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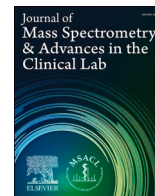
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Research Article

Imaging mass spectrometry reveals complex lipid distributions across *Staphylococcus aureus* biofilm layers

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ABSTRACT

Introduction: Although *Staphylococcus aureus* is the leading cause of biofilm-related infections, the lipidomic distributions within these biofilms is poorly understood. Here, lipidomic mapping of *S. aureus* biofilm cross-sections was performed to investigate heterogeneity between horizontal biofilm layers.

Methods: *S. aureus* biofilms were grown statically, embedded in a mixture of carboxymethylcellulose/gelatin, and prepared for downstream matrix-assisted laser desorption/ionization imaging mass spectrometry (MALDI IMS). Trapped ion mobility spectrometry (TIMS) was also applied prior to mass analysis.

Results: Implementation of TIMS led to a ~ threefold increase in the number of lipid species detected. Washing biofilm samples with ammonium formate (150 mM) increased signal intensity for some bacterial lipids by as much as tenfold, with minimal disruption of the biofilm structure. MALDI TIMS IMS revealed that most lipids localize primarily to a single biofilm layer, and species from the same lipid class such as cardiolipins CL(57:0) – CL(66:0) display starkly different localizations, exhibiting between 1.5 and 6.3-fold intensity differences between layers ($n = 3$, $p < 0.03$). No horizontal layers were observed within biofilms grown anaerobically, and lipids were distributed homogeneously.

Conclusions: High spatial resolution analysis of *S. aureus* biofilm cross-sections by MALDI TIMS IMS revealed stark lipidomic heterogeneity between horizontal *S. aureus* biofilm layers demonstrating that each layer was molecularly distinct. Finally, this workflow uncovered an absence of layers in biofilms grown under anaerobic conditions, possibly indicating that oxygen contributes to the observed heterogeneity under aerobic conditions. Future applications of this workflow to study spatially localized molecular responses to antimicrobials could provide new therapeutic strategies.

Abbreviations: CASI, continuous accumulation of selected ions; CMC, carboxymethylcellulose; CL, Cardiolipin; CFU, colony forming units; DAN, 1',5'-Diaminonaphthalene; DAPI, 4',6-diamidino-2-phenylindole; DG, diacylglycerol; DGDG, digalactosyldiacylglycerol; DHA, 2',5'-Dihydroxyacetophenone; DsRed, red fluorescent protein from *Discosoma*; EFG, electric field gradient; FWHM, full-width half max; GFP, green fluorescent protein; IMS, imaging mass spectrometry; ITO, indium-tin oxide; L-PG, lysyl-phosphatidylglycerol; MALDI, matrix-assisted laser desorption/ionization; OCT, optimal cutting temperature; PG, phosphatidylglycerol; PIP, phosphatidylinositol phosphate; PC, phosphatidylcholine; S/N, Signal-to-noise; TIMS, trapped ion mobility spectrometry; TSA, tryptic soy agar; TSB, tryptic soy broth; TIC, total ion current; XIM, extracted ion mobilogram.

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Introduction

Staphylococcus aureus is a Gram-positive opportunistic pathogen responsible for over 10,000 deaths and 320,000 hospitalizations per year in the United States and is a leading cause of healthcare-associated infections, such as endocarditis and bacteremia [1,2]. In addition, *S. aureus* is also the leading pathogen in biofilm-related infections, further confounding its burden on human health [3]. Biofilms are complex bacterial communities that adhere to a surface, such as surgical prosthetics or biological tissue, in infection [4,5]. Bacteria within these communities display heterogeneous behavior, and *S. aureus* biofilms can develop spatially discrete horizontal layers comprised of bacteria in different phenotypic states. Biofilm development leads to changes in characteristics compared to planktonic bacteria, including increased stress tolerance and altered gene regulation [6–9], and can involve changes in the bacterial lipidome [10]. Lipids play important roles in *S. aureus* ranging from membrane composition to evasion of the host immune response [11–13]. Bacterial metabolic processes, stress response, and virulence factor production can also be affected by changes in lipid composition [14,15]. The *S. aureus* lipidome consists primarily of phospholipids, including phosphatidylglycerols (PG), phosphatidylinositol phosphates (PIP), cardiolipins (CL), and lysyl-phosphatidylglycerols (L-PG) [14,16–19], each with distinct roles in bacterial functions. While the functions of these lipids are well described within an *S. aureus* bacterium, little is known about the roles they may play in larger bacterial colonies, such as an *S. aureus* biofilm. Determining where bacterial lipids localize within horizontal layers of *S. aureus* biofilms under basal conditions is necessary to provide insight into how these bacterial colonies respond to external factors, such as differences in oxygen availability.

Many previous studies have used lipid extracts from cultured bacteria to investigate the lipidome of *S. aureus* for research interests ranging from membrane rigidity to antibiotic resistance [16,18,20,21]. These approaches provide quantitation of the overall lipid content, but come at the expense of spatial information that is especially important in biofilm formation. Matrix-assisted laser desorption/ionization imaging mass spectrometry (MALDI IMS) [22] is a technology that is effective for mapping hundreds-to-thousands of molecules, including small metabolites [23,24], glycans [25–27], lipids [28,29], peptides [30,31] and intact proteins [32,33] in tissue at biologically-relevant spatial resolutions [22,34]. The ability to interrogate molecular distributions within bacterial communities by MALDI IMS has been previously demonstrated for a wide range of microbial molecules including phenol-soluble modulins [35–37], metabolites [35,38–45], oligosaccharides [46] and some lipids [40,41,46–48]. However, most of this work has been focused on analytes other than lipids, or has been performed with bacteria other than *S. aureus* or a co-culture of *S. aureus* with other microbes. Lipidomic investigations of monomicrobial *S. aureus* biofilms, however, would offer insight to the basal phenotypic state of bacterial lipid distributions. In addition, most prior examples of MALDI IMS of microbial molecules have been performed by sampling the top of the biofilm, investigating molecular distributions on the surface of the biofilm. A few examples of biofilm cross-sections have been demonstrated for the investigation of microbial metabolites within the cross-sectional depth of the biofilm [38,39], but none have investigated *S. aureus* lipid distributions within horizontal biofilm layers.

