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Journal Title: Journal of Clinical Investigation
Volume: Volume 125, Number 4
Publisher: American Society for Clinical Investigation | 2015-04-01, Pages 1471-1484
Type of Work: Article | Final Publisher PDF
Publisher DOI: 10.1172/JCI79329
Permanent URL: https://pid.emory.edu/ark:/25593/rxp6h

Final published version: http://dx.doi.org/10.1172/JCI79329

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Accessed April 5, 2019 2:47 PM EDT
Caspase-1–mediated pathway promotes generation of thromboinflammatory microparticles

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Extracellular ATP is a signal of tissue damage and induces macrophage responses that amplify inflammation and coagulation. Here we demonstrate that ATP signaling through macrophage P2X7 receptors uncouples the thioredoxin (TRX)/TRX reductase (TRXR) system and activates the inflammasome through endosome-generated ROS. TRXR and inflammasome activity promoted filopodia formation, cellular release of reduced TRX, and generation of extracellular thiolo-dependent, procoagulant microparticles (MPs). Additionally, inflammasome-induced activation of an intracellular caspase-1/calpain cysteine protease cascade degraded filamin, thereby severing bonds between the cytoskeleton and tissue factor (TF), the cell surface receptor responsible for coagulation activation. This cascade enabled TF trafficking from rafts to filopodia and ultimately onto phosphatidylserine-positive, highly procoagulant MPs. Furthermore, caspase-1 specifically facilitated cell surface actin exposure, which was required for the final release of highly procoagulant MPs from filopodia. Together, the results of this study delineate a thromboinflammatory pathway and suggest that components of this pathway have potential as pharmacological targets to simultaneously attenuate inflammation and innate immune cell–induced thrombosis.

Introduction

Atherosclerosis and its thrombo-occlusive complications remain a major cause of morbidity and mortality from cardiovascular diseases worldwide. Atherosclerotic plaques are complex inflammatory lesions with variable stability that upon rupture can cause acute thrombosis, vessel occlusion, and infarct (1). A major source for inflammatory and prothrombotic mediators are atherosclerotic plaque macrophages that express the proinflammatory cytokine IL-1β (2, 3) and tissue factor (TF) (4), which initiates the coagulation cascade (5).

A large body of evidence indicates an important role for IL-1β in the pathogenesis and progression of atherosclerosis (6). Endogenous or exogenous ligands for Toll-like receptors (TLRs), particularly TLR4 in atherosclerosis (7), specifically induce macrophage synthesis of the inactive pro-form of IL-1β. Secondary signals, including cholesterol crystals, a hallmark of atherosclerotic lesions (8), are required for assembly of the functional inflammasome multiprotein complex of NOD-like receptor family, pyrin containing 3 (NLRP3), apoptosis-associated speck-like protein containing a CARD (ASC), and procaspase-1. Activation of caspase-1 in this complex leads to cytokine maturation and release (9), although the final steps of IL-1β secretion are poorly understood (10).

Thrombosis is dependent on TF that is also induced by endogenous TLR ligands promoting atherosclerosis (11). Analogous to cytokine processing, TF is kept in an inactive form on the surface of myelomonocytic cells and requires additional signals to become fully procoagulant (12, 13). Activation of TF prothrombotic function is a thiol-disulfide exchange–dependent and protein disulfide isomerase–regulated (PDI-regulated) process (14) that occurs following outer membrane leaflet exposure of procoagulant phosphatidylserine (PS) (13, 15) and may involve other cell surface redox partners in the context of thrombosis (16, 17).

P2X7 receptor (P2RX7 in mice) activation has well-established roles in proinflammatory effects of macrophages (18). The purinergic receptor responds to extracellular ATP released from inflamed or damaged cells and is one of several signals that can activate the inflammasome and caspase-1–mediated processing of IL-1β (19). P2RX7 signaling also induces activation of cell surface TF and release of procoagulant TF-bearing microparticles (MPs) (12, 20). ATP promotes experimental thrombosis through platelet and neutrophil P2RX1 activation (21, 22) and P2RX7 signaling in mice (12). Since new biosensors have revealed the presence of high pericellular concentrations of ATP in inflammatory conditions in vivo (23), the presence of P2RX7 on macrophages in human atherosclerotic plaques (24) may indicate roles in the pathogenesis of arterial thrombotic complications of atherosclerosis.

P2RX7-mediated activation of macrophage TF procoagulant function specifically leads to extracellular thiol-disulfide exchange–dependent generation of procoagulant MPs carrying TF and integrin β1 as well as P-selectin glycoprotein ligand 1 (PSGL1), a major counterligand for platelet P-selectin in thrombotic vessel occlusion (25, 26). Indeed, atherosclerotic lesions accumulate large amounts of MPs that may support thrombus propagation upon plaque rupture (27). Although emerging evidence indicates that MP-incorporated proteins vary depending on the releasing stimulus (28, 29), little is known about the cellular mechanisms that determine MP composition and biological function.
The search for potential MP redox-regulated proteins by mass spectrometry did not identify proteins with vicinal thiols as potential downstream targets of TRXR. Rather, a major 54-kDa MP-associated thiol-labeled protein was identified as γ-actin. Western blotting showed that γ-actin was markedly increased in the MP fraction after ATP stimulation as well as released into the MP-depleted supernatant (Figure 1C). Thus, potential markers for the MP release mechanism may also be found in the cell supernatant. MPB labeling altered γ-actin electrophoretic mobility, suggesting that at least 1 of the 6 cysteine residues (31) was reduced and solvent exposed in the extracellular space (Figure 1F).

