested (>85% as estimated by FACS analysis; data not shown), and quantification of net monocyte migration shows that GM6001 impairs migration by 70% (Fig. 1C).

Metalloproteinase blockade delays completion of monocyte transendothelial migration under both static and flow conditions

Time-lapse video microscopy was then used to monitor the interactions of purified monocytes with endothelial cells to dissect at which step(s) GM6001 is acting in transendothelial migration. To assess migration under static conditions, unactivated endothelial monolayers on collagen gels were coincubated with primary monocytes (18), an in vitro assay in which key mediators of transmigration have subsequently been confirmed in vivo (19–23), and almost all leukocytes cross the endothelial cells at the cell borders, undergoing paracellular migration (24). As shown in Fig. 2A, GM6001 postpones the frequency of monocyte completion of transendothelial migration, the point at which remnants of monocyte cell body disappear from the apical surface of endothelial monolayers in the process of diapedesis. Because the total frequency of transendothelial migration in the presence of GM6001 is comparable to vehicle control (Fig. 2A), GM6001 does not inhibit transendothelial migration. Rather, GM6001 delays monocyte transendothelial migration under static conditions.

Metalloproteinase activity facilitates progression of diapedesis but not other steps of monocyte transendothelial migration

Because the process of monocyte transendothelial migration involves multiple steps, each was quantified to identify the metalloproteinase–targeted step(s). As summarized in Table I, GM6001 does not affect the initial adhesion or locomotion across the endothelium. We also verified that GM6001 does not alter endothelial junctions during the period of the migration assay as compared with vehicle control (data not shown). To analyze diapedesis, a frame-by-frame analysis of vehicle control and GM6001 conditions was performed on unactivated endothelial monolayers under static conditions (Fig. 3A). In vehicle control (Fig. 3A, top panel; Supplemental Video 3) monocytes first squeeze a membrane protrusion, and then their main cell body, between the endothelial junctions, and within a short time their tails retract.
Metalloproteinase blockade by GM6001 delays monocyte transendothelial migration under static and flow conditions. (A) Human monocytes were added to unactivated HUVEC monolayers in the presence of DMSO or GM6001 under static conditions and observed using time-lapse video microscopy every 37.8 s. Individual monocytes migrating on the apical surface of the monolayers were tracked until they complete migration across the monolayers. Monocytes that complete their transendothelial migration up to each time point are expressed as percentage of total monocytes in the field. In one experiment, an average of 50 monocytes were evaluated for DMSO and 46 for GM6001. (B) Monocyte transendothelial migration under flow conditions in the presence of DMSO (Supplemental Video 1) or GM6001 (Supplemental Video 2) was analyzed as described in (A). Monocytes were drawn across preactivated HUVEC monolayers (10 ng/ml TNF-α for 4–6 h) at 0.5 dyn/cm² in flow medium containing DMSO or GM6001. Pretreatment with DMSO or GM6001 was 30 min for HUVECs and 5 min for monocytes. Time-lapse video microscopy was performed every 10 s for 30 min. Means ± SD of three different experiments are shown. *p < 0.05, **p < 0.01, ***p < 0.005 (paired t test).

from the apical surface into the junctions. In contrast, monocytes in the presence of GM6001 (Fig. 3A, bottom panels, Supplemental Video 4) take a longer time to squeeze their cell body between endothelial junctions and their remnant (tail) remains on the apical surface for extended periods while the main cell body passes through the endothelial monolayer. We quantified the duration of diapedesis as the time interval between the arrival of monocytes at the site of diapedesis and their completion of diapedesis as assessed from the apical surface. The distribution of diapedesis duration shows that under control conditions ~90% of monocytes complete diapedesis within 20 min, whereas only 40% do in the presence of GM6001 (Fig. 3B). Comparison of diapedesis duration for monocytes prepared from different donors reveals that GM6001 prolongs diapedesis 2-fold as compared with control conditions (Fig. 3C, left). When the endothelium was preactivated with TNF-α, similar results were obtained with a 1.6-fold increase in the duration of diapedesis by GM6001 (Fig. 3C, right). Under physiological flow conditions, GM6001 results in 2-fold increase even though the diapedesis duration in vehicle control is much shorter than observed under static conditions (Fig. 3D). Taken together, these data demonstrate that prolongation of diapedesis by metalloproteinase blockade is responsible for the delay in completion of monocyte transendothelial migration.

Monocytes shed CD18 integrin dimers in a metalloproteinase-dependent manner upon coincubation with endothelial cells and CD11b/CD18 is cleaved by transmembrane proteases ADAM17 and ADAM10.

