Platelet-derived S100 family member myeloid-related protein-14 regulates thrombosis

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Expression of the gene encoding the S100 calcium–modulated protein family member MRP-14 (also known as S100A9) is elevated in platelets from patients presenting with acute myocardial infarction (MI) compared with those from patients with stable coronary artery disease; however, a causal role for MRP-14 in acute coronary syndromes has not been established. Here, using multiple models of vascular injury, we found that time to arterial thrombotic occlusion was markedly prolonged in Mrp14−/− mice. We observed that MRP-14 and MRP-8/ MRp-14 heterodimers (S100A8/A9) are expressed in and secreted by platelets from WT mice and that thrombus formation was reduced in whole blood from Mrp14−/− mice. Infusion of WT platelets, purified MRP-14, or purified MRP-8/MRP-14 heterodimers into Mrp14−/− mice decreased the time to carotid artery occlusion after injury, indicating that platelet-derived MRP-14 directly regulates thrombosis. In contrast, infusion of purified MRP-14 into mice deficient for both MRP-14 and CD36 failed to reduce carotid occlusion times, indicating that CD36 is required for MRP-14–dependent thrombosis. Our data identify a molecular pathway of thrombosis that involves platelet MRP-14 and CD36 and suggest that targeting MRP-14 has potential for treating atherothrombotic disorders, including MI and stroke.

Introduction

Acute myocardial infarction (MI) commonly results from the atherosclerotic plaque disruption (1) and thrombosis that cause coronary artery occlusion (2). Angioscopic (3) and pathological (1) observations indicate that platelets constitute a major component of such thrombi, yet the precise platelet-related molecular events that immediately precede acute MI remain uncertain. Our previous study used a transcriptional profiling strategy to identify novel regulators of vascular inflammation and atherothrombosis by examining platelet mRNA transcripts that are differentially expressed in patients with ST-segment elevation myocardial infarction (STEMI) compared with those with stable coronary artery disease (4). Myeloid-related protein-14 (MRP-14, also referred to as S100A9) was one of the strongest predictors of STEMI that arose from the transcriptional profiling analysis. MRP-14, a member of the S100 family of calcium-modulated proteins, complexes with MRP-8 (S100A8), and together the MRP-8/14 heterodimer regulates myeloid cell function by modulating calcium signaling (5) and cytoskeletal reorganization (6), by operating as a chemoattractant (7), and by binding to cell surface receptors, including CD36 (8), toll-like receptor 4 (TLR4) (9), and receptor for advanced glycation end products (RAGE) (10). In two prospective, nested case-control studies, one in apparently healthy postmenopausal women (4) and the other in patients presenting with acute coronary syndromes (11), elevated plasma levels of MRP-8/14 predicted the risk of future cardiovascular events, independent of traditional cardiovascular risk factors and highsensitivity C-reactive protein (hs-CRP). Elevated plasma levels of MRP-8/14 also serve as an early and sensitive marker of myocardial necrosis in the setting of chest pain (12).

Despite the presence of Mrp8 mRNA transcripts, Mrp14−/− mice lack both MRP-8 and MRP-14 protein, possibly due to the instability of MRP-8 protein in the absence of MRP-14 (5, 13). In vitro studies with Mrp14−/− neutrophils showed markedly diminished migration through endothelial monolayers and attenuated chemokinesis in a three-dimensional collagen matrix (13). An essential role for MRP-8/14 in leukocyte recruitment in vivo is supported by evidence that Mrp14−/− mice have diminished granulocyte recruitment during tissue wound healing (6) and acute pancreatitis (14). Recent studies using Mrp14−/− mice have also demonstrated that MRP-8 and MRP-14 play a regulatory role in endotoxin-induced phagocyte function by binding to TLR4 and promoting myeloid MyD88–dependent activation of NF-κB (9). Last, we have reported that MRP-8/14 broadly regulates vascular inflammation and contributes to the biological response to vascular injury in murine models of atherosclerosis, vasculitis, and restenosis by promoting leukocyte recruitment (15).

Despite our identification of Mrp14 transcripts in platelets, freshly isolated human bone marrow megakaryocytes, and megakaryocytes generated in vitro by differentiation of human CD34-positive cells (4), the role of MRP-8/14 in thrombosis and hemostasis is unknown. In this study, we provide evidence that platelet-derived MRP-14 directly modulates platelet function...
and thrombosis without influence on tail bleeding time or other hemostatic parameters.

