Sterile Inflammation in Acetaminophen-induced Liver Injury Is Mediated by Cot/tpl2*

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Background: MAP3K8 (Cot/tpl2) activates MKK1/2-Erk1/2 upon stimulation of receptors from the Toll-like/interleukin-1 receptor superfamily.

Results: Cot/tpl2 plays an essential role in acetaminophen-induced liver injury by modulating the generation of inflammatory signals induced by necrotic cells.

Conclusion: Sterile inflammatory processes triggered by tissue damage are modulated by Cot/tpl2.

Significance: Cot/tpl2 contributes to the development of pathologies associated with inflammation triggered by damage-associated molecular patterns.

Cot/tpl2 (MAP3K8) activates MKK1/2-Erk1/2 following stimulation of the Toll-like/IL-1 receptor superfamily. Here, we investigated the role of Cot/tpl2 in sterile inflammation and drug-induced liver toxicity. Cot/tpl2 KO mice exhibited reduced hepatic injury after acetaminophen challenge, as evidenced by decreased serum levels of both alanine and aspartate aminotransferases, decreased hepatic necrosis, and increased survival relative to Wt mice. Serum levels of both alanine and aspartate aminotransferases decreased hepatic necrosis, and increased reduced hepatic injury after acetaminophen challenge, as evidenced by decreased serum levels of both alanine and aspartate aminotransferases, decreased hepatic necrosis, and increased survival relative to Wt mice. Serum levels of both alanine and aspartate aminotransferases were also lower after intraperitoneal injection of acetaminophen in mice expressing an inactive aspartate aminotransferase. Moreover, similar activation profiles of intracellular pathways were observed in Wt macrophages stimulated with Wt or Cot/tpl2 KO damage-associated molecular patterns. However, upon stimulation with damage-associated molecular patterns, the activation of Erk1/2 and JNK was deficient in Cot/tpl2 KO macrophages compared with their Wt counterparts; an effect accompanied by weaker release of several cytokines, including IL-1α, an important component in the development of sterile inflammation. Taken together, these findings indicate that Cot/tpl2 contributes to acetaminophen-induced liver injury, providing some insight into the underlying molecular mechanisms.

The liver is a crucial metabolic organ and is highly susceptible to drug toxicity. Acetaminophen (APAP) is one of the best-selling analgesics and antipyretics on the market in the United States and Western Europe, although it also accounts for ~50% of all cases of acute liver failure (1–3). APAP overdose generates N-acetyl-p-benzoquinone imine, which depletes hepatic glutathione and promotes oxidative stress, leading finally to hepatocyte necrosis (reviewed in Refs. 4, 5). Necrosis occurs when cells die rapidly in response to acute injury, ultimately provoking the release of intracellular constituents into the surrounding milieu (6). Some of these components released belong to a group of molecules known as damage-associated molecular patterns (DAMPs). Outside of their physiological environment, these molecules behave as “danger sensors” and are capable of triggering an inflammatory response in a similar way as previously described for the pathogen-associated molecular patterns (PAMPs) (reviewed in Refs. 7, 8). Similar to pathogen-induced inflammation, sterile inflammation initiated by necrotic cells alerts the host tissue to host damage by activating the innate immune system (9, 10). Initially, this process is manifested by the production of pro-inflammatory mediators and triggering the recruitment of leukocytes to the damaged tissue (7, 11–13). However, when excessive, this inflammatory response can contribute to severe organ damage and dysfunction (7, 11, 14). DAMPs include a wide variety of molecules, and the number of known DAMPs is growing continually (7, 15–18).

