Immunoregulation of Neutrophil Extracellular Trap Formation by Endothelial-Derived p33 (gC1q Receptor)

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Abstract
The formation of neutrophil extracellular traps (NETs) is a host defence mechanism, known to facilitate the entrapment and growth inhibition of many bacterial pathogens. It has been implicated that the translocation of myeloperoxidase (MPO) from neutrophilic granules to the nucleus is crucial to this process. Under disease conditions, however, excessive NET formation can trigger self-destructive complications by releasing pathologic levels of danger-associated molecular pattern molecules (DAMPs). To counteract such devastating immune reactions, the host has to rely on precautions that help circumvent these deleterious effects. Though the induction of DAMP responses has been intensively studied, the mechanisms that are used by the host to down-regulate them are still not understood. In this study, we show that p33 is an endothelial-derived protein that has the ability to annul NET formation. We found that the expression of human p33 is up-regulated in endothelial cells upon infections with \textit{Streptococcus pyogenes} bacteria. Using tissue biopsies from a patient with streptococcal necrotising fasciitis, we monitored co-localisation of p33 with MPO. Further in vitro studies revealed that p33 is able to block the formation of DAMP-induced NET formation by inhibiting the enzymatic activity of MPO. Additionally, mice challenged with \textit{S. pyogenes} bacteria demonstrated diminished MPO activity when treated with p33. Together, our results demonstrate that host-derived p33 has an important immunomodulating function that helps to counterbalance an overwhelming DAMP response.

Introduction
Activation of polymorphonuclear cells or neutrophils (PMNs) is one of the first host immune responses to combat invading microbes. To this end, PMNs are equipped with a manifold arsenal of antimicrobial tools including the degranulation of bacteriolytic agents [1], the phagocytosis of pathogens [2], and the formation of neutrophil extracellular traps (NETs) [3]. Within the last decade, this last mechanism has attracted much attention because...
blood-derived PMNs employ a sophisticated approach to expel web-like structures that can entrap and kill pathogens such as *Staphylococcus aureus*, *Salmonella typhimurium*, and *Shigella flexneri* [3]. Subsequent work by Fuchs et al. [4] showed that PMNs can undergo a special cell death program (NETosis) in order to release their DNA into the surroundings. During this process, antimicrobial agents such as histones, myeloperoxidase (MPO), neutrophil elastase, and the host defence peptide LL-37 become attached to DNA fibres [3, 5, 6] which can further increase the antimicrobial activity of the formed NETs [7].

Though PMN activation and the subsequent generation of NETs are considered important defence mechanisms, recent studies also report that excessive NET formation or impaired NET degradation can be detrimental to the host [8–12]. Along these lines, it has been described that the exaggerated mobilisation of some NET-associated molecules such as LL-37 and extracellular histones can evoke overwhelming inflammatory reactions [13]. Notably, the secretion of LL-37 under pathological conditions [14] can lyse bacterial but also mammalian cell membranes [15], and similar findings have been reported for the histones released from necrotic cells [16]. It therefore seems plausible that the immune system has to take special precautions to counteract the self-destructive activity of these danger-associated molecular patterns (DAMPs). However, the underlying regulatory mechanisms behind such down-regulating mechanisms are so far only poorly understood.

Recently, it was reported that p33, also known as the gC1q receptor or p32, is able to inhibit the cytotoxic activity of several antimicrobial peptides, including LL-37 and extracellular histones [17, 18]. Originally, p33 was discovered for its affinity to complement factor C1q [19], but, since then, proteins such as high-molecular-weight kinogen [20], thrombin [21], and lamin B receptor [22] as well as a number of pathogen-associated molecular pattern (PAMP) molecules [23, 24] have also been described as p33 binding partners. The expression of p33 is to be found in many cells types, including lymphocytes, dendritic cells, endothelial cells, and platelets [20, 25, 26]. In these cells, p33 has been detected in various compartments such as the nucleus, mitochondria, cell membrane, and extracellular matrix [27, 28]. The expression of p33 is up-regulated in the endothelial cells by pro-inflammatory stimuli including TNFa, IFNy, and lipopolysaccharide [29], or under hypoxic conditions [30]. The fact that p33 can protect host cells from a lytic attack by DAMPs such as LL-37 and extracellular histones and that it is up-regulated under pathological conditions points to an important role of the protein in dampening an over-accelerated host response under infectious disease conditions [31]. Thus, p33 is an interesting protein for studying its ability to down-regulate the toxic side effects evoked by the release of DAMPs exceeding pathological levels.

