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Penicillin V acylases from gram-negative bacteria degrade N-acylhomoserine lactones and attenuate virulence in Pseudomonas aeruginosa — Source link

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- 1 Penicillin V acylases from Gram-negative bacteria degrade N-acylhomoserine
- 2 lactones and attenuate virulence in Pseudomonas aeruginosa
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- 9 **Running head:** Penicillin V acylases as quorum quenching agents
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Abstract: 28 29 Virulence pathways in Gram-negative pathogenic bacteria are regulated by quorum-sensing mechanisms, through the production and sensing of N-acylhomoserine lactone (AHL) signal 30 molecules. Enzymatic degradation of AHLs leading to attenuation of virulence (quorum 31 quenching) could pave the way for the development of new antibacterials. Penicillin V acylases 32 (PVAs) belong to the Ntn hydrolase superfamily, together with AHL acylases. PVAs are 33 exploited widely in the pharmaceutical industry, but their role in the natural physiology of their 34 native microbes is not clearly understood. This report details the characterization of AHL 35 36 degradation activity by homotetrameric PVAs from two Gram-negative plant pathogenic bacteria, Pectobacterium atrosepticum (PaPVA) and Agrobacterium tumefaciens (AtPVA). Both 37 the PVAs exhibited substrate specificity for degrading long chain AHLs. Exogenous addition of 38 39 these enzymes into Pseudomonas aeruginosa greatly diminished the production of elastase and pyocyanin, biofilm formation and increased the survival rate in an insect model of acute 40 infection. Subtle structural differences in the PVA active site that regulate specificity for acyl 41 42 chain length have been characterized, which could reflect the evolution of AHL-degrading acylases in relation to the environment of the bacteria that produce them and also provide 43 strategies for enzyme engineering. The potential for using these enzymes as therapeutic agents in 44 clinical applications and a few ideas about their possible significance in microbial physiology 45 have also been discussed. 46 **Keywords:** 47 48 Penicillin V acylase, N-acylhomoserine lactone acylase, Ntn hydrolase, quorum quenching, pathogenesis 49 50 51 52 53 54

Introduction:

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Penicillin acylases are microbial enzymes that cleave the amide bond of natural penicillins 56 (Arroyo et al. 2003), finding industrial application in the manufacture of the pharmaceutical 57 intermediate 6-aminopenicillanic acid (6-APA). Penicillin acylases can show substrate 58 59 preference for benzyl penicillin (Pen G, PGAs) or phenoxymethyl penicillin (Pen V, PVAs). Although both enzymes belong to the Ntn hydrolase superfamily (Oinonen and Rouvinen 2000), 60 they differ in their catalytic N-terminal nucleophile residue (PGA-ser, PVA-cys) and their 61 subunit composition. While PGAs are heterodimeric enzymes, PVAs are homotetramers and are 62 evolutionarily related to bile salt hydrolases (BSHs) that deconjugate bile salts in the mammalian 63 64 gut (Kumar et al. 2006) forming the cholylglycine hydrolase (CGH) group. A recent study (Panigrahi et al. 2014) has explored the phylogenetic clustering of CGHs from Gram-positive 65 and Gram-negative bacteria into two different groups. 66 Quorum sensing (QS) allows the bacteria to perceive their population density (Rutherford and 67 Bassler 2012) through the secretion of auto-inducer signal molecules and modulate gene 68 expression to trigger specific metabolic pathways. QS has been linked to bioluminescence, 69 bacterial virulence and swarming motility among other physiological processes (Li and Nair 70 71 2012). Bacterial pathogens including Pseudomonas aeruginosa, Vibrio cholerae and Acinetobacter baumanii use QS to regulate virulence genes and formation of biofilms, thereby 72 73 increasing their persistence (Li and Tian 2012). Gram-negative proteobacteria use autoinducers 74 N-acylhomoserine lactones (AHLs) (Churchill and Chen 2011), with a homoserine lactone ring linked via an amide bond to an acyl side chain (C₄-C₁₈) which may be saturated or unsaturated, 75 or with a hydroxy, oxo or no substituent on the carbon at the 3-position of the N-linked acyl 76 77 chain. Synthesized AHLs diffuse into neighbouring cells, where they modulate gene expression 78 through binding to the LuxR family of regulators. While Pectobacterium carotovorum and Agrobacterium tumefaciens produce 3-oxo-C₆ and 3-oxo-C₈-HSLs, respectively (Uroz et al. 79 80 2009), P. aeruginosa utilizes C₄ and 3-oxo-C₁₂-HSLs as signals for auto-induction. Bacteria in mixed-species communities have also been known to respond to structurally related non-cognate 81 82 AHLs produced by other bacteria (Winson et al. 1998). The disruption of AHL-directed signaling (termed "quorum quenching", QQ) through inhibition 83 or enzymatic degradation is an attractive strategy for controlling bacterial pathogenesis and 84