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4 The abbreviations used are: APAP, acetaminophen; BMDM, bone marrow-derived macrophages; DAMP, damage-associated molecular pattern; PAMP, pathogen-associated molecular pattern; TLR, Toll-like receptor; LPS, lipopolysaccharide; i.p., intraperitoneal; LDH, lactate dehydrogenase; MFI, mean fluorescence intensity.
DAMPs activate a variety of receptor types in different cells, among which are the members of the pattern-recognition receptor (PRR) family, including the receptors of the Toll-like receptor (TLR) family (7, 10, 16, 18, 19). Upon activation, all TLRs (except TLR3) recruit the MyD88 adaptor protein, which mediates activation of the p38α and JNK MAP pathways, and the canonical IKKs, IKKa, and IKKβ (20, 21). Activated IKKβ phosphorylates p105 NFκB, marking it for partial proteolysis. In resting cells, Cot/tpl2 (MAP3K8) forms an inactive complex with p105 NFκB and ABIN2, from which Cot/tpl2 is released following proteolysis of p105 NFκB (reviewed in Refs. 22, 23). The dissociated and activated Cot/tpl2 stimulates MKK1/2 and consequently Erk1/2 (24–26), and it is subsequently rapidly degraded through the proteasome pathway (27, 28). Cot/tpl2 is the only MAP3K to activate the Erk1/2 pathway in response to both TLR activation and IL-1 or TNFα stimulation (29, 30) in different cell types, including macrophages, epithelial, and stellate cells (25, 29, 31). Moreover, Cot/tpl2 can also activate the MAP kinases JNK and p38α in certain conditions, in a cell type- and stimulus-specific manner (22, 30). Thus, Cot/tpl2 fulfills a role in the development of acute pancreatitis (37). However, the role of Cot/tpl2 in Acetaminophen-induced Liver Damage

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**EXPERIMENTAL PROCEDURES**

**Cells and Stimuli—**Immortalized Wt and Cot/tpl2 KO hepatocytes were generated from pools of 4–6 livers obtained from Wt or Cot/tpl2 KO neonatal mice (3.5–4-day-old) that were digested with collagenase and then cultured. The cells were subsequently immortalized with the SV40 Large T antigen, as described previously (39). Hepatocytes were incubated for 18 h with APAP (Sigma-Aldrich) and/or TNFα (Peprotech), and their viability was then assayed using the MTT kit (Roche), according to the manufacturer’s instructions. DAMPs were generated from Wt and Cot/tpl2 KO hepatocytes as described previously (12, 13) with minor modifications: prior to 1-h heat-shock at 60 °C, the hepatocytes were washed five times with PBS and resuspended in DMEM-HEPES. Bone marrow-derived macrophages (BMDM) or peritoneal macrophages were generated as described previously (26) and then stimulated with DAMPs (300 μg/ml), LPS (300 ng/ml, Sigma-Aldrich), or PMA (10 μM, Sigma-Aldrich). When necessary, the IKKβ inhibitor B1605906 (10 μM), the JNK inhibitor SP 600125 (12 μM), or the Erk1/2 inhibitor PD 0325901 (0.5 μM), gifts from Sir Philip Cohen (Dundee), were added prior to stimulation.

**Animals and Animal Treatments—**C57BL/6 Wt, C57BL/6 KO, and C57BL/6 Cot/tpl2 KD littermates were generated from the crossing of heterozygous mice (34), and they were used for experiments at 10–12 weeks of age. All animals were handled in accordance with institutional guidelines for the care and use of laboratory animals in research. The mice were fasted overnight before treatment, and the APAP solution administered (10 mg/ml) was prepared freshly in warmed (55 °C) PBS that was cooled to 35 °C before it was injected. For survival studies, 600 g/kg APAP (intraperitoneal) was administered to mice. A comprehensive laboratory animal monitoring system (TSE) was used to analyze metabolic behavior 4 h after injection, measuring O2 consumption and CO2 production every 10 min for 48 h. The time of death was determined as the point at which the respiratory exchange ratio (RER; O2/CO2) became zero or negative. To evaluate hepatotoxicity the animals were administered intraperitoneally (i.p.) either with PBS (control) or with APAP (450 g/kg), food was withheld for 4 h, and at the times indicated, the mice were sacrificed. Liver tissue was taken from each mouse, immediately ground into small fragments, frozen in liquid nitrogen, and stored at −80 °C. Blood was collected, and the serum was stored at −80 °C until use. The ALT, AST, and lactate dehydrogenase (LDH) levels were determined (at the UCM, Madrid), and cytokine levels were assessed by Luminex analysis (at the CNB, Madrid). Peritoneal leukocyte recruitment in mice was measured 18 h after injection of liver homogenate (32 mg, intraperitoneal) or of DAMPs generated from hepatocytes (1.7 mg, 35 × 10⁶ cells). Peritoneal cells were collected as described previously (12, 13, 34). Hepatic non-parenchymal cells were isolated as described previously (40). To analyze liver damage, tissue samples were rinsed with saline and fixed by immersion in 4% formalin for 24 h and treated with hematoxylin and eosin (H&E) staining as described previously (34). The analysis of the necrotic areas as well as the quantification (Image J) of the liver areas with congestive changes occupied by erythrocytes (including the intravascular and hemorrhagic areas) were performed in seven randomly selected microscopic fields from each sample using a 10× objective.

