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Ornithodoros moubata Complement Inhibitor Is an Equally Effective C5 Inhibitor in Pigs and Humans

Andreas Barratt-Due,* Ebbe Billmann Thorgersen,* Julie Katrine Lindstad,* Anne Pharo,* Olga Lissina,†‡ John D. Lambris,§ Miles A. Nunn,† and Tom Eirik Mollnes*

Experimental evidence suggests that C inhibition and more particularly combined inhibition of C and the TLR coreceptor CD14 may be of therapeutic benefit in sepsis and other inflammatory conditions. A barrier to the testing and further development of many inhibitors is that their activity is species specific. Pig is a relevant species for experimental models of human disease, and this study undertakes a comprehensive comparison of the inhibitory efficacy of the C5 inhibitor *Ornithodoros moubata* C inhibitor (OmCI) in human and porcine whole blood ex vivo models of *Escherichia coli*-induced sepsis. The effect of OmCI on complement activity in pigs undergoing *E. coli* sepsis was also examined. Porcine and human serum, and whole blood anticoagulated with lepirudin, was incubated with *E. coli* and the effect of OmCI investigated. The ex vivo results were virtually identical in pig and human. OmCI completely ablated the activity of all three C pathways at 0.64 μM. *E. coli*-induced C activation and expression of CD11b (wCD11R3 in the pig), was abolished ex vivo at 0.32 μM OmCI. Combining anti-CD14 and OmCI reduced the formation of IL-8 and TNF-α more potently than the single inhibitors. OmCI also efficiently bound *E. coli*-induced leukotriene B₄ in pig and human plasma. In support of our ex vivo findings, in vivo the activity of all C pathways was inhibited at 0.6 mg OmCI/kg pig. In conclusion, OmCI efficiently inhibited pig and human C activation, has accompanying anti-inflammatory effects and is a promising candidate inhibitor for further in vivo studies of sepsis. The Journal of Immunology, 2011, 187: 4913–4919.

C and the TLR family play essential parts in innate immune reactions. Upon pattern recognition, these danger sensors identify and eliminate pathogens as well as endogenous danger motifs, thereby protecting the host and maintaining homeostasis (1, 2).

The physiological effects of C are numerous and diverse (3). Activation is known to occur via three routes, the classical, the lectin, and the alternative pathways, all convergeing and leading to the cleavage of the central C factor C3. The C3 fragments C3β and iC3b are important opsonins in the defense against bacteria. Activation of the terminal pathway with the cleavage of C5 is biologically highly potent. Release of the anaphylatoxin C5a is known to induce upregulation of adhesion molecules, stimulate cytokine production, cause paralysis of neutrophils, increase vascular permeability and might lead to disseminated intravascular coagulation among other effects (4). C5b induces assembly of the terminal C5b–9 C complex (TCC), which can lyse certain pathogens and cells when incorporated into their lipid membranes (5).

The TLRs are an important class of pattern recognition receptors (6). TLR4 is a pattern recognition receptor for LPS of Gram-negative bacteria and is dependent on two important coreceptors, MD-2 and CD14 (7, 8). TLR signaling induces NF-κB, required for transcription of a wide variety of inflammatory and immune response genes (6).

Uncontrolled systemic activation of these upstream danger sensors may lead to a counterproductive response that endangers self as seen in septic shock (9, 10). Ex vivo studies with human whole blood have demonstrated that *Escherichia coli*-induced inflammation is strongly attenuated by a combined inhibition of C and CD14 (11, 12). Consequently we hypothesized that combined inhibition of C and CD14 might prove a useful therapeutic regimen in sepsis and other inflammatory conditions (13). Notably, inhibition of the classical proinflammatory cytokines TNF-α and IL-1β was shown not to have any impact on the *E. coli*-induced inflammatory reaction (14). We recently showed that inhibition of CD14 attenuated proinflammatory cytokines, granulocyte activation, and hypercoagulation in *E. coli*-induced sepsis in vivo in pigs (15). Furthermore, selective inhibition of C improved organ function in a baboon model of sepsis (16). Combined inhibition of C and CD14 in vivo has yet to be tested.