Results

Photochemical injury–induced arterial thrombosis is delayed in Mrp14−/− mice. To elucidate the effect of MRP-8/14 on the development of arterial thrombosis in real time, carotid arteries of WT and Mrp14−/− mice were subjected to the Rose Bengal model of thrombosis, an endothelial cell photochemical injury model caused by local free-radical release (16, 17). We then continuously monitored carotid artery blood flow with a vascular flow probe. Mean time to occlusive thrombus formation in WT mice was 27.1 ± 5.9 minutes and was significantly prolonged in Mrp14−/− mice to 46.6 ± 22.9 minutes (n = 16 per group, P = 0.004) (Figure 1A). We harvested carotid arteries 25 minutes after photochemical injury for histological analysis. Both perfusion and nonperfusion fixation techniques were used in order to visualize thrombus in situ. In nonperfused animals, a fibrin-platelet–rich thrombus with some red blood cells was evident in the lumen of WT arteries (Figure 1B). In contrast, the lumen of Mrp14−/− arteries was filled with blood at this 25-minute time point when the flow probe indicated that the vessel was widely patent. Nonocclusive thrombus (arrow) was visible along the wall of vessel. With perfusion fixation, we found that the occlusive thrombus within the lumen of injured WT arteries was still visible, whereas the lumen of Mrp14−/− arteries was devoid of blood elements, indicating delayed thrombus formation and the instability of nonocclusive thrombus formed in Mrp14−/− mice. Immunohistochemical analysis of serial sections of injured arteries from WT and Mrp14−/− mice with anti-MRP-14 and the platelet-specific marker GPIIb antibodies showed positive MRP-14 staining.
that colocalized with platelets in WT arteries (Figure 1C). We observed no staining for MRP-14 in Mrp14–/– arteries.

**Impaired thrombus formation after laser-induced injury of the cremaster microvasculature in Mrp14–/– mice.** We used intravital microscopy to compare thrombus formation in vivo after laser-induced injury to the arteriolar wall in the cremaster microcirculation of Mrp14–/– mice with that of WT mice (18). In WT mice, platelet accumulation in arterioles was evident within 30 seconds of laser injury (Figure 1, D and E, and Supplemental Videos 1 and 2; supplemental material available online with this article; doi:10.1172/JCI70966DS1). In contrast, platelet accumulation was markedly attenuated in Mrp14–/– mice (mean percentage of inhibition over time = 85.0% ± 5.1%, n = 10–15 arterioles per group). Initial platelet adhesion and small platelet aggregates were observed, but developing thrombi were unstable and embolized frequently.

**Platelet count and coagulation assays are similar in WT and Mrp14–/– mice.** Having observed delayed thrombosis in Mrp14–/– mice, we set out to determine the mechanism by first performing screening platelet and coagulation assays in WT and Mrp14–/– mice (Supplemental Figure 1). The platelet count was similar in WT (736,000 ± 307,000 platelets/μl) and Mrp14–/– (753,000 ± 241,000 platelets/μl, P = 0.90) mice. We assessed the coagulation activity of plasma using the activated partial thromboplastin time (aPTT) and a thrombin generation assay. The aPTT was not prolonged in Mrp14–/– mice (WT: 66 ± 28 seconds versus Mrp14–/–: 55 ± 16 seconds, P = 0.54). Tissue factor–induced total thrombin generation was similar in WT and Mrp14–/– plasma (WT: 21,055 ± 407 versus Mrp14–/–: 21,001 ± 4,041 arbitrary fluorescent units, P = 0.54). Moreover, multiple parameters of thrombin generation, including the lag time of thrombin generation, the maximum rate of thrombin generation, and the time to reach maximal thrombin activity, were comparable in WT and Mrp14–/– mice (data not shown). Taken together, these data indicate that neither platelet count nor coagulation parameters likely account for delayed thrombosis in Mrp14–/– mice.

**Platelets express MRP-8 and MRP-14 proteins.** Although we have detected MRP14 transcripts in human platelets, freshly-isolated...
MRP-14 deficiency attenuates thrombus formation under flow and is associated with defects in collagen-induced platelet activation. (A) Platelet thrombi on collagen-coated capillaries following perfusion of rhodamine 6G–labeled blood from WT and Mrp14–/– mice at an arterial shear rate of 625 s–1. Thrombus formation (B, area; C, volume) was quantified using computer-assisted imaging analysis (n = 3–5 per group). Flow cytometric analysis of P-selectin expression (D) and assessment of GPIIb/IIIa activation using staining with the JON/A antibody (E) following stimulation of washed platelets from WT (black bars) and Mrp14–/– (white bars) mice with 0 to 10 μg/ml collagen (n = 5 per group).

Platelet surface expression of MRP-14 (Figure 2C). To verify secretion of MRP-14, gel-filtered human platelets were stimulated with thrombin, pelleted, and the supernatant was harvested for determination of MRP-8/14 concentration by ELISA. The concentration of MRP-8/14 in 1 ml of supernatant from 400 million thrombin-stimulated platelets was 1.03 ± 0.56 μg/ml.

