1	Phylogroup-specific variation shapes the clustering of antimicrobial resistance genes and defence
2	systems across regions of genome plasticity
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29 Summary

Background *Pseudomonas aeruginosa* is an opportunistic pathogen consisting of three phylogroups
 (hereafter named A, B, and C) of unevenly distributed size. Here, we assessed phylogroup-specific
 evolutionary dynamics in a collection of *P. aeruginosa* genomes.

Methods In this genomic analysis, using phylogenomic and comparative genomic analyses, we generated hybrid assemblies from a phylogenetically diverse collection of clinical and environmental *P. aeruginosa* isolates, and contextualised this information with 1991 publicly available genomes of the same species. We explored to what extent antimicrobial resistance (AMR) genes, defence systems, and virulence genes vary in their distribution across regions of genome plasticity (RGPs) and "masked" (RGP-free) genomes, and to what extent this variation differs among the phylogroups.

Findings We found that members of phylogroup B possess larger genomes, contribute a comparatively 39 larger number of pangenome families, and show lower abundance of CRISPR-Cas systems. Furthermore, 40 41 AMR and defence systems are pervasive in RGPs and integrative and conjugative/mobilizable elements (ICEs/IMEs) from phylogroups A and B, and the abundance of these cargo genes is often significantly 42 43 correlated. Moreover, inter- and intra-phylogroup interactions occur at the accessory genome level, 44 suggesting frequent recombination events. Finally, we provide here a panel of diverse P. aeruginosa strains to be used as reference for functional analyses. 45 46 Interpretation Altogether, our results highlight distinct pangenome characteristics of the P. aeruginosa

phylogroups, which are possibly influenced by variation in the abundance of CRISPR-Cas systems and
that are shaped by the differential distribution of other defence systems and AMR genes.

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61 Research in context

62 Evidence before this study

To date, pangenome studies exploring the epidemiology and evolution dynamics of bacterial pathogens 63 64 have been limited due to the use of gene frequencies across whole species dataset without accounting for biased sampling or the population structure of the genomes in the dataset. We searched PubMed without 65 66 language restrictions for articles published before September 1, 2021, that investigated the phylogroup-67 specific evolutionary dynamics across bacterial species. In this literature search we used the search terms "pangenome" and "phylogroup" or "uneven", which returned 14 results. Of these, only one study used a 68 population structure-aware approach to explore pangenome dynamics in a bacterial species consisting of 69 70 multiple phylogroups with unevenly distributed members.

71 Added value of this study

72 To our knowledge, this study is the first to assess phylogroup-specific evolutionary dynamics in a 73 collection of genomes belonging to the nosocomial pathogen P. aeruginosa. Using a refined approach that challenges traditional pangenome analyses, we found specific signatures for each of the three 74 75 phylogroups, and we demonstrate that members of phylogroup B contribute a comparatively larger number of pangenome families, have larger genomes, and have a lower prevalence of CRISPR-Cas 76 77 systems. Additionally, we observed that antibiotic resistance and defence systems are pervasive in regions of genome plasticity and integrative and conjugative/mobilizable elements from phylogroups A and B, 78 and that antibiotic resistance and defence systems are often significantly correlated in these mobile 79 80 genetic elements.

81 Implications of all the available evidence

These results indicate that biases inherent to traditional pangenome approaches can obscure the real distribution of important cargo genes in a bacterial species with a complex population structure. Furthermore, our findings pave the way to new pangenome approaches that are currently under-explored in comparative genomics and, crucially, shed a new light on the role that integrative and conjugative/mobilizable elements may play in protecting the host against foreign DNA.

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94 Introduction

Pseudomonas aeruginosa is a ubiquitous metabolically versatile γ -proteobacterium. This Gram-negative 95 bacterium is also an opportunistic human pathogen commonly linked to life-threatening acute and chronic 96 97 infections ¹. It belongs to the ESKAPE pathogens collection ², highlighting its major contribution to nosocomial infections across the globe and its ability to "escape" antimicrobial therapy because of the 98 99 widespread evolution of antimicrobial resistance (AMR)³. This species is also often found to be multi- as well as extensively drug resistant (MDR and XDR, respectively)⁴, making it difficult and in some cases 100 even impossible to treat. For this reason, P. aeruginosa is placed by the World Health Organization 101 (WHO) in the top priority group of most critical human pathogens, for which new treatment options are 102 urgently required 5. These efforts rely on an in-depth understanding of the species biology and its 103 104 evolutionary potential, which may be improved through a functional analysis of whole genome 105 sequencing data.

106 The combined pool of genes belonging to the same bacterial species is commonly referred to as the pangenome. Frequently, only a small proportion of these genes is shared by all species members (the core 107 genome). On the contrary, a substantial proportion of the total pool of genes is heterogeneously 108 distributed across the members (the accessory genome). Following Koonin and Wolf ⁶, the pangenome 109 can be divided into 3 categories: i) the persistent or softcore genome, for gene families present in the 110 111 majority of the genomes; ii) the shell genome, for those present at intermediate frequencies and that are 112 gained and lost rather slowly; iii) the cloud genome, for gene families present at low frequency in all genomes and that are rapidly gained and lost ⁷. Clusters of genes that are part of the accessory genome 113 (i.e, the shell and cloud genome) are often located in so-called regions of genome plasticity (RGPs), 114 115 genomic loci apparently prone to insertion of foreign DNA. By harbouring divergent accessory DNA in different strains, these loci can represent highly variable genomic regions. The shell and cloud genomes 116 117 are also characterized by mobile genetic elements (MGEs) that are capable of being laterally transferred 118 between bacterial cells, including plasmids, integrative and conjugative/mobilizable elements 119 (ICEs/IMEs), and prophages ^{8,9}. These MGEs can mediate the shuffling of cargo genes that may provide a selective advantage to the recipient cell, such as resistance to antibiotics, increased pathogenicity, and 120 defence systems against foreign DNA¹⁰⁻¹². 121

122 Most pangenome studies described to date have characterized gene frequencies across the whole species dataset without accounting for biased sampling or the population structure of the genomes in the dataset. 123 124 This is particularly relevant in species consisting of multiple phylogroups with unevenly distributed members. As recently reported for Escherichia coli¹³, genes classified as part of the accessory genome 125 126 using traditional pangenome approaches are in fact core to specific phylogroups. Since P. aeruginosa is 127 composed of three different-sized phylogroups (hereafter referred to as phylogroups A, B, and C as per the nomenclature proposed by Ozer et al¹⁴; see also results), characterized by high intraspecies functional 128 129 variability ^{15,16}, it is likely that evolution in these phylogroups is driven by specific sets of genes found in 130 the majority of members within the groups, but not across groups.

131 The aim of the current study is to enhance our understanding of the pangenome of the human pathogen P. aeruginosa by specifically assessing phylogroup-specific characteristics and genome dynamics, including 132 data from more than 2000 genomes. We explore to what extent particular groups of cargo genes, such as 133 those encoding AMR, virulence, and defence systems, vary in their distribution across RGPs and 134 135 "masked" (RGP-free) genomes, and to what extent this variation differs among the phylogroups. Our data set includes new full genome sequences of a representative set of P. aeruginosa strains, the 'major P. 136 aeruginosa clone type' (mPact) strain panel. This set of strains was previously isolated by the Tümmler 137 lab (Hanover, Germany) from both clinical and environmental samples ¹⁷. This mPact panel encompasses 138 the most common clone types in the contemporary population ¹⁸⁻²⁰ and provides a manageable, focused 139 resource for in-depth functional analyses. 140

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142 Methods

143 Sequencing and hybrid assembly of the mPact strain panel

Genomic DNA from 18 strains of the mPact panel 17 were extracted using the Macherey-Nagel 144 NucleoSpin Tissue kit, according to the standard bacteria support protocol from the manufacturer. We 145 used Nanodrop 1000 for DNA quantification and quality control (260/280 and 260/230 ratios), followed 146 147 by measurements in Qubit for a more precise quantification. The Agilent TapeStation and the FragmentAnalyzer Genomic DNA 50KB kit served to control fragment size. Sequencing libraries were 148 prepared with the Illumina Nextera DNA flex and Pacific Bioscience (PacBio) SMRTbell express 149 template prep kit 2.0. Libraries were sequenced on the Illumina MiSeq at 2x300bp or the PacBio Sequel 150 II, respectively. Illumina reads were verified for quality using FastQC v0.11.9²¹ and trimmed with Trim 151 152 Galore v0.6.6²², using the paired-end mode with default parameters and a quality Phred score cutoff of 10. Both datasets were then combined using the Unicycler v0.4.8 assembly pipeline ²³. We used the 153 default normal mode in Unicycler to build the assembly graphs of most strains, except of the mPact 154 strains H02, H14, H15, H18, and H19, where we used the bold mode. The assemblies were visually 155 inspected using the assembly graph tool Bandage v0.8.1²⁴. 156

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158 Bacterial collection

We downloaded a total of 5468 P. aeruginosa genomes from RefSeq's NCBI database using PanACoTA 159 160 v1.2.0²⁵. After quality control to remove low-quality assemblies, 2704 were kept and 2764 genomes with more than 100 contigs were discarded (Table S1). Next, 713 genomes were discarded by the distance 161 filtering step, using minimum (1e-4) and maximum (0.05) mash distance cut-offs to remove duplicates 162 and misclassified assemblies at the species level ²⁶, respectively. This resulted in 1991 publicly available 163 genomes. The 18 genomes sequenced in this study from mPact panel 17 passed both filtering steps, 164 resulting in a pruned collection of 2009 genomes in total. Multi-locus sequence typing (MLST) profiles 165 166 were determined with mlst v2.19.0 (https://github.com/tseemann/mlst).

