Genome sequence of *Pseudomonas aeruginosa* PAO1161, a PAO1 derivative with the ICEFP2 integrative and conjugative element.

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ABSTRACT

Pseudomonas aeruginosa is a common cause of nosocomial infections, especially in patients with cystic fibrosis and burn wounds. An attenuated PAO1 strain and its derivatives are widely used to study the biology of this bacterium, however recent studies indicated an ongoing evolution of the laboratory strains, highlighted by changes in the genomes of PAO1 sublines and derivatives used in different laboratories worldwide.

Here we have sequenced the genome of *Pseudomonas aeruginosa* PAO1161 strain, a *leu*-, Rif^R, restriction-modification defective PAO1 derivative. This strain is described as the host of IncP-8 plasmid FP2, conferring the resistance to mercury and used in the past for preparation of genetic map of the *P. aeruginosa* chromosome. Comparison of PAO1161 genome sequence with PAO1-UW reference genome revealed an inversion of the large genome segment between *rrnA* and rrnB rRNA operons and more than 100 nucleotide polymorphisms and short insertions-deletions, many of which were also found within the recently re-sequenced PAO1 sublines. The PAO1161 specific sequence variants include E108K mutation in PA3792 (leuA), which we showed is responsible for leucine auxotrophy, H531L in PA4270 (rpoB), likely conferring rifampicin resistance, as well as nonsense mutations in PA2735 encoding a DNA methyltransferase as well as a putative OLD family endonuclease PA1939. Additionally, PAO1161 genome poses a 12 kb RPG42 prophage and a 108 kbp insertion within tRNA-Lys, encompassing a putative mercury resistance and showing similarity to PAPI-1 like integrative conjugative elements (ICE). Our data indicate that the ICE can excise from the bacterial chromosome and can be transferred to *Pseudomonas putida* cells where it integrates in the genome at three specific sites and confers the cells with mercury resistance. Overall this data provide insight into the genome variation between P. aeruginosa laboratory strains. Moreover, this data indicate that the FP2 element for years described as a IncP-8 plasmid is an integrative conjugative element.

INTRODUCTION

Pseudomonas aeruginosa is a Gram-negative gammaproteobacterium commonly found in various ecological niches and characterized by the ability to survive in very unfavourable, frequently changing environmental conditions. Being a human opportunistic pathogen, it is often the cause of nosocomial infections in immuno-compromised patients. In cystic fibrosis patients *P. aeruginosa* chronically colonizes the lungs and is a major factor causing mortality (Klockgether and Tümmler, 2017; Azam and Khan, 2018).

Majority of research on this metabolically versatile bacterium uses sublines or derivatives of P. aeruginosa PAO1 strain, originally isolated in prof. Bruce Holloway laboratory from a wound of a patient in Melbourne, Australia (Holloway, 1955) Over the years, the strain was shipped to laboratories worldwide and different derivatives, including auxotrophic strains and strains with mobile genetic elements were obtained (Stanisich and Holloway, 1969). In 1999 the genome of P. aeruginosa PAO1 stored at the University of Washington (PAO1-UW) was sequenced (Stover et al., 2000), providing a reference for studies on *P. aeruginosa* genomes. As of November 2018, the Pseudomonas Genome Database, a database devoted to the information on Pseudomonas species (Winsor et al., 2016), contained 2226 sequenced P. aeruginosa genomes, including 11 PAO1 derivatives (sublines). Remarkably, sequencing of PAO1 subline (MPAO1) as well as PAO1-DSM strain stored at the German Collection for Microorganisms and Cell Cultures (DSMZ) revealed presence of multiple short sequence variants which could lead to the variations in virulence and fitness between strains used in different laboratories (Klockgether et al., 2010). A major feature differing genomes of PAO1 derivatives is the occurrence of a large inversion resulting from a homologous recombination between two rRNA operons rrnA and rrnB (Stover et al., 2000), which is present in the reference PAO1-UW genome and not in MPAO1 and PAO1-DSM (Klockgether et al., 2010). Recent data indicate that, despite displacing the *dif* region, this inversion does not seem to affect chromosome segregation and such large rearrangements could be common among bacteria (Bhowmik et al., 2018). This data indicate an ongoing micro- and macro- evolution of bacterial genomes and suggests that sequence diversification in laboratory strains should be taken into consideration in the analysis of phenotypic data (Sidorenko et al., 2017; Luong et al., 2014).

In this work we focus on the genome of *P. aeruginosa* PAO1161 strain, a PAO1 derivative requiring leucine for growth on minimal media and selected as lacking the restriction-modification system (*rmo*-10 mutation) (Dunn and Holloway, 1971). This strain is described as the host of IncP-8 plasmid FP2, the only known member of this incompatibility group, conferring bacteria with the resistance to mercury (Finger and Krishnapillai, 1980; Shintani et al., 2015). The FP2 factor has chromosome-mobilizing ability (Cma) and was extensively used in interrupted mating technique for preparation of the genetic map of *P. aeruginosa* chromosome (Stanisich and Holloway, 1969; Holloway and Fargie, 1960).

PAO1161 derives from the PAO38 *leu-38* mutant obtained by treatment of PAO1 with manganus chloride and selection for leucine auxotrophy (Holloway, 1955) (Figure 1A). PAO38 acquired the FP2 element from PAT (*P. aeruginosa* strain 2) (Stanisich and Holloway, 1972) to yield strain PAO170 (Kung and Lee, 1975). Finally, PAO1161 was selected as defective in both restriction and modification systems, called previously host-controlled modification (HCM), following mutagenesis of PAO170 with N-methyl-N'-nitro-N-nitrosoguanidine and a selection, based on the altered susceptibility to phage infection (Rolfe and Holloway, 1968; Dunn and Holloway, 1971). To facilitate genetic manipulations in this strain, *eg.* pAKE600 based allele exchange (El-Sayed et al., 2001), a spontaneus PAO1161 rifampicin resistant clone was obtained (Lasocki et al., 2007). The PAO1161 strain is used in our studies of chromosome segregation and gene expression in *P. aeruginosa* using genome wide approaches (Kawalek et al., 2018; Bartosik et al., 2014; Kawalek et al., 2017) but it is also used in other physiological and genetic studies (Ciok et al. 2018; Komatsu et al., 1994; Manavathi et al., 2005; Kujawa et al., 2017; Papagiannitsis et al., 2013; Laudy et al., 2015; Romaniuk et al., 2017).