Working with the hypothesis that MP protein cargo is a unique fingerprint of underlying release mechanisms, we focused on defining the MP proteome susceptible to reduction during MP generation following prothrombotic P2RX7 activation. Here, we identify the thioredoxin (TRX)/TRXR system as a common upstream regulator for both procoagulant and proinflammatory responses of macrophages; and elucidate key steps of this pathway that ultimately triggers MP release through an unexpected function of caspase-1–dependent actin remodeling in primary cells.

Results

**P2RX7 activation induces extracellular reductive changes dependent on TRXR.** The mechanism through which P2RX7 activation by ATP leads to reductive changes in the extracellular proteome of macrophages and on released MPs (12), as detected by thiol labeling with the extracellular, water-soluble probe 3-(N-maleimido-propionyl)-biocytin (MPB), has remained unexplained to date. By using biotinylated PEG-maleimide with a membrane-impermeable molecular weight of 10 kDa, we confirmed that thiol labeling occurred extracellularly (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI79329DS1). Consistent with thiol-isomerase dependence of TF+ MP release, the vicinal thiol blocker phenylarsine oxide (PAO) prevented the appearance of solvent-accessible free thiols on the cell surface and on MPs (12). Although PAO might have blocked labeling of a variety of cell surface proteins carrying vicinal thiols, ATP-stimulated cells pretreated with 2,4-dinitrochlorobenzene (DNCB) also had markedly reduced cell surface free thiols. These data were surprising, since DNCB, unlike PAO, is not an inhibitor of reducing vicinal thiol groups, as confirmed for the reductive function of the thiol-isomerase PDI in an insulin reductase assay (Figure 1B).

As used here at low micromolar concentrations, DNCB rapidly and specifically inactivates the TRXR catalytic selenocysteine residue even in the presence of physiological intracellular concentrations (mM) of reduced glutathione (GSH), but not the vicinal thiols of TRXR required for NADPH oxidase activity (30). Prolonged inhibition of TRXR is known to affect the cellular redox equilibrium by depleting GSH. However, we measured only minimally lower levels of GSH after 20 minutes of ATP stimulation in DNCB-treated macrophages (85% ± 10% of ATP-stimulated controls). These results suggested that the effects of DNCB resulted primarily from TRXR inhibition and not global changes in redox equilibrium.

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**Figure 1. P2X7 receptor activation induces TRXR-dependent extracellular thiol-disulfide exchange.** (A) Effects of DNCB or PAO on ATP-induced exposure of solvent-accessible free thiols on the cell or MP surface were measured by thiol labeling with MPB and detected by streptavidin blotting. Detection of cellular γ-actin by Western blotting served as loading control. (B) Inhibition of PDI reductase activity measured by reduction of insulin by PDI with PAO or DNCB added at the indicated micromolar concentrations; mean ± SD, ***P < 0.001, ANOVA (Bonferroni). (C) Western blotting for γ-actin in MPs and MP-depleted supernatant from WT and P2rx7−/− macrophages stimulated with ATP in the presence or absence of MPB. (D) Streptavidin blot for thiol labeling with MPB detects a prominent protein band (arrowhead) of approximately 12 kDa – consistent with the molecular weight of TRX in the MP-free supernatant of ATP-stimulated cells. (E) Western blotting for TRX in cells, MPs, and MP-free supernatant of P2rx7−/− and WT macrophages. (F) MP modification of free thiols groups in TRX alters electrophoretic mobility. MPB was added at the beginning of (0 minutes) or 30 minutes after ATP stimulation. Proteins were detected by Western blot. (G) Effect of DNCB on ATP-induced release of γ-actin, TRX, or TRXR determined by Western blotting. Cell lysates were diluted 1:10 for γ-actin blot.
TRX contains 2 vicinal and 4 noncatalytic cysteine residues \((33)\). When free thiols were labeled with MPB (MW = 523) during (MPB added \(t = 0\) minutes) or after (\(t = 30\) minutes) MP release, TRX, but not \(\gamma\)-actin, showed differences in mobility (Figure 1F). The increased thiol content evidenced by the larger molecular mass of TRX in samples labeled continuously during the reaction 1C). Notably, actin is known to interact with TRX, the substrate of TRXR (32). The MP-depleted supernatant of ATP-stimulated cells was analyzed as an additional control. ATP, forward scatter. (B) Detection of TF and P5 on ATP-induced MP populations. (C) PS-dependent prothrombinase activity of macrophage MPs generated with or without DNCB. PS dependence was confirmed by blockade with annexin 5 (AS); mean \(\pm\) SD, \(n = 3\), ***\(P < 0.001\), ANOVA (Bonferroni). (D) Annexin 5 staining of ATP-stimulated WT cells after 30 minutes with and without DNCB treatment showed that blocking TRXR did not prevent cell surface PS exposure; scale bar: 100 \(\mu\)m; mean \(\pm\) SD, \(n = 3\), **\(P < 0.01\), t test. (E) Detection of PSGL1, TF, integrin \(\beta1\), PDI, and \(\gamma\)-actin on ATP-induced MPs from control and DNCB-treated TFK1 macrophages by Western blot. (F) Effect of DNCB on TF activity on cells and MPs from control (−) and ATP-stimulated (+) cells measured by FXa generation assay; mean \(\pm\) SD, \(n = 13\), *\(P < 0.05\), **\(P < 0.01\), ANOVA (Bonferroni). (G) TF activity on MPs collected 0–15 minutes and 15–30 minutes following ATP stimulation of control and DNCB-treated cells; mean \(\pm\) SD, \(n = 3\), *\(P < 0.05\), ANOVA (Bonferroni). (H) Coomassie blue–stained gels of MP-depleted supernatants of control and ATP-stimulated TFK1 macrophages with or without DNCB treatment. Six prominent protein bands were identified by mass spectrometry (see Supplemental Table 1 for results).
indicated that TRX was initially released in a more reduced form and then became oxidized in the extracellular space. Although TRX release required functional TRXR, TRXR was not released from P2RX7-stimulated cells (Figure 1G), indicating uncoupling of the TRXR/TRX system. Moreover, TRX secretion and extracellular MPB labeling occurred with remarkably similar kinetics (Supplemental Figure 1B), and inhibiting TRX directly following its release with the oxidizer PX-12 effectively diminished MPB labeling of cell surface proteins (Supplemental Figure 1C). Taken together, these data show that the reductive function of TRXR is required for the release of reduced TRX, which in turn promotes exposure of extracellular thiol groups following P2RX7 activation.