In this study, we investigated the influence of p33 on the DAMP-facilitated NET generation. We found that p33 is up-regulated in endothelial cells upon bacterial challenge. Moreover, our results show that p33 is able to dampen NET formation by binding to and inhibiting the enzymatic activity of MPO, a crucial protein in the neutrophil host defence.

### Materials and Methods

#### Bacterial Strain

*Streptococcus pyogenes* strain AP1 (40/58; WHO Collaborating Centre for Reference and Research on Streptococci, Institute of Hygiene and Epidemiology, Prague, Czech Republic) was grown in THY medium at 37°C to mid-log phase. For the infection experiments, the bacteria were washed with PBS and re-suspended in DMEM medium (Gibco) + 10% FCS (Thermo Fisher Scientific [Thermo]) or PBS. For the NET induction experiments, bacteria were heat-inactivated for 15 min at 65°C and stored at –80°C until further use.

#### Animals

Wild-type C57BL/6 mice (Charles River Laboratories) were housed under standard conditions and fed with laboratory chow. Water and food were given ad libitum. Experiments were performed with 22-week-old male animals and were approved by the local ethics committee, the Malmö-Lund Animal Care Ethics Committee (M138-13).

#### PMN Isolation and NET Induction

PMNs were isolated from blood donated by healthy donors (approved by the ethics committee at Lund University, Lund, Sweden) as described previously [32]. Briefly, heparinised blood was layered on top of PolymorphPrep™ (Axis-Shield) and centrifuged at 370 g for 30 min at room temperature (RT). Neutrophils were re-suspended in HBSS medium with Ca²⁺/Mg²⁺ (Gibco) and 2 × 10⁵ cells were seeded onto poly-L-lysine-coated (Sigma) glass slides. Samples were incubated for 3 h at 37°C with 5% CO₂ with 5 μM LL-37 (Schafer-N) or 50 μg/mL calf thymus histones (CTHs; Roche) +/- MBP-p33 (333.3 or 100 μg/mL, respectively; purified as described previously [18]). When 25 nM PMA (Sigma) was used, 100 μg/mL p33 was added. *S. pyogenes* AP1 was used at a multiplicity of infection (MOI) of 10. Cells were fixed with a final 4% PFA.

#### Fluorescence Microscopy

NET structures were visualised using an MPO antibody (Santa Cruz) or a DNA/histone 1 antibody (Millipore). Anti-p33 immune serum (Innovagen AB) and anti-β-actin (Abcam) were used in visualisation experiments of p33. Primary antibodies were incubated for 1 h at RT or overnight at 4°C. AlexaFluor 488A or AlexaFluor 633O (Invitrogen) were added as secondary antibodies for 45 min to detect the antibody conjugations. Samples were finally embedded in ProLong®.
incubated for 3 h at 37 °C with 5 µM LL-37, 50 µg/mL CTHs, or overnight at 4 °C, and then ProLong anti-MPO (Santa Cruz) and anti-p33 (Innovagen AB) took place [33], with 2 modifications. Incubation with primary antibodies, sectioned, fixed, and stained (essentially as described previously [1]) was performed. Slides were analysed using an Eclipse TE300 fluorescence microscope with a PlanFluor 40X/0.6 DIC N2/1.4 NA objective (all by Nikon) or a DeltaVision OMX V4 (GE Healthcare) with an 60x/1.42 NA Planapo objective (Olympus) and immersion oil with a refraction index of 1.524. Fifty-eight z-stacks, 0.125-µm-thick, were used for the displayed images.

**Tissue Preparation and Immunostaining**

Snap-frozen soft tissue collected from a patient with necrotising fasciitis infected with an emm1 S. pyogenes strain was cryosectioned, fixed, and stained (essentially as described previously [33], with 2 modifications). Incubation with primary antibodies, anti-MPO (Santa Cruz) and anti-p33 (Innovagen AB) took place overnight at 4 °C, and then ProLong Gold Antifade Mountant with DAPI (Molecular Probes) was added together with the secondary antibodies, AlexaFluor 488 and AlexaFluor 546 (Molecular Probes). A Nikon AIR confocal microscope was used for imaging and image evaluation (Nikon Instruments).

**Scanning Electron Microscopy**

For the scanning electron microscopy (SEM) analysis, 2 × 10^4/mL cells were treated with either 25 nM PMA alone or 25 nM PMA + 100 µg/mL p33 for 3 h at 37 °C and 5% CO_2. Samples were spun down with 370 g, re-suspended in 2.5% glutaraldehyde (Merck) in 15 mM Na-acid (Sigma) and transferred to poly-L-lysine coated titanium slides. After overnight fixation at RT, samples were dehydrated for 1 h and critical-point dried. Finally, samples were coated with 20 nm Au-palladium and analysed using a Phenom ProX desktop scanning electron microscope (PhenomWorld) with an acceleration voltage of 10 kV.