Here we report the genome sequence of *P. aeruginosa* PAO1161 strain. Comparison with PAO1-UW reference sequence revealed the presence of a large number of single-nucleotide polymorphisms (SNPs), insertions-deletions (indels), as well as inversion of large genome segment between rRNA genes. Moreover a functional PAPI-1 like integrative conjugative element (ICE), containing a mercury resistance operon was identified in PAO1161 genome, indicating that the FP2 factor is not a plasmid but an ICE named ICEFP2.

RESULTS AND DISCUSSION

Pseudomonas aeruginosa PAO1161 genome was sequenced and assembled, resulting in a single circular chromosome of 6383803 bp. The GC content (66.42 %) was similar to those of other *P. aeruginosa* genomes (Mathee et al., 2008). A comparison with the reference PAO1-UW genome (NC_002516) revealed three major structural differences (Figure 1B). PAO1161 genome lacks the large inversion between ribosomal RNA operons *rrnA* and *rrnB* observed in PAO1-UW but not in other PAO1 derivatives including MPAO1 and PAO1-DSM (Klockgether et al., 2010). Additionally, PAO1161 poses two large insertions. The 107796 bp insertion in tRNA-Lys gene between *lepA* (*PA4541*) and *clpB* (*PA4542*), flanked by 48 bp repeated sequences, displays a significant similarity to PAPI-1 like integrative conjugative elements (discussed below) (Liu et al., 2018; Qiu et al., 2006). The second 11981 bp insertion between tRNA-Met (*PA4673.1*) and *higA* (*PA4674*), flanked by 82 repeated bp, is identical to the prophage-like RGP42 element identified in MPAO1 and PAO1-DSM (Klockgether et al., 2010). Additionally, PAO11-DSM (Klockgether et al., 2010). Additionally, PAO11-DSM (Klockgether et al., 2010). Additionally, PAO1161 lacks a 280 bp fragment containing PA1796.3 and PA1796.4 tRNA genes and has an 107 bp insertion downstream of PA2327.

Impact of PAO1161 / PAO1-UW sequence variation on gene structure

A comparisons of PAO1161 and PAO1-UW genome sequences using Nucdiff (Khelik et al., 2017) revealed 251 single nucleotide polymorphisms (SNPs), multiple nucleotide polymorphisms (MNPs) and short insertions / deletions (indels). Since a direct sequence comparison does not provide information about quality of predicted variants the outcome can be greatly affected by errors during genome sequence consensus calling, caused for instance by mapping of reads derived from highly similar sequences to another, very similar, parts of the genome. To identify high quality variants we aligned the reads used in genome assembly to the consensus sequence and checked the coverage of the region with reads, their mapping quality as well as frequency of the variant at a given genome position. The variants present in more than 80% of the reads were considered as homozygous. Overall 100 SNPs/MNPs/indels were called with a high confidence between the PAO1161 and the PAO1-UW reference. The remaining (heterozygous) variants were mostly SNPs

(149/151) located in coding sequences of genes belonging to Pseudomonas Orthologous Groups (POG) possessing multiple same strain members (multiple orthologs within PAO1 genome) (Winsor et al., 2016). This group included 17 SNPs in PA0723-PA0726 genes from the Pf1 cryptic prophage region syntenic to the genes within RGP42 element also present in PAO1161 genome (Klockgether et al., 2010). Eight of these variants were also identified previously in MPAO1 and PAO1-DSM (Klockgether et al., 2010), nevertheless these did not meet our quality criteria.

The high-confidence variants encompassed 52 SNPs, 6 MNPs, 15 deletions and 27 insertions. Of these 44 were mapping to intergenic regions in PAO1-UW genome and 9 were synonymous (silent) mutations (Supplementary Table S1). Five identified variants lead to frame shifts, resulting in expression of truncated proteins (Table 1). Three of these mapped to the 3' part of the genes, resulting in short sequence alterations, unlikely to affect the function of corresponding protein products (Table 1). Moreover similar shorter versions of the encoded proteins were found for other representatives of related POG (POG001263, POG003705, POG003333, respectively) indicating that such shortened version of PA1327, PA1685 or PA2141 homologues are more frequent among *Pseudomonas* species. Relative to PAO1, PAO1161 genome contains a frame shift in the PA0683 (*hxcY*) gene encoding a component of the Hxc system, a type II secretion system active under phosphate-limiting growth conditions and dedicated to the secretion of low-molecular-weight alkaline phosphatases LapA and LapB (Cianciotto and White, 2017; Ball et al., 2002, 2012).

Two more mutations resulted in an introduction of premature stop codons (Table 1). These are in PA2735 gene, recently shown to encode a N6-adenosine DNA methyltransferase acting on a conserved sequence (Doberenz et al., 2017), and in PA1939, encoding a putative overcoming lysogenization defect (OLD) family nuclease containing an N-terminal ATPase domain and a C-terminal TOPRIM domain (Aravind et al., 1998). Since *P. aeruginosa* PAO1161 strain was selected as defective in its restriction system (Dunn and Holloway, 1971; Rolfe and Holloway, 1968), it is tempting to speculate that PA1939 could be the protein involved in degradation of foreign DNA.

Together with PA2735 this protein could constitute the chromosomal restriction-modification

system of *P. aeruginosa*.

In contrast, 14/15 of the identified sequence variants could lead to an extension of the predicted protein product (Table 2). These include 413850 T \rightarrow C mutation upstream of PA0369, removing an in frame stop-codon and possibly leading to a translation of a longer protein. Similar, start codon changing role can be attributed to nucleotide differences/changes within PA2046-PA2047 and PA4874-PA4875 intergenic regions, as well as to 5 other indels (Table 2). The effect of three sequence variants was a fusion of proteins encoded by neighbouring genes in PAO1-UW genome into one protein in PAO1161. Additionally, the frameshift observed in PAO1-UW in PA0748 gene, encoding an AraC type transcriptional regulator, is not observed in PA01161. Notably, analysis of the sequences composing Pseudomonas Orthologs Groups (www.pseudomonas.com) of corresponding PAO1 genes showed that the modified (PAO1161-like) protein products constitute a vast majority of all sequences (in comparison to PAO1 reference like sequences), indicating that for these genes PAO1-UW reference may not represent protein products of other strains and Pseudomonas species.

