Changes to the resistome of Pseudomonas aeruginosa clone ST308 associated with corneal infection over time Mahjabeen Khan¹, Mark D P Willcox¹, Scott A Rice², Savitri Sharma³, Fiona Stapleton¹ ¹School of Optometry and Vision Science, UNSW, Sydney, Australia ²The Singapore Centre for Environment Life Sciences Engineering (SCELSE) and The School of Biological Sciences, Nanyang Technological University Singapore ³LV Prasad Eye Institute Hyderabad, India **Corresponding author** Email: mahjabeen.khan58@yahoo.com School of Optometry and Vision Science UNSW Sydney Australia.

Highlights

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- Recent clonal ocular isolates of *Pseudomonas aeruginosa* from India have acquired a
- 31 number of resistance genes compared to historical clones
- Consequently, resistance to antibiotics particularly fluoroquinolones in recent clones
- of *P. aeruginosa* appears to have increased.
- The acquired resistance genes found in the recent *P. aeruginosa* isolates were related
- 35 to mobile genetic elements.
- 36 Abstract

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- **Objectives**
- 38 This study compared the resistomes of isolates of *Pseudomonas aeruginosa* clone ST308
- 39 from 2018 and 1997 from India.
- 40 Methods
- 41 Two ocular clonal type ST308 isolates of *Pseudomonas aeruginosa* (198 and 219) isolated in
- 42 2018 and five historical isolates (31, 32, 33, 35 and 37) isolated in 1997 at the LV Prasad Eve
- 43 Institute in India were analysed for their susceptibilities to ciprofloxacin, levofloxacin,
- 44 gentamicin, tobramycin, piperacillin, imipenem, ceftazidime and polymyxin B. DNA was
- 45 extracted using the DNeasy® Blood and Tissue. Paired-end library was prepared using
- Nextera XT DNA library preparation kit. Libraries were sequenced on Illumina® MiSeq
- bench top sequencer generating 300 bp paired-end reads. Spades v3.12.0 was used for
- assembly, Resfinder v3.1. for acquired resistance genes and Snippy V2 for variants calling.
- 49 Integron finder v1.5.1 was used to identify the integrons present in the genomes.
- 50 **Results**

The recent isolate 219 was resistant to all tested antibiotics except polymyxin while isolate 198 was resistant to ciprofloxacin, levofloxacin, gentamicin and tobramycin. Among historical isolates five were resistant to gentamicin, tobramycin and ciprofloxacin, four were resistant to levofloxacin while two were resistant to polymyxin. Twenty-four acquired resistance genes were present in the 2018 isolates compared to 11 in the historical isolates. All isolates contained the following genes encoding for aminoglycoside aph(6)-Id, aph(3')aph(3'')-Ib), beta-lactam (blaPAO), tetracycline (tet(G)), fosfomycin (fosA), chloramphenicol (catB7), sulphonamide (sul1), quaternary ammonium (qacEdelta1) and fluoroquinolone (crpP) resistance. Isolate 198 possessed aph(3')-VI, rmtD2, qnrVC1, blaOXA-488, blaPME-1, while 219 possessed aadA1, rmtB, aac(6')-Ib-cr, blaTEM-1B, blaVIM-2, mph(E), mph(A), msr(E). In the isolate 219 genes blaTEM-1b, blaVIM-2, sul1, qnrvc1, rmtB and aadA1 were carried on class 1 integron. While an incomplete class 1 integron was also found in isolate 198 which was located on the genome where gene rmtB, blaPME-1, qnrVC1 and sul1 genes were positioned. There were no notable differences in the number of single nucleotide polymorphisms, but recent isolates carried more insertions and deletions in their genes.

Conclusion

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- 69 P. aeruginosa ocular clonal isolates have changed over time, with strains acquiring genes and
- 70 having more insertions and deletions in their chromosomal genes that confirm resistance to
- 71 antibiotics.

