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Induction of Complement C3a Receptor Responses by Kallikrein-Related Peptidase 14

Katerina Oikonomopoulou,* Robert A. DeAngelis,* Hui Chen,* Eleftherios P. Diamandis,†,* Morley D. Hollenberg,§,* Daniel Ricklin,*+† and John D. Lambris*+†

Activation of the complement system is primarily initiated by pathogen- and damage-associated molecular patterns on cellular surfaces. However, there is increasing evidence for direct activation of individual complement components by extrinsic proteinases as part of an intricate crosstalk between physiological effector systems. We hypothesized that kallikrein-related peptidases (KLKs), previously known to regulate inflammation via proteinase-activated receptors, can also play a substantial role in innate immune responses via complement. Indeed, KLKs exemplified by KLK14 were efficiently able to cleave C3, the point of convergence of the complement cascade, indicating a potential modulation of C3-mediated functions. By using in vitro fragmentation assays, mass spectrometric analysis, and cell signaling measurements, we pinpointed the generation of the C3a fragment of C3 as a product with potential biological activity released by the proteolytic action of KLK14. Using mice with various complement deficiencies, we demonstrated that the intraplantar administration of KLK14 results in C3-associated paw edema. The edema response was dependent on the presence of the receptor for C3a but was not associated with the receptor for the downstream complement effector C5a. Our findings point to C3 as one of the potential substrates of KLKs during inflammation. Given the wide distribution of the KLKs in tissues and biological fluids where complement components may also be expressed, we suggest that via C3 processing, tissue-localized KLKs can play an extrinsic complement-related role during activation of the innate immune response. The Journal of Immunology, 2013, 191: 3858–3866.

The complement network, consisting of >50 plasma-borne and membrane-bound proteins, is an essential component of the innate immune system, which forms a first line of defense against microbial invaders and exerts important functions in immune surveillance and homeostasis (1). The constituents of the network are organized into three major pathways (classical, lectin, and alternative) characterized by distinct “molecular recognition motifs” that trigger activation of the cascade upon detection of damage- or pathogen-associated surface patterns (1, 2).

All three pathways converge on the proteolytic release of active fragments of C3, the C3a anaphylatoxin and C3b, the latter of which is an important opsonin and vital component of the C3 convertase. This convertase can in turn amplify the generation of opsonic C3b, as well as contribute to cleavage of C5 and generation of the C5a anaphylatoxin and C5b. The main biological activities of complement activation are as follows: 1) the opsonization of pathogens; 2) the anaphylatoxin-derived recruitment of inflammatory cells; 3) microbial cell lysis as a result of the formation of the membrane-attack complex downstream of C5; and 4) the regulation of adaptive immunity by stimulation of B and T cells (1, 2). C3a and C5a, and their desargminated derivatives, have received particular attention as major signaling effectors that exert a broad spectrum of biological effects ranging from chemotaxis to innate cell activation, thereby contributing to innate and adaptive immune functions and inflammation. They act primarily via their G protein–coupled receptors C3aR and C5aR (CD88), respectively; a third anaphylatoxin receptor, C5L2, has been described but its role and specificity are still a matter of debate (3).

Apart from the traditional view of activation within the three branches of the complement cascade, a potential extrinsic pathway of initiation of complement activation, mainly at the level of C3 and/or C5, by noncomplement proteinases also has been suggested (1, 2). For example, generation of potentially biologically active fragments from C3 and C5 by the prototype serine proteinase trypsin has been reported previously (4, 5). Direct activation and/or degradation of C3 or C5 have additionally been shown for other proteinases, including plasma kallikrein (6, 7), components of the coagulation cascade such as factor Xa and thrombin (8), the neutrophil-released proteinases elastase and cathepsin-G (9), and mast cell tryptase (10). A role for microbial proteinases released during infection in the regulation of anaphylatoxin activity also has been suggested (11).
Despite the presence of circulating enzymes described above that may be able to modulate complement activity in the periphery, little is known about local complement activation in inflamed tissues and tumors, where proteases in addition to thrombin and tissue-trypsin would be activated. Kallikrein-related peptidases (KLKs) are a family of tissue serine proteinases, distinct from plasma kallikrein (12, 13). KLKs have trypsin or chymotrypsin-like activity and belong biochemically to the same enzyme class as trypsin. The human KLK family consists of 15 enzymes, which are upregulated in tissues, mainly of epithelial origin, and in biological fluids, such as ascites fluid and sera from cancer patients. Cascade enzymatic activity of KLKs has been hypothesized to occur in the CNS and seminal plasma. In the skin, several KLKs have been isolated from the outermost layers of the stratum corneum and overall epidermis and are implicated in the physiology and pathology of the skin (14). Among the KLK enzymes, KLKs 5, 6, and 14, are characterized by wide tissue distribution, high biological activity, and significant association with the clinical progression and outcome of several types of cancer, such as breast, ovarian, prostate, and skin carcinoma (12). Specifically KLK14, possibly as part of a KLK catalytic cascade, has been linked to epidermal desquamation (i.e., shedding) because of a degradation of the intercellular (corneo)desmosomal adhesion molecules that link adjacent corneocytes (14). Preliminary analyses have also connected KLK14 expression in inflamed skin tissues to the pathogenesis of rosacea, whereas other KLKs have been associated with atop dermatitis, psoriasis vulgaris, and Netherton syndrome (13, 14). Despite data establishing a role for KLKs in skin homeostasis (i.e., desquamation and matrix remodeling) (12–14), knowledge of their specific contributions to skin inflammatory pathologies is still largely unknown.

