Rapid Neutrophil Destruction following Phagocytosis of *Staphylococcus aureus*

Scott D. Kobayashi\(^a\)  Kevin R. Braughton\(^a\)  Amy M. Palazzolo-Ballance\(^a\)
Adam D. Kennedy\(^a\)  Elizabeth Sampaio\(^d,g\)  Ervand Kristosturyan\(^d\)
Adeline R. Whitney\(^a\)  Daniel E. Sturdevant\(^b\)  David W. Dorward\(^c\)
Steven M. Holland\(^d\)  Barry N. Kreiswirth\(^e\)  James M. Musser\(^f\)  Frank R. DeLeo\(^a\)

\(^a\) Laboratory of Human Bacterial Pathogenesis, and Research Technologies Section, \(^b\) Genomics and \(^c\) Microscopy Units, Rocky Mountain Laboratories, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Hamilton, Mont. \(^d\) Laboratory of Clinical Infectious Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Md., \(^e\) Public Health Research Institute and the University of Medicine and Dentistry of New Jersey, Newark, N.J., and \(^f\) Center for Molecular and Translational Human Infectious Diseases Research, Department of Pathology, The Methodist Hospital Research Institute, Houston, Tex., USA; \(^g\) Leprosy Laboratory, Oswaldo Cruz Institute, FIOCRUZ, Rio de Janeiro, Brazil

**Key Words**
Neutrophils  ·  Bacterial infections  ·  Host defense  ·  *Staphylococcus aureus*

**Abstract**
Mechanisms underlying the enhanced virulence phenotype of community-associated methicillin-resistant *Staphylococcus aureus* (CA-MRSA) are incompletely defined, but presumably include evasion of killing by human polymorphonuclear leukocytes (PMNs or neutrophils). To better understand this phenomenon, we investigated the basis of rapid PMN lysis after phagocytosis of USA300, a prominent CA-MRSA strain. Survival of USA300 clinical isolates after phagocytosis ultimately resulted in neutrophil lysis. PMNs containing ingested USA300 underwent morphological changes consistent with apoptosis, but lysed rapidly thereafter (within 6 h), whereas cells undergoing FAS-mediated apoptosis or phagocytosis-induced cell death remained intact. Phagosome membranes remained intact until the point of PMN destruction, suggesting lysis was not caused by escape of *S. aureus* from phagosomes or the cytolytic action of pore-forming toxins. Microarray analysis of the PMN transcriptome after phagocytosis of representative community-associated *S. aureus* and healthcare-associated MRSA strains revealed changes unique to community-associated *S. aureus* strains, such as upregulation of transcripts involved in regulation of calcium homeostasis. Collectively, the data suggest that neutrophil destruction after phagocytosis of USA300 is in part a form of programmed necrosis rather than direct lysis by *S. aureus* pore-forming toxins. We propose that the ability of CA-MRSA strains to induce programmed necrosis of neutrophils is a component of enhanced virulence.

S.D.K. and K.R.B. contributed equally to this paper.

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Dr. Frank R. DeLeo
Laboratory of Human Bacterial Pathogenesis, Rocky Mountain Laboratories National Institute of Allergy and Infectious Diseases, National Institutes of Health 903 South 4th Street, Hamilton, MT 59840 (USA) Tel. +1 406 363 9448, Fax +1 406 363 9394, E-Mail fdeleo@niaid.nih.gov
Introduction

Human polymorphonuclear leukocytes (PMNs or neutrophils) are critical for host defense against invading bacteria. The acute inflammatory response to infection is characterized by the rapid recruitment of neutrophils. After ingestion of microorganisms, PMNs generate intraphagosomal reactive oxygen species (ROS) and nascent phagosomes are enriched with microbicidal proteins [1]. Collectively, these antimicrobial processes are highly effective at eliminating most invading bacteria. On the other hand, pathogens such as Staphylococcus aureus moderate and/or circumvent these processes and ultimately cause disease [reviewed in 2]. Although progress has been made, the mechanisms used by bacterial pathogens to evade killing by neutrophils are incompletely defined.