Ticks produce a plethora of proteins that interfere with host immune and hemostatic defense responses (17). OmCI is a 16.8-kDa saliva protein from the soft tick *Ornithodoros moubata* (18). This small protein binds directly to C5 and prevents cleavage by both the classical and the alternative C5 convertases, and thus, C5a and TCC is not formed (19). C5 inhibition by OmCI thus has vast anti-inflammatory potential, without affecting beneficial immunoprotective and immunoregulatory functions of upstream C activity (20). OmCI has cross-species C-inhibitory activity that has been demonstrated but not explicitly quantified in human, rat, mouse, and guinea pig (18, 21). The protein is protective in experimental passive and active autoimmune myasthenia gravis models in rats. OmCI-treated animals exhibited significantly fewer symptoms, and the inflammatory response was substantially attenuated (21, 22). Like some of its closest known homologs (23), OmCI has bifunctional properties, capturing the inflammatory...
mediator leukotriene B_4 (LTB_4) in an internal binding pocket in addition to binding to C5 (M.A. Nunn, unpublished observation). LTB_4 is a potent chemotactic agent and activator of white cells in particular neutrophils (24). Despite its low molecular mass, the half-life of OmCI is ~30 h in rats due to stable binding to C5, making OmCI an interesting agent for use in vivo (21).

The aim of the current study was to compare the efficacy of OmCI in pigs and humans, using functional assays designed for these species. By using a unique whole blood ex vivo model for investigation of the cross-talk of the inflammatory network, C inhibitory efficacy as well as anti-inflammatory properties of OmCI was elucidated. In addition, the C inhibitory effect of OmCI was explored in pilot experiments using an in vivo porcine model of E. coli-induced sepsis.

Materials and Methods

Reagents and equipment

Sterile PBS (pH 7.2) was purchased from Sigma-Aldrich (St. Louis, MO), Cryo Tubes (polypropylene) from Nunc (Roskilde, Denmark), Lepirudin (Refludan) from Pharmion (Hamburg, Germany), and 96-well plates (Costar 3590) from Corning (New York, NY). Zymosan A (Z-4250) was purchased from Sigma-Aldrich. Human γ globulin (160 mg/ml; Beriglobin) was purchased from Behring (Marburg, Germany) and converted to heat-aggregated IgG (HAIGG) by heating a 10 mg/ml stock to 63˚C for 15 min. Albumin (200 mg/ml) was purchased from Octapharma (Hundal, Norway).

Bacteria

E. coli strain LE392 (ATCC 35372) from the American Type Culture Collection (Manassas, VA) was used in all experiments. E. coli was grown over night in LB medium (tryptone 1%, yeast extract 0.5% and sodium chloride 0.9%), then centrifuged and washed in PBS. After resuspension in PBS, E. coli was immediately heat-inactivated (1 h, 60˚C) for use in the ex vivo experiments. Growth plates confirmed that all bacteria were killed. Thereafter, the bacteria were washed nine times to remove extrabacterial LPS and counted by flow cytometry. Live E. coli strain LE392 was used in vivo. Sepsis was induced with an increasing i.v. infusion of E. coli, as described previously (15). Each pig received a total of 1.075 × 10^9 E. coli/kg during the time course of the experiment, corresponding to 1.1 × 10^6 bacteria/ml blood.

