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Within-host whole genome analysis of an antibiotic resistant *Pseudomonas aeruginosa* strain sub-type in cystic fibrosis

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**Abstract**

*A Pseudomonas aeruginosa* AUST-02 strain sub-type (M3L7) has been identified in Australia, infects the lungs of some people with cystic fibrosis and is associated with antibiotic resistance. Multiple clonal lineages may emerge during treatment with mutations in chromosomally encoded antibiotic resistance genes commonly observed. Here we describe the within-host diversity and antibiotic resistance of M3L7 during and after antibiotic treatment of an acute pulmonary exacerbation using whole genome sequencing and show both variation and shared mutations in important genes. Eleven isolates from an M3L7 population (*n* = 134) isolated over 3 months from an individual with cystic fibrosis underwent whole genome sequencing. A phylogeny based on core genome SNPs identified three distinct phylogenetic groups comprising two groups with higher rates of mutation (hypermutators) and one non-hypermutator group. Genomes were screened for acquired antibiotic resistance genes with the result suggesting that M3L7 resistance is principally driven by chromosomal mutations as no acquired mechanisms were detected. Small genetic variations, shared by all 11 isolates, were found in 49 genes associated with antibiotic resistance including frame-shift mutations (*mexA*, *mexT*), premature stop codons (*oprD*, *mexB*) and mutations in quinolone-resistance determining regions (*gyrA*, *parE*). However, whole genome sequencing also revealed mutations in 21 genes that were acquired following divergence of groups, which may also impact the activity of antibiotics and multi-drug efflux pumps. Comparison of mutations with minimum inhibitory concentrations of anti-pseudomonal antibiotics could not easily explain all resistance profiles observed. These data further demonstrate the complexity of chronic and antibiotic resistant *P. aeruginosa* infection where a multitude of co-existing genotypically diverse sub-lineages might co-exist during and after intravenous antibiotic treatment.
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**Competing interests:** The authors have declared that no competing interests exist.

**Introduction**

Cystic fibrosis (CF) is the most common lethal recessively inherited disease in Caucasians. Respiratory disease secondary to chronic airway infection is the major complication in CF, which causes the majority of mortality and morbidity in patients. *Pseudomonas aeruginosa* is the most common respiratory pathogen isolated from people with CF [1]. Person-to-person transmission of *P. aeruginosa* has been documented in CF clinics globally with adverse clinical outcomes correlated with some shared strains [2–5]. In Queensland, Australia, a predominant *P. aeruginosa* shared strain, AUST-02, has been detected in patients at six clinics and in 16 clinics nationally (18% of patients with *P. aeruginosa* infection) [2,6–8]. We recently reported the emergence of an AUST-02 strain sub-type, M3L7, at The Prince Charles Hospital (TPCH, Brisbane, Queensland, Australia) characterized by the *mexZ*-M3 and *lasR*-L7 alleles [9]. This strain infected 7.6% adults with CF in 2007–2009 and 6.4% adults with CF in 2011 [9]. Patients infected with the M3L7 sub-type had greater treatment requirements and a higher 3-year risk of death or lung transplantation compared to patients infected with other AUST-02 sub-types and *P. aeruginosa* strains [9].

It is widely recognized that there is substantial phenotypic and genetic intra-strain *P. aeruginosa* population diversity within the chronically infected CF airway with the emergence of multiple clonal lineages and strains with antibiotic resistance, hypermutation (caused by mutations in DNA mismatch repair [MMR] genes) and pathoadaptive mutations that enable adaptation to the CF airways all demonstrated [10–18].

Whilst the M3L7 sub-type has been associated with increased antibiotic resistance compared to other *P. aeruginosa* strains including other sub-types of AUST-02, the diversity within an individual is unknown [9]. Furthermore, antibiotic resistance mechanisms are recognized as diverse amongst *P. aeruginosa* but the within-host variation is less clear [19]. Here we used whole genome sequencing (WGS) to investigate the M3L7 sub-type isolated during and after intravenous antibiotic treatment of a pulmonary exacerbation with the principle aim of constructing a genomic analysis of antibiotic resistance gene mutations.