S. aureus remains a significant health problem worldwide and is the leading cause of bacterial infections in the United States. The high prevalence of these infections is compounded by antibiotic resistance among S. aureus. Notably, methicillin-resistant S. aureus (MRSA) are endemic in most healthcare settings [reviewed in 3]. Although historically associated with healthcare facilities, MRSA has emerged as a leading cause of bacterial infections in the community over the past 10–15 years [4]. An important distinction between healthcare-associated MRSA (HA-MRSA) and community-associated MRSA (CA-MRSA) is with the groups of individuals at risk for infection. HA-MRSA infections occur in individuals with risk factors for infection, such as surgery or immunodeficiency, whereas CA-MRSA cause infections in otherwise healthy individuals [4]. CA-MRSA strains prevalent in the United States, including pulsed-field gel electrophoresis types USA300 and USA400, have enhanced virulence by comparison [5]. Although S. aureus in general have capacity to survive after phagocytosis by neutrophils [6, 7], a phenomenon due in part to modulation of the effects of ROS and antimicrobial peptides [8–10], USA300 and USA400 have increased capacity to evade killing by human PMNs compared to representative HA-MRSA strains [5]. Inasmuch as neutrophils are the primary cellular defense against S. aureus infections, increased survival of USA300 and USA400 after uptake may be linked to the ability of these strains to cause rapid PMN lysis after phagocytosis [5]. Therefore, neutrophil lysis after phagocytosis is a process potentially important in the pathogenesis of CA-MRSA infection. The mechanism for this process is not known.

To better understand CA-MRSA virulence, we used functional assays, video and transmission electron microscopy, and human oligonucleotide microarrays to investigate the mechanism underlying enhanced PMN lysis that occurs after phagocytosis of USA300. Our results suggest that the rapid neutrophil death occurring after phagocytosis of S. aureus is in part a host cell-mediated process.

Materials and Methods

Bacterial Strains and Culture

S. aureus strains LAC (USA300) [5, 11] and other USA300 clinical isolates [12], MW2 (USA400) [13], MnCop (USA400, MSSA) [14], COL [15] and MRSA252 (USA200) [16] were grown overnight in tryptase soy broth (TSB; BD Biosciences). Overnight cultures of S. aureus were diluted in fresh TSB at 1:200 for USA300 strains, MW2, MnCop, 1:150 for MRSA252 and 1:100 for COL. S. aureus were cultured at 37 °C with shaking (250 rpm) to mid-exponential growth phase (OD600 = 0.75), at which time they were washed with Dulbecco’s phosphate-buffered saline (DPBS; Sigma-Aldrich), opsonized using 50% autologous human serum, washed twice with DPBS and resuspended in RPMI 1640 medium (Invitrogen) buffered with 10 mM HEPES (RPMI/H, pH 7.2). Serum-opsonized bacteria were used immediately for experiments. For selected experiments, bacteria were heat-killed at 95 °C for 5 min or killed by UV irradiation for 1 min at 2,000 × 100 μm/cm² using a Hoefer ultraviolet crosslinker (Thermo Fisher Scientific). These conditions were determined empirically to be optimal for killing S. aureus.

Serum complement-opsonized and IgG-opsonized latex beads (C3bi/IgG-LB) were prepared as described previously [17].

Isolation of Human PMNs

Human neutrophils were isolated from heparinized venous blood of healthy donors or those with chronic granulomatous disease using dextran sedimentation and Hypaque-Ficoll gradient centrifugation as described previously [17]. This work was performed in accordance with protocols approved by the Institutional Review Board for Human Subjects, National Institute of Allergy and Infectious Diseases. All human subjects gave informed consent prior to participation in the study. Purity of PMN preparations and cell viability were determined by flow cytometry (FACSCalibur; BD Biosciences) as described previously [17]. Using this method, cell preparations typically contain approximately 98–99% granulocytes; primarily neutrophils with approximately 2–5% eosinophils.

Phagocytosis of S. aureus by Human PMNs

For experiments that evaluated post-phagocytosis processes, that is, cell lysis, apoptosis, ROS production, microarray analyses, transmission electron microscopy (TEM) and live-cell imaging, synchronized phagocytosis was performed in the same manner for each experiment essentially as described [17–19]. In general, bacteria were washed in DPBS and opsonized with 50% autologous normal human serum at 37 °C for 30 min. Bacteria were washed twice with DPBS and resuspended at the desired concentration in RPMI/H. Human neutrophils were aliquoted at the desired concentration in tissue culture plates precoated with normal...
human serum (to prevent PMN activation). Neutrophils were allowed to adhere to the plate for 30 min on ice (if no inhibitors were used for the experiment) or at room temperature (with the desired inhibitor added) before bacteria were added to the plates. Bacteria were added to the desired wells and centrifuged at approximately 350 relative centrifugal force for 10 min at 4°C to synchronize phagocytosis. A ratio of 10 bacteria per PMN was used for all experiments unless otherwise indicated. In some experiments, PMNs were allowed to ingest S. aureus for 30 min and gentamicin (5 μg/ml final concentration) or lysostaphin (6.25 U/ml) were subsequently added to assay wells to prevent potential contribution of any bacteria remaining uningested. S. aureus survival and permeabilization of neutrophil plasma membranes (pore formation) were determined using published methods [5, 20].