85	biofilm formation (Dong et al. 2007). Enzymes that degrade AHL include lactonases (ring
86	cleavage) and acylases (amide bond cleavage), which have been characterized from a variety of
87	bacteria. An exhaustive list has been provided by Grandclément et al. (2016). Penicillin acylases
88	are known to share similar structural fold and mechanistic features with AHL acylases, and the
89	probability of substrate cross-reactivity has been suggested earlier (Kreszlak et al. 2007).
90	Although recent studies have demonstrated activity of Kluyvera citrophila PGA (Mukherji et al.
91	2014) and aliphatic penicillin acylase from <i>Streptomyces lavendulae</i> (Torres-Bacete et al. 2015)
92	on AHLs, both these enzymes are ser-Ntn hydrolases with heterodimeric structure. A new AHL
93	acylase from P. aeruginosa (HacB) (Wahjudi et al. 2011) cleaves Pen V to a small extent;
94	however, AHL degradation by PVA enzymes or any other cys-Ntn hydrolase has not been
95	explored in detail so far. Moreover, the role of PVAs in microbial physiology is not been clearly
96	understood till date, but a few possible links to quorum sensing and pathogenesis have been
97	suggested (Avinash et al. 2016b).
98	In earlier reports, we have characterized the unique biochemical (Avinash et al. 2015) and
99	structural (Avinash et al. 2016a) features of a highly active PVA from the Gram-negative
100	Pectobacterium atrosepticum (PaPVA). The present study describes the characterization of PVA
101	from another related plant pathogen A. tumefaciens (AtPVA, 62% sequence identity with
102	PaPVA) and elucidates the subtle structural differences between the enzymes. Further, we report
103	the promiscuous deacylation of AHLs by these PVAs, and explore the structural interactions
104	involved in AHL binding. The application of PVA enzymes also led to reduction in QS-
105	regulated biofilm formation in <i>P. aeruginosa</i> PAO1 culture and the attenuation of <i>P. aeruginosa</i>
106	virulence in Galleria mellonella infection models, making them attractive options for novel QQ-
107	based therapeutic formulations.
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Bacterial strains and plasmids:

Materials and Methods:

The bacterial strains and plasmids used in this study are listed in Table 1. E. coli DH5a and BL21

star strains were maintained on Luria-Bertani (LB) medium supplemented on appropriate

113	antibiotics and cultured at 37°C. Antibiotics were added (100 μ g/ml ampicillin, 35 μ g/ml					
114	kanamycin or 10 μg/ml tetracycline) as required.					
115	Preparation of AtPVA and PaPVA:					
116	The pva gene from A. tumefaciens (GenBank GI:159185562) was cloned in pET22b vector					
117	between NdeI and XhoI restriction sites using the primers AtuF					
118	(gcttgacatatgtgcacgcgtttcgtttatatag) and AtuR (ctgaatctcgagaaagcccgagaaacttgaaag), and					
119	expressed in E. coli BL21 star cells with a C-terminal His-tag. The enzyme was purified to					
120	homogeneity using a HIS Select Ni ²⁺ affinity column (Sigma) and ENrich TM 650 (BioRad) size					
121	exclusion column. The protein was dialyzed against 10 mM Tris-Cl buffer pH 7.4 containing 100					
122	mM NaCl and 1mM DTT and stored in aliquots at -20°C. PaPVA was purified from					
123	recombinant E. coli as described earlier (Avinash et al. 2015).					
124	PVA enzyme activity assay					
125	Pen V hydrolysis activity was estimated by studying the formation of Schiff's conjugate with the					
126	product 6-APA and p-dimethyl amino benzaldehyde (Shewale et al. 1987). One unit (IU) of					
127	enzyme activity was defined as the amount of enzyme producing 1 μ mol 6-APA in 1 min.					
128	Biochemical characterization of AtPVA					
129	The Pen V hydrolysis activity was assayed at different pH (4-9) and temperatures (20-70°C) to					
130	ascertain the optimum conditions. AtPVA stability was studied by incubating the protein in 10					
131	mM Tris-Cl buffer pH 7.4 for 2 h at different temperatures between 30-90°C, and assaying for					
132	PVA activity at 45°C after different time intervals. Effect of pH on enzyme stability was studied					
133	by incubating the protein in 100 mM buffers of different pH (1-11) for 4 h at 25°C and assaying					
134	the residual activity. Kinetic parameters were determined by assaying the enzyme activity with					
135	increasing concentrations (5-240 mM) of penicillin V (potassium salt, Sigma) as substrate. The					
136	data were fitted using non-linear regression as detailed for <i>PaPVA</i> earlier (Avinash et al. 2015).					
137	AtPVA crystallization and structure determination					
138	Crystallization trials were set up with AtPVA (15 mg ml ⁻¹) using the sitting drop vapour diffusion					
139	technique. The protein crystallized in the 0.1M HEPES pH 7.5 and 12% PEG 3350 condition of					

