Shedding of the Mer Tyrosine Kinase Receptor Is Mediated by ADAM17 Protein through a Pathway Involving Reactive Oxygen Species, Protein Kinase Cδ, and p38 Mitogen-activated Protein Kinase (MAPK)*

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Edward Thorp†‡, Tomas Vaisar‡, Manikandan Subramanian§, Lauren Mautner¶, Carl Blobel‡, and Ira Tabas†‡

From the †Departments of Medicine, Pathology and Cell Biology, and Physiology, and Cellular Biophysics, Columbia University, New York, New York 10032, the ‡Department of Medicine, University of Washington, Seattle, Washington 98195, and the ¶Hospital for Special Surgery, New York, New York 10021

Background: Proteolytic cleavage of MerTK leads to inhibition of thrombosis and efferocytosis.

Results: In macrophages, lipopolysaccharide required reactive oxygen species to activate protein kinase Cδ and then p38 MAPK, culminating in ADAM17-mediated proteolysis of MerTK at proline 485.

Conclusion: ADAM17 is a key protease required during pattern recognition receptor-induced MerTK cleavage.

Significance: These findings uncover targets to test the consequences of MerTK cleavage in vivo.

Mer tyrosine kinase (MerTK) is an integral membrane protein that is preferentially expressed by phagocytic cells, where it promotes efferocytosis and inhibits inflammatory signaling. Proteolytic cleavage of MerTK at an unidentified site leads to shedding of its soluble ectodomain (soluble MER; sMER), which can inhibit thrombosis in mice and efferocytosis in vitro. Herein, we show that MerTK is cleaved at proline 485 in murine macrophages. Site-directed deletion of 6 amino acids spanning proline 485 rendered MerTK resistant to proteolysis and suppression of efferocytosis by cleavage-inducing stimuli. LPS is a known inducer of MerTK cleavage, and the intracellular signaling pathways required for this action are unknown. LPS/TLR4-mediated generation of sMER required disintegrin and metalloproteinase ADAM17 and was independent of Myd88, instead requiring TRIF adaptor signaling. LPS-induced cleavage was suppressed by deficiency of NADPH oxidase 2 (Nox2) and PKCδ. The addition of the antioxidant N-acetyl cysteine inhibited PKCδ, and silencing of PKCδ inhibited MAPK p38, which was also required. In a mouse model of endotoxemia, we discovered that LPS induced plasma sMER, and this was suppressed by Adam17 deficiency. Thus, a TRIF-mediated pattern recognition receptor signaling cascade requires NADPH oxidase to activate PKCδ and then p38, culminating in ADAM17-mediated proteolysis of MerTK. These findings link innate pattern recognition receptor signaling to proteolytic inactivation of MerTK and generation of sMER and uncover targets to test how MerTK cleavage affects efferocytosis efficiency and inflammation resolution in vivo.

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† To whom correspondence may be addressed: Northwestern University, Feinberg School of Medicine, Dept. of Pathology, 303 E. Chicago Ave., Tarry Bldg. 3-705, Chicago, IL 60611. E-mail: ethorp@northwestern.edu.

‡ To whom correspondence may be addressed. E-mail: iat1@columbia.edu.

MerTK (also known as c-Eyk, Nyk, and Tyro12) is a tyrosine kinase receptor for the growth arrest-specific protein GAS6 and anticoagulant Protein S (1, 2). Engagement of MerTK with either GAS6 or Protein S has been linked to numerous functions, including cell survival, thrombosis, and the phagocytosis of apoptotic cells (effectorcysis) (3–5). In the case of efferocytosis, both GAS6 and Protein S serve as bridging molecules that link MerTK to phosphatidylserine on dying cells (6). This leads to activation of intracellular signaling pathways that culminate in actin-driven apoptotic cell engulfment (7). MerTK is expressed predominantly in monocytic, epithelial, and reproductive tissue (8). In epithelial cells of the eye, naturally occurring mutations in MerTk are associated with onset of autosomal recessive retinitis pigmentosa (9). This is due to a defect of retinal pigment epithelial cells to promote clearance of adjacent light-sensing photoreceptor outer segments (10). Defects in MerTK are linked to other disease phenotypes. For example, in rodents, apoptotic thymocyte removal is defective in mice carrying a kinase-dead Merk (MerTkK226R) (3). MerTK deficiency in turn promotes autoantibody production and can stimulate lupus-like autoimmunity (11). Our group has shown that Merk deficiency promotes defective efferocytosis that is associated with increased vascular wall necrosis in advanced atherosclerotic plaque (12). Thus, MerTK has a critical anti-inflammatory role in a number of clinically relevant disease states.