**Cell Lysis Experiments**

PMNs (1 × 10⁶) were aliquoted into wells of 96-well plates and Z-Y-VAD-FMK (Y-VAD, 250 μM; EMD Chemicals), Z-VAD-FMK (Z-VAD, 10–300 μM; EMD Chemicals), actinomycin D (0.5–10 μg/ml; Sigma), peptatin A (10–200 μM) and puromycin (50–1,000 μg/ml; Sigma) were added to appropriate wells and plates were incubated at room temperature for 30 min before bacteria were added unless indicated otherwise. Serum-opsonized bacteria (1 × 10⁵) were added to the desired wells and phagocytosis was synchronized as described above. Following centrifugation, plates were incubated at 37°C with 5% CO₂ for up to 6 h. PMN lysis was determined with a standard assay for release of lactate dehydrogenase (LDH) as described by the manufacturer (Cytotoxicity Detection kit; Roche Applied Sciences). Statistics were performed using repeated-measures ANOVA (GraphPad Prism Version 5.0 for Windows; GraphPad Software).

**PMN Apoptosis Assays**

To estimate early-stage apoptosis, neutrophils were stained with annexin-V-FITC and propidium iodide as described previously [17]. Stained cells were analyzed on a FACSCalibur flow cytometry (BD Biosciences). Percentage of PMNs with condensed nuclei was assessed by nuclear morphology after staining with Hoechst 33342 (5 μg/ml for 20 min at ambient temperature) as described previously [21]. A total of 250 PMNs from at least 5 fields of view were scored for nuclear morphology consistent with apoptosis using a Zeiss Axioskop2 Plus fluorescence microscope (Carl Zeiss). This method does not distinguish live from dead cells, but separate experiments performed using identical conditions evaluated PMN lysis (dead cells). Representative images were obtained using an Axiocam HR digital camera and AxioVision version 4.4 software (Carl Zeiss).

**Live-Cell Imaging**

For live-cell imaging experiments, 6 × 10⁴ PMNs were aliquoted into serum-coated wells of a Lab-Tek® 8-chamber No. 1 borosilicate coverglass system (Thermo Fisher Scientific) and allowed to adhere at room temperature for 30 min. Serum-opsonized bacteria (6 × 10⁴) were labeled with Alexa 568 (10 μM final concentration) for 30 min at 37°C, washed twice with DPBS and resuspended at the desired concentration in RPMI/H. Following addition of bacteria, wells were filled with RPMI/H and phagocytosis was synchronized as described above. Chambers were sealed using a coverslip and vacuum grease and samples were kept on ice until the start of the experiment.

Live-cell imaging was performed on a Zeiss model LSM 510 Meta NLO laser scanning confocal system equipped with an Axiovert 200M microscope, an environmental chamber and AIM® version 4.2 SPI imaging software (Carl Zeiss). The environmental chamber and enclosed microscope parts were pre-equilibrated to 37°C for at least 3 h prior to imaging. Samples were imaged with a 40× 0.12 NA water-immersion objective lens (Carl Zeiss). Images were collected at 1-min intervals for up to 6 h. For most experiments, z-stacks were collected at each time point. Samples were visualized using differential interference contrast microscopy coupled with a 543-nm HeNe laser (to visualize labeled bacteria). Images were adjusted for optimal brightness and contrast in Adobe PhotoShop CS3 (Adobe Systems).