Inhibitors

Recombinant bacterial OmCI (18), a 16.8-kDa protein, was produced in the laboratory of one of the authors (M.A.N.). Before use it was stored in single use aliquots at −70˚C. The mouse anti-human CD14 (clone 18D11) F(ab′)_2 and control F(ab′)_2 (clone BH1), was purchased from Diatec (Oslo, Norway). Mouse anti-porcine CD14 mAb clone MIL-2, isotype IgG2b, was purchased from AbD Serotec (Oxford, U.K.). A mouse anti-human IgG2b (clone BH1, product no. 2070) was purchased from Diace monoclonals (Oslo, Norway). Mouse anti-porcine CD14 mAb clone MIL-2, isotype IgG2b, was purchased from AbD Serotec (Oxford, U.K.). A mouse anti-human IgG2b (clone BH1, product no. 2070) was purchased from Diace monoclonals and used as an isotype-matched control. The compstatin analog Ac-I[CV(1MeW)QDWGAHRC]T, which binds to and inhibits cleavage of C3, and the cyclic hexapeptide Ac[FOPdChaWR], a C5a receptor antagonist (C5aRa), were produced as described previously (25, 26). A total of 10 mg/ml eculizumab (Soliris), a humanized monoclonal IgG2/4k-Ab that binds and inhibits cleavage of C5, was from Alexion Pharmaceuticals (Cheshire, CT), provided by Dr. Y. Fløysand (Oslo University Rikshospitalet).

Ex vivo whole-blood experiments

The whole blood model has previously been described in detail (27). Briefly, whole blood was collected from 15- to 60-kg pigs (Sus scrofa domesticus; Landrace: outbred stock) and healthy human donors. Blood was drawn into tubes containing the anticoagulant lepirudin (Refludan) at a final concentration of 50 μg/ml whole blood. The blood was immediately preincubated with inhibitors, controls, or PBS for 5 min at 37˚C. PBS or E. coli, at a final concentration of 10^9, 10^8, or 10^7 E. coli/ml whole blood, was then added and incubated for 2 h at 37˚C. After incubation, the tubes were put on ice and EDTA (final concentration, 20 mM) was added to stop the C activation, and the tubes were centrifuged for 15 min at 1400 × g at 4˚C. The plasma was stored at −70˚C until analyzed.

In vivo experiments

Norwegian Landrace pigs (n = 7) of either sex, with a mean weight of 14.4 kg (range, 14.0–16.0 kg) were used. Premedication, anesthesia, surgery, hemodynamic monitoring, induction of sepsis, and euthanasia were performed as described previously (15). Pigs received OmCI i.v. as a bolus (22.5, 15, 7.5, and 3.75 mg) and thereafter as a continuous i.v. infusion (0.5 mg/h) for 4 h. The total amounts of OmCI given were 24.5 mg (n = 1), 17 mg (n = 1), 9.5 mg (n = 2), and 5.75 mg (n = 1). Two pigs served as positive controls, receiving saline only (9 mg/ml). Blood samples were drawn and physiological data were registered after surgery at baseline and thereafter at 0 (T0), 30, 60, 120, 180, and 240 min.

Engine immunoassays

The commercially available enzyme immunoassay (EIA) (Wienisa; Wielis, Lund, Sweden) was used to test functional activity of the classical, lectin, and alternative C pathways. The test was designed for human C activity (28) but was later shown to cross-react with pig (29). Human C5a was analyzed by a commercial EIA (BD Biosciences, San Jose, CA). TCC was measured in an EIA, previously described in detail (30, 31). Briefly, the mAb aE11, specific for a C9 neoepitope in TCC, was used as capture Ab and a biotinylated monoclonal anti-C6 (Quidel, San Diego, CA) was used as detection Ab. Both Abs cross-react with pig epitopes; thus, the assay can be used to detect porcine as well as human TCC. The porcine cytokines TNF-α, IL-1β, and IL-8 were analyzed by Quantikine Porcine Immunoassay kits from R&D Systems (Minneapolis, MN). Human plasma samples were analyzed using multiplex technology (Luminex 100; Bio-Rad Laboratories, Hercules, CA) — an immunoassay based on colored beads, each with a unique cytokine detection Ab that permits simultaneous measurement of a wide range of human biomarkers. A 27-plex kit purchased from Bio-Rad Laboratories (Hercules, CA) was used containing the following cytokines: IL-1α, IL-1β, IL-2, IL-4, IL-5, IL-6, IL-7, IL-8, IL-9, IL-10, IL-12, IL-13, IL-15, IL-17, TNF-α, IFN-γ, eotaxin, fibroblast growth factor-basic, G-CSF, GM-CSF, IFN-γ-inducible protein-10, MCP-1, MIP-1α, MIP-1β, platelet-derived growth factor-BB, RANTES, and vascular endothelial growth factor. The analyses were performed according to the instructions from the manufacturer. Results are presented for the cytokines that exhibited responses in the inhibitors in comparison with the negative controls. LTB_4 from pig and human plasma was measured using a competitive enzyme immunoassay from R&D Systems.
Flow cytometry