P2RX7-induced release of highly procoagulant MPs is dependent on TRXR. We then evaluated by flow cytometry how blockade of the TRXR system with DNCB influenced the thiol-disulfide exchange–dependent release of MPs. We used macrophages expressing human TF (TF knock-in [TFKI] mice) (34) to facilitate TF detection by FACS and confirmed that these cells were indistinguishable from macrophages expressing mouse TF in P2RX7-dependent MP release and response to inhibitor treatment (Supplemental Figure 2A–D).

ATP stimulation generated a population of large MPs (Figure 2A) with sizes ranging from 200 to 500 nm (Supplemental Figure 2, E and F). These MPs were no longer generated from DNCB-treated cells, whereas smaller vesicles were still released (Figure 2A). Collecting MPs from control and DNCB-treated cells by centrifugation at different speeds further supported different physical properties of these MPs (Supplemental Figure 3A). Detection of procoagulant PS by lactadherin (35) and of TF by a specific mAb (Supplemental Figure 3B) showed that only the large MP population carried both PS and TF, whereas the smaller MPs released from DNCB-treated cells incorporated TF, but little PS (Figure 2B). Accordingly, MPs released from DNCB-treated cells had low procoagulant activity in a prothrombinase assay, whereas high prothrombinase activity of MPs generated in the absence of DNCB was confirmed to be dependent on functional PS by blockade with annexin 5 (Figure 2C). Based on these findings, we defined the large MP population carrying TF and PS as highly procoagulant MPs. Notably, blocking TRXR did not prevent PS exposure detected by annexin 5 staining on the cell surface (Figure 2D), demonstrating that outer leaflet presence of PS alone does not promote highly procoagulant MP generation. To exclude compound-specific effects of DNCB, we reproduced the results of key experiments with relevant concentrations of another well-established inhibitor of TRXR, auranofin (36), as well as PX-12, which at high concentrations inhibits the TRX/TRXR system by a distinct mechanism not primarily targeting the catalytic selenocysteine residue of TRXR (37) (Supplemental Figure 4).

Further characterization of MPs released from TRXR-blocked cells (Figure 2C and Supplemental Figure 2A) confirmed the presence of membrane receptors — TF, PSGL1, and integrin β1 — but not of the TRX-interacting components PDI and γ-actin. Control experiments showed that ROS production, which was not blocked by DNCB, was responsible for the release of membrane receptors on MPs (Supplemental Figure 5, A and B). Despite similar TF antigen levels, TF-FVIIa–mediated FX activation was significantly diminished on MPs from DNCB-treated cells (Figure 2D), demonstrating that outer leaflet presence of PS alone was dispensable for TF procoagulant activity. Further, TF-FVIIa–mediated FX activation was significantly diminished on MPs from DNCB-treated cells (Figure 2D), demonstrating that outer leaflet presence of PS alone was dispensable for TF procoagulant activity. Further, TF-FVIIa–mediated FX activation was significantly diminished on MPs from DNCB-treated cells (Figure 2D), demonstrating that outer leaflet presence of PS alone was dispensable for TF procoagulant activity. Further, TF-FVIIa–mediated FX activation was significantly diminished on MPs from DNCB-treated cells (Figure 2D), demonstrating that outer leaflet presence of PS alone was dispensable for TF procoagulant activity.
ulant MPs for extended time periods. This pathway also released various soluble proteins into the extracellular space (Figure 2F), including cytoskeleton-associated proteins and annexins 1 and 5 (Supplemental Table 1), which are known to interact with the MP components actin and PS (38, 39).