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168 Pangenome and phylogenomics

The average nucleotide identity (ANI) between the 2009 genomes was calculated with fastANI v1.33²⁶. 169 We used the genome sequences to generate a pangenome with the panrgp subcommand of PPanGGOLiN 170 v1.1.136 ^{27,28}. We built a softcore-genome alignment (threshold 95%), followed by inference of a 171 maximum likelihood tree with the General Time Reversible model of nucleotide substitution in IQ-TREE 172 v2.1.2²⁹. To detect recombination events in our collection and account for them in phylogenetic 173 reconstruction, we used ClonalFrameML v1.12³⁰. Phylogenetic trees were plotted in iTOL v6 174 (https://itol.embl.de/)³¹ and explored to cluster genomes according to the phylogroup. Due to the sample 175 size difference, we subsequently focused the analysis on each phylogroup separately. Pangenome analysis 176 177 was performed for each phylogroup, using the panrgp subcommand from PPanGGOLiN. Core and accessory genes were classified across genomes from different phylogroups with a publicly available R 178 script (https://github.com/ghoresh11/twilight)¹³. We used the gene presence/absence output from the 179 whole collection's pangenome and the grouping of our genomes according to the phylogroup. 180

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182 Identification of RGPs and ICEs/IMEs

To mask all the genomes, we used the RGPs coordinates determined by panrgp for each individual genome as input in bedtools maskfasta v2.30.0 ³². We extracted the RGP nucleotide sequences with the help of bedtools getfasta. All genomes were annotated with prokka v1.4.6 ³³. To look for ICEs/IMEs on complete genomes, we used the genbank files created by prokka as input in the standalone-version of ICEfinder ³⁴.

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189 Identification of ICEs and functional categories

We retrieved the annotated proteins for the RGPs and masked genomes across the three phylogroups. We 190 clustered each of the six groups of proteins with MMseqs2 v13.45111³⁵ and an identity cut-off of 80%. 191 These clustered proteins were scanned for functional categories in eggNOG-mapper v2³⁶, using the built-192 in database for clusters of orthologous groups ³⁷. We calculated the relative frequency of these categories 193 by dividing the absolute counts for each category by the total number of clustered proteins found in each 194 of the six groups. CRISPR-Cas systems were identified with the help of CRISPRCasTyper v1.2.3³⁸. 195 AMRFinder v3.10.18 ³⁹ served to locate AMR genes and resistance-associated point mutations. 196 Virulence genes were characterized with the pre-downloaded database from VFDB ⁴⁰ (updated on the 12-197 198 05-2021 and including 3867 virulence factors) in abricate v1.0.1 (https://github.com/tseemann/abricate). 199 Finally, we searched for defence systems using the protein sequences generated by prokka as input in 200 defense-finder v0.0.11⁴¹, a tool developed to identify known defence systems in prokaryotic genomes, for 201 which at least one experimental evidence of the defence function is available.

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203 Network-based analysis of RGPs and ICEs/IMEs

As a first step, we calculated the Jaccard Index between the RGPs with the help of BinDash v0.2.1 42 with 204 205 k-mer size equal to 21 bp. In detail, we used the sketch subcommand to reduce multiple sequences into 206 one sketch, followed by the dist subcommand, to estimate distance (and relevant statistics) between RGPs 207 in query sketch and RGPs in target-sketch. The Jaccard Index between ICEs/IMEs was similarly obtained 208 with BinDash. We used the mean() function in R to calculate the arithmetic mean of the Jaccard Index. 209 Only Jaccard Index values equal to or above the mean were considered, and the mutation distances served 210 as edge attributes to plot the networks with Cytoscape v3.9.1 under the prefuse force directed layout (https://cytoscape.org/). Based on the Analyzer function in Cytoscape, we computed a comprehensive set 211 212 of topological parameters, such as the clustering coefficient, the network density, the centralization, and 213 the heterogeneity. Clusters in our networks were identified with the AutAnnotate and clusterMaker apps 214 available in Cytoscape, using the connected components as the clustering algorithm.

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216 Statistical analysis

217 The correlation matrix was ordered using the helust function in R. Statistical comparison of the variation 218 between groups was always based on non-parametric tests, thereby taking into account that the compared 219 groups varied in data distributions (e.g., at least one group with a skewed distribution) and/or showed 220 unequal variances. Moreover, as non-parametric tests are usually considered to be conservative, the thus 221 identified significant test results should indicate trustworthy differences between groups. In particular, the 222 three phylogroups (e.g., genome size, GC content) were generally compared using the Kruskal-Wallis 223 test. The unpaired two-sample Wilcoxon test was used in multiple comparisons between two independent groups of samples (RGPs vs. masked genomes, CRISPR-Cas positive vs negative genomes). In both tests, 224 225 p-values were adjusted using the Holm-Bonferroni method. Values above 0.05 were considered as nonsignificant (ns). We used the following convention for symbols indicating statistical significance: * for p 226 ≤ 0.05 , ** for p ≤ 0.01 , *** for p ≤ 0.001 , and **** for p ≤ 0.0001 . 227

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239 Results

240 The *P. aeruginosa* phylogeny is composed of three phylogroups

Our phylogenomic characterization was based on 2009 assembled P. aeruginosa genomes belonging to 241 242 519 MLST profiles and including 1991 publicly available genomes (following quality control and 243 distance filtering, Table S2) and additionally 18 genomes for the mPact strain panel (Table S3)¹⁷. Analysis of the ANI values (Figure S1) and the softcore-genome alignment of these genomes identified 244 three phylogroups, as previously reported ^{14,43} (Figure 1). The two major reference isolates are part of the 245 larger phylogroups: PAO1⁴⁴ is part of phylogroup A (n=1531), while the PA14 strain falls into 246 phylogroup B (n=435). Phylogroup C includes a substantially smaller number of members (n=43) (Table 247 **S2**). Members of the phylogroup C were recently subdivided into either 2 14 or 3 clusters, including the 248 distantly related PA7 cluster ⁴³. In this work, however, the PA7 cluster was excluded, and we focused our 249 analysis on only the remainder of phylogroup C, since genomes from the PA7 cluster were too distantly 250 related to the other genomes. In fact, the PA7 strain was first described as a taxonomic outlier of this 251 252 species ⁴⁵, and genomes belonging to this cluster were recently proposed to belong to a new *Pseudomonas* 253 species ⁴⁶. To test the impact of recombination on the softcore-genome alignment, we used 254 ClonalFrameML to reconstruct the phylogenomic tree with corrected branch lengths. The segregation of P. aeruginosa into three phylogroups was maintained, resulting in a tree with decreased branch lengths 255 256 and with identical number of members assigned to each phylogroup (Figure S2). Genomes from the mPact panel sequenced in this study were widely distributed across the *P. aeruginosa* phylogeny, with 12 257 258 strains in phylogroup A, 5 in phylogroup B, and 1 in phylogroup C (Figure 1 and Table S3). Our results show that P. aeruginosa consists of three asymmetrical phylogroups and that the segregation of the 2009 259 260 genomes into phylogenetically distinct groups is not an artefact of recombination.

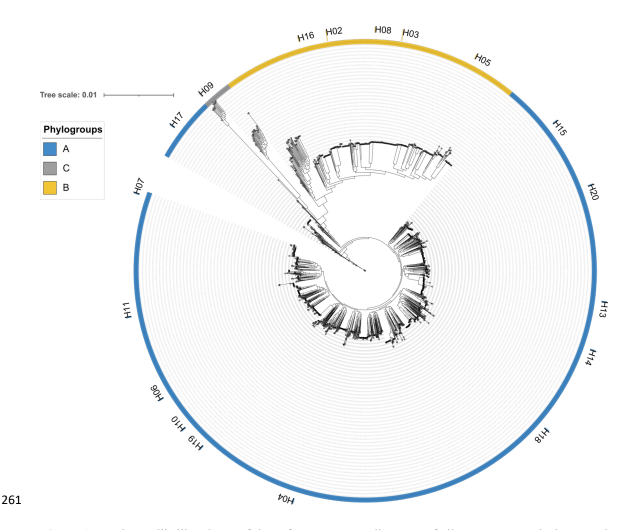


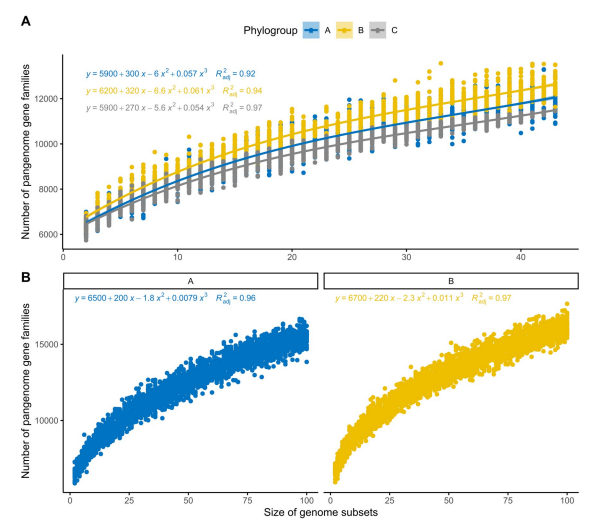
Figure 1. Maximum-likelihood tree of the softcore-genome alignment of all *P. aeruginosa* isolates used in this study (n=2009). The scale bar represents the genetic distance. Arcs in blue represent phylogroup A, yellow B, and grey C. The phylogenetic placement of the major *P. aeruginosa* clone type (mPact) strain panel, sequenced in this study, are highlighted in the tree, with the strain name (the "H" before each number stands for Hanover, referring to the location of the Tuemmler lab and the study that first described this collection ¹⁷) next to strips coloured according to the phylogroup.