Keywords

73 DNA extraction, genome sequencing, acquired resistance, single nucleotide polymorphism

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Introduction Pseudomonas aeruginosa causes a variety of infections including lung infections in patients with cystic fibrosis, skin infections after burns and corneal infections (microbial keratitis). The increasing prevalence of multidrug resistant (MDR) P. aeruginosa reduces the treatment options and complicates management of these infections. Antibiotic resistance occurs mainly due to chromosomal gene mutations and possession of transferrable resistance determinants. [1] MDR isolates can be clonal, particularly those associated with hospital acquired infections. [2] Clones of *P. aeruginosa* may vary based on the environments [3], and may cause infection outbreaks when these clones enter a new environment. For example, P. aeruginosa isolated from water sources can also be isolated from cystic fibrosis patients [4]. Only a few studies have identified clones of ocular isolates of *P. aeruginosa* [5, 6]. Five multi-drug resistant *P.* aeruginosa isolates from corneal infections have been reported to be clonal and of sequence type 308. [6] The isolates were collected in 1997 from microbial keratitis cases in India. The current study investigated the genomes of more recently collected MDR P. aeruginosa corneal isolates recovered from the same location in India to investigate whether this clonal variant had persisted and whether it had acquired or lost antibiotic resistance genes. Materials and methods P. aeruginosa genomic sequencing DNA was extracted using DNeasy Blood and Tissue Ki (Qiagen, Hilden Germany) as per the manufacturer's recommendations from two keratitis P. aeruginosa strains 198 and 219 isolated in India in 2018. A paired-end library was prepared using Nextera XT DNA library preparation kit (Illumina, San Diego, CA, USA). All the libraries were multiplexed on one MiSeq run. The raw reads of the sequenced genomes were analysed for their quality using **FastQC** version 0.117 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc).

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Version 0.38 of the Trimmomatic [7] was used for trimming the adapters from the reads following de-novo assembly using Spades v3.13.0 [8]. Genomes were annotated using Prokka v1.12 [9]. Sequence types were investigated using PubMLST https://pubmlst.org/. Resistance genes were identified using online database Resfinder v3.1 (Centre for Genomic Epidemiology, DTU, Denmark) [10]. Mutations in the genes were detected using Snippy V2 [11] using PAO1 as a reference genome. Core genome and pan genomes were analysed using Harvest Suite Parsnp v1.2 and Roary v3.11.2 respectively. Integrons were located using Integron finder v1.5.1. The genes possessed by strains 198 and 219 were then compared to those from other ST308 isolates that had been previously examined [6]. Antibiotic resistance Strains 198 and 219 (isolated in 2018) and five strains isolated in 1997 PA31, PA32, PA33, PA35 AND PA37 [6] were screened for resistance to a variety of antibiotics which are commonly used to treat microbial keratitis [12]. The minimum inhibitory concentration (MIC) and minimum bactericidal concentration (MBC) of ciprofloxacin, levofloxacin, gentamicin, ceftazidime (Sigma-Aldrich, St. Louis Missouri, USA), polymyxin B (Sigma-Aldrich, Vandtårnsvej, Søborg, Denmark) tobramycin, piperacillin (Cayman Chemical Company, Ann Arbor, Michigan, USA) and imipenem (LKT Laboratories Inc, Minnesota, USA) were determined using the broth microdilution method in 96-wells plates following CLSI guidelines. The concentrations of antibiotics tested ranged from 5120 µg/ml to 0.25 µg/ml. The susceptibility results were interpreted using EUCAST v9 [13] and CLSI [14] breakpoints for antibiotics. Results Antibiotic susceptibility and Sequence type analysis Isolates 198 and 219 had sequence type 308 indicating that these strains were clonally related

to the ST308 strains isolated in 1997 at the same hospital.