**Transmission Electron Microscopy**

For TEM, PMNs (3 × 10⁵) were aliquoted into wells of a 24-well tissue culture plate containing serum-coated Thermaxon coverslips (Thermo Fisher Scientific) and allowed to adhere at room temperature for 20 min. Cells were chilled on ice for 10 min and 3 × 10⁶ serum-opsonized S. aureus were added to the desired wells and plates were centrifuged as described above to synchronize phagocytosis. Assays were incubated at 37°C with 5% CO₂ for up to 6 h. At regular intervals, samples were fixed by immersion in a mixture of 4% glutaraldehyde, 4% paraformaldehyde and 0.1 M sodium phosphate buffer pH 7.2 (Electron Microscopy Sciences), and kept overnight at 4°C. Subsequent steps were performed using a Model 3451 microwave processor equipped with a vacuum chamber at approximately 250 Torr; a ColdSpot® microwave attenuator and a load cooler (Ted Pella). Vacuum was applied to all steps except solvent dehydration. Rinses and dehydration steps were performed for 1 min at 80 W. Postfixation and staining were conducted at 80 W with cycles of 2 min on, 2 min off and 2 min on. Resin infiltration steps were performed for 3 min at 240 W. The chamber temperature ranged from 24 to 32°C during processing. Following primary fixation as above, the samples were rinsed in phosphate buffer, treated with two post-fixation cycles using 1% osmium tetroxide in phosphate buffer, rinsed twice with water, stained with 1% uranyl acetate, rinsed again with water, dehydrated with ethanol, and infiltrated and embedded in Spurr’s resin. Sections were examined at 80 kV on a model H7500 transmission electron microscope (Hitachi High Technologies America). Digital images were captured using a model XR-100 CCD camera (Advanced Microscopy Techniques), and adjusted for brightness and contrast using Adobe PhotoShop CS3 (Adobe Systems).

**Assays for PMN ROS**

PMNs were incubated with 25 μM 2',7'-dihydrodichlorofluorescein diacetate (DCF; Invitrogen) for 20–30 min at room temperature in RPMI/H. DCF-containing PMNs (10⁶) and serum-opsonized S. aureus were combined in wells of a 96-well microtiter plate, centrifuged to synchronize phagocytosis as described above, and transferred to a microplate fluorometer (Spectramax Gemini; Molecular Devices). ROS production was measured as described previously [5, 19].

**RNA Preparation and Analysis of Gene Expression**

For microarray experiments, PMNs (1 × 10⁶) were combined on ice with 1 × 10⁶ serum-opsonized LAC, MW2, MnCop, COL, MRSA252 or IgG/C3bi-LB in wells of a 12-well tissue culture plate.
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and phagocytosis was synchronized as described above. Following centrifugation, plates were incubated at 37°C in a CO2 incubator for up to 6 h. At the designated times, the supernatant was removed and PMNs were lysed directly using RLT buffer (Qiagen). Purification of PMN RNA and subsequent preparation of labeled cRNA target (12 μg) was performed as described previously [19]. Labeling of samples, hybridization of cRNA to HU133 Plus 2.0 oligonucleotide arrays (Affymetrix), and scanning were performed according to Affymetrix protocols (http://www.affymetrix.com/pdf/expression_manual.pdf) as described. Experiments were performed with 4 separate donors at each time point using a separate oligonucleotide array for each donor.

Microarray analysis was performed as described [22] but with modifications. In addition to quality filters described previously [22] (combined calls and signal intensity), to be included in the final gene list transcript levels must have been changed at least 2-fold for one of the treatments and the fold-change must have been considered significant by p value obtained from a three-way ANOVA. Genes with a test/control ratio that resulted in a control 2-fold for one of the treatments and the fold-change must have been considered significant by p value obtained from ANOVA and Dunnett’s posttest or paired t test (GraphPad Prism 5; GraphPad Software).

Results

Lysis of Human PMNs during S. aureus Interaction Occurs after Phagocytosis

Survival of S. aureus after neutrophil phagocytosis is likely an important component of pathogenesis. To determine whether survival of S. aureus after phagocytosis correlates precisely with eventual PMN lysis, we evaluated these phenomena using previously described USA300 clinical isolates, including several USA300 epidemic clones [12]. There was varied survival of USA300 isolates after PMN phagocytosis (range is approx. 30–70% survival), even among closely related organisms (e.g. LAC and 18810 differ by 23 single nucleotide polymorphisms) (fig. 1a). Although there was not necessarily a precise correlation between percent S. aureus survival and PMN lysis, survival of USA300 isolates after phagocytosis resulted ultimately in the destruction of human neutrophils (fig. 1b). By contrast, permeabilization of the plasma membrane by culture supernatants of these isolates, that is, permeabilization that should be mediated by secreted cytolytic toxins, did not always correlate with PMN lysis after phagocytosis (fig. 1c). For example, culture supernatants from isolates 18805 and 18811 had little or no capacity to permeabilize plasma membranes of human neutrophils (pore formation was 2.4 ± 0.8 and 10.2 ± 4.1% at 30 min for these isolates vs. 3.5 ± 0.6% using YCP culture media alone), yet these isolates caused sig-
nificant PMN lysis after phagocytosis (42.2 ± 6.0 and 48.8 ± 6.1% at 6 h, respectively) (fig. 1b, c). Previous studies demonstrated that isolate 18811 has a defect in accessory gene regulator and thus fails to secrete α- or δ-hemolysin [12]. Taken together, these data suggest that neutrophil destruction after phagocytosis of *S. aureus* is at least in part independent of molecules typically secreted into culture medium.