It has been reported that C3aR, responsive to C3a signals, can be found on cells that can be localized in skin tissue (i.e., mast cells, neutrophils, and monocytes/macrophages) (15), where KLKs would also be present (14). Keratinocytes, a major source of KLKs (14), are key components of the innate immune response (16), and abundantly secrete C3 (17, 18) and express C3aR (19) in the settings of inflammation. Similar to the KLKs (13, 14), an association between complement activation and skin inflammation (e.g., dermatitis and psoriasis) has also been made (20). In addition to inflammation, both the complement system and KLKs have independently been attributed roles in the regulation of oncogenesis (12, 21). Specifically for skin carcinogenesis, proteolytic activation of C3 may contribute to tumorigenesis and metastasis of human melanoma cells in nude mice (22).

Given the similar tissue distribution and role of the complement and KLK families, we hypothesized that KLKs can have a major contribution to inflammatory responses taking place in peripheral tissues, including the skin, by modulating the availability and activity of complement components. Our analyses of the C3-targeting function of KLKs, exemplified in this study by the activity of KLK14, show that this interaction can lead to biologically significant responses in ex vivo and in vivo models of peripheral tissue complement activation.

Materials and Methods
Reagents and protein preparations
Chemicals were purchased from Fisher Scientific (Hampton, NH), unless indicated otherwise. Human complement proteins C3a, C3a-desArg, and C5 were from Complement Technology (Tyler, TX). C3 was purified from human plasma as previously described (23) with addition of an extra gel filtration step (Superdex 200, flow rate of 1 ml/min in PBS; GE Healthcare). Fractions were analyzed by SDS-PAGE under reducing conditions, total protein assay, and ELISA. More specifically, regarding ELISA, our purified preparations of unknown C3 content were diluted 2.5-fold with PBS (pH 7.5) and incubated in 96-well plates for 2 h at room temperature. Plates were subsequently blocked with 1% BSA in PBS and allowed to sit for 30 min at room temperature. Wells were washed three times with PBS-Tween 20 (0.05%) before addition of an HRP-conjugated goat anti-human C3 polyclonal detection Ab (50 μl, 1/1000 in PBS; catalog number 0855237; MP Biomedicals) for 30 min at room temperature. The plate was washed three times with PBS-Tween 20 (0.05%) and developing solution (50 μl/well of a 100 mM sodium citrate buffer supplemented with 0.5 mg/ml ABTS [catalog number 10102946001; Roche Applied Science, Indianapolis, IN] and 0.03% H2O2 [catalog number 7722-84-1; Fisher]) was added to each well and incubated for 2 min at room temperature. Absorbance of light was measured at a wavelength of 405 nm.

KLK6 was expressed in a mammalian human embryonic kidney cell system (24), and KLK3 was expressed in Pichia pastoris (25); enzymes were purified and activated as described previously (24, 25). Active KLK5 was purified and activated as described previously (25). In brief, concentrated yeast supernatant, diluted 1:2 with running buffer (10 mM MES [pH 5.3]), was loaded onto a 5-ml HiTrap SP HP-Sepharose cation exchange column (Amersham Biosciences). The column was eluted with a linear gradient of 0–1 M KCl in running buffer. Fractions were analyzed by a KLK14-specific ELISA, pooled, and concentrated 10 times in 10 mM MES (pH 5.3). The concentration of the proteins was quantitatively and analytically validated by using total and total protein assay. The purity and integrity of the protein (>90%) was verified by SDS-PAGE.

Estimation of trypsin-like serine proteinase activity
The enzymatic activity of the trypsin-like kallikrein-related serine peptidases was measured using the fluorogenic t-butoxycarbonyl-tripeptide-7-amino-4-methylcoumarin (AMC) synthetic substrates glucoseamine-alanine-arginine (QAR)-AMC (KLK14) or valine-proline-arginine (VPR)-AMC (KLK5 and 6) (Bachem Biosciences, King of Prussia, PA). In brief, each KLK was incubated at 37°C in a microtiter plate, with the optimal activity assay buffer and varying concentrations (within the linear range) of fluorescent substrates in a final volume of 100 μl. The initial rate of AMC release was measured on a Wallac Victor fluorometer (PerkinElmer, Waltham, MA) at 355 nm for excitation and 406 nm for emission. The fluorescence values of enzyme-free reactions were subtracted from each vial. All experiments were performed in triplicates. The activity was quantified by monitoring the rates of tripeptide–AMC hydrolysis (slopes of the fluorescence curves expressed as fluorescence units per minute). A standard curve with known concentrations of AMC was used to calculate the rate of product formation. Kinetic analysis was performed by nonlinear regression analysis using the Enzyme Kinetics Module 1.1 (Sigma Plot; SPSS, Chicago, IL), and activity of each enzyme was estimated by means of nanomolar amounts of AMC-substrate cleaved per minute of reaction per milligram of enzyme used. The activity of a given concentration of KLK was compared with that of a standard concentration of pure trypsin (5 nM or 2.5 U/ml; catalog number T-7418; Sigma-Aldrich), measured using the same AMC substrate used for assessing each KLK activity and subsequently expressed by means of “trypsin-like units of activity.” This approach allowed for the direct comparison of activity levels between the different proteinases.