**Flow Cytometry—**Cells (0.3–0.5 × 10⁶ cells/test) were incubated with CD16/32 (2.4G2, Caltek) for 20 min at room temperature, and they were subsequently stained for 1 h at 4 °C in the dark with the following antibodies (5 μg/ml): CD3-PECy7 (Hamster IgG, ebioscience), CD11a-PECy7 (rat IgG2ak, Pharmingen), CD11b-(Mac1)-PECy7 (rat IgG2bk anti-mouse, ebioscience), CD45-FITC (rat IgG, Beckman), F4/80-APC (rat IgG2ak, ebioscience), Ly6G-PE (rat IgG2ak, Pharmingen), and NK1.1-APC (mouse IgG2ak anti-mouse, Pharmingen), or their corresponding isotype controls (Pharmingen, ebioscience). After three washes, Perfect-Count microspheres (Cytognos) were added to quantify the exact number of cells. Flow cytometry analysis was performed using the CXP program. To determine the oxidative burst of peritoneal leukocytes, 3 × 10⁶ cells were resuspended in PBS, incubated in the dark with 10 μM...
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2’,7’-dichlorofluorescin diacetate (Invitrogen), and subsequently incubated in the presence or absence of 10 μM PMA for 15 min at 37 °C. After several washes, the cells were analyzed by flow cytometry.

**Western Blot and RT-PCR Analysis**—Homogenized liver extracts and cell extracts were analyzed in Western blots (41), probed with primary antibodies raised against the following proteins: Cot/tpl2, Erk2, p38α, and p52 JNK2 (Santa Cruz Biotechnology); P-S933 p105 NFkB, P-T202/Y204 Erk1/2, and P-T180/Y182 p38α (Cell Signaling); and P-T183/Y185 p48/p52 JNK (Invitrogen). Secondary antibodies raised against rabbit (Cell Signaling), goat (DAKO), and mouse (Amersham Biosciences) were used to detect the primary antibodies. RNA extraction and RT-PCR analysis were performed as described previously (42). The specific TQA MN primers (Applied Biosystems) IL-1β, IL-10, TNFα, and β-actin were used.

**Statistical Analysis**—Data are presented as the mean ± S.D., and they were analyzed using the Student’s t test. Values were considered statistically significant at p < 0.05: *, p < 0.01, **, p < 0.001.

**RESULTS**

**Cot/tpl2 Participates in APAP-induced Liver Damage**—Following APAP administration (600 g/kg, intraperitoneal), 12% survival was observed in Wt mice in the 15 to 30 h following injection, whereas all the Cot/tpl2 KO mice survived for the duration of the experiment (Fig. 1A). The serum levels of ALT, AST, and LDH were all lower in APAP-treated (450 g/kg, intraperitoneal) Cot/tpl2 KO mice than in their Wt counterparts (Fig. 1B and supplemental Fig. S1). Increased hepatic P-JNK levels, as a marker of liver damage, were observed 6 h after APAP-induced liver injury (4, 43), and Cot/tpl2 deficiency decreased JNK phosphorylation in APAP-injected mice (supplemental Fig. S1). Moreover, the liver of Cot/tpl2 KO mice revealed a marked histological attenuation of liver injury compared with the severe centrilobular necrosis and hemorrhage observed in Wt animals (Fig. 1C), indicating that Cot/tpl2 deficiency protected against APAP-induced hepatotoxicity. In Cot/tpl2 KO mice that express an inactive version of Cot/tpl2 (42), the levels of AST and ALT after APAP treatment were also diminished compared with those in Wt mice (Fig. 1D). These data indicate that APAP-induced liver injury is modulated by Cot/tpl2 activity rather than by modifications in the Cot/tpl2-ABIN2-p105 NFkB complex, because of Cot/tpl2 knockdown.

