Cholesterol Crystals Induce Coagulation Activation through Complement-Dependent Expression of Monocytic Tissue Factor

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Cholesterol crystals (CC) are strong activators of complement and could potentially be involved in thromboinflammation through complement–coagulation cross-talk. To explore the coagulation-inducing potential of CC, we performed studies in leprinubin-based human whole blood and plasma models. In addition, immunohistological examinations of brain thrombi and vulnerable plaque material from patients with advanced carotid atherosclerosis were performed using polarization filter reflected light microscopy to identify CC. In whole blood, CC exposure induced a time- and concentration-dependent generation of prothrombin fragment 1+2 (PTF1.2), tissue factor (TF) mRNA synthesis, and monocyte TF expression. Blocking Abs against TF abolished CC-mediated coagulation, thus indicating involvement of the TF-dependent pathway. Blockade of FXII by corn trypsin inhibitor had a significant inhibitory effect on CC-induced PTF1.2 in platelet-free plasma, although the overall activation potential was low. CC exposure did not induce platelet aggregation, TF microparticle induction, or TF on granulocytes or eosinophils. Inhibition of complement C3 by CP40 (compstatin), C5 by eculizumab, or C5aR1 by PMX53 blocked CC-induced PTF1.2 by 90% and reduced TF+ monocytes from 18–20 to 1–2%. The physiologic relevance was supported by birefringent CC structures adjacent to monocytes (CD14), TF, and activated complement iC3b and C5b–9 in a human brain thrombus. Furthermore, monocyte influx and TF induction in close proximity to CC-rich regions with activated complement were found in a vulnerable plaque. In conclusion, CC could be active, releasable contributors to thrombosis by inducing monocyte TF secondary to complement C5aR1 signaling. The Journal of Immunology, 2019, 203: 853–863.

The World Health Organization estimates that cardiovascular disease causes more than 17 million deaths every year. Of these, 80% are due to myocardial infarction or ischemic stroke as a result of a thrombus obstructing blood flow to the heart or brain, respectively (1). Thrombosis occurs following atherosclerotic plaque rupture and involves lesion-derived tissue factor (TF) from endothelial cells (2), plaque macrophages (3, 4), and blood monocytes following exposure to oxidized LDL and C-reactive protein (5, 6). Cholesterol crystals (CC) formed upon oxidized LDL accumulation at the lesion sites are hallmarks of atherosclerotic plaques (7). Their sharp, needle-like structure promotes erosion of the fibrous cap and thereby plaque rupture with release into blood (8). Except for their role in promoting plaque rupture, CC are considered passive participants in the thrombotic events.

CC activate the complement system (9, 10), whereas their role as direct promoters of coagulation has not yet been investigated. The recent finding of CC being strong inducers of complement-dependent inflammatory processes (11) raises the question of whether...
CC could activate coagulation through thromboinflammation. This question is further justified in light of evidence of cross-talk between the complement and coagulation cascades (12–16).

The complement and coagulation cascades are first-line defense systems mediating inflammation and clot formation, respectively. Both consist of circulatingzymogens converting to reactive proteases upon proteolytic activation (12). Complement can be activated through the classical, lectin, or alternative pathways, converging in the formation of C3 convertases. Downstream formation of C5 convertases cleaves C5 and leads to formation of C5a, a highly potent proinflammatory mediator (17), and the terminal complement complex (TCC) C5b–9. Of note, CC induce activation of all three complement pathways (18).

The coagulation system is activated either through the intrinsic pathway, activating FXII to FXIIa on a surface, or through the extrinsic pathway by binding of FVIIa to cellular expressed TF. Both pathways converge in the activation of FX to FXa, with subsequent cleavage of prothrombin to thrombin. This leads to the conversion of fibrinogen to insoluble fibrin. Together with activated platelets, this process builds up a stable thrombus. TF is the major initiator of blood coagulation in vivo and is, under normal conditions, only expressed on extravascular cells. However, circulating monocytes are capable of expressing TF on their surface when exposed to proinflammatory stimuli, including LPS, TNF, C-reactive protein, and C5a (19). The latter demonstrates one of the most important points of cross-talk between the complement and coagulation systems (12).

Activated factors of complement and coagulation are evident within aspirates of human atherosclerotic plaques and thrombi. More specifically, TCC as well as mRNA of complement C1r, C1s, and C4 are found within atherosclerotic lesions (20). C3 has been detected in thrombi (21), and C3a and C5a have been shown to be increased in surrounding plasma at the site of thrombus formation (22). This suggests local complement activation within the plaque and thrombi. Leukocyte-associated TF has also been found in thrombi and is thought to contribute to clot propagation and growth (23).

Because CC activate complement, and there is an increased recognition of complement–coagulation cross-talk, the aim of the current study was to investigate the role of CC in thrombosis by studying their activation of coagulation and the dependency of complement activation in this process.

