

1 **Identifying and exploiting genes that potentiate the evolution of antibiotic resistance**

2 Danna Gifford^{1*§}, Victoria Furió^{1,§}, Andrei Papkou¹, Tom Vogwill¹, Antonio Oliver² and R.
3 Craig MacLean^{1*}

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5 **Affiliations:**

6 ¹University of Oxford, Department of Zoology
7 New Radcliffe House
8 Woodstock Road
9 Oxford OX2 6GG

10 ² Servicio de Microbiología and Unidad de Investigación, Hospital Universitario Son Espases
11 Instituto de Investigación Sanitaria de Palma (IdISPa)
12 Palma de Mallorca, Spain

13 [§]Equal author contributions

14 * Corresponding authors:

15 Email: danna.gifford@manchester.ac.uk

16 Phone: +44(0)7534715187

17 Present address: The University of Manchester

18 School of Biological Sciences

19 Faculty of Biology, Medicine and Health

20 Michael Smith Building

21 Oxford Road

22 Manchester M13 9PT, UK

23 Email: craig.macleon@zoo.ox.ac.uk

24 Phone: +44(0)7703327882

25 Address: University Of Oxford

26 Department Of Zoology

27 New Radcliffe House, Woodstock Road

28 Oxford OX2 6GG

29 **One Sentence Summary:** Here we identify potentiator genes and pathways that make bacteria
30 prone to evolving antibiotic resistance, and we exploit this to design treatment strategies for
31 preventing resistance evolution.

32 **Introductory paragraph:** There is an urgent need to develop novel approaches for predicting
33 and preventing the evolution of antibiotic resistance. Here we show that the ability to evolve *de*
34 *novo* resistance to a clinically important β -lactam antibiotic, ceftazidime, varies drastically across
35 the genus *Pseudomonas*. This variation arises because strains possessing the *ampR* global
36 transcriptional regulator evolve resistance at a high rate. This does not arise because of mutations
37 in *ampR*. Instead, this regulator potentiates evolution by allowing mutations in conserved
38 peptidoglycan biosynthesis genes to induce high levels of β -lactamase expression. Crucially,
39 blocking this evolutionary pathway by co-administering ceftazidime with the β -lactamase
40 inhibitor avibactam can be used to eliminate pathogenic *P. aeruginosa* populations before they
41 can evolve resistance. In summary, our study shows that identifying potentiator genes that act as

42 evolutionary catalysts can be used to both predict and prevent the evolution of antibiotic
43 resistance.

44

45 **Main text:** Antibiotic resistance in pathogenic bacteria poses a growing threat to human health,
46 by increasing the mortality rate and economic burden associated with bacterial infections¹. In
47 light of this threat, there is an urgent need to develop new tools for predicting when resistance is
48 likely to evolve in pathogen populations². Research in this area has largely focused on
49 understanding how differing antibiotic treatment strategies, such as mixtures and cycles,
50 influence the evolutionary dynamics of resistance³⁻⁵. An alternative approach is to ask if there are
51 specific genes that make bacteria more likely to evolve resistance to antibiotics⁶. Whole genome
52 sequencing has highlighted the incredible genetic diversity of pathogenic bacteria⁷, but the
53 impact of this diversity on the evolution of antibiotic resistance remains poorly understood. For
54 example, recent work in *Streptococcus pneumoniae* has shown that genes that are important for
55 resistance in one strain may be completely dispensable in another⁸. Although many genes are
56 associated with clinical resistance, it is unclear to what extent other genes in the genome
57 influence the evolution of resistance. For example, recent work has shown that some genes
58 ‘potentiate’ the evolution of novel bacterial phenotypes by opening otherwise inaccessible routes
59 to adaptation^{9,10}. The existence of potentiator genes suggests that genomic background may play
60 a key role in the evolution of antibiotic resistance.

61 *In vitro* selection experiments have emerged as an important tool for studying the
62 evolution of antibiotic resistance^{1,3,5}. However, these studies typically use selection lines derived
63 from a single ancestral clone, making it difficult to understand the role that genetic background
64 itself plays in the evolution of resistance. One approach to circumvent this difficulty is to use
65 comparative experimental evolution, where a diverse collection of strains are challenged with
66 adapting to a common selective pressure⁶. Using this approach, we recently demonstrated that
67 genetic background influences the evolution of resistance to rifampicin by altering the spectrum
68 and fitness effects of mutations in a highly conserved domain of RNA polymerase that confer
69 resistance to rifampicin^{6,11}. In this paper, we extend this approach to uncover resistance
70 potentiator genes by challenging 8 strains that span the genus *Pseudomonas* with the β -lactam
71 antibiotic ceftazidime.

