



**DTU Library** 

# Itaconimides as Novel Quorum Sensing Inhibitors of Pseudomonas aeroginosa

Fong, July; Mortensen, Kim Thollund; Nørskov, Amalie; Qvortrup, Katrine; Yang, Liang; Tan, Choon Hong; Nielsen, Thomas E.; Givskov, Michael Published in: Frontiers in Cellular and Infection Microbiology

Link to article, DOI: 10.3389/fcimb.2018.00443

Publication date: 2019

Document Version Version created as part of publication process; publisher's layout; not normally made publicly available

### Link back to DTU Orbit

Citation (APA):

Fong, J., Mortensen, K. T., Nørskov, A., Qvortrup, K., Yang, L., Tan, C. H., ... Givskov, M. (2019). Itaconimides as Novel Quorum Sensing Inhibitors of Pseudomonas aeroginosa. Frontiers in Cellular and Infection Microbiology, 8, [443]. DOI: 10.3389/fcimb.2018.00443

#### **General rights**

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

• Users may download and print one copy of any publication from the public portal for the purpose of private study or research.

- · You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.



59

60 61

62

63

64 65

66

67

68

69

70

72

73

74

75 Q11

76

77 78

Q1 Q3 71

Q10

# Itaconimides as Novel Quorum Sensing Inhibitors of Pseudomonas aeruginosa

July Fong<sup>1†</sup>, Kim T. Mortensen<sup>2†</sup>, Amalie Nørskov<sup>2</sup>, Katrine Qvortrup<sup>2</sup>, Liang Yang<sup>1,3\*</sup>, Choon Hong Tan<sup>1,4</sup>, Thomas E. Nielsen<sup>1,5</sup> and Michael Givskov<sup>1,5\*</sup>

<sup>1</sup> Singapore Centre for Environmental Life Sciences Engineering, Nanyang Technological University, Singapore, Singapore, <sup>2</sup> Department of Chemistry, Technical University of Denmark, Lyngby, Denmark, <sup>3</sup> Southern University of Science and Technology, Guangdong, China, <sup>4</sup> Division of Chemistry and Biological Chemistry, School of Physical and Mathematical Sciences, Nanyang Technological University, Singapore, Singapore, <sup>5</sup> Department of Immunology and Microbiology, Costerton Biofilm Center, University of Copenhagen, Copenhagen, Denmark

#### **OPEN ACCESS**

#### Edited by:

You-Hee Cho. CHA University, South Korea

2

3

4 5

6

7

0

10

11

12

13

14

15

16

17

18

19

20

21 22

23

24

25

26

27

28

29

30

31

32

33

34

36

37

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

56

57

Q2 35 Reviewed by:

Eric Déziel. Institut National de la Recherche Scientifique (INRS), Canada Daniel Angel Ortiz, Laboratory Corporation of America Holdings (LabCorp), United States

#### \*Correspondence:

Liang Yang yangliang@ntu.edu.sg Michael Givskov mgivskov@sund.ku.dk

<sup>†</sup>These authors have contributed equally to this work

#### Specialty section:

This article was submitted to Clinical Microbiology, a section of the journal Frontiers in Cellular and Infection Microbiology

Received: 06 October 2018 Accepted: 11 December 2018 Published: xx December 2018

Citation:

#### Fong J, Mortensen KT, Nørskov A, Qvortrup K, Yang L, Tan CH, <mark>lsen TE</mark> and <mark>Givskov M</mark> (2018) Itaconimides as Novel Quorum

Sensing Inhibitors of Pseudomonas aeruginosa. Front. Cell. Infect. Microbiol. 8:443. doi: 10.3389/fcimb.2018.00443 Pseudomonas aeruginosa is known as an opportunistic pathogen that often causes 79 80 persistent infections associated with highly antibiotic-resistance and biofilms formation. 81 Chemical interference with bacterial cell-cell communication, termed quorum sensing 82 (QS), has been recognized as an attractive approach to control infections and address 83 the drug resistance problems currently observed worldwide. Instead of imposing direct 84 85 selective pressures on bacterial growth, the right bioactive compounds can preferentially 86 block QS-based communication and attenuate cascades of bacterial gene expression 87 and production of virulence factors, thus leading to reduced pathogenicity. Herein, 88 we report on the potential of itaconimides as quorum sensing inhibitors (QSI) of 89 90 P. aeruginosa. An initial hit was discovered in a screening program of an in-house 91 compound collection, and subsequent structure-activity relationship (SAR) studies 92 provided analogs that could reduce expression of central QS-regulated virulence factors 93 (elastase, rhamnolipid, and pyocyanin), and also successfully lead to the eradication of P. 94 aeruginosa biofilms in combination with tobramycin. Further studies on the cytotoxicity 95 96 of compounds using murine macrophages indicated no toxicity at common working 97 concentrations, thereby pointing to the potential of these small molecules as promising 98 entities for antimicrobial drug development. 99

#### Keywords: quorum sensing, biofilm, itaconimides, antivirulence, chemical biology

# INTRODUCTION

Pseudomonas aeruginosa is an opportunistic pathogen and a major cause of nosocomical infections 105 in patients with pneumonia, chronic wounds, urinary tract infections, and intensive care units (ICUs) (Vincent et al., 1995). As an opportunistic pathogen, this organism is highly adaptive, 107 versatile and exhibits remarkable resistance toward many antimicrobial agents. Resistance is 108 a distinctive characteristic of *P. aeruginosa*, due to its ability to express multiple resistance 109 mechanisms, including enzymes and efflux pumps (Poole, 2001; Lister et al., 2009). (U.S. Centers 110 for Disease Control and Prevention (CDC), 2013) estimated that more than 20,000 deaths per year 111 are attributed to antibiotic resistance cases. It has become a global issue, and we are threatened with 112 the slow progress of new antibiotics development and lack of preventive measure for the spread of 113 resistance. 114

194

195

196

210

211

Q8

In addition, P. aeruginosa is well-known to form biofilms, 115 which have been identified as a major underlying cause of 116 persistent infections in immunocompromised patients, chronic 117 wounds as well as on medical devices like implants, catheters, 118 tubes, artificial hip, and many more (Costerton et al., 1999). 119 Biofilms infections are often characterized by their broad 120 range resistance toward host defense mechanisms and antibiotic 121 therapy. This results in prolonged treatment, complications in 122 clinical outcomes, and additional socio-economic burdens. In 123 cystic fibrosis patients, chronic biofilm infections of P. aeruginosa 124 can cause premature death despite intensive antibiotic therapy 125 care (Bjarnsholt et al., 2009). 126

The bacteria in biofilms often exhibit different phenotypic and 127 genetic variants as compared to their planktonic counterparts. 128 In the biofilm mode of life, bacterial cells are enclosed within 129 a matrix of extracellular polymeric substances (EPS) comprises 130 of exopolysaccharides, proteins, deoxyribonucleic acid (DNA), 131 lipids or surfactants, and macromolecules that are self-produced 132 by the cells (Flemming and Wingender, 2010). All of these could 133 render antibiotics impenetrable, chelated or sequestered, and 134 diminish the efficacy of the treatment. The presence of persister 135 cells in biofilms also contributes to multidrug resistance property 136 of biofilms (Lewis, 2007). Overall, the complex biology of biofilms 137 represents a tremendous challenge to develop therapeutic agents 138 that could successfully prevent or eradicate biofilms-associated 139 infections 140