One challenge posed by microbial imaging is the small size of an individual bacterium. Most MALDI IMS studies of bacterial communities have been performed at spatial resolutions greater than 50 μm , with rare exceptions [47,45]. Because these *S. aureus* biofilm layers are approximately 100 μm thick, higher spatial resolutions are necessary to effectively investigate inter-layer variations. To this end, we sought to increase the spatial resolution of our lipidomic study (10 μm), which better enables visualization of these multicellular biofilm layers. While 10 μm spatial resolution is achievable, it is often at the expense of sensitivity. This limitation has been addressed via various enrichment

technologies, such as continuous accumulation of selected ions (CASI) [49] or MALDI-2 postionization [50,51]. However, the utility of each of these enrichment technologies can present new challenges, such as limited mass range and spatial resolution for ion accumulation technologies, or increased ion suppression by chemical interferences produced in the post-ionization process. Incorporation of ion mobility separation prior to mass analysis has also been demonstrated to increase sensitivity by reducing chemical noise and increasing peak capacity [52] and S/N [53,54] in an IMS experiment. Ion mobility utilizes opposing forces of an inert gas to direct analytes in one direction and a voltage gradient directing analytes in the opposite direction to achieve molecular separation as a function of the size-to-charge ratio of an analyte [57,58]. Numerous types of ion mobility have been integrated with IMS including drift-tube [59,60], travelling-wave [53], high-field asymmetric waveform [61], and we recently reported the integration of trapped ion mobility spectrometry (TIMS) with IMS [52,62]. Through the implementation of TIMS for increased sensitivity, here we perform 10 μm spatial resolution MALDI IMS of cross-sections for the investigation of lipids within distinct, horizontal *S. aureus* biofilm layers to uncover stark inter-layer heterogeneity at the lipidomic level.

Methods

Materials

2',5'-Dihydroxyacetophenone (DHA), 1',5'-Diaminonaphthalene (DAN), ammonium formate, ammonium acetate, and poly-L-lysine solution were purchased from Sigma-Aldrich Chemical Co. (St. Louis, MO, USA). HPLC-grade acetonitrile, methanol, and gelatin powder were purchased from Fisher Scientific (Pittsburgh, PA, USA). Carboxymethylcellulose (CMC) was purchased from PM MP Biomedicals (Salon, OH). Tryptic soy agar (TSA) and tryptic soy broth (TSB) were purchased from Becton Dickinson (Franklin Lakes, NJ) and all lab plasticware was from USA Scientific (Ocala, FL) or Corning (Corning, NY). Whatman Nucleopore polycarbonate filter discs (0.1 μm pore size) were acquired from Millipore Sigma (Burlington, MA).

Sample preparation

S. aureus bacterial cultures and biofilm growth. *S. aureus* strains USA300 LAC [73] or USA300 LAC modified to constitutively express GFP [74] were streaked on TSA and incubated at 37 °C. Single colonies were picked and used to inoculate 5 mL of TSB in 15 mL polypropylene tubes. Tubes were agitated for 16 hr at 37 °C. Twenty microliters of the resulting stationary phase culture were then pipetted onto filter discs on TSA plates and incubated at 37 °C for a total of 72 h. During this time, filter discs were transferred to fresh TSA plates every 24 h. Biofilm transfers were performed using antiseptic technique with disinfected forceps to grasp the edge of the filter disc, with care to maintain the biofilm in a horizontal orientation, and to not make contact with the biofilm. In order to measure *S. aureus* biofilm growth over time CFU counts were obtained every 24 h after inoculation. To measure CFUs at the appropriate time points individual filter discs containing biofilms were placed into 1.5 mL Navy Eppendorf Bead Lysis Kit tubes containing 1 mL of Phosphate Buffered Saline and homogenized at 4 °C for three successive 4 min cycles using a Bullet Blender Tissue Homogenizer (Next Advance, Inc., Troy, NY) on the maximum speed. Serial dilutions of the homogenates were plated on TSA to determine CFUs.

Anaerobic biofilms were grown by inoculating filter discs as described above and then placing the petri dishes containing the filter discs in a 37 °C incubator housed inside an anaerobic chamber (Coy Lab Products, Inc., Grass Lake, MI) filled with a mixture of 90 % nitrogen, 5 % hydrogen, 5 % carbon dioxide. Filter discs were transferred every 24 h to fresh TSA plates that had been allowed to equilibrate to the anaerobic chamber for at least 24 h prior to transfer.