Materials and Methods

Reagents

This work includes the following reagents: anticoagulant lepirudin (Refludan) from Celgene Europe (Boudry, Switzerland) and sterile PBS (PBS with or without Ca2+ and Mg2+) and EDTA from Sigma-Aldrich (St. Louis, MO). Human serum albumin (HSA) was from Octapharma (Jessheim, Norway). Heat-inactivated Escherichia coli strain LE392 (ATCC 33572) was from the American Type Culture Collection (Manassas, VA). Cryo Tubes (polypropylene) were from Nunc (Roskilde, Denmark). Complement inhibitors were as follows: compstatin analog CP40 (sequence: 14-aa cyclic peptide [D]Tyr-Ile-[Cys-Val-Trp(Me)Gln-Asp-Trp-Sar-Ala-His-Ary-Cys]-melle-NH2) and a control peptide (Sar-Trp(Sar-Asp(Me)-Ala-Ala-Asp-Ile-His-Val-Gln-Ary-melle-Trp-Ala-NH2), synthesized as described (24); C5 inhibitor ecluzolimab (Soliris) from Alexion Pharmaceuticals (New Haven, CT) and control rituximab (Roche, Basel, Switzerland); and cyclic peptide CSaR1 antagonist PMX53 (AcPhe-Om-Pro-Cha-Trp-Arg), synthesized as described (25). TF inhibitors used included the functional grade purified inhibitory Ab against human TF (HTF-1) from eBioscience (San Diego, CA) and Ultra-LEAF Purified Mouse IgG1 (MGI-41, 400165) from BioLegend (San Diego, CA). Corn trypsin inhibitor (CTI; CTI-01) from Haematologic Technologies (Essex Junction, VT) was used as a factor XII inhibitor. Flow cytometry Abs included anti-CD44 FITC (M499) from BD Biosciences (345784), anti-HLA-DR allophyocyanin–Cy7 (L243) from BioLegend (307602), anti–Siglec-8 allophyocyanin–C6 (7C9) from BioLegend (347106), anti-TF PE (CD142, NY2) from BioLegend (365204), and anti-mouse IgG1, κ PE (MOPC-21) BioLegend (981804). FACS Lysing Solution was from BD Biosciences. Abs for histology included murine monoclonal neoseptopeptide anti-human C9 (DIA011-01, clone E11, isotype IgG2a) from BioPorto Diagnostics (DK-2820, Gentofte, Denmark), neoseptopeptide anti-human C3b (A209, isotype IgG1) from Quidel (San Diego, CA), mAb against human TF (CD142, ADG4509, clone IID8, isotype IgG1) from Sekisui (American Diagnostica, Pungstdorf, Germany), a monoclonal anti-human CD141 (18D11, isotype IgG1) synthesized in-house, monoclonal mouse anti-human CD68 (clone KP1, M0814, isotype IgG1), and anti-human CD61 (clone 92/51, M10753, isotype IgG1) from Dako (Glostrup, Denmark). Isotype controls were purified mouse IgGs IgG1 (clone MOPC-21), IgG2a (clone MOPC-172), and IgG2b (clone MOPC-11) from BioLegend.

Preparation of CC and endotoxin measurement

CC were prepared as described by Smadst et al. (10). Briefly, ultrapure cholesterol (100 mg) was dissolved in 1-propanol (50 ml) and aggregated by the addition of distilled water. CC were air-dried and suspended in 0.05% (v/v) HSA/PBS. After preparation, the endotoxin contamination was measured in a Chromogenic LAL Assay (Lonza). The LPS was below the detection limit (0.1 endotoxin units/ml) in a sample size of 50 mg/ml CC, and thus, the endotoxin content would be below 0.18 pg/ml CC.

Experiments in human whole blood

Human whole blood from healthy volunteers, anticoagulated with lepirudin (50 μg/ml), was used as first described by Mollnes et al. (26). Final concentrations of CC or controls were as follows: CC (500, 1000, and 2000 μg/ml), E. coli (1 × 107 particles/ml), or glass control (200 μl of PBS with Ca2+/Mg2+ in BD Vacutainer glass tubes). Final concentrations of inhibitors or peptide controls was added to plasma 10 min before the CC or controls (20 μM). After plasma harvesting, the pellets were added to 350 μl of PBS (equal to plasma harvesting) and further processed as described in TF mRNA below.

Experiments in platelet-free plasma

Platelet-free plasma (PPP) anticoagulated with lepirudin (50 μg/ml) was made by centrifugation at 2000 × g for 15 min and stored in aliquots at −20°C for analyses of prothrombin fragment 1+2. After plasma harvesting, the pellets were added to 350 μl of PBS (equal to plasma harvesting) and further processed as described in TF mRNA below.

PFTF1.2 quantification

PFTF1.2 levels were quantified in duplicate using the Enzygnost Prothrombin F1+2 ELISA kit (Monoclonal OPBD03; Siemens Healthcare, Marburg, Germany) following the manufacturer’s instructions. OD was measured using a PolarStar OMEGA plate reader (BMG Labtech, Ortenberg, Germany).