The cell-to-cell communication system called quorum sensing 141 (QS) has been reported to play major roles for establishing 142 persistent, biofilm based infections (Hentzer et al., 2003; Alhede 143 et al., 2009; Van Gennip et al., 2009; Chiang et al., 2013). The 144 QS system in P. aeruginosa utilizes acyl homoserine lactones 145 (AHLs) as signal molecules and comprises the Lux homologs 146 LasRI and RhlRI. LasI synthase is responsible for the synthesis 147 of N-(3-oxododecanoyl) homoserine lactone (3-oxo-C12-HSL), 148 which will bind to its receptor LasR and activate transcription 149 of genes responsible for virulence such as lasB, apr, and toxA 150 (Gambello et al., 1993; Passador et al., 1993). The las system also 151 positively regulates *rhl* system, where RhlI directs the synthesis of 152 N-butanoylhomoserine lactone (C4-HSL) that would bind to its 153 receptor RhlR and subsequently activate gene expression of QS 154 target genes (Brint and Ohman, 1995; Pearson et al., 1995; Pesci 155 et al., 1997). In addition, there is also a third signaling molecule 156 "pseudomonas quinolone signal" (PQS) which is intertwined 157 between the las and rhl systems (Schertzer et al., 2009). Recently, 158 a fourth signal molecules called Integrative Quorum Sensing 159 Signal (IQS) has been reported, which could overtake the 160 central las system under phosphate depletion condition (Lee 161 et al., 2013). QS defective P. aeruginosa mutants are attenuated 162 as compared to the wild-type strain, and their biofilms are 163 more susceptible toward antibiotics treatment and host immune 164 system as compared to the wild-type (Pearson et al., 2000; 165 Hentzer et al., 2002, 2003). 166

As QS governs various patterns of genes expression to control
virulence and biofilm formation, it has been proposed that
interfering with the communication system could be a promising
strategy for the control and prevention of bacterial infections
(Hentzer et al., 2003). Quorum sensing inhibitors (QSI) are

compounds that interfere with QS pathways, reduce expression 172 of QS-controlled genes and attenuate infecting bacteria. As such 173 compounds do not affect the growth of bacteria, these molecular 174 entities pose lower selective pressure on bacteria and lower the 175 risk of resistance development. Recent exploration of new classes 176 of QSI comprises natural products, synthetic molecules, and 177 enzymes that may quench or inactivate QS signals (Dong et al., 178 2001; Hentzer et al., 2002; Jakobsen et al., 2012; Fong et al., 2017). 179 Unfortunately, no clinical candidates have yet been developed for 180 therapy. 181

In the present study, we report a new class of small molecules 182 that disrupt QS pathways in P. aeruginosa. The structurally 183 related itaconimides and citraconimides have previously been 184 reported to inhibit growth of mycobacteria (Balganesh et al., 185 1999). We synthesized a range of small molecules by an 186 iterative structure-activity relationship (SAR) study and found 187 two promising candidates for further biological investigation. 188 The efficacies of these compounds were tested on P. aeruginosa 189 QS bioreporter strains (lasB-gfp; rhlA-gfp; pqsA-gfp) and also on 190 QS-controlled virulence phenotypes, such as elastase, pyocyanin, 191 and rhamnolipid production. 192

# MATERIALS AND METHODS

## **General Information**

All chemicals were purchased from Sigma Aldrich and used 197 without further purification. For biological studies, synthesized 198 compounds were prepared in DMSO as 10 mM stock solution 199 and stored at  $-20^{\circ}$ C until further usage. Overnight culture of 200 bacteria was grown in Lysogeny broth (LB) which consisted 201 of 1% tryptone, 0.5% yeast extract, 0.5% NaCl. For bioreporter 202 assay, strains were grown in ABTGC (AB minimal medium 203 supplemented with 0.2% glucose and 0.2% casamino acids) 204 (Clark and Maaloe, 1967) to minimize fluorescence interference. 205 ABTG (with no casamino acid) medium was used for biofilms 206 study in flow chambers. Strains used in this study can be found 207 in Table 1. Summary of chemical synthesis and spectroscopy data 208 can be found in Supplementary Material. 209

# **Bioreporter QS Assay**

From its frozen stock, compounds were diluted appropriately to 212 their working concentration in ABTGC medium. Experiments 213 were done as previously reported (Fong et al., 2017). Briefly, 200 214 µL of compounds was pipetted into the first rows of 96-well 215 plates (Nunc, Denmark), followed by two-fold serial dilution to 216 the rest of the rows. The last two rows were allocated to solvent 217 control (DMSO 0.1%) and blank (media control). Overnight 218 cultures of P. aeruginosa bioreporter strains were diluted to 219 optical density at 600 nm (OD<sub>600</sub>) of 0.02 ( $\sim 2.5 \times 10^8$  CFU/mL). 220 Next, 100 µL of the bacteria culture was added into each well 221 to make final OD<sub>600</sub> of 0.01. The plate was incubated at 37°C 222 for 16 h, with time-point measurement of GFP fluorescence 223 (excitation 485 nm, emission 535 nm) and OD<sub>600</sub> recorded at 224 every 15 min using Tecan Infinite 200 Pro plate reader (Tecan 225 Group Ltd, Männedorf, Switzerland). The data were exported 226 into excel files, and IC50 value calculation was determined using 227 GraphPad Prism 6 software. For IC<sub>50</sub> values determination, the 228

Q5 229

251

256

282

#### TABLE 1 | Bacterial strains used in this study.

Strains or plasmids	Relevant genotype and/or characteristics <sup>a</sup>
Strains	
PAO1	ATCC Pseudomonas aeruginosa Hentzer et al., 2002
PAO1-gfp	GFP-tagged wild-type Pseudomonas aeruginosa Yang et al., 2007
PAO1 <i>-lasB-gfp</i>	PAO1 containing <i>lasB-gfp</i> (ASV) reporter fusion Hentzer et al., 2002
PAO1-rhlA-gfp	PAO1 containing <i>rhlA-gfp</i> (ASV) reporter fusion Yang et al., 2007
PAO1-pqsA-gfp	PAO1 containing pqsA-gfp(ASV) reporter fusion Yang et al., 2009
PAO1 $\Delta lasl \Delta rhll$	Gm <sup>a</sup> ; PAO1 lasl and rhll mutant Hentzer et al., 2003
PAO1 <i>∆lasR</i>	PAO1 lasR mutant Hentzer et al., 2003
$\Delta lasR-rhlA-gfp$	PAO1 <i>lasR</i> mutant containing <i>rhlA-gfp</i> (ASV) reporter fusion Tan et al., 2013
∆lasR-pqsA-gfp	PAO1 <i>lasR</i> mutant containing <i>pqsA-gfp</i> (ASV) reporter fusion Tan et al., 2013

<sup>a</sup>Description of the strains' antibiotic resistance. Gm, gentamicin; Carb, carbenicillin resistance. 250

GFP/OD<sub>600</sub> values were taken at the time point between 4 and
6 h, where inhibition started to occur. All experiments were done
in triplicate manner and repeated at least twice to confirm the
results.