After 72 h growth, *S. aureus* biofilms were embedded in a CMC/

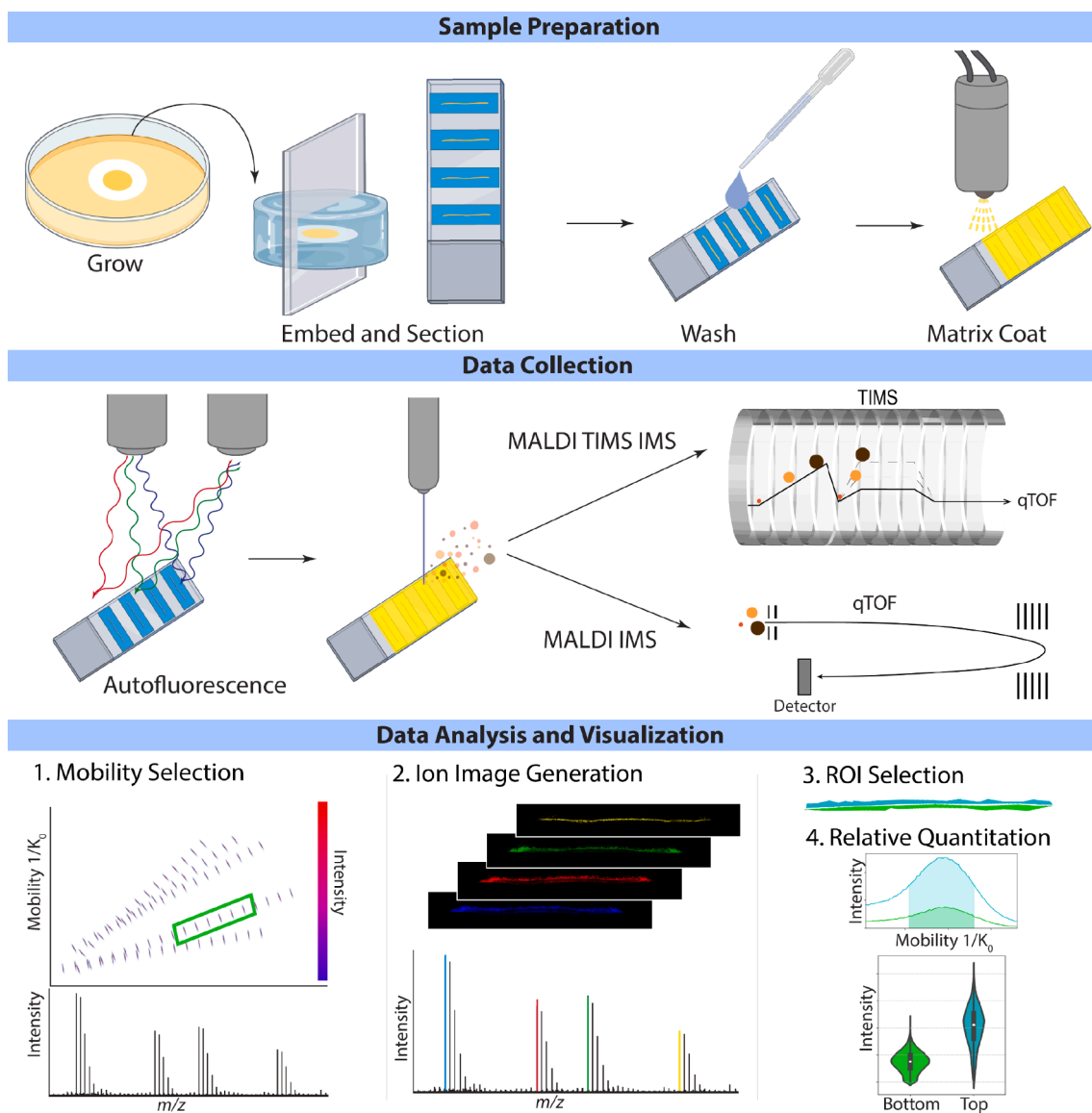


Fig. 1. Workflow of methodology for IMS of *S. aureus* biofilms including sample preparation demonstrating the orientation of the biofilm cross-section followed by data collection of both autofluorescence and IMS and finally data analysis and visualization where data are first filtered by TIMS mobility selection. Ion images are created, and relative quantitation is performed between layered subpopulations within the biofilm.

gelatin mix and frozen at -80°C . Samples were then cryosectioned to a $10\ \mu\text{m}$ thickness using a CM3050 S cryostat (Leica Biosystems, Wetzlar, Germany), and biofilm sections were then transferred to a conductive indium tin oxide (ITO) coated glass slide (Delta Technologies, Loveland, CO) coated with poly-L-lysine. A Teflon-coated slide was used to press down on top of the biofilm section, sandwiching it between the ITO slide and the Teflon-coated slide, resulting in even adherence of the section to the ITO slide. Sections were then thaw-mounted as normal. Washing was performed using an eyedropper 3X for 30 s and dried in a vacuum desiccator. Autofluorescence and fluorescence microscopy images were acquired using enhanced green fluorescent protein (EGFP), filter when imaging only green fluorescent protein (GFP), or EGFP, 4',6-diamidino-2-phenylindole (DAPI), and red fluorescent protein from *Discosoma* (DsRed) filters for autofluorescence on a Zeiss AxioScan.Z1 slide scanner (Carl Zeiss Microscopy GmbH, Oberkochen, Germany) before matrix application. Briefly, $150\ \text{mg}$ of 2',5'-Dihydroxyacetophenone (DHA) was dissolved in $10\ \text{mL}$ of a methanol/acetonitrile/water (80:10:10) solution and applied to a density of $2.08\ \mu\text{g}/\text{mm}^2$ by an M5 TM sprayer (HTX Technologies, Chapel Hill, NC). 1',5'-Diaminonaphthalene (DAN) was applied similarly by dissolving $100\ \text{mg}$ into acetonitrile/water (90:10)

and spraying to a density of $1.67\ \mu\text{g}/\text{mm}^2$.

Imaging mass spectrometry

All experiments were performed on a prototype timsTOF fleX mass spectrometer (Bruker Daltonik, Bremen, Germany) [52]. Images were acquired in TIMS mode of operation with an ion transfer time of $110\ \mu\text{s}$ and a prepulse storage time of $5\ \mu\text{s}$. Biofilm imaging data were collected using the Bruker Smartbeam 3D laser capable of steering the laser beam to ablate an adjustable square area, termed beam scanning. The beam scan was set to $6 \times 6\ \mu\text{m}$ with a pitch of $10\ \mu\text{m}$ using 70 shots per pixel and 59 % laser power. MALDI IMS burn patterns are provided to demonstrate the resultant laser ablation (**Supplemental Figure 14**). Positive and negative ionization mode data were collected on biofilm serial sections scanning from $m/z\ 300 - 1,300$ and $m/z\ 100 - 2000$, respectively. The TIMS EFG scan time was set to $500\ \text{ms}$, with a tunnel-in pressure of $2.1\ \text{mbar}$ and reduced mobility ($1/K_0$) range of $0.8 - 2.2\ (\text{V}\cdot\text{s})/\text{cm}^2$. TIMS imaging data were visualized via SCiLS Lab (Bruker Daltonik, Bremen, Germany) where hotspot removal was applied. In brief, each signal intensity scale was normalized to the maximum intensity detected