**MPO Release and Enzyme Activity Detection**

For the measurement of MPO activity, 1 × 10^5 cells per well were incubated for 3 h at 37 °C with 5 µM LL-37, 50 µg/mL CTHs, or S. pyogenes (MOI 10 +/- 333.3 or 100 µg/mL p33). After incubation, supernatants were transferred into a new plate (Nunc). Collected air-pouch fluids were handled accordingly. For enzyme activity inhibition, 5 µM of recombinant MPO (Thermo) were incubated +/- 333.3 µg/mL of full p33 or 48 µg/mL p33-derived peptides for 30 min at RT; 3’-(p-aminophenyl) fluorescein (Invitrogen) was used as substrate. Fluorescence intensity was measured according to the manufacturer’s recommendations at ex/em 485/530 nm.

**MPO-p33 ELISA**

The interaction of MPO with p33 or p33-derived peptides was tested using an ELISA as described earlier [17]. Briefly, plates (Nunc) were coated overnight at 4°C with 2 µg/mL p33 or 0.48 µg/mL peptide in coating buffer (15.9 mM Na_2CO_3, 30 mM NaHCO_3, pH 9.6) and blocked in PBS-Tween for 60 min at 37°C. Subsequently, plates were incubated with 1 µg/mL MPO for 1 h at 37°C. Binding of MPO was detected using an MPO antibody (Santa Cruz) and a (H + L)-HRP conjugated antibody (BioRad) each, at 37°C for 1 h. Stabilised TMB chromogen (Thermo) was used as a substrate. The reaction was terminated with 10% H_2SO_4 and measured at 450 nm.

**Western Blot Analysis of p33**

EA.hy926 cells were grown to 80% confluency and infected with S. pyogenes strain AP1 for 3 h with a MOI of 10. Cells were lysed with RIPA buffer (Thermo) and SDS PAGE was run for 35 min at constant 200 V. Samples were transferred onto PVDF membrane (Millipore) for 30 min at constant 15 V. Membranes were incubated overnight at 4 °C with primary antibodies (anti-p33 and anti-β-actin, Abcam). Secondary HRP-conjugated antibody (BioRad) was added for 1 h at RT. Bands were visualised using West Pico chemiluminescent substrate (Thermo).

**p33 Transfection and Co-Incubation Experiments**

EA.hy926 cells were transfected using 25 nM p33 siRNA (Hs_C1QBP_6) or All Stars Negative Control (Qiagen) using Oligo-fectamine in OptiMEM medium (Life Technologies). Cells were incubated with the siRNA for 48 h at 37 °C and 5% CO_2. PMNs were pre-incubated with 25 nM PMA for 20 min and then added onto the endothelial cells with 1 × 10^6 cells/mL and co-incubated for another 3 h at 37 °C and 5% CO_2. Supernatants were collected and the release of MPO was measured as described above.

**Chemotaxis Assay**

PMNs were isolated as described above and seeded with 200 µL 2 × 10^5 cells in the upper compartment of a transwell insert with a pore size of 3.0 µm; 500 µL of 10, 50, or 100 µg/mL p33, or 10 nM fMLP was used as chemo-attractant. HBSS medium was used as a negative control. Following incubation for 1, 2, or 3 h at 37 °C, the cells in the lower compartment were counted using a Nikon bright-field microscope with a ×20 bright-field objective and ImageJ software.

**Animal Infection Experiment**

Male C57BL/6 mice (aged 22 weeks) were subcutaneously infected with 10^6 CFU/mL S. pyogenes AP1 into an air pouch, as described earlier [34]. PBS was used as the vehicle control. After 24 h, 50 µg/mL p33 or PBS was added for an additional 8 h. Air pouches were re-filled with PBS and the fluids were collected. MPO release was analysed as described above. Cell number was determined using a LUNA™ automated cell counter (Logos Biosystems).

**Statistical Analysis**

Data was analysed by using GraphPad Prism v7.0 (GraphPad Software). Differences between 2 groups were analysed by using a well chemotaxis assay. Neutrophils were incubated for 1, 2, and 3 h with 10, 50, and 100 µg/mL p33 and 10 nM fMLP. e Total cell count of fluids collected from S. pyogenes air-pouch infection. Data represents mean +/- SEM of 3 independent experiments. Ctr, control/non-treated cells; Inf, infected cells. * p ≤ 0.05, ** p ≤ 0.01, *** p ≤ 0.001, **** p ≤ 0.0001.

(For figure see next page.)