TF mRNA

Cell pellets (550 μl) were added to 1.53 ml of PaxGene Blood RNA tubes solution (PreAnalytiX, Hombrechtikon, Switzerland) and stored at −80°C until RNA extraction. Total RNA was extracted using the MagNA Pure Pure 96. Cellular RNA large volume kit (Roche Diagnostics, Mannheim, Germany) according to the manufacturer’s instructions, and the concentration was measured using the NanoDrop 2000c spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). cDNA was synthesized using the High-Capacity cDNA Reverse Transcription Kit and 2720 Thermal Cycler (Applied Biosystems, Foster City, CA) and stored at −80°C. TF mRNA levels were measured using the StepOne Plus Real-Time PCR System (Applied Biosystems). Gene expression was detected by TaqMan Fast Universal PCR Master Mix reagents and predeveloped TaqMan gene expression assays using the target gene TF (Tf, Hs 0017225) with reference gene human β-2-microglobulin (Tf, Mm 00444296_m1). The LPS was below the detection limit (0.1 endotoxin units/ml) in the presence of CC (2000 μg/ml) or in glass vials with incubation lengths of 60 and 240 min.

TF-bearing microparticle quantification

TF-bearing microparticles (TF-MP) were isolated from blood plasma using the Nanopore Drop 2000e spectrophotometer (Thermo Fisher Scientific, Wilmington, DE). cDNA was synthesized using the High-Capacity cDNA Reverse Transcription Kit and 2720 Thermal Cycler (Applied Biosystems, Foster City, CA) and stored at −80°C. TF mRNA levels were measured using the StepOne Plus Real-Time PCR System (Applied Biosystems). Gene expression was measured by TaqMan Fast Universal PCR Master Mix reagents and predeveloped TaqMan gene expression assays using the target gene TF (Tf, Hs 0017225) with reference gene human β-2-microglobulin (Tf, Mm 00444296_m1). The LPS was below the detection limit (0.1 endotoxin units/ml) in the presence of CC (2000 μg/ml) or in glass vials with incubation lengths of 60 and 240 min.
thereafter centrifuged at 1500 × g 15 min at RT before the plasma fractions were further centrifuged at 13,000 × g for 2 min at RT. PPF was then harvested by retracting 400 μl from the uppermost plasma fraction. Samples were stored at −80°C until analysis.

**Flow cytometry**

EDTA-inactivated blood samples (50 μl) were added to a mixture of anti-CD14 FITC (2.5 μl), anti–Siglec-8 allophycocyanin (1.25 μl), anti–HLA-DR allophycocyanin–Cy7 (1.25 μl), and anti-TF PE (IgG1; 1.25 μl) or isotype control (0.625 μl, concentration-matched) for anti-TF (anti-mouse IgG1c PE), and RBCs were lysed by 1 ml of FACS Lysing Solution (diluted 1:10 in H2O). Samples were analyzed using a BD FACSCanto II flow cytometer with data processing in FlowJo software (version 10; Tree Star). To separate the monocytes from granulocytes and eosinophils emitting autofluorescence following CC exposure, a flow cytometric gating strategy based on the following settings was used: monocytes defined by side scatter (SSC)high/CD14low/Siglec-8−/MHCII+ population (based on settings from nonstimulated blood) and a positive selection of the MHCII+ cells (red color in fourth picture). The percentages of TF+ leukocytes were determined in a histogram plot set by the control Ab. The gating strategy is given in Fig. 1.

**Platelet aggregation assay**

Platelet aggregation was analyzed on a Multiplate electrical impedance aggregometry analyzer (Roche). Blood was collected in 4.5-ml plastic blood collection tubes containing the anticoagulant hirudin (25 μg/ml; Roche). After a 30-min rest at 20°C, the samples were diluted with 5% (v/v) PBS without Ca2+ and Mg2+ with 0.05% (w/v) HSA (Octapharma) or CC (500–2000 μg/ml) or PBS with Ca2+ and Mg2+ with 0.05% (w/v) HSA (Octapharma) or CC (500–2000 μg/ml). The samples were analyzed as recommended by the manufacturer, adding 0.3 ml of preheated (37°C) isotonic saline (0.9%) to 0.3 ml of whole blood in the Multiplate test cell, followed by incubation for 3 min at 37°C. The aggregation response over 6 min was expressed by the area under the curve as arbitrary units × min, either by the CC or its background control alone or following additional stimulation with 20 μl of ADP or thrombin receptor activating peptide-6 (TRAP) test as described by the producer (Roche).