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Isolate 219 was resistant to all antibiotics (Table 1) other than polymyxin (MIC= 0.5 µg/ml, MBC=1 µg/ml). Isolate 198 was resistant to ciprofloxacin, levofloxacin, tobramycin and gentamicin, and showed intermediate susceptibility to polymyxin (Table 1). The isolates from 1997 were all resistant to gentamicin and tobramycin and showed intermediate or definite resistance to imipenem (Table 1). All five isolates from 1997 were resistant or had intermediate resistance to ciprofloxacin and four were resistant to levofloxacin (Table 1). Two isolates from 1997 showed intermediate resistance to polymyxin (Table 1). Overall, the MIC and MBC values to ciprofloxacin and levofloxacin of 198 and 219 were higher than those recorded for the historical isolates (Table 1). Possession of horizontally-acquired resistance genes In total 24 acquired resistance genes were present in the ST308 isolates of *P. aeruginosa* (Table 2). The isolates from 1997 all possessed the same 11 resistance genes. However, the isolates from 2018 had acquired additional resistance genes. Isolate 198 carried 15 and 219 carried 20 resistance genes. Ten resistance genes were common to all seven isolates (Table 2). These ten genes were three aminoglycoside resistance genes (aph(6)-Id, aph(3')-IIb,aph(3'')-Ib), a beta-lactam resistance gene (blaPAO), a tetracycline resistance gene (tet(G)), a fosfomycin resistance gene (fosA), a chloramphenicol resistance gene (catB7) a sulphonamide resistance gene (sul1), and a quaternary ammonium compound resistance gene (qacEdelta1). The recent isolates lacked one beta lactam gene (blaOXA-50) which was present in all the historical isolates. Aminoglycoside resistance genes Strain 198 had acquired a 16S rRNA methylase gene (rmtD2) carried on class 1 integron and three aminoglycoside modifying enzyme genes aph(6)-Id, aph(3')-Ilb, and aph(3'')-Ib). Strain 219 had acquired three different aminoglycoside resistance genes, a 16S rRNA methylase

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(rmtB), a streptomycin adenylyltransferase gene (aadA1) carried on class 1 integron and an aminoglycoside acetyltransferase gene (aac(6')-Ib-cr). Strain 219 had also acquired the plasmid related aminoglycoside and fluoroquinolone resistance gene *aac*(6')-*Ib-cr*. Fluoroquinolones resistance genes One fluoroquinolone resistance gene, *crpP*, was present in all isolates from 1997 and 2018. An integron-related fluoroquinolone resistance gene *qnrVC1* was only present in isolates from 2018 (Table 1) and was carried on class 1 integron on both isolates 198 and 219. The plasmid related aminoglycoside and fluoroquinolone resistance gene aac(6')-Ib-cr was found in strain 219 and this strain had higher MICs for ciprofloxacin and levofloxacin compared to strain 198 and the 1997 isolates. **Beta-lactam resistance genes** The metallo-beta-lactamase gene class B metallo-b-lactamase blaVIM-2 and a transposon (Tn2) encoded gene blaTEM-1B had been acquired by isolate 219 and were carried on class 1 integron. An extended spectrum plasmid-related class A beta lactamase gene blaPME-1 had been acquired by 198 and were carried on class 1 integron. A class-D beta lactamase gene blaOXA-488 had been acquired by both 198 and 219. Non-synonymous mutations in the ST308 resistome Table 3 details the non-synonymous mutations leading to changes in the nucleic acid sequence in the resistance genes of these *P. aeruginosa* isolates, including those related to efflux pumps, antibiotic-inactivating enzymes and drug target alterations. These nonsynonymous mutations were made in comparison to the reference genome of strain PAO1. The number of mutations in almost all of the genes remained same in the 1997 and 2018 isolates. However, the efflux pump gene opmH contained 10 SNPs in the isolates from 2018, but only 1-5 SNPS in the isolates from 1997 (Table 3). Similarly, oprD also contained 8

SNPs in the two 2018 isolates. Furthermore, non-synonymous insertions/deletions [15] and frame-shift mutations were found in the two isolates from 2018 (Table 4) whereas the ST308 isolates from 1997 had no insertions/deletions or frame-shift mutations [6].

Table 1. Antibiotic susceptibility of *P. aeruginosa* isolates

Breakpoints	Ciproflo μg/ml ≤1		Levoflo μg/ml : ≥8	≤2, 4,	Gentamicii ≤4, 8, 3		Tobramyci ≤4, 8, 3		Piperao μg/ml		lmipe μg/ml ≥٤	≤2, 4,	Ceftaz µg/ml : ≥3	≤8, 16,	μg/n	myxin nl ≤2, ≥8
P. aeruginosa Strains of	MIC	МВС	MIC	МВС	MIC	МВС	MIC	МВС	MIC	МВС	MIC	МВС	МІС	МВС	МІС	МВС
198	1280 (R)	2560	320 (R)	1280	2560 (R)	5120	16 (R)	16	8	8	1	2	8	8	4 (I)	4
219	≥5120 (R)	≥5120	640 (R)	1280	≥5120 (R)	≥5120	1280 (R)	2560	2560 (R)	5120	32 (R)	64	16 (I)	32	0.25	1
31	32 (R)	64	32 (R)	32	5120 (R)	5120	640 (R)	1280	4	8	4 (I)	16	16 (I)	32	4 (I)	4
32	64 (R)	128	32 (R)	32	2560 (R)	5120	640 (R)	1280	16	32	4 (I)	4	16 (I)	16	4 (I)	16
33	128 (R)	128	32 (R)	64	2560 (R)	5120	≥5120 (R)	≥5120	32 (R)	64	8 (R)	16	32 (R)	64	2	4
35	2 (1)	4	2	4	2560 (R)	2560	1280 (R)	2560	8	16	16 (R)	16	4	8	2	2
37	64 (R)	128	32 (R)	32	2560 (R)	2560	1280 (R)	2560	8	16	8 (R)	8	16 (I)	64	2	4