At the structural level, MerTK is a type I transmembrane (TM) protein that encodes four extracellular domains: two fibronectin type-III domains and two extracellular immunoglobulin-like domains (13). Its cytoplasmic tail encodes a tyrosine kinase and controls distinct and separable effects that promote efferocytosis and inflammation resolution (8, 14). This domain homology is shared by two other molecules, Axl and...
Tyro3, to make up the TAM receptor family of tyrosine kinases (15). In some cases, a truncated TM-less isoform of MerTK can be generated by alternative mRNA splicing. A recent report by Sather et al. (16) indicates that a soluble form of MER (sMER) can also be induced through proteolytic cleavage of its ectodomain, leaving behind a carboxyl-terminal portion of the cleaved MER that remains cell-associated. In addition, sMER that is shed can act as a competitive inhibitor of MerTK during effec- rocytosis and platelet aggregation by acting as a decoy for its ligand GAS6. In our own hands (17) and others (18), sMER has been identified in inflammatory cardiovascular lesions, a dis- ease linked to defective effecrocytosis.

The identification of sMER seats MerTK in a growing list of TM-anchored protein receptors that are regulated by proteo- lytic shedding (19). Shedding of these cell surface proteins is often catalyzed by metalloproteinases. In the case of TAM receptor tyrosine kinases, mass spectrometric analysis indicates that closely related AXL is cleaved by the metallopro- tease linked to defective efferocytosis. In vivo, shedding of these cell surface proteins is implicated in the context of inflammatory cardiovascular lesions, a disease linked to defective effecrocytosis.

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MATERIALS AND METHODS

**Reagents**

**Antibodies**—Polyclonal goat anti-mouse MERTK was from R&D (catalogue no. AF591). For immunoblots, anti-goat IgG-HRP was also from R&D (catalogue no. HAF109). Rabbit polyclonal antibody to Adam17 was from ABCAM (ab2051). Rabbit antibody to Adam17 (phospho-Thr735) at 1 ng/ml stock was from ABCAM (ab60996). Adam10 antibody was ab1997. Total PKCβ antibody to SC-937 (C-20). Rabbit anti-phospho-PKCβ (Thr505) 9374S was from Cell signaling. Phospho-p38 MAPK (Thr180/Tyr182) 12F8 Rabbit was from Cell Signaling. Phospho-MKK3 (Ser189)/MKK6 (Ser186) 22A8 Rabbit monoclonal antibody was from Cell Signaling (catalogue no. 9236).

**Cleavage Inducers**—Lipopolysaccharide (LPS) purified by gel filtration chromatography was from Sigma (product number L4391) from *Escherichia coli* 0111:B4. Lipoteichoic acid and poly(I:C) were from InvivoGen. PMA and 4α-phorbol 12-myristate 13-acetate were from Sigma.

**Chemical Inhibitors**—Gö 6976 and Gö 6983 were from EMD Biosciences. p38 inhibitor, SB 202190, was from Sigma (cata- logue no. S7067) and used at 10 μM. TAPI-0 was from Calbiochem (catalogue no. 579050). N-acetyl cysteine was prepared fresh before use and was from Sigma.

Bryostatin 1 (catalogue no. B 7431) was from Sigma. 1,10-Phenanthroline monohydrate, reagent grade, in methanol was from Sigma (catalogue no. P9375).

**Detection Reagents**—5-((and -6)-chloromethyl-2',7'-di- chloro dihydrofluorescein diacetate, acetyl ester (CM- H2DCFDA) was from Invitrogen. Human Mer sandwich ELISA reagents from R&D DuoSet IC (DYC891-2).

**siRNA**

TLR4 siRNA (mouse) was from Santa Cruz Biotechnology, Inc. (Santa Cruz, CA) (catalogue no. sc-40261). PKCβ siRNA1 is Mm_Prkcd_2 (SI01388730) (target sequence CCG GGT GGA CAC ACC ACA CTA), and PKCβ siRNA2 is Mm_Prkcd_3 (SI01388737) (target sequence TTG AAT GTA GTT ATT GAA ATA) (Qiagen). Adam17 siRNA was from Qiagen (Mm_Adam17_6 SI02689190 target sequence CCG GGT GGA CAC ACC ACA CTA, and Mm_Adam17_5 SI02669261 target sequence CCC GGG TAT TAT TAT GGT TCA CAA, and Mm_Adam17_6 SI02689190 target sequence CCG GGT GGA CAC ACC ACA CTA, where Mm_Adam17_6 and Mm_Adam17_5 are isoforms of Adam17). Adam10 siRNA was from Qiagen (Mm_Adam10_5 SI02666062 target sequence CAC AGT GTG CAT TCA AGT CAA and Mm_Adam10_1 SI00165760 target sequence CCA GGA GAG AGA TAC ATT AAA). siRNAs were added to primary macro- phages and J774 cells with Lipofectamine 2000 from Invitrogen.