APAP-induced liver injury triggers the recruitment of leukocytes to the liver (8, 44, 45). Upon intraperitoneal injection of PBS, similar numbers of hepatic leukocytes were detected in Wt and Cot/tpl2 KO mice; yet APAP injection produced a 3-fold increase in the number of hepatic leukocytes (CD45+) in Wt mice, whereas only a 1.5-fold increase was observed in APAP-treated Cot/tpl2 KO mice (Fig. 2A). Analysis of the leukocytes recruited revealed that Cot/tpl2 deficiency mainly decreased the number of infiltrated neutrophils (Ly6G+) and macrophages (F4/80+; Fig. 2, B and C). In the neutrophils, neither Cot/tpl2 deficiency nor APAP treatment altered the mean fluorescence intensity (MFI) of the CD11a and CD11b activation markers. However, the macrophages recruited to the liver following APAP administration had a higher CD11b MFI (44), an effect that was less pronounced in Cot/tpl2 KO macrophages (Fig. 2D). Following APAP challenge, Cot/tpl2 also triggered the recruitment of NKT (CD3' NK1.1) cells to the liver (Fig. 2B). Moreover, the serum IL-1α, IL-1β, and IL-6 levels were lower in APAP-treated Cot/tpl2 KO mice than in Wt mice, while Cot/tpl2 deficiency promoted an increase in IL-10 levels (Fig. 3A). Similarly, 6 h upon APAP challenge, the liver of Cot/tpl2 KO mice showed decreased mRNA levels of IL-1α, IL-1β, IL-6, and TNFα, but higher IL-10 levels compared with Wt-treated mice (Fig. 3B).

To determine whether Cot/tpl2 also modulates APAP-induced hepatotoxicity in isolated hepatocytes, immortalized Wt and Cot/tpl2 KO hepatocytes were incubated with different concentrations of APAP for 18 h. Similar survival rates were observed for both Wt and Cot/tpl2 KO cells in the presence of APAP (supplemental Fig. S2), and although the presence of TNFα along with APAP further increased cell toxicity, Cot/tpl2 did not appear to modulate hepatocyte cell death in either of these conditions (Fig. 4A). Accordingly, similar LDH levels, as a consequence of cell necrosis, were detected in the supernatant of both Wt and Cot/tpl2 KO hepatocytes following incubation for 18 h with APAP plus TNFα (Fig. 4B). Furthermore, in APAP plus TNFα-treated hepatocytes Cot/tpl2 deficiency did not modulate the activity of JNK (Fig. 4C).