PMN lysis occurred at a relatively low multiplicity of infection (for example, lysis at 6 h was 27.6 ± 4.1% using...
5 bacteria per PMN) and, as reported previously [5], there was a significant difference between LAC and strain COL, one of the earliest hospital MRSA isolates, to cause neutrophil destruction within 6 h after phagocytosis (fig. 2a). Although these S. aureus strains have relatively similar DNA content in the core genome [11, 12], there are differences in mobile genetic elements, including presence of a prophage in LAC that encodes Panton-Valentine leukocidin (PVL). Since bacterial culture media can affect the level of secreted toxins produced by S. aureus in vitro, we tested whether bacterial culture media impacts the ability of S. aureus to cause PMN lysis. Destruction of PMNs using LAC initially cultured in CCY or YCP media, which promote production and secretion of high lev-

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**Fig. 3.** Analysis of USA300-mediated PMN lysis by microscopy. Phagocytosis of USA300 and subsequent lysis of human neutrophils were monitored by confocal laser scanning microscopy over a 6-hour time period. Black arrows indicate a neutrophil that has phagocytosed USA300. n = Nucleus.
ponents of PVL [23], was similar to that caused by bacteria grown in TSB (fig. 2b). This observation is consistent with our current and previous findings using wild-type and isogenic PVL-negative USA300 strains that demonstrate PMN lysis after phagocytosis is PVL independent [20] (fig. 2c). Notably, lysis of human neutrophils required live S. aureus (fig. 2b), was caused largely by internalized bacteria (74.7% of the total) rather than organisms potentially remaining in the extracellular milieu or a toxin secreted by external bacteria (7.8% of the total lysis) (fig. 2d), and was significantly increased in PMNs that had phagocytosed serum-opsonized LAC (fig. 2e).

**Live-Cell Imaging of S. aureus-Induced PMN Lysis**

We next used confocal laser scanning microscopy to perform real-time imaging of S. aureus-induced PMN lysis (fig. 3). In accordance with our previous findings [5], uptake of USA300 was rapid and within 30 min most bact-
**Fig. 5.** USA300 induces changes in nuclear morphology consistent with PMN apoptosis.  

(a) PMN apoptosis was determined by nuclear morphology as described in Methods. Serum-opsonized LAC was cultured with human PMNs at a ratio of 10 bacteria per PMN. Alternatively, PMNs were cultured with 500 ng/ml anti-FAS antibody (αFAS Ab) as indicated.  

(b) Images from a representative experiment. Arrows indicate cells with condensed nuclei consistent with PMNs undergoing apoptosis.  

PMN lysis 6 h after phagocytosis of LAC or IgG/C3bi-LB, or addition of anti-FAS antibody (αFAS Ab) as indicated.  

d Caspase inhibitors were combined with PMNs 30 min before addition of *S. aureus* strains LAC or COL, and PMN lysis was measured at the indicated times.  

e Pepstatin A was combined with PMNs 30 min prior to addition of USA300 strain LAC (10 bacteria per PMN) and LDH release was measured as described in Methods. *p < 0.05 vs. PMNs alone (white bars) using a paired t test (2 and 4 h) or one-way ANOVA and Dunnett’s post-test (6 h). Results are the mean ± standard deviation of 6 (a), 4 (c), 3–7 (d), or 3 (e) PMN donors.
bacteria (>90%) were phagocytosed. Approximately 3–4 h after phagocytosis, PMNs underwent significant morphological change characterized by blebbing of the plasma membrane (see cells at 2 h 56 min in fig. 3; online suppl. video; for all supplementary material, see www.karger.com/doi/10.1159/000317134). Immediately following membrane blebbing, PMNs became quiescent and the nucleus typically coalesced and/or condensed in a manner consistent with PMN apoptosis (4 h 17 min; fig. 3). All PMNs undergoing this process lysed within 2 h after nuclear condensation, thus releasing any surviving bacteria (see 4 h 53 min to 5 h 27 min in fig. 3; online suppl. video). Initial morphological changes of the nucleus are consistent with those occurring during PMN apoptosis and apoptotic neutrophils will ultimately lyse after extended time in culture (approx. 48 h). However, the observation that PMNs lysed within 6 h after uptake of USA300 indicates death occurs by necrosis, albeit this process could be secondary to rapid apoptosis.