To determine the relative abundance of MRP-8/14 compared with other intracellular platelet proteins/agonists, we assayed the protein content of human platelet MRP-8/14 compared with that of platelet CD40L, a platelet α-granule agonist that binds to GPIIb/IIIa and promotes thrombosis in an autocrine manner (19). We found that human platelet MRP-8/14 protein was more abundant than human platelet CD40L (Figure 2D). To verify that platelet activation and thrombosis are associated with secretion of MRP-14, we hypothesized that MRP-14 levels would increase after carotid artery photochemical injury. Although the source of MRP-14 is uncertain (i.e., platelet, leukocyte, or endothelial cell derived), plasma MRP-14 levels increased 10 minutes after injury compared with levels in noninjured animals (Figure 2E). In addition, when plasma and serum levels of MRP-8/14 in paired samples from 5 normal human donors were compared, we found that MRP-8/14 levels were significantly higher in serum than in plasma (Figure 2F).

Taken together, these observations indicate that platelets express both MRP-8 and MRP-14 protein and are capable of secreting MRP-8/14 after agonist stimulation and thrombus formation.

Deficiency of MRP-14 attenuates platelet thrombus formation under flow ex vivo. Having demonstrated that thrombus formation is attenuated in Mrp14–/– mice and that platelets express MRP-8/14, we next determined whether platelet MRP-14 itself regulates platelet function. We examined the role of MRP-14 in platelet thrombus formation under physiologic arterial shear conditions using a highly automated dynamic flow system (20, 21). Thrombus formation was achieved by perfusion of anticoagulated blood labeled with rhodamine 6G through collagen-coated rectangular capillaries at an arterial shear rate of 625 s–1. We quantified the thrombus area and volume in real time using computer-assisted imaging analysis (Figure 3A). Perfusion of blood resulted in the rapid formation of platelet thrombi that were significantly reduced in Mrp14–/– (Figure 3A). Perfusion of blood resulted in the rapid formation of platelet thrombi that were significantly reduced in Mrp14–/– mouse platelets compared with WT mice (percentage of inhibition of thrombus area = 49, P < 0.001; percentage of inhibition of thrombus volume = 60, P < 0.001) (Figure 3, B and C).

Given the defect in platelet thrombus formation on collagen under flow, we assessed platelet activation by monitoring the expression of P-selectin and activated GPIIb/IIIa (JON/A-positive staining) in response to agonist stimulation. Washed platelets from WT and Mrp14–/– mice were stimulated with collagen, thrombin, arachidonic acid, or ionomycin. P-selectin expression...
was significantly decreased in Mrp14–/– compared with that in WT platelets following stimulation with collagen at 5 and 10 μg/ml (P = 0.005 and P = 0.048, respectively; Figure 3D) and with 800 μM arachidonic acid (P = 0.027) (Supplemental Figure 3), but not with thrombin or ionomycin (Supplemental Figure 3). Collagen-induced activation of GPIIb/IIIa was also significantly reduced (P = 0.010) in Mrp14–/– versus WT platelets (Figure 3E). Importantly, we verified that the expression levels of the platelet receptors GPIIbα, GPIVI, and αIIbβ3, were comparable on WT and Mrp14–/– platelets (Supplemental Figure 4A).

We also performed platelet secretion and aggregation studies to further characterize the nature of the Mrp14–/– platelet defect. We assessed dense granule secretion by measuring agonist-induced secretion of [14C] 5-hydroxytryptamine and observed no significant difference in the uptake of [14C] 5-hydroxytryptamine between WT and Mrp14–/– platelets (44% and 47%, respectively). After collagen stimulation, Mrp14–/– platelets secreted 45% ± 19% of [14C] 5-hydroxytryptamine compared with 33% ± 3% for WT platelets (P = 0.32; Supplemental Figure 4E). Similarly, there was no difference in α-thrombin–induced platelet secretion (Mrp14–/– platelets secreted 98% ± 1% of [14C] 5-hydroxytryptamine versus 95% ± 4% for WT platelets, P = 0.28; Supplemental Figure 4E). We monitored platelet aggregation during secretion experiments and found no difference in collagen- or thrombin-induced platelet aggregation between WT and Mrp14–/– platelets (Supplemental Figure 4F). Finally, to determine whether Mrp14–/– modulates the platelet aggregation threshold, we evaluated ADP-induced fibrinogen binding by flow cytometry over a range of ADP concentrations and found that Mrp14–/– deficiency had no effect on ADP-stimulated fibrinogen binding (Supplemental Figure 4G).

Hemostasis as determined by tail vein bleeding time is unimpaired in Mrp14–/– mice. To assess the role of Mrp14 in hemostasis, we examined tail vein bleeding times. There was no difference in tail bleeding times between WT and Mrp14–/– mice using complete cessation of bleeding either for 3 minutes (A) or 30 seconds (B) as the criterion for determination of bleeding time (mean ± SD, n = 16 per group).