**Mice**

Wild-type macrophages were obtained from 8–10-week-old female C57Bl6/J mice (Jackson Laboratories). For Adam17-deficient studies, macrophages were from 8–10-week-old female Adam17fl/fl mice, which have normal ADAM17 expression, or from control littermate Adam17+/− mice, which have normal ADAM17 expression (26). 8–10-week-old female Nax2 mice were from Jackson, Strain B6.129S6-Cybb, stock number 002364. B6 Myd88−/− (009088) and Tfr−/− mice were also from Jackson.

**Isolation of sMER and Synthesis of MER Ectodomain Peptide for Mass Spectrometry**

sMER was isolated from serum-free medium over 80% con- fluent J774 cells after a 1-h treatment with 50 nM PMA. Cell supernatant was clarified by centrifugation to remove cellular and membrane debris. Clarified supernatant was immunopre- cipitated with polyclonal anti-MER (AF591) and protein A/G plus-agarose (sc-2003), and the sample was resolved by reducing SDS-PAGE. Confirmation of capture was performed by treating glycosylated MER extracellular domain with glycanase PNGase F. Post-PNGase treatment, sMER resolved at a molecular mass of 65 kDa, similar to the predicted size of the MerTK ectodomain. For the synthetic peptide, a peptide matching the
sequence of the semispecific Arg-C proteolytic fragment of MERTK (as described below) was custom synthesized by NeoBioscience (Cambridge, MA). The sequence is as follows: IAA ITK GGI GFP SEP VNI II P EHS KVD YAP. Its identity was confirmed by accurate mass and tandem mass spectrum.

**Mass Spectrometry**

In-gel sMER was subjected to proteolytic digestion. The gel was rinsed, reduced with DTT, and alkylated with iodoacetamide and digested with trypsin, chymotrypsin, or Arg-C overnight at 37 °C. Supernatant from the gel was collected, and the gel pieces were further extracted with 50% acetonitrile, 0.1% formic acid, and followed by 10% acetonitrile and 0.1% formic acid. The two washes were combined with the original supernatant, dried down, and suspended in 15 μl of 5% acetonitrile, 0.1% formic acid.

**LC-ESI-MS/MS**—Extracted in-gel digests were injected onto a C18 trap column (Magic AQ C18 200A, 5 μm, 0.1 × 20 mm, Michrom Biosources, Inc.), desalted for 15 min with water, 0.1% formic acid (4 μl/min), eluted onto an analytical column (Magic AQ C18 90A, 5 μm, 0.1 × 200 mm, Michrom Biosources, Inc.), and separated at a flow rate of 0.4 μl/min over 90 min, using a linear gradient of 5–35% acetonitrile, 0.1% formic acid in 0.1% formic acid on a NanoAquity HPLC (Waters, Milford, MA). Positive ion mass spectra were acquired with electrospray ionization in a hybrid linear ion trap-Orbitrap mass spectrometer (LTQ Orbitrap XL, Thermo Fisher, San Jose, CA) with data-dependent acquisition of MS/MS scans (linear ion trap) on the eight most abundant ions in the survey scan (Orbitrap, resolution 30,000). An exclusion window of 45 s was used after each repeated acquisition of the same precursor ion. Extracted in-gel digests and the synthetic peptide were further analyzed by targeted LC-ESI-MS/MS on the Arg-C semispecific proteolytic fragment at m/z 1045.5 (3+)+ and 784.4 (4+). A high resolution full scan MS in the Orbitrap (resolution 30,000) was altered with two targeted MS/MS scans with precursor selection window 2.5 Da in the linear trap and two high resolution MS/MS scans in the Orbitrap (resolution 10,000).