For analyses of CD11b (human) or wCD11R3 (pig ortholog), whole blood was preincubated with inhibitor, control, or PBS for 7 min at 37°C. PBS or E. coli at a final concentration of 10⁸ E. coli/ml whole blood was added and incubated for 10 min before the cells were fixed with 0.5% (v/v) paraformaldehyde for 4 min at 37°C. Human fixed samples were stained with PE-conjugated anti-human CD11b (catalog no. 333142) or an isotype-matched PE-conjugated IgG1 control Ab (catalog no. 349053) (both BD Biosciences Pharmingen, San Jose, CA). Pig-fixed samples and EDTA-anticoagulated blood from the in vivo experiments were stained with FITC-conjugated anti-porcine wCD11R3 (catalog no. MCA2309F) or an isotype-matched FITC-conjugated IgG1 control Ab (catalog no. MCA920F) (both AbD Serotec) and incubated for 15 min at room temperature in the dark. The RBCs were then lysed for 7 min thereafter centrifuged and washed before being resuspended in PBS containing 1% albumin. Samples were analyzed by FACS (FACScan; BD Biosciences, Franklin Lakes, NJ). Granulocytes were gated in a forward light scatter/side scatter (of light) dot plot, and CD11b and wCD11R3 expression was given as median fluorescence intensity.

Statistics

GraphPad Prism version 5 (GraphPad Software, San Diego, CA) was used for the statistical analysis. The data were analyzed by one-way ANOVA, followed by Dunnett’s multiple comparison test. A two-tailed p value < 0.05 was considered statistically significant.

Ethics

Informed written consent was obtained from each blood donor, and the study was approved by the local ethical committee. The animals used were treated in adherence to the Norwegian laboratory animal regulations, and the study was approved by the Norwegian Animal Research Authority.

Results

In vitro

The effect of OmCI on the different complement pathways. In serum, OmCI dose-dependently inhibited all three C activation pathways similarly, and the inhibition was complete at a dose of 0.64 μM OmCI in both pig and human (Fig. 1). Concentrations between 1.28 and 20 μM were also completely inhibitory (data not shown).

The effect of OmCI on the formation of C5a and fluid-phase TCC. Pig and human whole blood was incubated with 10⁸ E. coli/ml for 30 min. OmCI dose-dependently inhibited human C5a formation (Fig. 2, left panel). At concentrations of OmCI ≥ 0.32 μM, the formation of human C5a was completely ablated.

OmCI also dose-dependently inhibited TCC formation in both pig and human whole blood (Fig. 2, middle panels). At concentrations of OmCI ≥ 0.32 μM, the formation of TCC was completely inhibited in both species. When serum was activated with either HAIGG or zymosan, which are potent activators of the classical and lectin/alternative C path way respectively, TCC formation was completely inhibited by 0.64 μM OmCI in both species (Fig. 2, right panels).

The effect of OmCI on E. coli-induced CD11b/wCD11R3 upregulation on granulocytes. Whole blood was incubated with 10⁸ E. coli/ml for 10 min, and the upregulation of CD11b (human) and wCD11R3 (pig ortholog) on granulocytes was examined. OmCI dose-dependently inhibited the expression of the cell surface marker in both species (Fig. 3). In human granulocytes the CD11b upregulation was reduced by >60% and in pigs the wCD11R3 upregulation was completely abolished at 0.32 μM OmCI.