Diapedesis is dependent upon both endothelial and monocyte adhesion molecules, several of which have been shown to be shed by metalloproteinases, including leukocyte-specific integrin CD18 (6, 25) and its partner α integrins (CD11a, CD11b (6), CD11c and CD11d), ICAM-1 (12, 26), activated leukocyte cell adhesion molecule (ALCAM) (27), platelet/endothelial cell adhesion molecule (PECAM)-1 (28), and JAM-A (9). To identify specific molecules(s) involved in metalloproteinase-mediated regulation of diapedesis, we initially screened these adhesion molecules by flow cytometry to determine whether monocyte surface levels were increased in the presence of GM6001 under the same conditions used for evaluation of transendothelial migration (Fig. 4A, 4B, Supplemental Table II). CD14⁺ monocytes were evaluated following PBMC incubation with or without TNF-activated HUVEC monolayers in the presence of vehicle and GM6001 for 2 h, a time period during which the majority of monocytes complete transendothelial migration (Fig. 2A). As highlighted in Fig. 4A and 4B, only surface levels of CD11b are upregulated (~1.4-fold) following coincubation with HUVECs in the presence of GM6001 (Fig. 4B). ICAM-1 levels were also increased by GM6001 treatment but both with and without HUVEC coincubation (Supplemental Table II). Monocyte ICAM-1 does not seem relevant to transendothelial migration because monocyte-targeted functional blockade of ICAM-1 did not significantly affect net Transwell migration of mononuclear cells (data not shown). CD11a, ALCAM-1, PECAM-1, and JAM-A levels were not altered by GM6001 (Fig. 4A, Supplemental Table II). Taken together, these data demonstrate a metalloproteinase-dependent and selective increase in CD11b/CD18 levels on monocytes upon coincubation with endothelial cells.

To confirm that CD18 is shed from primary monocytes in a metalloproteinase-dependent manner, we analyzed cell lysates and
conditioned media (CM) prepared from monocytes treated with GM6001 for 16 h. Soluble integrin CD18 is decreased in 16-h CM with GM6001 treatment as determined by Western blot analysis (Fig. 4C), and this is associated with increased integrin CD18 in cell lysates. The modest difference in m.w. of soluble and cellular integrin CD18 is similar to previous reports (6, 7, 29), indicating that metalloproteinase cleavage within the ectodomain close to the cell membrane results in loss of the transmembrane and the small cytoplasmic domains. The relatively small increase in cellular CD18 with GM6001 is consistent with significant intracellular stores of CD18 (30). Further analysis of monocyte CMs demonstrates that both CD11b/CD18 (Mac-1) and CD11a/CD18 (LFA-1) are shed by metalloproteinases, but the amount of shed Mac-1 was ∼7.8 times higher than that of LFA-1, suggesting a difference in the extent of constitutive shedding (Fig. 4D).

Because shedding of a number of cell surface adhesion molecules can be mediated by the constitutively expressed transmembrane proteases ADAM10 and ADAM17 (4, 5), we tested whether one or both of these enzymes contribute to cleavage of LFA-1 and Mac-1. Primary monocytes were transfected with siRNAs to deplete ADAM17 or ADAM10 using the Nucleofector system, and maximal knockdown was achieved 16 h after transfection (Fig. 5A) with a transfection efficiency >95% (data not shown) and comparable viabilities (~75%; data not shown). Although levels of soluble LFA-1 in CM collected from posttransfection culture (16 h) are not altered by knockdown of ADAM10 or ADAM17, soluble Mac-1 levels are reduced 20 and 30% by ADAM10 and ADAM17 siRNA, respectively, as compared with control-transfected cells (Fig. 5B). Thus, both monocyte ADAM17 and ADAM10 can contribute to ectodomain shedding of Mac-1.

