| 1 | Phosphorus stress induces the synthesis of novel glycolipids in <i>Pseudomonas</i> |
|----|---|
| 2 | aeruginosa that confer protection against a last-resort antibiotic |
| 3 | |
| 4 | Running title: ecophysiology of lipid renovation in Pseudomonas aeruginosa |
| 5 | |
| 6 | Rebekah A. Jones ^{1,2} , Holly Shropshire ² , Caimeng Zhao ³ , Andrew Murphy ² , Ian Lidbury ^{2,4} , |
| 7 | Tao Wei ³ , David J Scanlan ² , Yin Chen ² |
| 8 | |
| 9 | |
| 10 | ¹ MRC Doctoral Training Partnership, University of Warwick, Coventry, CV4 7AL, UK. |
| 11 | |
| 12 | ² School of Life Sciences, University of Warwick, Coventry, CV7, 7AL, UK |
| 13 | |
| 14 | ³ School of Food and Biological Engineering, Zhengzhou University of Light Industry, |
| 15 | Zhengzhou, 450002, China |
| 16 | |
| 17 | ⁴ Department of Animal and Plant Sciences, University of Sheffield, Sheffield, S10 1AE, |
| 18 | UK |
| 19 | |
| 20 | |
| 21 | Correspondence to Yin Chen, (email, <u>y.chen.25@warwick.ac.uk</u> , phone 00 44 24 |
| 22 | 76528976) |
| 23 | |
| 24 | Classification: Integrated genomics and post-genomics approaches in microbial ecology |

25 Abstract

26 Pseudomonas *aeruginosa* is a nosocomial pathogen with a prevalence in 27 immunocompromised individuals and is particularly abundant in the lung microbiome of 28 cystic fibrosis patients. A clinically important adaptation for bacterial pathogens during 29 infection is their ability to survive and proliferate under phosphorus limited growth 30 conditions. Here, we demonstrate that P. aeruginosa adapts to P-limitation by substituting 31 membrane glycerophospholipids with sugar-containing glycolipids through a lipid 32 renovation pathway involving a phospholipase and two glycosyltransferases. Combining 33 bacterial genetics and multi-omics (proteomics, lipidomics and metatranscriptomic 34 analyses), we show that the surrogate glycolipids monoglucosyldiacylglycerol and 35 glucuronic acid-diacylglycerol are synthesised through the action of a new phospholipase 36 (PA3219) and two glycosyltransferases (PA3218 and PA0842). Comparative genomic 37 analyses revealed that this pathway is strictly conserved in all P. aeruginosa strains 38 isolated from a range of clinical and environmental settings and actively expressed in the 39 metatranscriptome of cystic fibrosis patients. Importantly, this phospholipid-to-glycolipid 40 transition comes with significant ecophysiological consequence in terms of antibiotic 41 sensitivity. Mutants defective in glycolipid synthesis survive poorly when challenged with 42 polymyxin B, a last-resort antibiotic for treating multi-drug resistant *P. aeruginosa*. Thus, 43 we demonstrate an intriguing link between adaptation to environmental stress (nutrient 44 availability) and antibiotic resistance, mediated through membrane lipid renovation that is 45 an important new facet in our understanding of the ecophysiology of this bacterium in the 46 lung microbiome of cystic fibrosis patients.

47 Introduction

48 P. aeruginosa is a significant nosocomial pathogen in intensive care units causing 49 pneumonia, surgical wound site infections and sepsis (1-2). It is now recognised as a 50 leading cause of morbidity and mortality in chronically infected cystic fibrosis (CF) patients 51 and immunocompromised individuals due to the surge of carbapenem resistant strains, a 52 key group of first line antibiotics for treating P. aeruginosa infections (3). For these drug-53 resistant *P. aeruginosa* strains, a viable but not ideal treatment option are polymyxins, 54 considered to be last resort antibiotics. Although polymyxins are active against P. 55 aeruginosa, their use was originally discontinued due to concerns over toxicity (4). Indeed, 56 P. aeruginosa has started to develop mechanisms of resistance to polymyxins due to an 57 increase in their use globally. These primarily include modifications to the 58 lipopolysaccharide (LPS) layer of the outer membrane through the addition of 4-amino-4-59 deoxy-L-arabinose (L-Ara4N) or phosphoethanolamine (pEtN) (5-6). These changes 60 perturb the electrostatic interaction between cationic polymyxins and the normally 61 negatively charged LPS.

62

63 Glycerophospholipids, such as phosphatidylglycerol (PG) and phosphatidylethanolamine 64 (PE), are the major lipids forming the membrane lipid bilayer in bacteria, archaea and 65 eukaryotes (7-11). They play a fundamental role in the evolution of the cell and it is widely 66 accepted that the last universal common ancestor possessed a phospholipid membrane (12-13). Although it is uncertain why evolution selected glycerophospholipids as the 67 68 building blocks for maintaining cellular membranes (13), it is known that organisms can 69 alter their membrane lipid composition in response to nutrient stress or environmental 70 changes (7, 14). Previous studies have firmly established the link between nutrient stress, 71 particularly phosphorus availability, and the expression of a variety of virulence factors in 72 P. aeruginosa (15-19). However, it is unclear whether adaptation to phosphorus limitation

73 in this bacterium causes a change in membrane lipid composition and, if so, whether lipid 74 remodelling comes with unforeseen ecophysiological consequences. Using a synthesis of multi-omics approaches, here we show that *P. aeruginosa* produces surrogate glycolipids 75 76 to replace phospholipids in response to phosphorus limitation. These glycoglycerolipids 77 have not previously been reported in *P. aeruginosa*. This lipid renovation pathway is strictly conserved in all P. aeruginosa strains isolated from a range of clinical settings and 78 79 actively expressed in the metatranscriptome of cystic fibrosis patients. Importantly, such 80 a phospholipid-to-glycolipid transition comes with a significant consequence in antibiotic 81 sensitivity, in that glycolipids confer protection when challenged with the antimicrobial 82 peptide polymyxin B. As such, glycolipid-mediated resistance to polymyxin B represents 83 a new resistance mechanism that is quite different from the previously documented 84 modification of LPS (5-6). This work highlights how the physiological adaptation of 85 Pseudomonas aeruginosa to phosphorus limitation can mediate a physiological response 86 that may have profound implications for the survival of the bacteria in the lung microbiome.

- 87
- 88

89 Materials and methods

90 Cultivation of *P. aeruginosa* and mutants

91 P. aeruginosa strain PAO1 was obtained from the DSMZ culture collection (Germany) and 92 routinely cultured in lysogeny broth (LB). A defined medium previously outlined for 93 Pseudomonas species to control phosphate levels was also used (20). This modified 94 minimal media A comprised: Na-succinate 20 mM, NaCl 200 mg L⁻¹, NH₄Cl 450 mg L⁻¹, CaCl₂ 200 mg L⁻¹, KCl 200 mg L⁻¹, MgCl₂ 450 mg L⁻¹, with trace metals FeCl₂ 10 mg L⁻¹ 95 and MnCl₂ 10 mg L⁻¹, with 10 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid 96 97 (HEPES) buffer used at pH 7. Na₂HPO₄ was then added to a final concentration of 50 µM 98 (low P) or 1 mM (high P). An intermediate phosphate source of 400 µM Na₂HPO₄ was 99 used for overnight cultures in some experiments to prevent any excess storage of 100 phosphate that could hamper results. All components were filter sterilised using 0.22 μ m 101 pore-size filters, and made up using double deionised H₂O. Mutants were obtained from 102 the *P. aeruginosa* strain PAO1 transposon mutant library at the University of Washington, 103 and confirmed using PCR and subsequent sequencing.