Effect of SNPs, MNPs and indels

Except sequence variants affecting the gene structure, numerous SNPs and indels resulting in amino acid changes relative to corresponding PAO1-UW genes/proteins were identified in PAO1161 genome (Table 3). One special case is PA2492 (*mexT*), where both a deletion and a SNP were observed in PAO1161 relative to PAO1-UW genome (Table 1, 3). MexT is a LysR type transcriptional regulator activating expression of the MexEF-OprN multidrug efflux system, extensively studied in the context of quorum sensing signalling and resistance to antimicrobial agents (Köhler et al., 1999; Uwate et al., 2013; Köhler et al., 2001; Lamarche and Déziel, 2011). PAO1161 does not possesses the 8-nt insertion within *mexT* (Table 1), observed in some PAO1 sublines (Maseda et al., 2000). Concomitantly, PAO1161 contains a SNP in the *mexT* gene, leading to F172I change which is also found in most of the strains lacking the 8-nt insertion in *mexT*

(Maseda et al., 2000). A sequence variation was also observed in *mexS*, encoding a negative regulator of MexT (Uwate et al., 2013; Sobel et al., 2005; Fargier et al., 2012), resulting in D249N amino acid change. Recent data indicate that the MexS protein with this amino acid change appears to be fully functional (Uwate et al., 2013). The presence of functional *mexT/mexS* system in PAO1161 is strengthened by the observation that this strain shows low resistance to chloramphenicol (MIC < 64 µg/ml), whereas the *mexT* defective PAO1 sublines are extremely resistant to this antimicrobial (Luong et al., 2014; Maseda et al., 2000).

PAO1161 strain derives from the strain PAO38 which was selected as the leucine auxotroph (Figure 1A). Genome sequencing of PAO1161 revealed that this strain possess E108K mutation in PA3792 (*leuA*), encoding a putative 2-isopropylmalate synthase. Analysis of Pseudomoans Orthologue Group of the *leuA* (POG001874) showed that, the only other *P. aeruginosa* strains carrying this mutation are PAO579 (Ryan Withers et al., 2013) and PAO581 (Yin et al., 2013), both derived from PAO38. To validate that this mutation is responsible for/causing the leucine auxotrophy, we reversed the mutation. The exchange of the mutant allele by PAO1 gene fully restored the ability of PAO1161 strain to grow on minimal medium without leucine (Figure 2AB), confirming that the E108K mutation in PA3792 (*leuA*) caused leucine auxotrophy.

P. aeruginosa PAO1161 used in our study was a spontaneous rifampicin resistant clone (Lasocki et al., 2007). Rifampicin acts by binding to a conserved pocket on the β -subunit of RNA polymerase therefore blocking RNA transcript elongation (Campbell et al., 2001). Resistance to this drug results from mutations on the *rpoB* gene that change the structure of the pocket (Campbell et al., 2001; Severinov et al., 1993; Hall et al., 2011). Our SNP analysis revealed presence of H531L mutation in PA4270 (*rpoB*) encoding *P. aeruginosa* DNA-directed RNA polymerase subunit beta (Table 3). The *rpoB* H531L mutation was frequently observed in spontaneous *P. aeruginosa* Rifr mutants (Jatsenko et al., 2010), strongly indicating that this SNP confers PAO1161 strain with rifampicin resistance.

From 26 SNPs and indels identified in PAO1161 resulting in amino acid changes relative to corresponding PAO1-UW proteins, 15 seem to be strain (PAO1161) specific (Table 3, bolded gene names). Examples include codons deletion identified in PAO1161 genes encoding a pimeloyl-CoA synthetase PauA (PA1017) involved in biotin synthesis (Binieda A et al., 1999), a transcriptional regulator PA3067 from MarR family or a sensor histidine kinase KinB. The KinB as a part of two-component system is involved in phosphorylation/dephosphorylation of the transcriptional regulator AlgB and control of alginate and virulence factors production in *P. aeruginosa* (Ma S. et al., 1997; Damron et al., 2009; Chand et al., 2012; Balasubramanian D et al. 2013). The point mutation changing amino acid sequence R221H was also detected in PAO1161 *algB* (Table 3).

A start codon shift in *relA* gene encoding a GTP pyrophosphokinase in PAO1161 relative to PAO1-UW was identified (Table 3). It raised the question, if this protein is expressed in PAO1161. Performed sequence analysis showed a possibility of translation starting from a valine codon generated at the beginning of the changed *relA* in PAO1161. Indeed, proteomic analysis of PAO1161 strain using mass spectrometry and iTRAQ quantification demonstrated that the RelA is produced in PAO1161 (Malinowska A. et al. 2012). This protein together with SpoT plays an important role in controlling the level of the alarmone guanosine 5'-triphosphate-3'-diphosphate (ppGpp) involved in bacterial adaptive responses and virulence (Xu X Yu H et al., 2016),

Among identified indels comparing predicted gene products of PAO1161 and PAO1 the codon insertion change in PA5024 locus encoding conserved hypothetical protein with transmembrane domain was identified. Deeper analysis of homologues of this protein from POG004631 showed that the modified shortened version of PA5024 are characteristic only for few close relatives of PAO1 (e.g. PAO579, PAO581, PAO1-VE2), while the vast majority of sequences represents the longer version of the protein similar to that of PA5024 from PAO1161.

Several PAO1161 specific mutations corresponding to a single amino acid changes were also identified (Table 3). Examples include aa substitutions in proteins involved in different metabolic processes e.g. the change T103I in the transcriptional regulator GcdR engaged in

regulation of glutarate utilization and lysine catabolism (Indurthi et al., 2016), A232T in the carbamate kinase ArcC involved in fermentative arginine degradation (Schreiber et al., 2006), V126M in the ribonuclease E (*rne*) involved in mRNA turnover (Kushner, 2004). The G108D change was identified in Tsi4 of PAO1161 relative to corresponding protein of PAO1. This protein is a part of antibacterial effector-immunity pair together with type VI secretion exported toxin Tse4 (PA2774) (Whitney et al., 2014). These could influence the structure and/or function of the corresponding protein and therfore care should be taken in analyses of these genes using PAO1161 strain.