**TRXR is required for lipid raft–dependent translocation of TF onto filopodia.** To clarify how membrane receptors are incorporated into highly procoagulant MPs, we tracked TF during the course of stimulation with antibodies (40) on TFKI macrophages. Control experiments excluded antibody-induced TF internalization following specific staining of the human protein, and demonstrated that the specifically labeled cell surface pool of TF, and not potential Golgi-mobilized TF (41), became activated and released on MPs (Supplemental Figure 6, A–C).

ATP stimulation promoted TF translocation onto filopodia (Figure 3) that formed concomitantly with measurable TF activity and release on MPs (Supplemental Figure 7, A–D). Blocking TRXR markedly reduced the length of filopodia in ATP-stimulated cells, and, most significantly, TF accumulated on the cell surface in the distal cellular cortex but was not further transported onto extending filopodia (Figure 3 and 3-dimensional reconstructions in Supplemental Videos 1–4). Confirming the requirement for filopodia formation to generate highly procoagulant MPs, we showed that pretreatment with the membrane-permeable actin filament stabilizer phalloidin oleate or the CDC42 inhibitor CASIN resulted in stunted filopodia outgrowth (Supplemental Figure 8A), reduced release of the large MP population (Supplemental Figure 8B), and decreased TF-specific activity on MPs to 46.5% ± 3.4% and 38.0% ± 1.4% of control, respectively.

Since lipid rafts regulate localization and function of TF and integrin β1 as well as PSGL1 (42–44), we asked whether rafts are required for receptor trafficking and incorporation into highly procoagulant MPs. In order to avoid effects of chronic depletion of cholesterol on TF induction (45, 46), we rapidly perturbed lipid raft dynamics by chelating cholesterol with a low concentration of filipin. Filipin had no significant effects on cellular TF activity, consistent with previous reports (47), but potently blocked ATP-induced release of TF activity on MPs (Figure 4A). PSGL1, TF, and integrin β1 were no longer incorporated into MPs in the presence of filipin (Figure 4B), but MP marker proteins γ-actin and PDI, which are targets for TRX, were still released and TRX and γ-actin were recovered in the cell supernatant (Figure 4, B and C). FACS analysis confirmed the unaltered release of the large PS + MP population (95% ± 3% of control) in the presence of filipin, which also had no effect on functional PS-dependent MP prothrombinase activity (78% ± 36% of control). However, filipin markedly reduced TF expression on these MPs (20% ± 2% of control) (Figure 4D). Filipin did not interfere with ATP-induced formation of filopodia, but prevented the appearance of TF on filopodia (Figure 4E). Taken together, these data demonstrate raft-dependent trafficking of membrane receptors onto filopodia in the course of MP generation.
P2RX7-induced TF⁺ MPs promote fibrin strand formation in whole blood under flow. We verified the procoagulant function of these macrophage-derived MPs in a more physiologically relevant whole blood environment. On the surface of LPS-primed WT macrophages exposed to recalcified flowing blood, limited amounts of fibrin formed that was associated primarily with platelet aggregates (Figure 5A, top panels). Following ATP stimulation of macrophages, fibrin associated with platelet aggregates was still present, but in addition long fibrin strands formed that extended in the direction of flow (Figure 5A, bottom panels). ATP simulation of LPS-primed but TF⁻ macrophages induced negligible fibrin formation, although platelet clusters were still recruited around some cells (Figure 5B).

Adding isolated MPs derived from ATP-stimulated cells to blood before perfusion resulted in fibrin strand formation on the surface of TF-deficient, unstimulated macrophages (Figure 5C, top), in striking resemblance to the strands formed on ATP-stimulated WT cells except that there were no platelet aggregates with associated fibrin (compare with Figure 5A, bottom panels). Since the added MPs were generated from humanized TFKI mice, and species-selective antibodies against human TF blocked fibrin formation (Figure 5C, middle), these results showed that human TF on MPs and not blood-borne mouse TF activated by added MPs triggered prothrombotic activity. In addition, PS-expressing procoagulant MPs generated in the presence of filipin with severely reduced levels of raft-associated receptors, including TF, failed to form fibrin strands comparable to those seen with TF⁺ MPs (Figure 5C, bottom). The similar pattern of fibrin deposition in flow-oriented strands that formed on ATP-stimulated WT macrophages or on TF⁺ macrophages when blood contained procoagulant MPs suggests that de novo MP release from cells is a crucial factor for coagulation propagation in flowing blood.