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269 Phylogroup B contributes comparatively more gene families to the pangenome than the other two270 phylogroups

We next built a pangenome for the whole species, and separate pangenomes for each of the three phylogroups. This latter approach is important to take phylogenetic subdivisions of the species into account, which is additionally critical because the three phylogroups in our collection have substantially different sample sizes. We observed that the number of persistent gene families in the larger phylogroups A and B were similar to those found in the whole species, while the phylogroup C contained a substantially smaller number of persistent gene families (**Table S4**).

277 The pangenome of bacterial species is usually classified in two types: open pangenomes and closed ones 278 ⁴⁷. Since *P. aeruginosa* is an example of a bacterial species with open pangenome ¹⁴, i.e., the sequencing of new genomes will increase pangenome size, we explored the contribution of each phylogroup to the 279 280 pangenome. To ensure comparability among the three phylogroups in our first analysis, we randomly 281 drew 43 genomes from each phylogroup (thus, including the total sample size of the smallest phylogroup C), and observed that there is more diversity in the accessory genes of phylogroup B as to the functions 282 contributed by the acquired genes (Figure 2A and Table S5). In our second analysis, we focused only on 283 284 the two larger phylogroups A and B, for which we randomly drew in each case 100 genomes and found the trend unchanged (Figure 2B and Table S6). 285



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Figure 2. Rarefaction curves of the pangenome gene families for each phylogroup. All curves were
inferred using polynomial regression lines. Curves in blue represent phylogroup A, yellow B, and grey C.
A) The curves were generated by randomly re-sampling 43 genomes from each phylogroup several times
and then plotting the average number of pangenome families found on each genome. B) Rarefaction
curves were plotted with 100 random genomes from phylogroups A and B.

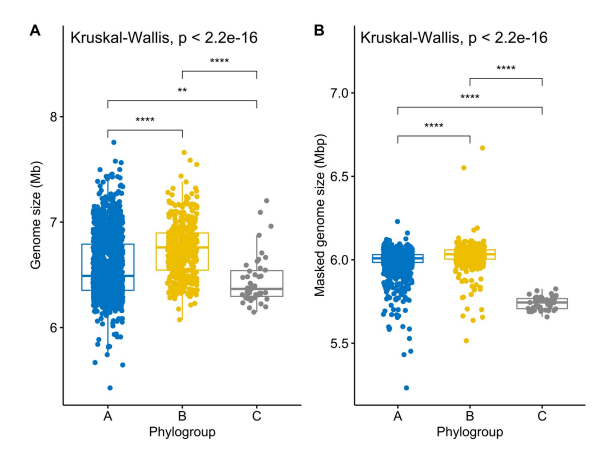
We then explored if specific gene families were pervasive across single or multiple phylogroups. We found 14 phylogroup-specific softcore gene families in phylogroup C, and one gene family each was

294 exclusively found in the softcore genomes of phylogroups A and B, respectively (Figure S3 and Table 295 S7). Most gene families uniquely found on the softcore genome of phylogroup C were part of the Xcp 296 type-II secretion system (T2SS), which is one of two complete and functionally distinct T2SS present in 297 this species (**Table S8**). The Xcp system is encoded in a cluster containing 11 genes (*xcpP–Z*), as well as an additional xcpA/pilD gene found elsewhere in the genome 48. These genes were also found in the 298 majority of the genomes from phylogroups A and B (Table S9), but the encoded proteins were too 299 distantly related to those from phylogroup C. A similar pattern was observed for the two gene families 300 301 indicated exclusively for either phylogroup A or B, for which we also found distantly related orthologues in phylogroup C. Altogether, these results highlight that phylogroup B differs from the other two by 302 contributing a comparatively larger number of gene families to the pangenome, possibly suggesting that 303 304 phylogroup B members have larger genomes.

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306 Phylogroup B genomes are significantly larger and most carry no CRISPR-Cas systems

A comparison of genome lengths revealed significantly larger genome sizes for phylogroup B than the 307 308 other two phylogroups (Figure 3A, p-value < 2.2e-16). We then extracted the RGPs from each 309 phylogroup, and found a total of 57901 RGPs across the three phylogroups. The RGPs from phylogroup B were significantly larger than that of phylogroup A (Figure S4), thus at least contributing to the overall 310 311 size difference. Nevertheless, after removing the RGPs, the resulting "masked" genomes from phylogroup 312 B were still significantly larger than those from the other two phylogroups (Figure 3B, p-value < 2.2e-16). Additionally, we found that genomes from phylogroup B are still significantly larger than those from 313 the other two phylogroups, even if phylogroup sample sizes were adjusted to sample size of the smallest 314 group, phylogroup C (with 43 genomes; Figure S5, p-value 3.2e-07). These results point to a potentially 315 316 higher number of genes conserved across genomes from phylogroup B. Still, the difference in genome 317 size between phylogroups A and B is mainly explained by differences in accessory genome size (Figure 318 S4). Masked genomes from phylogroup C are significantly smaller than genomes from the other two 319 phylogroups, which is consistent with the smaller number of persistent gene families identified in this 320 phylogroup (Table S4). We further explored variation in GC content and observed that the GC content 321 from phylogroup B genomes is significantly lower than those from other phylogroups (Figure S6, p-322 value < 2.2e-16).



Phylogroup 喜 A 喜 B 喜 C

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Figure 3. Boxplots representing the variation in genome size (A) and masked genome size (B) across the three phylogroups. Values above 0.05 were considered as non-significant (ns). Stars indicate significance level: * $p \le 0.05$, ** $p \le 0.01$, *** $p \le 0.001$, and **** $p \le 0.0001$. Boxplots in blue represent phylogroup A, yellow B, and grey C.

328 We next assessed whether presence of the defence CRISPR-Cas system is associated with genome size 329 variation. Since CRISPR-Cas systems are important to defend bacteria against foreign DNA ^{12,49}, we expected that genomes carrying these systems would be smaller, while those devoid of these systems 330 would accumulate mobile elements and hence be larger. We subdivided genomes from each of the three 331 332 phylogroups into two groups depending on whether they contain or lack CRISPR-Cas systems, 333 respectively (CRISPR-Cas^{pos}, CRISPR-Cas^{neg}). We indeed found that genomes with CRISPR-Cas systems are significantly smaller than those without (Figure 4A, p-values 8.3e-05 and 0.00025 for the phylogroup 334 A and B comparisons, respectively), supporting the hypothesis that CRISPR-Cas systems can constrain 335 horizontal gene transfer in P. aeruginosa 50-52. While the number of CRISPR-Cas^{pos} and CRISPR-Cas^{neg} 336 337 genomes in phylogroups A and C is evenly distributed, phylogroup B genomes without CRISPR-Cas 338 (n=279) were nearly two times more prevalent than those that carried these systems (n=156, Table S2). Interestingly, masked genome size of CRISPR-Cas^{pos} and CRISPR-Cas^{neg} phylogroup B isolates was no 339 340 longer significantly different from one another (Figure 4B). In line with this finding, we observed that the

- 341 cumulative size of all RGPs was higher in genomes without CRISPR-Cas systems across phylogroups A
- 342 and B (Figure S7). The absence of these defence systems in most genomes from phylogroup B may help

³⁴³ to explain the observed larger size.

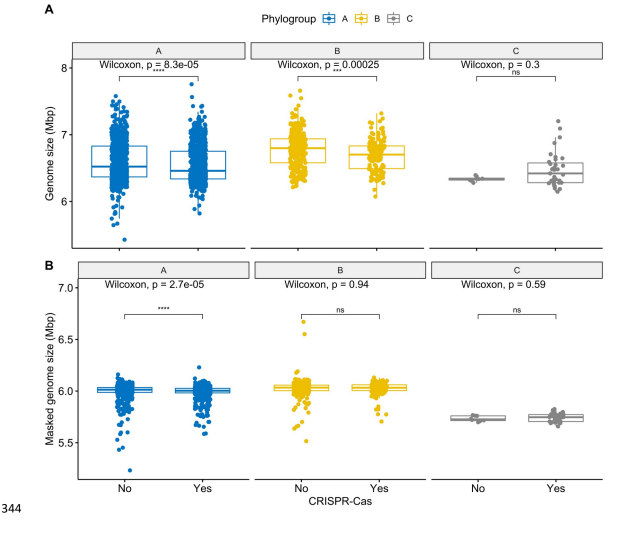


Figure 4. Boxplots representing the variation in genome size (A) and masked genome size (B) across pairs of conspecific genomes from the same phylogroup with and without CRISPR-Cas systems. Values above 0.05 were considered as non-significant (ns). Stars indicate significance level: * $p \le 0.05$, ** $p \le$ 0.01, *** $p \le 0.001$, and **** $p \le 0.0001$. Boxplots in blue represent phylogroup A, yellow B, and grey C.