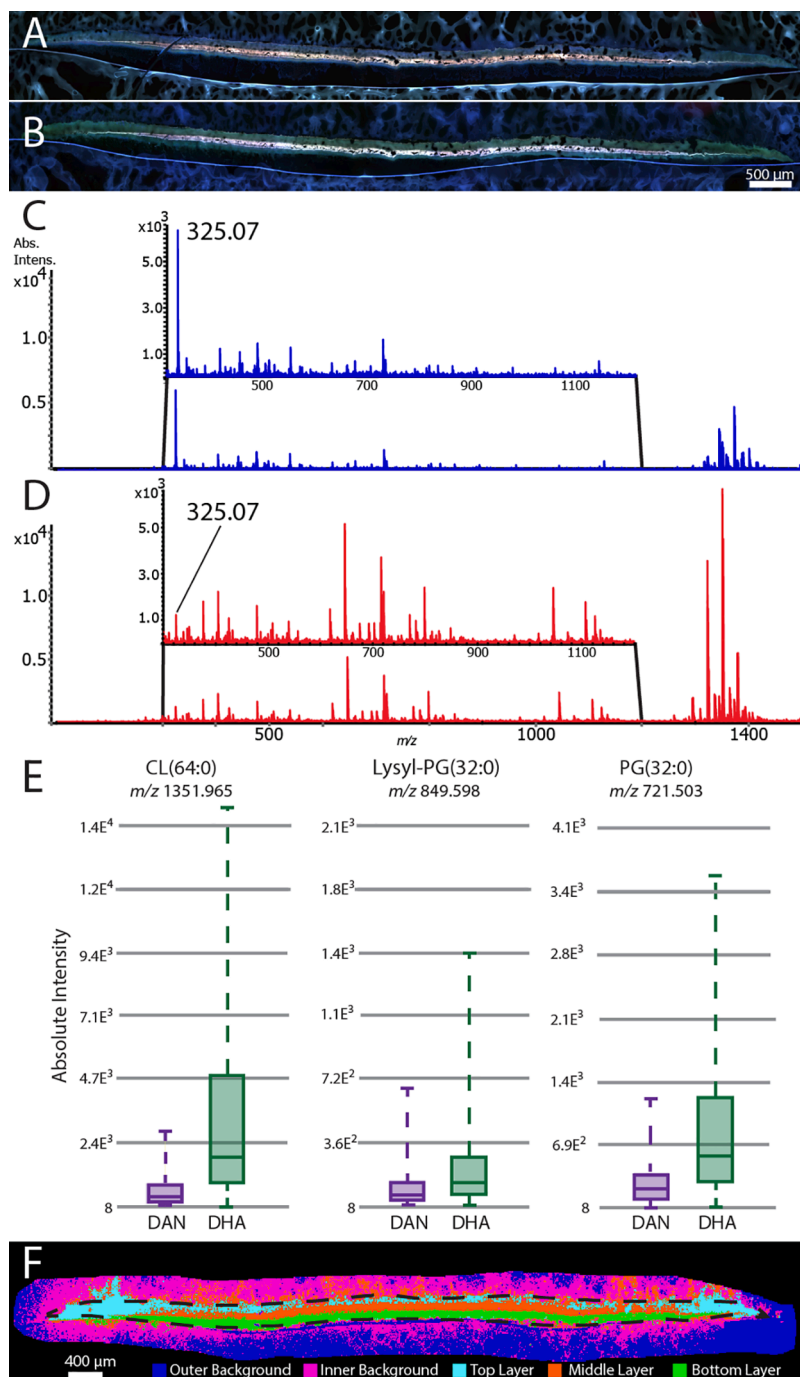


Fig. 2. Washing of *S. aureus* biofilms and MALDI matrix selection enhance lipid signal abundance in an IMS experiment. Autofluorescence microscopy shows minimal sample loss before (A), and after (B) washing with 150 mM ammonium formate. Negative mode mass spectra before (C), and after (D) washing show that washing also enhances signal of lipids, while reducing signal of chemical noise. Signal intensity box plots of CL(64:0), L-PG(31:0) and PG(32:0) (E) compare DHA and DAN matrices, demonstrating higher signal intensity for each lipid with DHA. *k*-means clustering performed on IMS data of a biofilm demonstrates biofilm layers cluster independently (F).

for a given ion and the color map is extended for enhanced perception of texture and dynamic range. *k*-means clustering was also performed in SCiLS using total ion current (TIC) normalization, $k = 5$ and a correlation distance metric. Lipid identifications were determined with high mass accuracy and LIPIDMAPS lipidomics gateway (lipidmaps.org) [75,76] and confirmed by MS/MS fragmentation where possible. Experiments for lipid abundance comparisons were run in technical triplicate and violin plots were generated using in-house developed software capable of considering both m/z and TIMS dimensions of the data.

Data processing

The data were exported into a custom binary format optimized for storage and speed of analysis of the ion mobility-IMS data. Each frame/

pixel contains 10^4 - 10^6 centroid peaks that span the acquisition range of m/z 100 – 2000 with 443,878 and 3,818 bins in the MS and ion mobility dimensions, respectively. The individual centroid peaks were inserted at their correct bin position along the MS and ion mobility dimensions and missing values were set to zero. Data was mass aligned using seven automatically selected m/z features and mass calibrated to several known standard molecules. The datasets were also ion mobility aligned to remove any drift time shift acquired during the experiment. Ion mobility-aware ion images were extracted using a ± 3 ppm mass window around each studied mass to produce a matrix of *the number of pixels* * *the number of ion mobility bins*. Such images can be used to create an ion image (summing all mobility information into a single vector), a subset of mobility bins can be selected (*e.g.*, to select one or multiple conformations) or all pixels can be summed to produce an extracted ion

mobilogram (XIM). In all cases, ion mobility images are normalized using TIC normalization.

Statistics

The XIM was fitted with a Gaussian curve to determine the full-width half maximum (FWHM) of the single conformation of the ion. The FWHM boundaries were used to select a subset of ion mobility bins within the ion mobility image and the 2D matrix was summed along the ion mobility dimension to produce a vector of intensities. The top and bottom masks were used to sub-select a set of pixels in the vector of intensities; each subset of pixels was then collected into a table and padded with NaNs to make sure the two vectors were identical in size and shape. Padding was necessary due to slight differences in the number of pixels in the top and bottom layer masks. All statistics were computed while excluding NaNs using the pandas Python package while the violin plots were generated using the seaborn visualization library.

Results and discussion

Biofilm growth and sample preparation for IMS

S. aureus biofilm preparation consisted of growth, embedding, cryo-sectioning, washing, autofluorescence microscopy, matrix deposition, and MALDI IMS analysis (Fig. 1). *S. aureus* biofilms were grown on tryptic soy agar (TSA) for 72 hrs to generate biofilms of ~ 400 μm thickness. Biofilms grown in this manner reproducibly reached maturity by 24 h in terms of the number of bacteria, as counted by colony forming units (CFUs), and maintained a constant CFU count over the next 48 h (Supplemental Fig. 1A). Optical images of biofilms were acquired at each transfer timepoint to provide additional morphological context throughout development (Supplemental Fig. 1B). Unlike other bacterial species such as *P. aeruginosa* [63], *S. aureus* biofilms grown *in vitro* often lack structural rigidity, making it challenging to retain their morphology during IMS sample preparation and necessitating embedding of the biofilm. Traditional embedding materials, such as OCT, interfere with MALDI IMS analysis. As a result, we elected to embed the biofilms in a mixture of CMC and gelatin (5 % and 10 % w/w, respectively) to provide structural support during cryosectioning [64]. Notably, we found that adhering the biofilm section evenly to the microscope slide by pressing down with a Teflon-coated microscope slide before thaw-mounting was critical for maintaining biofilm morphology with discrete layers (Supplemental Fig. 2A). Washing with ammonium formate and ammonium acetate solutions has been previously reported for MALDI IMS of tissue [65], but has not been investigated for use in *S. aureus* biofilms to our knowledge. For these reasons, we assessed HPLC grade water and both 50 mM and 150 mM solutions of ammonium formate and ammonium acetate for removing the salt and other additives present in growth media (Supplemental Fig. 2B). Sample loss resulting from each wash procedure was determined by autofluorescence microscopy by comparing before and after wash images. Of all the washes, 150 mM ammonium formate minimally perturbed sample morphology (Fig. 2A and 2B), while water caused the most damage. Furthermore, we found that washing with 150 mM ammonium formate yielded increased lipid signal for lipid species above m/z 300, especially enhancing CL lipid signal (Fig. 2C and 2D) compared to unwashed samples. For example, CL(64:0) displayed a more than tenfold enhancement after washing when analyzed in quadruplicate (Supplemental Fig. 3A – B). The identity of the predominant peak measured at m/z 325.07 in the unwashed samples is unknown, however it localizes to the matrix around the biofilm and primarily to the filter disc. It is likely a chemical interference from the bacterial culture or sample preparation process being remediated by the washing step (Supplemental Fig. 2C). For lipid analysis, we assessed both DAN and DHA matrices as they enable dual polarity imaging, thereby increasing the number and type of lipid species that can be detected. For effective comparison, PG(32:0), L-

