Staphylococcus aureus Releases Proinflammatory Membrane Vesicles To Resist Antimicrobial Fatty Acids

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ABSTRACT Staphylococcus aureus is a major pathogen, which colonizes one in three otherwise healthy humans. This significant spread of S. aureus is largely due to its ability to circumvent innate immune responses, including antimicrobial fatty acids (AFAs) on the skin and in nasal secretions. In response to AFAs, S. aureus swiftly induces resistance mechanisms, which have yet to be completely elucidated. Here, we identify membrane vesicle (MV) release as a resistance strategy used by S. aureus to sequester host-specific AFAs. MVS protect S. aureus against a wide array of AFAs. Strikingly, beside MV production, S. aureus modulates MV composition upon exposure to AFAs. MVS purified from bacteria grown in the presence of linoleic acid display a distinct protein content and are enriched in lipoproteins, which strongly activate Toll-like receptor 2 (TLR2). Cumulatively, our findings reveal the protective capacities of MVS against AFAs, which are counteracted by an increased TLR2-mediated innate immune response.

IMPORTANCE The nares of one in three humans are colonized by Staphylococcus aureus. In these environments, and arguably on all mucosal surfaces, bacteria encounter fatty acids with antimicrobial properties. Our study uncovers that S. aureus releases membrane vesicles (MVS) that act as decoys to protect the bacterium against antimicrobial fatty acids (AFAs). The AFA-neutralizing effects of MVS were neither strain specific nor restricted to one particular AFA. Hence, MVS may represent “public goods” playing an overlooked role in shaping bacterial communities in AFA-rich environments such as the skin and nose. Intriguingly, in addition to MV biogenesis, S. aureus modulates MV composition in response to exposure to AFAs, including an increased release of lipoproteins. These MVS strongly stimulate the innate immunity via Toll-like receptor 2 (TLR2). TLR2-mediated inflammation, which helps to fight infections, may exacerbate inflammatory disorders like atopic dermatitis. Our study highlights intricate immune responses preventing infections from colonizing bacteria.

KEYWORDS Staphylococcus aureus, Toll-like receptors, antimicrobial fatty acids, lipoproteins, membrane vesicles

Staphylococcus aureus is a Gram-positive bacterium and the causative agent of numerous infections ranging from mild skin and soft tissue infections to invasive infections, such as bacteremia, endocarditis, and pneumonia (1). The morbidity, mortality, and health costs of these infections are exacerbated by the high prevalence of multidrug-resistant strains (2). In contrast, S. aureus colonizes asymptptomatically the nares of ~30% of the human population (3). For this bacterium, the skin carriage differs sharply between healthy individuals (5 to 20%) and patients with skin disorders such as atopic dermatitis (80 to 100%) (4). Both skin and nasal environments are rich in long-chain unsaturated fatty acids with antimicrobial properties (5, 6). These antimicrobial fatty acids (AFAs) also contribute to an important defense mechanism against
pathogens by maintaining the low pH of the skin, whose alkalization correlates with microbial dysbiosis, increased colonization with \textit{S. aureus}, and atopic dermatitis (7, 8).

In mice, topical application, intraperitoneal injection, and AFA-rich diet lead to decreased bacterial load and increased survival upon \textit{S. aureus} infections (9, 10). AFAs do not inhibit only \textit{S. aureus} and numerous Gram-positive species but also Gram-negative bacteria (11, 12). However, the role of AFAs in the innate immune response to bacterial infections goes beyond direct toxicity. Indeed, AFAs also possess immune-modulatory properties. For instance, upon incubation with sebum AFAs, human sebocytes considerably enhance their expression and secretion of beta-defensin 2, one of the predominant antimicrobial peptides found in the skin (13). Neutrophil release of the antimicrobial peptide LL-37 and alpha-defensins is also stimulated by AFAs (14). Furthermore, AFA incorporation into \textit{S. aureus} lipoproteins potentiates TLR2 (Toll-like receptor 2)-dependent innate immune activation (15).