140 the PEG Rx crystallization screen (Hampton Research, USA). The crystals were frozen in liquid nitrogen with 25% (w/v) 2, 5-hexanediol as cryoprotectant. Diffraction data were collected at 141 2.8 Å resolution at the SSRL-BL12-2 beamline at the Stanford Synchrotron Light Source (USA). 142 Investigation and scaling of the diffraction data was performed using XDS (Kabsch 2010) and 143 144 SCALA (Evans 2006). The AtPVA structural model was built using molecular replacement on Phaser ver. 2.5.6 (McCoy et al. 2007) and Autobuild (Phenix), with the refined structure of 145 PaPVA (PDB ID: 4WL2) as the template. Further model building and refinement was done 146 147 using Coot and Refmac5 (CCP4 software suite) respectively. AtPVA crystallized in P2₁2₁2₁ space group with a single tetramer per asymmetric unit (Table S1, Online Resource 1). 148 Bioluminescence assay for detection of AHL degradation 149 AHL degradation activity was monitored by employing biosensors that exhibit luminescence in 150 the presence of specific AHLs (Winson et al. 1998). Quenching of luminescence levels can be 151 used as an indication of AHL hydrolysis by the acylase enzymes (Steindler and Venturi 2007). 152 153 0.5 µL of 5 mM AHL stock solution in DMSO was spotted to a flat-bottom µClear white microplate (Greiner Bio-One) and dissolved in 50 µL reaction mixture containing 5 µg enzyme 154 155 in 100 mM NaCl, 1 mM DTT and 25 mM Tris HCl buffer pH 7.4 (for AtPVA) or 20 mM sodium acetate buffer pH 5.2 (for PaPVA). After 4 h incubation at 25°C, the enzyme was heat 156 inactivated (80°C for 15 min), and an equal volume of modified PBS (137 mM NaCl, 2.7 mM 157 158 KCl, 100 mM Na₂HPO₄, 1.8 mM KH₂PO₄) was added to each well, followed by 100 μl of 1:100 diluted overnight biosensor. Luminescence of the biosensors was measured at 30°C during a 12 h 159 160 time-course using FLUOstar Omega (BMG Labtech) as described previously (Papaioannou et al. 161 2009). Control reactions were performed in the same manner using heat-inactivated enzyme. E. 162 coli (pSB536) was used to analyze C₄-HSL degradation, E. coli (pSB401) for C₆- to C₈-HSL (Swift et al. 1997), and *E. coli* (pSB1075) for C₁₀- to (3-OH- and 3-oxo-) C₁₂-HSL (Winson et al. 163 164 1998). To determine the enzyme activity on AHLs, the ratio of luminescence unit to biosensor growth in OD₆₀₀ (relative luminescence unit, RLU) from active enzymes was compared to those 165 166 from inactive enzymes. 167

HPLC analysis

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To confirm the deacylase activity of PVA enzyme on long chain AHLs, the reaction with C₁₀-170 171 HSL was analyzed by HPLC (Uroz et al. 2008). The enzymes (25 µg in 3ml of same buffer as the bioluminescence assay) were incubated with 0.4 mM C₁₀-HSL for 4 h at 25°C (heat-172 inactivated enzyme was used as a control). Samples of 750 µl from time 0 and 4 h were 173 processed for detection of residual substrate, HSL and decanoic acid (Wahjudi et al. 2011). 174 For detection of the substrate, residual C₁₀-HSL in the reaction mixture was extracted twice with 175 176 an equal volume of acidified ethyl acetate. The free HSL released during the reaction was dansylated with an equal volume of 2.5 mg ml⁻¹ dansyl chloride (in acetone) and incubated 177 overnight at 37°C (Lin et al. 2003). After SpeedVac evaporation, the sample was neutralized 178 with 50 µl of 0.2 M HCl and diluted with acetonitrile. Decanoic acid in the sample was extracted 179 180 thrice with an equal volume of hexane followed by drying under a nitrogen stream and derivatization with 4-bromomethoxy-7-methyl coumarin (BrMMC) reagent was performed as 181 described previously (Wolf and Korf 1990). 182 HPLC was carried out in a Shimadzu LC-10AT VP system using a Phenomenex Luna C18 183 184 reverse-phase column (250 x 4.60 mm, 5 µm) coupled with a SPD-M10AVP PDA detector. The 185 column was washed with 5% acetonitrile in water (solvent A), and the sample was eluted in a linear gradient to 100% acetonitrile (solvent B). C₁₀-HSL was detected at 219 nm, dansylated 186 HSL at 267 nm, and BrMMC-derivatized decanoic acid at 328 nm (Uroz et al. 2008). Reaction 187 188 control of reference substrate and products showed that the dansylation and BrMMC 189 derivatization was specific to HSL and decanoic acid, respectively (data not shown). 190 Kinetics of AHL degradation by PVAs 191 The kinetic behavior of AtPVA and PaPVA on 3-oxo-C₁₂-HSL was determined by an end-point 192 assay using ortho-phthalaldehyde (OPA) derivatization of the HSL product. Eight different 193 concentrations of 3-oxo-C₁₂-HSL in which the substrate was completely soluble (0.01-0.25 mM) 194 were prepared from DMSO stock. The reaction mixture consisted of 100 mM NaCl, 1 mM DTT and 25 mM sodium phosphate buffer pH 7.4 (for AtPVA) or 20 mM sodium acetate buffer pH 195 5.2 (for PaPVA). The DMSO concentration was kept at 0.8% for each reaction. Enzyme (2 µg 196 AtPVA or 0.5 μg PaPVA) was added into the 1 ml reaction mixture; a 90 μL sample was taken 197