104

105 Alkaline phosphatase assay

106 Alkaline phosphatase activity was monitored as a measure of Pi stress. Liquid *P.* 107 *aeruginosa* culture samples were incubated with 10 mM *para*-nitrophenol phosphate 108 (*p*NPP) to a final concentration of 1 mM. Yellow-*p*NP supernatant was measured in 109 triplicate at 407 nm (BioRad iMark microplate reader). Readings were normalised using 110 both a Tris-only incubation control and further by bacterial density (optical density reading 111 at 600 nm (OD₆₀₀)).

112

113 Over-expression of Agt1 and Agt2 in *E. coli*

114 P. aeruginosa genes PA3218 (agt1) and PA0842 (agt2) were codon optimised for E. coli and chemically synthesised (GenScript) into plasmid pET-28a(+). E. coli BLR(DE3) 115 116 competent cells were thawed for 5 minutes before incubation with 10 ng pET-28a Agt 117 plasmid, and placed on ice for 5 minutes. Cells were then subjected to heat shock at 42°C 118 for 30 seconds, placed back on ice for 2 minutes. Recovery SOC media was added, with 119 samples incubated at 37°C shaking, for 1 hour. Transformed cells were then plated onto 120 kanamycin-LB agar and grown overnight at 37°C. To harvest cells for lipid extraction, 121 single colonies were picked to grow in small volume LB-Kan to 0.6 OD₆₀₀ before induction 122 with 0.4 mM IPTG overnight at 25°C. 1 mL samples were then pelleted at 10,000 x q for 123 5 minutes. Pellets were stored at -80°C until lipid extraction and subsequent analysis on 124 HPLC-MS.

125 To purify the Agt1 and Agt2 proteins from recombinant E. coli, IPTG was added to 126 a final concentration of 0.5 mM once the cultures reached an OD₆₀₀ of 0.6. After a further 127 12 h of growth at 30°C, cells were harvested by centrifugation and resuspended in buffer 128 A containing 50 mM Tris-HCl, pH 7.9, 50 mM NaCl. Cells were disrupted by sonication 129 and 1% (w/v) triton X-100 was then added and the cells were then incubated for 2.5 hr at 130 4°C. The cells were then centrifuged at 12,000 × g for 20 min, and the soluble fraction 131 was loaded onto a nickel column (GE Healthcare, USA) pre-equilibrated with buffer A. The 132 recombinant Agt1 and Agt2 enzymes were eluted with an elution buffer (20 mM Tris-HCl, 133 pH 7.9, 500 mM NaCl. 300 mM imidazole) and dialyzed overnight into buffer A to remove 134 imidazole. For further purification, the samples were dialyzed overnight into buffer B 135 containing 50 mM Tris-HCl, pH 7.9, 200 mM NaCl, concentrated by ultrafiltration using a 136 30-kDa membrane (Millipore), and loaded onto a Superdex 200 (16/60) gel filtration 137 column (GE Healthcare, USA), which was pre-equilibrated with buffer B (50 mM Tris-HCl, 138 pH 7.9, 200 mM NaCl). The fraction size was 0.5 ml, and the flowrate was 0.5 ml/min. 139 Purified protein was analysed by SDS-PAGE, and protein concentrations were determined 140 using the Bradford assay.

141

142 Membrane lipid extraction and HPLC-MS analysis

143 Intact polar membrane lipids were extracted using a modified version of the typically used 144 Folch extraction method (21, 22). Liquid P. aeruginosa cultures growing in high and low 145 phosphate modified minimal medium A were sampled after 8 hrs. collecting the equivalent 146 of 0.5 OD₆₀₀ into a 2 mL glass chromacol vial (Thermo Scientific), pelleted at 4°C, 4,000 147 rpm for 15 minutes. For lipid extraction, a ratio of 500:300:1000 µL of methanol:water:chloroform (all LC-MS grade) was used. The lipid fraction was collected 148 149 from the lower phase using a glass Pasteur pipette. This chloroform extract was then dried 150 under a stream of nitrogen (Techne sample concentrator) and resuspended in 1 mL 95%

151 (v/v) acetonitrile (HPLC grade): 5% (w/v) ammonium acetate (10 mM, pH 9.2) for analysis. 152 Extracted lipid samples were analysed using an UltiMate 3000 HPLC (Thermo Scientific) 153 system coupled to an AmazonSL quadrupole ion trap (Bruker) mass spectrometer (MS), 154 using electrospray ionisation. Hydrophilic interaction chromatography (HILIC) using a 155 BEH amide XP column (Waters) was utilised to separate lipid classes based on their head 156 group (35). The column chamber was maintained at 30°C and the samples passed 157 through at a 150 µL min⁻¹ flow rate. The mobile phase of acetonitrile:ammonium acetate 158 (pH 9.2) was used to elute the sample in a 15 minute per sample gradient, from 95% to 159 28% ammonium acetate. The lipid d17:1/12:0 sphingosylphosphoethanolamine (Sigma-160 Aldrich, 50 nM) was added to the samples and used as an internal standard. Tandem MS 161 (or MSⁿ) was used to fragment the intact lipids for identification. The data were analysed 162 using the Bruker Compass software package (DataAnalysis and QuantAnalysis).

163

164 Enzyme activity assays

165 The glycosyltransferase activity of Agt1 and Agt2 was measured using uridine diphosphate (UDP)-glucose or UDP-glucuronic acid and 0.1 mM C16:0/C18:1 166 167 diacylglycerol (DAG) as the substrate. 2.0 µM purified enzyme was used in 10 mM 168 Tricine/KOH buffer, pH 8.5 with 2 mM dithiothreitol. The resulting mixture (500 µl) was 169 incubated at 30°C for 60 min with constant shaking at 200 rpm. The lipid products were 170 extracted using the Folch method as described above. The lipid extracts were further 171 analysed by LC-MS for the identification of MGDG/GADG though MSⁿ fragmentation and 172 for the quantification of DAG against standards. The K_m and V_{max} values were calculated 173 using Michaelis-Menten plots with various concentrations of UDP-sugars (0.1 to 1.0 mM) 174 in three replicates.