**Materials and methods**

**Overview**

This study focused on a single patient with CF identified as harboring the AUST-02 shared strain sub-type, M3L7 (hereafter referred to as M3L7), on the basis of *mexZ* and *lasR* DNA sequencing [9]. In brief, sputum samples were collected at five time-points during and after treatment of an acute pulmonary exacerbation (a 3 month period). All AUST-02 isolates identified were screened for the M3L7 sub-type. M3L7 isolates were then selected for WGS on the basis of antibiotic susceptibility data.

**Ethical approval**

Ethics approval for this project was granted under HREC/13/QPCH/127 by TPCH Human and Research Ethics Committee, Metro North Hospital and Health Service, Brisbane, Queensland, Australia and the participant provided written, informed consent.

**Patient details**

The patient was a 40-year old male with severe lung disease (forced expiratory volume in one second of 44% predicted), who was admitted to TPCH for treatment of an acute pulmonary exacerbation in February 2014. The patient previously tested M3L7 positive in 2007 (isolate ID, AUS951) [9].
Hypersensitivity reactions had previously complicated courses of extended activity anti-
pseudomonal penicillin (ticarcillin/clavulanate) and cephalosporin antibiotics (ceftazidime)
thus limiting the choice of β-lactam antibiotics to treat pulmonary exacerbations. Based on the
patient’s previous response to treatment, meropenem (1 g three times daily) and tobramycin
(300 mg once daily), which have different mechanisms of action, were administered intrave-
nously on admission. However, on this occasion meropenem was ceased and substituted with
intravenous aztreonam (3 g three times daily) on day 14 due to a suboptimal clinical response
and was continued until hospital discharge (day 23). Clinical stability was maintained until
outpatient follow-up 8 weeks later (from date of discharge). The timeline of antibiotic treat-
ment is shown in Fig 1.

The patient received long-term oral azithromycin (250 mg once daily) throughout the
study (Fig 1). The patient did not receive alternate month inhaled therapies (intolerance
with nebulized colistin and patient declined nebulized tobramycin due to perceived lack of
efficacy).

Patient isolates

Expectorated sputum was collected (Fig 1) and processed to isolate single colonies (method
described in S1 File). Presumptive *P. aeruginosa* isolates (*n* = 48) were randomly selected from
each sputa collected at five time-points: day 1 (start of intravenous antibiotics), day 7 (week 1),
day 14 (week 2), day 23 (completion of intravenous antibiotics) and outpatient follow-up
(8-weeks later). All isolates underwent strain typing using an allele-specific AUST-02 PCR (S1
File) and Sequenom iPLEX SNP-based strain typing [20]. All confirmed AUST-02 isolates
were further differentiated by mexZ-specific PCR and lasR gene sequencing as described previ-
ously [9]. Susceptibility testing to 11 anti-pseudomonal antibiotics (Table 1) was performed
using the disc diffusion assay with isolates categorized as resistant, intermediate resistant or
susceptible to each antibiotic according to Clinical and Laboratory Standards Institute (CLSI)
breakpoint guidelines [21].

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The timeline of antibiotic treatment and sputum collection is shown in Fig 1. The length and timing of a specific antibiotic treatment is shown by different line-styles. The days of sputum collection are shown by filled-in circles (days 1, 7, 14, 23) and 8 weeks following discharge (day 79). Isolates cultured at each time-point and used in this study are also indicated.

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Selection of isolates for whole genome sequencing

Based on the disc diffusion susceptibility data, 11 isolates were selected for WGS (Table 1 and Fig 1). Two M3L7 isolates with a different susceptibility profile were chosen at each time-point (day 1, AUS959 and AUS960; day 7, AUS961 and AUS962; day 14, AUS963 and AUS964; day 23, AUS965 and AUS966; Follow-up, AUS967 and AUS968; Table 1). An additional isolate (AUS969; M3L43) that was cultured at day 23, harboring a different lasR sequence, also underwent WGS (Table 1).