Fragmentation analysis
In vitro fragmentation experiments were performed by incubating native C3 or C5 in PBS (pH 7.5) in the absence or presence of KLKs or trypsin at 37°C in a concentration- and time-dependent manner. The enzyme to substrate ratio was expressed in w/w units; in the case of KLK14 the 1% w/w enzyme to C3 ratio corresponds to a 1:16 molar ratio. Given the comparable molecular masses of KLKs and trypsin, the molar ratios of enzyme to substrate was expressed in w/w units; in the case of KLK14 the 1% w/w enzyme to C3 ratio corresponds to a 1:16 molar ratio. In brief, purified C3 (20 μg) was incubated with KLK14 or trypsin at indicated concentrations and times at 37°C in PBS (pH 7.5). Samples were passed through C3a ZipTip (Millipore, Billerica, MA) and analyzed by MALDI-TOF. Spectra were acquired on a MALDI-Micro MX mass spectrometer (Waters, Milford, MA) in linear mode as described recently (8).

C3 (7.4 μg) or C5a (0.5 μg) were also subjected to KLK14 incubation (equal molar ratio at the level of 1:30 of enzyme to C3/C5a, corresponding to a w/w ratio of 0.5% for C3 and 10% for C3a) directly at 37°C or at room temperature in PBS (pH 7.5), followed by liquid chromatography–mass spectrometry (LC-MS) analysis. Analytes were separated on a 1.7 μm
ultraperformance liquid chromatography (UPLC) BEH130 C18 column (2.1 mm × 150 mm; catalog number 18603556; Waters). The analytical column temperature was maintained at 40 ºC. Peptides were separated for 8 min using a flow rate of 0.15 ml/min and a linear gradient of 10–60% of solvent B (0.1% formic acid in acetonitrile [ACN], v/v) in combination with solvent A (0.1% formic acid in water, v/v). LC-MS analysis was performed on a SYNAPT G2-S instrument (Waters) equipped with an electrospray ionization source controlled by MassLynx 4.1 software (Waters). The capillary voltage was 3.2 kV, the cone voltage was 30 V, and the source temperature was 120 ºC. [Glu1]-fibrinopeptide B (Sigma-Aldrich) was used for lock-mass correction with a sampling rate of 30 s. Mass spectra were acquired in positive mode over an m/z range of 100–2000 Da at a scan rate of 1 s. Fragment sequences were verified by collision-induced dissociation fragmentation.

The C3a degradation by KLK14 was also followed by studying the proteolytic release of downstream fragments by HPLC (Ettan LC; GE Healthcare). In particular, 4 µg C3a was incubated at room temperature with 0.4 µg KLK14 (10% enzyme to C3a ratio in PBS; 1:30 molar ratio) and the reaction was followed for up to 60 min. Reactions were terminated by using 0.02% TFA, and samples were run on an analytical column (Vydac and the reaction was followed for up to 60 min. Reactions were terminated by using 0.02% TFA, and samples were run on an analytical column (Vydac MS C3 300 A, 5 µm × 50 mm, ID 1 mm; catalog number 218MS5105; Grace, Deerfield, IL) using a gradient of 15–65% solvent B (0.1% trifluoroacetic acid in ACN, v/v) for 60 min and a flow rate of 100 µl/min (0.1% TFA in water was used as solvent A). Peak fractions (absorbance at 215 nm) were collected and identified of the KLK14-mediated C3a fragments was performed by MALDI-TOF and/or electrospray ionization analysis as detailed above.

Detection of C3a by ELISA

Estimation of C3a released by the KLK14-mediated C3 cleavage relative to the one released by trypsin was achieved by incubating KLK14 with purified human C3 (1% w/w for the indicated time points at 37 ºC, in PBS [pH 7.5]) and by detecting C3a using a rabbit polyclonal Ab (Up1896), which was raised against recombinant human C3a and produced in-house according to standard procedures, and a HRP-conjugated goat anti-rabbit IgG (catalog number 172-1019; Bio-Rad, Hercules, CA) in a direct ELISA approach.

C3aR-mediated cell activation assays

Rat basophilic leukemia (RBL) cells stably expressing the C3aR were obtained as described previously (26); cell dynamics in response to C3a were monitored with the SRU BIND platform (SRU Biosystems, Woburn, MA), which uses photonics crystal technology to sense activation-induced changes in cell morphology and adhesion (27, 28). In brief, the wells of a 96-well BIND C2A biosensor plate (C2A is a matrix surface that promotes cell attachment; SRU Biosystems) were hydrated with 50 µl deionized water for 30 min at room temperature. Water was then substituted with 25 µl of complete medium (DMEM supplemented with 10% heat-inactivated FCS) for 30 min at room temperature. The baseline signal was recorded using the BIND PROFILER instrument (SRU Biosystems). RBL-C3aR cells (25,000 cells/well) were seeded on the plate in DMEM (Invitrogen, Grand Island, NY), supplemented with 10% heat-inactivated FCS, and maintained overnight at 37 ºC (5% CO2). The plate was allowed to equilibrate at room temperature for 30 min and the cell attachment signal was obtained. Cells were stimulated with C3, C3a, or the KLK14-mediated C3 fragments. Incubation of KLK14 with C3 (0.5% w/w; 1:30 molar ratio) occurred in the absence of serum to avoid the KLK inhibitory effects of serum-contained serpins (28). Cells were subsequently treated with serial dilutions of the original incubation mixture in the FCS-supplemented medium and the changes in cell dynamics (recorded as peak wavelength values [PWV] in picometers [pm]) were followed for up to 30 min post-ligand incubation. Recorded values were analyzed using the EMS software (SRU Biosystems) and the EC50 of cell responses were calculated.