Does PMN Lysis following Phagocytosis of USA300 Involve Antecedent Apoptosis?

Inasmuch as the live-cell imaging suggested PMNs undergo apoptosis prior to lysis, we measured surface expression of annexin-V-FITC and nuclear condensation in PMNs following phagocytosis of USA300 as a step toward determining whether apoptosis precedes and/or accompanies neutrophil lysis. Although PMN annexin-V-FITC staining was relatively high (43.0 ± 3.8%) within 15 min after start of the assay, we cannot exclude the possibility that annexin-V-FITC bound nonspecifically to bacteria (fig. 4a). In contrast to PMNs cultured with anti-FAS antibody, which stained only with annexin-V-FITC, those stimulated with LAC were permeabilized rapidly and stained also with propidium iodide (fig. 4a–c).

In accordance with the live-cell imaging data, 68.3 ± 24.9% of PMNs remaining intact at 6 h had morphological consistent with cells undergoing apoptosis, and morphology was essentially identical to that of cells undergoing FAS-mediated apoptosis (fig. 4a, b). However, there was little or no lysis of PMNs at this time point during FAS-mediated apoptosis or following phagocytosis of IgG/C3Bi-LB, which has been shown previously to cause approximately 30–40% neutrophil apoptosis at 6 h (fig. 4b) [17, 24, 25]. Collectively, these data suggest that PMN lysis following phagocytosis of USA300 is not directly linked to apoptosis.

To determine whether caspases contribute to S. aureus-induced PMN lysis, we tested the ability of Z-VAD (irreversible inhibitor of caspases 1, 3, 4 and 7) and Y-VAD (irreversible inhibitor of caspases 1, 4 and 5) to inhibit PMN lysis occurring after phagocytosis of LAC or COL (fig. 5d). Neither Z-VAD (30 μM) nor Y-VAD (250 μM) inhibited PMN lysis at concentrations determined empirically to block FAS-induced apoptosis (fig. 5d; data not shown). Higher concentrations of Z-VAD (up to 300 μM) caused minimal inhibition of PMN lysis at 6 h, indicating caspases are not involved in this cell death process (data not shown). These findings provide additional support to the notion that the PMN lysis reported herein is not linked to apoptosis, since previous studies by Zhang et al. [26] demonstrated that phagocytosis-induced cell death is caspase-dependent. Pepstatin A, an inhibitor of cathepsin D, which has been shown

**Fig. 6.** Phagocytosis-induced PMN lysis is not dependent on ROS. a Live or heat-killed LAC elicit robust production of PMN ROS. b Lysis of PMNs from chronic granulomatous disease (CGD) patients or healthy control donors after phagocytosis of USA300. Results in panel a are the mean of 6 PMN donors. Results in panel b are the mean ± standard deviation of 4 healthy and 5 chronic granulomatous disease PMN donors. n.s. = Not significant.
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recently to initiate PMN apoptosis through activation of caspase 8 [27], also failed to block *S. aureus*-induced PMN lysis (fig. 5e).

**PMN Lysis after Phagocytosis of USA300 Is NADPH Oxidase Independent**

NADPH oxidase-derived ROS are required for phagocytosis-induced apoptosis [24, 28], a process suggested by data in figure 5 to be distinct from the rapid lysis caused by phagocytosis of USA300. If ROS are sufficient for *S. aureus*-induced PMN lysis, one would predict heat-killed LAC, which fails to cause PMN lysis (fig. 2b), would correspondingly fail to elicit production of ROS. However, heat-killed LAC elicited robust production of PMN ROS, thus indicating that ROS are not sufficient to cause lysis after uptake of *S. aureus* (fig. 6a). More notably, lysis of PMNs from patients with chronic granulomatous disease was similar to that of healthy control donors (73.9 ± 8.2% vs. 72.6 ± 6.4%; fig. 6b). Taken together, these observations demonstrate that ROS generated after phagocytosis of *S. aureus* do not contribute to PMN lysis.