We observed a wider diversity of CRISPR-Cas systems in genomes from phylogroup A, including I-C, I-E, I-F, IV-A1, and IV-A2 (**Figure S8** and **Table S10**). These CRISPR-Cas subtypes were all found in genomes from phylogroup B, with the exception of the IV-A2. Curiously, only subtypes I-E and I-F were present in phylogroup C. Type IV CRISPR-Cas systems were found almost exclusively on plasmids, and recent work revealed that they participate in plasmid–plasmid warfare ^{12,53}. The type I-C CRISPR–Cas subtype is typically encoded on ICEs and is also involved in competition dynamics between mobile elements ^{51,54}. Overall, our findings show that phylogroup B genomes are significantly larger and have a

357 wider pool of accessory genes than those from the other two phylogroups, possibly driven by the lower

358 prevalence of CRISPR-Cas systems in phylogroup B.

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360 AMR and defence systems are overrepresented in RGPs from phylogroups A and B

We next assessed variation in the relative frequency of proteins encoded in RGPs from different phylogroups. We observed that most functional categories are conserved across phylogroups. However, proteins coding for replication, recombination and repair functions are more prevalent in phylogroups A and B RGPs than those from phylogroup C (**Figure 5A**). Since these proteins are frequently involved in mobilization, this finding may suggest that genomes in these phylogroups have more functional mobile elements, with the ability to be horizontally transferred, while the RGPs in phylogroup C may be derived from remnants of mobile elements that can no longer be mobilized.

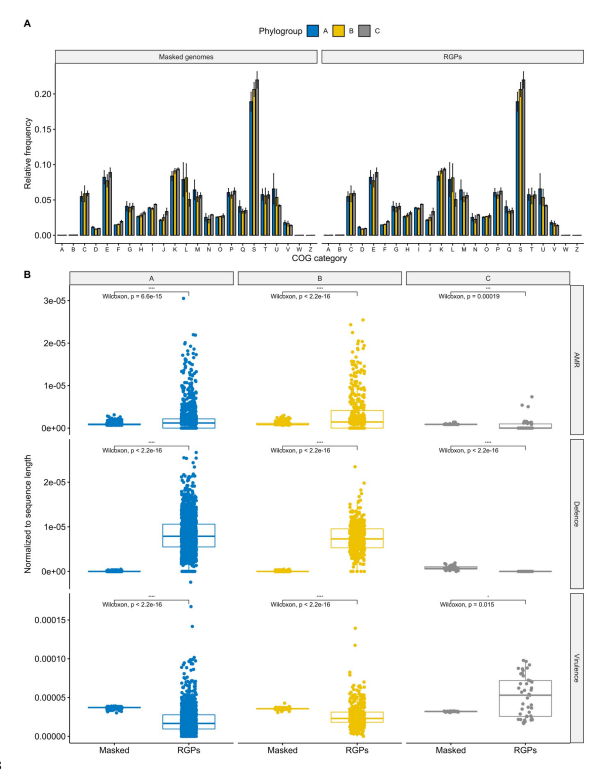


Figure 5. Distribution of functional categories across RGPs and masked genomes from the different phylogroups. Bar and boxplots in blue represent phylogroup A, yellow B, and grey C. **A.** Relative frequencies of cluster of orthologous groups categories. The relative frequencies were calculated by dividing the absolute counts for each category by the total number of clustered proteins found in each of the six groups. Error bars indicate the degree of variation across each COG category from each phylogroup across RGPs and masked genomes. The functional categories are indicated by capital letters, including: A, RNA processing and modification; B, chromatin structure and dynamics; C, energy

376 production and conversion; D, cell cycle control and mitosis; E, amino acid metabolism and transport; F, 377 nucleotide metabolism and transport; G, carbohydrate metabolism and transport; H, coenzyme 378 metabolism; I, lipid metabolism; J, translation; K, transcription; L, replication, recombination and repair; 379 M, cell wall/membrane/envelop biogenesis; N, cell motility; O, post-translational modification, protein 380 turnover, chaperone functions; P, inorganic ion transport and metabolism; Q, secondary structure; R, general functional prediction only; S, function unknown; T, signal transduction; U, intracellular 381 trafficking and secretion; V, defence mechanisms; W, extracellular structures; Z, cytoskeleton. B 382 383 Boxplots of the variation in the number of AMR genes, defence systems, and virulence genes found in RGPs and masked genomes across the three phylogroups. Absolute counts of genes and systems were 384 normalized to RGP and masked genome sequence lengths in each strain. Values above 0.05 were 385 considered as non-significant (ns). Stars indicate significance level: * p <= 0.05, ** p <= 0.01, *** p <= 386 387 0.001, and **** p <= 0.0001.

We next assessed to what extent RGPs and masked genomes vary in prevalence of genes for three types of functions, which are often encoded on MGEs, including virulence, defence systems, and AMR. Since the cumulative size of all RGPs is substantially smaller than that of masked genomes (**Table S2**), the number of virulence genes, defence systems, and AMR genes were normalized to the sequence length of the RGPs and masked genomes for each strain. We observed that the gene prevalence for these functions is conserved across masked genomes from different phylogroups, while they are unevenly distributed in RGPs. (**Figure 5B**).

Two important virulence factors were only present in some genomes from phylogroup C, and absent from 395 the other two phylogroups (Table S9). These genes (exlA and exlB) encode hemolysins, and when 396 397 genomes from phylogroup C carry these genes, the typical type-III secretion system (T3SS) machinery found in most bacteria (encoding the toxins ExoS, ExoY, ExoT, and ExoU) is absent from these genomes, 398 supporting previous reports that these are mutually exclusive 55. In agreement with previous findings 14, 399 we further found that two important genes encoding T3SS effector proteins (exoS and exoU) were 400 401 unevenly distributed across the phylogroups: the exoS gene was pervasive among genomes from 402 phylogroup A (99.5%, 1524/1531) and the majority of phylogroup C strains (28/43), while the exoU gene 403 was overrepresented in genomes from phylogroup B (408/435) and nearly absent in genomes from the other two phylogroups (Table S9). Surprisingly, we also found 23 genomes with the atypical 404 $exoS^{+}/exoU^{+}$ genotype, all belonging to phylogroup A (**Table S9**). A high frequency of this genotype has 405 recently been reported in patients from the Brazilian Amazon and Peruvian hospitals 56,57. As expected 58, 406 407 some virulence genes were exclusively found on RGPs (i.e., absent from masked genomes): flagellarassociated proteins *flel/flag*, *flgL*, *fliC* and *fliD*, as well as wzy, which codes for an O-antigen chain length 408 regulator. All these virulence genes were found in RGPs from both phylogroups A and B. 409

In agreement with the important role of MGEs as vectors for AMR genes in *P. aeruginosa*^{9,59}, we found that AMR genes were overrepresented in RGPs from phylogroups A and B (Figures 5B and S9). We then calculated the relative proportion of different AMR classes across RGPs from the three phylogroups, revealing that most AMR classes were overrepresented across RGPs from phylogroup B (Figure S10).

414 This result is consistent with our finding that RGPs play a significant role in the larger genome sizes from

this phylogroup (Figure S4). Point mutations linked to resistance to beta-lactams and quinolones were
observed for all phylogroups (Table S11).

A wide array of defence systems with a patchy distribution in closely related and distantly related strains 417 418 was recently characterized in P. aeruginosa, suggesting high rates of horizontal gene transfer 41. According to this hypothesis, we would expect to observe an abundance of defence systems in RGPs, 419 420 when compared with masked genomes. Similar to our results for AMR genes, we found that defence 421 systems are indeed overrepresented in RGPs from phylogroups A and B (Figure 5B). Defence systems 422 such as the globally distributed restriction-modification and CRISPR-Cas systems were common in RGPs 423 from both phylogroups. Some rarer systems such as cyclic-oligonucleotide-based anti-phage signalling systems (CBASS) 60, Zorya, Gabija, Druantia 61, abortive infection 62, and bacteriophage exclusion 424 (BREX) ⁶³ were also observed in RGPs from phylogroups A and B (Figure S11 and Table S12). In 425 426 contrast, dGTPases were absent from both phylogroups. Finally, we also observed that AMR and defence 427 systems are overrepresented in specific MLST profiles, including the high-risk clones ST111 and ST233 (Figure S12)¹. Our results revealed that AMR and defence systems are pervasive in RGPs from 428 phylogroups A and B, and the majority of AMR classes are overrepresented in RGPs from phylogroup B. 429

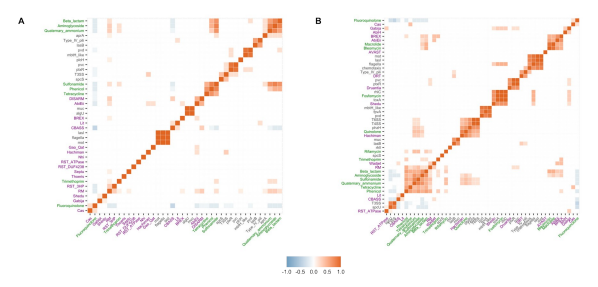
430

431 AMR and defence systems are prevalent in ICEs/IMEs from phylogroups A and B

Given that the distribution and clustering of defence systems in P. aeruginosa is not dependent on the 432 phylogenetic distance between all strains ⁴¹, and considering the high prevalence of ICEs/IMEs in this 433 434 species ⁶⁴, we explored the potential role of these elements as defence islands. To accurately detect these MGEs, we focused our analysis on complete genomes. We noted that 12.6% of our collection consisted 435 of complete genomes (254/2009), including 172 genomes from phylogroup A, 78 from phylogroup B, 436 437 and 4 genomes from phylogroup C (Table S2). 215 out of the 254 complete genomes harboured a total of 477 ICEs and 76 IMEs (Table S13). These ICEs/IMEs were present in 136 genomes from phylogroup A, 438 77 from phylogroup B, and 2 from phylogroup C. Thus, ICEs/IMEs were pervasive in strains from 439 440 phylogroup B (77/78) and in the majority of strains from phylogroup A (136/172).