**Involvement of Cot/tpl2 in Macrophage Activation by DAMPs**—We evaluated the role of Cot/tpl2 in the activation of thioglycolate-elicited peritoneal macrophages by DAMPs obtained from Wt necrotic hepatocytes. After DAMPs stimulation, a decrease in IL-10, IL-1α, IL-1β, IL-6, and TNFα was detected in the supernatant of Cot/tpl2 KO with respect to Wt peritoneal macrophages (Fig. 5A). When the activation state of different intracardial pathways was analyzed in thioglycolate-elicited peritoneal macrophages stimulated with DAMPs, p105 NFkB phosphorylation, an upstream effector of Cot/tpl2, was similarly increased in both Wt and Cot/tpl2 KO macrophages (Fig. 5B). Moreover, in Wt macrophages, we observed the degradation of both the high and low molecular weight forms of Cot/tpl2, suggesting that it had dissociated from the p105 NFkB. Moreover, in APAP-treated Wt KO macrophages resulted in an increase in Erk1/2 phosphorylation over 7 min that decreased rapidly in Cot/tpl2-deficient cells. In Wt peritoneal macrophages but not in Cot/tpl2 KO cells, a second increase of P-Erk1/2 levels was observed 30 min after stimulation (Fig. 5B). This early Erk1/2 activation peak following macrophage stimulation with DAMPs was not due to a possible FBS contamination contained in the DAMPs (supplemental Fig. S3). Activation of different TLRs in Wt and Cot/tpl2 KO peritoneal macrophages, as described previously in BMDM (46), triggered the phosphorylation of p105 NFkB by IKKβ, whereas Erk1/2 phosphorylation was observed only in Wt macrophages, evident as a single activation peak (supplemental Fig. S4). Cot/tpl2 deficiency decreased JNK activation, but not p38α, in peritoneal macrophages stimulated with Wt DAMPs (Fig. 5B). The loss of Cot/tpl2 expression in Cot/tpl2 KO hepatocytes did not affect the capacity of the obtained DAMPs to phosphorylate p105 NFkB or the different MAP kinases, as observed when BMDM were stimulated with Wt or...
Cot/tpl2 KO DAMPs (supplemental Fig. S5). As detected in peritoneal macrophages (Fig. 5B), the deficient expression of Cot/tpl2 in BMDM abolished the 30–40 min Erk1/2 activation and decreased JNK phosphorylation following the stimulation with Wt or Cot/tpl2 KO DAMPs (Fig. 5C and supplemental Fig. S5). Furthermore, in Wt BMDM stimulated with DAMPs, the IKKβ inhibitor B1605906 blocked the activation of Erk1/2 at 40 min but not that at 5 min (Fig. 5D). We previously demon-
strated that Erk1/2 activation by LPS is entirely dependent on Cot/tpl2, whereas Erk1/2 activation by PMA is Cot/tpl2-independent (24). Indeed, IKK-H9252 inhibition blocked LPS- but not PMA-induced Erk1/2 activation (supplemental Fig. S6). Together, these data indicate that two distinct intracellular signal pathways regulate Erk1/2 activation upon DAMPs stimulation of macrophages, one IKK-H9252-Cot/tpl2-dependent and another IKK-H9252-Cot/tpl2-independent. Furthermore, taking into account that Cot/tpl2 deficiency impaired both Erk1/2 and JNK MAP kinases in the production of different cytokines. To this end, Wt peritoneal macrophages were stimulated for 18 h with DAMPs in the presence of the JNK inhibitor SP 600125 or in the presence of the Erk1/2 inhibitor PD 0325901. The measurement of IL-10, IL-1α, IL-1β, IL-6, and TNFα levels in the different cell supernatants suggested that Cot/tpl2 mainly controls the production of the different cytokines by its capacity to mediate Erk1/2 activation in DAMPs-stimulated macrophages (supplemental Fig. S7).

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FIGURE 2. Leukocyte recruitment in the liver of Wt and Cot/tpl2 KO mice following APAP injection. A, mice were injected intraperitoneally with PBS (Control) or APAP (450 g/kg), and CD45+ liver cells were isolated 18 h later and analyzed by flow cytometry. B, analysis of Ly6G−, F4/80+, CD3+ (CD3−NK1.1−), NK1.1+ (NK1.1+CD3−), and NKT (CD3−NK1.1−) cells from the CD45+−gated cells in A. C, representative Ly6G+ versus F4/80+ staining FACS profile from the CD45+−gated cells obtained in A. D, CD11b and CD11a MFI of F4/80+ and Ly6G+ cells isolated from the livers of Wt and Cot/tpl2 KO mice 18 h after intraperitoneal injection with PBS (Control) or APAP (450 mg/kg). Graphs represent the mean ± S.D. of one experiment performed in quadruplicate and similar data were obtained in three additional experiments. Data in A and B represent the mean ± S.D. of four independent experiments performed in quadruplicate.

Cot/tpl2 Modulates Sterile Inflammation in Vivo—Intrapерitoneal injection of DAMPs induces the recruitment of leukocytes to the peritoneum (12, 13). Cot/tpl2 deficiency in mice significantly decreased the peritoneal recruitment of both macrophages and neutrophils 18 h after intraperitoneal injection of Wt or Cot/tpl2 KO DAMPs (Fig. 6, A and B). However, Cot/tpl2 deficiency did not alter the CD11a and Cd11b MFI values of the cells recruited (supplemental Fig. S8). Similarly, compared with Wt mice, Cot/tpl2 KD mice showed reduced peritoneal recruitment of both macrophages and neutrophils 18 h after intraperitoneal injection of Wt liver homogenate (supplemental Fig. S9). Oxidative burst analysis of the Wt and Cot/tpl2 KO peritoneal leukocytes recruited indicated that Cot/tpl2 deficiency slightly reduced their oxidative burst capacity, a difference that augmented in Cot/tpl2 KO leukocytes upon PMA stimulation (Fig. 6C). On the other hand, following intraperitoneal injection of Wt DAMPs, peritoneal levels of IL-1α, IL-1β, and IL-6 diminished in Cot/tpl2 KO mice compared with their Wt counterparts (Fig. 6D).
APAP overdoses are the number 1 cause of acute liver failure in the Western world, accounting for 50% of them (1–3). The present study demonstrates the fundamental role of Cot/tpl2 activity in APAP-induced liver damage. APAP-induced liver injury is associated with sterile inflammation (8, 44, 45), and Cot/tpl2 participates in its development by modulating the production of cytokines and the liver recruitment of leukocytes. In this context, it has been previously shown that TLR9 KO mice, with reduced levels of inflammatory cytokines, show resistance.