198 immediately and thereafter regularly at 1 min intervals. The enzyme was inactivated with 10 µL of 1M NaOH; this step did not interfere with the subsequent derivatization. After removal of 199 200 enzyme by centrifugation, 50 µL was transferred into a black Fluotrac microplate (Greiner Bio-One) and mixed with 50 µL OPA reagent (Sigma-Aldrich), followed by 20 min incubation at 201 25°C. Fluorescence was measured on a FLUOstar Omega, BMG Labtech with an excitation at 202 355 nm and emission at 460 nm. A standard curve using 0-0.25 mM HSL standard prepared in 203 reaction mixture showed a straight line that can be fitted to the following equation: y = 77290x +204 490.5 (R²=0.9996). Initial velocity was limited in the range of 15% substrate conversion and 205 calculated from the standard curve. The enzyme kinetics model was analyzed by fitting the v/[S] 206 207 curves in GraphPad Prism software. Docking of AHLs to PaPVA and AtPVA 208 209 The 3D structures of C₆-HSL, C₁₀-HSL and 3-oxo-C₁₂-HSL used in the docking study were obtained from PubChem compound database. Partial atomic charges of each ligand atom were 210 determined from OPLS_2005 all-atom force field using LigPrep. Grid based ligand docking 211 program Glide was used for docking these ligands in the binding site of PaPVA and AtPVA. The 212 binding site was defined as a grid box of dimension 26x26x26 Å, centered on the Cys1 residue. 213 Receptor grid generation was followed by ligand docking where the ligands were docked flexibly 214 using Glide's extra precision. Free energy of binding was roughly estimated by using an 215 216 empirical scoring function called GlideScore, which includes electrostatic, van der Waals interaction and other terms for rewarding or penalizing interactions that are known to influence 217 ligand binding. All structural figures were prepared using PyMol or CCP4MG. 218 219 Disruption of quorum sensing in Pseudomonas aeruginosa PAO1 by PVAs Purified AtPVA (0.08 mg ml⁻¹) or PaPVA (0.4 mg ml⁻¹) was added to a 1:100 diluted overnight 220 culture of P. aeruginosa PAO1 in 100 ml LB. Samples were taken at 6 and 24 h post inoculation, 221 centrifuged for 5 min and supernatant was stored at -20°C until further analysis. 222 223 (i) AHLs measurement. The levels of 3-oxo-C₁₂-HSL and C₄-HSL were measured by bioluminescence assay using biosensor E. coli pSB1075 and pSB536 respectively (Winson et al. 224

1998; Swift et al. 1997). Cell-free supernatant was filtered through a 0.2 µm pore filter, and 20

- 226 μL of the sample was mixed with 180 μL of 1:100 diluted overnight biosensor culture. Light
- production was monitored at 30°C for 12 h.
- 228 (ii) Elastase assay. Cell-free supernatant (100 μL) was added to 900 μL of elastase buffer (100
- 229 mM Tris HCl pH 7.5; 1 mM CaCl₂) containing 20 mg of Elastin Congo Red (ECR, Sigma
- Aldrich) (Ohman et al. 1980). After 2h at 37°C, elastase activity of the supernatant was measured
- 231 as A_{495}/A_{600} .
- 232 (iii) Pyocyanin assay. Cell-free supernatant (5 ml) was extracted with 3 ml chloroform, and re-
- extracted with 1 ml of 0.2 M HCl (Essar et al. 1990). After centrifugation, the absorbance of HCl
- layer was measured at 520 nm. Production of pyocyanin (µg ml⁻¹ culture) was calculated as
- 235 $(A_{520}/A_{600}) \times 17.072$.
- 236 (iv) Biofilm formation assay. The static biofilm assay was performed in a round-bottom
- polystyrene 96-well plate (Greiner Bio-One) using a method by Merrit et al. (2005) with
- modification. 0.5 mg ml⁻¹ AtPVA or 0.66 mg ml⁻¹ PaPVA was added to an overnight culture of
- 239 *P. aeruginosa* PAO1 (0.01OD) in M9 medium (47.7 mM Na₂HPO₄.7H₂O; 22 mM KH₂PO₄; 8.5
- 240 mM NaCl; 18.7 mM NH₄Cl; 2 mM MgSO₄; 0.1 mM CaCl₂; 0.01 mM glucose). A minimum of
- 241 20 wells per treatment were used with an aliquot of 110 µL in each well. Biomass quantification
- was performed using a crystal violet method (Chow et al. 2014) after 18 h at 30°C.
- 243 (v) Galleria mellonella killing assay. Larvae of G. mellonella were obtained from Frits Kuiper
- (Groningen, The Netherlands) and kept in a dark container at 15°C. Animals of 2.5-3 cm size
- were selected for the assay, with a minimum of 15 animals per treatment. An overnight culture of
- 246 P. aeruginosa PAO1 was diluted 1:100 in LB medium, grown into an early logarithmic phase
- $(A_{600} 0.3-0.4)$, and the CFU count was determined from a standard curve of CFUs/ A_{600} . The
- cells were then washed with sterile 10 mM MgSO₄, and diluted into 10³ CFU/mL. Afterwards,
- 100 μL of enzyme (0.5 mg ml⁻¹ AtPVA or 0.66 mg ml⁻¹ PaPVA) or reaction buffer was added to
- 250 900 μL of bacteria and incubated at 30°C for 1 hour. An insulin pen (HumaPen Luxura; Lilly
- Nederland) was used to inject 10 µL of the culture to the last proleg of the larvae. Animals
- 252 injected with 10 mM MgSO₄ served as a control for physical trauma. Infection development was
- followed for 24 hours at 30°C (Beeton et al. 2015; Koch et al. 2014b). The animals were
- considered dead when not reacting to touch or have turned black.