Paw edema inflammation model

Complement-deficient mice lacking C3 (C3(-/-)), the C3aR (C3aR(-/-), or the C5aR1 (C5aR1(-/-)) have been described previously (29–31). Deficient mice had been backcrossed for at least nine generations onto a C57BL/6J background, and their wild-type littermates or C57BL/6J mice (The Jackson Laboratory, Bar Harbor, ME) were used as controls where indicated. Complement-deficient and control animals (6–8 wk old; 20–25 g) were obtained and housed at the University of Pennsylvania animal facility under carefully regulated conditions with a 12-h light-dark cycle and provided with food and water. All experimental procedures were approved by the Institutional Animal Care and Use Committee of the University of Pennsylvania and were performed in accordance with guidelines established by the Animal Welfare Act. Prior to the intraplantar administration of KLK14, a basal measurement of the paw thickness of each mouse was recorded using an electronic micrometer caliper (catalog number 500-196-20; Mitutoyo, Aurora, IL). The compounds used for treatment were diluted in sterile Dulpbecko’s PBS and administered via intraplantar injection into the hind paw at a final volume of 10 µl/mouse (1.2 µg or 22 trypsin-like units of KLK14 per paw). Control mice were treated with KLK14 inactivated with soybean trypsin inhibitor (STI; Sigma-Aldrich; incubation for 30 min at 37 ºC using 1.5 molar kV the enzyme to inhibitor) (32) or buffer alone (n = 4–5 mice/group; three independent experiments). STI was used to indicate that the edema response is mediated because of the proteolytic action of KLK14. As an index of edema formation, paw thickness monitored with the micrometer caliper (Mitutoyo) was followed for up to 4 h postinjection. Paw thickness values at baseline were subtracted from the values obtained at each time point. All values obtained from the paw thickness measurements were analyzed by two-way ANOVA (Prism; GraphPad Software, La Jolla, CA) and the χ2 test. p ≤ 0.05 was considered to indicate a significant difference between groups.

Results

KLKs cleave C3

KLK5, KLK6, and KLK14 caused a limited proteolysis of C3 (Figs. 1, 2, Supplemental Fig. 1). The cleavage of C3 resulted in the formation of a molecular mass species similar to that of C3b that is composed of the fragmented α-chain (α’-chain) and intact β-chain fragments. The proteolytic processing pattern for KLKs (Fig. 1A–C) displayed similarities with that of the enzymatically-related serine proteinase trypsin (Fig. 1D) (4, 33) that resulted in the disappearance of the C3 α-chain and appearance of an α’-chain fragment. Coomassie staining of the C3 digestion products detected in reducing gels that separate the α- and β-chains indicated that all three KLK enzymes could cleave C3, but with different efficiencies. For instance, at comparable ratios of enzyme to substrate, both KLK5 and KLK14 cleaved the α-chain (Fig. 1A, 1C, Supplemental Fig. 1), whereas much higher enzyme to substrate ratios were required for KLK6-mediated cleavage (Fig. 1B, Supplemental Fig. 1). In contrast to trypsin, which at higher ratios of enzyme to substrate caused a disappearance of the α’-chain to yield a constituent migrating just above the β-chain (Fig. 1D, far-right lane), KLK14, even at relatively high enzyme to substrate ratios, was not able to efficiently cleave the α’-chain further (Fig. 1A).

We elected to focus on KLK14 as a prototype member of the KLK family with expression and/or activity documented in variable disease settings. To standardize the activity of KLK14, its proteolytic efficiency was compared with the one exhibited by a standard concentration of pure trypsin (5 nM or 2.5 U/ml), measured using the same QAR-AMC substrate used for assessing the KLK activity (25). By performing fluorogenic substrate activity analysis for all KLKs and trypsin, we found that our KLK14 preparation had a trypsin-like activity of ~18 U/µg KLK (25) when compared with the preparation of trypsin we used for our analysis, 1% of KLK14 (0.05 µg; w/w of C3) corresponded to 0.9 U trypsin-like activity against the same QAR-AMC substrate. By performing similar calculations, and by using the VPR-AMC substrate, the levels of trypsin-like activity exhibited by 0.05 µg KLKs 5 and 6 were found to be lower than KLK14 (decreased by ~5- and 22-fold, respectively, when compared with KLK14; data not shown).

A comparative time course analysis of the C3 cleavage at 1% enzyme concentration revealed a faint band of a low molecular mass species corresponding to C3a that rapidly appeared after 1 min and remained visible throughout the 2-h incubation (Fig. 2A, white arrowhead). In agreement, the α-chain was rapidly converted to the α’ fragment; the visible weakening of the α’ band after 1 h with appearance of additional faint bands may indicate a slow yet progressive degradation of the α’-chain. Cleavage of the β-chain appears possible but would likely occur at a slow rate.
mass C3 fragments are indicated with gray arrowheads.

As both C3 and C5 are degraded by trypsin (4, 5, 33), the similar levels of C3-fragmentation activity between KLK14 and trypsin raised the possibility for KLK14-mediated cleavage of C5. However, our data indicated that the two enzymes displayed significantly different efficiency for cleaving the downstream component C5 (Fig. 2C, 2D). More specifically, C5 cleavage by KLK14 was limited compared with the extensive cleavage caused by trypsin. Furthermore, the fragments that were generated by the proteolytic action of KLK14 under the same conditions as used for C3 were not indicative of the release of fragments with molecular weights expected for activated C5 (i.e., C5b of ∼170 kDa) (Fig. 2C). This is in contrast to what has been reported for C5 fragmentation by several proteinases of the coagulation system, which lead to the generation of active C5 fragments (8). Although these studies on the KLK-mediated cleavage of C5 may be extended in the future, our data for the cleavage specificity of KLK14 revealed a cleavage pattern distinct from trypsin and indicated that C3 was a preferred substrate when compared with C5 (Fig. 2E). Consequently, our focus for subsequent work remained on C3.