PG(32:0) and CL(64:0) lipids were selected as representative of three unique and biologically important lipid classes. Absolute signal intensity for each of these lipids was compared for both MALDI matrices and in each case, DHA provided higher signal intensity than DAN (Fig. 2E). Moreover, DAN matrix clusters were detected across the measured m/z range in both polarities (Supplemental Fig. 2D), increasing overall background and conflating with lipid species. For example, lysyl-PG (30:0) was detected with an interfering and overlapping ion using DAN, but fully resolved using DHA (Supplemental Fig. 4A – B). To assess the effect of this on downstream lipid identifications, a LipidMAPS search of these datasets was performed comparing DAN and DHA where the number of annotated peaks using DHA was 388 compared to only 185 when DAN was used. For these reasons, DHA was selected for all future experiments.

With the goal of visualizing lipidomically distinct layers within the biofilms, we next collected 10 μm spatial resolution IMS data of an *S. aureus* biofilm and performed segmentation by *k*-means clustering (Fig. 2F). This analysis provided an unsupervised approach for clustering pixels whose spectral profiles are similar and then segmenting the imaging data by color coding each pixel by its cluster membership. Using five clusters ($k = 5$), we found that the matrix background clustered into two classes (outer and inner background, outside dotted line, Fig. 2F). We attributed this independent clustering of outer and inner background to slight delocalization of bacterial molecules or whole bacteria around the main structure of the biofilm. Interestingly, we observed three distinct layers within the biofilm to cluster independently, suggesting that the lipidomic profiles differ significantly in these distinct regions.

We elected to improve sensitivity and coverage of detected lipids using ion mobility. The contributions of TIMS were assessed by comparing peak capacity and signal-to-noise (S/N) with, and without TIMS separation implemented. To do this, we averaged the spectrum of a 10 μm spatial resolution imaging experiment without considering the TIMS dimension, resulting in the detection of 550 features (Supplemental Fig. 5C). Low signal intensity for various lipids, such as L-PG species was observed with S/N as low as 3. Considering the TIMS dimension however, S/N was increased for most detected lipid species. For example, lysyl-PG(28:0) (m/z 793.535) was detected with a S/N of 88 (Supplemental Fig. 5A), but by excluding noise at $1/K_0$ values, other than the lipid of interest, we found that the S/N improved by more than twofold (S/N 193, Supplemental Fig. 5B). Furthermore, when the TIMS dimension was considered for the dataset as a whole, ion mobility separation revealed ~ 1,000 additional detected features (1,545), likely consisting of isobars, isomers, and conformers of detected analytes (Supplemental Fig. 5D). Chemical specificity was also enhanced by TIMS, as we found that many ions of interest were observed with notable peak tailing in the TIMS dimension, which could be due to slight separation of additional confounding species from the ion of interest at similar m/z . By selecting a narrow window (0.015) around the peak of interest (Supplemental Fig. 6), extraneous species were excluded to enhance chemical specificity. As a result, all downstream analyses were performed with TIMS to reduce the overall spectral noise and increase chemical specificity, and all ion images depict ions within a $1/K_0$ and m/z range.

S. aureus biofilm imaging

Biofilms are known to comprise distinctive layers with inter-layer heterogeneity in gene regulation [8]. Here, we aimed to assess the lipid distributions within these layers. This growth pattern is thought to arise from a series of steps including attachment, multiplication, and detachment that *S. aureus* undergoes during the biofilm generation process [8,66]. Due to their small size, high spatial resolution IMS is required to obtain the necessary spatial fidelity for investigating these discrete layers. We performed 10 μm spatial resolution IMS in both positive and negative ion modes. TIMS was utilized to maximize

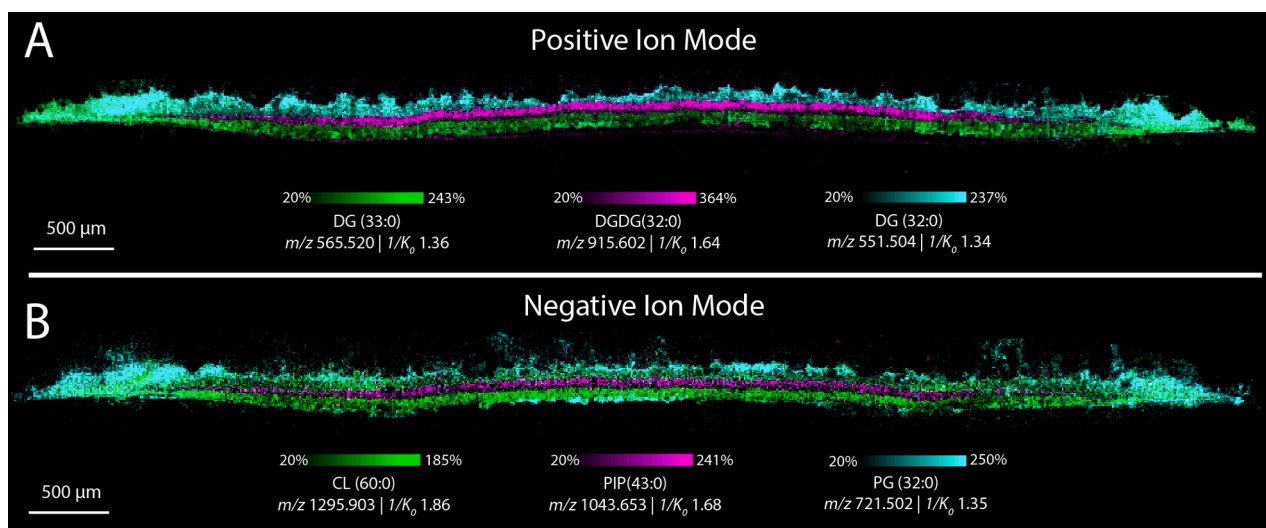


Fig. 3. High spatial resolution IMS (10 μ m) collected from a cross-section of an *S. aureus* biofilm in positive ion mode (A) and negative ion mode (B) showing delineation of three distinct layers. DG(33:0) (m/z 565.520, $1/K_0$ 1.36, 1.77 ppm error) (and CL(60:0) (m/z 1295.903, $1/K_0$ 1.86, 0.46 ppm error) appear to localize primarily to the bottom layer, while DGDG(32:0) (m/z 915.64, $1/K_0$ 1.637, 0.55 ppm error) and PIP(43:0) (m/z 1043.653, $1/K_0$ 1.68, 3.83 ppm error) localize to the middle layer. Finally, DG(32:0) (m/z 551.504, $1/K_0$ 1.34, 1.09 ppm error) and PG(32:0) (m/z 721.502, $1/K_0$ 1.35, 0.14 ppm error) are more abundant in the top layer.