The effect of OmCI, anti-CD14, and the combination thereof, on E. coli-induced cytokine formation in porcine and human whole blood. In pig, OmCI significantly reduced the formation of IL-8 compared with the positive control, whereas TNF-α and IL-1β were nonsignificantly reduced (Fig. 4, upper panels). Inhibition of CD14 significantly reduced the formation of pig TNF-α and IL-1β, whereas combined inhibition by anti-CD14 and OmCI had no additional effect (Fig. 4, upper panels). In human, OmCI sig-

![Figure 2](http://www.jimmunol.org/)

**FIGURE 2.** Effect of OmCI on complement induced formation of C5a and TCC in vitro. **Left panel,** Human whole blood preincubated with OmCI or equimolar amounts of albumin was incubated with 10⁸ E. coli/ml for 30 min. Data are expressed as nanograms per milliliter and presented as mean ± SEM (n = 3). **Middle panels,** Human and pig whole blood preincubated with OmCI or equimolar amounts of albumin was incubated with 10⁸ E. coli/ml for 30 min at 37°C. Data are expressed as arbitrary units (AU) per milliliter and presented as mean ± SEM (n = 3 for human and n = 4 for pig). **Right panels,** Human and pig serum preincubated with 0.64 μM OmCI or albumin was activated with HAIGG or zymosan, both at a final concentration of 1 mg/ml. In all experiments, zymosan-induced TCC formation in pig whole blood was higher than the upper standard, so these are set to 1.0. Data are expressed as TCC ratio, where 1.0 is defined as the amount of TCC formation induced by HAIGG and zymosan, respectively. The data are presented as mean ± SD (n = 5 for human and n = 3 for pig).
significantly reduced the formation of IL-1β and IL-8, whereas TNF-α was nonsignificantly reduced (Fig. 4, lower panels). Inhibition of CD14 significantly reduced the formation of human TNF-α, IL-1β, and IL-8 (Fig. 4, lower panels). In contrast to pig, combined inhibition in human by anti-CD14 and OmCI appeared to enhance the inhibitory effect observed for cytokines (Fig. 4, lower panels). In human, the combined effect of OmCI and anti-CD14 (i.e., near complete ablation of TNF-α, IL-1β, and IL-8) was similar to that of compstatin (C3 inhibitor) or C5 receptor antagonist combined with anti-CD14 (data not shown).

The effect of OmCI on LTB₄. Pig blood was preincubated with OmCI, and human blood was preincubated with OmCI and the human specific complement inhibitors eculizumab and compstatin and then activated with 10⁶ E. coli/ml and incubated for 2 h at 37°C. OmCI, as well as eculizumab and compstatin, significantly reduced the formation of LTB₄ (Fig. 5, left panels). The data from

FIGURE 3. Effect of OmCI on E. coli-induced expression of CD11b and wCD11R3. Human and pig whole blood was preincubated with OmCI or equimolar amounts of albumin and then incubated with 10⁵ E. coli/ml for 10 min at 37°C. Expression of human CD11b and pig scCD11R3 was measured using flow cytometry. Median fluorescence intensity (MFI) is expressed as mean ± SD (n = 3 for humans and n = 4 for pigs). To the right, histograms and contour plots of the effect of 1.28 μM OmCI for pig and human.