### 257 QS-Regulated Virulence Factor Assays

Elastase activity was measured using EnzChekElastase kit 258 (Invitrogen, USA), following the manufacturer's instruction. 259 260 Rhamnolipid was extracted and quantified using method reported by Koch et al. with modifications (Koch et al., 261 1991). Briefly, overnight cultures of P. aeruginosa were diluted 262 into ABTGC medium (OD<sub>600</sub> = 0.01), with and without the 263 presence of compounds (DMSO as control). Cultures were grown 264 overnight at 37°C, 200 rpm. Rhamnolipid was extracted from 265 the supernatant with diethyl ether (twice), and organic fractions 266 were concentrated to yield yellowish-white solids. The solids were 267 re-suspended in deionized water and added with 0.19% (w/v) 268 orcinol in 50% H<sub>2</sub>SO<sub>4</sub>. It was then heated at  $80^{\circ}$ C for 20–30 min 269 to give dark orange color. Absorbance was measured at 421 nm 270 and the values were normalized with cell density at  $OD_{600}$ . 271 Pyocyanin was extracted from overnight culture of P. aeruginosa 272 grown in Kings Medium A Base [MilliQ water supplemented 273 274 with proteose peptone (20 g/L), potassium sulfate (10 g/L), 275 magnesium chloride, anhydrous (1.640 g/L), and glycerol (10% v/v)]. Supernatants were collected and extracted with chloroform 276 and 0.2 M HCl. The presence of pyocyanin would turn the HCl 277 278 solution into pinkish color. Absorbance was measured at 520 nm and normalized with cell density OD<sub>600</sub> values. Experiments were 279 done in triplicate manner and repeated at least twice to confirm 280 the results. 281

## 283 **Biofilm Experiments**

GFP-tagged *P. aeruginosa* were grown in ABTG medium and flowed through flow chambers as previously described 302

303

322

323

324

325

(Sternberg and Tolker-Nielsen, 2006). Each flow chamber is 286 consisted of three-channel flow cells that were supplied with 287 a flow of medium and oxygen, while waste medium would be 288 directed into a waste flask. Briefly, overnight cultures were diluted 289 1,000 times in ABTG medium and injected into each channel for 290 1 h incubation time without flow. Next, the medium was allowed 291 to flow into the flow cells and the velocity was maintained at 0.2 292 mm/s using Cole-Palmer peristaltic pump. Biofilms were grown 293 for 72 h before treatment with compounds for further 48 h. To 294 visualize dead cells, 300 µL of propidium iodide (PI) stain was 295 injected into each flow cells. Biofilm images were taken with LSM 296 confocal laser scanning microscope (Carl Zeiss, Germany) at 20x 297 objective lens. Microscopy images were processed with IMARIS 298 software (Bitplane AG, Zurich, Switzerland). Experiments were 299 done in triplicate manner and repeated at least twice to confirm 300 the results. 301

## **Cytotoxicity Assay**

Toxicity assay was done as previously reported (Fong et al., 304 2017). Murine macrophage RAW264.7 cell lines were 305 grown in Dulbecco's Modified Eagle's Medium (DMEM, 306 Life Technologies), supplemented with 10% fetal bovine 307 serum (Gibco) at 37°C and 5% CO2. The cells were passaged 308 into 96-well microplates, with each well containing 1  $\times$ 309  $10^4$  macrophages. After 16 h, the cells were washed with 310 phosphate-buffered saline (PBS) and treated with compounds 311 at varying concentration in 100 µL DMEM. The plate was 312 incubated at 37°C and 5% CO2 for further 16 h. Resazurin 313 was then added into each well to reach final concentration 314 of 10 µM for cell viability measurement. The live cells would 315 be able to convert the dye to red color, whereby dead cells 316 would remain as blue color. Absorbance was taken at 595 nm 317 using Tecan Infinite 200 Pro plate reader (Tecan Group 318 Ltd, Männedorf, Switzerland). Experiments were done in 319 triplicate manner and repeated at least twice to confirm the 320 results. 321

## RESULTS

## Synthesis and SAR Study

Recently, we discovered that 3-methylene-1-326 tetradecylpyrrolidine-2,5-dione (Table 2, 1a) displayed QSI 327 activity against P. aeruginosa QS reporter strain (carrying a 328 lasB-gfp fusion). Different variation on the left-hand side of 329 1a proved that the exo-cyclic double bond was essential for 330 its biological activity. This led us to focus on the synthesis 331 and biological evaluation of N-substituted itaconimide analogs 332 against P. aeruginosa (Scheme 1). The procedure for the 333 synthesis of the itaconimides have been described by Cava 334 et al. (1961) and Leow et al. (2008). The commercially available 335 itaconic anhydride was treated with anilines (1.0 equiv.) 336 in CHCl<sub>3</sub> to afford the corresponding  $\alpha$ -itaconamic acids 337 (Kyung et al., 1974). The resulting acids were subsequently 338 treated with Ac<sub>2</sub>O (3.5 equiv.) and NaOAc (0.5 equiv.) at 339 elevated temperature to afford a mixture of itaconimide and 340 the isomerized product citraconimide. Generally, the yield for 341 the aniline derivatives was higher than that of the aliphatic 342

TABLE 2   Summary of	the synthesized	compounds and the	corresponding yields.
----------------------	-----------------	-------------------	-----------------------

Compound	R	<b>a (%)</b> <sup>a</sup>	b (%) <sup>a</sup>	Compound	R	a (%) <sup>a</sup>	b (%)
1		50	14	10	CI	35	15
2		37	_b	11		61	9
3		17	_b	12	Br	37	20
4	K (Y <sub>2</sub>	10	_b	13	F	40	19
5		47	_b	14	X 44	8	14
6		49	29	15	V 46	16	b
7	ОМе	_c		16	X 47 8	13	10
3	-CI	31	40	17	V 44 10	4	12
9	FCI	24	14	18	V 47 12	2	11

<sup>b</sup>Not isolated. <sup>c</sup>Commercial sample.

391 392

393 394

395

amines. An overview of the synthesized compounds is provided below (Table 2).

#### 396 **QSI Activities of Itaconimides** 397

The synthesized compounds were tested for their QS inhibitory 398 activity against the P. aeruginosa lasB-gfp reporter strain. None 399

of our synthesized compounds showed antibiotic properties 451 (Supplementary Figures 1, 2). We found that both the p-452 bromophenyl and tetradecyl-substituted itaconimide (12a and 453 18a) showed strong QS inhibition activity against the lasB-454 gfp reporter strain (Figure 1A). Both compounds inhibit the 455 expression of *lasB-gfp* in a dose-dependent manner (Figure 1B). 456

448

449

450

Fong et al.



**18a** was proved to be the most active compound synthesized in this series, with almost 25-fold higher activity ( $IC_{50}$ ) compared to our hit compound (**Figures 1C,D**).