**Human thrombus and plaque material**

Intracranial thrombus material was obtained by arterial recanalization or thrombectomy (27) from a single-center cohort study (28). Carotid plaques were obtained by carotid endarterectomy. Study inclusion criteria were acute stroke with clinical ischemic symptoms corresponding to an angiographically-proven large intracranial vessel occlusion, absence of intracranial hemorrhage on computed tomography scan, and a clearly defined time of symptom onset within 6 h of inclusion. Mechanical thrombectomy involved arterial catheterization from the groin artery to the level of the intracranial vessel occlusion and thrombus retraction with the use of a retrievable stent delivered through a microcatheter. After retraction, the thrombi were stored at 4°C in Allprotect tissue reagent (Qiagen) or snap-frozen.

**Histology**

Histology processing was performed at the Cellular and Molecular Imaging Core Facility Histology laboratory at Norwegian University of Science and Technology. We chose frozen tissue material without preservations because of the possibility of preserving neoepitopes of activated complement and preserving the CC structures. Briefly, frozen thrombus tissue sections (4 μm) were stained for hematoxylin erythrosine saffron (HES) or with specific Abs. The HES procedure, in brief, was as follows: rehydration through descending grades of ethanol to water (1 min), staining in aqueous hematoxylin (5 min) followed by bluing in water (8 min), erythrosine treatment (5 min) and rinsing in water (1 min), dehydration in ethanol (80, 96, and 100%), and staining in saffron (5 min) before rinsing in absolute ethanol and clearing in Tissue Clear. Cover-slipping used Sakura.
Results

CC induce PTF1.2 in human lepirudin whole blood

Lepirudin specifically inhibits thrombin and thus avoids fibrinogen activation and downstream coagulation while still allowing the coagulation system to be activated upstream. Despite the inhibition of the thrombin feedback loop, the upstream coagulation proteins together with all complement proteins are able to be activated, thereby allowing interaction with leukocytes. The conversion of prothrombin to thrombin results in a cleavage product, PTF1.2, that serves as an indirect estimate of the initial coagulation activation by either of the two coagulation pathways. CC 2000 µg/ml induced a significant increase (p < 0.001) of PTF1.2 compared with the PBS/HSA control following 240 min of incubation (Fig. 2A). Significant elevation was achieved for CC 1000 µg/ml (p < 0.05), whereas a nonsignificant (p > 0.05) elevation was obtained under exposure to CC 500 µg/ml (data not shown). Glass significantly elevated PTF1.2 levels by 60 min (p < 0.0001) and E. coli by 120 min (p < 0.001) (Fig. 2A). Glass is a highly potent activator of coagulation FXII and can be inhibited by CTI (29), whereas E. coli activates TF that can be explored by TF-inhibiting Abs (14). Inhibiting TF prior to addition of CC as a TF-blocking Ab efficiently and significantly (p < 0.0001) blocked the CC- and E. coli–induced PTF1.2 (Fig. 2B). These data pointed to the TF-driven pathway as most important for the CC-induced PTF1.2. To test for the FXII dependency, CC or glass were incubated in lepirudin PFP with or without blockage of FXII using CTI. In PFP, the overall PTF1.2 induction by CC or glass was low (just above baseline) but could still be significantly (p < 0.05) inhibited by CTI (Fig. 2C). The ability of CC to induce platelet
aggregation was investigated by electrical impedance aggregometry (Fig. 2D–F). CC did not induce platelet aggregation as compared with its background control (HSA/PBS) and positive control, TRAP (Fig. 2D). The CC did not induce additional activation or inhibition in activated platelets, as investigated by the ADP test (Fig. 2E) or the TRAP test (Fig. 2F). Altogether, these data point to the TF pathway as the main pathway triggered by CC, with a small contribution from FXII.

**CC induce coagulation through monocytic TF**

The procoagulant activity of TF detected in the inhibition studies (Fig. 2B) could be a consequence of cellular expressed TF or TF-MP. Measuring TF at the transcriptional level (TF mRNA) after CC 2000 µg/ml exposure in the whole blood resulted in a highly significant \( p < 0.0001 \) increase as compared with the PBS/HSA control, peaking after 60 min of incubation (Fig. 3A). Similar patterns, although less prominent, were found following exposure of CC 1000 µg/ml and CC 500 µg/ml (Supplemental Table I). In comparison, *E. coli* induced a significant \( p < 0.0001 \) and stronger response but also peaking after 60 min (Fig. 3A).

Measuring the TF-MP activity in human whole blood (Fig. 3B) resulted in the lack of activity following CC exposure, although the *E. coli* induction was consistent with previous data (14).