PAO1161 genome contains an ICE conferring resistance to mercury

PAO1161 was described as a strain containing the IncP-8 plasmid FP2, which confers the cells with mercury resistance. Indeed, the strain used in our lab was exceptionally resistant to mercury, showing growth in LB medium with up to 200 µM HgCl₂ (Ciok et al., 2018) (and data not shown), confirming presence of the element. Surprisingly, during genome assembly no extrachromosomal elements could be identified. Instead, an almost 108 kbp insertion in the chromosome, with a putative mercury resistance operon, was found (Figure 1B, Figure 3A). The insertion shows similarities (in sequence and organization of operons flanking the putative integration site) to the PAPI-1 family of integrative conjugative elements (ICE) (Liu et al, 2018; Qiu et al., 2006; Johnson and Grossman, 2015). ICEs are mobile genetic elements, with modular structure, encoding complete conjugation machinery (usually a type IV secretion system) allowing transfer of their genome to another host. They are reversibly integrated into a host genome and can be passively propagated during bacterial chromosome segregation and cell division (Johnson and Grossman, 2015; Delavat et al., 2017; Wozniak and Waldor, 2010). PAPI-1 was first described in the genome of highly virulent *P. aeruginosa* PA14 strain (He et al., 2004). It has a size of 108 kbp, encodes 115 predicted genes, integrates into tRNA_{Lvs} gene and represents an abundant family of ICEs with a similar gene structure, present in *Pseudomonas* genomes (Liu et al., 2018). Integrated PAPI-1-like ICEs contain an operon starting with a gene encoding an orphan F-type ATPase, with high

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similarity to Soj/ParA proteins involved in segregation of chromosomes and plasmids, adjacent to the integration site (Qiu et al., 2006). At the other end of the integrated ICE, an operon encoding a putative relaxase (*tral*) and a tyrosine type site specific recombinase is encoded. The element identified within PAO1161 genome has an integration site within a tRNALys and PAPI-1 like organization of boundary operons (Figure 3A). According to the ICEberg database, this family encompasses PAPI-1(He et al., 2004), PAGI-5 (Battle et al., 2008), PFGI-1 (Mavrodi et al., 2009), pKLC102 (Klockgether et al., 2004), ICE*Pae*2192-1, ICE*Pae*C3719-1 and ICE*Pae*PACS2-1(Liu et al., 2018). Analysis of gene content (Supplementary Table S2), revealed that out of 120 predicted genes within the putative ICE, 102 genes are found in at least of the 5 other PAPI-1 like elements, whereas orthologs of 41 genes are found in all ICE analysed (Supplementary Table S2). These data indicate high similarity of the putative ICE identified in PAO1161 genome with PAPI-1 like elements.

Integration of ICE into the chromosome as well as its excision is mediated by a an ICE encoded site directed recombinase / integrase (Grindley et al., 2006). Recombination between a attachment site in the chromosome (*attB*) and the corresponding site on an circular ICE (*attP*) leads to integration of the element in the genome, now flanked by identical *attL* and *attR* sequences (Figure 3B). Excision of the putative ICE from the PAO1161 chromosome, as well as presence of a stable circular form was analysed using PCR with primers flanking the *att* sequences (Figure 3BC). The analysis confirmed presence of the circular ICE in PAO1161 cells (Figure 3C), indicating that the element can be excised from the chromosome.

To verify that the ICE can be transferred to another host, we inserted a streptomycin resistance cassette within the ICE. Subsequently, PAO1161::ICE-Sm^R strain was used as an ICE donor for mating in static liquid cultures with *Pseudomonas putida* KT2440 as a recipient. Conjugants were selected on M9 plates supplemented with streptomycin, but lacking leucine to block the growth of donor cells. Streptomycin resistant *P. putida* clones were obtained at an efficiency of $2x10^{-7}$ conjugants / donor cell.

To confirm that the *P. putida* conjugants show also enhanced resistance to mercury, attributed to the presence of *mer* operon within the ICE, we analysed the growth of conjugants in medium containing HgCl₂. The recipient *P. putida* KT2440 cells were unable to grow at a HgCl₂ concentration higher than 2 μ M (Figure 3D and data not shown). In contrast, the growth of streptomycin resistant *P. putida* conjugants was not inhibited by addition of 40 μ M HgCl₂ to the medium, confirming acquisition of mercury resistance (Figure 3D).

Finally, to confirm that the ICE, transferred from PAO1161, integrated in *P. putida* KT2440 chromosome we searched for putative *attB* sites in the genome using the 48 bp TGGTGGGTCGTGTAGGATTCGAACCTACGACCAATTGGTTAAAAGCCA sequence flanking the ICE in PAO1161 chromosome. *P. putida* KT2440 genome contains three potential attachment sites called *attB1-3*, with the *attB1* and *attB2* located adjacent to each other (Figure 3E). A PCR analysis, using primer pairs specific to the ICE/chromosome junctions revealed a highly oriented integration of the putative ICE in the genome (Figure 3F). In individual clones, the specific PCR product was observed for one or more of the 3 attachment sites (Figure 3EF, blue arrows), as the PCR analysis only confirmed integration in an orientation resembling the one observed in PAO1161 genome (Figure 3C and data not shown). Overall, these data confirm the ability of the ICE identified in the PAO1161 genome to transfer to another host.

Conclusions

In this work we show that *P. aeruginosa* PAO1161 strain carries a PAPI-1 family integrative conjugative element capable of excision, transfer and integration in the genome of a closely related strain. The ICE contains an operon conferring mercury resistance, in the past attributed to the FP2 plasmid, the only member of IncP-8 incompatibility group. Our data indicate that FP2 is an integrative conjugative element and we propose to rename this element to ICEFP2.

The genomic features of *P. aeruginosa* PAO1161, a close derivative of reference PAO1 strain, were identified based on the complete genome sequence. Our data indicate a number of

sequence variants, likely affecting different aspects of cell physiology. Thus genome sequencing of laboratory strains is highly recommended to provide full information of genotype and help in interpretation of observed phenotypes under the conditions studied.