**Protein Identification**—For identification of MERTK, MS/MS spectra were matched against the mouse Uniprot/Swiss-Prot database (mouse version 3.54, April 2010), using the SEQUEST (version 2.7) search engine with fixed Cys carbamidomethylation and variable Met oxidation modifications and no enzyme specificity (semispecific restriction was applied to the results of the database search). The mass tolerance for precursor ions was 50 ppm (LTQ-Orbitrap data); SEQUEST default tolerance was accepted for product ions. SEQUEST results were further validated with PeptideProphet and ProteinProphet, using an adjusted probability of ≥0.90 for peptides and ≥0.95 for proteins. Each charge state of a peptide was considered a unique identification. Identity of the semispecific Arg-C proteolytic fragment was further confirmed by a Mascot database search (version 2.1, mouse SwissProt data base, v.XX, Matrix Science) on the MS/MS spectrum of the m/z 1045.5 (semitrystptic specificity, mass tolerance 50 ppm precursor, 0.4-Da fragments, modifications: fixed Cys +57.021, variable Met +15.99).

**Site-directed Mutagenesis and Analysis of Mutant MerTK Post-transfection**

Mutant MerTKs were generated from pIRES2-EGFP Mer from Addgene. Site-directed mutagenesis was performed using the QuikChange Lightning site-directed mutagenesis kit from Stratagene and in cooperation with Genewiz. Successful deletion and sequence integrity were confirmed by sequencing analysis at the Columbia University Core Sequencing Facility. For transfection assays, pIRES2-EGFP was used as a control. DNAs were transfected with Lipofectamine 2000 reagent from Invitrogen into HEK-293A cells (Invitrogen). Cellular MerTK and sMER were assessed post-transfection as described below. For efferocytosis analysis, apoptotic cells were labeled with Cell Tracker Orange from Invitrogen to measure internalization of apoptotic cells by fluorescent microscopy.

**Fluorescent Analysis of Peroxide Accumulation**

Macrophages were loaded with 5-(and 6)-chloromethyl-2',7'-dihydrodichlorofluorescein (DCF) diacetate ester (Invitrogen). After 30 min, the cells were washed and viewed immediately at room temperature with an inverted fluorescent microscope (IX-70) equipped with filters appropriate for fluorescein, and images were obtained with a charge-coupled device camera (Cool Snap) equipped with imaging software. Three fields of 700 cells/field were photographed for each condition, and the number of DCF-positive cells in each field was counted and expressed as a percentage of the total number of cells.

**Subcellular Fractionation**

To measure membrane translocation of PKCδ, primary macrophages were resuspended in 20 mm Tris-HCl (pH 7.5), 0.25 μM sucrose, 2 mM EGTA, 2 mM EDTA, and protease and phosphatase inhibitor mixture. Cells were subjected for sonication at 4 °C for 5 s. Cell lysates were subjected to centrifugation at 600 × g to remove nuclei and cellular debris. Supernatant was next spun at 100,000 × g for 1 h. Supernatant was soluble fraction, and pellet was membrane fraction. Proteins were resolved via reducing SDS-PAGE.
**Mechanism of MerTK Cleavage**

**Immunoblots**

Cell extracts were electrophoresed on 4–20% gradient SDS-polyacrylamide gels and transferred to 0.45-μm nitrocellulose membranes. The membrane was blocked in Tris-buffered saline, 0.1% Tween 20 (TBST) containing 5% (w/v) nonfat milk at room temperature for 1 h and then incubated with the primary antibody in TBST containing 5% (w/v) nonfat milk or 5% bovine serum albumin at 4 °C overnight, followed by incubation with the appropriate secondary antibody coupled to horseradish peroxidase. Proteins were detected by ECL Supersignal West Pico chemiluminescence (Pierce).

**Plasma Analysis**

Plasma was collected from the left ventricle of the heart after intraperitoneal injection of LPS, and ELISA was performed for sMER. Capture antibody (mouse MER affinity-purified polyclonal antibody, goat IgG, catalogue no. AF591, R&D) was overlaid onto ELISA plates at 0.2 μg/ml. Detection antibody (0.2 μg/ml) was mouse MER affinity-purified polyclonal antibody goat IgG (catalogue no. BAF591, R&D). Signal from MerTKKD mice was not above background. For TNFα, measurements were performed at the University of Maryland Cytokine Core Laboratory (Baltimore, MD). All experiments were performed in triplicate, and results were extrapolated from a standard curve.

**Statistical Analysis**

Results are presented as means ± S.E. Differences between multiple groups were compared by analysis of variance (one- or two-way), and differences between two groups were compared by paired or unpaired Student’s t test. p < 0.05 was considered significant. Stated n values are biological replicates.