Role of intracellular versus extracellular MRP-8/14 in thrombosis. Transfusion of WT gel-filtered platelets nearly corrected the thrombotic defect in Mrp14–/– mice, indicating that platelet MRP-8/14 content regulates thrombus formation (Figure 5A). To determine whether extracellular MRP-8/14 action modulates thrombosis, we infused purified, recombinant human MRP-8, MRP-14, or MRP-8/14 (0.08 μg/g mouse) into Mrp14–/– mice (Figure 5B). Intravenous infusion of purified MRP-14 (30.9 ± 10.5 versus saline control 46.4 ± 20.5 minutes, P = 0.0025) or MRP-8/14 (25.7 ± 13.3 versus saline control 46.4 ± 20.5 minutes, P = 0.007) into Mrp14–/– mice shortened thrombotic occlusion time to that observed in WT mice (27.1 ± 5.9 minutes) (Figure 5B). In contrast, infusion of purified MRP-8 alone had no significant effect on thrombotic occlusion time (MRP-8: 41.1 ± 24.4 minutes, P = 0.625), strongly suggesting that MRP-14 is responsible for thrombotic action of the MRP-8/14 heterodimer complex.

To verify the importance of extracellular MRP-8/14 in modulating platelet function, we assessed the effect of purified MRP-8/14 on platelet thrombus formation under flow conditions. Purified MRP-8/14 enhanced thrombus formation of Mrp14–/– whole blood perfused through collagen-coated capillaries (Figure 5, C and D). Finally, to extend these findings to human platelets, we examined the effect of targeting extracellular MRP-8/14 on platelet aggregate formation by assessing the effect of anti–MRP-14 monoclonal antibody on platelet thrombus formation under flow of anticoagulated human whole blood and found that anti–MRP-14 antibody inhibited platelet thrombus formation compared with control antibody (Figure 5E).

Identification of the candidate platelet receptor for MRP-14. Having observed that extracellular MRP-14 promotes thrombosis, we next sought to identify candidate platelet receptor(s). Putative receptors for MRP-8/14 on target cells include CD36 (8), RAGE (10), and TLR4 (9). Interestingly, both CD36 (22) and RAGE (23) signaling have been directly implicated in platelet activation and thrombosis. Platelets also express functional levels of toll-like...
receptor 4 (TLR4) (24). We evaluated shear-induced platelet aggregation and thrombus formation in anticoagulated human whole blood in the presence of blocking antibodies against CD36, RAGE, or TLR4 (Figure 6). Anti-CD36 monoclonal antibody inhibited shear-induced thrombus formation (percentage of inhibition = 53.8 ± 16.9, \( P = 0.012 \)) (Figure 6, A–C), and the extent of inhibition was comparable to that observed with anti–MRP-14 antibody (Figure 5E). In contrast, blocking antibodies against either RAGE or TLR4 had no effect on platelet aggregate formation under these experimental conditions (Figure 6, A–C).

Next, to determine whether CD36 is a candidate receptor for MRP-14, we examined the direct binding of MRP-14 to purified soluble CD36 in a plate binding assay. MRP-14 bound to soluble CD36-coated, but not BSA-coated, wells (Figure 7A).

To establish whether CD36 is required for MRP-14 action, we crossed 

\( \text{Mrp14}^{/-/-} \)

mice with CD36-deficient (\( \text{Cd36}^{/-/-} \)) mice to generate compound mutants (\( \text{Mrp14}^{/-/-} \text{Cd36}^{/-/-} \)) and then subjected them to carotid photochemical injury. Deficiency of CD36 alone had no effect on thrombotic occlusion time (CD36: 23.1 ± 8.3 minutes versus WT: 27.1 ± 5.9 minutes, \( P = 0.167 \)) in this model (Figure 7B), a finding similar to that observed with the FeCl\(_3\) carotid injury model using 12.5% FeCl\(_3\) (25). Doubly deficient 

\( \text{Mrp14}^{/-/-} \text{Cd36}^{/-/-} \)

mice have prolonged thrombotic occlusion time that is no different than that in singly deficient 

\( \text{Mrp14}^{/-/-} \)

mice (44.7 ± 16.2 minutes versus 46.6 ± 22.9 minutes, respectively, \( P = 0.803 \); Figure 7B). In direct contrast to experiments performed with 

\( \text{Mrp14}^{/-/-} \)

mice, in which infusion of purified MRP-14 (0.08 μg/g mouse) shortened the prolonged time to thrombotic occlusion (saline: 46.4 ± 20.5 minutes versus purified MRP-14: 30.9 ± 10.5 minutes, \( P = 0.026 \); Figure 7B), infusion of MRP-14 into doubly deficient 

\( \text{Mrp14}^{/-/-} \text{Cd36}^{/-/-} \)

mice had no significant effect on the prolonged occlusion time (saline: 48.2 ± 15.3 minutes versus purified MRP-14: 47.2 ± 18.8 minutes, \( P = 0.884 \); Figure 7B).