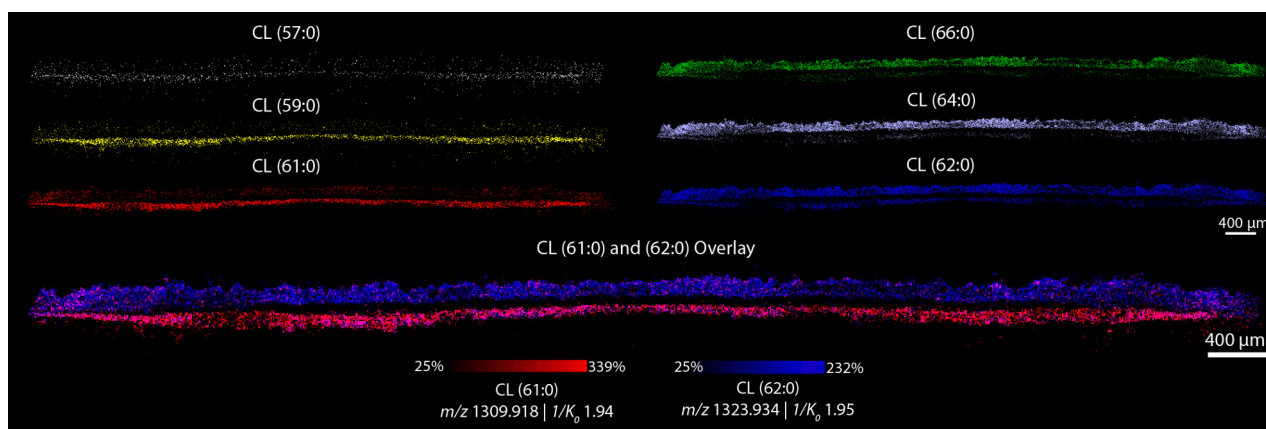


Fig. 4. Bacterial cardiolipin species distribute heterogeneously within biofilm layers. CL(57:0), CL(59:0) and CL(61:0) distribute primarily to the bottom layer, despite being from the same lipid class. Similarly, CL(62:0), CL(64:0) and CL(66:0) distribute primarily to the top layer of the biofilm. An overlay of CL(61:0) and CL(62:0) clearly demonstrates the delineation.

sensitivity and molecular coverage. Overall, we detected both odd-chain and even-chain PG, Lysyl-PG and CL, as well as phosphatidylcholine (PC), phosphatidylinositol-phosphate (PIP), diacylglycerols (DG) and digalactosyldiacylglycerols (DGDG), underscoring the sensitivity of our method. Furthermore, the spatial and molecular specificity of IMS uncovered three discrete layers within the biofilm. While horizontal *S. aureus* biofilm layers are known to exist, these results are among the first to visualize this gross structural information on the lipidomic level. As an example, DG(33:0) (m/z 565.520, $1/K_0$ 1.36), DG(32:0) (m/z 551.504, $1/K_0$ 1.34), and DGDG(32:0) (m/z 915.602, $1/K_0$ 1.64) localize to the bottom, top and middle layers, respectively (Fig. 3). Notably, DG(32:0) and DG(33:0) localize to opposite layers of the biofilm, despite differing only by the addition of a single methyl group on the acyl chain, signifying that lipid localization is not based solely on lipid class. Similar trends were observed in other classes of bacterial lipids such as cardiolipins, as discussed below. We also found discrete localizations of lipids to specific biofilm layers in negative ion mode. For example, CL(60:0) (m/z 1295.903, $1/K_0$ 1.86), PIP(43:0) (m/z 1043.653, $1/K_0$ 1.68), and PG(32:0) (m/z 721.502, $1/K_0$ 1.35) localized to the bottom,

and top layers, respectively. All ions depicted were putatively identified by accurate mass measurement and MS/MS fragmentation where possible (Table S1 and Supplemental Figure 7). More lipid classes ionize in negative ion mode, contributing to the higher diversity of lipids detected. Comparable to previous studies performed via liquid-chromatography tandem mass spectrometry (LC-MS/MS) [16], at least five lipid classes were detected in negative ion mode with most ions localizing primarily to a single layer. The lipidomic profile within each of these regions is distinct, and the delineation between each region is visually clear. These results coincide with other work on *B. subtilis* that demonstrated bacteria in different phenotypic states in varying locations within a biofilm [39,67], and suggest that *S. aureus* biofilm layers also act as discrete communities with a complex system of lipid distribution between them. We again found spatial heterogeneity of lipids from within the same class, such as CL(57:0), CL(59:0) and CL(61:0) (m/z 1253.853, $1/K_0$ 1.88, m/z 1281.887, $1/K_0$ 1.91, and m/z 1309.918, $1/K_0$ 1.94), that are all observed primarily in the bottom layer of the biofilm (Fig. 4). Conversely, CL(62:0), CL(64:0) and CL(66:0) (m/z 1323.934, $1/K_0$ 1.95, m/z 1351.965, $1/K_0$ 1.98 and m/z 1379.996, $1/K_0$ 2.01) were