Whole-genome sequencing

Preparation of genomic DNA for WGS was undertaken using the UltraClean R⃝ Microbial DNA Isolation Kit (Mo Bio) with several modifications as follows: prior to DNA extraction, samples were placed on ice (1 hour); the pelleted sample was washed with 0.9% saline and centrifuged at 14,000 x g for 1 min; during cell lysis, the samples were heated to 70°C for 10 mins with bump vortexing for 15 s every 2 mins; during protein removal, the centrifugation speed
used was 12,000 x g; during re-suspension, the DNA was eluted to a final volume of 100 μL. Quantity and quality of the genomic DNA preparations was determined using NanoDrop Spectrophotometry, Quant-iT™ PicoGreen® dsDNA Reagent chemistry, and 0.8% visual gel analysis. Library preparation (Truseq), qPCR (TapeStation, Agilent Genomics) and WGS using the Illumina HiSeq 2500 platform with 100 bp paired-end read chemistry were carried out at the Australian Genome Research Facility, Melbourne, Australia.

**Genome mapping and assembly**

Reads were examined for contamination using Kraken (v0.10.4), quality filtered with Nesoni (v0.128), and mapped to the PAO1 reference genome (Accession number: NC_002516), using SHRiMP2, as implemented in Nesoni [22–25]. SNPs and small insertions or deletions were called using Nesoni. WGS reads were assembled with Velvetoptimiser (v2.2.5) and Velvet (v1.2.10) [26,27]. Resulting contigs were reordered against PAO1 using Mauve (v.2.4.0) and annotated using Prokka (v1.10) [28,29]. Gene annotations from PAO1 were used as the primary reference.

**Phylogenetic analysis**

The core genome alignment was generated from the consensus sequence of the mapped reads against PAO1 comprising a total of 1753 polymorphic positions. Recombination filtering was carried out using Gubbins (v1.4.9), with 1534 polymorphic positions remaining after filtering [30].

Maximum-Likelihood phylogenetic trees were reconstructed from the pre-filtered and recombination filtered core genome alignment using RAxML (v8.1.15), with the rapid bootstrap option (-f a) and the general time reversible model with a gamma distribution of invariant sites [31]. The Lewis correction for ascertainment bias (-m ASC_GTRGAMMA –asc_corr lewis) was used for the recombination filtered SNP-only alignment. The genome of a different AUST-02 strain sub-type (M3L1; isolate ID, AUS970) identified as part of an ongoing study by our group from another CF patient was used as an outgroup to root the tree.

**Etest® susceptibility testing**

In order to obtain minimum inhibitory concentrations (MICs) of the 11 anti-pseudomonal antibiotics (that were used to select isolates for WGS), each M3L7 isolate that underwent WGS was inoculated onto Mueller-Hinton Agar (Thermo Fisher Scientific) with susceptibility testing determined by Etest® (BioMérieux) according to the manufacturer’s instructions. Breakpoints were categorized as susceptible, intermediate resistant or resistant as defined by the CLSI guidelines [21].

**Growth curve analysis**

The fitness of the M3L7 sub-type was assessed by determining the total viable count of M3L7 to that of PAO1 over 24 hours and comparing the doubling time during the logarithmic phase of growth. The initial inoculum of each M3L7 isolate (n = 11) and PAO1 (n = 3, biological replicates) was prepared to ~1 X 10^5 colony-forming units (CFU)/mL in Luria-Bertani Broth (20 mL). Bacterial cultures were incubated aerobically with shaking (200 rpm) at 37˚C and total viable counts were enumerated on Mueller-Hinton agar at 0, 2, 4, 6, 10 and 24 hours.

**Identification of genes associated with antibiotic resistance, hypermutation and pathoadaptation**

Screening for antimicrobial resistance genes was performed by a BLAST alignment of all predicted coding regions against CARD (Comprehensive Antimicrobial Resistance Database),
MERGEM (Mobile Elements and Resistance Genes database Enhanced for Metagenomics) and ResFinder databases [32–34].