Infusion of a 5-fold increased amount of MRP-14 (0.4 μg/g mouse weight) into 

\( \text{Mrp14}^{/-/-} \text{Cd36}^{/-/-} \)

mice also failed to shorten the prolonged occlusion time (43.0 ± 8.7 minutes, \( P = 0.40 \)).
To verify the importance of CD36 in modulating platelet function in response to extracellular MRP-14, we assessed the effect of purified MRP-14 on platelet thrombus formation under flow using Mrp14–/– Cd36–/– whole blood. While purified MRP-14 restored platelet thrombus formation of Mrp14–/– whole blood perfused through collagen-coated capillaries (Figure 7, C and D), purified MRP-14 had minimal enhancement of platelet thrombus formation of doubly deficient Mrp14–/– Cd36–/– whole blood (Figure 7, C and D). Further, we also examined whether MRP-14 is capable of activating platelets in a CD36-dependent manner. Oxidized LDL (oxLDL) initiates a CD36-mediated signaling cascade involving recruitment of Src family kinases (VAV, FYN, and LYN) and activation of JNK (26). Similarly to oxLDL, MRP-14 induces phosphorylation of VAV and JNK (Figure 7, E and F). Since hyperlipidemia increases both plasma oxLDL and MRP-14 concentrations (15), we examined the phosphorylation of VAV and JNK after stimulating platelets with both oxLDL and MRP-14. Interestingly, MRP-14 potentiated oxLDL-induced phosphorylation of both VAV and JNK. Taken together, these observations indicate that platelet CD36 is required for MRP-14 action (Figure 7, E and F).

Platelet MRP-8/14 in human coronary thrombus. MI is most commonly caused by atherosclerotic plaque rupture and occlusive thrombus formation (1). To begin to examine the pathophysiological relevance of our findings of MRP-8/14 expression in platelets, we obtained coronary artery thrombi from patients (n = 4) presenting to the cardiac catheterization laboratory with acute STEMI. Angiography performed on one patient demonstrated thrombotic occlusion of the proximal right coronary artery (Figure 8A) that was treated with aspiration thrombectomy followed by balloon angioplasty and stent deployment (Figure 8B), resulting in a widely patent right coronary artery with no significant luminal narrowing (Figure 8C). The thrombectomy catheter retrieved multiple coronary artery thrombi (Figure 8D) that were then stained for platelets and MRP-8/14 using immunofluorescence microscopy. Platelet and MRP-8/14 staining were abundant and colocalized in this human coronary artery thrombus (Figure 8, E–H). These findings were confirmed in the examination of intracoronary thrombi from three additional STEMI patients (Supplemental Figure 5).

**Discussion**

In this study, we identified a new pathway of thrombosis involving platelet MRP-14 and CD36 that does not affect bleeding time. This conclusion is supported by the following data: (a) the time to thrombotic occlusion was significantly prolonged in Mrp14–/– mice; (b) laser-induced platelet thrombi in the cremaster microvasculature were reduced and less stable in Mrp14–/– mice; (c) platelet thrombus formation under flow was reduced in whole blood from Mrp14–/– mice; (d) MRP-8 and MRP-14 were expressed in...
Figure 7
CD36 is required for MRP-14 action. (A) Binding of purified MRP-14 (0–2.5 μg/ml) to purified soluble CD36-coated or BSA-coated wells. (B) Thrombotic occlusion time after carotid artery photochemical injury in indicated mouse strains and occlusion time with intravenous infusion of saline or purified human MRP-14 (0.08 μg/g mouse) into Mrp14−/− or Mrp14−/− Cd36−/− recipient mice prior to photochemical injury. (C) Purified MRP-14 restores platelet thrombus formation under flow in Mrp14−/−, but not Mrp14−/− Cd36−/−, murine whole blood. Platelet thrombi on collagen-coated capillaries following perfusion (shear rate of 625 s⁻¹) of rhodamine 6G–labeled Mrp14−/− blood from WT, Mrp14−/−, or Mrp14−/− Cd36−/− mice that was treated with purified human MRP-14 (5 μg/ml) or control buffer. Original magnification, ×40; observation area, 360 × 270 μm. (D) Continuous, real-time thrombosis profiles of the average fluorescence of three independent experiments. MRP-14 induced phosphorylation of VAV (E) and JNK (F) in platelets. Gel-filtered human platelets (2 × 10⁹/ml) containing 2 mM CaCl₂ and 1 mM MgCl₂ were incubated with 50 μg/ml oxLDL, 1 μg/ml MRP-14, or a combination of these for 10 minutes, and platelet lysates were analyzed by immunoblotting with anti-phosphoprotein antibodies. The membranes were then stripped and reprobed with antibodies against the total relevant protein and actin. Results are representative of three independent experiments from different donors.
platelets, and agonist stimulation led to increased platelet surface expression and secretion of MRP-8/14; (e) platelet count, aPTT, thrombin generation, and bleeding time were similar in WT and Mrp14–/– mice; (f) transfusion of WT platelets or infusion of purified MRP-14 or MRP-8/14 into Mrp14–/– mice shortened the prolonged carotid artery occlusion time in Mrp14–/– mice; (g) compound deficiency of MRP-14 and CD36 resulted in a prolonged carotid artery occlusion time despite infusion of purified MRP-14; (h) MRP-14 was capable of activating platelets in a CD36-dependent manner (i.e., induces phosphorylation of VAV and JNK); and (i) robust expression of MRP-8/14 was evident in platelet-rich coronary artery thrombi, causing acute MI.

