Neonatal NET-inhibitory factor and related peptides inhibit neutrophil extracellular trap formation

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Neutrophil granulocytes, also called polymorphonuclear leukocytes (PMNs), extrude molecular lattices of decondensed chromatin studded with histones, granule enzymes, and antimicrobial peptides that are referred to as neutrophil extracellular traps (NETs). NETs capture and contain bacteria, viruses, and other pathogens. Nevertheless, experimental evidence indicates that NETs also cause inflammatory vascular and tissue damage, suggesting that identifying pathways that inhibit NET formation may have therapeutic implications. Here, we determined that neonatal NET-inhibitory factor (nNIF) is an inhibitor of NET formation in umbilical cord blood. In human neonatal and adult neutrophils, nNIF inhibits key terminal events in NET formation, including peptidyl arginine deiminase 4 (PAD4) activity, neutrophil nuclear histone citrullination, and nuclear decondensation. We also identified additional nNIF-related peptides (NRPs) that inhibit NET formation. nNIFs and NRPs blocked NET formation induced by pathogens, microbial toxins, and pharmacologic agonists in vitro and in mouse models of infection and systemic inflammation, and they improved mortality in murine models of systemic inflammation, which are associated with NET-induced collateral tissue injury. The identification of NRPs as neutrophil modulators that selectively interrupt NET generation at critical steps suggests their potential as therapeutic agents. Furthermore, our results indicate that nNIF may be an important regulator of NET formation in fetal and neonatal inflammation.

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Neonatal NET-inhibitory factor and related peptides inhibit neutrophil extracellular trap formation

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Introduction

Formation of neutrophil extracellular traps (NETs) may be an important component in the defensive armamentarium of neutrophils (polymorphonuclear leukocytes [PMNs]) that allows them to capture, immobilize, and putatively kill microbes in the extracellular space (1–4). NET formation occurs by a novel cell death process often called NETosis, although “vital” NETosis, in which the neutrophils do not immediately die, has also been described (3, 5). The molecular mechanisms leading to NET formation have not been completely dissected and may depend in part on the stimulus (1–3, 6). Nevertheless, decondensation of chromatin and extrusion of DNA together with histones and granule contents, are central events (1–3, 5, 6). Deamination of histones mediated by peptidyl arginine deiminase 4 (PAD4) (7–9) is thought to be a sine qua non for nuclear decondensation and NET formation (1).

NET-mediated capture and elimination of pathogens may complement traditional PMN antimicrobial activities including phagocytosis and intracellular killing (2, 10). Clinical observations indicate that defects in NET formation contribute to intractable infections in some instances (2, 11), but the importance of NETs in pathogen killing in vivo remains unclear and controversial (1–3). Conversely, there is substantial evidence that NETs and NET-associated factors, including histones and granule proteases, mediate vascular and tissue injury and that NET-mediated injury is a previously unrecognized mechanism of innate immune collateral damage to the host (1–3, 9, 12). Experimental models and limited clinical observations suggest that intra- or extravascular NET formation contributes to tissue injury in bacteremia (9, 13, 14), transfusion-related acute lung injury (15), primary graft dysfunction after lung transplantation (16), sterile vasculopathies and immune inflammation (17, 18), thrombosis (19), and influenza (20). Thus, NET formation may be an important maladaptive activity of neutrophils (1) if it is triggered inappropriately or is unregulated in infection and inflammation.

Human neonates have unique and complicated immune regulation, susceptibility to infection, and inflammatory pathology. Although the infant is in a sterile environment in utero, it can be challenged by pathogens and their products before or during labor (21). Furthermore, newborns are rapidly colonized with bacteria after delivery, a process associated with increases in circulating and bone marrow neutrophils (22–24). Complex adaptations appear to have evolved that prevent excessive, injurious inflammation in the perinatal period and in the abrupt neonatal transition from the protected intruterine environment to continuous microbial colonization and exposure (25–27). These adaptations may, however, be accompanied by increased susceptibility to infection (26, 27). Earlier, we found that PMNs isolated from umbilical cord blood of preterm and term infants do not form NETs when stimulated and have a defect in NET-mediated bacterial killing, suggesting such an adaptation (28). Other investigators subsequently reported temporally delayed NET formation when isolated neonatal neutrophils were stimulated in vitro (29). In exploring the
and from peripheral blood of infants collected at later days of life. NET formation was assessed qualitatively using live cell imaging (NETs, red fluorescence, yellow arrows; nuclear DNA, gray) and release of NET-associated histone H3 (fold change over baseline; mean ± SEM). Original magnification, ×20. Scale bar: 100 μm.

Figure 1. A NET-inhibitory factor is present in human umbilical cord blood. (A) Neutrophils from 7 preterm neonates were longitudinally examined over the first 28 days after birth for NET formation in response to LPS (100 ng/ml, 1 hour) assessed by live cell imaging (NETs, red fluorescence, yellow arrows; nuclear DNA, gray) and release of NET-associated histone H3 (fold change over baseline; mean ± SEM). Original magnification, ×20. Scale bar: 100 μm. One-way ANOVA with Tukey’s post hoc testing. *P < 0.05; **P < 0.01, compared with control histone H3 release arbitrarily set at 1 (red dashed line). (B) Neutrophils isolated from cord blood of a healthy term neonate on the day of delivery (left panel) or from venous blood on day 2 after birth (right panel) were stimulated with LPS (100 ng/ml, 1 hour) and imaged as in A. Analysis of NET formation by neutrophils from a second-term neonate yielded the same pattern. (C) Neutrophils isolated from venous blood of a healthy pregnant woman on the day of delivery were incubated in medium alone or stimulated with LPS (100 ng/ml) for 1 hour and imaged as in A. Neutrophils from a second healthy term mother also robustly formed NETs in response to LPS. Original magnification, ×60. Scale bar: 100 μm. (D) Neutrophils were isolated from venous blood of 60-day-old preterm neonates (n = 5), preincubated for 1 hour with day-60 autologous plasma or with stored autologous cord blood plasma, stimulated with LPS, and assessed for NET formation as in A. Original magnification, ×60. Scale bar: 100 μm. Neutrophils isolated from venous blood of healthy adults and preincubated in autologous or stored cord blood plasma were studied in parallel. One-way ANOVA with Tukey’s post hoc testing. *P < 0.05, LPS/adult vs. LPS/neonatal; **P < 0.01, neonatal PMNs in autologous plasma vs. cord blood plasma; †P < 0.001, adult PMNs in autologous plasma vs. cord blood plasma.