175

176 Antibiotic sensitivity assays

177 *P. aeruginosa* cultures were grown to an OD₆₀₀ of 0.6 in high or low phosphate minimal 178 media A (see above). Cultures were then diluted 1:100 in prewarmed minimal media A 179 containing 4 µg mL⁻¹ polymyxin B sulfate (Sigma). Samples were incubated at 37°C, 180 180 rpm, and assayed for survivors at specified time points by serial dilution plating onto LB. 181 *E. coli* cultures containing pET-28a-Aqt1 or pET-28a-Aqt2 were grown to an OD_{600} of 0.6 182 in LB broth, and the expression of Agt1 and Agt2 was induced by incubation with 0.4 mM 183 IPTG overnight at 25°C. A negative control of *E. coli* containing the pET-28a vector only 184 was also set up. Overnight cultures were diluted 1:100 in prewarmed LB broth containing 0.4 mM IPTG and 20 µg mL⁻¹ polymyxin B sulfate (Sigma). Samples were incubated at 185 37°C, 180 rpm, and assayed for survivors at specified time points by serial dilution plating 186 187 onto LB agar + kanamycin 25 μ g mL⁻¹.

188

189 Comparative proteomic analysis

190 P. aeruginosa PAO1 WT (1 mM phosphate, 50 µM phosphate) and PIcP mutant (50 µM 191 phosphate) cell pellets in three biological replicates were resuspended in LDS (lithium 192 dodecyl sulfate) sample buffer containing 1% ß-mercaptoethanol before lysing at 95°C 193 and vortexing. 30 µL of each sample were run on NuPAGE 10% Bis-Tris protein gel 194 (Invitrogen) for a short time before staining with SafeStain (Thermo Fisher) and excising 195 the whole protein band. In-gel proteins were de-stained using 50% (v/v) ethanol, 50 mM 196 ammonium bicarbonate (ABC), before being reduced and alkylated for 5 min at 70°C using 197 10 mM TCEP (tris(2-carboxyethyl)phosphine) and 40 mM CAA (2-chloroacetamide), 198 respectively. After washing with 50% (v/v) ethanol 50 mM ABC, peptides were lysed 199 overnight using trypsin. Finally, peptides were extracted by sonication in a water bath (10 200 min at room temperature), concentrated using a Speed-Vac (50 mins) and resuspended 201 in 2.5% acetonitrile 0.05% formic acid. Extracted peptides were analysed by nanoLC-ESI-202 MS/MS using the Ultimate 3000/Orbitrap Fusion instrumentation (Thermo Scientific). The 203 UniProt proteome for *P. aeruginosa* strain PAO1 was used for peptide analysis. Further 204 data analysis was carried out using MaxQuant and Perseus software as described 205 previously; peptides without triplicate measures were filtered out (23). Comparative 206 proteomics data are presented in Supplementary Tables S1 and S3.

207

208 **Phylogenomics and metatranscriptomics analyses**

209 The protein sequences of PA3219, PA3218 and PA0842 were used to search genome 210 Pseudomonas sequences of clades in the JGI IMG genome portal 211 (https://img.jgi.doe.gov/). Note that the PA3218 protein is incorrectly annotated in the 212 genome of PAO1. The putative glycosyltransferase located immediately downstream of 213 PA3219 was manually inspected by aligning to the corresponding gene (PA14 22600) in 214 the genome of *P. aeruginosa* PA14. To identify PA3218 in misannotated *P. aeruginosa* 215 genomes, the nucleotide sequence immediately downstream of PA3219 was aligned with 216 agt1 in strain PA14. (locus tag PA14 22600). The phylogeny of Pseudomonas clades was 217 determined using the nucleotide sequences of six housekeeping genes (rpoB, rpoD, 218 dnaE. recA. atpD. gyrB) retrieved from each genome using IQ-Tree with the parameters -219 m TEST -bb 1000 -alrt 1000. The most suitable model was chosen by the software. 220 Evolutionary distances were inferred using maximum-likelihood analysis. Relationships 221 were visualised using the online platform the Interactive Tree of Life viewer 222 (https://itol.embl.de/). The conserved Pho box sequence was predicted using the MEME 223 server (24).

The metatranscriptomics datasets of sputum samples obtained from a CF patient 7-days (SRX5145606) and 8-days (SRX5145605) before death (25), CF patient G (SRR6833349) from Denmark (26) and a patient (SRR6833340) with chronic wound infection (26) were retrieved from the short reads archive (SRA) database (https://www.ncbi.nlm.nih.gov/sra). The reads were downloaded using fastq-dump and

229 mapped using the BBMap aligner as described previously (27). Briefly, the SRA reads 230 were mapped to the genome sequence of *P. aeruginosa* PAO1 using a stringent cut-off 231 of minid=0.97. Relative abundance data were compared using RPKM (reads per kilobase 232 of transcript, per million mapped reads). The list of RPKM abundance of individual genes 233 of *P. aeruginosa* PAO1 is shown in **Suppl. Table S4**.

- 234
- 235

236 **Results and discussion**

237 *P. aeruginosa* produces novel glycolipids in response to Pi stress

238 To determine changes in the membrane lipidome in response to P-stress, the model P. 239 aeruginosa strain PAO1 was grown in minimal medium under high (1 mM) or low Pi (50 240 µM) conditions (Figure 1a). The latter condition elicited strong alkaline phosphatase activity, measured through the liberation of *para*-nitrophenol (*pNP*) from *para*-nitrophenol 241 242 phosphate (pNPP) (Figure 1b), this being a strong indication that cells were P-stressed. 243 Analysis of membrane lipid profiles using high performance liquid chromatography 244 coupled to mass spectrometry (HPLC-MS) revealed the presence of several new lipids 245 under Pi stress conditions (Figure 1c). Thus, during Pi-replete growth (1 mM phosphate), 246 the lipidome is dominated by two glycerophospholipids: phosphatidylglycerol (PG, eluted 247 at 6.8 min) and phosphatidylethanolamine (PE, eluted at 12.2 min). During Pi-stress a lipid 248 species with mass to charge ratio (m/z) of 623 and 649 were also found, with MS 249 fragmentation resulting in a 131 m/z peak, a diagnostic ion for the amino-acid containing 250 ornithine lipid. This is consistent with previous reports of ornithine lipids in the P. 251 aeruginosa membrane in response to Pi stress (28-29).

Eurther to ornithine lipids, three unknown lipids eluting at 7.7 min, 8.7 min and 9.8 min, were only present under Pi stress conditions (**Figure 1c**). Using several rounds of MS fragmentation (MSⁿ), with a quadrupole ion trap MS, fragmentation patterns

255 characteristic of glycolipids were found for all three peaks. For each peak of interest, the 256 most predominant lipid masses of 774.7 m/z, 786.8 m/z and 788.6 m/z were analysed by 257 MSⁿ in positive ionisation mode (**Figure 1d**). In each case, an initial head group was lost 258 leaving a significant signal of 595.6 m/z, the mass of the glycolipid building block 259 diacylglycerol (DAG). Further fragmentation leads to the loss of either fatty acyl chain from 260 DAG, leaving monoacylglycerols of 313.2 m/z and 339.3 m/z. Two monoacylglycerols with 261 different masses are produced as a result of the original lipid containing 16:0 and 18:1 262 fatty acids (313.2 m/z and 339.3 m/z monoacylglycerols, respectively). To further elucidate 263 the identity of the peaks, a search for a neutral loss of a polar head group was carried out. 264 Thus, the intact masses of 774.7 m/z and 788.6 m/z in positive ionisation mode leads to 265 the loss of a head group of -179 and -193 m/z, which corresponds to a hexose- and a 266 glucuronate- group, respectively (Figure 1d), suggesting the occurrence of novel 267 monoglucosyldiacylglycerol (MGDG) and glucuronic acid diacylglycerol (GADG) 268 alvcolipids in *P. aeruginosa*. The third glycolipid peak at 8.7 min remains an unknown lipid 269 with intact mass of 786.8 m/z (hereafter designated as a putative unknown glycolipid, 270 UGL). Together, these data confirm the production of new glycolipids in *P. aeruginosa* in 271 response to Pi stress.