MRP-8/14 complexes are also present in mouse cells, and extensive biochemical characterization has confirmed that mouse MRP-14 is functionally equivalent to its human counterpart (27). Analyses of mice that lack MRP-8 and MRP-14 have provided important insights into the function of these proteins. Although homozygous deletion of MRP-8 results in embryonic lethality (28), deletion of the MRP14 gene does not affect viability and results in the additional loss of MRP-8 protein (5, 13). Failure to produce mature MRP-8 protein in the presence of normal MRP8 mRNA production likely results from an instability of MRP-8 in the absence of MRP-14. Thus, the Mrp14–/– mice used in this and previous studies lack both MRP-8 and MRP-14 protein and MRP-8/14 complexes.

MRP-8/14 is known to be expressed by various cell types, including neutrophils (29), monocytes (29), tissue macrophages under conditions of chronic inflammation (30), mucosal epithelium (31), and involved epithemis in psoriasis (32). Immunofluorescence, immunoblotting, and ELISA data from the present study now indicate that platelets also express and secrete MRP-8/14 protein. Immunofluorescence staining of purified platelets and coronary artery thrombi as well as flow cytometry of purified platelets indicate that MRP-8/14 protein is expressed in platelets. However, not all platelets appeared to express MRP-8/14. The basis for MRP-8/14 high versus low/negative platelet staining is unknown. Platelet age (young versus old) and activation status are known to influence platelet protein expression (33) and are the focus of ongoing studies.

The established roles of intracellular and extracellular MRP-8/14 in leukocyte function provide important clues to the putative action of MRP-8/14 in platelet function and thrombosis. Experiments with leukocytes isolated from Mrp14–/– mice have demonstrated aberrant calcium signaling and blunted calcium responses following chemokine stimulation (34). MRP-8/14 further appears to modulate calcium-coupled arachidonic acid signaling by binding to arachidonic acid in a calcium-dependent manner (35, 36) and by facilitating translocation of arachidonic acid to the cytoskeleton (37, 38). Intracellular MRP-8/14 also modulates cytoskeletal reorganization by promoting polymerization of microtubules (6).