To address the specificity of our compounds, we also tested our compounds against PAO1 wild-type (WT) and  $\Delta lasR$ harboring either *rhlA-gfp* or *pqsA-gfp* fusions. In this case, we would be able to determine if the compounds could affect other QS pathways in *las*-dependent or independent manner. Both *rhlA* and *pqsA* are the first genes of *rhl* operon and *pqs* operon that code for the production of the rhamnolipid and PQS precursor molecules (Ochsner et al., 1994; Gallagher et al., 2002). To eliminate false positive, we also tested both compounds with a *gfp*-tagged *P. aeruginosa* (expresses GFP constitutively) as control and did not observe any reduction in the fluorescence signals (**Supplementary Figure 3**).

Compounds 12a and 18a were found to inhibit expression of rhlA-gfp and pqsA-gfp in PAO1 WT. IC<sub>50</sub> values calculated for 12a are 6.67  $\pm$  0.27  $\mu$ M for *rhlA-gfp* and 2.51  $\pm$  0.19  $\mu$ M for pqsA-qfp, whereby compound 18a provides IC<sub>50</sub> values of  $0.61 \pm 0.04 \,\mu\text{M}$  for *rhlA-gfp* and  $0.143 \pm 0.13$  for *pqsA-gfp* (Figure 2). At  $10 \,\mu$ M, compound 12a was observed to inhibit *rhlA-gfp* more strongly in  $\Delta lasR$  (78% inhibition) as compared to the wild-type (58% inhibition). In  $\Delta lasR \ pgsA-gfp$  reporter strain, the inhibition only happened in the later stage. Meanwhile, compound 18a was found to inhibit both rhlA-gfp and pqsA-gfp efficiently in  $\triangle lasR$  (Figure 3). Therefore, it is likely that two compounds have different mechanisms, and compound 18a affect multiple QS pathways in las-independent pathway. 

# Effects of Itaconimides on Virulence Production

Next, we investigated the effects of synthesized compounds on virulence factors produced by P. aeruginosa, notably elastase, rhamnolipid, and pyocyanin. The three virulence factors are under QS-regulation, therefore they could be a good indicator for evaluating antivirulence activities of our compounds. We also included QS mutants as control strains, where they are defective in producing quorum sensing signals hence lower level of virulence production. At  $10 \,\mu$ M, both compounds were able to reduce all three virulence factors production. Elastase level was reduced almost half, and rhamnolipid and pyocyanin productions were abolished almost to the same level as mutant strains (Figure 4). The results showed that both compounds could indeed lower the production of virulence factors. 

# Effects of Itaconimides on *P. aeruginosa* Biofilms

QS has been shown to play important roles in biofilms matrix formation (Davey et al., 2003; Sakuragi and Kolter, 2007). Mutants lacking QS often form flat and undifferentiated biofilms that could be easily cleared by antimicrobial agents and host immune system (Wu et al., 2001; Christensen et al., 2007). Here, we investigated if the itaconimides could eradicate biofilms formation when used in combination with tobramycin antibiotic.

From **Figure 5**, it can be seen that treatment with tobramycin alone  $(10 \,\mu g/ml)$  couldn't clear the whole population of *P. aeruginosa* biofilms. The dead cells only appeared on the surface of the biofilms, which indicated that the antibiotic could not penetrate into the biofilms. In addition, biofilms often contain extracellular DNA (eDNA) which could chelate aminoglycoside antibiotics and render it inactive (Chiang et al., 2013). At  $10 \,\mu$ M, itaconimide **12a** could remarkably kill the base population of the biofilms, but not the surface. When added together with tobramycin, the combination treatment successfully eradicated the whole population of biofilms, which appeared red. Compound **18a** did not show any synergistic effect when used together with tobramycin, which could be due to poor solubility in the aqueous media.

# Cytotoxic Effects of Itaconimides on Macrophages

We next evaluated the potential application of compounds **12a** and **18a** for therapeutic application. To test the cytotoxic effects, both compounds were added into murine macrophage RAW2647 cell lines, and the cell viability was measured after 16 h incubation time. Both compounds were observed to be toxic at 40  $\mu$ M, but not cytotoxic at their working concentration (10  $\mu$ M and lower, **Figure 6**). This indicates that they could be further used for subsequent *in vivo* experiments at their relevant concentration.

# DISCUSSION

In this study, we present the investigation of itaconimides as a novel QSI against *P. aeruginosa*. The approach of using QSI molecules to attenuate the virulence of pathogenic bacteria has advantages over conventional antibiotics therapy. Instead of targeting the growth of bacteria (bactericidal or bacteriostatic strategies), QSI compounds pose lesser selective 570

E 0 E

 66'



FIGURE 1 | (A) Effects of compounds 12a and 18a on PAO1-lasB-gfp reporter strain. DMSO 0.1% was used as solvent control. (B) Dose-response effects of compounds on P. aeruginosa lasB-gfp. (C) IC50 values of 12a and 18a (D) on lasB-gfp. Calculation was done using GraphPad Prism 6 software. taken at time point between 4 and 6 h when QS inhibition started to occur. All experiments were done in triplicate manner (technical and biological replicates), only representative data are shown



FIGURE 2 | QSI inhibition, dose-response and IC50 values of compounds 12a and 18a on P. aeruginosa rhIA-gfp (A) and pqsA-gfp (B). DMSO 0.1% was used as solvent control. For the IC50 values, calculation was done using GraphPad Prism 6 software, taken at time point between 4 and 6 h when QS inhibition started to occur. All experiments were done in triplicate manner (technical and biological replicates), only representative data are shown.

pressure for the development of resistant mutants. In several studies, it has been shown that mice treated with QSI afford better survival profile and lower bacteria count loads when compared to the control group (Hentzer et al., 2002, 2003; Jakobsen et al., 2013; Fong et al., 2017). With limited options of treatment available, antivirulence therapy could be a viable approach to mitigate the future crisis of antibiotic resistance.

In this study, a structure-activity relationship (SAR) study was performed to investigate the QSI activities of itaconimides and citraconimide analogs. We screened the compounds against QS reporter strain lasB-gfp, where gene expression is under control of the QS las system. We first investigated the inhibitory properties of the two isomers with respect to *lasB-gfp* expression. Interestingly, the itaconimide scaffold was found to be more active than the citaconimide. Next, we investigated the influence 

Fong et al.



4-chloro derivative induced a 1.4-fold inhibition of PAO1-*lasB*-*gfp*. This guided us to explore anilines with different EWGs.
The most potent compound in this series was 12a from 4bromoaniline that results in a 4.4-fold increase over the parent
aniline analog. To our surprise, the 4-fluoroaniline analog 13a
did not afford a more potent inhibitor. Alongside the findings
that 4-bromoaniline provided a more potent analog, we also
found that the octylamine analog 14a was more active than our
hit compound 1a. Lastly, we also included aliphatic chains to

available aliphatic amines with variating carbon lengths (n = 4, 6, 8, 10, 12, 14). A clear trend was observed, where longer alkyl chain resulted in lower IC<sub>50</sub> values. The outcome was quite interesting, as IC<sub>50</sub> values of the citraconimide substituents of these analogs (**16b**, **17b**, and **18b**) were significantly lowered. The most potent compound was found to be the tetradecylamine analog, **18a**, being the most active of both series. By substituting the naphthalene group with either 4-bromoaniline or long chain alkyl amines, IC<sub>50</sub> values against PAO1-*lasB-gfp* were significantly reduced to the low micromolar range (2.41  $\pm$ 

792

793

794

795

796

797

798

Fong et al.