We further investigated the TF expression on the surface of the monocyte and granulocyte populations using flow cytometry. The monocytic population (CD14+/Siglec-8-2/MHCII+) histogram showed a clear shift to the right for TF following CC exposure (Fig. 3C) and stronger for *E. coli*. Quantification from six donors indicated an average of 23% of the monocytic population expressing TF at the surface upon CC 2000 µg/ml exposure and between 18 and 20% at the lower concentrations (Supplemental Table II). Of note, a significant increase was detected already after 60 min for CC 2000 µg/ml \( p < 0.0001 \) and CC 1000 µg/ml \( p < 0.0001 \) and after 120 and 240 min for all concentrations of CC (Supplemental Table II). *E. coli* induced the fastest and strongest TF expression, involving between 70 and 80% of the monocytic population, in agreement with previous findings. In contrast, no shift in TF was found on the surface of the granulocytes (SSChigh/CD14+/Siglec-8-) for either CC or *E. coli* stimulation (Fig. 3D) (14). Equally, no TF induction could be detected on the eosinophilic population (SSChigh/MHCII-2/Siglec-8+; data not shown). In conclusion, CC induces synthesis of TF mRNA, whereas only monocytes express TF at the surface.

**CC induce coagulation through an initial complement activation**

CC are strong activators of complement, leading to TCC formation (10). Complement-induced TF has previously been demonstrated (13–15), suggesting a possibility also for CC-induced coagulation. Inhibiting complement by blocking C3 (CP40) or C5 (eculizumab) significantly \( p < 0.0001 \) reduced the CC-induced PTF1.2 and monocyte TF expression close to baseline (Fig. 4A–D). Of note,
the percentage of monocytes expressing TF was <1% upon inhibition of C3 and <2% upon inhibition of C5. Upon E. coli exposure, the inhibition was less efficient, in accordance with previous data demonstrating the need for dual blockage of CD14 and complement C5 to avoid E. coli–induced coagulation (14). Because the inhibition of CC was efficient also at the level of C5, we further investigated the involvement of the anaphylatoxin C5a receptor, C5aR1, using the inhibitor PMX53. The effective blocking of the CC-induced PTF1.2 and monocyte TF by PMX53 points to an important role of the C5a–C5aR1 axis in the CC-induced coagulation (Fig. 4E, 4F). In conclusion, the CC-induced coagulation activation measured by PTF1.2 and monotypic TF is dependent on initial complement activation, in which the anaphylatoxin receptor C5aR1 is central.

CC are found within human intracranial vessel thrombus material and are associated with activated complement, monocytes, and TF+ leukocytes

Our studies in whole blood pointed to complement and monocytes as essential participants in CC-induced coagulation. To further explore the physiological relevance of these findings, we next explored thrombus material retracted from the intracranial vessel of patients with advanced carotid atherosclerosis. First, we investigated the possibility of detecting CC structures within the thrombus material and next in the association with complement and monocytes. Because CC are one of few materials showing birefringence in human tissue (30), it was possible to detect CC structures by polarization filter reflected light microscopy. Birefringent structures similar to CC were detected within hematoxylin-stained samples (Fig. 5A, 5B). The corresponding HES sections, the birefringent structures were absent (Fig. 5C, 5D). Crystalized cholesterol dissolves in ethanol (31), which is one of the constituents of HES. The disappearance of birefringent structures upon HES staining suggested that the structures could be of cholesterol origin.

Of note, the CC areas were richly populated with cells having bean-shaped nuclei typical for monocytes (Fig. 5). Immunohistochemistry staining confirmed the presence of monocytes within the CC-rich areas, as marked by yellow arrows (Fig. 6A, 6B). The corresponding adjacent section stained prominently for TF (Fig. 6C, middle picture). Furthermore, complement neoepitopes C5b–9 (Fig. 5D) were densely present and strongly associated with the birefringent CC structures (yellow arrows). The neoepitope iC3b was also found within the CC-rich areas (Fig. 5E), in addition to being present in the outermost regions of the thrombus (Fig. 5E, left picture, blue arrows). The presence of monocytes/macrophages in the thrombus was further confirmed by positive staining for CD68 in areas rich in CC (Supplemental Fig. 1A). Platelets (CD61), in contrast, were present throughout the thrombus (Supplemental Fig. 1B). The Ab specificities were manifested by the areas free of staining and further confirmed by the lack of staining by their respective isotype control, IgG1 (Supplemental Fig. 1C), IgG2a, and IgG2b (data not shown). In conclusion, the histological examinations pointed to a physiological

**FIGURE 4.** CC induce coagulation activation dependent on an initial complement activation. Whole blood was incubated with CC (500, 1000, or 2000 μg/ml), PBS/HSA, or E. coli (1 × 10^7 particles/ml) for 240 min with the C3 inhibitor CP40, the C5 inhibitor eculizumab, or their respective controls (scrambled peptide for CP40 or rituximab as control for eculizumab). (A) PTF1.2 production following C3 inhibition (CP40; n = 7 donors). (B) Monocyte TF expression at 120 min of incubation following C3 inhibition (n = 6 donors). (C) PTF1.2 production following complement C5 inhibition (eculizumab; n = 7 donors). (D) Monocyte TF expression at 120 min of incubation following C5 inhibition (n = 6 donors). (E) PTF1.2 following inhibition of complement receptor C5aR1 with PMX53 after 240 min (n = 5 donors). (F) Monocyte TF expression after 120 min of incubation following inhibition of C5aR1 with PMX53 (n = 5 donors). Data are given as means ± SEM. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.
relevance of CC and complement in thrombosis through the activation of monocytic TF.