MATERIALS AND METHODS

Bacterial strains and growth conditions

P. aeruginosa strains used were PAO1161 (*leu*^{*}, r^{*}, m⁺), kindly provided by B. M. Holloway (Monash University, Clayton, Victoria, Australia) and its derivative PAO1161 Rif^R (Lasocki et al., 2007), PAO1161 Rif^R ICERP2^Sm^R (this work). *Escherichia coli* strain DH5 α [F(ϕ 80d*lac*Z\DeltaM15) *recA1 endA1 gyrA96 thi-1 hsdR17*(r_Km_K) *supE44 relA1 deoR* Δ (*lacZYA-argF*)*U196*] was used for all plasmid constructions and S17-1 (*pro hsdR hsdM recA* Tp^R Sm^R Ω RP4-Tc : :Mu-Km : : Tn7) (Simon et al., 1986) was used to mate plasmids into *P. aeruginosa*. Details of plasmid and strain constructions are provided in Supplementary Material.

Bacteria were grown in Luria broth (Kahn et al., 1979) at 37°C or on Luria agar (Luria broth with 1.5% w/v agar) supplemented with antibiotics as appropriate: benzylpenicillin sodium salt at 150 μ g ml⁻¹ in liquid medium and 300 μ g ml⁻¹ on agar plates for penicillin resistance in *E. coli*, streptomycin at 30 μ g ml⁻¹ for streptomycin resistance in *E. coli*, streptomycin at 150 μ g ml⁻¹ for streptomycin resistance in *E. coli*, streptomycin at 150 μ g ml⁻¹ for carbenicillin resistance in *P. aeruginosa* and *P. putida*, carbenicillin at 300 μ g ml⁻¹ for carbenicillin resistance in *P. aeruginosa*, rifampicin at 300 μ g ml⁻¹ for rifampicin resistance in P. *aeruginosa*. Some experiments were performed in M9 minimal medium (Sambrook et al., 1989) supplemented when indicated with leucine (10 μ g ml⁻¹) or glucose (0.1%).

Bacterial growth was monitored by measurements of optical density at 600 nm (OD_{600}) using spectrophotometer UV1000 (Shimadzu) or Varioskan Lux Multimode Microplate Reader with SkanIt RE 5.0 software (Thermo Fisher Scientific).

Plasmids, oligonucleotides and DNA manipulations

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The plasmids used and constructed during this study as well as primers are listed in Supplementary Table S1. Plasmid DNA was isolated by alkaline method and standard genetic procedures were used as recommended (Sambrook et al., 1989). Chromosomal DNA templates for PCRs were prepared from 100 μ l overnight cultures of appropriate strains. Pelleted cells were washed with sterile water, re-suspended in 100 μ l of sterile ddH₂0, boiled and centrifuged 10 min at 4°C at maximum speed. Standard PCR was performed using 1 μ l of boiled cell suspension as a template, 5 pmoles of each primer and DreamTaq polymerase (Thermo Fisher Scientific) in recommended buffer. Fidelity of the amplified DNA sequence was verified by DNA sequencing in the internal sequencing facility (DNA Sequencing and Oligonucleotides Synthesis Laboratory, IBB PAS, Warsaw, Poland).

PCR site-directed mutagenesis

The QuickChange site-directed mutagenesis method was applied (Stratagene) with the pAKE600 plasmid (El-Sayed et al., 2001) derivatives containing *pa3972* gene and ICE fragment, respectively. The presence of change/mutation was verified by DNA sequencing of PCR-amplified fragments.

Bacterial transformation

Competent E. coli cells were prepared by the standard CaCl₂ method (Sambrook et al., 1989).

Introduction of *pa3972* and ICE::*addA* [Sm^{R]} fragment into *P. aeruginosa* PAO1161 chromosome by homologous recombination

The competent *E. coli* S17-1 cells were transformed with pAKE600 suicide vector derivatives to construct donor strains for conjugation. Bacterial conjugation was done on L-agar by mixing 100 μ l of overnight cultures of *E. coli* S-17 (pAKE600 derivatives) donors and *P. aeruginosa* PAO1161 [Rif^R] recipient strain and incubation for 24 h at 37°C. The bacterial mixtures were washed off the plates with 2 ml of L-broth, and diluted cell suspensions were plated on L-agar with rifampicin and carbenicillin for *pa3972* allele exchange and with rifampicin, carbenicillin and streptomycin for ICE::*addA* [Sm^R] selection of transconjugants/integrants.

The integrants of PAO1161 Rif^R (pAKE600 derivatives) were treated as described previously (Lasocki et al., 2007). The allele integration and then allele exchange was verified by PCR using

chromosomal DNA as a template and the adequate pairs of primers and by confirmation of streptomycin resistance for PAPO1161 ICE::*addA* [Sm^R] derivatives.

ICE transfer to *P. putida*

P. aeruginosa PAO1161- ICE[Sm^R] was used as a donor and *P. putida* KT2442 [Rif^R] was used as the recipient strain in ICE transfer/mobility testing. 1 ml of overnight cultures of the donor and recipient strains in liquid LB were centrifuged (1 min, 6500 rpm), the supernatant was discarded and cells suspended in 1 ml of fresh liquid rich medium. The donor and recipient strains were mixed in Eppendorf tubes in different ratio of donor to recipient (1:2; 1:4; 1:8) adding 100 µl of donor strain and appropriate volume of recipient cells. The mixtures were incubated for 2 h at 37°C with shaking and then 3 h without shaking. Then the mixtures were centrifuged at 6500 rpm for 1 min, the supernatant pour out and the cells suspended in the initial volume of 0.9% NaCl. To estimate the number of transconjugants a serial dilution of bacteria in 0.9% NaCl solution was performed and aliquots of serial 10-fold dilutions were plated onto M9 minimal medium agar plates with 0.1% glucose and 150 µg ml⁻¹ streptomycin without leucine. Lack of leucine in plates with M9 medium allows selection of *P. putida* KT2442 transconjugants (leu^+) and elimination of PAO1161 *leu*⁻ donor cells since they need supplementation of leucine on minimal media. In parallel 100 µl of donor or recipient strain cultures, respectively were treated the same way to establish the titer of donor and recipient cells in the conjugation mixture. The transfer frequency was calculated as number of transconjugants per donor cell.