The \textit{S. aureus} membrane is the primary target of AFAs, and their effects include increased fluidity, compromised integrity, and depolarization (16, 17). Recently, arachidonic acid, a polyunsaturated AFA, was shown to kill \textit{S. aureus} via lipid peroxidation (18). In response to the pleiotropy of AFA toxicity, \textit{S. aureus} has developed numerous resistance strategies. For instance, under iron-limiting conditions, the surface protein IsdA increases \textit{S. aureus} cellular hydrophilicity, precluding the bacterial binding to hydrophobic AFAs (9). Owing to similar properties, wall teichoic acids shield \textit{S. aureus} against AFAs (19). However, efflux pumps FarE (20) and Tet38 (21) prevent the cellular accumulation of AFAs, which are still able to bind to \textit{S. aureus}. Additionally, this bacterium possesses a functional oleate hydratase, which hydrates and thereby detoxifies AFAs containing cis-9 double bonds (22).

Strikingly, \textit{S. aureus} grown for a few hours in the presence of subinhibitory amounts of AFAs survives subsequent exposures to otherwise bactericidal AFA concentrations (20, 23), suggesting that the bacteria activate an AFA stress response program. High-throughput transcriptomic and proteomic studies on \textit{S. aureus} primed with AFAs revealed more than 100 differentially expressed genes (17, 23–25) but could not identify an inducible AFA resistance mechanism common to all \textit{S. aureus} strains and efficient against various AFAs. However, \textit{S. aureus} primed with several AFAs secretes distinct proteins, including triacylglycerol lipase 2 and several proteases (23, 26). Intriguingly, the contribution of secreted factors to \textit{S. aureus} resistance has not been thoroughly investigated. In addition to oleate hydratase, we sought to identify further factors released by \textit{S. aureus} to neutralize AFAs.

Here, using a clickable AFA analogue, we show that \textit{S. aureus}-conditioned medium sequesters AFAs and prevents their binding to the bacteria. Furthermore, we characterize the AFA-binding capacity of \textit{S. aureus} membrane vesicles (MVs), which enable the bacteria to grow in the presence of otherwise toxic amounts of AFAs. In response to AFAs, \textit{S. aureus} modulates its MV production and composition. MVs released in the presence of linoleic acid (LA) are enriched in lipoproteins and induce a potent TLR2 stimulation. Thus, the protective effects of MVs against AFAs are counteracted by a stronger innate immune response.

**RESULTS**

\textbf{\textit{S. aureus} release decoys that reduce AFA bacterial binding.} To gain new insights into \textit{S. aureus} interaction with AFAs, we used a LA analogue, linoleic acid alkyne (Fig. 1A), and click chemistry with azide fluor 488 for AFA-binding studies. Importantly, LA retained its capacity to inhibit \textit{S. aureus} growth upon addition of the alkyne group compared to LA (see Fig. S1A and B in the supplemental material). LA alkyne binding to \textit{S. aureus} USA300 LAC and Newman strains, which are resistant and sensitive to methicillin, respectively, could be readily quantified by flow cytometry (Fig. 1B). Strikingly, bacteria stained with LA alkyne in the presence of \textit{S. aureus}-conditioned culture supernatants exhibited markedly decreased signals compared to bacteria resuspended in fresh medium, suggesting that \textit{S. aureus} releases a secreted factor to its culture supernatant that sequesters LA or interferes otherwise with LA binding. We extended
our LA-binding studies to two other S. aureus strains: USA400 MW2 (community-acquired methicillin-resistant S. aureus [MRSA]) and SH1000 (methicillin-susceptible S. aureus [MSSA]). Clearly, a strong decrease in LA staining upon click chemistry in the presence of bacterial supernatants was noticed for both strains compared to fresh medium (Fig. S2). Taken together, our data suggest that S. aureus impedes binding by AFAs by a new strategy involving a secreted factor.

MVs promote S. aureus growth in the presence of AFAs. Recently, Andreoni and coworkers showed that the release of MVs helps S. aureus to survive exposure to daptomycin, a membrane-targeting antibiotic (27). Our finding that S. aureus releases a secreted factor that prevents bacterial accumulation of membrane targeting, labeled LA (Fig. 1) raised the question whether MVs may be responsible for sequestering AFAs in culture supernatants. First, we isolated MVs from MSSA and MRSA. S. aureus Newman MVs had the highest protein content (Fig. 2A), while all S. aureus MVs had similar lipid amounts, as measured with the lipophilic dye FM4-64 (Fig. 2B). As exemplified with the Newman strain, S. aureus MVs were highly hydrophobic and able to bind LA alkyne within a few minutes (Fig. 2C and D).