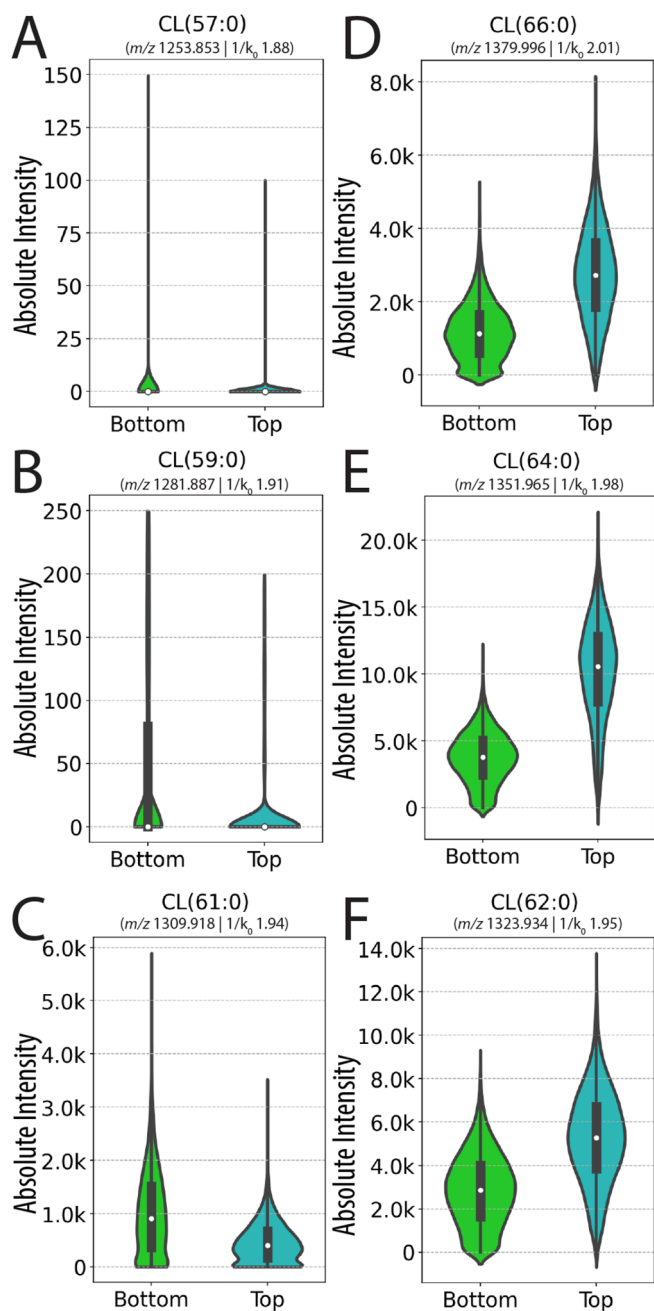


Fig. 5. *S. aureus* biofilms exhibit lipidomic heterogeneity between layers. Violin plots generated from one of three replicate images show that CL(57:0), CL(59:0) and CL(61:0) are more abundant in the spectrum extracted from the bottom layer (green) when compared to the top layer (cyan) (A – C), and that CL(62:0), CL(64:0) and CL(66:0) are more abundant in the top layer compared to the bottom (D – F). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

all observed primarily in the top layer of the biofilm. All CL lipids were confidently identified by MS/MS (Supplemental Figure 7B – G) (except CL(57:0) due to low signal intensity). If these spatial differences were only observed between different classes of lipid such as PG and CL lipids, this phenomenon might be the result of different growth phases between top and bottom layers. PG lipids are the predominant species in growing *S. aureus*, switching to CL lipids in the stationary phase [68,69]. However, there are no known functional differences for species within the cardiolipin class, potentially suggesting distinct functions of CL species within a spatial context. Interestingly, CL signal was below the limit of

detection in the middle layer of the biofilm. To determine if this is due to a difference in the cell density in that layer, a biofilm was grown with a strain of USA300 LAC that was genetically modified to constitutively express GFP. We then collected fluorescence microscopy and MALDI TIMS IMS on the same cross-section and found lower GFP signal in the middle layer by fluorescence microscopy, and corresponding absence of CL signal in the same region by MALDI TIMS IMS (Supplemental Figure 8). These results suggest that the absence of CL signal observed in the middle layer may be due to fewer bacteria present in that region of the biofilm. Most odd-chain CL lipids localize primarily to the bottom layer of the biofilm, and even-chain CL lipids are more abundant in the top layer (Fig. 4). However, the trend did not hold in every case in that CL(58:0) and CL(60:0) were observed primarily in the bottom layer of the biofilm (Supplemental Figure 9A and Fig. 3B). Cardiolipin chain length does not fully explain the separation observed between biofilm layers. Although it appears that longer chain length cardiolipins localize to the top layer and shorter chain length to the bottom, several cardiolipins with longer acyl chain length than 61:0, such as 65:0 have similar abundances between top and bottom layers (Supplemental Figure 9B). While the biological drivers of these observations are still unclear, these results show that CL lipids differing by only one methyl group have starkly different localizations within the microbial colony and further underscores the complexity of the lipid distribution patterns between layers within *S. aureus* biofilms.

We wanted to further assess the heterogeneity of lipidomic profiles between biofilm layers by making relative comparisons of lipid abundances between them. By acquiring IMS of a biofilm in triplicate, we first extracted spectra from regions of interest of top and bottom layers (Supplemental Fig. 6A–C). After selecting ion mobility ($1/K_0$) windows (Supplemental Fig. 6A–C) to remove chemical noise and any extraneous species, we compared mean signal intensities for each CL species shown in the ion images above (Fig. 5 and Supplemental Figure 10A – B). We found that CL(57:0) CL(59:0) and CL(61:0) were 3.7-, 6.3-, and 2.3-fold more intense in the bottom layer than in the top layer (Fig. 5A – C and Supplemental Figure 10A – B). In contrast, CL(62:0), CL(64:0), and CL(66:0), which primarily localize to the top layer, demonstrated a 1.8, 2.6, and 2.2-fold difference between layers (Fig. 5D – F and Supplemental Figure 10A – B). These findings are also clear in the extracted spectra from each layer (Supplemental Figure 11A – F). The means for these six ions were significantly different ($n = 3$, $p < 0.03$) between the top and the bottom layers. In addition to the quantitative analysis performed in technical triplicate, we confirmed that these trends were consistent in biological replicate (USA300 LAC), as well as a different isolate of USA300 (JE2), (Supplemental Figure 12A – B). Although this analysis is an example that analyzes only one class of bacterial lipid species, it demonstrates that the lipidomic differences between biofilm layers are significant and further supports that they are distinct at the lipidomic level.

This pronounced lipidomic heterogeneity between *S. aureus* biofilm layers led us to probe the biological drivers of this phenomenon. Notably, *S. aureus* is a facultative anaerobe with the ability to survive in both aerobic and anaerobic conditions [70]. *S. aureus* growth under anaerobic conditions leads to significant differences in fatty acid content. Specifically, anaerobic metabolism results in a shift from long to short isobranched fatty acids [71]. Changes in fatty acids, a major building block of lipids, could result in a shift in lipid content, as each observed lipid class is synthesized from the same pool of available fatty acids. We hypothesized that these layered subpopulations experience different amounts of accessible oxygen, and resultant shifts in fatty acid content could explain differences in CL lipid content between them. We investigated the role of oxygen in the spatial distributions of lipids in biofilms grown under anaerobic conditions compared to those grown under atmospheric conditions for the same duration. To investigate the growth of aerobic vs anaerobic *S. aureus* biofilms, we compared CFUs between anaerobic biofilms to their aerobic counterparts during biofilm development. At 72 hrs, when the MALDI TIMS IMS was performed,