Accession code: 255 256 The structural coordinates for AtPVA have been deposited in the PDB under the accession codes 5.J9R. 257 258 **Results:** 259 Biochemical characterization of AtPVA 260 261 AtPVA was expressed as a tetramer of molecular mass 148 kDa; the enzyme exhibited a specific activity of 205 μmolmin⁻¹mg⁻¹with high specificity for Pen V over bile salts and other β-lactam 262 antibiotics (Fig. S1, Online Resource 1). Maximum Pen V hydrolysis was observed at 45°C in 263 264 optimum pH 6 - 7 (Fig. 1). AtPVA was stable in the pH range 5-8, while PaPVA (Avinash et al. 2015) was more stable in acidic pH (3-6). There was also a drastic reduction in AtPVA activity 265 266 and loss of tertiary structure at 60°C (Fig. 1). AtPVA was observed to exhibit complex kinetic behaviour similar to PaPVA, showing positive 267 cooperativity and substrate inhibition with Pen V and modulation of PVA activity in the presence 268 269 of bile salts (Fig. 2a). The major difference between AtPVA and PaPVA lies in the extent of substrate inhibition; AtPVA showed a K_i of 47.2 mM, compared to 163.1 mM for PaPVA. Near 270 complete reduction of AtPVA activity was observed at 240 mM Pen V, while PaPVA still had 271 considerable activity (20% of V_{max}) at the same concentration (Avinash et al. 2015). Drastic 272 reduction in Pen V hydrolysis with AtPVA was also observed in the presence of high GDCA 273 (glycodeoxycholate, a bile salt) concentration (Fig. 2b). 274 Structural analysis of AtPVA 275 The structural features of AtPVA closely resemble the PaPVA structure (PDB ID 4WL2) with a 276 few subtle differences. Although the AtPVA tetramer (Fig. 3) possesses a similar non-planar 277 278 orientation and distance between subunits as PaPVA (Avinash et al. 2016a), the angle between the opposite subunits (169.6°) was closer to the planar shape of the PVA from *Bacillus* 279 sphaericus (171°) than PaPVA (158°). AtPVA shares many similar active site residues with 280 PaPVA including the nucleophilic N-terminal cysteine (C1), and the presence of two Trp 281

282 residues (W21, W80) in the active site participating in substrate binding. Superposition of the two structures revealed that AtPVA (and other PVAs) lack the 5-residue insertion in the loop 283 region (61-74) near the active site in contrast to PaPVA (Avinash et al. 2015). It is possible that 284 the length of this loop might play a role in modulating the substrate inhibition in PVAs from 285 Gram-negative bacteria. Finally, AtPVA and BtBSH (BSH from Gram-negative Bacteroides 286 287 thetaiotamicron, PDB ID 3HBC) also lack a solvent-exposed loop covering the region 228-239 that is present in PaPVA. 288 **AHL degradation by PVAs** 289 The ability of PVAs from Gram-negative bacteria (PaPVA and AtPVA) to hydrolyze AHL 290 291 signals was evaluated to explore their possible association with quorum sensing. Incubation (4 h) of long chain AHLs with pure PVA enzymes showed reduction in bioluminescence compared to 292 the heat-inactivated control, indicating AHL degradation. Activity of PaPVA was restricted to 293 C₁₀ and C₁₂-HSL. AtPVA was active on a broader substrate spectrum (C₆ to C₁₂-HSL), although 294 significant quenching was observed with the long chain AHLs, with moderate activity on C₆ and 295 296 C₈-HSLs (Table 2). Both enzymes were observed to be distinctly more active on straight chain 297 AHLs, with only moderate quenching in case of oxo- or hydroxy- substituted AHLs. The activity of the PVA enzymes on long chain AHLs was further confirmed by monitoring the degradation 298 299 of C₁₀-HSL using HPLC (Fig. 4). 300 **Kinetics of AHL degradation** For kinetic analysis, 3-oxo-C₁₂-HSL was chosen as a representative substrate as it is a highly 301 studied signal produced by *P. aeruginosa* and has significant clinical relevance (Cooley et al. 302 2010; Miyari et al. 2006). PaPVA (18.9 μmolmin⁻¹mg⁻¹) exhibited 4-fold higher activity over 303 AtPVA (4 μmolmin⁻¹mg⁻¹) with 0.2 mM 3-oxo-C₁₂HSL as substrate, similar to the trend for Pen 304 V as substrate (Avinash et al. 2015). 305 AtPVA and PaPVA showed sigmoid v/[S] curves with increasing concentrations of 3-oxo-C₁₂-306 HSL, exhibiting a better fit for allosteric behaviour. However, saturation could not be achieved 307 308 for both the enzymes as the low solubility of 3-oxo-C₁₂-HSL in aqueous buffer did not permit

rate measurements at concentrations higher than 0.25 mM. A reasonable estimate of kinetic

parameters calculated by applying initial values as constraints to the allosteric sigmoidal

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- equation revealed similar $K_{0.5}$ values but a significantly higher V_{max} for PaPVA (Fig. 5).
- 312 Apparent k_{cat}/K_m values for PaPVA (13.5x10⁴ M⁻¹s⁻¹) and AtPVA (2.68x10⁴ M⁻¹s⁻¹) were
- comparable to the available value for HacB acylase $(7.8 \times 10^4 \, \text{M}^{-1} \, \text{s}^{-1})$ (Wahjudi et al. 2011) and
- 314 10 fold higher than PvdQ acylase $(5.8 \times 10^3 \text{ M}^{-1} \text{s}^{-1})$ (Koch et al. 2014a).