272

273 Comparative proteomics uncover the lipid renovation pathway in *P. aeruginosa*

To determine the proteomic response of *P. aeruginosa* to phosphorus limitation, and identify the genes involved in glycolipid formation, strain PAO1 was cultivated under high and low Pi conditions for 8 hours and the cellular proteome then analysed. A total of 2844 proteins were detected, 175 of which were found to be differentially regulated by Pi availability (**Figure 2b, Suppl. Table S1**). In line with previous transcriptomic studies of strain PAO1 (18), major phosphorus acquisition mechanisms were highly expressed under Pi stress conditions, *e.g.* the Pi-specific transporter PstSCAB, the two-component
 regulator PhoBR (Suppl. Table S1) (30).

282 Comparative proteomics also identified several genes which are likely important 283 for membrane lipid remodelling (Figure 2a) including PA3219 (4.6-fold increase under Pi-284 depleted conditions, FDR<0.01), encoding a putative phospholipase C protein, and 285 PA0842 (4-fold increase under Pi-depleted conditions, FDR<0.01), encoding a putative 286 glycosyltransferase (Figure 2b). PA3219 has 47% protein sequence identity to PIcP from 287 Phaeobacter sp. MED193 and 46% identity to PIcP from Sinorhizobium meliloti (25-27). 288 In these bacteria, PICP is essential in the lipid remodelling pathway for the formation of the 289 diacylglycerol (DAG) backbone, representing the essential intermediate for the production 290 of glycolipids (31-32). In P. aeruginosa PAO1, PA3219 appears to form an operon with 291 PA3218, a putative glycosyltransferase likely under the control of the PhoBR two 292 component system, as a highly conserved Pho box sequence was recognisable in the 293 promoter region (Figure 2c), PA3218 (hereafter referred to as Agt1) has 41% protein 294 sequence identity to the Agt of *Phaeobacter* sp. MED193. PA0842 showed 35% identity 295 to the Aqt of *Phaeobacter* sp. MED193 and a Pho box sequence is also found in its 296 promoter region. This corroborates the finding that the PA0842 protein (hereafter referred 297 to as Agt2) was significantly upregulated under Pi-depleted conditions (Figure 2b). In 298 summary, comparative proteomic analysis suggests that P. aeruginosa PAO1 adopts this 299 PIcP-Agt lipid remodelling pathway for the production of glycolipids in response to Pi-300 stress (Figure 2a).

301

The PIcP-Agt mediated lipid renovation pathway is strictly conserved in *P. aeruginosa* and actively transcribed in the metatranscriptomes of cystic fibrosis patients

305 To uncover how widespread this predicted PIcP-Agt lipid remodelling pathway is amongst 306 the genus *Pseudomonas*, including *P. aeruginosa* strains, we conducted a thorough 307 comparative genomics analysis of these lipid renovating loci. PICP-Agt is strictly conserved 308 in all 770 genome-sequenced *P. aeruginosa* strains in the IMG/M database, including all 309 three-previously recognised P. aeruginosa lineages (33-34), group 1 represented by strain 310 PAO1, group 2 represented by strain PA14 and group 3 represented by strain PA7 (Figure 311 **3, Suppl. Table S2**). Indeed, this remodelling pathway is prevalent in many *Pseudomonas* 312 groups, including the plant pathogen *Pseudomonas syringae*. To investigate whether the 313 PlcP-Aqt lipid remodelling pathway is involved in host-pathogen interactions, we analysed 314 metatranscriptomic datasets from cystic fibrosis patients, where *P. aeruginosa* is known 315 to be prevalent in the fatal exacerbation period before patient death (25). To the best of 316 our knowledge, only two studies have reported the metatranscriptome of the bacterial 317 community present in CF sputum (25, 26). Indeed, *phoBR* and *pstS* are amongst the most 318 highly expressed genes, confirming previous observations that P. aeruginosa is Pi-limited 319 during human airway epithelia infection (36, 37). Interestingly, the alkaline phosphatase 320 phoA (38) was highly expressed in sputum from CF patients but not from wound samples 321 which was also dominated by *P. aeruginosa*. Importantly, the transcripts of *P. aeruginosa* 322 agt1/plcP/agt2 are highly expressed in CF sputum during the fatal exacerbation period 323 before death (Figure 2d). Therefore, our phylogenomic and metatranscriptomic analyses 324 suggest that not only is the PIcP-Agt lipid remodelling pathway strictly conserved and 325 prevalent in *P. aeruginosa*, but also the corresponding genes are also highly expressed 326 during CF patient infection, suggesting a potential role for lipid renovation in host-327 pathogen interactions.

328

329 Experimental validation of the lipid renovation pathway for glycolipid formation in
 330 *P. aeruginosa*

331 To validate the function of these two putative glycosyltransferases (Agt1, Agt2) in the 332 formation of glycolipids, we synthesized the codon-optimized genes (PA3218 and 333 PA0842, respectively) for recombinant expression in *Escherichia coli*. The total lipidomes 334 from the recombinant E. coli strains were then analysed by HPLC-MS to determine the 335 presence of glycolipids in a gain-of-function assay. Expression of P. aeruginosa Agt1 336 (PA3218) was sufficient for the production of MGDG (eluted at 7.7 min) in E. coli, 337 confirmed through MSⁿ fragmentation (**Figure 4a**). No UGL nor GADG was observed in 338 the lipidome of this Agt1-overexpressing E. coli strain. Expressing Agt2 (PA0842) from P. 339 aeruginosa in E. coli was sufficient for the accumulation of the GADG glycolipid (eluted at 9.8 min), also confirmed through the MSⁿ fragmentation pattern (**Figure 4b**). Equally, no 340 341 UGL nor MGDG was observed in the Agt2-overexpressing E. coli strain. Production of 342 these glycolipids was not observed in the same strain of *E. coli* transformed with an empty 343 vector control (pET28a). It is therefore likely that UGL production is carried out by another 344 alvcosvltransferase, the identity of which remains to be discovered.