AFA-binding capacity of MVs prompted us to test whether MVs promote S. aureus growth in the presence of toxic amounts of AFAs. The growth of S. aureus strain LAC was inhibited by 125 or 200 μM LA, but growth was not impeded when bacteria were supplemented with MVs from the same strain (Fig. 3A). Importantly, S. aureus LAC MVs were also able to support S. aureus Newman growth in the presence of inhibitory amounts (100 μM) of LA, as revealed by optical density monitoring (Fig. 3B). In addition to optical density, CFU counts showed that LA toxicity was alleviated by LAC MVs for

FIG 1 Culture supernatants impede S. aureus targeting by LA alkyne. (A) Chemical structure of alkyne functionalized linoleic acid (LA alkyne; 9Z,12Z-octadecadien-17-ynoic acid). (B) S. aureus bacteria grown for 6 h in TSB were incubated at 37°C for 20 min with or without LA alkyne prior to labeling with azide fluor 488 and flow cytometry analyses. Click chemistry was performed in the absence (-) or presence of culture supernatants (+ supe). Data shown are bacterial mean fluorescence intensities (MFI) plus standard errors of the means (SEM) (error bars) (n = 3). Values that are significantly different by one-way ANOVA with Tukey’s test are indicated by asterisks as follows: **, P < 0.01; ****, P < 0.0001.
and Newman (Fig. 3D) strains. Likewise, Newman MVs abrogated the LA-induced growth inhibition of both LAC and Newman strains (Fig. S3A and B).

To make sure that the protective effect of MV preparations against LA was not due to some residual components of the complex media used to grow bacteria prior to MV isolation, LAC MVs were purified from bacteria grown in four different broths (basic medium, lysogeny broth, Mueller-Hinton broth [MHB], or tryptic soy broth [TSB]). These four types of LAC MVs equally helped the LAC strain grow in the presence of otherwise inhibitory amounts of LA (Fig. S3C).

To elucidate whether the AFA-neutralizing capacity of MVs is specific for LA, we investigated whether MVs could enable S. aureus growth in the presence of other AFAs in addition to LA. Accordingly, the LAC strain was grown with 100 μM palmitoleic, sapienic, α-linolenic, or arachidonic acid. These AFAs were all able to inhibit bacterial growth, which resumed in the presence of LAC or Newman MVs (Fig. 3E). However, MVs did not completely shield bacteria against α-linolenic acid (Fig. 3E and Fig. S4). Collectively, our data strongly suggest MVs as a resistance mechanism against a broad range of different AFAs.

Linoleic acid boosts S. aureus release of MVs with strong TLR2-stimulating capacities. S. aureus resistance to AFAs is known to be induced by subinhibitory amounts of AFAs (20, 23), which is concomitant with an altered secretome (23, 26). Our discovery that MVs protect S. aureus against AFA toxicity prompted us to analyze...
bacterial MV release in the presence of LA. As the presence of membrane lipids is a hallmark for MVs, MVs from *S. aureus* grown in the presence of 0, 20, or 40 μM LA were stained with the lipophilic dye FM4-64, and the amount of lipids in MV preparations (surrogate for MV amount) was quantified with a plate reader. *S. aureus* strains USA300 LAC (Fig. 4A), USA400 MW2 (Fig. 4B), and SH1000 (Fig. 4C) responded to LA exposure with a significantly increased release (20 to 60%) of AFA-neutralizing MVs. Thus, *S. aureus* appears to produce MVs as inducible decoys for the sequestration of harmful AFAs.

Recently, we demonstrated that the release of *S. aureus* MVs was driven by surfactant-like small peptides, phenol-soluble modulins (PSMs), which are controlled by...
Importantly, Agr-deficient mutants, which are defective in MV biogenesis, are also more susceptible to AFAs (24). LAC Δagr significantly augmented its MV release (70 to 110%) in the presence of LA (Fig. 4D). However, irrespective of LA treatment, Δagr MVs were residual compared to those of the wild type (Fig. 4D). Although a detailed mechanism of AFA-triggered increase in MV release is lacking, it is apparent that AFAs only enhance preexisting bacterial capacity to vesiculate.