Mercury resistance testing of P. putida ICE transconjugants

Analytical grade salt HgCl2 was used in a resistance assay performed in 96-well plates employing Varioskan Lux Multimode Microplate Reader with SkanIt RE 5.0 software (Thermo Fisher Scientific). Triplicate cultures of each strain/transconjugant were challenged with a range of concentrations of mercury chloride in liquid rich medium (2-400 μ M). The derivatives of *P. putid*a KT2442 that grew in the presence of HgCl2 at concentrations higher that 2 μ M were considered resistant.

Genome sequencing and assembly of sequence reads

To unravel the genomic properties of *P. aerugionsa* PAO1161 strain whole genome sequencing (WGS) was carried out. Bacterial total genomic DNA was isolated using CTAB/lysozyme method (Wilson et al., 1987) with addition of cell wall degrading enzyme: lysozyme (20 mg/ml; Sigma-Aldrich, Dorset, UK). DNA quality was checked on standard 0.8% agarose gel, 1% PFGE gel on Biorad CHEF-III apparatus and template quantity was measured using Qubit v3.0 fluorimeter. Genomic bacterial DNA was mechanically sheared to appropriate size and used for Paired-End TruSeq-like libraries construction using KAPA Library preparation kit (KAPA/Roche, Basel, Switzerland) following manufacturer's instructions. The bacterial genome was sequenced in paired-end mode (v3, 600 cycle chemistry kit) using MiSeq instrument (Illumina, San Diego, CA). Illumina sequencing yielded 23 380 926 reads and 4 676 185 200 nucleotides of sequence data. Obtained sequence reads were filtered by quality using FastX toolkit (http://hannonlab.cshl.edu/fastx_toolkit/) and residual Illumina adapters were removed using Cutadapt (https://github.com/marcelm/cutadapt). Quality filtered and subsampled to 700 million nucleotides Illumina data (7 000 000 paired reads) was assembled using Spades v3.11.1 (http://cab.spbu.ru/software/spades/) to estimate approximate size of draft bacterial genome. In the next stage long reads were generated using MinION nanopore sequencing instrument (Oxford Nanopore Technologies, Oxford, UK). Bacterial DNA was sheared into approximately 20 kb fragments using Covaris gTube (Covaris, Ltd., Brighton, United Kingdom) and the library was prepared using the ONT 1D ligation sequencing kit (SQK-LSK108) with the native barcoding kit (EXP-NBD103). sequencing performed expansion Nanopore was using the NC_48 h_Sequencing_Run_FLO-MIN106_SQK-LSK108 protocol, R9.4.1 MinION flowcell and MinION MkIB instrument. Raw nanopore data was basecalled using Albacore v2.3.1 (Oxford Nanopore Technologies, Oxford, UK). After quality filtering and sequencing adapter removal using Porechop (https://github.com/rrwick/Porechop) and NanoFilt (De Coster et al., 2018) 161877 barcoded reads remained. The median read length of obtained dataset was 12489 nucleotides and 2

160 720 766 of total bases. Long nanopore reads were assembled in a hybrid mode with Illumina data using Unicycler v.0.4.6 (Wick et al. 2017). Genome hybrid assembly resulted in 1 circular replicon of the size 6 383 803 bp. The remaining sequence errors in the genome assembly were verified by the PCR amplification of DNA fragments, followed by Sanger sequencing with an ABI3730x1 Genetic Analyzer (Life Technologies, USA) using BigDye Terminator Mix v. 3.1 chemistry (Life Technologies, USA). All of the sequence errors and missassemblies were further corrected using Seqman software (DNAStar, USA) to obtain complete nucleotide sequence of bacterial genome.

Genome annotation

The assemblies were mapped against *P. aeruginosa* reference genome POA1 to evaluate the core genome average identities and completeness. The finalized draft genome was submitted to NCBI Prokaryotic Genome Annotation Pipeline (PGAP) server for gene predictions and annotations. Prokka Version 1.11 (Prokaryotic annotation) was used to perform the gene prediction (Seemann, 2014). The prediction was relied on the existing annotation resources such coding DNA sequences (CDS) and proteins.

Analysis of single nucleotide polymorphism (SNP)

To identify genetic differences at genome level, the sequencing reads of PAO1161 strain were mapped to its corresponding reference genome *P. aeruginosa* POA1 using Bowtie v0.12.0 software (Langmead et al., 2009). SNPs and indells between the 2 consensus sequences were called using Nucdiff (Khelik et al., 2017). The effect of SNP was predicted using snpEff (Cingolani et al.,

2012).Genome comparison

Protein Basic Local Alignment Search Tool (BLASTp) was used to match the sequence similarities between PAO1161 genome and the *P. aeruginosa* reference genome PAO1 (Altschul et al., 1997).

Nucleotide sequence accession numbers. The nucleotide sequence of the PAO1161

described in this paper has been deposited in the GenBank database (accession number CP032126).

SUPPLEMENTARY MATERIALS

Supplementary Material S1. Details of plasmid and strain constructions.

Supplementary Table S1. SNPs identified in PAO1161 genome. Non-synonymous SNP's are bolded.

Supplementary Table S2. Gene content of ICEFP2 from PAO1161.

Supplementary Table S3. Plasmids, primers used in this study.

ACKNOWLEDGEMENTS

This work was funded by National Science Centre, Poland [2015/2015/18/E/NZ2/00675 granted to A.A.B. and in part by 2013/11/B/NZ2/02555 granted to G.J.B.]. DNA sequencing was performed by DNA Sequencing and Oligonucleotides Synthesis Laboratory, IBB PAS, Warsaw, Poland. Conflict of interest statement. None declared.

FIGURE LEGENDS

Figure 1. (**A**) How PAO1 became PAO1161? History of PAO1 derivatives. (**B**) Pairwise nucleotide sequence comparison of the *P. aeruginosa* PAO1-UW reference strain and genome sequence of PAO1161. An inversion of the large genome segment between *rrnA* and *rrnB* rRNA operons is demonstrated. Positions and schematic gene organization of identified in PAO1161genome mobile/integrative elements: ICEFP2 and RGP42 are showed.

Figure 2. Growth of PAO1161 leu^+ revertant on minimal medium (**A**) on solid agar and (**B**) in liquid cultures. The PAO1161 leu^- is showed for comparison including growth on minimal medium with leucine.