Our observations that transfusion of WT platelets or infusion of purified MRP-14 or MRP-8/14 into Mrp14–/– mice shortened the prolonged carotid artery occlusion time of Mrp14–/– mice suggest that extracellular, rather than intracellular, MRP-14 is largely responsible for MRP-14 action in thrombosis. In response to cytokines or during contact with activated endothelium, myeloid cells secrete heterodimeric MRP-8/14, which is the dominant extracellular form (39–41), through a tubulin-dependent “alternative” secretion pathway (42). Extracellular MRP-8/14 is then able to bind to receptors on target cells, including CD36 (8), RAGE (10), TLR4 (9), special carboxylated N-glycans (43), and heparin-like glycoaminoglycans (44). Interestingly, both CD36 (22) and RAGE (23) signaling have been directly implicated in platelet activation and thrombosis. Platelets also express functional levels of TLR4 (24), which may contribute to thrombocytopenia through neutrophil-dependent pulmonary sequestration in response to LPS (45). Antibody-blocking experiments performed in the present study revealed CD36 as the most likely platelet receptor for MRP-14.
CD36 is an 88-kDa integral membrane protein expressed on platelets, monocytes/macrophages, microvascular endothelium, adipocytes, muscle cells, and specialized epithelium (e.g., retinal pigment epithelium) (46, 47). CD36 was first isolated and structurally characterized from platelets (48, 49), and a series of studies indicate that platelet CD36 binding to oxLDL (25) or endothelial cell–derived microparticles (22) contributes to platelet activation and thrombosis in mice. Despite having short intracytoplasmic domains, CD36 is capable of serving as a signaling receptor (50). oxLDL binding to CD36 induces platelet activation via a signaling cascade involving MAP kinase kinase 4 (MKK4), the MAP kinase JNK2, and Src kinases (e.g., VAV, FYN, LYN) (26). Interestingly, C36Δ mice have normal tail vein bleeding times and do not exhibit a bleeding diathesis (22). However, CD36 deficiency is associated with variable protection against thrombosis in the FeCl3 injury model that was dependent on the FeCl3 dose — namely, using a lower dose of FeCl3 (7.5% for 1 minute compared with 12.5% for 3 minutes, used by Podrez et al., ref. 25). Ghosh and colleagues concluded that CD36 ligands are generated during vascular injury and that the signals induced by these ligands contribute to thrombus formation by paracrine-like action (22). The results of our study indicate that MRP-14 is likely one such ligand. Our findings that transfusion of WT platelets into Mrp14−/− mice or infusion of purified MRP-14 into Mrp14−/−, but not Mrp14−/− C36Δ−/− (Figure 7B), mice shortened the prolonged carotid artery occlusion time in Mrp14−/− mice strongly suggest that platelet MRP-14 influences thrombosis secondary to secretion and binding to platelet CD36. Furthermore, while purified MRP-14 restored platelet thrombus formation in Mrp14−/− whole blood perfused through collagen-coated capillaries (Figure 7, C and D), purified MRP-14 had little effect on platelet thrombus formation in doubly deficient Mrp14−/− C36Δ−/− whole blood (Figure 7, C and D). We performed a plate binding assay, which demonstrated that purified MRP-14 is capable of binding directly to soluble CD36 (Figure 7A) and of activating platelets in a CD36-dependent manner, as indicated by the phosphorylation of VAV and JNK (Figure 7, E and F). Similarly to Podrez and coworkers, who used 12.5% FeCl3 (25), we did not observe a thrombosis defect in C36Δ−/− mice using the photochemical injury model. In contrast, Ghosh and colleagues reported a thrombosis defect using 7.5% FeCl3 (22). The basis for the discrepant findings in C36Δ−/− mice is uncertain, but they suggest that there may be model-dependent factors. The fact that the time to thrombogenic occlusion after photochemical injury was prolonged in Mrp14−/−, but not C36Δ−/−, mice raises the possibility that intracellular (e.g., cytoskeletal reorganization, calcium-coupled arachidonic acid signaling) as well as extracellular MRp-14 action modulates platelet function. Indeed, a role for intracellular MRP-14 is supported by Figure 7D, which shows that the addition of purified MRP-14 restored platelet thrombus formation in Mrp14−/− whole blood perfused through collagen-coated capillaries to a level below that of WT whole blood. Its role is also supported by the finding that arachidonic acid–induced platelet activation was attenuated in Mrp14−/− platelets (Supplemental Figure 3C). The possibility that MRP-14 binds to additional platelet receptors (e.g., TLR4, RAGE) to influence platelet and thrombosis functions cannot be definitively excluded in the present studies and will require the generation of doubly deficient MRP-14 and TLR4 or RAGE mice, respectively.

The binding of MRP-14 to platelet CD36 is analogous to other ligand-receptor interactions that act within the platelet-platelet contact zone after the initial aggregation event, including, among others, CD40L and its binding to platelet GPIIb/IIIa (19), GAS6 and its tyrosine kinase receptors MER, TYRO3, and AXL (51), and ephrins and their EPH kinase receptors (52). Indeed, it has been proposed that after initial aggregation, platelets form a synapse that facilitates signaling by membrane-tethered receptor-ligand pairs and localizes secreted and shed ligands that ultimately promote thrombus growth and stability (53). MRP-14 appears to function in this synapse.

Our studies show that MRP-14 deficiency did not interfere with tail bleeding time, platelet adhesion to and spreading on vWF or collagen, or plasma coagulation activity (i.e., aPTT and thrombin generation). The identification of a new platelet-dependent pathway for thrombosis that does not affect hemostatic parameters, such as bleeding time and platelet adhesion and spreading, has possible clinical implications. Thrombotic cardiovascular diseases, including MI and stroke, are the leading cause of death in developed countries (54). Total U.S. health care expenditures in 2009 for coronary heart disease and stroke were a staggering $165.4 billion and $68.9 billion, respectively (54), with the cost of pharmacologic therapies estimated to exceed $20 billion worldwide (54). Primary drug therapies include antiplatelet and anticoagulant agents. Antiplatelet agents and anticoagulants are used in the treatment of acute coronary syndromes and in primary and secondary prevention of coronary artery disease and stroke (55, 56). Current drugs are subject to significant bleeding risk, which is associated with increased mortality (57–59). While new antiplatelet (prasugrel, ticagrelor) and anticoagulant (dabigatran, rivaroxaban, apixaban) agents are being developed on the basis of superior efficacy, these therapeutic advances are associated with a 25% to 30% increase in the rate of bleeding (prasugrel, ticagrelor, apixaban) or transfusion (rivaroxaban, apixaban) in the treatment of acute coronary syndromes (60–62). There is emerging experimental evidence distinguishing the molecular and cellular mechanisms of hemostasis and thrombosis (63). Extracellular MRP-14 is now positioned as a novel and targetable mediator of thrombosis, but not hemostasis (i.e., reduced bleeding risk).