Candidate endothelial substrates are not altered by metalloproteinase blockade and neither endothelial ADAM10 nor ADAM17 promotes monocyte diapedesis

Because endothelial junctions may serve as gatekeepers for leukocyte transmigration (31), we also assessed changes in surface levels of endothelial candidate substrates during monocyte transendothelial migration in the presence of GM6001 (Supplemental Table III) as shown in the analysis of monocyte surface adhesion molecules (Supplemental Table II). Four candidate substrates were screened, including VCAM-1 (32), ICAM-1 (12, 26), ALCAM (27), and JAM-A (9). However, GM6001 treatment has no effect on surface levels of any of the four candidate substrates, with or without coincubation (Supplemental Table III).
Diapedesis duration was also assessed following separate knockdown of endothelial ADAM10 and ADAM17 from HUVECs using siRNA (Supplemental Fig. 1). Depletion of ADAM10 consistently leads to a significant decrease in endothelial–junctional permeability (Supplemental Fig. 1C), which agrees with a previous report (8). However, neither ADAM10 nor ADAM17 depletion from HUVECs reproduces GM6001-mediated prolongation of monocyte diapedesis (Supplemental Fig. 1A, 1B). Taken together, our data demonstrate that cleavage of metalloproteinase substrates on endothelial cells and endothelial ADAM10 and ADAM17 proteolytic activity are not major contributors to metalloproteinase-mediated facilitation of monocyte diapedesis.

**ADAM17 expressed on monocytes, but not ADAM10, promotes their diapedesis**

To determine the monocyte metalloproteinases that contribute to the regulation of diapedesis, we first analyzed MT1-MMP previously reported to promote monocyte transmigration on activated endothelium based on partial inhibition by an MT1-MMP Ab, but only if MCP-1 was present as a chemoattractant (11). Using the same Ab, we observed no effect on monocyte transmigration under our conditions lacking MCP-1 (data not shown). Thus, MT1-MMP does not appear to be a major regulatory mechanism for diapedesis. We then used siRNA knockdown to evaluate whether monocyte ADAM17 and/or ADAM10 are involved in metalloproteinase-mediated regulation of diapedesis. GM6001-dependent prolongation of diapedesis was verified for control-transfected monocytes (Fig. 6A), and a similar extent of inhibition (1.5-fold delay) to that with freshly prepared monocytes (compare with Fig. 2C) was observed. In contrast, depletion of ADAM17 (Supplemental Video 6), but not ADAM10 (Fig. 6B), extends the duration of monocyte diapedesis compared with control-transfected monocytes (Supplemental Video 5), although the extent of inhibition (1.3-fold) was less than that observed with GM6001 (Fig. 6A). Taken together, these data demonstrate that monocyte ADAM17, but not monocyte ADAM10, makes a significant contribution to metalloproteinase-dependent promotion of diapedesis.

**Discussion**

Although metalloproteinase-mediated “ectodomain shedding” has been implicated in the regulation of leukocyte trafficking (4–9), the steps targeted, enzymes responsible, and key substrates are still unclear, particularly for monocytes. The present study shifts the paradigms regarding metalloproteinase regulation of monocyte
Duration of monocyte diapedesis was determined for monocytes transfected with control or transfected with ADAM10 or ADAM17 siRNA. Means ± SD from three different experiments are shown. All monocyte preparations show comparable viability (>95% as determined by trypan blue exclusion). *p < 0.01 (paired t test), **p < 0.0005 (unpaired t test with Welch correction; n = 19, 37, and 27 monocytes for untransfected, DMSO, and GM6001, respectively).

FIGURE 6. Depletion of ADAM17, but not ADAM10, from monocytes prolongs the diapedesis step. (A) Untransfected monocytes and monocytes transfected with control siRNA were generated as described in Materials and Methods and subjected to time-lapse video analysis on TNF-activated endothelium in the presence of DMSO or GM6001. Representative data from two independent experiments are expressed as means ± SEM. (B) Duration of monocyte diapedesis was determined for monocytes transfected with control, ADAM10, or ADAM17 siRNA. Means ± SD from three different experiments are shown. All monocyte preparations show compatible viability (>95% as determined by trypan blue exclusion). *p < 0.01 (paired t test), **p < 0.0005 (unpaired t test with Welch correction; n = 19, 37, and 27 monocytes for untransfected, DMSO, and GM6001, respectively).

transendothelial cell migration at several levels. First, to our knowledge, we demonstrate for the first time that metalloproteinases promote monocyte transendothelial migration through facilitation of the diapedesis step. Second, our studies reveal that blockade of metalloproteinases elevates monocyte surface Mac-1 (CD11b/CD18) levels specifically upon coincubation with activated endothelial cells, conditions that mimic transendothelial cell migration. Furthermore, despite previous documentation of metalloproteinase-mediated shedding of multiple other adhesion molecules on both monocytes and endothelial cells, none of these shows elevated surface levels dependent on monocyte–endothelial coincubation and metalloproteinases. Third, we show two enzymes, ADAM10 and ADAM17, contribute to metalloproteinase-dependent constitutive shedding of Mac-1. Finally, depletion of monocyte ADAM17 significantly impairs the diapedesis step, whereas neither monocyte ADAM10 nor endothelial ADAM10 or ADAM17 are involved in diapedesis regulation.