**DISCUSSION**

![Graphs showing the modulation of cytokine levels.](image1)

**FIGURE 3.** *Cot/tpl2 modulates cytokine production associated with APAP-induced liver injury.* Wt and Cot/tpl2 KO mice were challenged with APAP (450 g/kg, intraperitoneal), and cytokines levels were determined. A, serum levels of IL-1α, IL-1β, IL-6, and IL-10 were determined 18 h later using a Luminex assay. TNFα could not be detected. B, IL-1α, IL-1β, IL-6, IL-10, and TNFα mRNA expression levels upon 6 h of APAP challenge were determined by RT-PCR analysis. A and B, data represent the mean ± S.D. of four independent experiments performed in quadruplicate.

![Graphs showing cytokine expression.](image2)

**FIGURE 4.** *Toxicity induced by APAP plus TNFα in Wt and Cot/tpl2 KO hepatocytes.* A, Wt and Cot/tpl2 KO hepatocytes were incubated with 10 ng/ml of TNFα in combination with different concentrations of APAP, and their viability was assessed 18 h later through a MTT assay. The value obtained for Wt and Cot/tpl2 KO hepatocytes incubated in the absence of APAP was considered as 100%. B, LDH was determined in the cell supernatant of Wt and Cot/tpl2 KO hepatocytes incubated with 10 ng/ml of TNFα plus 20 mM of APAP for 18 h. The 100% value is given to the one obtained after Triton X-100 permeabilization. C, hepatocytes were incubated with 20 mM APAP plus 20 ng/ml TNFα for 0, 2, and 3 h, and P-JNK and JNK2 levels were measured by Western blot. The relative relation of P-JNK levels/JNK2 levels was determined by densitometric quantification of the radiographs, given the value of 1 to the one obtained with Wt hepatocytes at time 0. Similar data have been obtained in three independent experiments. A and B, graphs show the mean ± S.D. of three independent experiments performed in triplicate.
to acetaminophen toxicity (47). While some controversy surrounds the role of macrophages in the pathogenesis of APAP-mediated liver damage, neutrophil recruitment has been implicated in this process (40, 48–50). Indeed, neutrophil infiltration is associated with increased tissue necrosis because of the release of cytotoxic agents such as reactive oxygen species (51).

APAP overdose generates in hepatocytes the N-acetyl-p-benzoquinone imine that depletes glutathione and then generates oxidative stress reactions, leading to hepatocyte necrosis (4). DAMPs, once released into the surrounding tissue as a consequence of cell necrosis, orchestrate an inflammatory response, which in many cases can cause a second wave of destruction contributing to the pathogenesis of many damaging conditions (7, 11, 14). DAMPs are recognized by different receptors of the PPR family, including different TLRs (7, 10, 16). However, the i.p. recruitment of inflammatory cells upon intraperitoneal injection of DAMPs is not affected by the loss of a single TLR (12), although the inflammatory response to DAMPs is significantly attenuated in IL-1R and MyD88 KO mice (12). Here, we show that Cot/tpl2 participates in the i.p. recruitment of leukocytes in response to stimulation with DAMPs. IL-1R and virtually all TLRs (except TLR3) signal via MyD88 (20). Indeed, Cot/tpl2, activated via MyD88 or Trif, triggers the MKK1-Erk1/2 pathway following stimulation of IL-1R and TNFα receptors and also of all the different TLRs, including TLR3 (29, 30, 46).