**RESULTS**

**Identification of the MerTK Proteolysis Site by Mass Spectrometry**—To identify the site at which MerTK is susceptible to proteolysis, we induced cleavage and then immunopurified sMER from murine macrophage cell supernatants. Under these conditions, cleavage resulted in both generation of sMER and a reduction of cell surface MER as determined by flow cytometry (data not shown). SDS-PAGE-purified sMER was subjected in parallel to trypsin, chymotrypsin, and endoproteinase-Arg-C (clostripain) digestion and LC-MS/MS to identify MerTK. As an initial test of purity of the immunoprecipitated material, the combined results from the three separate proteolytic digests identified only peptides originating from the MerTK ectodomain. All three protease digests identified peptides in close proximity of the putative transmembrane domain (Fig. 1A). Significantly, the trypsin (K ↓ GGIGPFPSEPV-NIIIEHSK ↓ V) and chymotrypsin (F ↓ SEPVNIIEPEHS-KVDY ↓ A) proteolytic peptides were cleaved at enzyme-specific sites at both termini, whereas the Arg-C proteolytic peptide C terminus (R ↓ IAAITKGGIGPFSEPVNIIEPEHS-KVDYAP ↓ S) was after a proline and not after the usual site of Arg-C cleavage, arginine (Fig. 1B). Two independent MS search engines (Sequest and Mascot) identified the peptides with high confidence. We further synthesized the peptide IAAIT KGGIG PFSEP VNIII PEHSH VDYAP and subjected it to LC-MS/MS under the same conditions as above. Both synthetic and cell-derived Arg-C semispecific proteolytic peptides showed identical retention time, accurate mass at both 4+ and 3+ charge states (mass accuracy of <3 ppm), and MS/MS spectra of both 4+ and 3+ ions (Fig. 1C). Collectively, these data identify Pro485-Ser486 as the induced MerTK proteolytic site.

**Deletion of Six Amino Acids Spanning Proline 485 Renders MerTK Resistant to Induced Proteolysis and Efferocytosis Suppression by Cleavage Stimuli**—If amino acids including and proximal to proline 485 encode susceptibility to proteolysis, then targeted deletion of these residues could confer MerTK resistance to cleavage inducers. By site-directed mutagenesis, we engineered a six-amino acid deletion mutant of MerTK lacking amino acids 483–488 (Fig. 2A). Mutant MERTKK483–488 was transfected into HEK-293 cells, and shedding was induced by the addition of PMA, a known inducer of MerTK proteolysis (16). As indicated in Fig. 2B, expression of MerTKK483–488 was equal to WT expression by immunoblot, whereas generation of sMER was nearly completely abrogated in the mutant MerTK post-PMA treatment. A principle function of MERTK is to promote efferocytosis (3). To determine if MerTKK483–488 was functional, we transfected cleavage-resistant MertkK483–488 into HEK 293 cells, which do not express MerTK and do not engulf apoptotic cells (Fig. 2C, nontransfected cells). As indicated in Fig. 2C, and consistent with previous findings (7), transfection of WT Mertk induced the capacity of HEK cells to promote efferocytosis of UV-irradiated apoptotic Jurkat cells. The cleavage-resistant MerTK promoted efferocytosis in HEK cells to a comparable extent. We next measured efferocytosis after adding cleavage inducer PMA. Efferocytosis was significantly reduced in cells transfected with WT cDNA post-PMA; however, the mutant was resistant to PMA-induced efferocytosis suppression (p < 0.05). Thus, deletion of MerTK amino acids 483–488 confers resistance to induced MerTK cleavage and to suppression of efferocytosis by cleavage stimuli.

**LPS-induced MerTK Cleavage Requires TLR4-TRIF Signaling Independent of Myd88 and Is Inhibited by NADPH Deficiency**—We next sought to elucidate the signaling pathway that leads to generation of sMER. The two known inducers of MerTK cleavage are PMA and LPS. PMA (50 nm)-induced cleavage of MerTK from macrophages can be detected by immunoblot as early as 15 min and 1–2 h after 50 ng/ml LPS (16). Besides LPS, we asked if other prototypic inflammatory stimuli could acutely induce MerTK cleavage; however, sMER was not detected in cell supernatants after treating primary macrophages with TNFα or IFNγ for 1 h (Fig. 3A). Generation of sMER was concomitant with reductions in cell surface MerTK as determined by surface biotinylation (Fig. 3B). As expected, silencing of TLR4 with siRNA significantly inhibited LPS-induced sMER generation (Fig. 3C). Interestingly, LPS-mediated cleavage of sMER was not affected by Myd88 deficiency as indicated both by generation of sMER and reductions in full-length cell-associated MerTK (Fig. 3D). Similarly, inhibition of LPS-mediated NF-κB activation, which is downstream of MYD88 signaling, also failed to inhibit formation of sMER (data not shown). Instead, cleavage was suppressed by deficiency of the TLR4...
adaptor Trif (Fig. 3E). Consistent with a TRIF-mediated signaling pathway, the TLR3 ligand poly(I:C) (28) also was capable of inducing MerTK shedding (Fig. 3F).