Figure 3. Integrative and conjugative element of PAO1161 – ICEFP2. (**A**) Comparison of gene organization of PAO1161 ICEFP2 with other representative integrative and conjugative elements identified in *P. aeruginosa* strains. The boxes with arrows indicate individual ORFs and their transcriptional orientation. ORF colour corresponds to the predicted protein function in each mobile

element. Gene operons/clusters involved in maintenance and regulation functions, conjugation and excision are marked. The coloured regions show the homology between analysed integrative elements. (**B**) Model of ICEFP2 excision from the chromosome. Thin, orange arrows indicate the primers used for detection of integrated and circular ICEFP2 form. The genes are not drawn to scale. (**C**) PCR products obtained with indicated primer pairs detecting the integrated and circular ICEFP2 as well as empty integration site in PAO1161, respectively. (**D**) Growth of *P. putida* KT2440 in rich medium containing 40 μ M HgCl₂. Growth curves of *P. putida* KT2440 recipient strain and *P. putida* KT2440 carrying ICEFP2 from PAO1161 in L broth and L broth supplemented with HgCl₂ at 37°C, respectively. (**E**) Three potential integration sites *attB* (in blue) in *P. putida* KT2440 genome and ICEFP2 in *P. putida* KT2440 conjugants. Six independently obtained transconjugants was analysed. PCR products obtained with indicated primer pairs detecting the integrated transcord ICEFP2 in *P. putida* KT2440 are presented.

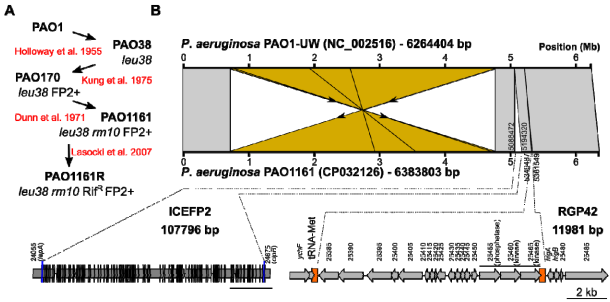
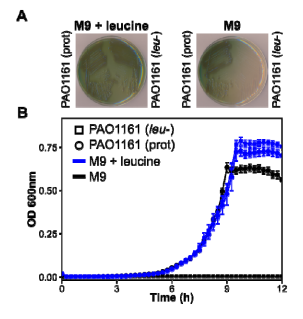


Figure 1





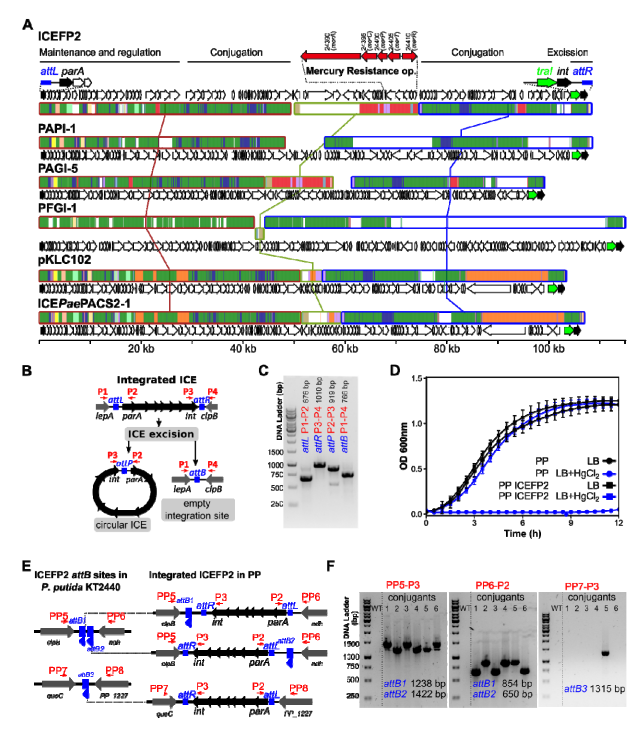


Figure 3

of truncated pr	oterns.						
Mutationeffect	PAO1- UW position	Nucleotide change	AA change	length PAO1/ PAO1161	PAO1 gene	PAO1161 ID	Description
stop gained	2121203	$C \rightarrow T$	W340*	665 /339	PA1939	D3C65_15950	Probable ATP- dependent endonuclease of the OLD family
	3097884	$G \rightarrow A$	Q209*	792 / 208	PA2735	D3C65_11725	SAM-dependent DNA methyltransferase
frame-shift	740419	$G \rightarrow GC$	V73A?	381 / 124	PA0683 (<i>hxcY</i>)	D3C65_22610	probable type II secretion system protein
	1440623	$AA \rightarrow A$	K640	656/642	PA1327	D3C65_19175	probable protease
	1835045	$G \to GC^\dagger$	S218S?	249 / 226	PA1685 (<i>masA</i>)	D3C65_17305	enolase- phosphatase E-1
	2356682	$CC \rightarrow C$	L173	182 /172	PA2141	D3C65_14865	CinA family protein
	2807706	$\begin{array}{c} CAGCCGGCC\\ \rightarrow C \end{array}$	aa1-78/ 35aa	347 / 304	PA2492 (<i>mexT</i>)	D3C65_13040	transcriptional regulator MexT

Table 1 SNPs and indels identified in *P. aeruginosa* PAO1161 genome, possibly resulting in expression of truncated proteins.

 † - SNP at this position in PAO1DSM / MPAO-1 (Klockgether et al., 2010) but a nucleotide insertion in our study

Table 2 SNPs and indels identified in *P. aeruginosa* PAO1161 genome influencing the reading frames of genes and possibly leading to longer protein products. The effect of a mutation is predicted using the PAO1-UW genome as a reference. The possibility of the production of a longer protein was validated by comparison of the predicted sequence with sequences from corresponding Pseudomonas Ortholog Group (POG) at pseudomonas.com.