Activation of the inflammasome is required for extracellular thiol-dependent MP release. TRX dissociation from TRX interacting protein (TXNIP) activates the inflammasome (48). Since TXNIP, unlike TRX, remained cell associated in P2RX7-stimulated cells (Supplemental Figure 9A), we next asked how inflammasome activation is coupled to extracellular thiol pathways and MP release. ROS of different sources can promote the dissociation of TRX and TXNIP, subsequent assembly of the NLRP3 inflammasome, and caspase-1 activation (49). Strikingly, we found that treating macrophages with niflumic acid (NFA), a selective inhibitor of endosomal ROS of different sources can promote the dissociation of TRX and TXNIP, subsequent assembly of the NLRP3 inflammasome, and caspase-1 activation (49). Strikingly, we found that treating macrophages with niflumic acid (NFA), a selective inhibitor of endosomal ROS production (50), abrogated P2RX7-induced caspase-1 activation and TRX release (Figure 6A). Consistently, extracellular thiol appearance (Figure 6B), filopodia formation (Supplemental Figure 9B), and the downstream release of thromboinflammatory MPs (Supplemental Figure 9C) were blocked by NFA treatment. Since NFA also inhibits COX2, we excluded contributions of COX2 with the COX2 inhibitor rofecoxib and confirmed with another endosomal ROS inhibitor, 4,4’-diisothiocyanato-2,2’-stilbenedisulfonic acid (DIDS) (51), the specificity of the NFA effects on TRX release, caspase-1 activation, and secretion of IL-1β (Supplemental Figure 9, B–D). NFA did not block total cellular ROS production (Supplemental Figure 9E), and cell surface receptors were consequently still released (Supplemental Figure 9F), pointing to a specific role of endosomal ROS for P2RX7-induced inflammasome activation and thromboinflammatory MP release.

We further evaluated the roles of inflammasome activation in MP release with the pharmacological inhibitors parthenolide...
and BAY 11-7082 (52). As expected, both blocked ATP stimulation–induced caspase-1 activation and the subsequent processing and release of IL-1β (Figure 6C). Importantly, inhibitors of inflammasome activation also markedly attenuated the release of TRX and extracellular thiol appearance (Figure 6D), the formation of filopodia (Figure 6E), the generation of the typical large procoagulant MPs (Figure 6F) with TRX-target marker proteins (Supplemental Figure 9G), and release of MP TF activity (58.5% ± 7.7% in parthenolide-treated and 32.0% ± 1.7% in BAY 11-7082–treated cells compared with uninhibited control; n = 3, P < 0.05 and P < 0.01). These data demonstrate a novel role for inflammasome activation in promoting extracellular thiol modifications and coupling of thromboinflammatory responses of macrophages.

A caspase-1/calpain cysteine protease cascade releases TF from cytoskeletal anchoring during thiol-dependent MP release. We investigated additional contributions of inflammasome activation to the generation of highly procoagulant MPs. The cortical localization of surface TF in DNCB-treated cells suggested that a retention signal impaired trafficking of TF onto filopodia. TF is associated with cytoskeletal structures (53), and the cytoplasmic domain of TF interacts with filamin A (54) that anchors membrane proteins to the actin cytoskeleton (55). Proteolytic cleavage of filamin A by calpain (56) separates the 190-kDa actin-interacting domain from a 90-kDa carboxyl-terminal fragment involved in TF binding (54). We hypothesized that TF is retained by cytoskeletal anchoring and requires calpain-mediated filamin degradation for incorporation into MPs.

The endogenous inhibitor of calpain, calpastatin (56, 57), is degraded by cysteine proteases calpain and caspase-1 (58). ATP stimulation indeed induced calpastatin and filamin A degradation in WT macrophages, but not in P2rx7−/− macrophages (data not shown) or in the presence of DNCB (Figure 7A). As expected, the calpain inhibitor ALLN blocked cleavage of filamin A and calpastatin (Figure 7A). Calpain inhibition with ALLN had no effect...
Caspase-1 is required for extracellular actin-dependent MP generation. The following experiments uncovered additional calpain-independent roles of caspase-1 in the thromboinflammatory release pathway of macrophages. As expected, incubation with the caspase-1 inhibitor YVAD completely prevented P2RX7-induced processing of IL-1β but had no effect on inflammasome activation, as evidenced by generation of the caspase-1 p10 proteolytic fragment (Figure 8A). Furthermore, incubating cells with YVAD did not affect ATP-induced release of TRX into the extracellular space (Figure 8A) and reductive changes on the cell surface (Figure 8B). As predicted from the position of caspase-1 upstream of calpain, caspase-1 blockade caused retention of TF in the cell body; unexpectedly, it also changed the appearance of filopodia, increasing their length (Figure 8C), and led to an overall flatter shape of ATP-stimulated macrophages (Supplemental Videos 5 and 6).

Surprisingly, caspase-1 inhibition prevented the appearance of thiol-labeled proteins in the MP fraction (Figure 8B), including γ-actin and PDI (Supplemental Figure 11A, B), but ALLN treatment prevented the ATP-induced translocation of TF onto filopodia (Figure 7B). ALLN caused significantly reduced TF-specific activity on MPs (Figure 7C), particularly the activity on MPs that were released for prolonged times (Supplemental Figure 10D), and decreased the number of MPs incorporating both TF and PS (Supplemental Figure 9E).

In sharp contrast, deletion of the TF cytoplasmic domain linking TF to filamin A (54) prevented effects of ALLN and restored efficient translocation of TF onto filopodia (Figure 7B), and ATP caused significantly reduced TF-specific activity on MPs (Figure 7C), particularly the activity on MPs that were released for prolonged times (Supplemental Figure 10D), and decreased the number of MPs incorporating both TF and PS (Supplemental Figure 9E).