**Fig. 1.** Upon bacterial infection, p33 is up-regulated in endothelial cells. a Representative fluorescence micrographs of EA.hy926 cells control or infected with S. pyogenes strain AP1 after 3 h. DNA, blue; p33, purple; β-actin, green. Scale bar, 10 µm. b Western blot of EA.hy926 cell lysates in non-treated and infected cells after 3 h. c p33 bands were quantified and normalised to β-actin. d Transfection EA.hy926 cells were grown to 80% confluency and infected with S. pyogenes strain AP1 for 3 h with a MOI of 10. Cells were lysed with RIPA buffer (Thermo) and SDS PAGE was run for 35 min at constant 200 V. Samples were transferred onto PVDF membrane (Millipore) for 30 min at constant 15 V. Membranes were incubated overnight at 4 °C with primary antibodies (anti-p33 and anti-β-actin, Abcam). Secondary HRP-conjugated antibody (BioRad) was added for 1 h at RT. Bands were visualised using West Pico chemiluminescent substrate (Thermo).
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**Figure a**
- Merge, β-Actin, p33, DAPI

**Figure b**
- Ctr, Inf
- p33, β-Actin

**Figure c**
- p33 protein expression, % (vs. control)
- Ctr, Inf

**Figure d**
- Cells/power field
- Ctr, 10 µg/mL, 50 µg/mL, 100 µg/mL
- MBP-p33
- 1 h, 2 h, 3 h

**Figure e**
- Total cell count
- Mock, AP1, AP1 + MBP-p33

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Fig. 2. p33 impairs DAMP-mediated NET release. a Representative immuno-fluorescent micrographs display the results shown in b and c. NET structures were visualised with MPO (green) and DAPI (blue). Scale bar, 50 µm. b, c Human neutrophils were treated for 3 h with 5 µM LL-37 and 50 µg/ml CTHs, respectively. Recombinant p33 was added simultaneously in a 2:1 ratio. 3 independent experiments +/- SEM. Ctr, control. ** p ≤ 0.01, **** p ≤ 0.0001.
Fig. 3. a Representative fluorescence microscopy images of the PMA-triggered NET release. Histone H1, green; nuclei, blue. Scale bar, 50 µm. b Inhibitory effect of 100 µg/mL p33 on human PMNs incubated with 25 nM PMA for 3 h. c, d Scanning electron micrographs of 25 nM PMA-induced NETs in the presence or absence of 100 µg/mL p33. ×1,850. Scale bar, 30 µm. e, f Representative fluorescence images of NET formation induced by S. pyogenes AP1. Histone H1, green; nuclei, blue. Scale bar, 30 µm. g The addition of 100 µg/mL p33 shows significant inhibition of S. pyogenes-induced NET release. Data is for 3 independent experiments; unpaired one-tailed Student t test +/- SEM. *** p < 0.001, **** p ≤ 0.0001.
Results

Under Infectious Conditions: p33 Is Up-Regulated and Attracts PMNs

Recent studies have shown that MrgClqR, the p33 homologue in fresh-water prawns, is significantly elevated upon bacterial challenge [35]. To study whether human p33 is also up-regulated upon infection with gram-positive bacteria, we treated human umbilical vein endothelial EA.hy926 cells (ATCC® CRL-2922™) for 3 h with the S. pyogenes AP1 strain. Protein expression under infectious conditions was investigated by employing fluorescence microscopy and Western blot analysis. A clear up-regulation of p33 by fluorescence microscopy (Fig. 1a) was noted in infected cells compared to the uninfected control, and confirmed by Western blot analysis (Fig. 1b, c).

During infection or tissue damage, DAMPs and other cellular components can activate neutrophils and attract them to migrate to the site of bacterial colonisation (review [36]). A cell migration assay revealed that after 1 h of incubation, 100 µg/mL MBP-p33 acts as a chemo-attractant for PMNs in vitro (Fig. 1d). Based on these findings, we decided to use this concentration for further experiments. To investigate this phenotype in an animal model, we infected 22-week-old male C57BL/6 mice with S. pyogenes AP1 strain. Protein expression under infectious conditions was investigated by employing fluorescence microscopy and Western blot analysis. A clear up-regulation of p33 by fluorescence microscopy (Fig. 1a) was noted in infected cells compared to the uninfected control, and confirmed by Western blot analysis (Fig. 1b, c).