Single or multiple nucleotide substitutions, insertions or deletions in chromosomally encoded genes \((n = 136)\) associated with antibiotic resistance, hypermutation and pathoadaptive genes (identified based on a literature search; S2 Table), and intergenic regions were determined using both read mapping and sequence alignment of assembled contigs. Variants were processed and visualized using Nesoni’s nway function and the Harvest suite of tools (Parsnp and Gingr) (v1.1.2) [35]. Non-synonymous mutations were manually inspected and only unambiguous SNP calls were included in the analysis. Functionally important mutations were defined as premature stop codons or frame-shift mutations and recognized point mutations within the quinolone resistance-determining regions (QRDRs) of \(\text{gyrA, gyrB, parC}\) and \(\text{parE}\) [19,36]. The effect of amino acid changes on protein function (effect or no effect) can be predicted using computational algorithms [37]. PROVEAN (Protein Variation Effect Analyzer) was used to filter the remaining missense and in-frame mutations into those that are predicted to be functionally important or neutral using the default score thresholds [38]. A ternary plot of amino acid variants was constructed using EvolView [39].

Whole genome comparisons

Comparative genomic analyses were also performed to identify larger genomic variations using Parsnp, Gingr, BRIG (BLAST Ring Image Generator), Roary, ACT (Artemis Comparison Tool), bandage (a Bioinformatics Application for Navigating De novo Assembly Graphs Easily) and BLAST [35,40–43]. Prophage sequences were annotated with PHAST (PHAge Search Tool) [44].

Accession numbers

Genome sequence data was deposited at the European Nucleotide Archive under study PRJEB14771 with accession identifiers ERS1245422 to ERS1245432. Samples AUS940, AUS951 and AUS970 are available as part of a separate study (PRJEB14781).

Results and discussion

The \(P.\ aeruginosa\) AUST-02 strain sub-type M3L7 infects the airways of some patients with CF in Queensland and has been associated with antibiotic resistance [9]. We performed whole genome analyses of 11 M3L7 isolates isolated over a 3 month period, from a single patient and subsequently, investigated mutations in genes associated with antibiotic resistance, hypermutation and pathoadaptation.

In this study we demonstrate that the M3L7 population was heterogeneous when observed during treatment of a pulmonary exacerbation and subsequent recovery period highlighting the complexity of chronic M3L7 airway infection in CF. We detected variation in chromosomally encoded genes associated with antibiotic resistance, which may reflect adaptation of M3L7 isolates to different antibiotic selection pressures in spatially heterogeneous regions of the airway [14,45]. Moreover, our study design allowed us to identify mutations in target genes, shared by each isolate sequenced, that might also contribute to the high-level antibiotic resistance of the M3L7 sub-type.

Notably, despite the patient harboring multi-drug resistant M3L7 isolates, clinical response to the antibiotic regimen administered was achieved after a lengthy course of intravenous antibiotics (23 days). This corroborates with others who showed a lack of correlation between predicted susceptibility and clinical outcomes in CF [46–48].
Phenotypic variation in ciprofloxacin and tobramycin resistance in the M3L7 population sampled

Genotyping confirmed 234/240 (97.5%) isolates as *P. aeruginosa* and showed that 134 (57%) belonged to the AUST-02 strain. Of these, 131 (98%) showed an M3L7 sub-type and three were of the M3L43 sub-type (2%). The M3L43 isolate (isolate ID, AUS969) was derived from the M3L7 sub-type having acquired an additional SNP in the *lasR* gene (S1 Table).

Disc diffusion susceptibility results for the 134 isolates are presented in Table 1. Studies elsewhere demonstrated that antibiotic susceptibility profiles might differ between isolates of the same strain, even when isolated from individual samples or different regions of the lung [13,45]. Such differences were noted here (19 different antibiotic susceptibility profiles observed; Table 1), with the greatest variation in antibiotic susceptibility detected for ciprofloxacin (resistant: n = 39/134, 29%; intermediate resistant: n = 35/134, 26%; susceptible: n = 60/134, 45%; Table 1) and tobramycin (resistant: n = 48/134, 36%; intermediate resistant: n = 12/134, 9%; susceptible: n = 74/134, 55%; Table 1). Between day 1 of treatment of a pulmonary exacerbation and at follow-up during clinical stability, resistance to ciprofloxacin and tobramycin also varied. Antibiotic resistance to both antibiotics within the *P. aeruginosa* population sampled was greatest at day 14 of treatment (ciprofloxacin resistant: n = 22/37, 59%; tobramycin resistant: n = 22/37, 59%; Table 1). Of interest, the patient was not prescribed ciprofloxacin during the 3 month assessment period; however, several courses of ciprofloxacin (oral, 750 mg twice daily for two weeks) were prescribed at an outpatient clinic in the prior 12 months, including one month prior to recruitment, for treatment of mild exacerbation episodes. Therefore, resistance may have been acquired previously in some isolates and maintained in the absence of ongoing ciprofloxacin exposure.