Figure 7. P2RX7 signaling-activated cysteine proteases regulate TF trafficking. (A) Effects of inhibitors of TRXR (DNCB), caspase-1 (YVAD), calpain (ALLN), or lipid rafts (filipin) on ATP-induced degradation of filamin and calpastatin detected by Western blotting. Detection of cellular γ-actin served as loading control. (B) Effects of ALLN on TF localization in control and following ATP stimulation of WT (TFWT) or TF cytoplasmic domain-deleted (TFΔCT) macrophages. Cell surface TF was labeled with immunopurified rabbit anti–mouse TF antibody before the MP release reaction. Fixed cells were counterstained with Alexa 488-conjugated anti-rabbit IgG (green), phalloidin–Alexa 546 (red), and Hoechst (blue); scale bar: 10 μm. (C) Quantification of TF localized on filopodia; mean ± SD, n ≥ 12, ***P < 0.001, t test. TF-specific activity on MPs from ATP-stimulated TFWT and TFΔCT macrophages in the absence or presence of ALLN was determined based on normalization of MP-released TF quantified by Western blotting; mean ± SD, n = 3, **P < 0.01, t test. (D) Western blot analysis of a time course of ATP-induced filamin and calpastatin degradation in control and in caspase-1 inhibitor-treated cells.
inflammasome component NLRP3 was cell associated and in the supernatant, the released MPs contained activated caspase-1, but not processed IL-1β, which was predominantly detected in the MP-depleted supernatant (Supplemental Figure 13A). In DNCB-blocked cells, no uncleaved caspase-1 was seen on the released MPs (Supplemental Figure 13B). These findings are consistent with specific physical association of active caspase-1 with budding MPs during thromboinflammatory MP release.

Consistent with the concept that the release of soluble proteins is mechanistically linked to the generation of thromboinflammatory MPs, inhibition of caspase-1, but not of calpain, markedly reduced the appearance of γ-actin and other proteins in the extracellular space (Figure 8F). The small amount of actin detected in caspase-1–blocked supernatants had slower mobility as compared with actin released from ATP-stimulated cells with a functional caspase-1, indicating caspase-1–dependent cleavage. Importantly, staining of nonpermeabilized cells with cell-impermeable, fluorophore-labeled phalloidin showed that ATP stimulation induced a TRXR-dependent cell surface appearance of actin that particularly decorated filopodia. In contrast, and despite actin remodeling doc-

**Figure 8. Caspase-1 activity is required for thromboinflammatory MP generation.** (A) Effects of inhibitors of TRXR (DNCB), caspase-1 (YVAD), and calpain (ALLN) on ATP-induced activation of caspase-1 and IL-1β, and secretion of TRX detected by Western blotting. (B) Streptavidin blots of MPB-labeled free thiol groups on cells and MPs generated in the presence or absence of YVAD or DNCB. (C) Effects of YVAD on ATP-induced filopodia formation and TF translocation. TF was stained with anti-TF–Alexa 647 conjugate (red) before stimulation. Fixed and permeabilized cells were counterstained for F-actin (green) and nuclei (blue); scale bar: 10 μm. Length of filopodia and appearance of TF on filopodia were assessed; mean ± SD, n = 4, ***P < 0.001, t test. (D) ATP-induced MPs from control and YVAD-pretreated cells detected by FACS. (E) Confocal imaging of continuously MPB-labeled cells stimulated with ATP with or without YVAD. MPB was detected with Texas Red–labeled streptavidin (red) and nuclei with DAPI (blue) after fixation; scale bar: 10 μm. (F) Coomassie blue staining of proteins in the MP-depleted supernatants of ATP-stimulated cells with or without YVAD or ALLN treatment. TFKI macrophages were used for FACS analysis and confocal imaging.
face receptors into MPs with a distinct composition of proteins that are targets of TRX and a high surface density of procoagulant PS. Together with released soluble proteins, these proteomes are reflective of mechanistic details of the MP release reaction and may provide diagnostic fingerprints for detecting thromboinflammatory activation of macrophages in complex biological samples.

Following P2RX7 activation, functional TRXR promotes the cellular release of its substrate TRX in a reduced form and thereby prevents intracellular TRX regulatory functions and causes extracellular reductive changes required for generation of a distinct MP population. Actin residue Cys374 is reversibly modified by TRX and controls actin dynamics (59). Furthermore, the TRX substrate PDI promotes actin polymerization at membrane protrusions (60), providing an explanation for TRXR-dependent filopodia formation and the incorporation of these marker proteins into MPs generated following actin remodeling.

Dissociation of TRX from TXNIP furthermore leads to inflammasome activation. Although ROS derived from damaged mitochondria (49) have been shown to mediate spatially controlled activation of the inflammasome, our experiments with chloride channel inhibitors that block superoxide efflux from the endosome indicate that endosomal ROS are specifically required for P2RX7-mediated rapid activation of the inflammasome. This

Figure 9. F-actin is exposed on the cell surface during MP generation. (A) Surface F-actin was detected by phalloidin staining of nonpermeabilized ATP-stimulated cells (middle panel) using phalloidin–Alexa 633 conjugates, and counterstained for total F-actin (right panel) with phalloidin–Alexa 488 conjugates and nuclei with Hoechst after fixation. Cells were pretreated with DNCB or YVAD, as indicated; scale bar: 20 μm. (B) Detection of F-actin on ATP-induced MP populations by FACS using phalloidin–Alexa 633 conjugates. (C) Confocal images of ATP-stimulated cells in the presence of phalloidin–Alexa 546 concentrations used for cell staining (66 nM) or MP release blocking experiments (500 nM) demonstrate that higher concentration of phalloidin did not inhibit filopodia formation. (D) Abundance of ATP-induced MPs detected by FACS from control cells and from cells stimulated in the presence of cell-impermeable phalloidin to block extracellular actin. TFKI macrophages were used for FACS analysis and confocal imaging.