FIGURE 4. Effect of inhibiting complement, CD14, and a combination thereof, on E. coli-induced cytokine release. Upper panels. Pig whole blood was preincubated with 0.64 μM OmCl (O), 25 μg/ml anti-CD14 (whole IgG) (α), and the combination of both and then incubated with 10⁶ E. coli/ml for 2 h (TNF-α and IL-8) and/or 10⁷ bacteria/ml for 4 h (IL-1β and IL-8). The effect of anti-CD14 on pig IL-8 could not be measured because of assay interference. Albumin and an isotype-matched control Ab (IgG2b) were used as controls (Ctr) when analyzing TNF-α and IL-1β, whereas albumin only was used as control when analyzing IL-8. For IL-8, 2 and 4 h sample data were pooled. Data are normalized to the E. coli group, which is defined as 100%. Data are presented as mean ± SEM (n = 3, 4, and 5 for TNF-α, IL-1β, and IL-8, respectively). Lower panels. Human whole blood was preincubated with 0.64 μM OmCI, 10 μg/ml anti-CD14 [F(ab’)₂], combinations thereof, and controls for 5 min and then incubated for 2 h with 10⁶ E. coli/ml at 37°C. As control (Ctr), albumin was used in combination with a control F(ab’)₂, both in equimolar amounts to OmCI and anti-CD14, respectively. Data are normalized to the E. coli group, which is defined as 100% and presented as mean ± SEM (n = 3). Statistical comparisons were performed between the effect of OmCI, anti-CD14, and the combinations of both versus no inhibition. *p < 0.05.
the specific complement inhibitors indicates that LTB4 formation is C dependent. To demonstrate direct binding of OmCI to LTB4, pig and human whole blood were incubated with \textit{E. coli} to induce LTB4-enriched plasma, and then increasing doses of OmCI were added. By sequestering the LTB4 within the \beta-barrel of the lipo-
calin structure (23), OmCI dose-dependently decreased the signal in the LTB4 assay by preventing the leukotriene interacting with the capture Ab (Fig. 5, right panels). As expected, eculizumab did not have any neutralizing effect on LTB4. In vivo

The effect of OmCI on complement activity in an experimental sepsis model in pigs. To investigate the effect of OmCI on functional C activity in vivo, we used our established pig model of Gram-negative sepsis. The effect of different bolus doses of OmCI with uniform continuous infusion of the protein (0.5 mg/h) was assessed in five 15-kg pigs. No pathological side effects were observed when administering OmCI. Two control animals received \textit{E. coli} only. The three largest bolus doses of OmCI (22.5, 15, and 7.5 mg) completely ablated the classical, lectin, and alternative pathways immediately after administration and remained effective throughout the experiment (Fig. 6). The lowest bolus dose of OmCI (3.75 mg) did not totally ablate C activity at the beginning of the experiment but ablation was almost complete by the end of the experiment.

Discussion

In this study, to our knowledge, we show for the first time that OmCI is an excellent C inhibitor of humans and is as least as effective in pigs. Additionally, we demonstrated that OmCI attenuated C-induced proinflammatory mediators in pig and human. In the Wielisa analysis, OmCI was an equally potent inhibitor of the catalytic activity of the classical/lectin (C4bC3bC2a) and alternative (C3bBb3b) C5 convertases of humans and pigs. These effects were confirmed by dose-dependent inhibition of C5a and TCC formation in whole blood incubated with \textit{E. coli}. The potency...
of OmCI as a complement inhibitor was supported by total abrogation of TCC formation in serum activated with HAIGG or zymosan, which are very strong activators of the classical and alternative/lectin pathways, respectively. OmCI is known to bind directly and tightly to the C5 α-chain in the vicinity of the C5-C345C domain (18, 20, 32). OmCI binding stabilizes the overall conformation of C5 and inhibits the effect of the C5 convertases without directly blocking the cleavage site on C5 (19). At present, we lack detailed information on the precise locality and binding details of OmCI to C5, so the reason why OmCI is an equally potent inhibitor of both human and pig C5 is unknown.

OmCI significantly decreased the expression of the human cell surface marker CD11b and the pig ortholog wCD11R3, which together with CD18 constitute phagocytic C receptor 3. Because OmCI blocks the formation of C5a (18), this result is consistent with previous work showing that upregulation of this important inflammatory receptor is mainly dependent on the formation of C5a (33).