799

800

801

802

803

804

805 806

807

808

809 810

811

812

813

814

815

816 817 818



**FIGURE 6** | Cytotoxic effect of compounds **12a** and **18a** on murine macrophages. DMSO was used as vehicle control. Compounds were tested at various concentrations (40  $\mu$ M and subsequent 2x dilution). Experiments were done in triplicate manner (technical and biological replicates). Error bars are means  $\pm$  SDs.

819 0.99  $\mu$ M for **12a** and 0.17  $\pm$  0.04  $\mu$ M for **18a**). In summary, 820 our SAR studies revealed two structurally important variations 821 for the itaconimides that are highly important for QSI activity 822 (**Table 3**).

Previous studies have emphasized the advantage of 823 therapeutics that suppress multiple QS pathways (Fong et al., 824 2018). Indeed, lasR mutants are commonly found in patients 825 suffering from cystic fibrosis and other clinical setting (Hamood 826 et al., 1996; Cabrol et al., 2003; Marvig et al., 2014). Nevertheless, 827 the loss-of-function lasR mutants continuously express virulence 828 traits so P. aeruginosa could use other pathways to bypass LasR 829 in controlling pathogenicity (Dekimpe and Déziel, 2009; Lee 830 et al., 2013). Using different QS reporter strains in both PAO1 831 WT and  $\triangle lasR$ , it can be deduced that our compounds do not 832 specifically inhibit one QS pathway. Both compounds could 833 inhibit expression of *rhlA-gfp* and *pqsA-gfp* in *lasR* mutant. The 834 inhibition of *pqsA-gfp* in  $\Delta lasR$  was less apparent for compound 835 12a, which could indicate that the compound inhibits PQS 836 system through las-dependent manner. The itaconimide analogs 837 presented here hold unique potential as broad target QSIs to 838 control infections via an anti-virulence strategy. Future work will 839 aim to elucidate the mechanism on how both compounds inhibit 840 QS in P. aeruginosa. 841

Next, we also tested the effects of our compounds on 842 the production of various virulence factors, such as elastase, 843 rhamnolipid, and pyocyanin production. Elastase is one of the 844 major proteases produced by P. aeruginosa, involved in host 845 tissue damage and host immune responses (Kamath et al., 2002). 846 Rhamnolipid is also an essential virulence factor and plays several 847 key roles in biofilms formation, swarming, and in particular 848 host immune evasion. It promotes rapid necrotic killing of 849 polymorphonuclear (PMNs) leukocytes and also infiltration of 850 respiratory epithelial cells (Zulianello et al., 2006; Jensen et al., 851 2007). P. aeruginosa also secretes pyocyanin, blue redox-active 852 853 secondary metabolite that has several deleterious effects on mammalian cells (Lau et al., 2004), which is also regulated by QS 854 under *pqs* system. When tested at  $10 \,\mu$ M, the compounds could 855

856

857

879

880

881

882

883

884

**TABLE 3** | Summary of IC\_{50} values of itaconimide (**a**) and citraconimide (**b**) analogs against PAO1-*lasB-gfp*.

Compound	а	b
1	7.37 ± 0.71	-
2	$23.52 \pm 0.81$	-
3	$10.27 \pm 0.05$	-
4	-	-
5	$17.95\pm0.49$	-
6	$10.67 \pm 0.49$	-
7	-	NA
8	$7.45\pm0.65$	-
9	$7.65 \pm 0.37$	-
10	$6.53\pm0.61$	-
11	$17.68\pm0.16$	-
12	$2.41\pm0.99$	-
13	$11.89\pm0.34$	-
14	$26.66\pm0.98$	-
15	$1.71 \pm 0.34$	-
16	$0.30\pm0.05$	$2.01\pm0.39$
17	$0.30\pm0.17$	$0.91\pm0.20$
18	$0.17 \pm 0.04$	$0.53\pm0.12$

All IC<sub>50</sub> values were reported as  $\mu$ M. Experiments were done in triplicate manner and repeated at least twice to confirm the results. NA, Not available.

reduce production of various virulence factors controlled by QS. This shows potential of our compounds as antivirulence agent of *P. aeruginosa.* 

P. aeruginosa biofilms are highly resistant to most antibiotics, 885 including the last-resort polymyxin antibiotic available, colistin 886 (Chua et al., 2016). As QS has definite role in biofilms 887 development, it has been proposed that QSI compounds could 888 be used as prophylactic treatment for biofilms infections. One 889 such case is using azithromycin, which inhibits QS and biofilm 890 formation at sub-MIC concentration. Promising results were 891 observed in pulmonary infections and CF patients in many 892 clinical trials data upon treatment of low-dose of AZM (2 µg/ml) 893 (Hansen et al., 2005; Fleet et al., 2013). In this study, we utilized 894 a combination of our compounds with the aminoglycoside 895 tobramycin to treat P. aeruginosa biofilms grown on flow 896 chambers for 3 days. We chose to study tobramycin because 897 of its clinical relevance to cystic fibrosis (CF) patients. The 898 clinical isolates from lung patients often confer resistance to 899 aminoglycoside antibiotics (Hurley et al., 1995; Saiman et al., 900 1996). Our data also shows that treatment with tobramycin alone 901 only kills the upper layer of biofilms. Combined with 12a, we 902 observed that the whole population of biofilms was eradicated. 903 The results offer promising application of QSI in combination 904 with antibiotics as a control for biofilm-associated infections. 905

It has been reported that stringent response, which provides rapid adaptation to environmental stresses, regulates QS network and also bacteria's survival in biofilms of *P. aeruginosa* (van Delden et al., 2001; Nguyen et al., 2011; Schafhauser et al., 2014). QS-deficient mutants also have lower catalase and superoxide dismutase activities, and therefore more sensitive to oxidative stress (Hassett et al., 1999). Through our biofilms experiment, we

974 975

976 977

978

979

980

981

982

983

984

985

986

987

988

989

990

991

992 09

993

994 995

996

997

998

999

1000

1001

1004

1005

1006

1013

1014

Q12

could observe some cells death upon treatment with compound 913 12a in the base of the biofilms. This raises the possibility that our 914 compound could also target stringent response and other stress-915 response genes, which resulted in the observed killing effect. 916 For compound 18a, it contains long alkyl chain, which may 917 render its solubility and penetration into the biofilms matrix. 918 Future study will investigate how our compounds could synergize 919 with other antibiotics to treat biofilms from other clinical 920 isolates. 921

Lastly, we investigated the cytotoxic profile of our compounds 922 on macrophages. The cytotoxicity of the compounds is important 923 if they are to be used in animal studies for subsequent drug 924 development. Results indicated that compound 12a and 18a 925 are not toxic up to 40 µM concentration. Still, further studies 926 are needed to qualify the efficacy of the compounds in mice 927 models for their potential as anti-biofilm agents, as well as their 928 pharmacodynamic and pharmacokinetic profiles. 929

In conclusion, we report the novel use of itaconimides as 930 antivirulence compounds for P. aeruginosa. These compounds 931 suppress the las, rhl, and pas QS systems of P. aeruginosa, and 932 effectively abolish virulence expression activities. Compounds 933 12a and 18a showed low micromolar IC50 values against all three 934 QS reporter strains with only little toxicity against macrophages 935 at the administrated concentration. Moreover, a synergistic effect 936 with tobramycin was observed for the killing of P. aeruginosa 937