Given the pleiotropic effects of AFAs on *S. aureus*, which include increased membrane fluidity, altered proteome, and reduced surface hydrophobicity (16, 23, 24), we reasoned that these differences would be reflected in the composition of MVs released in the presence of AFAs. Accordingly, flow cytometry analysis of FM4-64-stained LAC MVs uncovered that MVs purified in the presence of LA displayed a twofold increase in lipid amounts (Fig. 5A). These MVs also had twofold-increased nucleic acid cargos (presumably RNA), as revealed by SYTO 9 staining (Fig. 5B). Taken together, these results indicate that *S. aureus* does not only increase MV production but also modulates MV composition in response to LA.

We further probed the altered composition of *S. aureus* MVs following bacterial exposure to LA by label-free proteomics. MVs purified from *S. aureus* USA300 LAC grown with no LA (control MVs) or 40 μM LA (LA-MVs) significantly differed in their protein content. We detected 414 and 442 proteins in control MVs and LA-MVs, respectively, of which 308 were common to both types of MVs, and roughly one in five proteins was detected exclusively in control MVs or LA-MVs (Fig. 6A). Furthermore,
compared to control MVs, more membrane proteins were identified in LA-MVs (Fig. S5A). Seven of these membrane proteins appeared to be lipoproteins, which were absent in control MVs (Fig. 6B). Quantitative analysis of protein abundance revealed that more than one in two proteins were differentially abundant (adjusted \( P \) value \( \leq 0.05 \) and \(|\text{fold change}| \geq 3\)) in LAC-MVs compared to control MVs (see Data Set S1 and Fig. S5B in the supplemental material). Strikingly, most of the detected lipoproteins were more abundant in LA-MVs (Fig. 6C). For the lipoprotein SitC (also referred to as MntC), the proteomic results were confirmed with control and LA-MVs isolated from a LAC strain expressing SitC with a C-terminally linked His tag. SitC-His was mildly but consistently more abundant in LA-MVs compared to control MVs (Fig. S6). These data collectively demonstrate that LA-MVs comprise an increased amount of lipoproteins.

As LA-MVs were enriched in TLR2-activating lipoproteins, we investigated their capacities to stimulate TLR2-transfected HEK293 cells (HEK-TLR2). In agreement with their lipoprotein content, LA-MVs had a substantially higher capacity to stimulate HEK-TLR2 cells compared to the same amounts of control MVs, as determined by interleukin 8 (IL-8) release in response to USA300 LAC MVs (Fig. 6D) and USA400 MW2 and SH1000 MVs (Fig. 6E). Taken together, these data reveal that the TLR2-activating capacities of MVs are exacerbated, when lipoprotein-rich MVs are released in the presence of subinhibitory amounts of AFAs.

**DISCUSSION**

Host-specific AFAs are important colonization barriers deployed by the innate immune system. Indeed, AFAs can inhibit growth or kill several opportunistic or pathogenic bacteria (12), including \( S. \) *aureus*. This opportunistic pathogen often colonizes the skin of patients with atopic dermatitis (AD), a chronic inflammatory skin disease predisposing to recurrent skin infections (7). Intriguingly, \( S. \) *aureus*-colonized AD patients have decreased amounts of antimicrobial sapienic and oleic acids (5, 29), suggesting that the reduced exposure of \( S. \) *aureus* to AFAs contributes to AD pathophysiology. The importance of AFAs at the host-pathogen interface is further demonstrated by the wide variety of resistance strategies used by \( S. \) *aureus* against AFAs (9, 18–21, 24). Moreover, subinhibitory amounts of AFAs induce increased resistance by mechanisms that have not been fully understood (20, 23). In the present study, we demonstrate that \( S. \) *aureus* responds to AFA exposure by boosting the release of MVs that protect against AFA toxicity. The MV-mediated resistance to AFAs was neither strain specific nor restricted to a limited set of AFAs. However, MVs released in response to AFAs provoked an increased TLR2-mediated immune response.

Mice immunized with \( S. \) *aureus* MVs are protected against otherwise lethal \( S. \) *aureus* lung infections in a TLR2-dependent manner (30). Beside this protective role, TLR2 activation is thought to contribute to the exacerbation and persistence of skin inflam-
mation during AD (31, 32). In keeping with this, *S. aureus* MVs can cause or worsen AD-like skin inflammation in mice (33–35). Therefore, it is enticing to speculate that reduced AFA concentrations in AD skin are too low to kill *S. aureus* but increase the release of TLR2 agonists, which further exacerbate skin inflammation.