Position in PAO1-UW	Nucleotide change	PAO1 loci	PAO1161 loci	Effect of mutation	
413850	$T \rightarrow C$	IG: PA0369- PA0370	D3C65_01955	TGA codon -12 to -10 of PA0369 start codon is removed. Longer (30 aa) PA0369 can be translated. Similar extension observed for 297/303 members of POG000360.	
816531	$C \rightarrow CC$	PA0748	D3C65_22250	Corrects the reading frame of PA0748 (still frameshift orf).	
1116213	$G \rightarrow GC$	PA1029	D3C65_20745	Removes start codon of PA1029. Predicted product has additional 24 aa fused to N- terminus of PA1029. Extension observed for 310/316 members of POG000991.	
1215658	$G \rightarrow GG$	PA1122	D3C65_20255	Reading frame is changed. Predicted product has 125 as of PA1122 fused to 54as instead of 22as (PAO1). Similar change observed in 498/504 proteins of POG001082.	
1275766	$GA \rightarrow G$	PA1174	D3C65_19970	First 11 aa of PA1174 (napA) replaced by 16aa. Similar N-terminus is present in most proteins from POG001132.	
2239555	$A \rightarrow AG$	IG: PA2046- PA2047	D3C65_15395	Predicted product has PA2046 sequence extended for 316 aa (N – terminus). The sequence is similar to most of PA2046 orthologs (POG003399).	
2355771	$A \rightarrow AG$	PA2139	D3C65_14875	Reading frame is changed. Predicted product has 29 aa of PA2139 fused with 42aa instead of 11 aa (PAO1). Similar change in 6/12 members of POG008397.	
2753522	$G \rightarrow GC$	PA2451	D3C65_13240	PA2452 fused with PA2451. Similar fusion observed for most proteins from POG003088.	
3016844	$G \rightarrow GC$	PA2668	D3C65_12075	Reading frame and start codon changed leading to a product with last 56 aa identical to C-teminus of PA2668. Extended sequences in 295/301 orthologs from POG008397.	
3083196	$A \rightarrow AG$	PA2727	D3C65_11765	PA2727 fused with PA2728. Similar fusion observed for most proteins from POG002843.	
3919508	$G \rightarrow GC$	PA3503	D3C65_07595	Reading frame and start codon changed. Proposed product has the last 210aa identical with PA3503. Extended sequences in 69/73 orthologs from POG005142.	
4539468	$G \rightarrow GC$	PA4058	D3C65_04690	PA4059 fused with PA4060. Similar fusion observed for all proteins but PA4059/PA4060 from POG001625.	
4888194	$A \rightarrow AG$	PA4360	D3C65_23120	Removes start codon of PA4360. Proposed product has additional 122 aa fused to N- terminus of PA4360. This extension is observed for most members of POG003953.	
5472415	$C \rightarrow CG$	IG: PA4874- PA4875	D3C65_26560	Frameshift in sequence preceding PA4875. Predicted product is PA4875 extended for 81 aa. Longer sequence is typical for most orthologs from POG004503.	

Table 3 SNPs and indels identified in *P. aeruginosa* PAO1161 genome resulting in amino acid changes relative to corresponding PAO1-UW proteins. Strain specific aa changes identified in PAO1161 are marked by bolded gene names.

Effect	PAO1- UW position	Nucleotide change	Amino acid change	PAO1 gene	Description
codon change and codon deletion	1102459	CGAGGCGGTGCTGGGCAGTGCCGAA $\rightarrow C$	VLGSAEEAV513V	PA1017 (pauA)	pimeloyl-CoA synthetase
codon deletion	3436070	$CGTGCATGGCACG \rightarrow C$	RAMH117	PA3067	transcriptional regulator
	6176156	AGCTCGATCGCCTGCA→A	LDRLQ489	PA5484 (kinB)	histidine kinase
codon insertion	5655229	$G \rightarrow GCGG$	W226GW	PA5024	conserved hypothetical protein
start codon shift	1023047	$GGCAAGAT \rightarrow G$	M1MVKGR	PA0934 (relA)	GTP pyrophosphokinase
missense	183697	$T \to G^*$	C310W	PA0159	probable transcriptional regulator
	504428	$C \rightarrow T$	T103I	PA0448 (gcdR)	transcriptional regulator
	1589438	$G \to C^{\#}$	G34A	PA1459	probable methyltransferase
	2669175	$G \to C^{\#}$	P819A	PA2400 (<i>pvdJ</i>)	pyoverdine side chain peptide synthetase
	2806625	$C \rightarrow T$	D249N	PA2491 (<i>mexS</i>)	probable oxidoreductase
	2807982	$T \rightarrow A^*$	F172I	PA2492 (<i>mexT</i>)	transcriptional regulator
	3132936	$C \rightarrow T$	G108D	PA2775 (<i>tsi4</i>)	immunity protein
	3133912	$C \rightarrow T$	S68F	PA2776 (<i>pauB3</i>)	FAD-dependent oxidoreductase
	3333256	$G \rightarrow A$	V126M	PA2976 (rne)	ribonuclease E
	4212201	$A \rightarrow G^*$	H636R	PA3760	N-Acetyl-D- Glucosamine phosphotransferase transporter
	4251149	$G \rightarrow A$	E108K	PA3792 (<i>leuA</i>)	2-isopropylmalate synthase
	4779026	$T \rightarrow A$	H531L	PA4270 (<i>rpoB</i>)	DNA-directed RNA polymerase
	4869855	$T \rightarrow G$	E158D	PA4341	probable transcriptional regulator

4924552	$CG \rightarrow GC^*$	PV177PL	PA4394	mechanosensitive ion channel protein
5336487	$C \rightarrow G$	M405I	PA4751 (ftsH)	cell division protein
5743461	$CG \rightarrow GC^*$	T431S	PA5100 (<i>hutU</i>)	urocanase
5825480	$G \rightarrow A$	A232T	PA5173 (<i>arcC</i>)	carbamate kinase
6115455	$T \to G^{\#}$	K286N	PA5434 (<i>mtr</i>)	tryptophan permease
6131592	$C \rightarrow T$	L169F	PA5443 (<i>uvrD</i>)	DNA helicase II
6174009	$G \rightarrow A$	R221H	PA5483 (algB)	two-component response regulator
6233039	$G \rightarrow A$	S162N	PA5540	hypothetical protein

* - observed in PAO1DSM / MPAO-1 (Klockgether et al., 2010)
* - observed in PAO1DSM / MPAO-1 and validated experimentally (Klockgether et al., 2010)

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