Administration of p33 Inhibits DAMP- and PMA-Triggered NET Formation

Next, we investigated the effect of p33 on NET formation triggered by DAMPs like LL-37 and histones. These antimicrobial molecules, when excessively released during inflammatory conditions [14] or tissue damage [37, 38], can have a self-destructive impact [39–41]. It has been shown that the interaction of MBP-p33 with LL-37 and histones can abrogate the cytolytic activity of these agents, which in turn helps to counteract endogenous destruction [17, 18]. In addition, it has also been reported that both LL-37 and histones can trigger the release of NETs [32, 42]. To address the hypothesis whether p33 can modulate NET formation, we stimulated human PMNs with LL-37 and CTHs for 3 h in the presence or absence of MBP-p33. Co-incubation of PMNs with 5 µM LL-37 for 3 h led to a greater release of NETs (Fig. 2a, b). Similar findings were recorded when CTHs (50 µg/mL) were used, even though NET formation was less pronounced (Fig. 2a, c). The simultaneous administration of MBP-p33, in a 2:1 ratio to either LL-37 (Fig. 2b) or CTHs (Fig. 2c), almost completely abolished the formation of NETs. We therefore concluded that p33 is involved in the regulation of NET formation triggered by DAMPs like LL-37 and histones. Next, we stimulated human PMNs with 25 nM PMA (a well-described NET inducer [3, 43]) for 3 h, which resulted in a significant release of extracellular DNA fibres. This was significantly diminished by the simultaneous application of 100 µg/mL MBP-p33 (Fig. 3a, b). SEM confirmed this observation. PMA treatment led to extensive, widespread fibrous structures (Fig. 3c) but under the same experimental conditions, co-stimulation with MBP-p33 resulted in less-pronounced NET structures (Fig. 3d).

It has also been reported that NETs are induced by bacteria such as S. pyogenes [44]. We wanted to establish whether p33 could affect bacteria-induced NET formation in a manner similar to that described above for DAMPs and PMA. To this end, we incubated PMNs for
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**Graphs and Figures:**

- **Figure a:** Comparing DNA, MPO, and Merge images under different conditions: Ctr, PMA, PMA + p33.
- **Figure b:** Merge images showing DNA, MPO, and p33.
- **Figure c:** Representative images under various conditions.
- **Figure d:** Graph showing release of MPO, relative fluorescence units, with bars for Ctr, PMA, neg Ctr siRNA, and p33 siRNA.
- **Figure e:** Graph showing LDH cytotoxicity (A490 nm) with bars for EA.hy926 alone and EA.hy926 + PMNs.
- **Figure f:** Graph showing release of MPO, relative fluorescence units, with bars for LL-37 and LL-37 + MBP-p33.
- **Figure g:** Graph showing release of MPO, relative fluorescence units, with bars for Histone, Histone + MBP-p33.
3 h with *S. pyogenes* AP1. To prevent immediate degradation of NETs by bacterial nucleases [45], the bacteria were heat-inactivated. As depicted in Figure 3e–g, *S. pyogenes* AP1 triggered NET formation, which was then almost completely abolished by the addition of MBP-p33.

**p33 Co-Localises with MPO in Infected Tissue and Interferes with MPO Activity**

In the next series of experiments, we wished to unravel the molecular mechanisms employed by p33 to modulate NET formation. Microscopic analysis revealed MPO translocation to the nucleus and chromatin loosening only in the PMA-treated PMNs (Fig. 4a, middle panel), which is in line with reports by others [46]. However, the addition of MBP-p33 to PMA-stimulated PMNs led to a resting of MPO in the granules and condensed chromatin (Fig. 4a, lower panel). This finding was supported by confocal microscopy over a 3-h time course, showing that in PMA-treated PMNs, nuclei start to decondense after 30 min of incubation but that in PMA/p33-treated PMNs, MPO remains in the granules around the nucleus. After 180 min, in the PMA-treated cells, almost all nuclei (80–90%) displayed a mix of chromatin and MPO (online suppl. Fig. 1; for all online suppl. material, see www.karger.com/doi/10.1159/000480386) when compared to PMA/p33-treated cells (approx. 15%). Super-resolution microscopy revealed that externally added MBP-p33 is located in close proximity to the neutrophil protein MPO (Fig. 4b). Consequently, we analysed biopsies from a patient with soft-tissue necrotising fasciitis caused by *emm*1 *S. pyogenes* infection to test whether this interaction has a patho-physiological implication. As shown in Figure 4c, on confocal fluorescence microscopy, we noted co-localisation of endogenous p33 and MPO in severely inflamed tissue.