72 *Pseudomonas* is a diverse genus of bacteria that includes *P. aeruginosa*, an important
73 opportunistic pathogen of humans that is the primary cause of mortality in patients who suffer
74 from cystic fibrosis. Crucially, it is possible to culture a wide range of *Pseudomonas* strains
75 under a common set of lab conditions, making it possible to study evolutionary responses to
76 antibiotics in these bacteria using tightly controlled and replicated experiments. We chose to
77 study the evolution of resistance to ceftazidime for two reasons. First, ceftazidime is a clinically
78 relevant antibiotic that is commonly used to treat *Pseudomonas* infections¹² and ceftazidime
79 resistance is common in clinical isolates of *P. aeruginosa*. Second, the mechanisms of
80 ceftazidime action and resistance are well characterized. Ceftazidime inhibits cell wall
81 biosynthesis by irreversibly binding to periplasmic penicillin-binding proteins, ultimately leading
82 to cell death. In spite of this simple mechanism of action, *Pseudomonas* can use at least 4 routes
83 to evolve resistance to ceftazidime: altering the structure of penicillin-binding proteins,
84 upregulating the expression of efflux pumps, reducing permeability of the outer membrane and
85 upregulating the expression of β -lactamase enzymes that break down the antibiotic¹³⁻¹⁵ (Figure

86 S1). Mutations altering the structure of the β -lactamase enzyme itself do occur, but provide much
87 lower increases in resistance¹⁶.

88 Here we use a serial passage experiment to challenge close to 1,000 populations of
89 *Pseudomonas* with doses of ceftazidime that increased from sub-lethal to lethal concentrations
90 over the course of 1 week. We then use extensive whole genome re-sequencing of evolved
91 clones to identify genes and pathways that contribute to the rapid evolution of elevated
92 ceftazidime resistance. Using selection experiments and competition assays with defined
93 mutants, we directly test the evolutionary impact of key pathways to resistance identified from
94 whole genome sequencing. Finally, we demonstrate that understanding the genetic drivers of
95 resistance evolution can be used to design a simple drug mixture, consisting of ceftazidime
96 coupled to a β -lactamase inhibitor, to prevent the evolution of resistance *in vitro*.

97

98 Results and discussion

99 **Strain-specific variation in resistance evolution** To test the role of genetic background in the
100 evolution of antibiotic resistance, we challenged 120 populations of each of 8 strains that span
101 the diversity of the genus *Pseudomonas* with ceftazidime (Figure 1a). This breadth of
102 phylogenetic coverage allowed us to explore the impact of genome content on resistance
103 evolution, and strains were chosen on the basis of variation in genome size, experimental
104 tractability, and the availability of high-quality published reference genomes. Populations were
105 serially passaged in standard lab culture medium supplemented with ceftazidime, the
106 concentration of which was doubled daily from sub-lethal (1/8 minimal inhibitory concentration,
107 or ‘MIC’) to super-lethal (8 \times MIC) levels over a 7 day selection experiment. The MIC of the
108 parental strains varies (0.65-8 mg/L) and we controlled for this variation by standardizing
109 antibiotic doses of selection lines to their appropriate parental strains. In this experimental
110 design, populations can only avoid extinction if they evolve elevated antibiotic resistance, and
111 we measured population survival at each day of the experiment. We define the rate of population
112 extinction within strains as a measure of adaptive potential for resistance evolution, or
113 ‘evolvability’. The rate of population extinction varied profoundly between strains (Figure 1b;
114 Cox’s proportional hazard, likelihood ratio=1930, df=7, $P<10^{-6}$). For example, all of the replicate
115 populations went extinct in some strains, such as *P. mendocina* CCUG1781 and *P. fulva*
116 CCUG12537, while at the other extreme, every population of *P. protegens* Pf-5 survived at up to
117 8 \times the MIC of the parental strain. Given that resistance evolved by selection on spontaneous
118 mutations, one potential explanation for this result is that the ability to evolve ceftazidime
119 resistance correlates to the mutation rate. However, evolvability does not correlate with mutation
120 rate ($r=0.33$, $F_{1,6}=0.74$, $P=0.42$, see supplementary table S1 for calculations) or mutation supply
121 rate, which is the product of initial population size and mutation rate ($r=0.22$, $F_{1,5}=0.27$, $P=0.62$).
122 Additionally, there was no correlation between survival and the absolute difference between the
123 temperature of the selection experiment (30° C) and published optimal growth temperatures for
124 each strain ($r=0.06$, $F_{1,6}=0.027$, $P=0.88$).