**M3L7 hypermutators and non-hypermutators co-exist in the CF airway**

Our results corroborate with other recent studies, which have reported that multiple related *P. aeruginosa* lineages can co-exist within a single patient [10,12,15,16,45]. A previous study also reported the co-existence of hypermutators and non-hypermutators over nine years within a single patient with CF [17]. Although only 8% (n = 11/134) of the total AUST-02 population sampled over 3 months was sequenced, our data provide further evidence of this occurrence; the draft genomes of the M3L7 sub-type comprised 96–133 contigs with a total length between 6.19 to 6.24 Mbp (S1 Fig and S3 Table). Isolates could be stratified into 1 of 3 phylogenetic groups (Group A; Group B; Group C) as shown in Fig 2 and isolates within these distinct groups co-existed throughout the sampling period. Longer branch lengths were found to separate isolates within Groups B and C from Group A. The higher number of SNPs that were observed for isolates of Groups B and C is suggestive of a hypermutable phenotype that may be caused by mutations in MMR genes [16,49]. In fact, a frame-shift mutation caused by a 1 bp deletion (1660delT) within the MMR gene, *mutL*, was identified for Groups B and C, but not Group A, consistent with the acquisition of this mutation prior to the divergence of Groups B and C (Figs 2 and 3).

Of note, the M3L7 isolate (isolate ID, AUS951) cultured from the patient in 2007 clustered with Group A non-hypermutator isolates and was most closely related to AUS961 (differed by 14 SNPs, data not shown).

**Variation in chromosomally encoded genes was observed**

Based on the CLSI guidelines, all isolates were categorized as resistant (S3 Fig) to aztreonam (MIC range 64 to >256 mg/L), ceftazidime (MIC >256 mg/L), cefepime (MIC >256 mg/L),...
imipenem (MIC > 32 mg/L), meropenem (MIC > 32 mg/L) and ticarcillin/clavulanate (MIC > 256 mg/L) and resistant or intermediate resistant to amikacin (range MIC 32 to 256 mg/L). In contrast, all isolates were categorized as susceptible (S3 Fig) to colistin (MIC range 0.125–0.5 mg/L) and polymyxin B (MIC range 0.25–1 mg/L). Ciprofloxacin (MIC range 0.5–8 mg/L) and tobramycin (MIC range 2–64 mg/L) showed the greatest variability in MICs (S3 Fig) with isolates categorized as resistant, intermediate resistant and susceptible (similar to disc diffusion results that were used to select isolates for WGS, Table 1). The S2 File contains the data set of the individual isolate MICs.

Analysis of each M3L7 genome via multiple antibiotic resistance gene databases failed to identify acquired antibiotic resistance genes suggesting M3L7 resistance is principally driven...
M3L7 isolates were cultured from sputum samples during intravenous treatment of an acute pulmonary exacerbation (Days 1, 7, 14 and 23) and at outpatient follow-up 8 weeks later. Protein sequences (compared to PAO1) were grouped according to the resistance they confer or to their biological function and three colors are used to filter the proteins. The complete plot with the full set of chromosomally encoded proteins investigated, genetic mutations identified and PROVEAN scores are available in S1 Table.

*Premature stop codon; \*FtsI (G63C) identified in all other M3L7 isolates (Group A and C); ^Additional frame-shift mutation detected in AUS964.