Discussion

Here, we delineate key events required for prothrombotic MP generation following injury signal–induced activation of macrophage P2X7 receptors (Figure 10). In this pathway, TRXR is crucial for several steps that ultimately lead to the incorporation of cell surface receptors into MPs with a distinct composition of proteins that are targets of TRX and a high surface density of procoagulant PS. Together with released soluble proteins, these proteomes are reflective of mechanistic details of the MP release reaction and may provide diagnostic fingerprints for detecting thromboinflammatory activation of macrophages in complex biological samples.

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Dissociation of TRX from TXNIP furthermore leads to inflammasome activation. Although ROS derived from damaged mitochondria (49) have been shown to mediate spatially controlled activation of the inflammasome, our experiments with chloride channel inhibitors that block superoxide efflux from the endosome indicate that endosomal ROS are specifically required for P2RX7-mediated rapid activation of the inflammasome. This
The extracellular proteome composition is governed by specific agonist pathways, such as those involved in inflammatory trans-signaling (65, 67). The current identification of vascular wall cells promoting inflammation (29, 66) and proangiogenic literature documents active uptake of MPs and exosomes by these cells can increase TF-bearing particle uptake and moreover promote prothrombotic effects, interaction of MP TF with its signaling receptor on host cell surfaces. While TF was required for these prothrombotic processes, the effector protease of the inflammasome pathway is at the center of the coupling of inflammation and thrombosis.

Unexpectedly, caspase-1 has a calpain-independent pivotal role in the final release of procoagulant MPs. In the presence of functional caspase-1, inflammasome activation causes the extracellular appearance of actin that decorates the surface of extending filopodia, suggesting considerable membrane instability allowing for actin translocation. Actin-associated proteins, including annexins 1 and 5, are also released dependent on caspase-1 activity and may stabilize the P2RX7-induced but TRXR-dependent flip of PS to the outer membrane. The extracellular appearance of these proteins in conjunction with the membrane curvature of filopodia may favor the concentration of negatively charged phospholipids on released MPs (62, 63). Thus, the effector protease of the inflammasome pathway is at the center of the coupling of inflammation and thrombosis.

We confirmed in flowing blood that MPs generated by this pathway are highly thrombogenic and promote fibrin strand formation extending even from otherwise unperturbed and quiescent cell surfaces. While TF was required for these prothrombotic effects, interaction of MP TF with its signaling receptor on host cells can increase TF-bearing particle uptake and moreover promotes intracellular signaling (64, 65). Consistently, an expanding literature documents active uptake of MPs and exosomes by vascular wall cells promoting inflammation (29, 66) and proangiogenic trans-signaling (65, 67). The current identification of MP proteome composition governed by specific agonist pathways represents a first step toward studies that define biological effects beyond thrombosis by distinct MP populations carrying TF.

Although not addressed experimentally here, extracellular thiol-disulfide exchange in the context of surface exposure of procoagulant PS may contribute to procoagulant activation of what is referred to as cryptic cell surface TF. Given the redox potential (68) and biochemical evidence for TRX-mediated reduction (69) of the TF allosteric disulfide bond (70, 71), extracellular TRX may initiate thiol-disulfide exchange leading to reshuffling of this bond in cell surface–localized but coagulation-inactive TF. The demonstrated progressive oxidation of extracellular TRX in turn may cause subsequent PDI chaperone activity–supported (72) TF reoxidation in a PS-localized conformation, in analogy to PDI-dependent TF activation in the context of complement activation (13). Extracellular TRX may influence additional redox targets (73) and thereby more generally promote coagulation reactions contributing to hemostasis and thrombosis.

The TRX/TRXR system plays important regulatory roles in biological responses to oxidative stress and redox signaling in cardiovascular diseases (74, 75), and serum levels of TRX are elevated significantly in patients with coronary syndromes with as-yet unclear pathogenetic implications (76). We provide novel mechanistic insight into the significance of extracellular TRX as a potential thromboinflammatory MP. The intracellular depletion of TRX abolishes regulatory functions of TRX and causes actin remodeling and filopodia formation as well as TXNIP-mediated inflammasome assembly and caspase-1 activation. Active caspase-1 mediates maturation and release of the proinflammatory cytokine IL-1β and facilitates activation of calpain, which in turn reverses TF retention by filamin (indicated as a red rod on the cytoplasmic side of TF) and thereby allows raft-dependent trafficking of TF onto filopodia. Membrane receptors TF, integrin (β3), and PSGL1 assemble with marker proteins of highly procoagulant MPs, γ-actin, and PDI when released from PS-rich filopodia, facilitated by caspase-1 and resulting in highly procoagulant thromboinflammatory MPs. Key pharmacological inhibitors used here are placed next to their targets to indicate where the thromboinflammatory pathway is inhibited.