Nearly half of the ICEs/IMEs carried at least one AMR gene (228/553), with the ciprofloxacin-modifying 441 crpP gene and the sulphonamide-resistance sull gene being most frequent (Table S14). Indeed, the crpP 442 gene was recently shown to be widely dispersed across ICEs from P. aeruginosa 65. Around one third of 443 the ICEs/IMEs (193/553) carried at least one defence system, resulting in a total of 250 defence systems 444 across the ICEs/IMEs and including 27 different types (Figure S13 and Table S14). The most frequent 445 defence subtypes were CBASS-III and restriction-modification type-II 60,62. Virulence genes were present 446 447 in a smaller proportion of the ICEs/IMEs (99/553) and showed higher variation in abundance across ICEs/IMEs than AMR genes and defence systems do (Figure S14). The exoU gene encoding for the 448 effector protein and the spcU gene encoding for its chaperone were the most frequent virulence genes, all 449 450 in ICEs/IMEs from phylogroup B (Table S14).

451 We next explored to what extent the prevalence of these three functional groups is correlated across ICEs/IMEs from the two larger phylogroups A and B. We observed that genes encoding resistance to 452 fluoroquinolones were negatively correlated with genes involved in resistance to other antibiotic classes, 453 and also with specific defence systems as restriction-modification and CBASS (Figure 6A). ICEs/IMEs 454 455 from phylogroup B carrying fluoroquinolone-encoding resistance genes were also negatively associated with genes from the type-III secretion systems (Figure 6B). In contrast, genes encoding resistance to 456 distinct antibiotic classes (e.g., beta-lactams, aminoglycosides, and sulphonamides) were often positively 457 458 correlated in the ICEs/IMEs from both phylogroups, consistent with the previous observations that these genes tend to be co-localized in genetic structures named integrons ⁶⁶. Virulence genes involved in 459 flagellar motility were also often correlated, either additionally with (phylogroup B) or without 460 (phylogroup A) genes involved in chemotaxis ⁶⁷. Defence systems BREX and AbiEii ^{62,63} were positively 461 462 correlated in ICEs/IMEs from phylogroup B. AMR and defence systems showed a high density in 463 ICEs/IMEs from phylogroups A and B, and their frequencies were positively correlated in both 464 phylogroups.



465

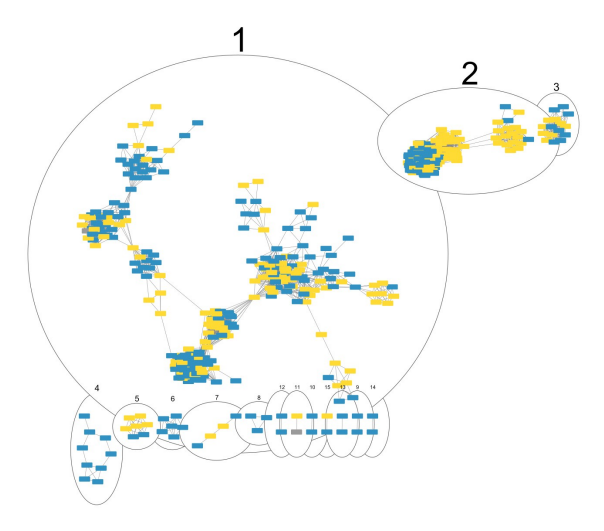
Figure 6. Correlation plots between AMR classes, virulence genes, and defence systems across 466 ICEs/IMEs from phylogroup A (A) and phylogroup B (B). The distribution of cargo genes across 467 ICEs/IMEs was converted into a presence/absence matrix. Correlation matrices were ordered using the 468 469 hierarchical clustering function. Positive correlations are shown in different shades of red, while negative 470 correlations are shown in different shades of blue. AMR genes and point mutations encoding resistance to particular AMR classes are part of the AMRFinder database ³⁹, defence systems of defense-finder ⁴¹, and 471 virulence genes of the VFDB 40. Virulence gene labels are coloured in black, AMR in green, and defence 472 473 systems in purple.

474

475 ICEs/IMEs and RGPs from different phylogroups share high genetic similarity

We next used an alignment-free sequence similarity comparison of the ICEs/IMEs to infer an undirectednetwork. The density plot showed a right-skewed distribution of pairwise distance similarities where the

- 478 vast majority of ICE/IME pairs shared little similarity, with a Jaccard Index value below 0.5 (Figure
- 479 **S15**), in accordance with the high diversity frequently observed across MGEs ⁶⁸. To reduce the density
- 480 and increase the sparsity of the network, we used the mean Jaccard Index between all pairs of RGPs as a
- threshold (0.12184). The network assigned 95.8% (530/553) of the ICEs/IMEs into 15 clusters (Figure
- 482 7). Nearly half of the ICEs/IMEs were grouped in cluster 1 (259/530, Table S15), which includes
- 483 representatives of the three phylogroups.



484

Figure 7. Network of clustered ICEs/IMEs from the three phylogroups, using the mean Jaccard Index between all pairs of ICEs/IMEs as a threshold. Each ICE/IME is represented by a node, connected by edges according to the pairwise distances between all ICE/IME pairs. Numbered ellipses represent ICEs/IMEs that belong to the same cluster. The network has a clustering coefficient of 0.794, a density of 0.099, a centralization of 0.217, and a heterogeneity of 0.785. ICEs/IMEs from phylogroup A are coloured in blue, from phylogroup B in yellow, and from phylogroup C in grey.

We then focused our analysis on the RGPs we extracted from all phylogroups (57901 RGPs in total). We filtered out RGPs smaller than 10kb, and calculated the Jaccard Index between all pairwise of the resulting 32744 RGPs. To reduce the density and increase the sparsity of the network, we used as a threshold the mean value (0.0919429) of the estimated pairwise distances between the 32744 RGPs identified in this study. The network assigned 99.7% (32651/32744) of the RGPs larger than 10kb into 51

496 clusters (Figure S16). While the majority of the RGP clusters were homogeneous for a given phylogroup,

497 we also observed DNA sharing events between different phylogroups. These findings suggest that RGPs

498 and ICEs/IMEs from different *P. aeruginosa* phylogroups share high genetic identity.

499

500 Discussion

501 In this work, we explored the pangenome of the opportunistic human pathogen P. aeruginosa in 502 consideration of its three main phylogroups. This approach allowed us to characterize defining properties of each phylogroup. In particular, we identified genes that are prevalent in the small phylogroup C and 503 504 absent from members of the two larger phylogroups. These genes would have been classified to be part of 505 the accessory genome in conventional analyses of the pangenome of the species as a whole. In contrast, 506 our refined approach suggests that these genes have an evolutionary advantage in a specific genetic context that is particular to this phylogroup ⁶⁹. Moreover, phylogroup C is also clearly distinct from the 507 508 other two phylogroups A and B in having a significantly smaller genome size and a low relative frequency of AMR and defence systems across RGPs. In addition, our results indicate an inverse 509 510 association between size of the phylogroup B accessory genome and presence of CRISPR-Cas systems. 511 This association could (but need not) be causal, such that a low prevalence of CRISPR-Cas defence 512 systems may possibly favour an increase in the size of the accessory genome. Remarkably, genomes 513 devoid of CRISPR-Cas systems in phylogroups A and B were generally significantly larger than those 514 with these systems, a trend that was no longer observed when only considering the non-RGP ("masked") genomes. This observation is consistent with the hypothesis that CRISPR-Cas systems can constrain 515 horizontal gene transfer in *P. aeruginosa* ^{50–52,70}, at least for genomes belonging to the larger phylogroups. 516

517 The three phylogroups vary substantially in the distribution of AMR genes, defence systems, and 518 virulence genes. This variation is particularly apparent in the separate analyses of RGPs and masked 519 genomes. While the length of RGPs is substantially smaller than that of masked genomes, the absolute 520 counts of most defence systems were higher in RGPs than in masked genomes across the three 521 phylogroups (Figure S11). Curiously, representatives of the recently described set of defence systems that are part of Doron's seminal study ⁶¹, such as Zorya, Wadjet, and Hachiman systems, were exclusively 522 found in RGPs across the three phylogroups. In Doron's study, the authors demonstrated that the Wadjet 523 system provided protection against plasmid transformation in Bacillus subtilis, while the Zorya and 524 525 Hachiman systems mediated defence against bacteriophages. These findings highlight the important role of defence systems encoded in RGPs in protecting genomes against infection by foreign DNA and their 526 contribution to MGE-MGE conflict. Moreover, AMR and defence systems are rare in RGPs from 527 528 phylogroup C, which may suggest that these strains are more often subjected to infection by foreign 529 DNA. Assuming that there is no sampling bias across the three phylogroups, then the smaller number of 530 phylogroup C members in public databases could thus be a consequence of the weaker arsenal of AMR and defence systems. Alternatively, phylogroup C strains may indeed be underrepresented, for example if 531 532 they mainly occur in non-clinical habitats, which are usually less well sampled. Collecting P. aeruginosa

samples from distinct geographic regions and environments may further help us reconstruct variation in
 metabolic competences and their connection to origin ⁷¹.