Next, we sought to investigate whether p33 is able to interfere with the mobilisation of MPO from its granular compartment into the nucleus, which is an established sign of NETosis [3, 47]. Previous work has shown that, upon PMN activation and subsequent ROS production, MPO, in concert with neutrophil elastase, can induce chromatin decondensation and histone degradation in a dose-dependent manner [48]. Based on these reports, we decided to study the effect of endogenous p33 secreted from the endothelial cell line on the release of MPO from PMNs. As a control, we employed EA.hy926 cells in which p33 levels were down-regulated using an siRNA approach. Figure 4d depicts that PMNs incubated with EA.hy926 cells, lacking normal p33 expression, mobilised more MPO than cells treated with negative-control siRNA. To exclude any cytotoxic effect of PMNs on the endothelial cells during co-incubation, we performed a lactate dehydrogenase assay. After a 3-h co-incubation period, we did not measure any increase in lactate dehydrogenase levels compared to endothelial cells alone (Fig. 4e). These results are in line with reports from others [49], and demonstrate that PMNs and their secretion products fail to influence the viability of endothelial cells after 3 h of co-incubation.

Next, we investigated the effect of recombinant MBP-p33 on the release of MPO from PMNs alone, induced by LL-37 and histones. MBP-p33 significantly impaired the degranulation of MPO triggered by LL-37 and by histones (Fig. 4f, g). When measuring the secretion of MPO, we found that PMA stimulation resulted in an increased release of MPO but that the addition of MBP-p33 to PMA-stimulated cells significantly blocked the mobilisation of MPO (Fig. 5a). These findings correlate well with the results obtained from the co-incubation studies (Fig. 4d).

To obtain more structural information about the sequence motif of p33, which is crucial for attenuating the release of MPO from the granules, we included a panel of synthetic peptides derived from p33 [17]. Our data shows that 3 p33-derived peptides (QEP20, TGE20, and FLA20, respectively) significantly blocked the mobilisation of MPO (Fig. 5a). The release of MPO was also measured when PMNs were treated with *S. pyogenes* AP1. Similar to the results obtained with DAMP/PMA-stimulated cells, p33 displayed an inhibitory effect on the se-
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**Binding to MPO, A450 nm**

**Release of MPO, relative fluorescence units**

**ROS production, relative fluorescence units**

**Release of MPO, relative fluorescence units**

**MPO activity, relative fluorescence units**

**Incubation Conditions**

- + 25 nm PMA
- + 5 U/mL MPO
- Mock AP1 AP1 + MBP-p33
- MBP-p33
- DEI LPK QEP VIK CHY TED FLA
- PMAPMA +
The secretion of MPO (Fig. 5b). The impact of MBP-p33 on MPO release was also measured in an animal model. In 22-week-old male C57BL/6 mice infected with S. pyogenes AP1, we found that the addition of MBP-p33 led to a reduced mobilisation of MPO into the air pouch when challenged with the bacteria (Fig. 5c). Next, we focused on the effect of p33 and p33-derived peptides on the enzyme activity of recombinant MPO. Figure 5d depicts strong competitive enzyme inhibition by MBP-p33, confirming our hypothesis that this protein can block the induction of NET via enzyme inhibition (Fig. 5d). In addition to p33, DEI20, LPK20, VIK20, CHY20, and TDS20 also significantly impair the activity of MPO (Fig. 5d). Active MPO enzyme plays an important, but not solitary part in the production of reactive oxygen species (ROS [50]). We thus tested the influence of p33 on ROS production in PMA-stimulated PMNs over time. As shown in Figure 5e, the production of ROS was impaired by the addition of 100 µg/mL of MBP-p33. Finally, we employed an indirect ELISA to test whether MBP-p33 or p33-derived peptides can bind to MPO. As shown in Figure 5f, direct interaction of MBP-p33 to MPO was detected, and the p33-derived peptides, VIK20 and TGE20, displayed a strong affinity for MPO comparable to the full-length protein. Using the Web-based iCn3D structure viewer [51], the structure of p33 was visualised (Fig. 5g). When looking at the interaction sites of p33 with MPO, it appears that peptides binding to (TGE20) or interfering with MPO enzyme activity (VIK20) are clustered, and located in the inside of the doughnut-shaped trimer (Fig. 5g). This data suggests that p33 interferes with the release of NETs due to the competitive inhibition of MPO enzyme activity. It provides new insights into the role of the multi-ligand protein p33 in innate immunity.

