The Membrane Composition Defines the Spatial Organization and Function of a Major Acinetobacter baumannii Drug Efflux System

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ACRENOTOBACTER BAUMANNII is one of the world’s most problematic nosocomial pathogens. The combination of its intrinsic resistance and ability to acquire resistance markers allow this organism to adjust to antibiotic treatment. Despite being the primary barrier against antibiotic stress, our understanding of the A. baumannii membrane composition and its impact on resistance remains limited. In this study, we explored how the incorporation of host-derived polyunsaturated fatty acids (PUFAs) is associated with increased antibiotic susceptibility. Functional analyses of primary A. baumannii efflux systems indicated that AdeB-mediated antibiotic resistance was impacted by PUFA treatment. Molecular dynamics simulations of AdeB identified a specific morphological disruption of AdeB when positioned in the PUFA-enriched membrane. Collectively, we have shown that PUFAs can impact antibiotic efficacy via a vital relationship with antibiotic efflux pumps. Furthermore, this work has revealed that A. baumannii’s unconditional desire for fatty acids may present a possible weakness in its multidrug resistance capacity.

IMPORTANCE Antimicrobial resistance is an emerging global health crisis. Consequently, we have a critical need to prolong our current arsenal of antibiotics, in addition to the development of novel treatment options. Due to their relatively high abundance at the host-pathogen interface, PUFAs and other fatty acid species not commonly synthesized by A. baumannii may be actively acquired by A. baumannii during infection and change the biophysical properties of the membrane beyond that studied in standard laboratory culturing media. Our work illustrates how the membrane phospholipid composition impacts membrane protein function, which includes an important multidrug efflux system in extensively-drug-resistant A. baumannii. This work emphasizes the need to consider including host-derived fatty acids in in vitro analyses of A. baumannii. On a broader scope, this study presents new findings on the potential health benefits of PUFA in individuals at risk of contracting A. baumannii infections or those undergoing antibiotic treatment.


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Acinetobacter baumannii is one of the world’s most notorious multidrug resistant pathogens (1, 2), yet how it responds to host-mediated stress is poorly understood. Previous reports have shown that this human pathogen displays susceptibility to host-derived omega-3 polyunsaturated fatty acids (PUFAs), such as docosahexaenoic acid (DHA) (3, 4). Here, we examined the transcriptomic responses to 0.25 mM DHA stress to identify possible genetic traits responsible for these outcomes. Although plasma DHA concentrations range from 0.1 to 0.2 mM in humans on a typical Western diet (5, 6), this can increase to levels greater than 0.4 mM DHA in populations with higher marine fish oil intake (7). Differential expression analyses revealed transcripts for 53 and 32 genes to be of higher and lower abundance, respectively ($\geq 2$-fold change, $P \leq 0.05$), in DHA treated compared to untreated A. baumannii AB5075_UW cells (see Table S1 in the supplemental material). Among the most downregulated genes were those coding for two putative oxidoreductases, ABUW_3843 (5.29-fold) and ABUW_1104 (2.44-fold) (Fig. 1A; see Fig. S1A to C in the supplemental material), which are likely to assist in the electron transport required for their cotranscribed fatty acid desaturases. This is likely to be a specific response to restrict the introduction of more double bonds in acyl chains following PUFA treatment. Consistently, an increase in DHA susceptibility in the repressor mutant of this cluster was observed (Fig. S1D). Despite minimizing the exposure to DHA, the transcriptomic analyses also revealed potential general stress responses. A homologue of the Escherichia coli stress tolerance gene ygiW was significantly downregulated 4.68-fold. Further, a total of 18 genes associated with iron utilization were significantly downregulated (Fig. 1A), which could not be linked directly to fatty acid homeostasis. A number of genes with putative roles in $\omega$-oxidation were found to be significantly upregulated upon DHA treatment (Fig. 1A). The resulting product of the $\omega$-oxidation pathway, fatty dicarboxylic acids, can be catabolized by $\beta$-oxidation or dicarboxylate catabolic (dca) pathways (8), and several putative components in both these pathways were upregulated under DHA stress. A predicted long-chain fatty acid transporter gene (fadL; ABUW_0724) was among the upregulated genes of the $\beta$-oxidation pathway. Importantly, fadL mutants in either the A. baumannii AB5075_UW or ATCC 17978 background displayed enhanced tolerance to DHA stress (Fig. 1B; Fig. S1). Although transcriptomic profiling showed the downregulation of two desaturases to restrict introduction of double bonds following DHA treatment, it also revealed the upregulation of fadL. This somewhat greedy and unconditional acquisition of energy-rich DHA has detrimental impacts on its fitness. Unlike A. baumannii, which can be represented in a vast array of environmental habitats, Streptococcus pneumoniae is fully host adapted and is attuned to the diligent acquisition of distinct fatty acids in the host environment, which is facilitated by the concerted action of having selective proteins (FakB1, saturated FAs; FakB2 monounsaturated FAs; FakB3, PUFAs), as well as appropriate transcriptional regulation of these systems (9, 10).