345 To confirm the role of Agt1 and Agt2 in the production of MGDG and GADG, we 346 purified Aqt1 and Aqt2 from recombinant E. coli (Figure 4c) and carried out enzyme 347 assays using UDP-glucose and UDP-glucuronic acid as the sugar donor and DAG as the 348 acceptor. Agt1 can only accept UDP-glucose as the substrate with an affinity of K_m=298.1 349 \pm 9.5 μ M (Figure 4c, middle panel) and produced MGDG as expected (Figure 4d, left 350 panel). Similarly, Agt2 can use UDP-glucuronic acid as the substrate (K_m = 373.0 ± 12.9 μ M (Figure 4c, right panel), producing the GADG lipid (Figure 4d, right panel). 351 352 Interestingly, the purified Agt2 enzyme can also use UDP-glucose to some extent with a 353 K_m of 480 μ M (data not shown) although the corresponding lipid MGDG was not observed 354 in the lipid extract from the lipidome of the recombinant host E. coli (Figure 4b).

355 To further confirm the role of these genes in *P. aeruginosa* glycolipid biosynthesis 356 we analysed the lipidomes of mutants in $\Delta plcP$, $\Delta agt1$ and $\Delta agt2$ in strain PAO1 (**Figure**

357 **5a**, **b**). Differences were analysed by searching for the intact masses of the glycolipids 358 MGDG and GADG: 774.6 m/z and 788.6 m/z in positive ionisation mode with an 359 ammonium adduct, respectively. As expected, under Pi stress MGDG is no longer 360 produced in the $\Delta agt1$ mutant and similarly GADG is no longer produced in the $\Delta agt2$ 361 mutant (**Figure 5a**). In the $\Delta plcP$ mutant, no MGDG was found and the GADG lipid was 362 significantly reduced but not entirely abolished (Figure 5b). The small amount of GADG 363 produced in the $\Delta p l c P$ mutant suggests that an alternative supply of DAG (independent of 364 the degradation of phospholipids by PIcP) is available in this mutant. Nevertheless, 365 lipidome analyses of the $\Delta plcP$, $\Delta aqt1$ and $\Delta aqt2$ mutants strongly supports the key role 366 of this PIcP-Agt pathway (Figure 2A) in lipid renovation in *P. aeruginosa*.

367

368 The protective role of glycolipids to antibiotic resistance in *Pseudomonas* 369 *aeruginosa*

370 To assess whether the growth of the glycolipid-deficient mutants ($\Delta p l c P$, $\Delta a q t 1$, $\Delta a q t 2$) 371 was affected by Pi stress, we grew the mutants in the defined minimal medium under high 372 and low Pi conditions. However, no significant difference in growth rates was found (Suppl. 373 Figure S1). The presence of glycolipids in the membrane may, however, have a profound 374 impact on the functioning of the membrane during Pi stress. For example, PG is an anionic 375 lipid with net negative charges whereas MGDG has a neutral charged sugar group. 376 Although a PG-to-GADG substitution may not necessarily change membrane charge (35), 377 it may affect membrane curvature and the packing density of lipids. Thus, subsequent 378 knock-on effects in membrane function might be expected (10). We therefore set out to 379 investigate whether membrane lipid composition may have an impact on antibiotic 380 resistance in *P. aeruginosa*. As cationic antimicrobial peptides directly interact with 381 bacterial cell membranes, we focused on the impact of lipid remodelling on the killing 382 activity of polymyxin B. We conducted the analyses under P-depleted conditions, since

Pi-stress is clinically important, already known to induce the expression of virulence factors (15, 17, 18, 29), and our own analysis confirmed that an array of genes involved in phosphate acquisition and lipid remodelling in *P. aeruginosa* are indeed highly expressed in sputum samples from lung microbiome of CF patients (**Figure 2d**). Polymyxins represent the drug-of-last resort for effectively treating carbapenem-resistant *P. aeruginosa* infections (3, 39).

389 To test the sensitivity of the mutants in the PIcP-Agt pathway to polymyxin B, we 390 compared WT and mutants using kill curve analyses as the typically used disk diffusion 391 method does not work efficiently for cationic antimicrobials (40). Indeed, there was a 392 significant decrease in the survival of all three PAO1 glycolipid synthesis mutants ($\Delta p l c P$, 393 $\Delta agt1$ and $\Delta agt2$) compared to the wild type when challenged with polymyxin B, 394 suggesting a protective role of glycolipids in polymyxin B resistance (Figure 5c). Such a 395 protective role of glycolipids in polymyxin B resistance was not observed for other 396 antibiotics, including ciprofloxacin, gentamicin, ceftazidime and meropenem (data not 397 shown). P. aeruginosa is known to enhance its resistance to polymyxins through 398 decoration of its lipopolysaccharide (LPS) layer using either 4-amino-4-deoxy-L-arabinose 399 (L-Ara4N) by arnB (5), or the addition of phosphoethanolamine (pEtN) by eptA (6). It is 400 thought that these changes perturb the electrostatic interaction between the cationic 401 polymyxin B and the normally negatively charged LPS. To investigate whether these 402 mechanisms play a role in the glycolipid-deficient mutants, we conducted a comparative 403 proteomics analysis of the $\Delta p l c P$ mutant and WT under Pi depleted conditions, which 404 revealed only a small number of differentially expressed proteins (Suppl. Table S3). The 405 majority of these differentially expressed proteins are uncharacterised. However, 406 importantly, LPS modification enzymes previously found to confer antimicrobial peptide 407 resistance, such as ArnB and EptA, were not differentially expressed between the WT and 408 $\Delta plcP$ mutant. Therefore, our data suggests that it is the glycolipids that are the major contributor to increased polymyxin B resistance, which constitutes a new biological
mechanism for polymyxin resistance. To this end, we tested the resistance to polymyxin
B of recombinant *E. coli* strains overexpressing *P. aeruginosa* Agt1 and Agt2, that produce
MGDG and GADG, respectively (Figure 4a, 4b). Indeed, in this gain-of-function assay,
both Agt1 and Agt2 overexpressing *E. coli* strains had enhanced resistance to polymyxin
B compared to the empty vector control (Figure 5d), supporting the protective role of
these glycolipids to antimicrobial peptides.

416

417 To conclude, we present here the discovery of novel alycolipids produced in P. 418 aeruginosa during adaption to phosphorus stress. This lipid renovation pathway is strictly 419 conserved in all P. aeruginosa isolates to date and highly expressed in the 420 metatranscriptome of CF patients, suggesting a key role of lipid remodelling in the 421 ecophysiology of this bacterium. Interestingly, lipid remodelling as a response to survive 422 phosphorus stress in turn comes with trade-offs in terms of antibiotic resistance: these 423 glycolipids may protect the bacterium from insult by cationic antimicrobial peptides, 424 highlighting a new resistance mechanism to polymyxin B which has been previously 425 overlooked. It remains to be seen whether the altered susceptibility to polymyxin B is the 426 sole trade-off following lipid remodelling of phospholipids to glycolipids. After all, evolution 427 appears to have selected phospholipids as the dominant lipids in the last universal 428 common ancestor (12).

429

430 Acknowledgements

This work was funded by an MRC Doctoral Training Partnership studentship in
Interdisciplinary Biomedical Research (MR/J003964/1) awarded to RAJ and by a Royal
Society International Exchanges 2017 Cost Share (China) award (IEC\NSFC\170213; grant
agreement no. 170213). AM and YC are supported by a European Research Council (ERC)

- 435 award under the European Union's Horizon 2020 research and innovation programme
- 436 (grant agreement no. 726116). We also thank the Proteomics Research Technology
- 437 Platform, University of Warwick, UK for their contribution.
- 438
- 439 **Competing interest**
- 440 The authors declare no competing interests.