**Discussion**

This study provides a novel insight into how p33 can modulate the generation of NETs. The release of NETs has been described as an immune response of the host to inflammation and infection [52]. However, in recent years, evidence has accumulated that excessive NET formation and the impaired degradation of the fibre-like structures is correlated to pathological conditions such as sepsis [53] and cell damage [54]. DAMPs, like LL-37 and histones, are cationic, antimicrobial components that are found entangled in the fibrous NET meshwork, and eventually cause considerable damage to the host [39, 40]. Our data shows that MBP-p33 can dampen the formation of NETs triggered by endogenous DAMPs, LL-37, and histones as well as by chemical stimuli, e.g., PMA.
The multi-ligand protein p33 can be detected on cell surfaces and in mitochondria [55, 56]. Previous work has shown that p33 is up-regulated by a variety of pro-inflammatory factors including interferons [29], suggesting that the protein has important functions during infections. As a proof of concept, we infected human umbilical vein endothelial cells EA.hy926 for 3 h with S. pyogenes, which led to an up-regulation of p33 as detected by fluorescence microscopy and Western blot analysis. These findings are in good correlation with earlier reports of increased p33 levels under inflammatory condition [30]. Having shown that p33 is up-regulated under infectious conditions in human endothelial cells, we next explored this in infected patients. To this end, we analysed tissue biopsies from a patient with severe S. pyogenes soft-tissue infections, which revealed the presence of p33 and MPO at the infected site, as well as co-localisation of the factors.

To unravel the role of p33 in regulating NETosis, a series of experiments were undertaken. Immunofluorescence microscopy revealed that the administration of p33 to PMA-stimulated PMNs results in the persistence of MPO in the granules, which did not mix with the still intact, not yet de-condensed chromatin. In addition, a co-incubation model of endothelial cells with neutrophils revealed increased MPO release when p33 was down-regulated with specific siRNA. For an in vivo proof, MPO release was followed in a mouse model, where S. pyogenes bacteria were injected subcutaneously into an air pouch. Here, we also noted a reduced mobilisation of MPO after the administration of MBP-p33 into the air pouch.

It has been suggested that the active MPO enzyme might be important for chromatin breakdown and subsequent NET formation [48, 57]. Our data show that the co-incubation of PMNs with MBP-p33 and its derived peptides resulted in the reduced release of MPO and blocked its enzyme activity. Chemical inhibition of MPO by aminobenzoic acid hydrazide (4-ABAH), which reduced enzyme activity irreversibly by 23% [58], had no effect on chromatin de-condensation [48], but did result in a decrease of NET formation [59]. We therefore speculated that a more physiological inhibitor like the endothelial protein p33 could indeed affect the chromatin de-condensation that is necessary to perform NET formation. This led us to conclude that our findings are of importance also under infectious disease conditions. Excess activation of PMNs by DAMPs or bacteria like S. pyogenes and the subsequent release of granular enzymes such as MPO at the site of infection can limit the repair of injury (review [36]) and boost the pathological conditions seen under sepsis or respiratory distress conditions [60]. Additionally, since MPO plays a crucial role in the production of ROS, which can have destructive effects on inflamed tissue, a diminished release of MPO could be beneficial to the host. Thus, our study adds new insights into how the delicate balance of an overwhelming immune response is regulated by endogenous proteins like p33 (Fig. 6). Future studies are needed to further elucidate a possible role of p33 as an effective therapeutic target to attenuate overwhelming PMN-evoked inflammatory responses.

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**Disclosure Statement**

The authors declare no competing financial interests.

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**Author Contributions**

A.N. designed the study, performed the experiments, analysed the data, and wrote the manuscript. P.P. performed the animal infection experiments. J.W. performed the ELISA experiment. O.H. collected the NSTI biopsy samples. J.S. and A.N.-T. performed the infection experiments. H.H. advised and wrote the manuscript. P.P. performed the animal infection experiments. J.W. performed the ELISA experiment. O.H. collected the NSTI biopsy samples. J.S. and A.N.-T. performed the NSTI tissue sectioning and staining. H.H. advised and wrote the manuscript.

**References**

1. Lehrer RI, Ganz T: Antimicrobial peptides in mammalian and insect host defence. Curr Opin Immunol 1999;11:23–27.
2. Mitschiniok E: Lecture on phagocytosis and immunity. BMJ 1891; 1: 213–217.
3. Brinkmann V, Reichard U, Goosmann C, Fauler B, Uhlemann Y, Weiss DS, Weinrauch Y, Zychlinsky A: Neutrophil extracellular traps kill bacteria. Science 2004;303:1532–1535.
41 Xu J, Zhang X, Pelayo R, Monestier M, Ammollo CT, Semeraro F, Taylor FB, Esmon NL, Lupp F, Esmon CT: Extracellular histones are major mediators of death in sepsis. Nat Med 2009;15:1318–1321.

42 Abrams ST, Zhang N, Manson J, Liu T, Dart C, Baluwa F, Wang SS, Brohi K, Kipar A, Yu W, Wang G, Toh CH: Circulating histones are mediators of trauma-associated lung injury. J Inflam 2013;10:160–169.