**Results**

**NET formation by human neonatal neutrophils is regulated by a peptide in umbilical cord blood.** We examined in vitro NET deployment by neutrophils from umbilical cord blood on the day of delivery and from peripheral blood of infants collected at later days of life. NET formation was assessed qualitatively using live cell imaging with SYTO Green (cell permeable) and SYTOX Orange (cell impermeable) DNA stains and quantitatively by supernatant NET-associated histone H3 measurement (30). PMNs isolated from cord blood (day 0), whether from preterm (n = 8) or healthy term infants (n = 2), did not form NETs when stimulated (Figure 1, A and B), consistent with our earlier observations (28). Nevertheless, term and preterm neonates rapidly developed durable capacity to form NETs (Figure 1, A and B). We serially assessed NET formation over the first 60 days of extrauterine life for 7 preterm neonates. Stimulated NET formation was demonstrable by day 3 ex utero for even the most prematurely born infants (Sup-
PMNs isolated from healthy pregnant women immediately before delivery robustly formed NETs (Figure 1C).

Rapid development of NET competency (Figure 1A) suggested that a factor in umbilical cord blood modulates NET formation. To complement the Table 1; supplemental material available online with this article; doi:10.1172/JCI83873DS1, and maximal NET-forming capacity was achieved between day 3 and day 14 (Figure 1A). Impaired perinatal NET formation is a feature of the neonate: PMNs isolated from healthy pregnant women immediately before delivery robustly formed NETs (Figure 1C).

Rapid development of NET competency (Figure 1A) suggested that a factor in umbilical cord blood modulates NET formation. To
test this possibility, we performed “switch” experiments in which PMNs from 60-day-old preterm neonates were preincubated with stored, day 0 autologous cord blood plasma or with freshly collected autologous day-60 venous blood plasma. PMNs from healthy adults were preincubated in day-0 cord blood plasma or, in parallel, in autologous adult plasma. Preincubation in day-0 cord blood plasma depressed NET formation by day-60 neonatal PMNs and control adult PMNs stimulated with LPS, whereas freshly isolated autologous plasma did not (Figure 1D). This result, and the time course of NET competency (Figure 1A), is consistent with a cord plasma factor that inhibits NET formation and that rapidly decreases in the circulation of the infant after delivery.

Preliminary experiments involving heat denaturation, proteinase K treatment, and lipid extraction of cord blood plasma to identify the NET inhibitory factor indicated that it is a protein (our unpublished data). We, therefore, examined the proteomes of day-0 cord blood plasma and day-28 venous blood plasma from a preterm infant whose NET-forming capacity was determined in experiments summarized in Figure 1A. 2D gel electrophoresis demonstrated protein and peptide clusters with differential representations in cord blood and day-28 plasma samples. Trypsin digest and tandem mass spectroscopic analysis of proteins from one of the clusters, using the NCBI human trypsin-specific database (ftp://ftp.ncbi.nlm.nih.gov/blast/db/Fasta/ncr.gz), yielded partial or complete sequences including a peptide in cord blood plasma with a predicted molecular mass of approximately 4 kDa. Its sequence was identical to the sequence in the carboxy terminus of α-1-antitrypsin (A1AT) (Figure 2A), a known 52-kDa plasma protease inhibitor with antiinflammatory and immunomodulatory properties (31, 32). Using Western blotting with a polyclonal antibody raised against the carboxy terminal 18 amino acids of A1AT, we found an approximately 4- to 6-kDa peptide in term infant cord blood plasma in much greater abundance than in venous plasma from healthy adults (Figure 2B) that we provisionally termed neonatal NET-inhibitory factor (nNIF). We then immunodepleted cord blood plasma using the anti-A1AT carboxy terminus antibody immobilized on affinity resin beads. Depleted plasma and peptides eluted from the affinity beads were examined for NET-inhibitory activity. Unaltered cord blood plasma inhibited NET formation by LPS-stimulated adult PMNs, as in previous experiments (Figure 1C), as did peptides eluted from the immunoaffinity beads, whereas immunodepleted plasma did not (Figure 2C). A 4- to 6-kDa candidate nNIF peptide was found in much higher quantity in the affinity purification eluate compared with the depleted plasma (Figure 2D). In parallel, we synthesized the 29-amino acid peptide detected in cord blood plasma (Figure 2A) and found that this synthetic nNIF had potent NET-inhibitory activity (Figure 2E). A scrambled control peptide (nNIF-SCR; Supplemental Table 2) did not. These results demonstrate that nNIF, or a larger protein that encompasses it, is an endogenous inhibitor of NET formation in cord blood plasma from preterm and term neonates (Figure 2B). Commercially available, active, full-length A1AT purified from human plasma and recombinant A1AT did not inhibit NET formation (Figure 2E), consistent with previous reports (33, 34), indicating that intact A1AT does not contribute to NET-inhibitory activity.

With an assay utilizing quantitative Western blotting with the anti-A1AT carboxy terminus antibody and a standard curve constructed with different concentrations of synthetic nNIF, we detected nNIF in preterm cord blood plasma samples, whereas it was undetectable, or detectable in only trace levels, in plasma from healthy adults (Figure 2B). Using the same assay, we did not detect a peptide of appropriate molecular mass in plasma samples from adult subjects (n = 10) with chronic inflammatory syndromes (granulomatosis with polyangiitis, giant cell arteritis, or rheumatoid arthritis) that might conceivably induce NET regulatory factors (our unpublished data). Thus, nNIF may be restricted to placental blood and blood of neonates in the first few days of life.

nNIF and nNIF-related peptides are a family of previously unrecognized PMN modulators. We identified additional NET inhibitory peptides based on sequence analysis of nNIF. nNIF has substantial similarity to cancer-associated SCM recognition, immune defense suppression, and serine protease protection peptide (CRISPP) (Figure 2A), a consensus peptide based on factors present in the blood of patients with cancer (35–37). We synthesized CRISPP and a scrambled control peptide (CRISPP-SCR) and found that CRISPP inhibits NET formation triggered by LPS, as does nNIF, whereas CRISPP-SCR does not (Figure 3A). nNIF and CRISPP inhibited NET formation by neutrophils isolated by Ficolli-Paque and differential centrifugation in addition to PMNs isolated by positive immunoselection. Posttreatment protocols demonstrated that nNIF and CRISPP do not degrade or dismantle previously formed NETs (Figure 4). Thus, they differ from DNases, which disrupt NETs after they are formed (2, 3, 9, 15, 38). We also examined a previously described 44-amino acid carboxy terminus cleavage fragment of A1AT, A1ATm358, which is bound to matrix in the human placenta and overlaps in sequence with nNIF (39, 40) (Supplemental Table 2). We synthesized A1ATm358 and found that it inhibits NET formation, although with lesser potency than nNIF (Figure 5).

Together, these observations indicate that nNIF, CRISPP, and A1ATm358 represent a previously unrecognized family of nNIF-related peptides (NRPs) that modulate NET formation (Figures 1–5). The presence of NRPs in umbilical blood (nNIF), placenta (A1ATm358), and, in some cases, adult plasma (CRISPP-related
peptides) suggests that NET inhibitory factors may be broadly distributed and that other NRPs may be identified.