Extracellular resistance mechanisms to antimicrobials have been reported for Gram-negative and Gram-positive bacteria and are collectively referred to as antibiotic
interceptors (36). For instance, a small protein known as lipocalin is released by *Burkholderia cenocepacia* to sequester hydrophobic antibiotics and enable bacterial growth in the presence of otherwise inhibitory concentrations of these drugs (37). Other protein interceptors are released as MV components and include β-lactamases from *Moraxella catarrhalis* and *S. aureus* (38, 39). Besides antimicrobial degradation, the outer MVs of Gram-negative bacteria are well characterized for their role as decoys for membrane-targeting agents like antimicrobial peptides (40–42). In contrast, not much is known about MV decoys from Gram-positive bacteria. Recently, MVs were shown to protect *S. aureus* against daptomycin, a membrane-targeting antibiotic (27). This bacterium releases membrane phospholipids and MVs in response to daptomycin (43). Importantly, daptomycin-induced lipid release by *S. aureus* is enhanced in the presence of AFAs (44), which is in agreement with our current findings that staphylococcal MV production is increased by AFAs. Notably, electron microscopic examinations of AFA-treated bacteria have consistently revealed abundant nanostructures reminiscent of MVs in Gram-positive staphylococci and streptococci (45, 46) as well as in Gram-negative *Porphyromonas gingivalis* and *Helicobacter pylori* (47, 48). Thus, MVs may represent “public goods” used by bacterial communities as a resistance mechanism against AFAs and lipophilic antibiotics such as daptomycin.

*S. aureus* responds to sublethal amounts of various AFAs by altering its secretome (23, 26). Interestingly, proteins secreted in response to AFAs are components of *S. aureus* MVs (28, 49). Our current data demonstrate that *S. aureus* indeed augments MV release in the presence of AFAs. The turgor pressure provides the energy for the budding of MVs (28), and this process is probably facilitated by altering membrane fluidity. In addition to AFAs, daptomycin has been reported to induce MV release in *S. aureus* (43). Furthermore, surfactant-like small peptides, phenol-soluble modulins (PSMs), have been shown to increase membrane fluidity of *S. aureus*, which favors MV budding (28). Similarly, it is likely that the fluidifying effect of AFAs on the *S. aureus* membrane (16, 17) would promote MV formation. Soaps and body lotions with surfactant-like properties may also promote MV release from skin bacteria and thereby increase TLR2 activation and inflammation with critical consequences in AD.

*S. aureus* oleate hydratase (OhyA) is another resistance strategy used by the bacterium to detoxify AFAs containing cis-9 double bonds (22). Intriguingly, our proteomic data revealed that OhyA was abundant in LA-MVs but absent in control MVs (see Data Set S1 in the supplemental material), suggesting that LA-MVs could detoxify AFAs not only by sequestration but also by inactivation. An additional link between MV and known anti-AFA defenses is provided by the effector and regulator of fatty acid resistance (FarE and FarR, respectively). FarE mediates AFA efflux under the control of its transcriptional regulator FarR (20). Constitutive activation of FarE confers increased resistance against AFAs and the membrane-targeting antibiotic rhodomyrtone (20, 50). Remarkably, high FarE levels also lead to an increased release of PSMs (50). Since PSMs are known to promote MV release (28, 51), we surmise an indirect contribution of FarE to this process. There is another precedent for an efflux pump-mediated lipid release in response to AFAs in *Acinetobacter baumannii* (11).

Besides their role as decoys for AFAs, *S. aureus* MVs are potent TLR2 activators by virtue of their lipoprotein cargos (28). Interestingly, mice respond to *S. aureus* skin infections by increasing their AFA production in a TLR2-dependent manner (52). Enhanced arachidonic acid blood levels have also been observed in mice nasally challenged with *Streptococcus pneumoniae* (53). *In vitro* studies have demonstrated an increased TLR2-mediated immune response to AFA-fed *S. aureus* (15). In light of our data, it seems likely that lipoprotein-enriched MVs released in the presence of AFAs enhanced TLR2 activation. It remains unclear why LA-MVs contain increased amounts of lipoproteins compared to control MVs. In line with our observation, *S. aureus* exposure to subinhibitory AFA concentrations has been found to increase the expression of several lipoproteins (23, 24). AFAs lead to upregulation of the virulence regulator SarA (23, 24), which was recently shown to control the expression of many lipoproteins (54). In keeping with this, SarA was more abundant in LA-MVs compared to the unchallenged control MVs (28, 51, 52).