P. aeruginosa isolates in the light shade are recent and those with dark shading are historical isolates. MIC and MBC values of historical isolates were included from previously published data [6] for all antibiotics except tobramycin and piperacillin. MICs and MBCs of these two antibiotics for the historical isolates and all antibiotics for the recent isolates have been are examined in this study.

Table 2. Presence of acquired antibiotic resistance genes in *P. aeruginosa* ocular isolates.

Antibiotic	Resistance	198	219	31	32	33	35	37
classes	genes				02			
	aph(6)-Id							
	<i>aph</i> (3')-VI							
Amino-	<i>aph</i> (3')- <i>lIb</i>							
glycoside	aph(3")-Ib							
grycosiae	aadA1							
	rmtD2							
	rmtB							
Fluoroquinolone +	aac(6')-Ib-							
Aminoglycoside	cr							
Fluoroquinolone	crpP							
Thuoroquinorone	qnrVC1							
	blaOXA- 488							
	blaPAO							
Beta lactam	blaOXA-50							
	blaTEM-1B							
	blaVIM-2							
	blaPME-1							
Quaternary ammonium compound	qacEdelta1							
Sulphonamide	sul1							
Tetracycline	tet(G)							
Macrolide	mph(E)							
	mph(A)							
Macrolide, Lincosamide and Streptogramin B	msr(E)							
Chloramphenicol	catB7							
Fosfomycin	fosA							

Red colour denotes gene presence and yellow colour shows gene absence

Table 3. Single nucleotide polymorphism due to non-synonymous mutations in the genes of *P. aeruginosa* genes

Gene Gene		P. aeruginosa/number of SNPs									
locus	name		31	32	33	35	37	198	219		
PA0156	triA		4	5	5	6	5	5 *	5 *		
PA0157	triB	- -	0	0	1	0	0	0	0		
PA0158	triC		2	2	2	2	2	2 *	2 *		
PA0424	mexR		2	1	1	1	1	1	1		
PA0426	mexB		1	1	2	1	4	2 *	2 *		
PA1236	farB		1	1	1	1	1	1	1		
PA1282	lrfA		6	6	9	6	8	8*	8*		
PA1316	lrfA		2	2	2	2	2	2	2		
PA1435	mexM		4	4	4	4	4	4	4		
PA1436	mdtC		2	2	2	2	2	2	2		
PA2018	mexY		5	5	5	5	5	5 *	5 *		
PA2019	mexX		4	4	4	4	4	4	4		
PA2389	macA		2	1	1	1	1	1	1		
PA2390	тасВ		1	1	1	1	1	1	1		
PA2391	opmQ	Antibiotic	6	5	6	6	6	5 *	5 *		
PA2491	mexS	Efflux	2	2	2	2	2	2	2		
PA2495	oprN		1	1	1	1	1	1	1		
PA2837	opmA		3	3	3	3	3	3	3		
PA3019	taeA		1	1	1	1	1	1	1		
PA3137	farB		1	1	1	1	1	0	0		
PA3521	opmE		3	3	3	3	3	3*	3*		
PA3522	mexQ		4	4	4	4	4	4	4		
PA3523	mexP		2	2	2	2	2	2	2		
PA3676	mexK		1	1	1	1	1	1	1		
PA3677	mexJ		2	2	2	2	2	2	2		
PA3678	mexL		1	1	1	1	1	1	1		
PA4205	mexG		1	1	1	1	1	1	1		
PA4206	mexH		1	1	1	1	1	1	1		
PA4207	mexI		1	1	1	1	1	2	1 *		
PA4208	opmD		3	3	3	3	3	2 *	2 *		
PA4374	mexV		2	2	2	2	2	2	2		
PA4375	mexW		2	2	2	2	2	2	2		
PA4598	mexD		2	2	2	2	2	2	2		
PA4599	mexC		7	8	8	8	8	7 *	7 *		