**CRISPP and nNIF inhibit NET formation induced by a spectrum of NET-triggering agonists.** We examined the inhibitory activity of NRPs when NET formation is induced by diverse agonists, focusing on nNIF and CRISPP. Both inhibited LPS-stimulated NET deployment in multiple experiments (Figure 2E, Figure 3A, and Supplemental Figure 2). PMA is commonly employed as a potent nonphysiologic agonist to induce NET formation in vitro (1, 6, 33, 41). nNIF and CRISPP, but not CRISPP-SCR, blocked PMA-stimulated NET deployment (Figure 3B). CRISPP also inhibited NET formation induced by live *Staphylococcus aureus* (2, 9, 28, 41) (Figure 3C). This result suggests that NRPs inhibit “vital” NETosis in addition to “suicidal” NETosis, as is triggered by PMA (3, 41), since *S. aureus* is reported to release NETs by chromatin decondensation and vesicular export without neutrophil lysis (3, 5, 42). We also examined the ability of NRPs to inhibit NETosis induced by other pathogens and found that CRISPP inhibited NET generation stimulated by dengue virus (Figure 3D). Several viruses trigger NET deployment (43–45), but dengue, which interacts with ligands on myeloid cells (46), has not previously been reported as having this activity. To further explore the inhibitory activities of nNIF and NRPs, we examined heme, an endogenous damage-associated molecular pattern (DAMP) and toxin (47) that induced NETs in a murine model of sickle cell vasculopathy (17). We found that heme triggers NET formation by human PMNs and that nNIF and CRISPP inhibit this response (Figure 3E). Thus, NRPs inhibit NET deployment induced by microbes and microbial toxins, host-derived DAMPs, and pharmacologic agonists.

**NRPs selectively inhibit NET formation without interrupting other key neutrophil antimicrobial functions or platelet responses.** We examined total, phagocytic, and NET-mediated PMN killing of a pathogenic strain of *Escherichia coli* using bacterial killing assays and found that CRISPP depressed extracellular NET-mediated and total bacterial killing, but that phagocytic intracellular killing was not altered (Figure 6A). In additional incubations employing nNIF or CRISPP, the NRPs did not inhibit generation of ROS, phagocytosis, or IL-8–induced chemotaxis in Boyden chambers (Figure 6, B–D). Although each peptide was not tested in all assays, these experiments indicate that the NRPs selectively inhibit NET formation while leaving other key antimicrobial activities of PMNs intact. In addition, we found, using flow cytometry, that CRISPP does not inhibit surface translocation of P-selectin by thrombin-stimulated platelets or formation of heterotypic aggregates by activated human platelets and PMNs (Figure 6, E and F). These and other functions of activated platelets are important in antimicrobial defense (48). In addition, interaction of activated platelets with neutrophils induces NET formation (13, 15). Figure 6, E and F, indicates that PMNs, but not platelets, are cellular targets for NRPs and that platelet inflammatory activities are not disrupted by NRPs.

**nNIF and CRISPP inhibit nuclear chromatin decondensation and histone citrullination in activated neutrophils.** We used nNIF and CRISPP to inhibit NETs induced by LPS and dengue virus (Figure 3C and 3D). In a separate experiment, we used live *S. aureus* to induce NETs in murine bone marrow-derived neutrophils (Figure 3E). We found that CRISPP and nNIF inhibited NET formation by (Figure 3E). Thus, NRPs inhibit NET deployment induced by microbes and microbial toxins, host-derived DAMPs, and pharmacologic agonists.

**Figure 4. NRPs do not dismantle NETs.** (A) Neutrophils isolated from venous blood of healthy adults were incubated in medium alone (control) or activated with LPS (100 ng/ml). CRISPP (1 nM) was added at 0, 30, or 60 minutes after LPS, and the presence of NETs (red fluorescence, yellow arrows) was assessed by live cell imaging as in Figure 1A after an additional 1 hour of incubation. The images are representative of 3 separate experiments. Original magnification, ×20. Scale bars: 100 μm. (B) Isolated adult neutrophils were stimulated with LPS (100 ng/ml, 1 hour). DNase (3.78 U/ml), nNIF (1 nM), or CRISPP (1 nM) was then added, and NETs were imaged as in A after an additional 1 hour of incubation. In a second experiment, NETs were also intact after treatment with nNIF or CRISPP, but dismantled by DNase. Original magnification, ×60. Scale bars: 20 μm.
and CRISPP as probes to dissect mechanisms of action for NRPs. Generation of ROS is thought to be essential in many, but not all, pathways that mediate NET formation (1, 2, 3, 6, 9, 10, 18, 28, 33, 49), but was not inhibited by CRISPP (Figure 6B). Consistent with this result, our previous observations indicated that ROS supplementation is not sufficient to restore NET competency to neonatal PMNs (28), suggesting that nNIF also acts at a regulatory step or steps different from those influenced by ROS.

Decondensation of nuclear chromatin is a pivotal event that is required for NET formation (1–3, 5, 6, 9, 10, 18, 28, 33, 41, 49). We found that PMA induces decondensation and loss of lobulation of PMN nuclei (Figure 7A), as previously reported (6). The number of decondensed nuclei was dramatically reduced by nNIF and CRISPP, but not by nNIF-SCR or CRISPP-SCR (Figure 7A). Neutrophil chromatin decondensation is mediated by PAD4, which weakens histone-DNA binding by catalyzing conversion of histone arginines to citrullines (1, 7, 8). Consistent with this, an irreversible inhibitor of PAD4, Cl-amidine (8), blocked nuclear decondensation under the conditions of our experiments (Figure 7A). We then employed a cell-free PAD4 assay and found that nNIF blocked its activity, as did Cl-amidine used as a control (Figure 7C). In an initial comparison of NRPs, the order of potency of inhibition of PAD4 was nNIF ≈ CRISPP > A1ATm358, which is the same as their relative inhibition of NETosis. In parallel, we examined nuclear histone H3 citrullination in activated neutrophils (1, 8) and found rapid citrullination detected within 15 minutes of activation with PMA. This was inhibited by nNIF and by Cl-amidine (Figure 7D), suggesting that NRPs act at this step to block nuclear decondensation (Figure 7A).

Neutrophil elastase (NE) is also implicated in nuclear decondensation and NET formation (1, 2, 3, 6, 9, 49). An NE inhibitor, sivelestat, blocks nuclear decondensation in vitro (Supplemental Figure 4). We synthesized CRISPP and CRISPP-SCR with FLAG tags added to the carboxy terminus of each peptide (CRISPP-F, CRISPP-SCR-F) and found that both are internalized by activated neutrophils (Figure 7E), that CRISPP-F inhibits NET formation (Supplemental Figure 3), and in a protein proximity assay, that CRISPP-F is initially localized within 40 nm of NE (our unpublished experiments). This suggested that NRPs may block actions of NE in NETosis. In in vitro assays, however, neither nNIF nor CRISPP inhibited NE activity (Supplemental Figure 4 and our unpublished experiments).