Enhanced arachidonic acid blood levels have also been observed in mice nasally challenged with *S. pneumoniae* (53). *In vitro* studies have demonstrated an increased TLR2-mediated immune response to AFA-fed *S. aureus* (15). In light of our data, it seems likely that lipoprotein-enriched MVs released in the presence of AFAs enhanced TLR2 activation. It remains unclear why LA-MVs contain increased amounts of lipoproteins compared to control MVs. In line with our observation, *S. aureus* exposure to subinhibitory AFA concentrations has been found to increase the expression of several lipoproteins (23, 24). AFAs lead to upregulation of the virulence regulator SarA (23, 24), which was recently shown to control the expression of many lipoproteins (54). In keeping with this, SarA was more abundant in LA-MVs compared to the unchallenged control MVs (28, 51, 52).
to control MVs. It is unclear whether certain lipoproteins have a protective role against AFAs as some of them do against copper (55) or classical antibiotics (56–58).

In sum, our data support the idea that MV release is not restricted to MRSA. As MV-conferring resistance to AFAs entails no strain specificity, it is enticing to speculate that MVs represent a conserved yet flexible strategy that *S. aureus* uses against structurally unrelated, hydrophobic, antimicrobial compounds. It is appropriate that our TLR2-mediated immune response appears to better recognize AFA-exposed bacteria (15) and MVs, which may help to protect healthy skin, but may exacerbate skin inflammation in AD patients. Thus, AFAs are major components of the intricate host defenses, which represent untapped resources for new antimicrobial therapeutic interventions.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** *S. aureus* strains used are listed in Table S1 in the supplemental material and were routinely grown aerobically in tryptic soy broth (TSB) overnight at 37°C prior to each experiment unless stated otherwise.

**Membrane vesicle purification.** MVs were isolated with the ExoQuickTC kit (EQPL10TC; System Bioscience) as described elsewhere (28). Briefly, overnight bacterial cultures diluted to an optical density at 600 nm (OD$_{600}$) of 0.1 in 20 ml plain TSB or TSB supplemented with AFAs or their solvent dimethyl sulfoxide (DMSO) were grown with shaking for 6 h (late exponential growth phase). Next, bacteria were pelleted by centrifugation, and supernatants were sterile filtered. MVs in these culture filtrates were concentrated with 100-kDa centrifugal concentrator cartridges (Vivaspin 20; Sartorius) prior to precipitation with the ExoQuickTC kit and resuspension in 500 μl phosphate-buffered saline (PBS).

**Protein, lipid, and nucleic acid quantification in MVs.** The quantification of the protein fraction in purified MVs was performed using a Bradford assay following the manufacturer’s recommendations (Quick Start Bradford protein assay kit; Bio-Rad). For lipids, the lipophilic dye FM4-64 (Life Technologies) was used to quantify MVs. For nucleic acid, the nucleic acid cargo of MVs was measured via staining with 10 μM SYTO 9 for 30 min. Samples were analyzed with a CLARIOStar microplate reader (BMG Labtech) or a BD LSRFortessa flow cytometer (BD).

**SitC detection.** SitC expression in *S. aureus* USA300 LAC Δspc pTX SitC-His was induced with 0.5% xylose added to TSB without glucose supplemented with DMSO or linoleic acid. After bacterial growth, MVs were purified and their protein content was quantified as described above. Thirty micrograms (protein amount) per MV sample were stained with either a phycoerythrin (PE) anti-His tag antibody (clone J095G46; BioLegend) or its appropriate mouse IgG2a, κ PE isotype control antibody (clone MOPC-173; BioLegend). PE-labeled MVs were then analyzed by flow cytometry.