PA4974	ортН		2	1	1	5	5	10 *	10 *
PA4990	emrE		1	1	1	1	2	1	1
PA4997	msbA		2	3	3	2	3	4 *	4
PA5158	adeC		3	3	3	3	3	3 *	3 *
PA5160	farB		4	3	3	4	3	3	3
PA5518	rosB		3	3	3	3	3	2	2
PA0706	catB7		4	4	4	4	4	4	4
PA4109	ampR	Antibiotic	2	2	2	2	2	2 *	2 *
PA4110	атрС	inactivation	5	5	5	5	5	5 *	5 *
PA4119	Aph(3')- IIb		2	2	2	2	2	2	2
PA5514	OXA-50		1	2		3	2	2	2
PA0903	alaS		1	1	1	1	1	1	1
PA1972	pmrC		3	3	3	3	3	3	3
PA3002	mfd	Antibiotic	1	2	2	2	2	2	2
PA3168	gyrA	target	1	1	1	1	1	1	1
PA3946	rosC	alteration	6	8	8	7	6	8*	8
PA4265	tufA		1	0	0	0	1	0	0
PA4560	ileS		2	2	2	2	2	3	2
PA4964	parC		2	2	2	2	2	2	2
PA4967	parE		1	1	1	1	1	1	1
PA3554	arnA		2	4	4	4	4	3*	3*
PA0920	mprF		6	6	6	6	6	6	6
PA0958	oprD		2	1	1	1	4	8	8
PA2492	mexT		3	2	3	5	4	2^{\dagger}	2 †
PA2020	mexZ	11.	0	1	1	1	1	1	1

210 (*) represents insertions or deletions in the genes, (†) represents frame-shift mutations in the genes

Table 4. Insertion/deletion and frame-shift mutation in the resistance genes of the 2018 isolates.

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Gene locus	Genes	Number of deletions and insertions in the genes				
		198	219			
PA0156	triA	2	2			
PA0158	triC	2	1			
PA0426	mexB	1	1			
PA1282	lrfA	2	2			

PA2018	mexY	1	1
PA2391	opmQ	3	3
PA3521	opmE	1	1
PA4208	opmD	1	1
PA4599	mexC	1	1
PA4974	ортН	7	8
PA4997	msbA	2	2
PA5518	adeC	1	1
PA4109	ampR	1	1
PA4110	ampC	2	2
PA3946	rosC	1	1
PA3554	arnA	2	2
PA0958	oprD	6	6
PA2492	mexT	1Frame-shift deletion	1Frame-shift deletion

Phylogeny of ST308 isolates

The genomes of these ST308 isolates were aligned using PAO1 as a reference for the core genome and pangenome phylogeny. In the core genome phylogenetic analysis, all the isolates were clustered together in a single group. The number of core genes and total genes were same for all isolates except isolate 219 which had a larger number of total genes but a similar number of core genes to all other isolates (Table 5).

Table 5. Number of genes present in Core and pan genomes of *P. aeruginosa* isolates.

P. aeruginosa isolates	Core genes	Total genes
31	5445	6937
32	5447	6927
33	5440	6932
35	5442	6932
37	5450	6958
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219	5451	7247

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Discussion This study examined whether the resistome of the ST308 clone of ocular isolates of P. aeruginosa had changed over time. Previously, P. aeruginosa ST308 clones had been reported as multidrug-resistant isolates of nosocomial [16] ocular [6] and canine origin [17]. For the *P. aeruginosa* isolates from three different sources the MIC to imipenem was high which was similar to the finding in the ocular isolates of *P. aeruginosa* in the present study. The ocular isolates of clone ST308 from 2018 had acquired additional resistance genes and had changes in the mutational patterns of the resistance genes compared to ocular isolates from 1997. Two different variants of 16S rRNA methylase, rmtD2 and rmtB, related to aminoglycoside resistance were found in the 2018 isolates. These genes have not been reported previously in ST308 but other variants of the same genes have been identified in clones ST316 and ST235 [18] the latter clone being identified as a widespread multi-drug resistance clone. The presence of a larger group of beta lactam resistance genes, specifically those acquired on mobile genetic elements including class A and B metallo-beta lactam genes including blaTEM-1B, blaVIM-2, blaPME-1 carried on integron is a unique finding related to ST308 in the current study. These beta lactam resistance genes have not been reported previously in strains of this clone [6]. However, the possession of *sul1* gene in isolates of present study was similar to the similar ST308 found previously [17]. The possession of blaVIM-2 and blaTEM-1B may have been responsible for the high MIC to piperacillin and imipenem of PA219. Previously, these genes were associated to increased MIC of imipenem and piperacillin/tazobactam in *P. aeruginosa* isolates [19]. Metallo-beta lactam genes are usually found on class-1 integrons along with other antibiotic resistance determinants [20] which is similar to the present study but identification of class 1