Phagocytes produce reactive oxygen species (ROS) during phagocytosis or after stimulation with a wide variety of agents, including LPS (29). DCF staining was used as a measure of intracellular peroxide/ROS accumulation. We found that LPS caused a substantial increase in the percentage of DCF-positive cells at times when sMER can be detected (Fig. 4A). The ROS scavenger, N-acetylcysteine, inhibited sMER shedding (Fig. 4B) and DCF-staining (data not shown). Generation of ROS can occur through plasma membrane assembly
and activation of nicotinamide adenine dinucleotide phosphate reduced (NADPH) oxidase (NOX) (30). Shedding was significantly inhibited by Nox2 deficiency in LPS-treated macrophages (Fig. 4C).

PKCδ Is Required for LPS-induced MerTK Shedding—PMA, a PKC activator, induces robust MerTK cleavage (16). In our own hands, sMER was robustly induced by PMA; however, the PKC-inactive analog of PMA, 4α-PMA, failed to induce MerTK cleavage (data not shown). PKCs are a family of serine-threonine kinases, which are classified into three major groups based on homology and cofactor requirements: “conventional” PKCs, “novel” PKCs, and “atypical” PKCs (31). LPS-induced cleavage was inhibited by the pan–PKC inhibitor Go6983 (32) but not by classical PKC inhibitor Go6976 (Fig. 5A). In addition, co-cultivation of the atypical PKCζ pseudopeptide failed to suppress MerTK cleavage (data not shown). These data suggested that PKC activation did not involve members of the classical or atypical PKCs during MerTK cleavage. We next considered the novel PKCs, particularly PKCδ. Knockdown of PKCδ by two separate siRNAs each yielded greater than 78% reduction of basal PKCδ levels (Fig. 5B). LPS-mediated cleavage, after knockdown with each siRNA, was reduced in both instances (p < 0.05 in each instance). Consistent with a role for PKCδ after LPS activation in macrophages, both membrane-bound and phospho–PKCδ (at Thr505) were elevated 45 min after adding LPS (Fig. 5C). In addition, both PKCδ phosphorylation and membrane translocation were reduced after adding the antioxidant NAC, implicating PKCδ action downstream of NADPH activation during signaling, leading to generation of sMER (Fig. 5C, right).

MerTK Proteolytic Cleavage Requires ADAM17 and MAPK p38—Shedding of MerTK is inhibitable by TAPI-0 (16), a hydroxamate-based inhibitor of collagenase, gelatinase, and the membrane-associated protease ADAM17/TACE (33). To determine if germ line Adam17 is required for proteolytic cleavage of MerTK, we measured sMER production in Adam17fl/fl and Myd88-deficient primary macrophages. Adam17fl/fl completely inhibited LPS-mediated sMER generation (Fig. 6A). As shown in Fig. 6A, gene inactivation of Adam17 completely inhibited LPS-mediated sMER generation. Similar findings were seen after acute knockdown of Adam17 with siRNA in both primary macrophages (483–488) from the MERTK stalk, between the fibronectin-III ectodomain and the TM domain. B, Western blot for cellular MERTK and supernatant sMER after overnight transfection of WT or MERTK483–488 into HEK-293 cells. Post-transfection, cells were treated with or without PMA (50 nM) for 1 h, and cell supernatants and cell extracts were harvested for analysis. C, efferocytosis of apoptotic cells by HEK293 cells was measured by fluorescence microscopy after transfecting with wild type or mutant MerTK cDNA with or without PMA. p < 0.05, as indicated, n.s., not significant. Error bars, S.E.
phages and J774 macrophages (data not shown). In some cases, for example, when ADAM17 is inactivated or, alternatively, if cells are activated by ionomycin, the structurally similar ADAM10 can also shed ADAM17 substrates (34). However, siRNA-mediated reduction in ADAM10 (73% knockdown efficiency as indicated in Fig. 6B) did not reduce MerTK cleavage in WT macrophages. Thus, ADAM17 is the primary and non-redundant sheddase of MerTK.