C3 cleavage by KLK14 results in generation of C3a-like species

The KLK14-generated release of a C3b-like molecular mass species (Figs. 1A, 2A) suggested the possible generation of a corresponding C3a-like anaphylatoxin fragment released from the α-chain of C3 (Fig. 2A, white arrowhead, 2E). Detection of a C3a-like species within 1 min of C3 incubation in the KLK14-containing reaction mixture using a direct ELISA approach further supported this hypothesis (Supplemental Fig. 2).

The release of the intact C3a fragment from C3 by KLK14 was verified by chromatographic separation of the proteolysis products followed by MALDI-TOF and/or LC-MS (Fig. 3). In particular, our proteome analyses identified a fragment with a molecular mass of 9087 Da and charge properties equivalent to those expected for human C3a (Fig. 3) (8). Furthermore, time-resolved analysis revealed that, after release from C3, intact C3a was processed further by KLK14 during incubation into a lower molecular mass species, which was resistant to subsequent proteolysis during the 30- to 60-min follow-up incubation time (Fig. 3A). Calculation of the peak areas for C3a and its fragmented peptide confirmed the fragmentation pattern and indicated a slower rate for the second cleavage (Fig. 3B). Notably, KLK14 was also able to cleave purified human C3a into the same truncation species (Fig. 3C); the time-dependent cleavage of C3a by KLK14 was also followed by HPLC (data not shown). The fragment secondarily produced by the proteolytic action of KLK14 had a molecular mass of 8282 Da. Using LC-MS analysis, it was shown that this shift in m.w. corresponds to the removal of the “ASHLGLAR” peptide, with a molecular mass of 824 Da, from the C terminus of C3a (Fig. 3C, 3D, Supplemental Fig. 3A). The peptide sequence was confirmed by collision-induced dissociation fragmentation analysis (Supplemental Fig. 3B). The KLK14-mediated proteolytic fragmentation of C3a may also explain the immunologic detection pattern in the C3a ELISA (Supplemental Fig. 2), which showed a decrease of the initial detection signal exhibited by the KLK14-incubated C3 fragments at and beyond 30 min of reaction; even though a polyclonal anti-C3a Ab has been used, the possibility that the dominant epitope involves the C-terminal region of C3a cannot be excluded. Sustained time-dependent detection of C3a released by KLK14 was enhanced further at lower ratios of enzyme to substrate, down to 0.01% of C3 (w/w; comparative MALDI-TOF analyses shown for KLK14 in Supplemental Fig. 3A), whereas in contrast, fragmentation by trypsin similar to that of KLK14 could only be observed when the enzyme to substrate ratio used was lower than 0.01% (w/w; data not shown). Notably, in our calculations, the comparable molecular masses of KLK14 and trypsin resulted in similar molar concentrations of either trypsin or KLK14 used at the equivalent enzyme to substrate w/w levels.

FIGURE 1. Proteolysis of complement component C3 by KLKs and trypsin. The concentration-dependent KLK14-mediated generation of C3b-like fragments from degradation of C3 is followed (A) in comparison with the proteolytic processing of C3 by KLKs 5 and 6 and trypsin (B–D) at the indicated enzyme to C3 ratios (w/w). Samples were run on 7.5% SDS-PAGE gels under reducing and denaturing conditions, and visualized with Coomassie staining. The C3 α- and β-chains are indicated with arrows, whereas the α'-chain is indicated with black arrowheads. Lower molecular mass C3 fragments are indicated with gray arrowheads.

as well (Fig. 2A). As observed above, digestion of C3 by trypsin is more complex and leads to the rapid generation of additional fragments at early time-points (Figs. 1D, 2B). Although detailed conclusions regarding the comparative enzyme kinetics of the proteolytic enzymes for cleaving C3 cannot be extrapolated from our qualitative degradation analysis, our data clearly show that KLK14 exhibits a proteolytic ability comparable to trypsin in performing the “initial” cleavage of C3 to a C3b-like fragment (Fig. 1); nevertheless, KLK14 does not subsequently cleave C3b further as extensively as trypsin.

KLK14 shows limited cleavage activity toward C5

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The release of the intact C3a fragment from C3 by KLK14 was verified by chromatographic separation of the proteolysis products followed by MALDI-TOF and/or LC-MS (Fig. 3). In particular, our proteome analyses identified a fragment with a molecular mass of 9087 Da and charge properties equivalent to those expected for human C3a (Fig. 3) (8). Furthermore, time-resolved analysis revealed that, after release from C3, intact C3a was processed further by KLK14 during incubation into a lower molecular mass species, which was resistant to subsequent proteolysis during the 30- to 60-min follow-up incubation time (Fig. 3A). Calculation of the peak areas for C3a and its fragmented peptide confirmed the fragmentation pattern and indicated a slower rate for the second cleavage (Fig. 3B). Notably, KLK14 was also able to cleave purified human C3a into the same truncation species (Fig. 3C); the time-dependent cleavage of C3a by KLK14 was also followed by HPLC (data not shown). The fragment secondarily produced by the proteolytic action of KLK14 had a molecular mass of 8282 Da. Using LC-MS analysis, it was shown that this shift in m.w. corresponds to the removal of the “ASHLGLAR” peptide, with a molecular mass of 824 Da, from the C terminus of C3a (Fig. 3C, 3D, Supplemental Fig. 3A). The peptide sequence was confirmed by collision-induced dissociation fragmentation analysis (Supplemental Fig. 3B). The KLK14-mediated proteolytic fragmentation of C3a may also explain the immunologic detection pattern in the C3a ELISA (Supplemental Fig. 2), which showed a decrease of the initial detection signal exhibited by the KLK14-incubated C3 fragments at and beyond 30 min of reaction; even though a polyclonal anti-C3a Ab has been used, the possibility that the dominant epitope involves the C-terminal region of C3a cannot be excluded. Sustained time-dependent detection of C3a released by KLK14 was enhanced further at lower ratios of enzyme to substrate, down to 0.01% of C3 (w/w; comparative MALDI-TOF analyses shown for KLK14 in Supplemental Fig. 3A), whereas in contrast, fragmentation by trypsin similar to that of KLK14 could only be observed when the enzyme to substrate ratio used was lower than 0.01% (w/w; data not shown). Notably, in our calculations, the comparable molecular masses of KLK14 and trypsin resulted in similar molar concentrations of either trypsin or KLK14 used at the equivalent enzyme to substrate w/w levels.
KLK14-mediated C3 fragmentation triggers cell activation