Binding of long chain AHLs to AtPVA and PaPVA

- Docking studies were performed to understand the structural interactions responsible for the
- 317 selective activity of PVAs on long chain AHLs. The mode of binding was almost identical in
- both PVAs, with the AHLs (C₆-HSL, C₁₀-HSL and 3-oxo-C₁₂-HSL) binding to the active site
- with similar amide bond orientation and favourable binding energy (Fig. S2, Online Resource 1).
- However, the extent of interaction of enzyme residues with the substrate molecule seemed to
- increase with the increase in length of acyl chain of the AHL molecule.
- The lactone ring was housed in the same pocket where the β -lactam moiety was bound in the
- case of Pen V (Avinash et al. 2016a) with an Asn residue (N250 in AtPVA or N271 in PaPVA)
- involved in hydrogen bonding with the NH group of the amide bond. The AHL acyl chain fits
- into a hydrophobic pocket lined primarily by the two Trp residues in the active site (W23, W87
- in AtPVA and W21, W80 in PaPVA respectively) and residues from loop 2 and loop3
- surrounding the active site (Fig. 6). It appears that longer hydrophobic chains in C_{10} -HSL and 3-
- 328 oxo-C₁₂-HSL enable greater number of hydrophobic interactions with the enzyme. The loop
- residues (Y61, L137, A138 in *At*PVA and F63, M69, L146 and A147 in *Pa*PVA respectively)
- form additional interactions with the hydrophobic acyl chain in these substrates, probably
- enhancing the strength of binding and favourably orienting the AHL molecule in the active site.
- Better binding affinity values (estimated as glidescores) and smaller nucleophilic attack distances
- from the N-terminal catalytic cysteine (C1) to the carbonyl carbon of the substrate were also
- observed in C₁₀-HSL and 3-oxo-C₁₂-HSL over C₆-HSL (Table 3). The presence of a (oxo- or
- 335 hydroxy-) substituent did not effect a significant change in binding orientation, although a
- reduction in activity was observed (Table 3). It is possible that a change in polarity due to the
- presence of a 3' substituent might have caused a binding impediment. A preference for
- unsubstituted AHLs has also been observed in AHL acylases from Shewanella sp. (Morohoshi et
- al. 2008) and Acinetobacter sp. (Ochiai et al. 2014).

Quorum quenching in P. aeruginosa by PVAs

340

Exogenous addition of the PVAs into P. aeruginosa PAO1 culture was followed by 341 measurement of AHL levels and monitoring of QS-regulated virulence factors and biofilm 342 343 formation, to study their quorum quenching activity. Decrease in 3-oxo-C₁₂-HSL levels was 344 apparent 6h post incubation (Fig. 7a), but the accumulation of C₄-HSL was unaffected (data not shown). This result corroborates the finding that both PVAs hydrolyze only long chain AHLs. 345 Elastase and pyocyanin levels were also negatively influenced at 6 h after acylase addition (early 346 stationary phase) (Fig. 7b). Interestingly, AtPVA almost completely blocked the production and 347 pyocyanin and elastolytic activity even at 5-fold lower concentration than PaPVA, despite 348 PaPVA exhibiting higher activity on AHLs in vitro. This might be explained by a probable loss 349 in enzyme (PaPVA) activity at pH 7 required for P. aeruginosa growth, or proteolytic 350 degradation of the enzymes in bacterial culture. Although there was comparable decrease in 3-351 oxo-C₁₂-HSL levels in both cases after 6 h, the insufficient stability of PaPVA might have 352 caused a delay in AHL degradation, giving the bacteria time to activate the QS circuit. Decrease 353 354 in QQ-mediated attenuation of virulence over time has been observed earlier in the case of AhlM 355 from *Streptomyces* sp. (Park et al. 2005). 356 PVA-mediated AHL degradation also led to a moderate reduction in biofilm formation by P. 357 aeruginosa (Fig. 7c). Weakening of biofilm structure in P. aeruginosa has been linked to the disruption of the 3-oxo-C₁₂HSL regulated lasI/R QS system (DeKievit et al. 2001). In addition, 358 359 the therapeutic effects of PVAs in attenuation of P. aeruginosa virulence could be ascertained by studies on G. mellonella larvae. Simplicity of use and a positive correlation between P. 360 361 aeruginosa virulence patterns in insects and mice make G. mellonella an attractive alternative infection model for anti-virulence experiments (Papaioannau et al. 2013; Jander et al. 2000). In 362 363 the present study, preincubation of P. aeruginosa culture (10 cfu) with PVAs was observed to increase the survival rates of G. mellonella larvae after 24 h from only 10.3±7.2% in untreated 364 infection to 73±5% (AtPVA) or 53.7±11% (PaPVA) (Fig. 7d). Control injection with only 365 MgSO₄ only did not affect the survival of the larvae. Here too as in the in vitro assay, AtPVA 366 turned out to be more efficient in attenuating virulence. Regardless, these results establish the 367 potential efficacy of PVAs as QQ therapeutic agents. 368

Discussion:

371	Enzymes active on AHLs hold great potential for application as QQ agents in clinical therapy as
372	they can reduce virulence without affecting the growth of the bacteria, thereby diminishing the
373	chance for emergence of resistant strains. Apart from the many AHL acylases and lactonases
374	characterized so far, it has recently come to light that other related enzymes can promiscuously
375	degrade the AHL signals as well, effecting QQ albeit at a lower rate. Examples include
376	mammalian paraoxanases (Dong et al. 2007), porcine acylase (Xu et al. 2003) and PGA from
377	Kluyvera citrophila (KcPGA) (Mukherji et al. 2014). Although PVAs and PGAs come under the
378	same functional ambit, they show significant differences in sequence and structural composition.
379	While AHL acylases are generally homologous to heterodimeric PGAs and share similar active
380	sites including an N-terminal catalytic serine, bacterial PVAs are homotetrameric and
381	evolutionarily related to BSHs with cysteine at the N-terminal. The heterodimeric acylase from
382	Streptomyces avendulae (SIPVA) active on aliphatic penicillins and Pen V has been recently
383	hinted to degrade AHLs (Torres-Bacete et al. 2015), but it shares significant sequence and
384	structural homology with the ser-Ntn hydrolases. In the present study, the ability of cys-Ntn
385	PVAs from Gram-negative bacteria to degrade long chain AHLs and attenuate QS-mediated
386	virulence in P. aeruginosa has been described for the first time. Both the organisms employed in
387	this study are also well-known plant pathogens that produce AHLs and employed as model
388	systems to study AHL-based QS mechanisms (Steindler and Venturi 2007). The AiiB (Liu et al.
389	2007) and BlcC/AttM (Carlier et al. 2003; White et al. 2009) lactonases from A. tumefaciens
390	have been implicated in QQ; however, no acylase active on AHLs has been reported so far from
391	these bacteria.
392	Acylases active on AHLs have been observed to vary in their substrate specificities, and separate
393	into different phylogenetic clusters (Ochiai et al. 2014). Enzymes of the AAC group (including
394	AAC from Shewanella sp., PvdQ from P. aeruginosa, AhlM from Streptomyces sp. and AiiD
395	from Ralstonia sp.) degrade only long chain AHLs, while some members of the penicillin G
396	acylase group (including QuiP and HacB from P. aeruginosa, and AiiC from Anabena sp.) group
397	can act on both long and short chain AHLs. A newly characterized AHL acylase AmiE of the
398	amidase family (Ochiai et al. 2014) possesses an activity preference for long chain unsubstituted

399 AHLs similar to PVAs. However, the PVA enzymes shared little sequence similarity (<15%) with any of the known acylases active on AHLs (Fig. S3, Online Resource 1). In addition, both 400 the PVAs explored in this study did not act on the AHL signals secreted by the bacteria that 401 produce these enzymes -3-oxo- C_8 -HSL of A. tumefaciens and 3-oxo- C_6 -HSL of P. 402 atrosepticum. It would be however, interesting to study whether the substrate spectrum of 403 penicillin acylases would include the non-canonical aryl HSLs (Ahlgren et al. 2011) as well, 404 given that penicillins also possess aryl side chains. 405 406 Docking analysis showed that the AHLs bind to PVA enzymes at the same site as Pen V, with 407 the acyl chain housed in a hydrophobic pocket lined by Trp residues and loop 2 and 3 while the lactone ring interacts with residues from loop 4. Accommodation of the AHL acyl chains in the 408 409 active site hydrophobic pocket has been illustrated in the AHL acylase PvdQ (Bokhove et al. 2010) and KcPGA (Mukherji et al. 2014), while the S. lavendulae acylase also contains a long 410 411 hydrophobic pocket to bind aliphatic penicillins that can accommodate AHLs. The size of the hydrophobic pocket and the conformational variations of a few critical residues in the binding 412 413 site have been suggested to modulate the activity of different PGAs on AHLs (Chand et al. 414 2015). Moreover, it has been demonstrated in PvdQ that mutagenesis of two residues (Lα146W, Fβ24Y) in the active site could change the size of the hydrophobic binding pocket thus effecting 415 a change in substrate specificity from long chain to medium chain AHLs (Koch et al. 2014a). 416 PVAs occur in a diverse range of bacteria and some fungi (Avinash et al. 2016b), and are usually 417 418 expressed constitutively. It has been demonstrated in V. cholerae (Kovacikova et al. 2003) that 419 the PVA expression is reduced during the induction of virulence genes by the AHL-based AphA/HapR QS system and expressed more at high cell densities. Moreover, long chain AHLs 420 have been known to antagonize QS in organisms that use C₆-C₈ HSLs as signals, including 421 Chromobacterium violaceum (McClean et al. 1997) and Aeromonas hydrophila (Swift et al. 422 423 1997). It is therefore possible that the PVAs could be employed in the environment to gain a competitive advantage in a mixed species community (Roche et al. 2004), while not interfering 424 425 with the bacterium's own QS system. Further genomic and knockout analyses of PVA producing 426 strains could help shed some light on the relevance of their QQ ability in microbial physiology. Nevertheless, the recent additions of many novel acylases to the list of AHL-degrading enzymes 427

seem to go hand in hand with the complexity of AHL-based signaling mechanisms in Gramnegative bacteria.

Importantly, the knowledge of AHL-hydrolysis activity of penicillin acylases adds them to the list of QQ enzymes that can be developed for clinical applications. PVA enzyme formulations could have great potential for the biocontrol of P. aeruginosa pulmonary infection in cystic fibrosis patients. A dry powder formulation of the enzyme could not only be directly delivered into the lungs, but also increases its shelf life (Wahjudi et al. 2011). With their broad spectrum activity, PVAs can also help attenuate virulence in *Acinetobacter baumanii* (Chow et al. 2014) and co-infections by other pathogens whose QS mechanisms are at least partly dependent on long chain AHLs. QQ enzymes have also been applied to disrupt bacterial biofilms on silicone surfaces (Ivanova et al. 2015). Sustained QQ activity can be ensured for clinical application by enhancing protein stability (via directed evolution) and the use of stabilizing excipients. It is also advantageous that many penicillin acylases have been already optimized for industrial use with methods for their production on large scale; this could help in reducing development times for their clinical application in QQ systems. However, their activity levels and specificity for AHL acyl chain length should also be studied to direct their application to specific pathogens. With the recent expansion in the volume of information about QS systems in pathogenic bacteria, the development of a battery of enzymes acting on a broad range of AHLs would definitely prove beneficial in tackling bacterial virulence. In addition to its potential clinical application, this result also encourages the further exploration of possible link between QQ and the natural role of PVAs for the bacteria.