**Ultrastructural Analysis of PMNs after Phagocytosis of USA300 Indicates Phagosomes Remain Intact until Cell Lysis**

We next performed transmission electron microscopy to elucidate whether USA300 phagosome membranes remain intact or if the pathogen produces molecules that compromise phagosome integrity and allow bacteria to escape into the cytoplasm to cause cell lysis (fig. 7). Early phagosomes were typically tight with little space between bacteria and phagosome membrane (0.5 h; fig. 7). By comparison, mature phagosomes (2 h and more after phagocytosis) were often spacious (see arrows at 3.5 and 5.5 h in fig. 7). Notably, phagosome membranes remained intact until the point of neutrophil lysis. In addition, there was little or no evidence for bacterial replication until after PMNs had lysed (see 6 h in fig. 7), an observation consistent with the kinetics of bacteriostatic activity toward USA300 after phagocytosis by neutrophils.

**Microarray Analysis Reveals Processes Potentially Linked to *S. aureus*-Induced PMN Death**

To better understand the molecular processes involved in *S. aureus*-induced PMN lysis, we used human oligonucleotide microarrays to measure changes in PMN gene expression that precede and/or accompany cell death. We compared PMN gene expression following phagocytosis of representative CA-MRSA (LAC and MW2), CA-MSSA (MnCop, a USA400 strain) and HA-MRSA (COL and MRSA252) strains. We reported previously that MW2 and MnCop each cause PMN lysis at levels comparable to that of LAC and significantly more than either COL or MRSA252 [5]. Therefore, changes in transcript levels that accompany and/or are linked to PMN lysis should in the-
ory be similar among the three community-associated *S. aureus* strains, but distinct from the two HA-MRSA strains.

Phagocytosis of *S. aureus* or IgG/C3bi-LB caused changes in gene expression either common to all phagocytic stimuli or all *S. aureus* strains for up to 6 h after ingestion (fig. 9). These findings are consistent with our previous studies demonstrating that phagocytosis in general triggers an apoptosis differentiation program in human neutrophils [17, 19, 29]. Although many genes were similarly differentially expressed following phagocytosis of CA-MRSA or HA-MRSA strains, there was a subset of transcripts that changed significantly only in neutrophils that had phagocytosed LAC, MW2 and/or MnCop (fig. 10). Principal component analysis of the microarray data indicated that PMN transcript levels in cells activated by community-associated *S. aureus* clearly segregated from those activated by COL or MRSA252 at 2 h after phagocytosis (fig. 10a). This observation is compatible with the finding that transcription and translation inhibitors block PMN lysis when added within the first 2 h after phagocytosis (fig. 8b). At this time point, there were approximately 110 transcripts significantly up- or downregulated only in PMNs activated by phagocytosis of community-associated *S. aureus* strains (fig. 10; online suppl. table). Sixteen transcripts encoding molecules involved in gene transcription or repression were differentially expressed, including *ETS2*, which encodes a transcription factor that regulates multiple genes that participate in cell death and senescence (fig. 10b). Five transcripts encoding calcium, potassium or proton channels, such as *ATPSF1*, *ATPI1B*, *CACNA1A*, *TRPV5* and *ENSA*, were upregulated only in cells that had ingested community-associated *S. aureus* strains (fig. 10b). Although it is tempting to speculate that cell osmolarity was negatively impacted and thereby resulted in disruption of the plasma membrane, there was no evidence of gradual swelling typical of changes in osmolarity (fig. 3). It will be important in future work to elucidate the possible involvement of these molecules in *S. aureus*-mediated neutrophil destruction following phagocytosis.