To investigate the biological function of the C3a-like fragments released from C3 by KLK14, we measured their signaling response in RBL-C3aR cells using the SRU BIND photonic crystal sensor platform, which has been previously used to characterize anaphylatoxin signaling (26). Treatment of the RBL-C3aR cells with serial dilutions of C3 previously incubated with KLK14 for two time points, which were selected based on the MS-derived time course (Fig. 3), resulted in detectable cell activation with an estimated EC50 in the low nanomolar range (~13 nM; Fig. 4A, 4B). In contrast, pure C3 at 100 nM, which was used as a control, did not induce a significant response (Fig. 4, grey hexagons). Importantly, the C3aR-mediated activity for the KLK14:C3 mixtures was in a similar range as for purified C3a (EC50 = 15.8 nM; Fig. 4C); C3a specifically acted on RBL-C3aR and did not stimulate RBL cells transfected with the C5a receptor (Supplemental Fig. 4A, 4B). Of note, the absolute concentration of biologically active C3a generated by KLK14 in our samples cannot be reliably determined and likely contains a mixture of intact and truncated C3a after 5 and 30 min of incubation with the enzyme (Fig. 3); given the importance of the arginated C terminus for C3aR signaling (3, 26), it is not expected that the truncated C3a fragments participate in active signaling. Indeed, incubation of C3a with KLK14 for 5 min led to visible decrease of the activation signal on RBL-C3aR cells when compared with time 0 (Supplemental Fig. 4C). Similarly, in the case of our transfected cell system, a role for other C3 species released during the proteolytic degradation of the remaining α-, α′-, and/or β-chains is improbable. Because serum-derived proteinase inhibitors are known to restrict KLK14 enzyme activity (28), and the KLK14-mediated cell stimulation ability of C3 was abrogated in the presence of serum (Supplemental Fig. 4D), no further significant degradation of C3 fragments by KLK14 during this cell assay is expected after addition to serum-supplemented media. Similarly, the use of heat-inactivated serum renders significant desargination of C3a by carboxypeptidase N within the time frame of our experiment rather unlikely. The reason and importance of the observed lower maximal signal exhibited by the KLK14-mediated C3 fragments (Fig. 4A, 4B) as opposed to the C3a control (Fig. 4C) is not yet determined; besides experimental variability in cell seeding, a potential interference of components in the more complex matrix of the KLK14:C3 mixture cannot be excluded. Experiments with purified or recombinantly produced KLK14-induced C3 fragments may be performed in the future to better explain the C3aR-mediated signaling pattern more quantitatively. Still, the current semiquantitative data on RBL-C3aR cells strongly indicate a biological activity for the C3 fragments that results from the proteolytic action of KLK14.

Administration of KLK14 results in complement-dependent paw edema in vivo

On the basis of our biochemical findings that KLK14 is a potential activator of human C3 (Figs. 1-3) that can induce C3aR-dependent cell signaling (Fig. 4), we hypothesized that the KLK14-released C3a-like fragments may exert proinflammatory roles. We therefore used a well-established mouse model of paw edema (34) in combination with different complement-deficient animals. As a first step, we confirmed that KLK14 was able to induce a similar cleavage pattern in vitro using mouse C3 as seen in human C3 incubations with KLK14 (above). Indeed, incubation of mouse C3 with KLK14 resulted in the cleavage of the α-chain with appearance of a C3a-sized band; this conversion was prevented in the presence of the STI-inhibited KLK14 (Fig. 5A). The intraplantar administration of KLK14 in paws of wild-type C57BL/6 mice or littermate controls of complement-deficient animals was able to induce a statistically significant edema response (Fig. 5B) comparable to that previously observed for trypsin (34) and formerly reported for KLK14 (35, 36). The KLK14-generated response was attenuated by preincubation with STI (Fig. 5B). Furthermore, this KLK14-mediated edema response was sustained at 120 min postinjection in contrast to inactivated KLK14 (Fig. 5B) and vehicle...
alone (KLK14-WT bars; Fig. 5B, 5C). Similarly, intraplantar C3a injections in wild-type C57BL/6J mice resulted in a paw edema response at 120 min postinjection equivalent to that induced by KLK14 (Fig. 5C, square-dotted bars). As expected, the tissue edema response to C3a-dsArg was significantly lower (Fig. 5C, light gray bar) when compared with C3a. The significance of the residual C3a-dsArg-mediated response is not yet clear; whereas C3a-dsArg does not bind to and activate C3aR (3, 26), activities of this fragment potentially independent of C3aR have been reported in similar settings (37, 38). In contrast, administration of KLK14 in C3-deficient animals resulted in a significantly decreased edema response 120 min postinjection and equivalent to edema caused by PBS alone (Fig. 5C, black bars). In agreement with these data, intraplantar KLK14 did not cause an edema response in the C3aR-deficient animals (Fig. 5C, gray bars), indicating that in this model the KLK14-mediated edema response is essentially C3a/C3aR dependent. Notably, the swelling responses to vehicle alone were comparable in all animals. The same was true for the inactivated KLK14 responses in wild-type (receptor-expressing) animals (data not shown). Furthermore, because genetic deletion of the major skin proteinase receptor, proteinase-activated receptor (PAR)2, cannot eliminate the murine paw edema response observed after intraplantar KLK14 administration (K. Oikonomopoulou, unpublished observations) (36), the paw swelling observed in this model was not attributed to the previously reported KLK14-PAR2 signaling (25, 35, 39). Of importance, in the C5aR-deficient animals the paw swelling in response to KLK14 was equivalent to that in the wild-type littermate mice (Fig. 5C, white bars). This result renders a role for C5a in generating the KLK14-mediated inflammatory response very unlikely and further supports a C3b/C5-independent KLK14 edema pathway. Notably, these in vivo data are also in agreement with our in vitro findings of limited human C5 degradation by KLK14 that does not generate C5-derived effector fragments (Fig. 2C).