Here we show that Cot/tpl2 once activated mediates, both in vivo and in vitro, IL-1α and IL-1β production. Sterile inflammation is largely dependent on IL-1α. Indeed, sterile inflammation requires this cytokine to a greater extent than pathogen-induced inflammation and its role in sterile inflammation cannot be substituted by IL-1β or any other cytokine (52–55).

Here we show that Cot/tpl2 once activated mediates, both in vivo and in vitro, IL-1α and IL-1β production. Sterile inflammation is largely dependent on IL-1α. Indeed, sterile inflammation requires this cytokine to a greater extent than pathogen-induced inflammation and its role in sterile inflammation cannot be substituted by IL-1β or any other cytokine (52–55).
that observed in individual isolated cells. Cot/tpl2 controls TNFα synthesis upon i.p. injection of LPS/d-galactosamine in mice (25) but not following zymosan-induced intraplantar inflammation or following an infection with Listeria monocytogenes (34, 57). Conversely, Cot/tpl2 deficiency in APC KO mice results in a decrease in serum IL-10 (58), while in LPS and CpG-DNA-treated mice, Cot/tpl2 blocks IL-10 production (59). Our findings demonstrate that Cot/tpl2 only mildly increases peritoneal IL-10 levels upon injection with DAMPs, and it decreases serum IL-10 levels in response to APAP-induced liver injury. Although IL-10 protects against APAP-induced liver injury (60), it remains to be determined whether the

**FIGURE 6. Leukocytes recruitment following DAMPs-induced peritonitis in Wt and Cot/tpl2 KO mice.** Animals were injected (intraperitoneal) with Wt or Cot/tpl2 KO DAMPs (1.7 mg) or with PBS (Control), and peritoneal cells were isolated 18 h later. A, number of isolated peritoneal macrophages (F4/80^+^) and neutrophils (LY6G^+^) recovered after injection of Wt or Cot/tpl2 KO DAMPs in Wt and Cot/tpl2 KO mice. B, one representative Ly6G versus F4/80^+^ staining FACS profile of Wt and Cot/tpl2 KO peritoneal cells isolated 18 h after injection of Wt DAMPs in Wt and Cot/tpl2 KO mice. C, oxidative burst activity of isolated Wt and Cot/tpl2 KO peritoneal cells incubated in the presence or absence of PMA (10 μM) in vitro. The relative induction is expressed in terms of that obtained in unstimulated Wt cells. D, levels of the indicated cytokines in the peritoneal cavity of Wt and Cot/tpl2 KO mice treated as described in B, A, C, and D, graphs represent the mean ± S.D. of three independent experiments performed in triplicate.
increased levels of IL-10 in Cot/tpl2 KO mice at least in part attenuate liver injury.

Tissue damage is recognized at the cell level. Different DAMPs are recognized by the different receptors of the PPR family. The failure of DAMPs from necrotic cells to express specific molecules modifies their capacity to develop sterile inflammation (7, 15–18). However, dampened Cot/tpl2 expression in hepatocytes does not affect the capacity of their obtained DAMPs to activate macrophages in vitro or to initiate the sterile inflammation response in vivo.

Sterile inflammation is initiated by DAMPs that upon stimulation of different cell types including macrophages (12), produce IL-1α a major regulator of sterile inflammation (52–55). In macrophages, DAMPs activate Erk1/2, and our data indicate that Cot/tpl2 mediates the activation of Erk1/2 after 30 min but not after 5 min following stimulation. This rapid increase in Erk1/2 phosphorylation is probably mediated by the activation of another MAP3K, most likely one of the RAF proteins. Cot/tpl2 is the only MAP3K to activate the Erk1/2 pathway in response to both TLR activation and IL-1 or TNFα stimulation (29, 30). However, cell stimulation with DAMPs results in activation of a variety of receptors (7), thus the two different intracellular pathways involved in Erk1/2 phosphorylation under this cellular condition. Cot/tpl2 can also activate JNK (22, 30), and here we show that Cot/tpl2 partially mediates JNK phosphorylation upon macrophage activation with DAMPs. However, the decreased production of cytokines in the macrophages from Cot/tpl2 KO mice is due to impaired activation of the Erk1/2 intracellular signal pathway. In conclusion, our data show that Cot/tpl2 participates in sterile inflammatory pathways triggered by damaged tissue and plays an essential role in APAP-induced liver injury.

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