**Discussion**

Although progress has been made, the basis for the enhanced virulence phenotype of CA-MRSA is incompletely defined. We previously demonstrated that prominent CA-MRSA strains have significant capacity to survive after phagocytosis by neutrophils and cause eventual host cell lysis [5]. Thus, the ability of USA300 and USA400 strains to cause relatively rapid destruction of neutrophils after phagocytosis may be linked in part to the observed high-virulence phenotype of CA-MRSA. As such, we investigated the mechanism used by USA300 to cause neutrophil destruction after phagocytosis.
Fig. 9. PMN genes differentially expressed following phagocytosis of S. aureus and/or IgG/C3bi-LB. Microarray results are presented as the mean fold change of transcripts significantly increased or decreased compared with unstimulated cells from 4 separate PMN donors. For transcripts in the category 'apoptosis and cell fate', green arrows, black X and the yellow diamond indicate the encoded protein promotes apoptosis, inhibits apoptosis and causes lysis, respectively. Gene names are provided in italics and common names are provided in parentheses where indicated.
CA-MRSA strains such as USA300 secrete multiple cytolytic toxins, including PVL, γ-hemolysin and α-type phenol-soluble modulins, which have cytolytic activity toward neutrophils [30–32]. However, the neutrophil lysis described herein appears distinct from the mechanism used by pore-forming toxins such as PVL to kill neutrophils. First, PMN lysis after phagocytosis did not always correlate with permeabilization of the neutrophil plasma membrane by molecules present in USA300 culture supernatants (fig. 1b, c). For example, culture supernatant from USA300 isolate 18811 had limited capacity to cause permeabilization of the neutrophil plasma but retained capacity to cause lysis after phagocytosis. Also, pore-forming toxins produced within phagosomes would need access to the plasma membrane and our TEM analyses suggest phagosome membranes remained intact to the point of host cell lysis (fig. 7). This finding is not compatible with cytolysis that requires disruption of the phagosome membrane by pore-forming toxins. In accordance with this idea, we demonstrated previously that PMN lysis after phagocytosis of USA300 or USA400 is PVL independent [20]. Further studies with additional isogenic
gene deletion strains are needed to determine whether cytolytic toxins other than PVL contribute to the PMN lysis described herein.

Another notable finding was that lysis of PMNs was significantly increased following uptake of serum opsonized of LAC compared with unopsonized bacteria (fig. 2e). Although differential kinetics of phagocytosis may account for some of the observed differences in PMN lysis, it is also possible that ligation of specific receptors during opsonophagocytosis of USA300 ultimately contributes to PMN lysis. This observation merits further investigation.

Within 4 h after phagocytosis of USA300, neutrophils underwent morphological changes consistent with apoptosis, including plasma membrane blebbing and nuclear condensation (fig. 3, 5; online suppl. video). However, neither FAS-mediated nor phagocytosis-induced apoptosis caused significant neutrophil lysis within the same time period (fig. 5b). Therefore, although phagocytosis-induced apoptosis may be occurring in PMNs containing ingested USA300, the rapid neutrophil lysis is likely unrelated to this process.

Inasmuch as PMN lysis occurred following phagocytosis of strains that produce little or no cytolytic activity in culture supernatants (cytolysis mediated by pore-forming toxins) and required new transcription and protein synthesis within the first 3 h after uptake, the process appears to be a form of programmed cell death. Pyroptosis is a type of programmed cell death that culminates in cell lysis, inflammation of the surrounding tissue, and phagocyte recruitment to the site of infection [33–35]. This process is caspase 1 dependent and involves fragmentation of DNA, activation of IL-1β and IL-18, and formation of pores in the plasma membrane [36, 37]. Pyroptosis is mediated by the inflammasome, which assembles after phagocytosed bacteria escape from phagosomes and bacterial products are detected by pattern-recognition receptors in the cytosol [35, 38]. Since USA300 failed to escape from neutrophil phagosomes in our current studies and PMN lysis was caspase independent (fig. 5d), S. aureus-mediated PMN destruction reported here was not pyroptosis.

Our data [current study, 5, 6] indicate S. aureus causes a form of neutrophil necrosis, albeit there are clearly strain-dependent differences in the magnitude of lysis (for example, lysis was high for USA300 and low for strain COL). This notion is strongly supported by the recommendations of the Nomenclature Committee on Cell Death in 2009, as the characteristics of neutrophil death after phagocytosis of S. aureus fit those described for necrosis [39]. Vandenabeele and colleagues [40, 41] recently provided compelling evidence that cell necrosis can be a form of programmed cell death involving multiple signal transduction pathways. Programmed necrotic cell death may involve alteration of calcium ion levels [41], findings compatible with our results. Although we did not measure alterations in intracellular calcium levels during S. aureus-mediated PMN lysis, transcripts encoding proteins involved in calcium homeostasis (CACA1A, TRPV5 and CALR) were significantly upregulated in PMNs 2 h following phagocytosis of community-associated S. aureus strains (fig. 10b). Further studies are required to elucidate the signal transduction pathways underlying necrotic neutrophil death after ingestion of S. aureus. Such studies will provide insight into the basis of enhanced PMN necrosis caused by CA-MRSA strains and thus lead to a better understanding of the enhanced virulence phenotype of CA-MRSA.

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