We next considered how LPS might activate ADAM17. Previous reports suggest a role for MAPKs, including ERK1/2 and p38 during the phosphorylation or activation of ADAM17 (35–38). Although the ERK inhibitor PD98059 failed to reduce sMER levels, the p38 inhibitor SB 202190 ADAM17 (35–38). Although the ERK inhibitor PD98059 was injected into the peritoneum, and plasma was harvested 3 h later. Plasma sMER levels were then measured by ELISA. As shown in Fig. 8A, sMER levels were significantly increased after LPS injection in control mice. sMER was not detected above background in MerTK-deficient mice before or after LPS injection. In addition, sMER generation was dependent on ADAM17, because Adam17fl/fl/Lysmcre mice failed to induce sMER post-LPS injection. Consistent with previous results, LPS induced robust TNFα production in WT mice, and this was elevated in Mertk-deficient mice and suppressed in Adam17-deficient mice (Fig. 8B). Thus, sMER is induced by LPS treatment in vivo, and this requires ADAM17.

DISCUSSION

As expected, the cleavage site of MerTK does not conform to any previously documented motif for ADAM17 substrates. In fact, mutational analysis of ADAM17 substrates, such as the IL-6 receptor, suggests relaxed sequence specificity proximal to the ADAM17 cleavage site (40). Instead, the length of the membrane-proximal stalk has been implicated as a factor that con-

FIGURE 4. LPS-mediated MerTK cleavage requires NADPH. A, monolayers of elicited primary peritoneal macrophages were treated with 50 ng/ml LPS for 15 min to directly add LPS. Corresponding cell extracts of full-length MERTK are shown below. C, sMER generation by LPS was measured from Nox2-deficient cells by Western blot. Densitometric analysis of the ratio of sMER (SOL) to full-length MERTK (FULL) is to the right. *, p < 0.05. Error bars, S.E.

FIGURE 5. PKCδ is required for MerTK cleavage. A, primary macrophages were pretreated for 30 min with 250 nM Go6976 or 250 nM Go6983 in complete medium and subsequently treated with 50 ng/ml LPS. Subsequently, levels of sMER from cell supernatants and levels of full-length (FULL) MERTK from cell extracts were measured by Western blot. B, primary macrophages were incubated with PKCδ siRNA for 48 h, and the top panel exhibits representative knockdown efficiency of two PKCδ siRNAs (δ1 and δ2) by Western blot. In parallel, macrophages were cultured with LPS in the presence of PKCδ siRNA and scrambled (sc) control and sMER measured from supernatants and full-length MERTK from cell extracts by immunoblot. Densitometric analysis is shown to the right after knockdown with both PKCδ siRNAs. C, membrane translocation of PKCδ and phospho-PKCδ (PKCδ-P) post-LPS was determined by immunoblot after isolation of membrane pellets as described under “Materials and Methods.” Membrane translocation was also measured after treatment with NAC (right). M, membranous fraction; C, cytosolic fraction; T, total cellular lysate. Error bars, S.E.
controls susceptibility to cleavage (41, 42). Based primarily on its structure, MerTK is grouped into the TAM receptor family of tyrosine kinases, which include Tyro3, Axl, and Mertk. Mass spectrometric analysis indicates that TAM family member AXL is cleaved by ADAM17 (20), and shed AXL has been identified in both human and murine serum. The cleavage site of human AXL has been mapped to a 14-amino acid region proximal to the predicted TM domain (43, 44). Cleavage of TYRO3 has not been reported. Based on our own sequence analysis, the stalk regions of murine TAMs fail to exhibit a consensus motif for cleavage. However, murine MerTK and human MerTK do share a significant number of proline residues within their stalk region, leading us to speculate that human MerTK could also be cleaved after a proline.

The degradome of ADAM17 indicates a wide range of susceptible substrates. Therefore, how ADAM17 activation is finely regulated or, alternatively, a preference for specific cleavage substrates may be key to understanding substrate specificity under disparate homeostatic and pathophysiological contexts. Previous reports indicate that Gram-positive bacteria can stimulate the transcription of ADAM17 (45, 46). However, LPS-mediated cleavage was not inhibited by actinomycin D treatment (data not shown) or cycloheximide (16), implicating a post-translational mechanism. Furthermore, generation of
Mechanism of MerTK Cleavage

sMER was specific to the TLR4 agonist, Gram-negative endotoxin. The Gram-positive cell wall component lipoteichoic acid and TLR2 agonist (47) was unable to induce sMER shedding (data not shown). Although the link between LPS and ADAM17/TACE is well established, surprisingly little is known about the intermediary signaling molecules required. A previous study showed that endotoxin-induced MYD88 was upstream of ADAM17 processing during generation of EGF receptor ligands in nonhematopoietic cells (48). Although MerTK cleavage required TLR4, it was independent of MYD88 and instead signaled through TRIF. Cleavage could also be activated by the TLR3 agonist poly(I:C). Indeed, in epithelial cells, multiple Toll-like receptors, including TLR3, have been implicated in ADAM17-mediated shedding (49).