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by chromosomal mutations. In total, mutations, shared by all 11 isolates, were identified in 49 chromosomal genes previously implicated in conferring *P. aeruginosa* antibiotic resistance (S1 Table). Whilst the impact of most SNPs is unknown, functionally important mutations are likely to include frame-shift mutations (*mexA, mexT*) and premature stop codons (*oprD, mexB*). Shared mutations that have been previously described were also identified including premature stop codons within *oprD* (E176 identified in M3L7) that could result in loss of the outer membrane porin and contribute to carbapenem resistance and an 8 bp deletion (226_233delCGGCCAGC) in *mexT* characteristic of nfxC-mutants that display mexEF-oprN overexpression but reduced oprD transcription [19,50–53]. A point mutation in *ampC* (T105A) that confers reduced susceptibility to imipenem, ceftazidime and cefepime when the enzyme is overexpressed was also identified [54]. Although no recognized mutations were observed that easily explain *ampC* derepression, a novel nucleotide substitution within the *ampC-ampR* intergenic region (PAO1, TAGAAT; M3L7, TATATT) was identified in all 11 isolates and was found to reside within the -10 promoter region of *ampC*, which could affect production of AmpC and anti-pseudomonal penicillin and cephalosporin resistance [55].

WGS revealed small genetic variations in 21 genes that may impact the activity of β-lactams (e.g. mutations in penicillin-binding proteins), aminoglycosides (e.g. mutations that may affect uptake) and multi-drug efflux pumps (e.g. mutations in cytoplasmic membrane components) between M3L7 groups or individual isolates (Figs 2 and 3). There were no mutations that could be easily correlated with ciprofloxacin and tobramycin MIC variability. A novel point mutation outside the QRDR was observed in *parC* (V48A) amongst Group C isolates (3/4 of these isolates were resistant to ciprofloxacin [4–8 mg/L] and the remaining isolate was intermediate resistant [2 mg/L]; Fig 2), which was acquired by this group after divergence from Group B and was predicted *in silico* to affect protein function (Fig 3). Other mutations identified that potentially contribute to ciprofloxacin resistance (e.g. *gyrA, T83I; parE, A473V; mexT, 8 bp deletion*) were shared by all isolates [MIC range, 0.5–8 mg/L] (S1 Table) [36,51]. Mutations only acquired by the tobramycin-resistant isolate (AUS961, 64 mg/L) and associated with aminoglycoside resistance were not identified and isolates with intermediate resistance to amikacin (AUS963, 32 mg/L) and tobramycin (AUS966, 12 mg/L) appeared genetically similar to isolates with resistant phenotypes when based on the panel of target chromosomally encoded genes investigated (Fig 2). Further work is required to investigate the effect of the mutations detected on antibiotic resistance and to determine their prevalence in a larger number of isolates [56].

It has been proposed that certain genes are involved in the parallel evolution and adaptation of different *P. aeruginosa* strains to the host [18]. Overall, 38/52 (73%) candidate pathoadaptive genes harbored small genetic variations in the M3L7 isolates sequenced (S1 Table) [18]. Some mutations found in these pathoadaptive genes (S1 Table) were acquired after divergence of groups (e.g. a frame-shift mutation in *mucA* was acquired by Group B isolates) [18]. Furthermore, growth of the M3L7 sub-type (mean doubling time, 59 mins) was slower than PAO1 (mean doubling time, 32 mins) suggesting that an *in vitro* fitness cost is associated with the sub-type (S4 Fig). The S2 File contains the data set of the total viable counts for each isolate.

Altogether the data emphasize the complexity of chronic M3L7 infection where genetically diverse sub-lineages, with variation in antibiotic resistance and pathoadaption genes, co-exist.

The patient was infected with multiple *Pseudomonas aeruginosa* strains

Recently there has been a substantial increase in the survival of people with CF, even in those with advanced pulmonary disease [57], and consequently the lifetime exposure to antibiotics is
rising. In parallel, the emergence of antibiotic-resistant CF pathogens are increasingly common and antibiotic toxicity challenges the effectiveness and limits the choice of antibiotic regimens, as described for the patient included here. In addition to infection with M3L7, the patient was co-infected with two other recognized shared strain-types, AUST-06 (n = 25/234; 11%) and AUST-07 (n = 75/234; 32%) that have been found in a number of patients attending TPCH Adult CF Centre [2]. This further demonstrates the challenges of treating chronic P. aeruginosa infection where multiple strain types and strain sub-lineages with differing susceptibility profiles may all be present [10,12,15,45].