E. coli-induced cytokine formation was more dependent on CD14 than on C. CD14 is a well known recognition molecule for LPS, cooperating with MD-2 in TLR4 signaling, but CD14 has been implicated in TLR2 and TLR3 signaling (8, 34, 35) and even acts as coreceptor for TLR7 and TLR9 (36). Thus, CD14 appears to be a promiscuous upstream recognition molecule, reacting with a number of ligands with low affinity, and transferring the ligand to receptors with a higher degree of specificity and affinity (37). The experiments showed that OmCI differentially attenuated E. coli-induced cytokine formation in pig and human, with a significant effect on IL-8 in both species and on IL-1β in humans. This is consistent with previous findings showing that these cytokines are relatively more dependent on C in Gram-negative–induced inflammation (33). Interestingly, combined inhibition using OmCI and anti-CD14 enhanced the attenuating effect on cytokine formation in human, compared with the individual inhibition by these two upstream recognition molecules of innate immunity. This effect was similar to that of compstatin or C5a receptor antagonist combined with anti-CD14 and fully agrees with previous results (12).

The leukotrienes are important multifunctional mediators of inflammation. LTB₄ promotes neutrophil chemotaxis, increases adherence of neutrophils to capillary walls and is a potent inducer of chemokinesis and neutrophil infiltration (38). In the current study we showed that inhibition of C profoundly attenuated the expression of E. coli-induced LTB₄. This finding supports existing evidence that LTB₄ synthesis is C5α dependent (39). However, because OmCI also captures LTB₄ in an internal hydrophobic binding pocket, it is difficult to attribute the inhibitory effect of OmCI on the formation on LTB₄ that is due to C inhibition or due to direct binding to the leukotriene. OmCIs dual inhibitory activity may provide anti-inflammatory advantages as recently shown in an experimental model of immune complex alveolitis (M. A. Num, unpublished observations). There is evidence that the level of LTB₄ significantly increases as an endotoxemic response in pigs (40), and our data indicate that OmCI will inhibit LTB₄-induced inflammatory responses.

Data from the experiments on pigs convincingly demonstrate that OmCI is an efficient inhibitor of C in vivo. The bolus doses that were used gave estimated blood concentrations of OmCI (∼0.31–1.83 μM) corresponding to the ranges that were used in the ex vivo experiments, where 0.64 μM OmCI in undiluted serum completely inhibited C, and 0.32 μM OmCI in whole blood ablated TCC formation in response to E. coli. This enables comparison between the ex vivo and the in vivo data. With the three highest bolus doses of OmCI (∼0.61, 1.22, and 1.83 μM), C was completely inhibited in vivo. Whereas using the smallest bolus dose of OmCI (∼0.31 μM in blood), we observed incomplete inhibition of all three C pathways at the beginning of the experiment but complete inhibition by the end of the experiment. The initial bolus dose was too low to ablate the activity of the standing pool of C5 completely, whereas the continuous infusion gradually blocked more C5. Continuous infusion of OmCI is needed to inhibit newly synthesized C5, because the unexpectedly long half-life of OmCI is dependent on stable binding to C5; thus, surplus OmCI is rapidly cleared from plasma (21).

In conclusion, the present data documents that OmCI is an effective inhibitor of C activation with anti-inflammatory properties in pig and human. The comparable potency of OmCI in the two species and correspondence between the amount of OmCI needed to inhibit C ex vivo and in vivo opens the door to further testing of OmCI in pig models and will facilitate progression toward clinical testing. This is of substantial interest as the list of conditions where C is involved in the disease pathology is extensive and increasing. Given the mortality and morbidity rates associated with sepsis and the existing data on the potential efficacy of combined inhibition of C and CD14 (12, 14, 33, 41), it seems imperative to proceed with animal studies of sepsis combining inhibition of these two important upstream danger sensors of innate immunity.

Disclosures

The authors have no financial conflicts of interest.

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