In general, our results underscore the role of ICEs/IMEs as vectors not only of AMR genes ⁵⁹, but also of 535 536 defence systems. Indeed, most of these systems show nonrandom clustering in defence islands and are often co-localized with mobilome genes 61,72-74. Co-occurrence of genes alone, however, does not infer an 537 ecological interaction between them ⁷⁵. Recently, it was proposed that the accessory genome of the genus 538 Pseudomonas is influenced by natural selection, showing a higher level of genetic structure than would be 539 expected if neutral processes governed the pangenome formation ⁷⁶. This suggests that coincident genes in 540 541 ICEs/IMEs are more likely to act together for the benefit of the host or to ensure their own maintenance 542 ^{9,11}. ICEs/IMEs, in particular, provide abundant material for the experimental study of bacterial defence 543 systems. For example, SXT ICEs in Vibrio cholerae, which are also involved in AMR, consistently 544 encode defence systems localized to a single hotspot of genetic shuffling 77. Additionally, ICEs in Acidithiobacillia carry type-IV CRISPR-Cas systems with remarkable evolutionary plasticity, which are 545 546 often involved in MGE-MGE warfare ⁷⁸. Moreover, a recent study proposed that size constraints may account for the low abundance of large defence systems on prophages ⁴¹. In turn, the comparatively larger 547 size of ICEs/IMEs (when compared with prophages) ⁷⁹ may then explain that they commonly harbor large 548 systems such as BREX and defence island system associated with restriction-modification (DISARM)⁸⁰ 549 550 across our dataset (Figure \$13). Even though the CBASS systems are not as prevalent as restriction-551 modification and CRISPR-Cas systems across the bacterial phylogeny ⁴¹, three types of this system were 552 found across ICEs/IMEs from the larger phylogroups.

553 For our analyses, we used complete and draft genome assemblies retrieved from public databases. 554 However, incomplete genome assemblies likely impact RGP definition, due to highly fragmented genomes, that might have inadvertently split RGPs into multiple contigs. With that in mind, we 555 subsampled the complete genomes from our collection and used these to accurately delineate ICEs/IMEs. 556 557 With the sequence similarity comparison between all pairs of ICEs/IMEs found in this study, as well as between all pairs of RGPs, we were able to explore interactions between these elements, suggesting that 558 559 members of the same and of different phylogroups frequently undergo DNA shuffling events. Importantly, this network-based approach using pairwise genetic distances of alignment-free k-mer 560 sequences between MGE pairs has bypassed the exclusion of non-coding elements, providing a more 561 comprehensive picture of MGE populations and dynamics ^{51,81}. Nevertheless, with the current progress in 562 sequence technology, especially including long-read sequencing, we envision a much larger number of 563 564 completely assembled P. aeruginosa genomes in the future, which will then improve reliable assessment of the RGP composition and the role of particular MGEs or gene functions in shaping this species' 565 566 genome characteristics.

To conclude, our work used a refined approach to explore phylogroup-specific and pangenome dynamics in *P. aeruginosa*. Members of phylogroup B contribute a comparatively larger number of pangenome families, have larger genomes, and have a lower prevalence of CRISPR-Cas systems. AMR and defence systems are pervasive in RGPs and ICEs/IMEs from phylogroups A and B, and these two functional groups are often significantly correlated, including both positive and negative correlations. We also

observed multiple interaction events between the accessory genome content both between and within 572 573 phylogroups, suggesting that recombination events are frequent. Our conclusions are contingent on the 574 current range of sequenced genomes for P. aeruginosa. We cannot exclude that some groups, for example phylogroup C and possibly its subgroups, are not fully represented in the currently available data. Future 575 576 sequencing efforts are likely to rectify such a possible problem, thus allowing to test the findings from our study. Finally, our work provides a representative set of phylogenetically diverse *P. aeruginosa* strains, 577 the mPact strain panel, which should prove useful as a reference set for future functional analyses. Such 578 579 functional analyses may help to experimentally assess the underlying reasons for some of the correlations identified in our study, for example the role of specific defence systems in RGP size expansion or in 580 mediating conflict between different MGE types. 581

582

583 Declaration of interests

584 We declare no competing interests.

585

586 Contributors

JB conceptualized and designed the work, acquired and analysed the data, interpreted the data, and wrote the original draft of the manuscript. LT conceptualized and designed the work, acquired the data, and interpreted the data. JF, FB, CU, JK, SN, and BT acquired and analysed the data. HS conceptualized and designed the work, interpreted the data, and contributed to writing of the original draft of the manuscript. HS, SN, and BT acquired funding for this work. All authors read, revised, and approved the final manuscript.

593 Data sharing

performed 594 Scripts for reproducing the analyses in this work available are at 595 https://gitlab.gwdg.de/botelho/pa_pangenome. The representative set of P. aeruginosa genomes and the input file used for the network analysis in Figure 7 are available at the Figshare project 596 https://figshare.com/projects/P aeruginosa pangenome/155021 . 597 Analyses were made with a combination of shell and R 4.0.3 scripting. Sequencing performed in this project were deposited in NCBI 598 599 under the Bioproject accession number PRJNA810040.

600

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- 612

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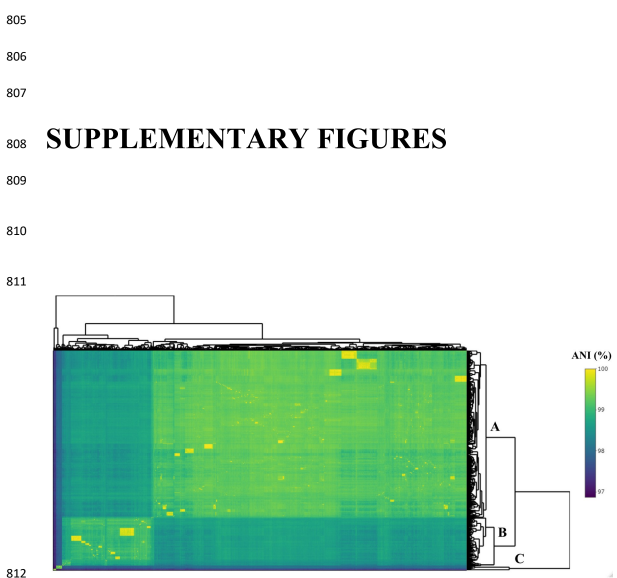
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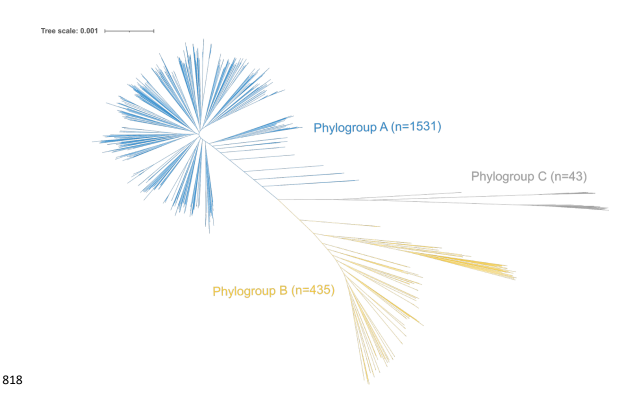
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813 Figure S1. Matrix of Average Nucleotide Identity (ANI) scores between the 2009 P. aeruginosa genomes 814 used in this study. Row and column dendrograms are displayed. Hierarchical clustering was performed 815 with the complete-linkage clustering method. The three phylogroups identified in this study are highlighted in the row dendrogram. 816



819 Figure S2. Maximum-likelihood tree of the softcore-genome alignment of all *P. aeruginosa* isolates used

- 820 in this study (n=2009), corrected for recombination. The scale bar represents the genetic distance.
- 821 Members of phylogroup A are coloured in blue, B in yellow, and C in grey.