125 **Genomics of resistance evolution** To determine the genetic basis of resistance evolution, we
126 sequenced the genomes of 100 independently evolved clones from populations that survived
127 selection for elevated resistance ($n=14-24$ clones/strain). We identified a total of 196 novel
128 mutations in 69 unique genes (i.e. orthologs across strains are each counted once). Mutations

129 included SNPs ($n=80$), short indels ($n=71$), insertion element insertions ($n=15$), larger insertions
130 and deletions ($n=7$), and intergenic mutations ($n=23$). Several lines of evidence indicate that the
131 mutations that we identified were predominantly beneficial. First, parallel evolution occurred
132 both within and across strains. We identified a total of 25 genes mutated in two or more
133 independent clones, and 76% of mutations occurred in these 25 genes. Second, all 80 SNPs
134 observed in coding regions were non-synonymous, which is a clear hallmark of positive
135 selection. A full list of the mutations we identified is given in supplementary data table S2. We
136 categorized mutations according to known resistance mechanisms: (i) porin genes, (ii) penicillin
137 binding proteins (PBPs), (iii) peptidoglycan biosynthesis genes and (iv) multidrug efflux
138 pumps¹³⁻¹⁵. Almost all of the evolved clones (88/100) carry mutations in previously established
139 ceftazidime resistance pathways. However, the distribution of mutations across these resistance
140 pathways differs profoundly between strains, demonstrating strain-specific mechanisms of
141 resistance evolution (Figure 2; $\chi^2=139$, $df=12$, $P<10^{-6}$). *P. protegens* Pf-5 and *P. fluorescens* Pf0-
142 1 adapt by mutations in genes involved in peptidoglycan biosynthesis and recycling (*ampD* and
143 *mpl*), knockouts of which are known to increase the expression of the chromosomal *ampC* β -
144 lactamase gene^{14,17}. In addition to mutations in *ampD* and *mpl*, 21 of 24 clones of *P. aeruginosa*
145 PAO1 carry mutations in a non-essential PBP (*dacB*/PBP4) that has also been shown to increase
146 *ampC* expression when knocked out¹⁸. Consistent with this genetic data, clones from these
147 strains have increased resistance to a broad spectrum of β -lactams, but retain sensitivity to
148 imipenem, which is a poor substrate for the AmpC β -lactamase. In contrast, *P. stutzeri*
149 ATCC17588 and *P. putida* KT2440 evolve resistance by mutations in efflux pump genes and, to
150 a lesser extent, porins. Mutations in efflux pumps are associated with small increases in
151 ceftazidime resistance and a multi-drug resistant phenotype, while porin mutations are
152 predominantly associated with elevated β -lactam resistance (Figure 2). A substantial fraction
153 (33.3%) of clones from these strains lack mutations in known resistance genes; however, these
154 clones have resistance profiles that are similar to those of clones carrying mutations in known
155 efflux pumps or porins.

156 ***The AmpR transcription factor increases evolvability*** The key insight from whole genome
157 sequencing, and phenotypic analysis of evolved clones, is that large increases in ceftazidime
158 resistance are associated with mutations in the peptidoglycan biosynthesis pathway associated
159 with increased β -lactamase production¹⁴. Importantly, the relevant peptidoglycan biosynthesis
160 genes (*ampD*, *mpl* and *dacB*) are present in all of the strains, and the *ampC* β -lactamase gene is
161 present in all of the strains except *P. stutzeri* ATCC17588 (which possesses another β -lactamase
162 gene, *blaZ*).