NRPs inhibit NET formation in vivo and alter outcomes in systemic inflammation. To determine whether nNIF and NRPs inhibit NET formation in vivo, we established a new model of in vivo NETosis using i.p. infection of C57BL/6 mice with a clinical isolate of E. coli. Three hours after inoculation, live cell imaging of peritoneal fluid samples demonstrated robust NET formation. In addition, we observed deposition of NETs on the serosal surface of the peritoneal membranes. nNIF and CRISPP, but not CRISPP-SCR, inhibited NET formation by peritoneal fluid PMNs (Figure 8A) and deployment of NETs on the peritoneal surface (Figure 8B). Active peritonitis was demonstrated with increased neutrophil numbers and bacterial counts (Figure 8, C and D). The number of PMNs was greater in samples from CRISPP-treated animals, and the trend was to greater numbers in nNIF-treated mice (Figure 8C), potentially due to inhibition of lytic NETosis (2, 3). The number of E. coli CFUs was also greater in samples from NRP-treated animals than in those treated with the CRISPP-SCR control (Figure 8D), suggesting decreased NET-mediated bacterial killing. nNIF and CRISPP also inhibited peritoneal NET formation in Swiss Webster mice infected with E. coli (Figure 8, E and F), suggesting that this result is generalizable across mouse backgrounds.

In a second model, we found that i.p. LPS triggers peritoneal NET formation in C57BL/6 mice, although not as robustly as do live E. coli, and that LPS-induced peritoneal NET formation was inhibited by nNIF and CRISPP, but not CRISPP-SCR (our unpublished
were alive when the experiment was terminated at 50 hours \( (P < 0.02) \). In a second experiment, in which nNIF was also examined and that was extended to 72 hours, there was reduced mortality in the NRP-treated groups challenged with LPS compared with mice treated with CRISPP-SCR or LPS alone \( (n = 10 \text{ for each group}) \) at 50 hours. Figure 9A illustrates combined data from the 2 experiments. At 72 hours in the second experiment, the survival advantage provided by nNIF was durable \( (P = 0.007 \text{ compared with LPS alone}) \), whereas that of CRISPP was not. This may be due to differences in pharmacokinetics or half-lives of nNIF and CRISPP under these conditions.

We also performed similar experiments using the cecal ligation and puncture (CLP) model of polymicrobial sepsis \( (51–53) \). Mice treated with nNIF had lower clinical illness scores \( (52) \) at 24 hours and significantly increased survival at 144 hours after CLP compared with nNIF-SCR–treated animals (Figure 9, B and C). Together, these experiments (Figure 8 and Figure 9) demonstrate that nNIF and CRISPP inhibit NET formation in vivo and provide initial evidence that they have beneficial effects in models of systemic sterile inflammation and infection in which NET formation may influence tissue injury and mortality \( (9, 13, 14, 53) \).

Discussion

Inflammatory pathways and immune mechanisms have checkpoints and modulatory brakes that prevent inappropriate initiation or unregulated propagation of effector events, which could otherwise cause pathologic collaterlal injury to the host \( (54, 55) \). Tight control and modulation of inflammatory responses appear to be particularly important in the fetus and neonate, but the cellular and molecular mechanisms involved remain incompletely defined \( (25–27, 56, 57) \). nNIF in umbilical cord blood and AIATm\textsuperscript{108} in the placental matrix may represent regulatory factors that modulate NETosis in the perinatal milieu. Placental IL-8, a NETosis-inducing chemokine \( (41) \), and syncytiotrophoblast microparticles trigger NET formation in vitro, and NETs are present in placentas of women with preeclampsia \( (58) \). Thus NET-inducing stimuli appear to be generated at the maternal-fetal...
interface, suggesting that unregulated NET formation can cause inflammatory pathology in the fetomaternal environment. Excessive intrapartum NET formation could also have additional negative consequences, including long-term neonatal immune dysregulation (2, 3, 25, 26, 56, 59). Immediately after delivery, the neonate is at risk for NET-mediated vascular injury and thrombosis (1–3, 9, 13, 19, 38) triggered by microbial colonization (22, 23) and consequent neutrophil mobilization (24) if NET formation is not tightly controlled. nNIF in neonatal plasma and the related NRP A1ATm358 in the placental interstitium (40) represent potential “stop signals” (54) that selectively limit NET formation before and immediately after birth. Rapid development of full NET competency by neonatal PMNs (Figure 1) and decreased nNIF in neonatal blood in the first few days of extrauterine life parallel establishment of the resident microbiota of the human infant (22, 23) and suggest that these are regulated features of immune development. In our initial screens, nNIF was not detected or was minimally present in plasma samples from healthy adults or adult patients with chronic inflammatory syndromes (Figure 2B; our unpublished data). This suggests that nNIF expression may largely be a feature of the fetus and neonate, as are certain other immunoregulatory mechanisms (25–27). Additional studies of healthy and diseased adults, potentially with more sensitive assay methods, will be required to resolve this issue.

Our studies and previous observations suggest that nNIF and A1ATm358 are generated by proteolytic cleavage of A1AT in the placenta. A1AT is abundant in human placental tissue compartments (60, 61). It has been proposed that progressive proteolytic cleavage of A1AT occurs in the placenta, and proteases that mediate enzymatic fragmentation of A1AT to A1ATm358 in vitro are known (39, 62, 63), although a specific placental protease has not been identified. A protease that is increased in human placental syncytiotrophoblasts in the third trimester of pregnancy, high temperature requirement protease 1, cleaves A1AT in the carboxy terminus (61). Together, these findings suggest a mechanism for generation of biologically active fragments of A1AT in the placenta that would no longer be active after delivery and separation of the neonate. Nevertheless, the mechanism of generation of nNIF in the fetomaternal environment and the factors that control levels of nNIF in cord and postnatal blood (Figure 2) remain to be defined.

In addition to nNIF and A1ATm358, we identified CRISPP as an NRP. CRISPP-related peptides have been detected in plasma from patients with multiple types of cancer (35–37), but have not been linked to regulation of NETosis. NET formation facilitates experimental metastasis (64) and may also contribute to outcomes in cancer-associated infection and sepsis. Thus, endogenous CRISPP-related peptides may have significant influences on neoplastic complications by inhibiting formation of NETs, an issue that merits investigation. In this study, however, we focused on characterizing CRISPP as a lead NRP that, together with nNIF, may have broad relevance and utility in exploration of NET biology.