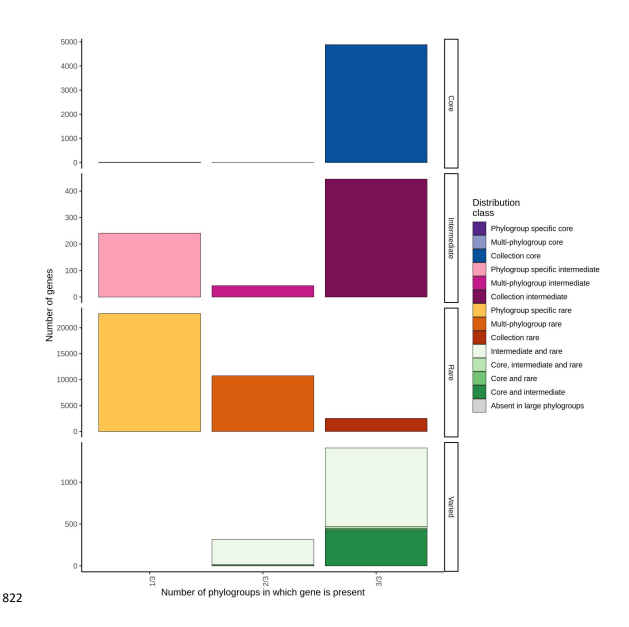
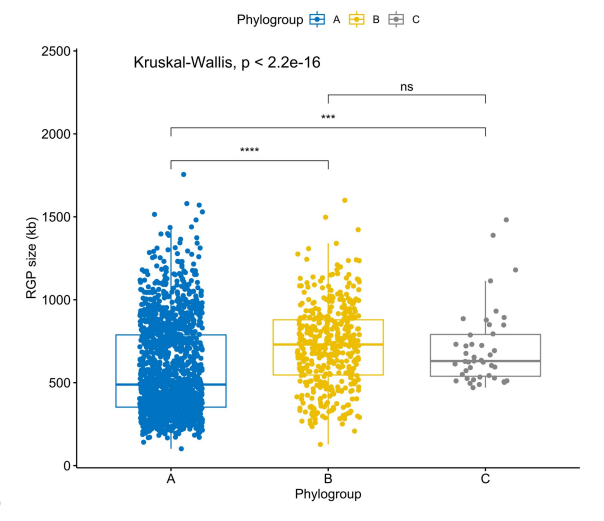
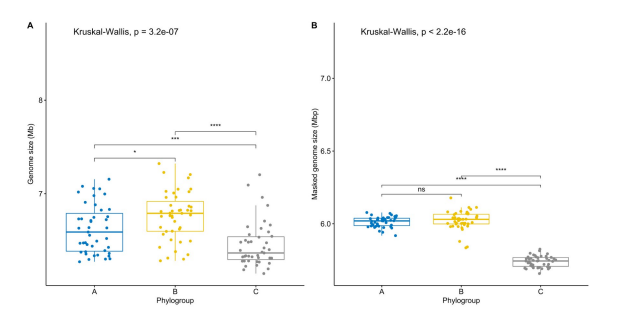


Figure S3. Barplots of the distribution of gene families into core, intermediate, rare, or varied parts of the pangenome across phylogroups. The first column shows genes that are specific to a given phylogroup, and further classified into core (\geq 95%), intermediate, rare (\leq 15%), or varied. The second column shows genes that are specific to two phylogroups, and their classification into core, intermediate, rare, or varied. The third column shows genes that are present across all three phylogroups, and their classification into core, intermediate, rare, or varied. A different colour is assigned to each classification. To create the plot, we modified the R script available in https://github.com/ghoresh11/twilight/blob/master/classify_genes.R.



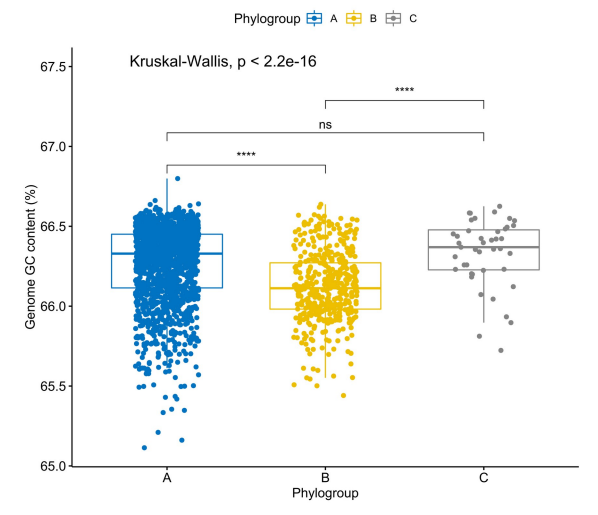
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Figure S4. Boxplots showing the variation in RGP size across the three phylogroups. Values above 0.05
were considered as non-significant (ns). Stars indicate significance level: * p <= 0.05, ** p <= 0.01, *** p
<= 0.001, and **** p <= 0.0001. Boxplots in blue represent phylogroup A, yellow B, and grey C.



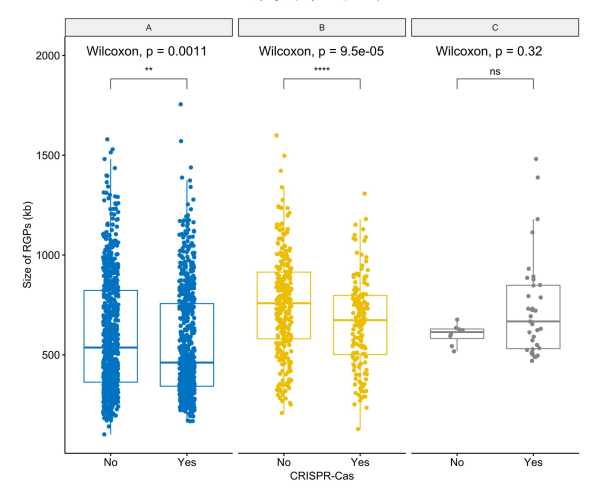
Phylogroup 軴 A 喜 B ڣ C

Figure S5. Boxplots representing the variation in genome size (A) and masked genome size (B) across the three phylogroups. Phylogroup sample sizes were adjusted to sample size of the smallest group, phylogroup C (with 43 genomes). Values above 0.05 were considered as non-significant (ns). Stars indicate significance level: * $p \le 0.05$, ** $p \le 0.01$, *** $p \le 0.001$, and **** $p \le 0.0001$. Boxplots in blue represent phylogroup A, yellow B, and grey C.



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Figure S6. Boxplots showing the variation in GC content across the three phylogroups. Values above
0.05 were considered as non-significant (ns). Stars indicate significance level: * p <= 0.05, ** p <= 0.01,
*** p <= 0.001, and **** p <= 0.0001. Boxplots in blue represent phylogroup A, yellow B, and grey C.



Phylogroup 喜 A 喜 B 喜 C

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Figure S7. Boxplots representing the variation in the size of RGPs across pairs of conspecific genomes from the same phylogroup with and without CRISPR-Cas systems. Values above 0.05 were considered as non-significant (ns). Stars indicate significance level: * $p \le 0.05$, ** $p \le 0.01$, *** $p \le 0.001$, and **** $p \le 0.0001$. Boxplots in blue represent phylogroup A, yellow B, and grey C.

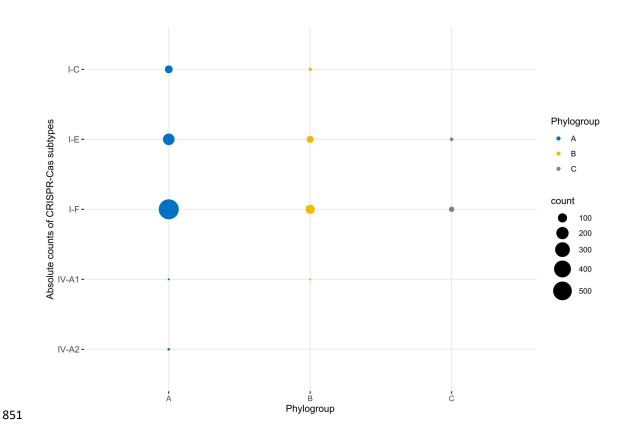
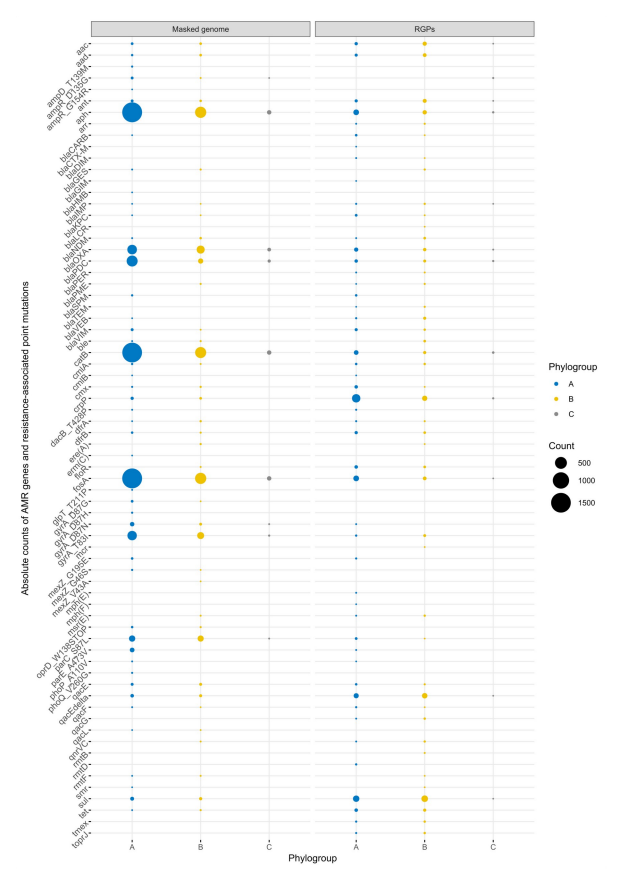


Figure S8. Absolute counts of CRISPR-Cas subtypes identified across genomes from the three
phylogroups. Circles in blue represent phylogroup A, yellow B, and grey C. Circle size is proportional to
the number of absolute counts.