441 References

- 442
 442
 443
 443 Murray, T. S., Egan, M., & Kazmierczak, B. I. (2007). *Pseudomonas aeruginosa* 443 chronic colonization in cystic fibrosis patients. *Curr. Opin. Pediatr.*, 19: 83-88.
- 444 **2.** Gaynes, R., Edwards, J. R., & System, N. N. I. S. (2005). Overview of nosocomial 445 infections caused by Gram-negative bacilli. *Clin. Infect. Dis.*, 41:848–854.
- 446
 447
 447 BMJ. 344:e3236.
- 448
 448
 448
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
 449
- 450 **5.** Chung, E. S., Lee, J. Y., Rhee, J. Y., & Ko, K. S. (2017). Colistin resistance in
 451 *Pseudomonas aeruginosa* that is not linked to *arnB. J. Med. Microbiol.*, 66:833–
 452 841.
- 453
 6. Nowicki, E. M., O'Brien, J. P., Brodbelt, J. S., & Trent, M. S. (2015). Extracellular
 454 zinc induces phosphoethanolamine addition to *Pseudomonas aeruginosa* lipid A
 455 via the CoIRS two-component system. *Mol. Microbiol.* 97:166–178.
- 456 **7.** Parsons JB, Rock CO. (2013) Bacterial lipids: metabolism and membrane
 457 homeostasis. *Prog. Lipid Res.* 52:249-276.
- 458 8. Zhang, Y.-M., & Rock, C. O. (2008). Membrane lipid homeostasis in bacteria. *Nat.*459 *Rev. Microbiol.* 6:222–233.
- 460 9. van Meer G, Voelker DR, Feigenson GW. (2008) Membrane lipids: where they are
 461 and how they behave. *Nat. Rev. Mol. Cell Biol.* 9:112-124.
- 462 **10.** Harayama T, Riezman H. (2018) Understanding the diversity of membrane lipid
 463 composition. *Nat. Rev. Mol. Cell Biol.* 19:281-296.
- 464 **11.** May, K.L., & Silhavy T.J. (2017) Making a membrane on the other side of the wall.
 465 *Biochim. et Biophy. Acta.* 1862:1386-1393.
- 466
 467
 467 membranes and the three domains of life. *Nat. Rev. Microbiol.* 10:507-515.
- 468
 468
 469
 469 membrane evolution. *Trends Biochem. Sci.* 29:469-477.
- 470
 470
 471 Agradation and lipid cycles in microbes. In *Aerobic Utilization of Hydrocarbons,*472 *Oils, and Lipids*, 10.1007/978-3-319-50418-6_38
- 473 **15.** Lamarche MG, Wanner BL, Crépin S, Harel J. (2008) The phosphate regulon and
 474 bacterial virulence: a regulatory network connecting phosphate homeostasis and
 475 pathogenesis. *FEMS Microbiol. Rev.* 32:461-473.

476
 16. Long, J., Zaborina, O., Holbrook, C., Zaborin, A., & Alverdy, J. (2008). Depletion of
 477 intestinal phosphate after operative injury activates the virulence of *P. aeruginosa* 478 causing lethal gut-derived sepsis. *Surgery*, *144*:189–197.

- 479 **17.** Francis VI, Stevenson EC, Porter SL. (2017) Two-component systems required for
 480 virulence in *Pseudomonas aeruginosa. FEMS Microbiol. Lett.* 364(11). doi:
 481 10.1093/femsle/fnx104.
- 482
 483 **18.** Bains, M., Fernández, L., & Hancock, R. E. W. (2012). Phosphate starvation
 483 promotes swarming motility and cytotoxicity of *Pseudomonas aeruginosa*. *Appl.*484 *Environ. Microbiol.*, 78:6762–6768.
- 485 **19.** Son, M. S., Matthews, W. J., Kang, Y., Nguyen, D. T., & Hoang, T. T. (2007). *In*486 *vivo* evidence of *Pseudomonas aeruginosa* nutrient acquisition and pathogenesis
 487 in the lungs of cystic fibrosis patients. *Infect. Immun.* 75:5313–5324.
- 20. Lidbury, I. D. E. A., Murphy, A. R. J., Scanlan, D. J., Bending, G. D., Jones, A. M.
 E., Moore, J. D., et al., (2016). Comparative genomic, proteomic and exoproteomic
 analyses of three *Pseudomonas* strains reveals novel insights into the phosphorus
 scavenging capabilities of soil bacteria. *Environ. Microbiol.* 18:3535–3549.
- 492 **21.** Sebastián, M., Smith, A. F., González, J. M., Fredricks, H. F., Van Mooy, B.,
 493 Koblížek, M., et al., (2016). Lipid remodelling is a widespread strategy in marine
 494 heterotrophic bacteria upon phosphorus deficiency. *ISME J.* 10:968–978.
- 495
 495
 496
 496
 496
 497
 497
 498
 498
 498
 499
 499
 490
 490
 491
 491
 492
 493
 494
 494
 495
 495
 496
 497
 497
 497
 498
 498
 499
 499
 499
 490
 490
 491
 491
 491
 492
 493
 494
 494
 495
 495
 496
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
 497
- 498
 498
 498
 499 Proteomics insights into the *Burkholderia cenocepacia* phosphorus stress
 500 response. doi.org/10.1111/1462-2920.15451.
- 501 24. Bailey TL, Elkan C (1994) Fitting a mixture model by expectation maximization to
 502 discover motifs in biopolymers. *Proceedings of the Second International* 503 *Conference on Intelligent Systems for Molecular Biology*, 28-36, AAAI Press,
 504 Menlo Park, Californias.
- 505 25. Cobián Güemes AG, Lim YW, Quinn RA, Conrad DJ, Benler S, Maughan H,
 506 Edwards R, et al. (2019) Cystic fibrosis rapid response: translating multi-omics data
 507 into clinically relevant information. *mBio.* 16;10(2):e00431-19.
- 508 26. Cornforth DM, Dees JL, Ibberson CB, Huse HK, Mathiesen IH, Kirketerp-Moller K,
 509 et al., (2018) *Pseudomonas aeruginosa* trascriptome during human infection. *Proc.* 510 *Nat. Acad. Sci. USA* 115:E5125-5134.