## REFERENCES

938 939

940

941

942

943

944

945

946

947

948

949

950

951

952

953

954

955

956

957

958

959

960

961

962

963

964

965

966

967

968

969

Q13

Q14

- Alhede, M., Bjarnsholt, T., Jensen, P. Ø., Phipps, R. K., Moser, C., Christophersen, L., et al. (2009). Pseudomonas aeruginosa recognizes and responds aggressively to the presence of polymorphonuclear leukocytes. Microbiology 155, 3500-3508. doi: 10.1099/mic.0.031443-0
- Balganesh, M., Ethirajulu, K., Ganguly, B. S., Janakiraman, R., Kaur, P., Kajipalya, R., et al. (1999). Mycobacterial inhibitors. WO1999065483A1.
- Bjarnsholt, T., Jensen, P. Ø., Fiandaca, M. J., Pedersen, J., Hansen, C. R., Andersen, C. B., et al. (2009). Pseudomonas aeruginosa biofilms in the respiratory tract of cystic fibrosis patients. Pediatr. Pulmonol. 44, 547-558. doi: 10.1002/ppul. 21011
- Brint, J. M., and Ohman, D. E. (1995). Synthesis of multiple exoproducts in Pseudomonas aeruginosa is under the control of RhlR-RhlI, another set of regulators in strain PAO1 with homology to the autoinducer-responsive LuxR-LuxI family. J. Bacteriol. 177. doi: 10.1128/jb.177.24.7155-7163.1995
- Cabrol, S., Olliver, A., Pier, G. B., Andremont, A., and Ruimy, R. (2003). Transcription of quorum-sensing system genes in clinical and environmental isolates of Pseudomonas aeruginosa. J. Bacteriol. 185, 7222-7230. doi: 10.1128/JB.185.24.7222-7230.2003
- Cava, M. P., Deana, A. A., Muth, K., and Mitchell, M. J. (1961). N-Phenylmaleimide. Org. Synth. 41:93. doi: 10.15227/orgsyn.041.0093
- Chiang, W.-C., Nilsson, M., Jensen, P. Ø., Høiby, N., Nielsen, T. E., Givskov, M., et al. (2013). Extracellular DNA shields against aminoglycosides in Pseudomonas aeruginosa biofilms. Antimicrob. Agents Chemother. 57, 2352-2361. doi: 10.1128/AAC.00001-13
- Christensen, L. D., Moser, C., Jensen, P. Ø., Rasmussen, T. B., Christophersen, L., Kjelleberg, S., et al. (2007). Impact of Pseudomonas aeruginosa quorum sensing on biofilm persistence in an in vivo intraperitoneal foreign-body infection model. Microbiology 153, 2312-2320. doi: 10.1099/mic.0.2007/006122-0
- Chua, S. L., Yam, J. K. H., Hao, P., Adav, S. S., Salido, M. M., Liu, Y., et al. (2016). Selective labelling and eradication of antibiotictolerant bacterial populations in Pseudomonas aeruginosa biofilms. Nat. Commun. 7:10750. doi: 10.1038/ncomms10750

biofilms, including otherwise the tolerant and hard to target sub-970 population cells. Overall, our findings point to a new class of hit compounds of relevance to the development of new drugs against the superbug P. aeruginosa.

# **AUTHOR CONTRIBUTIONS**

LY, TN, and MG designed methods and experiments. JF and KM performed the experiments and analyzed the data. AN, KQ, CT, and TN were in charge of the chemical synthesis. JF and KM wrote the paper and carefully revised by LY, TN, and MG. All authors have contributed to, read, and approved the manuscript.

# FUNDING

This research was supported by the National Research Foundation and Ministry of Education Singapore under its Research Centre of Excellence Program and AcRF Tier 2 (MOE2016-T2-1-010) from Ministry of Education, Singapore.

# SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb. 2018.00443/full#supplementary-material

- Clark, D. J., and Maaloe, O. (1967). DNA replication and the division cycle in Escherichia coli. J. Mol. Biol. 23, 99-112. doi: 10.1016/S0022-2836(67)80070-6
- Costerton, J. W., Stewart, P. S., and Greenberg, E. P. (1999). Bacterial biofilms: a common cause of persistent infections. Science 284, 1318-1322. doi: 10.1126/science.284.5418.1318
- Davey, M. E., Caiazza, N. C., and O'toole, G. A. (2003). Rhamnolipid surfactant 1002 production affects biofilm architecture in *Pseudomonas aeruginosa* PAO1. I. 1003 Bacteriol. 185, 1027-1036. doi: 10.1128/JB.185.3.1027-1036.2003
- Dekimpe, V., and Déziel, E. (2009). Revisiting the quorum-sensing hierarchy in Pseudomonas aeruginosa: the transcriptional regulator RhlR regulates LasRspecific factors. Microbiology 155, 712-723. doi: 10.1099/mic.0.022764-0
- Dong, Y.-H., Wang, L.-H., Xu, J.-L., Zhang, H.-B., Zhang, X.-F., and Zhang, 1007 L.-H. (2001). Quenching quorum-sensing-dependent bacterial infection by 1008 an N-acyl homoserine lactonase. Nature 144, 813-817. doi: 10.1038/350 1009 81101 1010
- Fleet, J. E., Guha, K., Piper, S., Banya, W., Bilton, D., and Hodson, M. E. (2013). A retrospective analysis of the impact of azithromycin maintenance therapy 1011 on adults attending a UK cystic fibrosis clinic. J. Cyst. Fibros. 2013, 49-53. 1012 doi: 10.1016/j.jcf.2012.05.010
- Flemming, H.-C., and Wingender, J. (2010). The biofilm matrix. Nat. Rev. Microbiol. 8, 623-633. doi: 10.1038/nrmicro2415
- 1015 Fong, J., Yuan, M., Jakobsen, T. H., Mortensen, K. T., Santos, M. M. S. D., Chua, S. 1016 L., et al. (2017). Disulfide bond-containing ajoene analogues as novel quorum sensing inhibitors of Pseudomonas aeruginosa. J. Med. Chem. 60, 215-227. 1017 doi: 10.1021/acs.jmedchem.6b01025 1018
- Fong, J., Zhang, C., Yang, R., Boo, Z. Z., Tan, S. K., Nielsen, T. E., 1019 et al. (2018). Combination therapy strategy of quorum quenching enzyme 1020 and quorum sensing inhibitor in suppressing multiple quorum sensing pathways of P. aeruginosa. Sci. Rep. 8:1155. doi: 10.1038/s41598-018-1021 19504-w 1022
- Gallagher, L. A., Mcknight, S. L., Kuznetsova, M. S., Pesci, E. C., and 1023 Manoil, C. (2002). Functions required for extracellular quinolone 1024 signaling by Pseudomonas aeruginosa. J. Bacteriol. 184, 6472-6480. 1025 doi: 10.1128/JB.184.23.6472-6480.2002 1026