We found that nNIF and CRISPP inhibit in vitro NET deployment induced by S. aureus, the bacterial toxin LPS, a previously unrecognized viral trigger, dengue, a host-derived DAMP, heme, and the potent pharmacologic agonist, PMA. While we have not yet examined fungal agonists (2, 11, 65), this analysis indicates that NRPs interrupt NET formation triggered by diverse stimuli that may be mediated by distinct activation pathways (1). Furthermore, nNIF and CRISPP inhibited in vivo NET formation in a murine model of E. coli peritonitis (Figure 8). Here, NET formation is likely induced by the bacteria, LPS, and host-derived mediators, suggesting that NRPs can inhibit NET deployment by neutrophils stimulated by multiple agonists that act in combinatorial fashion in a complex inflammatory milieu and that, perhaps, induce NET formation via more than one pathway simultaneously (3). Validation of nNIF and CRISPP as inhibitors of NET deployment in this model (Figure 8) also complements analysis of in vitro inhibition (Figure 3), since it is suggested that pathways to NET formation vary in vivo and in vitro (1).

We utilized nNIF and CRISPP as probes to explore the mechanisms by which NRPs inhibit NET formation. While NETosis induced by a number of agonists involves ROS generation (1–4, 6, 18, 49, 66), current (Figure 6B) and previous (28) experiments indicate that NRPs act at a different step or steps. Based on all studies to date, chromatin decondensation is critical in NET deployment regardless of the agonist (1–3, 5, 6, 33, 42, 49). We found that nNIF and CRISPP inhibit loss of lobules and expansion of nuclei in PMA-stimulated neutrophils (Figure 7A and Supplemental Figure 4) in an assay based on earlier studies of chromatin decondensation in NETosis (6). Neutrophil heterochromatin decondensation is mediated by PAD4, which catalyzes conversion of histone arginines to citrullines with consequent weakening of histone-DNA binding and unwinding of nucleosomes (1, 7–9). In a cell-free assay of PAD4 activity based on examination of a synthetic substrate, we found inhibition by nNIF in parallel with Cl-amidine, an established PAD4 inhibitor (Figure 7C). This is consistent with inhibition of nuclear decondensation by each agent (Figure 7A). In live, intact human neutrophils activated with PMA, nNIF and Cl-amidine inhibited nuclear histone citrullination, which, as expected, occurred before nuclear decondensation (Figure 7D). In preliminary assays, CRISPP also inhibited PAD4 activity and nuclear histone citrullination (our unpublished experiments). Together, these findings indicate that NRPs inhibit nuclear decondensation and NET formation at least in part by inhibiting PAD4 activity and nuclear histone examination. NE is also thought to have a critical role in NETosis (1). NE mediates nuclear histone cleavage in PMA-activated neutrophils (6), inhibitors of NE activity block nuclear decondensation (1, 6, 33) (Supplemental Figure 4) and NET formation in vivo (64), endogenous regulators of NE influence NETosis (33, 67), and NET generation is impaired in NE-deficient mice (9). Nevertheless, we found that NRPs do not directly inhibit NE activity in vitro assays (Supplemental Figure 4 and our unpublished experiments). We do not at this time exclude the possibility that NRPs interrupt NE-mediated events in NETosis pathways in other ways, however. A FLAG-tagged construct of CRISPP that inhibits NET formation (Supplemental Figure 3) was internalized by activated PMNs (Figure 7E) and was localized to NET components (unpublished experiments), suggesting that NRPs may have more than one site and mechanism of action.

A central question in current inflammatory biology is whether NETs primarily protect the host from pathogens, and, if so, under what circumstances, or whether they, in contrast, are primarily effectors or modulators of inflammatory injury (1–3, 9, 13–20, 66, 68). Selective inhibitors of NET formation may be par-
Figure 7. NRPs inhibit nuclear decondensation and histone citrullination in activated neutrophils. (A) Neutrophils were preincubated in medium alone, with nNIF, nNIF-SCR, CRISPP, or CRISPP-SCR (all 1 nM), or with the irreversible PAD4 inhibitor Cl-amidine (10 μM) for 1 hour, treated with PMA (20 nM), and then incubated on poly-L-lysine-coated coverslips for 2 hours, followed by examination for nuclear decondensation (arrows) by live cell imaging. Green fluorescence, nuclear DNA. Original magnification, ×60. Scale bars: 20 μm. (B) Nuclear areas were measured (n = 4) using ImageJ software (mean nuclear pixel area per cell ± SEM). Paired Student’s t test. *P < 0.05; **P < 0.01. (C) nNIF (1 nM), nNIF-SCR (1 nM), or Cl-amidine (10 μM) was examined in a cell-free deimination assay employing recombinant PAD4 and a synthetic substrate. One-way ANOVA with Tukey’s post hoc testing. **P < 0.001. (D) Left panels: neutrophils were preincubated for 30 minutes in medium, with nNIF or nNIF-SCR (both 1 nM) or with Cl-amidine (10 μM), activated for 15 minutes with PMA (20 nM), and citrullinated-histone H3 was detected by immunocytochemistry (n = 3). Green fluorescence, citrullinated-histone H3; magenta fluorescence, nuclear DNA. Original magnification, ×60. Scale bars: 20 μm. Right panel: histone H3 citrullination was quantified using ImageJ software (mean citrullinated-histone H3 pixel area per cell ± SEM) (n = 3). One-way ANOVA with Tukey’s post hoc testing. *P < 0.05. (E) Neutrophils were incubated with FLAG-tagged CRISPP-FLAG or CRISPP-SCR-FLAG (1 nM for both) for 1 hour, activated with LPS (100 ng/ml) for a further 2 hours, and then examined by confocal microscopy using an anti-FLAG antibody (n = 3). Yellow fluorescence, FLAG tag; blue fluorescence, nuclear counterstain. Original magnification, ×60. Scale bars: 20 μm. The FLAG-tagged peptides were not internalized by isolated human platelets (our unpublished experiments).

The results in both models suggest that NET generation, like other neutrophil effector functions (54), has evolved to contain and eliminate pathogens, but can also injure the host if it is activated by pathologic inflammatory signals in the absence of infection or by microbes in an uncontrolled fashion. Our findings also suggest that NRPs have potential as antiinflammatory therapies (71) in specific syndromes in which NET formation contributes to acute or progressive pathologic inflammation.

Methods

Animal studies. Swiss Webster and C57BL/6 male mice between the ages of 8 and 12 weeks were purchased from Charles River Laboratories or Jackson Laboratories for all experiments. Mice were housed in specific pathogen-free microisolator cages that were located in a room maintaining a constant temperature and on a 12-hour light/12-hour dark cycle. All treatment groups were weight matched and randomized to treatment at the initiation of an experiment. The researchers conducting the experiments were blinded to the experimental groups during testing. No inclusion or exclusion criteria were used in designing the experiments.