Discussion

The main finding of our work is that KLK14, a prominent member of the KLK family, can generate the C3a anaphylatoxin from C3 in vitro and, presumably due to a comparable cleavage, can trigger an inflammatory edema response via activation of the C3aR in vivo. Work in the past decade has revealed the tissue expression patterns of members of the KLK family. Generally, KLKs share a wide range of expression in different tissues and settings (12, 13) including the blood and skin, where complement components are also present (1, 15, 17). Our analyses presented in this study put a spotlight on C3 as a major target of KLKs. Even though we initially investigated the C3-processing activity of three members of the KLK family, we focused our subsequent studies on KLK14 as a prototype family member with high enzymatic activity and specificity for substrates such as C3 that contain an Arg or Lys at their P1 position (40).

FIGURE 3. Mass spectrometric identification of the C3a-like fragments released by KLK14. (A) The release by KLK14 of two distinct molecular mass species, C3a and a lower molecular mass C3a component, was investigated by LC-MS within the 60-min timeframe of incubation with the enzyme. (B) A graphical representation of the LC-MS spectrum area of the C3 fragments proteolytically released by KLK14 (A) is shown. Truncated C3a is indicated as tr.C3a. (C) Fragmentation of C3a by KLK14 under similar conditions is also shown for comparison purposes. The molar ratio of enzyme to C3 was similar to that of enzyme to C3a, corresponding to a w/w enzyme to C3a ratio of 0.5% for C3 and 10% for C3a. (D) The N-terminal part of the C3 α-chain is indicated, with the cleavage resulting in the release of active C3a fragment represented with a black arrow and the KLK14-mediated subsequent cleavage shown with a gray arrow. (A) and (C) show representative data from two independent experiments with similar results.
in adjuvant-induced arthritis (42), and to investigate the contribution of complements and C5aR to inflammatory reactions triggered after surgical incision (43). In agreement with our previous findings (35, 36), which singled out PARs as potential targets of the KLK14 degradative activity that can potentially result in cell signaling, tissue inflammation, and edema (25, 35, 39), our current work confirmed that KLK14 can trigger swelling when injected intraplantarly in mice. In addition, the observations in this study critically extend this model by suggesting a role for KLK action via proteolytic mechanisms in addition to the ability to signal via PARs. The data reported in this paper thereby implicate the targeting of the complement system by KLKs as an important regulator of inflammatory responses in peripheral tissues exemplified by our intraplantar model.

Specifically for the skin tissue, although several studies have investigated a role for the KLK cascade in skin homeostasis (i.e., desquamation and matrix remodeling) (13, 14), their involvement in skin pathologies is not yet clear. Upregulation of KLKs in several skin diseases has been reported (13, 14); for example, a role for PAR signaling by KLK14 in inflamed rosacea tissues (44) has been suggested. However, in a murine model of the Netherton syndrome, a pathological condition where activity of KLKs is highly increased, genetic deletion of the proteinase...
receptor was not able to abrogate cutaneous inflammation (45). Thus, our results offer a potential alternative mechanism of KLK-induced inflammation through complement activation that may contribute to skin pathologies.

Both the complement system and the PARs represent important innate immune defense mechanisms triggered by proteolytic activation of effector molecules (1, 2, 46). Complement activation in tissues in connection with local inflammatory processes is still poorly described and the absolute levels of its components in peripheral tissues, such as the skin, are largely unknown. The inflammatory actions of complement, primarily a result of anaphylatoxin signaling, can, among other functions, induce cytokine release by immune and nonimmune cells and regulate vascular permeability and cell adhesion (1, 47), functions that can be of importance in the paw edema inflammation model used in this work and by other groups (42, 43). Our study now shows that KLKs, and more specifically KLK14, can modulate C3a-mediated signaling in this established model of peripheral tissue inflammation.

More specifically, our data presented herein indicate that the C3aR can be a potential mediator of KLK-mediated localized inflammation in peripheral tissues including the skin. On the basis of the impaired reaction in C3aR−/− mice, the edema response is likely driven by the KLK-catalyzed generation of active complement C3a that stimulate C3aR. This is in agreement with previous work that suggested a role of C3aR in edema models of peripheral tissue inflammation (42). Although a role for secondary KLK14-mediated C3 fragmentation products in proinflammatory reactions cannot be fully excluded, a participation of the secondary C3a truncation fragment that was identified in our in vitro assays seems unlikely. In contrast to C5a, where the carboxypeptidase-processed product C5a-desArg still exerts signaling functions via C5aR, the corresponding truncation of C3a by even one C-terminal amino acid (i.e., to C3a-desArg) leads to complete loss of signaling activity (3, 26). In the current study, we did not find evidence to support activation of C5aR signaling by KLK14 in the same setting, a finding which corresponds well with the observation that KLK14 exerts very limited activity toward C5. Although the physiologically relevant C3 and KLK14 concentrations in peripheral tissues are not known, the C3 levels used by our in vitro assays were close to typically observed plasma concentrations of C3 (~1–1.5 mg/ml) (48). Notably, as dictated by our in vitro analyses, the levels of KLK14 needed for activation of 1.5 mg/ml C3 (~8 μM) can be as low as 0.15 μg/ml (~5 nM), an enzyme expression level that can be observed in tissue extracts and biological fluids (49).