Frontiers in Cellular and Infection Microbiology | www.frontiersin.org

1092

1093

1097

1098

1099

1100

1101

1102

1104

1105

1106

1107

1113

1114

1115

1119

1120

1103 Q15

- Gambello, M. J., Kaye, S., and Iglewski, B. H. (1993). LasR of Pseudomonas 1027 aeruginosa is a transcriptional activator of the alkaline protease gene 1028 (apr) and an enhancer of exotoxin a expression. Infect. Immun. 61, 1029 1180-1184.
- 1030 Hamood, A. N., Griswold, J., and Colmer, J. (1996). Characterization of elastase-1031 deficient clinical isolates of Pseudomonas aeruginosa. Infect. Immun. 64, 1032 3154-3160.
- Hansen, C. R., Pressler, T., Koch, C., and Høiby, N. (2005). Long-term 1033 azitromycin treatment of cystic fibrosis patients with chronic Pseudomonas 1034 aeruginosa infection; an observational cohort study. J. Cyst. Fibros 4, 35-40. 1035 doi: 10.1016/j.jcf.2004.09.001
- 1036 Hassett, D. J., Ma, J. F., Elkins, J. G., Mcdermott, T. R., Ochsner, U. A., West, S. E. H., et al. (1999). Quorum sensing in Pseudomonas aeruginosa 1037 controls expression of catalase and superoxide dismutase genes and mediates 1038 biofilm susceptibility to hydrogen peroxide. Mol. Microbiol. 34, 1082-1093. 1039 doi: 10.1046/j.1365-2958.1999.01672.x
- 1040 Hentzer, M., Riedel, K., Rasmussen, T. B., Heydorn, A., Andersen, J. B., Parsek, M. R., et al. (2002). Inhibition of quorum sensing in Pseudomonas aeruginosa 1041 biofilm bacteria by a halogenated furanone compound. Microbiology 148, 1042 87-102. doi: 10.1099/00221287-148-1-87 1043
- Hentzer, M., Wu, H., Andersen, J. B., Riedel, K., Rasmussen, T. B., Bagge, N., et al. 1044 (2003). Attenuation of Pseudomonas aeruginosa virulence by quorum sensing 1045 inhibitors. EMBO J. 22, 3803-3815. doi: 10.1093/emboj/cdg366
- Hurley, J. C., Miller, G. H., and Smith, A. L. (1995). Mechanism of amikacin 1046 resistance in Pseudomonas aeruginosa isolates from patients with cystic 1047 fibrosis. Diagn. Microbiol. Infect. Dis. 22, 331-336. doi: 10.1016/0732-8893(95) 1048 00138-6
- 1049 Jakobsen, T. H., Bjarnsholt, T., Jensen, P. Ø., Givskov, M., and Høiby, N. (2013). 1050 Targeting quorum sensing in Pseudomonas aeruginosa biofilms: current and emerging inhibitors. Future Microbiol. 8, 901-921. doi: 10.2217/fmb.13.57 1051
- Jakobsen, T. H., Gennip, M. V., Phipps, R. K., Shanmugham, M. S., Christensen, L. 1052 D., Alhede, M., et al. (2012). Ajoene, a sulfur-rich molecule from garlic, inhibits 1053 genes controlled by quorum sensing. Antimicrob. Agents Chemother. 56:2314. 1054 doi: 10.1128/AAC.05919-11
- 1055 Jensen, P. Ø., Bjarnsholt, T., Phipps, R., Rasmussen, T. B., Calum, H., Christoffersen, L., et al. (2007). Rapid necrotic killing of polymorphonuclear 1056 leukocytes is caused by quorum-sensing-controlled production of 1057 rhamnolipid by Pseudomonas aeruginosa. Microbiology 153, 1329-1338. 1058 doi: 10.1099/mic.0.2006/003863-0
- 1059 Kamath, S., Kapatral, V., and Chakrabarty, A. M. (2002). Cellular function 1060 of elastase in Pseudomonas aeruginosa: role in the cleavage of nucleoside diphosphate kinase and in alginate synthesis, Mol. Microbiol. 30, 933-941. 1061 doi: 10.1046/j.1365-2958.1998.01121.x 1062
- Koch, A. K., Kappeli, O., Fiechter, A., and Reiser, J. (1991). Hydrocarbon 1063 assimilation and biosurfactant production in Pseudomonas aeruginosa 1064 mutants. J. Bacteriol. 173, 4212-4219. doi: 10.1128/jb.173.13.4212-1065 4219.1991
- Kyung, J. H., Cha, S., and Clapp, L. B. (1974). Identification of isomeric amides of 1066 itaconic acid by proton magnetic resonance spectroscopy. Organ. Magnet. Res. 1067 6, 466-468, doi: 10.1002/mrc.1270060816
- 1068 Lau, G. W., Hassett, D. J., Ran, H., and Kong, F. (2004). The role of 1069 pyocyanin in Pseudomonas aeruginosa infection. Trends Mol. Med. 10, 599-606. doi: 10.1016/j.molmed.2004.10.002 1070
- Lee, J., Wu, J., Deng, Y., Wang, J., Wang, C., Wang, J., et al. (2013). A cell-cell 1071 communication signal integrates quorum sensing and stress response. Nat. 1072 Chem. Biol. 9, 339-343. doi: 10.1038/nchembio.1225
- 1073 Leow, D., Lin, S., Chittimalla, S. K., Fu, X., and Tan, C. H. (2008). Enantioselective 1074 protonation catalyzed by a chiral bicyclic guanidine derivative. Angew. Chem. Int. Ed. 47, 5641-5645, doi: 10.1002/anie.200801378 1075
- Lewis, K. (2007). Persister cells, dormancy and infectious disease. Nat. Rev. 1076 Microbiol. 5, 48-56. doi: 10.1038/nrmicro1557
- 1077 Lister, P. D., Wolter, D. J., and Hanson, N. D. (2009). Antibacterial-1078 resistant Pseudomonas aeruginosa: clinical impact and complex regulation of chromosomally encoded resistance mechanisms. Clin. Microbiol. Rev. 22, 1079 582-610. doi: 10.1128/CMR.00040-09 1080
- Marvig, R. L., Sommer, L. M., Molin, S., and Johansen, H. K. (2014). Convergent 1081 evolution and adaptation of Pseudomonas aeruginosa within patients with 1082 cystic fibrosis. Nat. Genet. 47, 1-9. doi: 10.1038/ng.3148 1083