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Figure S9. Absolute counts of AMR genes and resistance-associated point mutations across masked
genomes and RGPs from the three phylogroups. Genes and mutations are part of the AMRFinder
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858 database (45). Circle size is proportional to the number of absolute counts. Circles in blue represent

859 phylogroup A, yellow B, and grey C.



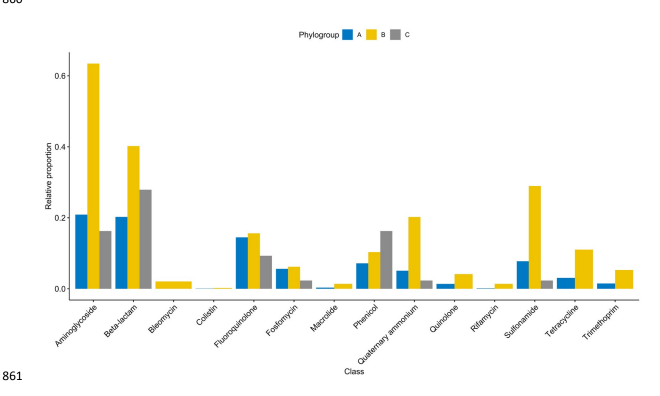


Figure S10. Barplots showing the relative proportion of genes encoding resistance to antibiotics from
different classes across RGPs from the three phylogroups. Genes were normalized to the total number of
genomes found in each phylogroup. Bars in blue represent phylogroup A, yellow B, and grey C.

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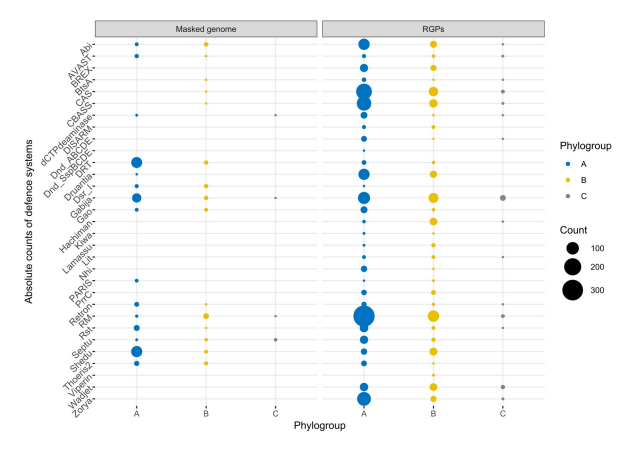


Figure S11. Absolute counts of defence systems across masked genomes and RGPs from the three
phylogroups. Defence systems are part of the defense-finder database (36). Circle size is proportional to
the number of absolute counts. Circles in blue represent phylogroup A, yellow B, and grey C.

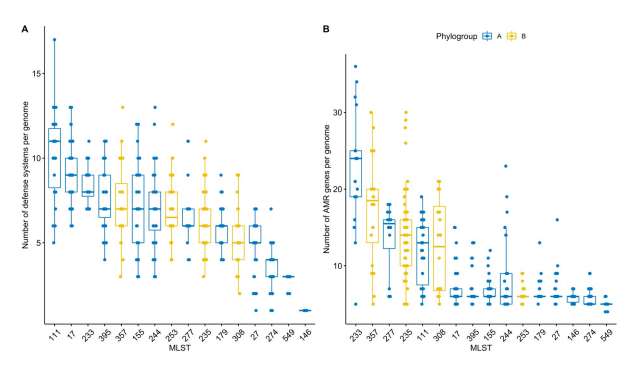
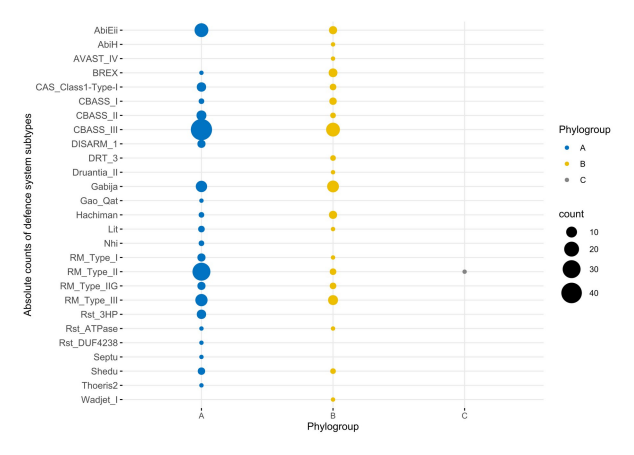




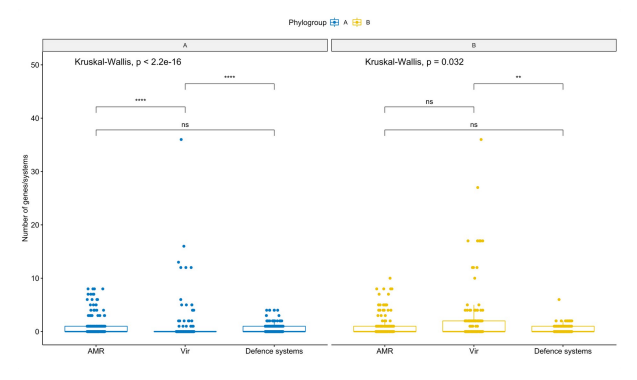
Figure S12. Boxplots representing the variation in the number of defense systems (A) and AMR genes
(B) found across the genomes from the main MLST profiles found in this study. Only MLST profiles
with at least 10 genomes are shown. Boxplots are ordered in descending order by the median values.



882 Figure S13. Absolute counts of defence systems across ICEs/IMEs from the three phylogroups. Defence

systems are part of the defense-finder database (36). Circle size is proportional to the number of absolute

884 counts. Circles in blue represent phylogroup A, yellow B, and grey C.



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Figure S14. Boxplots representing the variation in the number of AMR genes, defence systems, and virulence genes found in ICEs/IMEs across the two larger phylogroups A and B. Values above 0.05 were considered as non-significant (ns). Stars indicate significance level: * $p \le 0.05$, ** $p \le 0.01$, *** $p \le$ 0.001, and **** $p \le 0.0001$. Boxplots in blue represent phylogroup A, yellow B, and grey C.

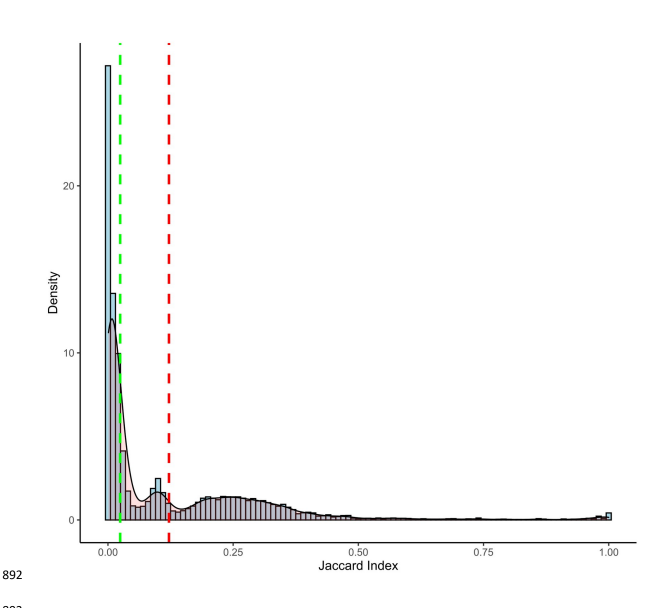
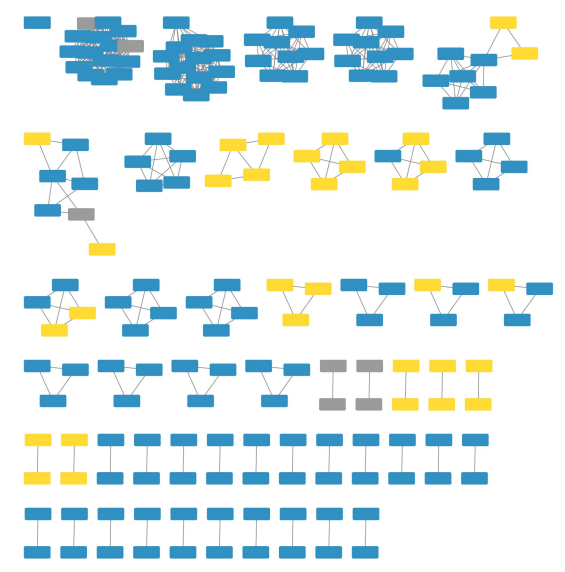


Figure S15. Histogram showing the right-skewed distribution of the Jaccard Index between all pairs of
ICEs/IMEs. Median and mean values are highlighted by vertical dashed lines in green and red,
respectively.



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900 Figure S16. Network of clustered RGPs from the three phylogroups, using the mean Jaccard Index 901 between all pairs of RGPs as a threshold. Each RGP is represented by a node, connected by green edges 902 according to the pairwise distances between all RGPs pairs. Numbered ellipses represent RGPs that 903 belong to the same cluster. The network has a clustering coefficient of 0.777, a density of 0.007, a 904 centralization of 0.026, and a heterogeneity of 0.755. RGPs from phylogroup A are coloured in blue, from 905 phylogroup B in yellow, and from phylogroup C in grey.

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