**Figure 10.** Schematic illustration of the proposed thromboinflammatory MP release pathway in macrophages. Macrophage priming by IFN-γ and TLR ligands induces transcriptional upregulation of TF and proinflammatory interleukins. Activation of the macrophage P2RX7 by ATP triggers TRXR-dependent release of TRX, leading to reductive changes of the extracellular proteome, as indicated by the extracellular free thiol (SH) groups. The intracellular depletion of TRX abolishes regulatory functions of TRX and causes actin remodeling and filopodia formation as well as TXNIP-mediated inflammasome assembly and caspase-1 activation. Active caspase-1 mediates maturation and release of the proinflammatory cytokine IL-1β and facilitates activation of calpain, which in turn reverses TF retention by filamin (indicated as a red rod on the cytoplasmic side of TF) and thereby allows raft-dependent trafficking of TF onto filopodia. Membrane receptors TF, integrin (β3), and PSGL1 assemble with marker proteins of highly procoagulant MPs, γ-actin, and PDI when released from PS-rich filopodia, facilitated by caspase-1 and resulting in highly procoagulant thromboinflammatory MPs. Key pharmacological inhibitors used here are placed next to their targets to indicate where the thromboinflammatory pathway is inhibited.
Although the P2RX7 thromboinflammatory pathway of TF activation and release is discussed primarily in the context of atherosclerosis, evidence suggests its implication in various diseases. For instance, cellular data indicate a critical role of P2RX7 in sepsis (78), which is particularly intriguing since bacterial LPS potently promotes TF upregulation. Indeed, increased TF⋅ MP levels correlate with hypercoagulability in endotoxemic mice (79), and extracellular TRX levels are significantly elevated in plasma of sepsis patients (80). Similarly, P2RX7 signaling is implicated in rheumatoid arthritis (81), and high levels of extracellular TRX (82, 83) as well as of TF⋅ MPs (84) are detected in the synovial fluids of patients. Notably, the TRXR inhibitor auranofin is efficacious in the treatment of rheumatoid arthritis. In light of the common role of caspase-1 in proinflammatory IL-1β activation and procoagulant MP release, it is tempting to speculate that the thromboinflammatory response pathway identified here is of broad clinical significance.

Methods

Extended experimental procedures and methods are described in the Supplemental Methods.

**MP release assay with primary bone marrow–derived macrophages.** Bone marrow–derived macrophages from C57BL/6, Tph1−/− Lysm-Cre mice (85), human TF knock-in (TFKI) mice (34), or caspase-1–deficient mice (86) were generated as previously described (12). Briefly, bone marrow cells were cultivated in DMEM containing 20% L cell medium, 10% FCS, 1 mM L-glutamine, and penicillin and streptomycin. The medium was exchanged on day 6, and on day 7 macrophages were seeded at 1 × 10^6 per well in a 12-well plate and primed with 100 ng/ml IFN-γ (PeproTech) overnight. On the following day, macrophages were treated with 1 mg/ml LPS (Salmonella abortus equi; Enzo Life Sciences) for 4 hours. Cells were pretreated for 30 minutes with DNCB (30 mM; Sigma-Aldrich), parthenolide (10 mM; Cayman Chemical), BAY-11-7082 (2 mM; Cayman Chemical), DPI (100 mM; Sigma-Aldrich), or NFA (30 mM; Sigma-Aldrich) or immediately before ATP stimulation with PAO (10 mM; Sigma-Aldrich), filipin (5 mg/ml; Sigma-Aldrich), ALLN (1 mM; EMB Millipore), or Ac-YVAD-CMK (10 mM; Enzo Life Sciences). Cells were rinsed once in BSS buffer (0.13 M Na-gluconate, 0.02 M HEPES, 5 mM glucose, 5 mM BSA, 0.45 mM NaCl) and stained with anti–human TF 9C3–Alexa 647 conjugate (5 mg/ml) and lactadherin-FITC (0.5 mg/ml; Haematologic Technologies), and data were analyzed using FlowJo software (Tree Star).

**Microscopy.** MP procoagulant activity on cell surfaces was evaluated in flow chamber experiments with whole blood using confocal microscopy and quantitative image analysis, as previously described (12). Cells stained and mounted as described in the Supplemental Methods were analyzed using a ×63 oil immersion objective on a Zeiss 710 LSM, and images were processed with Image Browser (Zeiss) and Imaris (Bitplane) image analysis software.

**Statistics.** Statistical analysis was performed using Prism 6 software (GraphPad Software). For statistics comparing 2 groups, the t test (2-sided) was used, and for statistical analysis of more than 2 groups, 1-way ANOVA with Bonferroni multiple comparison correction was used. For all tests at least 3 independent experiments were analyzed, and P values of less than P < 0.05 were considered significant.

**Study approval.** Animal procedures were approved by the IACUC of The Scripps Research Institute.

**Acknowledgments**

We greatly appreciate the excellent technical assistance by Cynthia Biazak, Pablito Tejada, and Jennifer Royce. This study was supported by NIH grants R01-HL031950 (to W. Ruf, Z.M. Rughger, and M.H. Ginsberg) and R01-HL077753 (to W. Ruf) and a fellowship of the Deutsche Forschungsgemeinschaft (RO 3795/2-1) and NovoNordisk (to A.S. Rothmeier).

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