- Nguyen, D., Joshi-Datar, A., Lepine, F., Bauerle, E., Olakanmi, O., Beer, K., et al. 1084 (2011). Active starvation responses mediate antibiotic tolerance in biofilms 1085 and nutrient-limited bacteria. Science 334, 982-986. doi: 10.1126/science.12 1086 11037 1087
- Ochsner, U. A., Fiechter, A., and Reiser, J. (1994). Isolation, characterization, 1088 and expression in Escherichia coli of the Pseudomonas aeruginosa rhlAB genes encoding a rhamnosyltransferase involved in rhamnolipid biosurfactant 1089 synthesis. I. Biol. Chem. 269, 19787-19795. 1090
- Passador, L., Cook, J. M., Gambello, M. J., Rust, L., and Iglewski, B. H. (1993). Expression of Pseudomonas aeruginosa virulence genes requires cell-to-cell communication. Science 260, 1127-1130. doi: 10.1126/science.84 93556
- Pearson, J. P., Feldman, M., Iglewski, B. H., and Prince, A. (2000). 1094 Pseudomonas aeruginosa cell-to-cell signaling is required for virulence 1095 in a model of acute pulmonary infection. Infect. Immun. 68, 4331-4334. 1096 doi: 10.1128/IAI.68.7.4331-4334.2000
- Pearson, J. P., Passadori, L., Iglewski, B. H., and Greenberg, E. P. (1995). A second N-acylhomoserine lactone signal produced by Pseudomonas aeruginosa. Proc. Natl. Acad. Sci. U.S.A. 92, 1490-1494. doi: 10.1073/pnas.92.5.1490
- Pesci, E. C., Pearson, J. P., Seed, P. C., and Iglewski, B. H. (1997). Regulation of las and rhl quorum sensing in Pseudomonas aeruginosa. J. Bacteriol. 179, 3127-3132. doi: 10.1128/jb.179.10.3127-3132.1997
- Poole, K. (2001). Multidrug efflux pumps and antimicrobial resistance in Pseudomonas aeruginosa and related organisms. J. Mol. Microbiol. Biotechnol. 3.255-264.
- Saiman, L., Mehar, F., Niu, W. W., Neu, H. C., Shaw, K. J., Miller, G., et al. (1996). Antibiotic susceptibility of multiply resistant Pseudomonas aeruginosa isolated from patients with cystic fibrosis, including candidates for transplantation. Clin. Infect. Dis. 23, 532-537. doi: 10.1093/clinids/23.3.532
- Sakuragi, Y., and Kolter, R. (2007). Quorum-sensing regulation of the biofilm 1108 matrix genes (pel) of Pseudomonas aeruginosa. J. Bacteriol. 189, 5383-5386. 1109 doi: 10.1128/JB.00137-07
- 1110 Schafhauser, J., Lepine, F., Mckay, G., Ahlgren, H. G., Khakimova, M., and Nguyen, 1111 D. (2014). The stringent response modulates 4-hydroxy-2-alkylquinoline biosynthesis and quorum-sensing hierarchy in Pseudomonas aeruginosa. J. 1112 Bacteriol. 196, 1641-1650. doi: 10.1128/JB.01086-13
- Schertzer, J. W., Boulette, M. L., and Whiteley, M. (2009). More than a signal: non-signaling properties of quorum sensing molecules. Trends Microbiol. 17, 189-195. doi: 10.1016/j.tim.2009.02.001
- 1116 Sternberg, C., and Tolker-Nielsen, T. (2006). Growing and analyzing biofilms in flow cells. Curr. Protoc. Microbiol. Chapter 1, Unit 1B 2. 1117 doi: 10 1002/9780471729259 mc01b02s00 1118
- Tan, S. Y., Chua, S.-L., Chen, Y., Rice, S. A., Kjelleberg, S., Nielsen, T. E., et al. (2013). Identification of five structurally unrelated quorum-sensing inhibitors of Pseudomonas aeruginosa from a natural-derivative database. Antimicrob. 1121 Agents Chemother. 57, 5629-5641. doi: 10.1128/AAC.00955-13
- U.S. Centers for Disease Control and Prevention (CDC) (2013). Antibiotic 1122 Resistance Threats in the United States. 1123
- van Delden, C., Comte, R., and Bally, M. (2001). Stringent response 1124 activates quorum sensing and modulates cell density-dependent gene 1125 expression in Pseudomonas aeruginosa. J. Bacteriol. 183, 5376-5384. doi: 10.1128/JB.183.18.5376-5384.2001 1126
- Van Gennip, M., Christensen, L. D., Alhede, M., Phipps, R., Jensen, P. 1127 Ø., Christophersen, L., et al. (2009). Inactivation of the rhlA gene in 1128 Pseudomonas aeruginosa prevents rhamnolipid production, disabling the 1129 protection against polymorphonuclear leukocytes. APMIS 117, 537-546. 1130 doi: 10.1111/j.1600-0463.2009.02466.x
- 1131 Vincent, J.-L., Bihari, D. J., Suter, P. M., Bruining, H. A., White, J., Nicolas-Chanoin, M. H., et al. (1995). The prevalence of nosocomial infection in 1132 intensive care units in Europe. results of the european prevalence of infection in 1133 intensive care (epic) study. epic international advisory committee. J. Am. Med. 1134 Assoc. 274, 639-644. doi: 10.1001/jama.1995.03530080055041
- 1135 Wu, H., Song, Z., Givskov, M., Doring, G., Worlitzsch, D., Mathee, K., et al. (2001). Pseudomonas aeruginosa mutations in lasI and rhlI quorum sensing 1136 systems result in milder chronic lung infection. Microbiology 147, 1105-1113. 1137 doi: 10.1099/00221287-147-5-1105 1138
- Yang, L., Barken, K. B., Skindersoe, M. E., Christensen, A. B., Givskov, M., 1139 and Tolker-Nielsen, T. (2007). Effects of iron on DNA release and biofilm

1140

1141	development by Pseudomonas aeruginosa. Microbiology 153, 1318-1328.	Conflict of Interest Statement: The authors declare that the research was	1198
1142	doi: 10.1099/mic.0.2006/004911-0	conducted in the absence of any commercial or financial relationships that could	1199
1143	Yang, L., Kybike, M. I., Jakobsen, I. H., Hentzer, M., Bjarnsholt, I., Givskov,	be construed as a potential conflict of interest.	1200
1144	M., et al. (2009). Computer-aided identification of recognized drugs as	Copyright © 2018 Fong Mortenson Nerskov Overtrup Vang Tan Nielsen and	1201
1145	Chemother 53, 2432–2443 doi: 10.1128/AAC.01283-08	Givskov This is an open-access article distributed under the terms of the Creative	1202
1146	Zulianello, L., Canard, C., Kohler, T., Caille, D., Lacroix, JS., and	Commons Attribution License (CC BY). The use, distribution or reproduction in	1203
1147	Meda, P. (2006). Rhamnolipids are virulence factors that promote	other forums is permitted, provided the original author(s) and the copyright owner(s)	1204
1148	early infiltration of primary human airway epithelia by Pseudomonas	are credited and that the original publication in this journal is cited, in accordance	1205
1140	aeruginosa. Infect. Immun. 74, 3134–3147. doi: 10.1128/IAI.	with accepted academic practice. No use, distribution or reproduction is permitted	1205
1149	01772-05	which does not comply with these terms.	1200
1150			1207
1151			1208
1152			1209
1153			1210
1154			1211
1155			1212
1156			1213
1157			1214
1158			1215
1159			1216
1160			1217
1161			1218
1162			1219
1163			1220
1164			1221
1165			1222
1166			1223
1167			1224
1168			1225
1169			1226
1170			1227
1171			1228
1172			1229
1173			1230
1174			1231
1175			1232
1176			1232
1177			1233
1178			1235
1170			1235
11/0			1230
1100			1237
1101			1230
1182			1239
1185			1240
1184			1241
1185			1242
1186			1243
1187			1244
1188			1245
1189			1246
1190			1247
1191			1248
1192			1249
1193			1250
1194			1251
1195			1252
1196			1253
1197			1254