**Click chemistry with linoleic acid alkyne.** Exponentially growing bacteria were centrifuged and resuspended in either sterile, fresh TSB or *S. aureus*-conditioned medium. These bacteria or purified MVs were incubated at 37°C for 5 to 20 min with 20 μM linoleic acid alkyne (Cayman Chemical). Samples were then centrifuged, and pellets were resuspended in Click-IT cell reaction buffer supplemented with copper(II) sulfate and Click-IT cell buffer additive, as recommended by the manufacturer (Click-IT cell reaction buffer kit; Invitrogen). Click chemistry was performed at 25°C for 30 min with 7 μM azide fluor 488 (Merck). After washing with PBS, LA-stained bacteria or MVs were analyzed by flow cytometry.

**Growth curves.** Overnight bacterial cultures were diluted to an OD$_{600}$ of 0.05 in plain MHB or MHB supplemented with AFAs and/or 3.75 to 60 μg/ml MVs. Bacteria were then grown in a 96-well plate (U-bottom) at 37°C with linear shaking at 567 rpm (3-mm excursion) for 20 h, and the OD$_{600}$ was measured every 15 min with an Epoch 2 plate reader (BioTek). Areas under the curves were computed with GraphPad Prism 8.4.2.

**Growth inhibition assays.** After dilution to an OD$_{600}$ of 0.001 in MHB and treatment with DMSO (no LA), LA, or LA plus 30 μg/ml MVs, bacteria were either directly plated or grown for 2 h at 37°C before plating on tryptic soy agar and CFU counting.

**Quantitative label-free proteomics.** For proteomic analysis, MVs were isolated as described above from *S. aureus* USA300 LAC strain grown in TSB supplemented with DMSO (control MVs) or 40 μM LA (LA-MVs). After protein quantification with a Bradford assay, 20 μg of protein from each sample was subjected to tryptic in-gel digestion, samples were analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS) as described elsewhere (28, 59). Briefly, a Q Exactive HF Hybrid Quadrupole-Orbitrap mass spectrometer (Thermo Scientific) and a 90-min LC separation with an EASY-nLC 1200 system (Thermo Scientific) were employed. The data were used to interrogate the UniProt Staphylococcus aureus USA300 database UP000001939, and the common contaminant database from MaxQuant (60). Protein identification and quantification were performed with the MaxQuant software using default settings. Intensities were log$_2$ transformed with the Perseus software, and proteins with only one or no valid value for every sample in triplicate were filtered. Missing values were then put in as the lowest intensity across all samples in R. Differential protein abundance was calculated with the limma R package (61).

**HEK-TLR2 cell culture and stimulation.** HEK293 cells stably transfected with the human TLR2 gene (HEK-TLR2) (Invivogen) were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum (FCS), 100 μg/ml Normocin, and 10 μg/ml blastidicin. For stimulation experiments, HEK-TLR2 cells were seeded into 24-well plates (2 × 10^5 cells/well) and cultivated until confluent.
ence was reached (2 to 3 days). Next, the growth medium was removed, cells were washed once with PBS before incubation for 20 h with MVs diluted in 500 µl DMEM per well. Human IL-8 release was used as a proxy for TLR2 activation and measured using an enzyme-linked immunosorbent assay (ELISA) kit (R&D Systems) according to the manufacturer’s instructions. The synthetic lipopeptide Pam2CysK (200 ng/ml) was used as a positive control.

**Statistical analysis.** Except for the proteomics data, statistical tests specified in the figure legends were performed with GraphPad Prism 8.4.2, and values of $P < 0.05$ were considered significant. Analysis of variance (ANOVA) with Dunnett’s or Tukey’s multiple-comparison test was used. The fold changes and $P$ values of the proteomics data were calculated with the R package limma ([61](#)), with control MVs as the reference.

**Data availability.** The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE ([62](#)) partner repository with the data set identifier PXD018809, where control MVs are labeled R01, R02, and R03. All other data generated are available within the paper and the supplemental material files.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 0.2 MB.

**FIG S2**, TIF file, 0.2 MB.

**FIG S3**, TIF file, 1 MB.

**FIG S4**, TIF file, 1.8 MB.

**FIG S5**, TIF file, 0.9 MB.

**FIG S6**, TIF file, 0.6 MB.

**TABLE S1**, PDF file, 0.1 MB.

**DATA SET S1**, XLSX file, 0.1 MB.

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