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integron carrying resistance genes in the ocular *P. aeruginosa* isolates is a novel finding. These metallo-beta lactam genes are easily transmissible on mobile genetic elements such as transposons, plasmid-integrative conjugative elements and genomic islands. These metallobeta lactam genes (blaTEM-1B, blaVIM-2, blaPME-1) have not been previously reported in ST308 but have been found in ST111 and ST235 [21] [22]. Although different variants of these genes were found in the similar ST308 before [16, 17]. Acquired genes within the mobile genetic elements of ST308 clones were not been identified in an earlier report [6]. Both recent isolates 198 and 219 had acquired genes associated with mobile genetic elements in the current study. The presence of the plasmid related fluoroquinolone resistance gene *ant* VC1 [23] and the recently reported plasmid related gene crpP [24] are also novel findings in the current study related to clonal ST308 P. aeruginosa isolates. All isolates contained the fluoroquinolone resistance gene crpP, but this had not been identified as a potential plasmid related fluoroquinolone resistance gene prior to the publication of resistance genes of the 1997 isolates [6]. Usually fluoroquinolone resistance is due to mutation in DNA gyrase and topoisomerase IV genes [25]. However, in the 2018 isolates of ST308 very high MICs to ciprofloxacin and levofloxacin might be due to the acquisition of *qnrVC1*. Strain 219 had also acquired the plasmid related fluoroquinolone resistance gene aac(6')-Ib-cr [26] which can confer resistance to both fluoroquinolones and aminoglycosides [27]. Previously this gene was found responsible for the 16 to 128-fold higher MICs for ciprofloxacin in the transconjugants bacteria of family Enterobacteriaceae [28] and MIC of 64 µg/ml of ciprofloxacin to MDR P. aeruginosa isolates [29]. These additional resistance imposing elements to fluoroquinolones suggest that alternative treatments for keratitis other than fluoroquinolone monotherapy should be considered. Acquisition of larger number of aminoglycoside and beta lactam resistance genes is alarming because, where first line therapy

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such as monotherapy with fluoroquinolones fails, fortified antibiotics [30] such as gentamicin plus cephalosporins are often prescribed. Among all the *P. aeruginosa* isolates, the core genome was composed of almost similar number of genes which was perhaps indicative of the collinear nature of conserved genome of *P. aeruginosa* isolates .[31, 32] However, a larger pan genome of isolate 219 indicates greater genomic diversity due to acquisition of genes from the same or different species or genera. This fact might relate to the larger number of acquired genes in isolate 219 by horizontal gene transfer. [31] Identification of indels (insertion/deletion polymorphisms due to non-synonymous mutations) in the 2018 keratitis isolates of *P. aeruginosa* which were not present in the strains isolated in 1997 [6] as well as the increased presence of certain SNPs suggest that there was an increase in selection pressure in the environment that has selected for these mutations. Increases in the resistance of keratitis isolates to the fluoroquinolone moxifloxacin have been associated with an increase in average diameter of the infiltrate or scar, a slower time to reepithelialization and decrease in final visual acuity. [33, 34] Therefore, the findings from the current study showing that strains of P. aeruginosa, at least in this Indian environment, have gained additional resistance genes and higher levels of resistance suggests that treatment of keratitis might be becoming more problematic. **Nucleotide accession** The nucleotide sequences are available in the GenBank under the Bio project accession number PRJNA590804. Acknowledgements The authors would like to acknowledge the Singapore Centre for Environmental Life Sciences Engineering (SCELSE), whose research is supported by the National Research Foundation Singapore, Ministry of Education, Nanyang Technological University and

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