163 These observations raise an interesting puzzle: if the key genes involved in adaptation are largely
164 maintained, then why does evolvability vary across strains? An alternative approach to
165 understanding why evolvability varies across strains is to take a functional approach to
166 characterizing the effects of beneficial mutations. Inactivation of the peptidoglycan biosynthesis
167 genes involved in adaptation in our experiment has been shown to increase *ampC* expression by
168 causing an intracellular accumulation of peptidoglycan catabolites^{14,17}. However, *ampC*
169 induction via this mechanism requires the AmpR transcription factor; inactivation of *ampR*
170 removes the ability to increase *ampC* expression in response to β -lactams¹⁷. Crucially, among
171 our strains *ampR* is only present in the genomes of *P. aeruginosa* PAO1, *P. protegens* Pf-5, and
172 the two *P. fluorescens* Pf0-1 and SBW25, and not found in the others. This simple association

173 between the presence of the AmpR transcription factor and the probability of survival to the end
174 of the experiment through adaptation suggests that regulation of *ampC* expression is key.

175 How does *ampR* increase evolvability? One simple possibility is that this regulator potentiates
176 evolution by opening up new genetic paths to evolving elevated ceftazidime resistance^{9,10}.
177 Specifically, *ampR* could potentiate the evolution of ceftazidime resistance by allowing
178 mutations in peptidoglycan biosynthesis genes, such as *ampD*, *mpl*, and *dacB* to increase levels
179 of *ampC* expression. Consistent with this hypothesis, mutations in peptidoglycan biosynthesis
180 genes and *dacB* are known to only increase resistance in the presence of *ampR*^{17,18}. This
181 hypothesis generates two simple predictions that can be tested using our method. First, if
182 elevated expression of *ampC* is a key mechanism for evolving ceftazidime resistance, then
183 deleting *ampC* should decrease evolvability. Second, if the AmpR regulator is required to drive
184 the evolution of increased *ampC* expression, then deleting *ampR* should reduce evolvability by
185 the same amount as deleting the *ampC*. To test these predictions, we challenged populations of
186 $\Delta ampR$ and $\Delta ampC$ mutants of *P. aeruginosa* PAO1 with increasing doses of ceftazidime, as in
187 our initial experiment (Figure 3a). Both of the mutants have dramatically reduced evolvability
188 compared to their isogenic *P. aeruginosa* PAO1 control (Cox's proportional hazard, likelihood
189 ratio=23.82, df=2, P=6×10⁻⁶) providing conclusive evidence that both the β -lactamase (*ampC*)
190 and its regulator (*ampR*) play key roles in driving the evolution of elevated ceftazidime
191 resistance.

192 The low survival probability of *P. aeruginosa* PAO1 in comparison with the other strains that
193 carry both *ampR* and *ampC* is also consistent with this hypothesis. Strains of *P. fluorescens* and
194 *P. protegens* carry 2 homologs of *ampD*, which represses the expression of *ampC*, whereas *P.*
195 *aeruginosa* PAO1 carries 3 homologs of this gene. The additional copy of *ampD* found in *P.*
196 *aeruginosa* ensures that *ampD* mutations lead to weaker de-repression of *ampC* expression, and
197 this is likely to translate into reduce evolvability in comparison to strains with only 2 *ampD*
198 homologs; the *ampD* dosage effect has been demonstrated experimentally²⁰. Consequently, most
199 surviving *P. aeruginosa* strains possessed two loss of function mutations in the peptidoglycan
200 biosynthesis pathway, in comparison with one only in the other *ampR/ampC* possessing strains
201 (Figure 2).

202 Additionally, it is possible that adaptive plasticity in *ampC* expression mediated by *ampR* could
203 increase evolvability²². Exposure to β -lactam antibiotics interferes with peptidoglycan
204 biosynthesis by inhibiting PBPs, causing an AmpR-mediated increase in *ampC* expression^{17,19}.
205 This, in turn, may accelerate the genetic evolution of resistance by providing bacterial
206 populations with the time to acquire ceftazidime resistance mutations. According to this
207 explanation, *ampR* increases evolvability through ecological potentiation. The key assumption of
208 this hypothesis is that the plasticity in *ampC* expression mediated by *ampR* must provide a
209 benefit in the presence of ceftazidime. To test this hypothesis, we measured the effect of deleting
210 *ampR* and *ampC* on fitness using short-term competition experiments (Figure 3b). Deleting
211 *ampC* leads to a decrease in fitness the presence of ceftazidime, demonstrating that induced
212 expression of this gene is beneficial. However, deleting *ampR* actually increases fitness in the
213 presence of sub-MIC concentrations of ceftazidime, demonstrating that plasticity in gene
214 expression cannot explain the link between *ampR* and increased evolvability. Indeed, as *ampR*
215 expression is not particularly strongly induced by ceftazidime²³, this suggests that *ampR* does not
216 simply allow populations to 'buy time' to wait for an adaptive mutation. Although this result is