Monocyte TF/CD14 expression patterns are connected to CC and activated complement in vulnerable plaque and in whole blood

Vulnerable plaques were examined for ruptures, the presence of CC, complement, and monocytes. One sample showed signs of an open lumen and had large CC present near the adventitia (Fig. 7A, Supplemental Fig. 2). Briefly, within the open lumen bright staining with the monocyte marker CD14 (CD14<sup>bright</sup>) suggested monocyte influx and blood exposure (Fig. 7A, yellow arrows). In the innermost region of the lumen, a layer of CD14<sup>low</sup>-staining monocytes was detected (Fig. 7A, first picture, blue arrows). The corresponding layer stained positive for TF (second picture, blue arrows). Of note, bright CC structures were observed in the

![Figure 5](image)

**FIGURE 5.** Histology from frozen sections of human intracranial thrombus retracted from a patient with advanced carotid atherosclerosis and explored by polarization filter reflected light microscopy. (A and B) Hematoxylin-stained sections with easily identifiable birefringent CC marked with orange arrows. (C and D) HES-stained sections of corresponding areas. Corresponding areas are (A) and (D), and (B) and (C). The thicknesses of the sections are 4 μm. Images were taken with original magnification ×40 objective by a polarization filter to visualize birefringent structures. Scale bar, 30 μm.

![Figure 6](image)

**FIGURE 6.** Immunohistochemistry of frozen sections of human intracranial thrombus retracted from a patient with advanced carotid atherosclerosis and explored with polarization filter reflected light microscopy. (A and B) Staining against the monocyte marker CD14. (C) Staining against TF. (D) Staining against the complement neoepitopes C5b–9. (E) Staining against the complement neoepitope iC3b. The thicknesses of the sections are 4 μm. Pictures in the left column are taken by original magnification ×4 objective, and the CC-rich area (blue frame) is further augmented with an original magnification ×40 objective (middle and right columns). Examples of birefringent structures are shown by yellow arrows, whereas blue arrows point to the outer thrombi sections stained heavily for neoepitope iC3b. Scale bars, 200 μm for pictures in left column and 30 μm for pictures in middle and right columns.
adjacent region (marked CC). The CC-rich areas stained heavily for complement iC3b and C5b–9 (Fig. 7A, third and fourth pictures). The monocytic origin of the TF+ cells was further confirmed by positive staining for the monocyte/macrophage marker CD68 and negative staining for CD34 and CD61 in corresponding areas (Supplemental Fig. 2A). Larger magnifications confirmed the monocytes’ origin by the presence of bean-shaped nuclei (Supplemental Fig. 2C, 2D). The presence of CC in the area adjacent to the CD14low/TF+ cells is clearly shown (Supplemental Fig. 2C–E). Endothelial staining was only found in the area of CD14bright monocytes, with irregular staining patterns typical for endothelial renewing (Supplemental Fig. 2E). The Ab specificities were confirmed by negative controls (data not shown).

The findings of CD14low/TF+ expressing leukocytes within the plaque could be explained by further observations in whole blood upon CC exposure (Fig. 7B–H). Briefly explained, in the monocytes, a reduction of CD14 from the initial expression level over time was observed, with a significant lowering for CC 2000 μg/ml after 120 and 240 min (p < 0.001–0.0001) and for CC 1000 μg/ml after 240 min (p < 0.01) (Fig. 7B). Moreover, a negative correlation (r = −0.697) between CD14 expression and the TF+ monocytes (percentage) was identified (Fig. 7C). The CC exposure led to a substantial and significant (p < 0.0001) increase in the granularity, indicating phagocytosis (10) (Fig. 7D). The reduction in CD14 (Fig. 7E, 7F) and increase in granularity (Fig. 7G, 7H) could be inhibited by blocking complement at the level of C3 and C5, thus linking both events to complement. From these findings, we would suggest that CC in a vulnerable plaque situation activates complement, with a subsequent influx of monocytes, phagocytosis of CC, and induction of TF in monocytes. During this process, CD14 is internalized, in part through complement-mediated activity.