Our study highlights the role of metalloproteinases, specifically monocyte ADAM17, in regulating the rate of diapedesis. The delay in diapedesis by metalloproteinase blockade is a form of regulation distinct from total blockade of the process as previously shown with blocking Abs to PECAM-1 (18) and CD99 (21). As our time-lapse imaging shows, the main cell body of the monocyte migrates into the endothelial junction, even in the presence of GM6001. Thus, blocking metalloproteinase-mediated cleavage does not impair initiation of diapedesis and functions related to movement of the cell body into the junction between endothelial cells. The marked effect of GM6001 treatment is that monocyte tails are persistent on the apical surface of endothelium, even after the main cell body is spreading in the subendothelial space. Therefore, we suggest that metalloproteinase inhibition impairs organized retraction of the monocyte cell body from the endothelial apical surface and thus diapedesis. The greater inhibitory efficacy of GM6001 shown in the Transwell assay as compared with apical video microscopy (Figs. 1, 3) further suggests that postdiapedesis mechanisms may additionally be targeted by metalloproteinases. Interesting possibilities for future studies include new steps that have been recently defined beneath the endothelium for leukocyte emigration into tissues in vivo in which uropod release from the endothelial basal surface (33) is followed by cell crawling and penetration of vascular basement membranes and pericyte sheaths (34).

A surprising finding in our study is the failure to observe any increase in endothelial surface expression of previously demonstrated substrates of metalloproteinases, including those of ADAM10 and ADAM17, upon coincubation of monocytes and endothelial cells in the presence of GM6001. Coincubation was limited to 2 h to focus on cleavage events during transendothelial migration. Using a similar detection system, Koenen et al. (9) showed that enhanced shedding of endothelial JAM-A upon 2-h coincubation with neutrophils is associated with significant decrease in their endothelial surface levels. Consistent with our inability to detect surface changes in endothelial ADAM substrates, which could contribute to the regulation of diapedesis, endothelial knockdown of ADAM10 or ADAM17 had no effect on the rate of diapedesis. These data are in contrast to previous reports that endothelial ADAM10 regulates T cell transendothelial migration, although the specific step targeted by ADAM10 was not determined (8). Endothelial ADAM10 and ADAM17 also control neutrophil transendothelial migration through generation of soluble JAM-A (9). These data strongly suggest that transendothelial migration of different leukocytes may be regulated by distinct mechanisms.

In contrast, analysis of expression levels of candidate monocyte substrates under conditions comparable to transendothelial migration identified a selective increase in surface levels of the integrin dimer Mac-1 (CD11b/CD18). Mac-1 has been shown to be involved in diapedesis (35) as well as other steps of leukocyte transendothelial migration (1). We were able to confirm that the broad inhibitor GM6001, as well as knockdown of ADAM17 and ADAM10, decreased shedding of soluble Mac-1. To our knowledge, this is the first report of ADAM-mediated cleavage of leukocyte integrins.

Our data further suggest that ADAM17 cleavage of Mac-1 may play a role in facilitation of monocyte diapedesis because knockdown of monocyte ADAM17 impairs diapedesis (Fig. 6B). Under normal conditions, soluble Mac-1 may also contribute to the facilitation of diapedesis because it retains its ability to bind to its endothelial ligand, ICAM-1 (6, 7). Interaction of leukocytes and endothelial cells triggers multiple signaling cascades and activation of leukocyte integrins (36). Because we have shown that integrin activation is not sufficient to induce cleavage of Mac-1 (6), we speculate that monocyte–endothelial contact may activate monocyte ADAM17 or promote its binding to Mac-1 and thus enhance temporally and spatially regulated cleavage. The dissociation of Mac-1/ICAM-1 interactions could then facilitate monocyte retraction from the apical side of endothelium, thereby achieving rapid progression of diapedesis (22). However, because Mac-1 functional blocking Abs impairs locomotion and thus prevent paracellular diapedesis (22), it is not possible to use this approach to define the contribution of Mac-1 cleavage to progression of monocyte diapedesis. Alternate strategies, such as uncleavable mutants, will be needed to eliminate the possible contribution of other effects of ADAM17 knockdown and to provide a direct link between ADAM17 cleavage of Mac-1 and facilitation of diapedesis. It also remains possible that additional metalloproteinases may be involved because GM6001 resulted in a greater suppression of diapedesis and soluble integrin CD18 dimer release than those following siRNA knockdown of ADAM17. Shedding of additional substrates not included in our screen may also contribute to the regulation of monocyte diapedesis.