Finally, platelet MRP-8/14 has the potential to serve as a useful biomarker of coronary artery disease activity. The major causes of acute coronary syndromes, including MI and unstable angina, are plaque rupture and thrombosis (1). These ischemic events are typically precipitous and without warning symptoms. New approaches are required to identify patients who are at risk for the transition from “stable/chronic” to “unstable/acute” disease. Using platelet expression profiling, we reported previously that platelet MRP14 mRNA is increased in patients with STEMI compared with stable coronary artery disease patients (4). Ongoing studies are planned to determine whether platelet MRP-8/14 expression is predictive of coronary artery disease activity (i.e., stable angina versus acute coronary syndromes).

**Methods**

**Materials.** Antibodies against human MRP-14, MRP-8, and MRP-8/14 were purchased from Abcam, BMA Biomedicals, and R&D Systems. Antibodies against mouse MRP-8 and MRP-14 were purchased from R&D Systems. WugE9 antibody against mouse P-selectin/CαD262 conjugated with fluorescein isothiocyanate and JON/A (antibody against mouse-activated integrin GPIIIb/IIIa) conjugated with R-phycocerythrin were purchased from emfret Analytics. Alexa Fluor secondary antibodies and Alexa Fluor 647–conjugated mouse anti-human CD42a were purchased from AbD Serotec. The following azide-free antibodies were used for the shear-induced platelet

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aggregation assays: control mouse IgG, anti-human CD36 (clone FA6-152), and TLR4 (clone HTA125) monoclonal antibodies from Abcam; anti-human MRP-14 (clone MRP1H9) from Biolegend; and anti-human RAGE (clone MAB11451) monoclonal antibodies from R&D Systems. Purified, recombinant human MRP-8 and MRP-14 were obtained from Novus Biologicals. Human α-thrombin was purchased from Haematologic Technologies and human fibrinogen from Enzyme Research Laboratories. Rose Bengal (4, 5, 6, 9-[9H] xanthan)-3-1 dipotassium salt) was purchased from Sigma-Aldrich.

Mice. MRP-14−/− mice were generated in the laboratory of Nancy Hogg (5), and CD36−/− mice were generated in the laboratory of Roy Silverstein (22). All mice had a congenic C57BL/6 background and were maintained in animal facilities at Case Western Reserve University School of Medicine. Eight- to 12-week-old male mice were used for all experiments.

Platelet preparation and coagulation assays. For platelet isolation, aPTT, and thrombin generation, see Supplemental Methods. MRP-8/14 expression in platelets. For immunofluorescence microscopy, immunoblotting, and ELISA assays, see Supplemental Methods. Platelet α-granule release and GPIIb/IIIa activation. See Supplemental Methods. Mouse bleeding times. Tail bleeding times were measured by transecting the tails of anesthetized mice (50 mg/kg sodium pentobarbital) 5 mm from the tip, as previously described (64). Briefly, the transected tail tip was placed into a beaker containing saline at 37°C, and the time to complete cessation of bleeding for 30 seconds and 3 minutes was determined with a stopwatch.

Thrombus formation under laminar flow conditions. The laminar flow chamber used in this assay has been described previously (20, 21). Factor Xa inhibitor (Portola Pharmaceutical Inc.) anticoagulated blood was incubated with 0.2 μg/ml rhodamine-6G (Sigma-Aldrich) and then perfused for 5 minutes through human type III collagen–coated rectangular capillaries at 625 s−1, resulting in the deposition of adherent platelets and platelet aggregates. Thrombus formation under flow was then analyzed in real time and quantified by measuring fluorescence, thrombus area, and thrombus volume over time using computer-assisted imaging analysis (original magnification, ×40 and observation area, 360 × 270 μm).


Plate binding assay. See Supplemental Methods. Adaptive transfer/plaquelet transfusion experiments. Blood from the inferior vena cava of 2 mice was collected directly into 3.8% sodium citrate (9:1 blood/citrate ratio) and diluted with an equal amount of Tyrode’s buffer. For isolation of plaquelets, anticoagulated and diluted blood was centrifuged to obtain plaquelet-rich plasma and then applied to a column packed with Sepharose 2B to obtain gel-filtered plaquelets. WT or Mrp14−/− plaquelets (1 × 10^9 in 250 μl) were injected into recipient Mrp14−/− mice via the tail vein.

Histology and immunohistochemistry of tissue samples. See Supplemental Methods.

Data. Statistics were presented as the mean ± SD. Comparisons between groups were performed by an unpaired, 2-tailed Student’s t test. P values of less than 0.05 were considered statistically significant.

Study approval. Animal care and procedures were reviewed and approved by the IACUC of Case Western Reserve University School of Medicine and were performed in accordance with the guidelines of the American Association for Accreditation of Laboratory Animal Care and the NIH.

Approval for use of human tissue for pathological examination was obtained by the University Hospitals Case Medical Center Institutional Review Board. Human platelets were prepared from the whole blood drawn from the antecubital vein of healthy volunteers after providing informed consent, in accordance with a University Hospitals Case Medical Institutional Review Board–approved protocol.

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