Discussion

CC have emerged as important contributors to atherosclerotic inflammation (11), but their potential role in thrombosis is unexplored. In this study, we show that CC induce coagulation through a complement-dependent expression of TF in monocytes by using an experimental whole blood model. The physiological relevance

FIGURE 7. Human vulnerable plaque and whole blood show corresponding patterns in monocyte TF/CD14 expression and complement activation to CC. (A) Immunohistochemistry of the vulnerable plaque with the monocyte marker CD14, TF, and activated complement neoepitope iC3b and neoepitopes C5b–9. Staining areas for CD14bright monocytes are marked with yellow arrows, and CD14low/TF+ expressing monocytes are marked by blue arrows; specified also are the CC-rich areas, lumen, and adventitia. Sections are 4 μm and are taken by original magnification ×4 objective and given for two adjacent areas in each staining. Scale bar, 200 μm. Whole blood following CC exposure showing effects on (B) monocyte CD14 expression, (C) correlation between CD14 and TF in monocytes, (D) the increase in monocyte granularity, (E) monocyte CD14 following inhibition of complement C3 (120 min), (F) monocyte CD14 following inhibition of complement C5 (120 min), (G) monocyte granularity following inhibition of complement C3 (120 min), and (H) monocyte granularity following inhibition of complement C5 (120 min). Data are given as mean fluorescence intensity (MFI; mean ± SEM, n = 6 donors). *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001.
of these findings is supported by human intracranial vessel thrombus and vulnerable plaque histology demonstrating CC in association with monocyte TF (CD14/TF$^+$ leukocytes) and activated complement (iC3b and C5b–9). The present data thus point to an active role of CC and complement in thrombosis.

Atherosclerotic plaques can contain dense amounts of CC (32), suggesting relatively high concentrations of CC at the releasing site. In this study, we show that CC in concentrations of 500–2000 μg/ml induce coagulation by cleavage product PTF1.2, mediated mainly through the TF pathway and TF surface expression by monocytes. Our data indicate a time span of CC-induced TF expression of 1–2 h, with TF mRNA peaking after 1 h and full surface expression after 2 h. Although endothelial damage could induce a faster response in concert with platelets, these findings point to releasable CC as active contributors in thrombus formation and growth. The demonstration of monocyte influx and TF-expressing monocytes in a vulnerable human plaque further support monocytes as important effectors in thrombosis. The time frame from plaque debris release to clinical symptoms is not well defined and depends on thrombus size and localization. According to clinical guidelines (33), patients with advanced carotid atherosclerosis should undergo carotid endarterectomy preferably within 48 h after a transient ischemic attack because of the high risk of recurrent infarction. The clinical picture with small ischemic occlusions such as transient ischemic attack promotes a high-risk period for up to 2 mo, in which patients are particularly vulnerable to new and often larger attacks. The release of CC during this period could thus promote thrombosis by inducing TF expression on monocytes.

Cholesterol embolization syndrome is a phenomenon characterized by a multitude of small emboli occurring over time in association with CC, with an incidence of 22–27% during surgical treatment of atherosclerosis (34). Of note, this phenomenon was connected to complement consumption, which fits well with the complement-dependent CC-induced coagulation activation demonstrated in this study. Our data might contribute to a mechanistic explanation by demonstrating that the coagulation activation is mediated predominantly by complement, with more than 90% inhibition at either the level of C3 or C5. The efficient prevention of coagulation by inhibiting the anaphylatoxin receptor C5aR1 points to an essential role of the C5a–C5aR1 signaling axis in CC-induced coagulation. This clear effect was in contrast to the partial inhibition of coagulation by C5aR1 under exposure to E. coli and clearly points to a specific complement-mediated activation by CC. Bacteria-induced coagulation is, in addition to complement, also dependent on CD14 (26) and, furthermore, connected to the TLR pathway (35). The lack of direct activation of CD14/TLRs by CC might explain the lower potency of CC to induce TF mRNA, the lower number of TF$^+$ monocytes, and the lack of TF-MP activity presently demonstrated. We observed, however, a rapid loss of CD14 in correlation with TF induction following CC exposure. This phenomenon was concentration-dependent and linked to complement and could possibly be the consequence of phagocytosis of CC. Whether CD14 is involved in the signal induction leading to TF expression remains to be elucidated. Previous exploration by our group has shown that the CC-induced complement activation mediates upregulation of CR3 (CD11b/CD18) through C5a/C5aR1 (10). The additional opsonization with complement fragments at the level of C3 leads to efficient phagocytosis of CC. This was also demonstrated in the current study by the concentration-dependent granulation in monocytes after CC exposure, which could be blocked by the complement inhibitors both at the level of C3 and C5. Although we did not explore the link between monocyte TF and the phagocytic events, the rapid phagocytosis prior to TF mRNA synthesis, as well as the complement dependency for both, could indicate that these events are connected.

The inflammatory cytokines TNF and IL-1β, among others, are triggered by CC exposure (10). The induction of TF has also been connected to TNF (36), which could suggest its involvement in CC-mediated coagulation. However, the time span between CC exposure and TF mRNA induction of 1 h points to a shorter induction time than is needed for activation of TNF. It is, thus, more likely that these processes are initiated in parallel in connection to complement than that TNF plays a central role in the initial TF induction by CC. The NLRP3 inflammasome has been linked to CC-induced IL-1β secretion mediated by C5a (10). Further on, NLRP3 has been linked to induction of TF-MP. However, our investigation in whole blood using the specific NLRP3 inhibitor MCC950, caspase-1 inhibitor VX-765, and pan-caspase inhibitor Z-VAD-FMK did not provide evidence for its involvement in CC-induced TF expression or downstream coagulation activation (data not shown). Additionally, no induction of TF-MP by CC was observed in our system. As regulation of TF has been shown to occur mainly on the transcriptional level (37), our findings imply that CC-induced generation of C5a activates the C5a–C5aR1 axis to drive synthesis of TF mRNA, which has been shown to occur both in monocytes (38) and in endothelial cells (39).