27. Jones, H.J., Krober, E., Stephenson, J., Mausz M.A., Jameson, E., Millard, A.,
Purdy, K.J., Chen, Y. (2019) A new family of uncultivated bacteria involved in
methanogenesis from the ubiquitous osmolyte glycine betaine in coastal saltmarsh
sediments. *Microbiome*. 7, 120. doi.org/10.1186/s40168-019-0732-4

- 28. Kim, S. K., Park, S. J., Li, X. H., Choi, Y. S., Im, D. S., & Lee, J. H. (2018). Bacterial
 ornithine lipid, a surrogate membrane lipid under phosphate-limiting conditions,
 plays important roles in bacterial persistence and interaction with host. *Environ. Microbiol.* 20:3992–4008.
- 29. Lewenza, S., Falsafi, R., Bains, M., Rohs, P., Stupak, J., Sprott, G. D., & Hancock,
 R. E. W. (2011). The *olsA* gene mediates the synthesis of an ornithine lipid in *Pseudomonas aeruginosa* during growth under phosphate-limiting conditions, but
 is not involved in antimicrobial peptide susceptibility. *FEMS Microbiol. Lett.* 320:
 95–102.
- 30. Wilton M, Halverson TW, Charron-Mazenod L, Parkins MD, Lewnza S. (2018)
 Secreted phosphatase and deoxyribonuclease are required by *Pseudomonas aeruginosa* to defend against neutrophil extracellular traps. *Infect. Immun* 86:e00403-18.
- 31. Wei, T., Quareshy, M., Zhang, Y. Z., Scanlan, D. J., & Chen, Y. (2018). Manganese
 is essential for PlcP metallophosphoesterase activity involved in lipid remodeling
 in abundant marine heterotrophic bacteria. *Appl. Environ. Microbiol.* 84, e0110918.
- 32. Zavaleta-Pastor, M., Sohlenkamp, C., Gao, J.-L., Guan, Z., Zaheer, R., Finan, T.
 M., et al., (2010). *Sinorhizobium meliloti* phospholipase C required for lipid
 remodeling during phosphorus limitation. *Proc. Nat. Acad. Sci. USA107*:302–307.
- 33. Freschi L, Jeukens J, Kukavica-Ibrulj I, Boyle B, Dupont MJ, Laroche J, et al.,
 (2015) Clinical utilization of genomics data produced by the international *Pseudomonas aeruginosa* consortium. *Front Microbiol.* 29;6:1036. doi:
 10.3389/fmicb.2015.01036.
- 34. Ozer EA, Nnah E, Didelot X, Whitaker RJ, Hauser AR. (2019) The population
 structure of *Pseudomonas aeruginosa* is characterized by genetic isolation of
 exoU+ and *exoS*+ lineages. *Genome Bio.I Evol.* 11:1780-1796.
- 542 **35.** Diercks H, Semeniuk A, Gisch N, Moll H, Duda KA, Hölzl G. (2015) Accumulation
 543 of novel glycolipids and ornithine lipids in *Mesorhizobium loti* under phosphate
 544 deprivation. *J. Bacteriol.* 197:497-509.
- 545 **36.** Chugani, SA and Greenberg, EP (2007) The influence of human respiratory

546 epithelia on *Pseudomonas aeruginosa* gene expression. *Microb. Pathog.* 42: 29–
547 35.

- 548 **37.** Frisk, A, Schurr, JR, Wang, G, Bertucci, DC, Marrero, L, Hwang, SH, et al. (2004)
 549 Transcriptome analysis of *Pseudomonas aeruginosa* after interaction with human
 550 airway epithelial cells. *Infect. Immun.* 72: 5433–5438.
- 38. Filloux A, Bally M, Soscia C, Murgier M, Lazdunski A (1988) Physhate regulation
 in *Pseudomonas aeruginosa*: Clonning of the alkaline phosphatase gene and
 identification of *phoB* and *phoR*-like gnes. *Mol. Gen. Genet*. 212(3):510-513.
- 39. Poirel L, Jayol A, Nordmann P. (2017) Polymyxins: antibacterial activity,
 susceptibility testing, and resistance mechanisms encoded by plasmids or
 chromosomes. *Clin. Microbiol. Rev.* 30:557-596.
- 40. Ezadi, F., Ardebili A., Mirnead R (2019) Antimicrobial susceptibility testing for
 polymyxins: challenges, issues, and recommendations. *J. Clin. Microbiol.* 57,
 e01390-18.
- 41. Monds RD, Newell PD, Schwartzman JA, O'Toole GA. (2006) Conservation of the
 Pho regulon in *Pseudomonas fluorescens* Pf0-1. *Appl. Environ. Microbiol.* 72:1910-1924.
- 42. Semeniuk A, Sohlenkamp C, Duda K, Hölzl G. (2014) A bifunctional
 glycosyltransferase from *Agrobacterium tumefaciens* synthesizes monoglucosyl
 and glucuronosyl diacylglycerol under phosphate deprivation. *J. Biol. Chem.*289(14):10104-10114.
- 43. Minh BQ, Schmidt HA, Chernomor O, Schrempf D, Woodhams MD, von
 Haeseler A, et al. (2020) IQ-TREE 2: New models and efficient methods for
 phylogenetic inference in the genomic era. *Mol. Biol. Evol.* 37:1530–1534.

| 5 | 7 | 1 |
|---|---|---|
| J | 1 | L |

Figure legends

572 Figure 1 Lipidomics analysis uncovers novel glycolipid formation in *Pseudomonas*

573 *aeruginosa* strain PAO1 in response to phosphorus limitation.

a) Growth of strain PAO1 WT in minimal medium A containing 1 mM phosphate (+Pi, blue)
or 50 µM phosphate (-Pi, black) over 12 hours. Data are the average of 3 independent
replicates.

577 **b)** Liberation of *para*-nitrophenol (*p*NP) from *para*-nitrophenol phosphate (*p*NPP) through 578 alkaline phosphatase activity, under Pi-repleted (1 mM, black) and Pi-depleted (50 μ M, 579 yellow) conditions. Error bars represent the standard deviation of three independent 580 replicates.

581 c) Representative chromatograms in negative ionisation mode of the P. aeruginosa 582 lipidome when grown under phosphorus stress (-Pi, black) compared to growth under 583 phosphorus sufficient conditions (+Pi, orange). PG, phosphatidylglycerol; PE, 584 phosphatidvlethanolamine. OL. ornithine lipids. Lower panel: extracted ion 585 chromatograms of three new glycolipid species in *P. aeruginosa* which are only produced 586 during Pi-limitation (black, 1 mM; orange, 50 µM). MGDG: monoglucosyldiacylglycerol, 587 GADG: glucuronic acid-diacylglycerol and UGL: unconfirmed glycolipid.

d) Mass spectrometry fragmentation spectra of three glycolipid species present under Pi stress in *P. aeruginosa*, at retention times of 7.7 (*m/z* 774.7), 8.7 (*m/z* 786.7) and 9.8 (*m/z* 788.6) minutes, respectively. Each spectrum depicts an intact lipid mass with an ammonium (NH₄⁺) adduct exhibiting neutral loss of a head group, yielding diacylglycerol (DAG) (595 *m/z*). Further fragmentation yields monoacylglycerols (MAG) with C16:0 or C18:1 fatty acyl chains.

595 Figure 2 Comparative multi-omic analyses for the identification of the PIcP-Agt 596 pathway responsible for glycolipid formation in *Pseudomonas aeruginosa* strain 597 PAO1.

a) The proposed pathway for lipid remodelling through the PlcP-Agt pathway. PlcP degrades membrane phospholipids such as PG, to generate diacylglycerol (DAG) intermediates for the formation of MGDG and GADG through the activity of glycosyltransferases, using either UDP-glucose or UDP-glucuronate as the co-substrate (42).

b) Volcano plot depicting differentially expressed proteins when comparing Pi-repleted and Pi-depleted conditions. Significantly upregulated proteins when under Pi stress are shown in red (left), and those that are significantly upregulated when Pi is sufficient are in green (right). Significance was accepted when the false discovery rate (FDR) was < 0.05, and a fold change \geq 2.