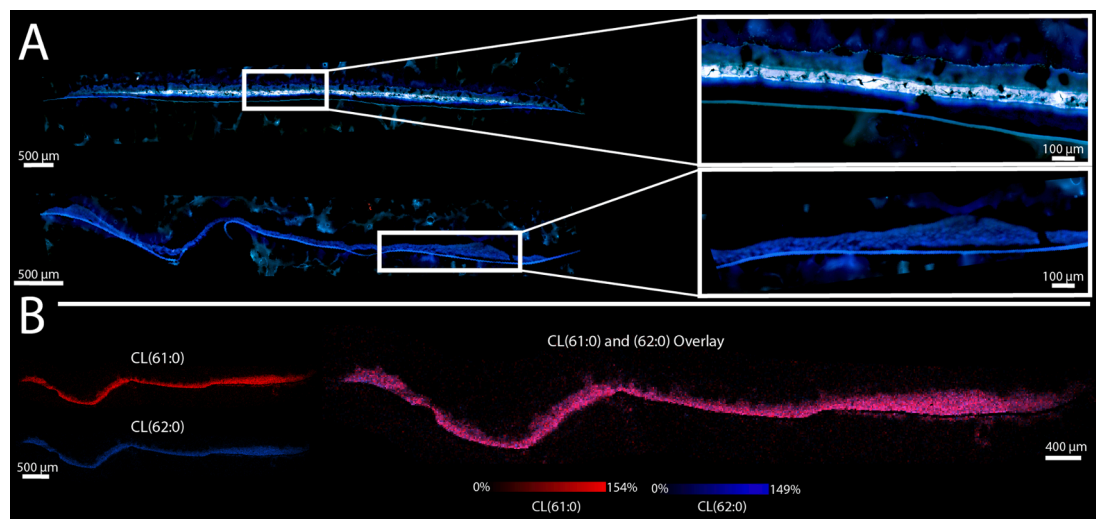


Fig. 6. *S. aureus* biofilms grown under anaerobic conditions generate thinner biofilms (bottom) comprised of a single layer compared to those grown under aerobic conditions (top) (A). IMS of a biofilm grown under anaerobic conditions reveals homogenous distribution of CL(61:0) and CL(62:0) (B) which have different distributions in biofilms grown under aerobic conditions.

biofilms grown in both conditions had reached densities greater than 1×10^9 CFUs per filter. While significantly different from one another, the difference between CFUs in the two conditions was still less than half a log (Supplemental Fig. 1A). Furthermore, after 24 hrs of development, both the aerobic and anaerobic biofilm populations were quite stable with no significant changes in their number of CFUs between 48 and 72 hrs. Autofluorescence of an anaerobic biofilm revealed considerably thinner single-layered biofilms, rather than three distinct layers observed in aerobic biofilms (Fig. 6A). We collected IMS data and performed *k*-means clustering with no discernable layers clustered separately in the IMS data. Instead, the entire biofilm clustered together, further suggesting the existence of a single layer (Supplemental Figure 13). The absence of layers indicates that oxygen may be necessary to induce *S. aureus* biofilm layer generation that could have implications for biofilm growth in environments of varying oxygen availability. In particular, if the layers observed in aerobic biofilms are the result of waves of bacterial detachment, a known step of the *S. aureus* biofilm generation process [66,72], these results suggest that oxygen may be required for biofilm detachment. Our findings also align with the hypothesis that there may exist a mechanism of biofilm structure formation involving metabolic differentiation of cells leading to metabolically distinct subpopulations within an *S. aureus* biofilm [66]. If this process depends on the presence of oxygen, this could lead to a lack of bacterial differentiation and subsequent absence of biofilm layers. In addition to the loss of layered subpopulations, we observed homogeneous lipid distributions spanning the cross-section of the anaerobic biofilm. For example, when comparing ion images of CL(61:0) and CL(62:0), both species appear to share the same spatial distribution across the entire biofilm (Fig. 6B). This is in contrast to that of the aerobic biofilm where these lipid species demonstrated distinct localizations to discrete biofilm layers (Fig. 4). We observed all CL species in the anaerobic biofilm, indicating that oxygen is not necessary for the production of these lipid species. However, this does not rule out that an oxygen gradient could be contributing to heterogeneity of lipid abundances between biofilm layers in aerobic biofilms.

Conclusions

Various sample preparation protocols including embedding in a mixture of CMC/gelatin and isotonic washing with ammonium formate resulted in preservation of *S. aureus* biofilm structure while removing spectral interferences. IMS and *k*-means clustering yielded unsupervised

visualization of *S. aureus* biofilm layers, providing an initial indication of the lipidomic heterogeneity between biofilm layers. Gas-phase separation by TIMS also contributed to increasing both the S/N and the overall peak capacity in 10 μ m spatial resolution IMS. Discrete, lipidomically distinct layered subpopulations were revealed by IMS of bacterial lipids in both positive and negative polarities facilitated by the methodologies developed herein. Not only were different lipids observed to localize primarily to discrete layers, but lipids from within the same class, in many cases only differing by one methyl group on the acyl chain, were observed to localize primarily to different layers. Relative quantitation from extracted mass spectra of discrete layers afforded inter-layer comparison of individual lipid abundances and demonstrated differences in select CL lipid species between layers. Further, our findings suggest that individual CL's that have historically been thought to contribute cooperatively toward certain functions, could be playing distinct biological roles. Future work growing *S. aureus* biofilms with access to defined fatty acids, or genetic alterations of various enzymes involved in CL synthesis could elucidate the biological drivers of these lipid distributions. Following the CL study, we found that *S. aureus* biofilms grown under anaerobic conditions developed thinner biofilms comprised of a single layer compared to thicker biofilms comprised of three layers when grown under aerobic conditions. This suggests that the generation of multiple layers in *S. aureus* biofilms may require aerobic growth conditions. Given that biofilms grown anaerobically are not able to reach the same CFU's as those grown under aerobic conditions, it remains possible that these differences are due to differences in size of the biofilm as opposed to oxygen concentration. However, because both growth conditions reached CFU's that remained stable over time, we would not expect development of horizontal layers in anaerobic biofilms over an extended time period. IMS of these anaerobic biofilms revealed that the lipids analyzed in this work localized homogeneously throughout the anaerobic biofilm. Although further analyses are required to fully understand the significance of lipid distributions within *S. aureus* biofilms, this work using IMS technology demonstrates significant lipidomic heterogeneity between biofilm layers. Furthermore, the methodologies developed herein have the potential to be applied towards understanding spatially discrete biofilm responses to external stimuli, such as exposure to antimicrobials. Studies of this nature could reveal localized molecular characteristics that could be exploited for new therapeutic strategies.

Associated content

The raw data are available at <https://doi.org/10.6084/m9.figshare.c.6054968.v1>.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.jmsacl.2022.09.003>.

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