Reagents. Lipopolysaccharide (E. coli serotype O111:134 and Salmonella enteritidis), poly-L-lysine, cytochalasin B, cytochalasin D, paraformaldehyde (PFA), sivelestat, NE, the NE substrate (MeOSuc)-AAPV-pNA, and thrombin were purchased from Sigma-Aldrich. Additional reagents were as follows: TOPRO-3, phalloidin, SYTO Green (cell-permeable DNA stain), and SYTOX Orange (cell-impermeable DNA stain) (Molecular Probes); Cl-amidine (Calbiochem); DNase (Promega); anti-CD15-microbeads (Miltenyi Biotec); Medium-199 (Lonza), and micrococcal DNase (Worthington).

nNIF and NRP synthesis. nNIF, NRPs, and their specific scrambled peptide controls (Supplemental Table 2) were synthesized by the DNA/Peptide Facility, a unit of the Health Sciences Center Cores at the University of Utah. The core facility also verified the sequence and purity of the provided peptides.

PMN and platelet isolation. PMNs were isolated from ACD or EDTA anticoagulated venous blood from healthy adults, healthy term infants, and prematurely born infants (28, 30). For the 8 prematurely born infants from whom we collected cord and peripheral blood samples, cord and peripheral blood plasma and PMN preparations were obtained at 5 separate time points throughout the first 2 months of life. PMN suspensions (>96% pure) were prepared by positive immunoselection using anti-CD15-coated microbeads and an auto-MACS cell sorter (Miltenyi Biotec) and were resuspended at 2 × 10⁶ cells/ml concentration in serum-free M-199 media at 37°C in 5% CO₂/95% air. Human platelets were isolated as described (72).

Live cell imaging of NET formation. Qualitative assessment of NET formation was performed as previously reported (28, 30). Briefly, primary PMNs isolated from preterm infants, healthy term infants, and healthy adults (2 × 10⁶ cells/ml) were incubated with control buffer or stimulated with indicated agonists or bacteria for 1 hour at 37°C in 5% CO₂/95% air on glass coverslips coated with poly-L-lysine. For selected experiments, primary PMNs were preincubated with autologous plasma, cord blood plasma, nNIF (0.2–70 nM), CRISPP (0.2–70 nM), nNIF-SCR (1 nM), or CRISPP-SCR (1 nM) for 1 hour prior to stimulation. After preincubation and/or stimulation, PMNs were gently washed with PBS and incubated with a mixture of cell-permeable (SYTO Green, Molecular Probes) and -impermeable (SYTOX Orange, Molecular Probes) DNA fluorescent dyes. Confocal microscopy was accomplished using a FV1000 1X81 confocal microscope and Fluoview software (Olympus). Both 20× and 60× objectives were used. Z-series images were obtained at a step size 1 μm over a range of 20 μm for each field. Olympus FluoView and Adobe Photoshop CS2 software were used for image processing.

Imaging of dengue virus-induced NET formation. Using BSL 2 safety protocols, primary PMNs isolated from healthy adults (2 × 10⁶ cells/ml) were incubated with mock infection buffer or live dengue virus (MOI 0.05) as for live cell imaging. After a 1-hour incubation, the infected PMNs were fixed with 2% PFA for 10 minutes prior to incubation with fluorescently labeled, cell-permeable, and cell-impermeable DNA dyes and imaged as for live cell imaging using confocal microscopy.
Figure 8. nNIF and CRISPP inhibit in vivo NET formation. (A and B) C57BL/6 mice not pretreated or pretreated with nNIF, CRISPP, or CRISPP-SCR (10 mg/kg i.p.; 1 hour) were inoculated with E. coli (4.5 × 10^7 bacteria i.p.). After 3 hours, animals were sacrificed and peritoneal fluid and membranes collected for analysis. (A) Peritoneal fluid NET formation (red fluorescence, yellow arrow) assessed by live cell imaging and histone H3 release (red dashed line, baseline arbitrarily set at 1). Three mice per group. Original magnification, ×60. Scale bars: 50 μm. One-way ANOVA with Tukey’s post hoc testing. ***P < 0.001, CRISPP and nNIF versus CRISPP-SCR. (B) NET formation on the surfaces of peritoneal membranes (red fluorescence, yellow arrows) was quantified by counting the number of NETs that crossed standardized grid lines in 4 random microscopic fields using ImageJ software. Original magnification, ×60. Scale bars: 50 μm. A second experiment yielded a similar pattern. (C) C57BL/6 mice not pretreated (left 2 bars) or pretreated with CRISPP, nNIF, or CRISPP-SCR were inoculated with E. coli i.p. as in A and B, and neutrophil numbers in peritoneal fluid were counted after 3 hours (3–5 mice/group). One-way ANOVA with Newman-Keul post hoc testing; ∗P < 0.05, CRISPP vs. CRISPP-SCR or not pretreated; **P < 0.05, control vs. all other groups. (D) C57BL/6 mice were pretreated with CRISPP or CRISPP-SCR and inoculated with E. coli i.p. as in A and B (5 animals/group). After 3 hours, bacteria CFU in the peritoneal fluid were measured. Single-tailed Mann-Whitney U test. *P < 0.05. (E and F) Swiss Webster mice not pretreated or pretreated with nNIF, CRISPP, or CRISPP-SCR were inoculated with E. coli i.p. as in A and B. After 3 hours, peritoneal fluid and membranes were collected. (E) Peritoneal fluid NET formation, image and measured as in A (10 mice/group). **P < 0.05, CRISPP/E. coli and nNIF/E. coli compared with CRISPP-SCR/E. coli and E. coli. (F) NET formation on peritoneal membrane surfaces, imaged and quantitated as in B (3 mice in each group). ∗P < 0.05, E. coli versus control (red dashed line); **P < 0.01, CRISPP/E. coli versus control; †P < 0.05, CRISPP/E. coli and nNIF/E. coli versus CRISPP-SCR/E. coli. One-way ANOVA with Tukey’s post-hoc testing applied in E and F.

Quantitation of NET formation — NET-associated histone H3 content. We determined NET-associated histone H3 content as previously described (30). After live cell imaging of control and stimulated primary PMNs (2 × 10^6/ml; various agonists), the cells were incubated with PBS containing DNase (40 U/ml) for 15 minutes at room temperature to break down and release NETs formed in response to stimulation. The supernatant was gently removed and centrifuged at 420 xg for 5 minutes. The cell-free supernatant was then mixed 3:1 with 4× Laemmli buffer prior to Western blotting. We used a polyclonal primary antibody against human histone H3 protein (Cell Signaling) and infrared-conjugated secondary antibodies (LI-COR). Imaging and densitometry were performed on the Odyssey infrared imaging system (LI-COR). We previously validated this assay as a surrogate for NET formation under in vitro conditions (30) as employed in the current studies.