Interestingly, depletion of monocyte ADAM10 did not alter diapedesis, suggesting that ADAM10-mediated cleavage of Mac-1 is not enhanced by interaction with endothelial cells. Indeed, it is
known that constitutive shedding by ADAM17 and ADAM10 can be enhanced through distinct stimuli and signaling pathways (37, 38). Previous experiments that defined the monocyte locomotion step also demonstrated a requirement for both monocyte CD11a/CD18 and CD11b/CD18 interaction with endothelial ICAM-1 and ICAM-2 (22). Because endothelial ICAM-1 has been identified as a substrate of ADAM17 (26) and MT1-MMP (12), inhibition of ICAM-1 cleavage has been suggested as a mechanism to regulate its interaction with CD18 integrins capable of altering the locomotion step. However, our inability to detect any effect of metalloproteinase blockade on the locomotion step, despite demonstrating metalloproteinase-dependent cleavage of both CD11a/CD18 and CD11b/CD18, suggests that other mechanisms to modulate their ICAM interactions, such as alteration of integrin affinity (39), may be more important for transitioning between locomotion and diapedesis.

Potential in vivo relevance for our prolongation of in vitro monocyte diapedesis is suggested by intriguing similarities with data from a recent study of in vivo neutrophil diapedesis. Woodfin et al. (40) described two modes of abnormal neutrophil diapedesis in ischemia–reperfusion stimulation cremaster venules: resistant diapedesis and reverse diapedesis. In resistant diapedesis, neutrophils move back and forth at endothelial junctions before finally completing migration into the subendothelial space as we observed with monocyte diapedesis in the presence of GM6001, most clearly seen under flow conditions (Supplemental Video 2). In vivo, neutrophils show delayed diapedesis similar to our studies. The delayed diapedesis of neutrophils is caused by relocalization of JAM-C to the venular surface from endothelial junctions (40), which is triggered by ischemia–reperfusion. JAM-C has been reported to mediate monocyte diapedesis in kidney models of ischemia–reperfusion (41) and to maintain monocyte unidirectional diapedesis in vitro (42), suggesting that monocytes may show delayed diapedesis in specific organs and conditions. Interestingly, JAM-C controls diapedesis through interaction with Mac-1 (35, 43, 44). Thus, it is possible that metalloproteinase cleavage of Mac-1 may be involved in JAM-C–regulated unidirectional diapedesis. Although we observed no delay in vivo in monocyte accumulation in the peritoneal cavity 24 h after injection of the sterile irritant thioglycollate in mice lacking ADAM17 in circulating cells (10), the potential role of Mac-1 and JAM-C has not been investigated in a 30% decrease in monocyte accumulation 48 h after thioglycollate seen in the same mice (J. Tang, C.L. Wilson, and E.W. Raines, unpublished observations). However, as discussed above, alternate approaches such as generation of uncleavable mutants are needed to address this question.

In conclusion, this investigation demonstrates that metalloproteinase activity promotes the diapedesis step of monocyte transendothelial migration under both static and physiological flow conditions in vitro by increasing the rate of diapedesis. During transendothelial migration, metalloproteinase activity leads to cleavage of monocyte Mac-1 but not other reported adhesion substrates expressed on monocytes and endothelial cells. Knockdown of monocyte ADAM17 impairs Mac-1 shedding and diapedesis, suggesting that the metalloproteinase-mediated promotion of diapedesis may involve monocyte ADAM17 cleavage of Mac-1. Because diapedesis is a central component of leukocyte recruitment, our study establishes cell-type specific regulatory mechanisms of diapedesis, which are critical for the development of cell-targeted therapy in chronic inflammatory diseases such as atherosclerosis.

Acknowledgments

We thank University of Washington colleagues Brian Fish (Nutrition Obe-

sity Research Center), Roderick Theobald and Chris Jordan-Squire (Depart-

ment of Statistics), and Nancy Temkin (Department of Biostatistics) for

statistical evaluation of the experiments. We thank Li-Chuan Huang for excellent technical assistance. We also thank Francis W. Luscinskas for help in establishing the flow assays and insightful discussions and José López and Junmei Chen (Puget Sound Blood Center) for assistance with the flow assays. We thank John M. Harlan, Carole L. Wilson, and Jingjing Tang for providing helpful comments and suggestions during the preparation of the manuscript.

Disclosures

The authors have no financial conflicts of interest.

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