Therefore, there is an urgent need for the development of new antibiotics to treat multi-drug resistant P. aeruginosa, new antibiotic formulations and adjuvant antimicrobial agents that can improve the action of current treatments [58–60]. In the future it may be possible to commence a personalized treatment approach that will consider the resistance profile of the complex P. aeruginosa population and limit pathoadaptation of bacteria residing within the lungs.

Limitations

A potential limitation of this study is the relatively small proportion of available isolates that we have sequenced. Further work is required to determine the prevalence of each M3L7 group (A, B, C) at different time-points or determine if one group dominated during or after treatment of an acute pulmonary exacerbation and if there is evidence of periodic selection [10]. Given the small sample size, we also could not directly correlate susceptibility profile and genotype but the results indicate that discrepancies are likely and that a cautious approach should be taken if susceptibility is predicted using genomic analyses of single isolates in chronic infections given the diversity observed [56].

Conclusions

We have provided a short-term within-host analysis of an antibiotic resistant P. aeruginosa strain sub-type in CF using whole genome data. The findings suggest that diversity exists in the M3L7 population and variation in chromosomally encoded genes associated with resistance, hypermutation and pathoadaptation were observed. Discrete mutational events might have accumulated to enable the M3L7 sub-type to continually evolve, diverge and adapt to the spatially heterogeneous niches of the CF airways. These results have broader implications for interpreting antimicrobial susceptibility profiles predicted by genomic analyses of individual isolates or metagenomic analysis of sputum samples during acute pulmonary exacerbations.

Supporting information

S1 Fig. Complete genome sequences of the M3L7 isolates (listed in the key) compared to the AUS940 reference genome (M3L7 isolate from a different patient identified from an ongoing study) using BLAST Ring Image Generator (BRIG) software. Each genome is grouped according to phylogeny. *Two strains (Group A, AUS961 and AUS962) have lost a 40 Kbp prophage between 3900 and 4000 Kbps. This prophage was not annotated with antibiotic resistance genes.

(TIF)

S2 Fig. Phylogeny of M3L7 isolates following recombination filtering. Recombination filtering was carried out using Gubbins. The phylogenetic tree was constructed based on a core SNP alignment of 1534 nucleotides generated from read mapping against PAO1. The sequenced genome of a M3L1 strain (isolate ID, AUS970), as part of an ongoing study, was
used as an out-group to root the tree. The scale bar represents 100 nucleotide substitutions. ** Indicates branches with 100% support from 1000 bootstrap replicates. **Abbreviation**: MRCA, most recent common ancestor.

(SIF)

S3 Fig. Minimum inhibitory concentrations (mg/L) of M3L7 isolates \((n = 11)\) against 11 anti-pseudomonal antibiotics. Isolates were categorized as resistant, intermediate or susceptible according to CLSI guidelines. The median and interquartile range are shown. Any isolates recorded as having a MIC greater than the maximum \((> 256 \text{ mg/L} \text{ or } > 32 \text{ mg/L})\) value on the Etest\textsuperscript{®} strip are shown as double the maximum concentration.

(PDF)

S4 Fig. Geometric mean (with 95% confidence intervals) total viable counts expressed as colony-forming units (CFU)/mL of the M3L7 sub-type \((n = 11)\) compared to PAO1 \((n = 3)\) grown in Luria-Bertani Broth.

(PDF)

S1 Table. Ternary plot of amino acid sequence variation.

(XLSX)

S2 Table. Literature search.

(DOCX)

S3 Table. General genome features of isolates within the M3L7 sub-type.

(DOCX)

S1 File. Collection and processing of sputum samples and Allele-specific PCR for AUST-02.

(DOCX)

S2 File. Data sets of individual data points relating to S3 Fig and S4 Fig.

(DOCX)

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