Isolation and identification of nNIF in umbilical cord blood plasma. Two plasma samples from a single preterm infant, one from the umbilical cord blood and one from a peripheral blood sample taken on ex utero day 28, were subjected to abundant plasma protein removal (Norgen) prior to 2D electrophoresis using separation first by isoelectric focusing (pH range 3–8) and then by size (TGX Precast Gel, Bio-Rad). The resulting gels were compared for differential protein content. Six differentially expressed protein clusters (“spots”) were sent to the University of Utah Proteomics Core for analysis. Following trypsin digestion and tandem mass spectroscopy using an LTQ-FT ion-trap/FTMS hybrid mass spectrometer (ThermoElectron), candidate proteins/peptides were identified as potential NET-inhibitory factors.

Affinity removal of nNIF. Plasma samples were subjected to abundant plasma protein removal (Norgen). A polyclonal antibody raised against the carboxy terminal 18 amino acids A1AT (LifeSpan BioSciences) coupled to resin beads from an immunoprecipitation kit purchased from Thermo Fisher Scientific was then used to immunodeplete the samples. Nonimmune IgG coupled to resin beads was used in parallel as a control. Plasma was diluted in lysis buffer from the kit and incubated with the anti-A1AT C terminus antibody–coupled beads or with control beads overnight at 4°C. The beads were then separated by centrifugation and the immunodepleted and control plasma samples collected. The A1AT C terminus antibody coupled and control beads were resuspended in kit-included elution buffer for 10 minutes at room temperature, followed by centrifugation and collection of the eluate and control supernatants. The eluate was analyzed by Western blotting (16.5% Tris-Tricine Gel, Bio-Rad) using the A1AT C terminus antibody and by tandem mass spectroscopy. Immunodepleted plasma and eluate samples were examined in assays of NET formation. Active full-length native and recombinant A1AT (both from Sigma-Aldrich) were suspended in elution buffer and tested in parallel.

Bacterial killing assay. We determined NET-mediated and phagocytic bacterial killing by primary human PMNs as previously described (28).

Chemotaxis assay. Chemotaxis by PMNs isolated from healthy adult donors was assessed using a modified Boyden chamber assay with or without a 1-hour preincubation with nNIF (1 nM), CRISPP (1 nM), or CRISPP-SCR (1 nM). Recombinant human IL-8 (2 ng/ml) was used as the chemotacticant. Chemotaxis through a 5-micron filter was determined by counting PMNs in 10 randomly selected high-power fields as previously described (73). In separate experiments, nNIF, CRISPP, or CRISPP-SCR (all at 1 nM) were evaluated for chemotacticant activity using the same system.

Phagocytosis assay. PMNs were isolated from blood of healthy adult donors and resuspended in M-199 at a concentration of 2 × 10^6 cells/ml. Leukocytes were preincubated for 60 minutes under standard conditions with cytochalasin D and B (10 μM), nNIF (1 nM), CRISPP (1 nM), or CRISPP-SCR (1 nM). Following preincubation, PMNs were incubated with 6 × 10^4 E. coli bioparticles (Molecular Probes) on a rotator for 4 hours at 37°C in 5% CO₂/95% air. The PMNs were then washed and resuspended in the starting volume of M-199 before being spun down onto glass coverslips and fixed with 2% PFA for 10 minutes and permeabilized with 0.1% Triton X-100 for 10 minutes. Leukocytes were stained with WGA 555 (Invitrogen) and TOPRO-3 (Molecular Probes), and randomly selected high-power visual field images were captured using confocal microscopy. We used ImageJ software (NIH) to determine the percentage of PMNs that were positive for fluorescently labeled E. coli bioparticles detected at 488 nm.

Reactive oxygen species generation. Human PMNs isolated from healthy adult whole blood were resuspended to a concentration of 2 × 10^6 cells/ml in M-199 media and preincubated with or without CRISPP (1 nm) or CRISPP-SCR (1 nm) peptide for 1 hour at 37°C in 5% CO₂/95% air. The PMNs were then stimulated with LPS (100 ng/ml) for 1 hour, washed, and resuspended with a dihydrodroadamine (7.25mM; Molecular Probes) and catalase (1000 units/ml; Sigma-Aldrich) mixture and incubated at 37°C for 10 minutes. After incubation, samples were placed at 4°C and analyzed for ROS-dependent fluorescence using flow cytometry as performed in the University of Utah core facility (BD, CellQuest software).

Platelet activation assays. P-selectin translocation and surface display by activated platelets (74) and formation of platelet-neutrophil aggregates (75) were measured as described.
was incubated with recombinant PAD4 and PAD4 enzyme sub-unity using a PAD4 Inhibitor Screening Assay Kit (Cayman). Briefly, nNIF incubated with PMA (20 nM) for 15 minutes at 37°C in 5% CO²/95% air, followed by stimulation with LPS (100 ng/ml) for 2 hours. The nuclear decondensation assay was identified as described (6). Five randomly selected high-power visual fields per sample were captured via confocal microscopy and analyzed for nuclear area using the cell-permeable, fluorescent DNA dye SYTO Green. The nuclear pixel areas of more than 100 individual cells per high-power field were determined using ImageJ software (NIH).

PAD4 activity assay. We determined nNIF inhibition of PAD4 activity using a PAD4 Inhibitor Screening Assay Kit (Cayman). Briefly, nNIF (1 nM) was incubated with recombinant PAD4 and PAD4 enzyme substrate (2 mM) in PAD4 assay reagent for 30 minutes at 37°C. The PAD4 inhibitor Cl-amidine (10 μM) for 1 hour at 37°C in 5% CO²/95% air, and treated with or without PMA (20 nM) on poly-L-lysine–coated glass coverslips for 2 hours. Nuclear decondensation was identified as described (6). Five randomly selected high-power visual fields per sample were captured via confocal microscopy and analyzed for nuclear area using the cell-permeable, fluorescent DNA dye SYTO Green. The nuclear pixel areas of more than 100 individual cells per high-power field were determined using ImageJ software (NIH).

Histone H3 citrullination determination. Adult PMNs were stimulated with PMA (20 nM) for 15 minutes at 37°C in 5% CO²/95% air following a 15-minute preincubation with nNIF, CRISPP, nNIF-SCR, or CRISPP-SCR (1 nM) or with Cl-amidine (10 μM), spun onto poly-L-lysine–coated slides, and examined by immunocytochemistry with a primary antibody used to detect human citrullinated histone H3 (Abcam). Imaging was accomplished via confocal microscopy using a FV1000 1X81 confocal microscope and Fluoview software (Olympus). Semi-quantification was accomplished using ImageJ software (NIH) to determine the average citrullinated histone H3 content per cell.