Our new data indicating that KLK14 is able to stimulate C3αR-mediated cell/tissue responses adds C3αR, in addition to PARs (25, 39), to the group of G protein-coupled receptors involved in KLK-mediated inflammatory processes. Therefore, future work should focus on elucidating the mechanisms of the inflammatory action of KLK14 in peripheral tissues by activating the two receptor systems simultaneously (i.e., both PARs and C3αR). It will be of much interest to determine the role this process may play in the innate immune response in tissues like the skin. Moreover, it would be important to further investigate whether any other C3/C3α cleavage products generated by KLK14 may contribute to biological effects, including C3αR-independent signaling or antimicrobial functions. A similar hypothesis has been recently put forward for the processing of C5 by thrombin, leading to a unique C5b fragment with enhanced cell lytic activity (50).

In view of the recent literature that draws links between the KLK and thrombostasis proteolytic cascades (51), our new work suggests dual targets for these serine proteinases: 1) the PARs, which are activated by both coagulation enzymes and the KLKs (46), and 2) the complement system, which is affected by several coagulation factors (8, 52) as well as by KLKs, as we now demonstrate. Thus, these three interconnected proteolytic cascades (thrombostasis, complement, and that of the KLKs) may well be considered as modulators and amplifiers of the innate immune response in the inflammatory microenvironment.

In conclusion, we provide the first evidence, to our knowledge, that KLKs, apart from signaling via PARs and having degradative roles important for extracellular matrix remodeling and cell shedding, can exert inflammatory signaling in peripheral tissues including the skin by targeting complement component C3 and triggering release of C3a that, in turn, activates C3αR. This role of KLKs in modulating the availability of C3a anaphylatoxin adds to the recent literature that proposes KLKs as inflammatory modulators of cathelicidin activity and PAR signaling (13). In the future, it may therefore be interesting to extend these investigations to include other tissue-localized KLKs, PARs, and complement components in our models of peripheral tissue inflammation to arrive at a more complete picture of this intriguing crosstalk. Our data may also be of relevance for developing new therapeutic strategies for the treatment of inflammatory or cancerous skin conditions, where both KLKs and complement have been implicated.

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Disclosures
The authors have no financial conflicts of interest.

References
Supplementary Figures

Figure S1: Rapid degradation of complement component C3 by KLK14. A-C) The time-dependent KLK14-mediated generation of C3b-like fragments by the degradation of C3 was studied in comparison to the proteolytic processing of C3 by KLKs 5 and 6 at the indicated enzyme to C3 ratios (1% w/w). The C3 α- and β-chains are indicated with arrows, while the α’-chain is indicated with black arrowheads. Samples were run on 7.5% SDS-PAGE gels under reducing and denaturing conditions and visualized with Coomassie staining.
Figure S2. Detection of the KLK14-mediated C3a release from degradation of C3. The time-course analysis of the C3a generation by the proteolytic action of KLK14 on human purified C3 is shown in comparison to that of the prototype serine proteinase trypsin. Quantification was performed by a direct ELISA approach utilizing an anti-C3a rabbit polyclonal antibody (UP1896). KLK14-mediated C3a activity is represented by black bars, while trypsin-released C3a is shown with white bars (mean±SEM). Analysis of the samples was performed in triplicate.
Figure S3: Identification of the C3a-like fragments released from C3 by KLK14. A) The time- and dose-dependent release after limited proteolysis of C3 by KLK14 was studied with MALDI-TOF. C3a generation is shown within the time frame of 5 min incubation of C3 with the enzyme. The w/w ratio of enzyme to C3 is indicated in each panel. B) The collision-induced dissociation (CID) fragmentation analysis that confirms the identity of the truncated C3a fragment generated by the sustained proteolytic activity of KLK14 on C3 is shown (see also Fig. 3). The y-axis represents the relative MS spectrum intensity.
Figure S4: Monitoring of the KLK14-mediated RBL-C3aR cell responses. A-B) The absence of C3a or C3a-desArg responses [peak wavelength value (PWV) shift in pm] by RBL-C5aR cells otherwise responsive to C5a is shown. C-D) Cell responses triggered by C3aR signals were recorded in response to the KLK14-mediated C3 and C3a fragments. C3 or C3a agonists were used at a concentration of 20 nM, alone (control) or treated with KLK14. KLK14 was used at a 0.5% w/w enzyme to C3 ratio or at a 10% enzyme to C3a ratio. Incubations were performed in DMEM media (C) or DMEM supplemented with heat-inactivated fetal calf serum (D), as indicated in the two panels, prior to cell treatment. Addition of agonist is indicated with an arrow in panels A, C and D. Note that the definition of incubation time “zero” in the case of medium not supplemented with FCS is not possible due to the potential inability to stop/slow down the reaction (generation of active fragments) from the time the enzyme/substrate mixture is put on the plate until the readings are performed by the SRU BIND\textsuperscript{®} platform.