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Compliance with ethical standards:

Conflict of interest: All the authors declare that they have no conflict of interest.

457 Ethical approval: All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. 458 459 460 **References:** Ahlgren NA, Harwood CS, Schaefer AL, Giraud E, Greenberg EP (2011) Aryl-homoserine 461 lactone quorum sensing in stem-nodulating photosynthetic bradyrhizobia. Proc Nat Acad Sci 462 463 108:7183-7188. Avinash VS, Panigrahi P, Suresh CG, Pundle AV, Ramasamy S (2016a) Structural analysis 464 of a penicillin V acylase from *Pectobacterium atrosepticum* confirms the involvement of two 465 466 Trp residues for activity and specificity. J Struct Biol 193:85-94. 467 Avinash VS, Pundle, AV, Suresh CG, Ramasamy S (2016b) Penicillin acylases revisited: Importance beyond their industrial utility. Crit Rev Biotechnol 36:303-316. 468 Avinash VS, Ramasamy S, Suresh CG, Pundle, AV (2015) Penicillin V acylase from 469 Petcobacterium atrosepticum shows high specific activity and unique kinetics. Int J Biol 470 471 Macromol 79:1-7. Arroyo M, de la Mata I, Acebal C, Castillon MP (2003). Biotechnological applications of 472 473 penicillin acylases, state-of-the-art. Appl Microbiol Biotechnol, 60:507-14. 474 Beeton ML, Alves DR, Enright MC, Jenkins ATA (2015) Assessing phage therapy against 475 Pseudomonas aeruginosa using a Galleria mellonella infection model. Int J Antimicrob Agents, 46:196-200. 476 477 Bokhove M, Jimenez PN, Quax WJ, Dijkstra BW (2010) The quorum-quenching N-acyl homoserine lactone acylase PvdQ is an Ntn N-hydrolase with an unusual substrate-binding 478 pocket. Proc Nat Acad Sci 107:686-691. 479 Carlier A, Uroz S, Smadja B, Fray R, Latour X, Dessaux Y, Faure D (2003) The Ti plasmid 480 of Agrobacterium tumefaciens harbors an attM paralogous gene, aiiB, also encoding N-acyl 481 homoserine lactonase activity. Appl Environ Microbiol 69:4989-4993. 482

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- **Fig. 1** AtPVA (a) pH and (b) temperature optima, stability at increasing (c) pH (after 4 h) and (d)
- temperature. Maximum activity was taken as 100%
- Fig. 2 (a) v/[S] plot of AtPVA with Pen V as substrate. Kinetic parameters are given in inset. (b)

623	Relative PVA activi	y in the presence	e of increasing co	ncentrations of GDCA.	Pen V
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- 624 concentration was kept constant at 50 mM
- Fig. 3 (a) Tetramer structure of AtPVA. Subunits are shown in different colours. (b)
- Superposition of monomer structures of AtPVA and PaPVA. The loop extensions in PaPVA are
- shown in green (residue numbering according to PaPVA). N-terminal cysteine (stick
- 628 representation) is shown in yellow
- 629 Fig. 4 HPLC analysis of residual C₁₀-HSL and released HSL and decanoic acid, for AtPVA
- (upper panels) and PaPVA (lower panels) after 4h incubation with C_{10} -HSL at 25 $^{\circ}$ C. Reduction
- of C₁₀-HSL levels was corroborated with the occurrence of free HSL and decanoic acid,
- confirming the acylase activity of PVAs on C_{10} -HSL
- Fig. 5 v/[S] curves for (a) AtPVA and (b) PaPVA showing sigmoid kinetics with 3-oxo- C_{12} -HSL
- as substrate. Kinetic parameters are given in inset
- Fig. 6 Mode of binding of 3-oxo- C_{12} -HSL in the binding site pocket of (upper) AtPVA and
- 636 (lower) PaPVA. The hydrophobic pocket in which the alkyl side chain fits is shown as mesh
- Fig. 7 Influence of AtPVA or PaPVA on P. aeruginosa PAO1 culture: (a) 3-oxo-C₁₂-HSL level,
- (b) Elastolytic activity and pyocyanin production 6 h after exogenous addition of enzyme, (c)
- Biofilm formation, (d) Survival rate in G. mellonella 24h after infection with P. aeruginosa
- PAO1. Larvae injected with MgSO₄ were taken as control. Error bars indicate standard deviation

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- Table 1 Bacterial strains and plasmids used in this study
- Table 2 Specificity of purified AtPVA and PaPVA for different AHL substrates. Remaining
- AHLs after degradation assay were detected by suitable Lux-based biosensor at 30°C for 12h.
- Bioluminescence (%RLU) is expressed relative to heat-inactivated enzyme (taken as 100%).
- Results are displayed as Mean \pm SD from three independent experiments.
- Table 3 Properties of different AHL substrates and results of docking with AtPVA and PaPVA
- structures (AlogP = hydrophobicity, SA = surface area, Nadist = Nucleophilic attack distance
- between SH group of cys1 and carbonyl carbon atom of AHL)