CRISPP peptide cellular localization. FLAG-tagged CRISPP (F-CRISPP) and FLAG-tagged CRISPP-SCR (F-CRISPP-SCR) peptides were synthesized by our core facility and detected using immunocytochemistry. Adult neutrophils were preincubated with either F-CRISPP (1 nM) or F-CRISPP-SCR (1 nM) for 1 hour at 37°C in 5% CO²/95% air, followed by stimulation with LPS (100 ng/ml) for 2 hours. The

PMNs were then spun down onto glass coverslips with 2% PFA fixation and 0.1% Triton X-100 permeabilization. FLAG-tagged peptide was detected using a monoclonal anti-FLAG antibody (Sigma-Aldrich) with TOPRO-3 as a nuclear counterstain.

Mouse models of E. coli and LPS-induced peritonitis. C57BL/6 or Swiss Webster mice were pretreated in blinded fashion with CRISPP (10 mg/kg), nNIF (10 mg/kg), or CRISPP-SCR (10 mg/kg) by i.p. injection 1 hour prior to infection (E. coli, 4.5 × 10⁷ CFU/mouse, i.p.) or inoculation (LPS, 25 mg/kg, i.p.) (Sigma-Aldrich). Control mice were injected with saline alone. The mice were sacrificed in a CO² chamber 3 hours after infection/inoculation and the peritoneal fluid and membrane samples harvested. Briefly, the abdomen was disinfected and opened in the midline without injuring the muscle. The peritoneal cavity was lavaged with sterile saline solution (1 ml) and analyzed for in vivo NET formation, bacteriology, and leukocyte accumulation. NETs in the peritoneal fluid were qualitatively and quantitatively analyzed using live cell imaging with confocal microscopy and NET-associated histone H3 release assays. NETs on the serosal surface of the peritoneal membrane were assessed quantitatively using live cell imaging, followed by standardized grid analysis of 5 randomly selected high-power visual fields per tissue sample (ImageJ, NIH). We quantified peritoneal bacterial CFU counts by permeabilizing all recovered leukocytes with 0.1% Triton X-100 for 10 minutes and performing serial dilutions and bacterial cultures on 5% sheep blood agar plates (Hardy Diagnostics). After a 24-hour incubation, bacterial counts were determined. Total leukocyte counts in the peritoneal lavage were determined in Neubauer chambers using an optical microscope after dilution in Turk’s solution (2% acetic acid). Differential leukocyte analysis was performed using a 60X oil immersion objective to assess morphology of cyt centrifuged cells stained with May-Gruenwald-Giemsa dye. All mice were included in the final analysis.

Mouse model of systemic inflammation induced by LPS (endotoxemia). C57BL/6 mice were pretreated in blinded fashion with CRISPP (10 mg/kg), nNIF (10 mg/kg), or CRISPP-SCR (10 mg/kg) by i.p. injection 1 hour prior to and 6 hours after inoculation with LPS (25 mg/kg, i.p. injection). Control mice were injected i.p. with saline alone. Fluid resuscitation and antibiotic treatment were not used in these experiments.

Figure 9. nNIF and CRISPP improve survival in experimental systemic inflammation. (A) C57BL/6 mice were challenged with LPS (25 mg/kg i.p.). CRISPP, nNIF, or CRISPP-SCR (10 mg/kg i.p.) was given 1 hour before and 6 hours after LPS. Animals with no treatments or given LPS alone were studied in parallel (n = 10 mice for each condition). **P < 0.01, log-rank (Mantel-Cox) statistical tool used. The survival difference between nNIF-LPS and CRISPP-LPS compared with CRISPP-SCR-LPS trended toward significance. *P < 0.051. (B) C57BL/6 mice were subjected to CLP. nNIF or nNIF-SCR (10 mg/kg i.p.) was given 1 hour before and 6 hours after surgery (n > 7 in each group). Mice subjected to sham surgery were studied in parallel (n - 3 in each group). Clinical illness severity scores (52) were determined at 24 hours. One-way ANOVA with Newman-Keul post hoc testing; **P < 0.01, nNIF vs. nNIF-SCR groups. (C) Mice assessed for severity of systemic illness in B were then followed daily and survivors sacrificed at 144 hours. Log-rank (Mantel-Cox) statistical tool used. **P < 0.01.
ments. Survival was assessed over 50- or 72-hour intervals. All mice were included in the final survival analysis.

Mouse model of polymicrobial sepsis using CLP. C57BL/6 mice were anesthetized with ketamine/xylazine (100 mg/kg and 10 mg/kg, i.p., respectively), and CLP was performed as previously described (52). nNIF (10 mg/kg) or nNIF-SCR (10 mg/kg) i.p. was given 1 hour prior to and 6 hours after CLP surgery. The animals received subcutaneous sterile isotonic saline (1 ml) for fluid resuscitation immediately after the surgery. Sham-operated mice were subjected to identical procedures except that CLP was not done. Twenty-four hours after CLP, all animals were scored for clinical illness severity as previously described (52). In this assessment, higher scores reflect increased illness severity. Survival of mice in the nNIF/CLP (n = 7), nNIF-SCR/CLP (n = 8), nNIF/sham surgery (n = 3), or nNIF-SCR/sham surgery (n = 3) groups was followed for 5 days after the surgical procedure.

NE activity assay. Synthetic fluorogenic substrate of NE (MeOSuc)-AAPV-(pNA) (160 μM) was incubated with bioactive NE (500 nM) with or without the NE inhibitor sivelstat (160 μM) or nNIF (50 nM) for 3 hours at 37°C. The reactions were quenched with 5% glacial acetic acid and centrifuged at 18,500 RCF for 5 minutes. Chromatograms were obtained using an Agilent 1100 Series HPLC and a Phenomenex 5 μm C18 Luna column (100 Å, 4.6 × 150 mm) over a 30-minute 10% to 90% B gradient (buffer A 0.1% TFA in H2O, buffer B 0.1% TFA in ACN). Mass spectra were obtained for secondary validation of the reaction products using an API 3500 triple quadrupole mass spectrometer. Chromatograms were offset on both the x and y axes (by 0.5 minutes and 0.1 A214, respectively) for greater visibility. Relative A214 was determined by normalizing all of the data to the tallest HPLC peak displayed in each graph.

Statistics. We used GraphPad Prism statistical software (version 5) to analyze results. Mean ± SEM was determined for each experimental variable. All data used in each statistical test met the assumption of the specific test and were normally distributed. P < 0.05 was considered statistically significant.

Study approval. The University of Utah Institutional Review Board approved this study (IRB nos. 0392, 11919, and 39621), and all human subjects provided informed consent in accordance with the Declaration of Helsinki. All murine experiments were performed by the University of Utah Institutional Animal Care and Use Committee (no. 12-11017) in a facility approved by the American Association for Laboratory Animal Care.

Author contributions

CCY, ASW, and GAZ conceived of the project and designed the experiments. CCY and CLK performed patient phenotyping. MTR provided statistical expertise. MJG, HS, JAW, RAC, AVDA, CVA, SS, and JMF conducted experiments. RAC, AVDA, CVA, ESH, and JWR developed experimental tools or mouse models. CCY and GAZ wrote the manuscript.

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