43 Mohany T, Sjogren J, Kahn F, Abu-Humaidan AH, Fisker N, Assing K, Morgelin M, Bentgsson AA, Borregaard N, Sorensen OE: A novel mechanism for NETosis provides anti-microbial defense at the oral mucosa. Blood 2015;126:2128–2137.

44 Li P, Li M, Lindberg MR, Kennett MJ, Xiong N, Wang Y: PAD4 is essential for antibacterial innate immunity mediated by neutrophil extracellular traps. J Exp Med 2010;207:1853–1862.

45 Sumby P, Barbian KD, Gardner DJ, Whitney AR, Welty DM, Long RD, Bailey JR, Parnell MJ, Hoe NP, Adams GG, Deleo FR, Musser JM: Extracellular deoxyribonuclease made by group A Streptococcus assists pathogenesis by enhancing evasion of the innate immune response. Proc Natl Acad Sci USA 2005;102:1679–1684.

46 Brinkmann V, Zychlinsky A: Beneficial suicide: why neutrophils die to make NETs. Nat Rev Microbiol 2007;5:577–582.

47 Urban CF, Ermert D, Schmid M, Abu-Abed U, Goossman C, Nacken W, Brinkmann V, Jungblut PR, Zychlinsky A: Neutrophil extracellular traps contain calprotectin, a cytosolic protein complex involved in host defense against Candida albicans. PLoS Pathog 2009;5:e1000639.

48 Papayannopoulos V, Metzler KD, Hakkim A, Zychlinsky A: Neutrophil elastase and myeloperoxidase regulate the formation of neutrophil extracellular traps. J Cell Biol 2010;191:677–691.

49 Gupta AK, Joshi MB, Philippova M, Erne P, Hasler P, Hahn S, Resink TJ: Activated endothelial cells induce neutrophil extracellular traps and are susceptible to NETosis-mediated cell death. FEBS Lett 2010;584:3193–3197.

50 Spickett CM, Jerlich A, Panasenko OM, Arnhold J, Pitt AR, Stelmaszynska T, Saur R: The reactions of hypochlorous acid, the reactive oxygen species produced by myeloperoxidase, with lipids. Acta Biochim Pol 2000;47:889–899.

51 Wang Y, Geer LY, Chappey C, Kans JA, Bryant SH: Ca3D: sequence and structure views for Entrez. Trends Biochem Sci 2000;25:300–302.

52 Goldmann O, Medina E: The expanding world of extracellular traps: not only neutrophils but much more. Front Immunol 2013;3:420.

53 Clark SR, Ma AC, Tavener SA, McDonald B, Goodarzi Z, Kelly MM, Patel KD, Chakrabarti S, McAvoy E, Sinclair GD, Keys EM, Allen-Vercoe E, Deviney R, Doig CJ, Green FH, Kubes P: Platelet TLR4 activates neutrophil extracellular traps to ensnare bacteria in septic blood. Nat Med 2007;13:463–469.

54 Saffarzadeh M, Juenemann C, Queisser MA, Lochnit G, Barreto G, Galuska SP, Lohmeyer J, Preisser KT: Neutrophil extracellular traps directly induce epithelial and endothelial cell death: a predominant role of histones. PLoS One 2012;7:e32366.

55 Soltys BJ, Kang D, Gupta RS: Localization of P32 protein (gC1q-R) in mitochondria and at specific extramitochondrial locations in normal tissues. Histochem Cell Biol 2000;114:245–255.

56 Waggoner SN, Cruise MW, Kassell R, Hahn YS: gC1q receptor ligation selectively down-regulates human IL-12 production through activation of the phosphoinositide 3-kinase pathway. J Immunol 2005;175:4706–4714.

57 Parker H, Dragunow M, Hampton MB, Kettle AJ, Winterbourn CC: Requirements for NADPH oxidase and myeloperoxidase in neutrophil extracellular trap formation differ depending on the stimulus. J Leukoc Biol 2012;92:841–849.

58 Kettle AJ, Gede CA, Winterbourn CC: Mechanism of inactivation of myeloperoxidase by 4-aminobenzoic acid hydrazide. Biochem J 1997;321:503–508.

59 Metzler KD, Fuchs TA, Nauseef WM, Reumaux D, Roels J, Schulze I, Wahn V, Papayannopoulos V, Zychlinsky A: Myeloperoxidase is required for neutrophil extracellular trap formation: implications for innate immunity. Blood 2011;117:953–959.

60 Lacy P: Mechanisms of degranulation in neutrophils. Allergy Asthma Clin Immunol 2006;2:98–108.

61 Jiang J, Zhang Y, Krainer AR, Xu RM: Crystal structure of human p32, a doughnut-shaped acidic mitochondrial matrix protein. Proc Natl Acad Sci USA 1999;96:3572–3577.