From the present and previous data, CC can be regarded as participators in thromboinflammation with a strong dependency on complement. Our investigation points to the monocytes as the only leukocytes able to induce TF upon exposure to CC, as surface expression on granulocytes and eosinophils was absent. CR3 expression is, in contrast, induced both on the monocytes and granulocytes (10), and our explorations also revealed increased granulocyte granularity upon CC exposure (data not shown). This further points to the ability of CC to activate the granulocytes but not induce surface expression of TF. Although there is agreement regarding the capability of monocytes to express TF, the expression of TF on other cell types is debated (40–42). These discrepancies in cellular sources could be related to the health status of the donors as well as the type of stimuli (36, 42, 43). Platelets contribute in the initial clot-formation process by binding to damaged vascular endothelium (44), and connections between platelets and monocyte TF are commonly accepted (45). We found no direct effect of CC on platelet aggregation or in the induction of PTF1.2 following CC exposure in platelet-containing plasma (data not shown). The tiny induction of PTF1.2 following CC exposure in PFP, which was still significantly inhibited by CTI, suggests a small contribution of FXI in CC-induced coagulation. This could also explain the ~10% of PTF1.2 activity remaining following complement inhibition. A limitation of the lepirudin anticoagulant is the inhibition of thrombin, the major platelet activator (46). However, because the platelets also contain complement receptors (47), a complement-dependent coagulation activation by the platelets could be possible. The presence of iC3b in the outermost parts of the thrombus also opens a role for complement–platelet interactions in thrombus growth, as previously described (22, 48).

CC are normally observed as imprints (CC clefts) in plaques because of their dissolution when specimens are treated with ethanol during regular immunohistochemical staining (31). CC accumulation over time gives rise to cleft structures within atherosclerotic plaques that are easily observed under the microscope. However, upon plaque rupture, CC will be subjected to blood flow and, therefore, more challenging to detect during thrombus formation. CC can be observed using polarization filter reflected light microscopy because of their high birefringence.
The detection of complement products is also more efficient in frozen samples, as used in this study. A recent paper suggests that freezing biopsy specimens from atherosclerotic plaques may produce birefringent structures in frozen section samples that do not represent CC (49). An exclusion of such artifacts from the present images is not possible, but still, several findings support the birefringent structures being of CC origin: 1) the birefringent structures were associated with activated complement, monocytes, and TF; 2) monocytes or TF² cells were adjacent to CC structures; 3) the shape of the birefringent structures resembled the square CC made in vitro for the smaller structures, whereas the larger needle-shaped or larger assemblies were similar to those seen in atherosclerotic plaques; and 4) upon HES staining of stepwise ethanol treatment, the birefringent structures disappeared, thereby supporting that the structures contained cholesterol. Altogether, these findings point to a CC nature of the birefringent structures.

Based on the efficiency of complement inhibition in reducing CC-induced coagulation, targeting complement warrants further exploration as a future treatment option for prevention of thrombosis. Currently, the complement C3 inhibitor CP40 is under clinical development (50), whereas the C5 inhibitor eculizumab is being used in clinical use for treatment of paroxysmal nocturnal hemoglobinuria (51). These patients often suffer from thromboembolism, and targeted treatment of C5 has proven to reduce the thrombotic events by 81% (51). In a preclinical porcine model of myocardial infarction, C5 treatment also reduced the infarction size (52). Note, complement activation is an upstream event to the inflammatory cytokines, including IL-1β (10), and thus, targeting complement could represent an alternative strategy. This might also avoid a compromised immune activation upon infections, as is the concern with anti-IL-1β therapy (53).

In conclusion, the current study reveals a role for CC in thromboinflammation. CC possess a procoagulant feature dependent on complement and monocytic TF, and following plaque rupture, releasable CC might contribute to thrombosis. The use of complement inhibitors targeting either C3, C5, or C5aR1 could be an alternative clinical strategy for prophylactic treatment of atherosclerotic plaques or after thrombosis and warrants further exploration.

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Disclosures

J.D.L. is the inventor of patents and/or patent applications that describe the use of complement inhibitors for therapeutic purposes, the founder of Amyndas Pharmaceuticals, which is developing complement inhibitors (i.e., third-generation compstatins) for clinical applications, and the inventor of the compstatin technology licensed to Apellis Pharmaceuticals (i.e., 4|MeW7]2/POT-4/APL-1 and PEGylated derivatives). The other authors have no financial conflicts of interest.

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