Considering PUFA incorporation affects A. baumannii membrane permeability (4), we examined antimicrobial susceptibility of A. baumannii strains AB5075_UW and ATCC 17978 with or without DHA supplementation (Fig. 1C). The presence of subinhibitory amounts of DHA resulted in a 4- to 8-fold decrease in aminoglycoside resistance (gentamicin and streptomycin, respectively) and 2-fold decrease in chloramphenicol resistance in strain AB5075_UW. These differences were less dramatic in the more-antibiotic-susceptible strain ATCC 17978, where only 2-fold reductions were observed for gentamicin and chloramphenicol following cotreatment with DHA. Although commonly known for their ability to interact with phospholipids, resistance to macrolides (erythromycin and azithromycin) was not affected by PUFA treatment. In addition to altering antibiotic resistance, we examined the impact of PUFA supplementation on oxidative stress tolerance in strain AB5075_UW and revealed that PUFA treatment impacted tolerance to paraquat, which induces the formation of intracellular...
superoxide stress (see Fig. S2 in the supplemental material). Contrastingly, PUFA-treated bacteria were not more susceptible to exogenously supplemented hydrogen peroxide compared to untreated bacteria (Fig. S2). Examination of the impact of DHA upon gentamicin or chloramphenicol susceptibility in a $fadL$ mutant in either an AB5075_UW or ATCC 17978 background revealed that increased susceptibility to these compounds occurs primarily following DHA uptake into the cell (Fig. 1D). In contrast to $A.\ baumannii$, no dramatic changes ($\geq 2$-fold) in antibiotic susceptibility following DHA treatment were seen in the Gram-positive bacterium, $Streptococcus\ pneumoniae$ (see Table S2 in the supplemental material). Hence, our study supports the identification of a plausible pathogen-specific Achilles’ heel, this being the active acquisition of DHA by $A.\ baumannii$ and the subsequent increase in antibiotic susceptibility.

Considering the roles of resistance-nodulation-cell division (RND) efflux systems in lipid homeostasis and DHA resistance (4, 11, 12), we studied the efflux activities of AdeB and AdeJ in $A.\ baumannii$ with and without omega-3 PUFA enrichment. We found that the specific roles of AdeB in gentamicin and pentamidine resistance were
impacted by DHA treatment, as the resistance decreased to a greater extent when AdeB was present (i.e., in wild-type or adeJ::T26 cells) than in the adeB::T26 mutant (Fig. 2A and B). To delineate the relative impact of DHA on AdeJ efflux activity, we examined the ethidium bromide (EtBr) efflux potential, by analyzing the cells with and without treatment with the protonophore carbonyl cyanide m-chlorophenyl hydrazine (CCCP), which can indirectly prevent efflux from RND pumps by collapsing the proton motive force. We found that active efflux by AdeJ is required for preventing the accumulation of EtBr, but DHA did not negatively impact this process (Fig. 2C). Instead, the increased membrane permeability as a result of DHA incorporation required greater EtBr efflux, which was reflected in the enhanced EtBr efflux potential of DHA-treated cells.
Since RND efflux relies upon the proton motive force across the cytoplasmic membrane, we ascertained the possibility of ion leakage in the *A. baumannii* membrane following DHA incorporation. Lipid samples extracted from actively growing *A. baumannii* cells were used to generate tethered bilayer lipid membranes (tBLMs). Both tBLMs displayed typical electrochemical properties (Fig. 2D) and similar responses to the incorporation of the ion carrier valinomycin (see Table S3 in the supplemental material). These analyses illustrate that the observed dysfunction of AdeB is not a result of the membrane being compromised in its ability to retain a proton motive force following the incorporation of DHA.

Evidence is emerging that phospholipids can influence folding, structure, and function of some membrane proteins, including bacterial RND systems (13–15). To identify possible differences in AdeB and AdeJ conformations in the membrane, we studied the dynamics of AdeB and AdeJ (both modeled on the AdeB cryo-electron microscopy [cryo-EM] structure; 6OWS) in the *A. baumannii* phospholipid environment with and without omega-3 PUFA enrichment in coarse-grained molecular dynamics (MD) simulations. After 15 µs replicate simulations, the lipid annulus surrounding both AdeB and AdeJ was enriched with unsaturated lipids in the untreated *A. baumannii* membrane (Fig. 2E). PUFA-containing phospholipids were heavily localized around AdeB and AdeJ when the protein complexes were embedded in a DHA-treated *A. baumannii* membrane (Fig. 2E). Although the conformation of the AdeJ trimer was largely unaffected by these changes in its phospholipid environment, the conformation of AdeB displayed a dramatic shift, with the complete loss of the protein-protein interface between adjacent transmembrane domains of two AdeB protomers (Fig. 2F). When assessed in conjunction with functional assays of AdeB activity in the presence of DHA, these changes suggest a possible mechanism for disruption of the conformational cycling of AdeB required for efflux activity. Consistently, previous reports on the membrane-disrupting biocides on AdeABC have also linked a role in cell envelope integrity and AdeABC efflux activity (16). Overall, the data presented here provide insights into the interplay between the membrane lipid composition and specific RND efflux activities. Hence, our work has established a molecular basis for how the lipid bilayer composition and its biophysical properties may affect antibiotic treatment success.

Overall, this study has presented a comprehensive analysis of the antimicrobial effects of host fatty acids upon *A. baumannii* membrane biology. Although, omega-3 PUFA supplementation is unlikely to affect healthy individuals, those that are at increased risk of contracting bacterial infections may benefit, in particular during antibiotic treatment.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**TEXT S1**, DOCX file, 0.1 MB.

**FIG S1**, TIF file, 0.7 MB.

**FIG S2**, TIF file, 0.1 MB.

**TABLE S1**, DOCX file, 0.03 MB.

**TABLE S2**, DOCX file, 0.02 MB.

**TABLE S3**, DOCX file, 0.02 MB.

**TABLE S4**, DOCX file, 0.02 MB.

**TABLE S5**, DOCX file, 0.02 MB.

**TABLE S6**, DOCX file, 0.03 MB.

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