608 c) Genomic organisation of predicted lipid remodelling genes in *P. aeruginosa*. 609 Glycosyltransferases (orange) PA3218 (Agt1) and PA0842 (Agt2) are predicted to be 610 involved in glycolipid synthesis. PA3219 is predicted to be PlcP in *P. aeruginosa*. 611 Predicted Pho box sequences in the promoter regions (represented in blue boxes) of each 612 glycosyltransferase operon from *P. aeruginosa* strains representing the PAO1 clade, the 613 PA7 clade and the PA14 clade are shown. The black dots represent residues which are 614 conserved in the Pho box consensus CTGTCATNNNNCTGTCAT (41).

d) Metatranscriptomic analysis of PlcP-Agt lipid remodelling genes in sputum samples from a cystic fibrosis patient 7-days (CF_D-7) and 8-days (CF_D-8) before death (25) and a Danish CF patient (CF_Person G) (26) as well as a wound sample from a burns patient from the USA (Burn patient) (26). Relative abundance is expressed as RPKM (reads per

kilobase of transcript, per million mapped reads). PhoA (PA3296) encodes an alkalinephosphatase (38).

621

622 Figure 3 Occurrence of plcP-agt genes in major Pseudomonas groups. The 623 phylogeny of *Pseudomonas* clades was determined using the nucleotide sequences of six 624 housekeeping genes (rpoB, rpoD, dnaE, recA, atpD, gyrB) retrieved from each genome 625 using IQ-Tree (43). The filled colour indicates the presence of the genes in the genomes 626 whereas a blank indicates the absence of the corresponding gene in the genomes. The 627 two-component system PhoBR (black circles) is found in all genomes and the PlcP-628 Agt1/Agt2 are strictly conserved in all 770 genome-sequenced P. aeruginosa strains that 629 form three clades represented by strain PA14, PA01 and PA7 respectively. Bootstrap 630 values >75% are shown.

631

Figure 4, Characterization of glycolipid formation from recombinant Agt1 and Agt2. a) Extracted ion chromatogram of the MGDG lipid from recombinant *E. coli* expressing Agt1. An empty vector control is also shown (red line). The identity of MGDG is further validated using mass spectrometry fragmentation showing the neutral loss of 179 corresponding to the loss of glucose and the formation of monoacylglycerols (MAG) with C16:0 or C18:1 (m/z 313.2, 339.3).

b) Extracted ion chromatogram of the GADG lipid from recombinant *E. coli* expressing
Agt2. An empty vector control is also shown (red line). The identity of GADG is further
validated using mass spectrometry fragmentation showing the neutral loss of 193
corresponding to the loss of glucose and the formation of monoacylglycerols (MAG) with
C16:0 or C18:1 (*m/z* 313.2, 339.2).

643 **c)** Purified Agt1 and Agt2 protein from recombinant *E. coli* (left panel) and Michaelis 644 Menten kinetics of Agt1 towards UDP-glucose (middle panel) and Agt2 towards UDP-645 glucuronic acid (right panel) as substrate, respectively.

d) Mass spectrometry identification of MGDG and GADG produced from purified Agt1 and
Agt2 using DAG and UDP-glucose and UDP-glucuronic acid as the substrate,
respectively.

649

Figure 5 Glycolipid formation in *P. aeruginosa* and mutants under Pi stress, showing a protective role of glycolipids to polymyxin B.

652 a-b) Relative abundance of the glycolipid MGDG (a) and GADG (b) in P. aeruginosa mutants plcP (purple trace), agt1 (blue trace) and agt2 (green, trace) compared to the wild 653 654 type (WT). Cells were cultivated under low Pi conditions (50 µM) and a representative 655 extracted ion chromatogram of MGDG/GADG is shown between the WT, (black trace) and 656 each mutant. The right most panel shows the abundance of MGDG or GADG calculated 657 relative to an internal lipid standard d17:1/12:0 sphingosylphosphoethanolamine (Sigma-658 Aldrich) in the wild-type and mutant strains of *P. aeruginosa*. Values are calculated from 659 three biological replicates and the error bars denote standard deviation. MGDG, 660 monoglucosyldiacylglycerol, GADG, glucuronic acid-diacylglycerol.

661 **c)** Survival of glycolipid remodelling mutants $\Delta plcP$ (purple), $\Delta agt1$ (green) and $\Delta agt2$ 662 (blue) when challenged with 4 µg mL⁻¹ polymyxin B compared to WT under Pi stress 663 (black). All experiments were conducted under Pi stress conditions and the results are the 664 average of three biological replicates; error bars denote standard deviation.

d) Survival of glycolipid producing *Escherichia coli* when challenged with 20 μg mL⁻¹
 polymyxin B. All experiments were conducted in three replicates and error bars denote

- 667 standard deviation. Black, E. coli containing the empty vector pET28a; green, E. coli
- 668 containing plasmid pET28a-Agt1; blue, *E. coli* containing plasmid pET28a-Agt2.

| 670 | Supplementary Figures and Tables |
|-----|---|
| 671 | |
| 672 | Suppl. Figure S1 Growth of the <i>P. aeruginosa</i> mutants in the minimal medium under Pi |
| 673 | repleted and Pi depleted conditions. |
| 674 | |
| 675 | Suppl. Table S1 Proteomic analysis of differentially expressed proteins in the wild-type |
| 676 | <i>P. aeruginosa</i> PAO1 in response to different Pi levels (1 mM versus 50 μ M). |
| 677 | |
| 678 | Suppl. Table S2 Protein BLAST identification of locus tags homologous to agt1 and agt2 |
| 679 | glycolipid synthesis genes in all genome sequenced P. aeruginosa strains at the JGI IMG |
| 680 | database. |
| 681 | |
| 682 | Suppl. Table S3 Proteomic analysis of wild type <i>P. aeruginosa</i> PAO1 versus the $\Delta plcP$ |
| 683 | mutant grown at 50 μM Pi. |
| 684 | |
| 685 | Suppl. Table S4 Whole genome mapping to Pseudomonas aeruginosa PAO1 of |
| 686 | metatranscriptomic datasets (SRX5145605, SRX5145606, SRR6833349) from sputum |
| 687 | samples taken from cystic fibrosis patients and a patient with chronic wound infection |
| 688 | (SRR6833340). S4A (SRX5145605), S4B (SRX5145606), S4C (SRR6833349), S4D |
| 689 | (SRR6833340). RPKM, reads per kilobase of transcript per million mapped reads; FPKM, |
| 690 | fragments per kilobase of transcript per million mapped reads. |
| | |
| | |





phoB phoR pstS phoA plcP agt1 agt2 rpoB









0.2





GADG in P. aeruginosa PAO1 and mutants