Mitochondrial ROS have been implicated in GPCR-induced TACE-dependent TGFβ shedding (50). In the case of MerTK shedding post-LPS, the ROS scavenger NAC and NADPH deficiency both blocked cleavage (Fig. 4). Although there are examples of TLR4 activation of NADPH through MYD88 (51), TRIF-mediated activation of NADPH is lacking. Interestingly, Park et al. (52) reported that NADPH oxidase subunits can directly interact with TLR4 to promote ROS generation. Furthermore, redox agents have been shown to regulate mature ADAM17 during neutrophil-mediated shedding of L-selectin (53). ROS has been suggested to activate ADAM17 (54, 55), in part through activation of PKCδ, and in some cell types, PKCδ is a redox-sensitive kinase (56, 57). A role for PKCδ in ADAM17 activation was previously implicated only based on data using the nonspecific PKC inhibitor Rottlerin. Once activated by ROS, PKCδ may in turn promote additional ROS activation (58).

Besides LPS, numerous other stimulators of ADAMs have been implicated. These include activators of protein kinase C, such as 12-O-tetradecanoylphorbol-13-acetate and PMA. Previously, a link between PKC signaling and LPS/TLR4 was shown for the PKC isozyme ε. PKCε was found to phosphorylate Trif-related adapter molecule downstream of LPS (59). In addition, PKCδ has been found to bind the TLR4/TLR2 adapter protein TIRAP/Mal (60). In vitro activation by LPS has been shown to induce rapid release of soluble FMS-like tyrosine kinase-1 receptor (sFlt-1), concomitant with phosphorylation of PKCδ (61). Furthermore, a PKCδ–p38 MAPK cascade has been identified in Caenorhabditis elegans (62). PKCδ-mediated activation of p38 may lead to p38 interaction with ADAM17 (37), and both p38 and ERK can phosphorylate ADAM17 at threonine 735 (35). However, activation of ADAM17 by PMA does not depend on its cytoplasmic domain, arguing against inside-out regulation via cytoplasmic phosphorylation as an underlying mechanism (24, 63). One possible explanation is that the ADAM17 cytoplasmic tail contains an inhibitory residue that must be phosphorylated for activation of ADAM17. Another possibility may be explained by differences in cell types utilized for the aforementioned studies. Finally, a still unidentified molecule may act to link PKCδ and p38 to ADAM17.

Proteolytic cleavage is known to regulate the activity of many transmembrane-anchored proteins. In the case of growth factors and cytokines, such as EGF, TFGα, and TNFα, proteolysis can lead to the biological activation of inactive precursors and their autocrine or paracrine release into the extracellular milieu. In the case of transmembrane receptors, such as TNFα receptor-I, TNF-α receptor-II, and L-selectin, proteolytic cleavage leading to ectodomain shedding can often lead to antagonist functions. Besides the loss of a cell surface signaling conduit, the shed ectodomains of cell surface receptors can also function as competitive decoys to bind receptor ligands. However, concentration is a critical factor. Low levels of soluble TNF receptor enhance TNFα action, whereas high concentrations are inhibitory (64). In the case of MerTK, recombinant sMER has been shown to be inhibitory by two accounts: first through suppression of efferocytosis in vitro and, second, through inhibition of thrombus formation in vivo (16). Interestingly, in the case of efferocytosis, LPS has been reported to inhibit the clearance of neutrophils in vitro, in part through induction of the ADAM17 target TNFα and suppression of macrophage-derived GAS6, the ligand for MerTK (65). These data suggest a coordinated response by macrophages to suppress MerTK function upon recognition of LPS. MerTK inactivation by cleavage would also suppress its anti-inflammatory function, thereby permitting the phagocyte to become fully activated. Future studies that seek to determine the in vivo/physiological relevance of MerTK cleavage, both in the context of bacterial challenge and during diseases of chronic inflammation and defective efferocytosis, will benefit from the identification of the cleavage site and signaling pathways revealed herein.

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