217 counter-intuitive, it is important to emphasize that *ampR* is a global transcriptional regulator that
218 affects the expression of 100s of genes^{24,25}, including repressing another chromosomal β -
219 lactamase, *poxB*²⁵. In particular, *ampR* is involved in regulating quorum sensing factors,
220 including *lasR*, several metabolic pathways, and the *rpoS*-mediated stress response pathway²⁵.
221 Although it is clear that inducing elevated levels of *ampC* expression in the presence of
222 ceftazidime is beneficial, the fitness cost associated with the *ampR* regulator implies that the net
223 fitness effect of all of the changes in gene expression caused by this regulator in the presence of
224 ceftazidime is deleterious. The importance of *ampR* as a global regulator of expression perhaps
225 explains why increased *ampC* expression did not arise through mutations in *ampR* itself, and
226 why *ampR* mutations are not typically observed in clinical *P. aeruginosa* isolates²⁶.

227 ***Inhibiting the evolution of ceftazidime resistance*** Given the important role that *ampR* mediated
228 induction of *ampC* expression plays in the evolution of resistance, our results suggest that one
229 possible strategy to prevent the evolution of cephalosporin resistance in *P. aeruginosa* infections
230 would be to co-administer ceftazidime with AmpC β -lactamase inhibitors²⁷. The rationale for
231 this strategy is that a combination of a β -lactam and β -lactamase inhibitor will be active against
232 both wild-type bacterial strains and mutants with elevated β -lactamase secretion. In other words,
233 this strategy should effectively block a major evolutionary path to elevated resistance. To test
234 this idea, we challenged *P. aeruginosa* PAO1 with ceftazidime in the presence of avibactam²⁸, a
235 recently developed AmpC inhibitor (Figure 4a). Unlike most β -lactamase inhibitors, avibactam
236 does not possess any toxic effects on *Pseudomonas*²⁸ and we did not detect any population
237 extinction in the avibactam treated control populations. In support of our hypothesis, avibactam
238 increased the rate of population extinction in the presence of ceftazidime compared to
239 ceftazidime treated control populations (Cox's proportional hazard, likelihood ratio test=78.968,
240 df=1, $P < 10^{-6}$). We failed to detect any viable cells in 59 out of 60 populations of *P. aeruginosa*
241 that were selected in 8 \times MIC ceftazidime supplemented with avibactam demonstrating that the
242 effect of avibactam suppresses the evolution of elevated β -lactamase secretion just as effectively
243 as knocking out *ampC* or *ampR* (Figure 3a). Importantly, this effect does not arise because
244 avibactam increases the potency of ceftazidime. Surprisingly, we found that avibactam treatment
245 actually increased the MIC of ceftazidime from 0.76 mg/L to 1.14 mg/L (Figure S2).

246 As a final test of the role of *ampR* in evolvability, we challenged populations of $\Delta ampR$
247 (Figure 4b) and $\Delta ampC$ (Figure 4c) mutants of *P. aeruginosa* PAO1 with a combination of
248 ceftazidime and avibactam, as in our experiment with wild-type *P. aeruginosa* PAO1. If our
249 hypothesis is correct, then avibactam should have no effect on evolvability in these mutant
250 strains, because they are effectively unable to increase *ampC* expression under our experimental
251 conditions. Consistent with this idea, we found that avibactam does not have an effect on
252 evolvability in either $\Delta ampR$ or $\Delta ampC$ mutants (Cox's proportional hazard, likelihood ratio
253 test=3.25, df=1, $P = 0.071$ and likelihood ratio test=0.02, df=1, $P = 0.876$, respectively).

254

255 Conclusion

256 Whole genome sequencing is revolutionizing our understanding of the evolution and
257 ecology of bacterial pathogens. One of the challenges that has arisen from this revolution is to
258 understand the consequences of genetic diversity in pathogen populations. Here we show that
259 comparative experimental evolution can be used to identify genes and pathways that influence

260 the rate and mechanisms of adaptation to antibiotics. Our experiment addressed this problem at a
261 fairly broad scale, by comparing the evolutionary responses of strains from different species. Our
262 initial reasoning for working at this scale was that comparing divergent strains effectively
263 maximizes the number of genes and SNPs that are included in the experiment, therefore
264 maximizing the likelihood of detecting an impact of genetic background on evolvability.
265 However, the sheer number of genetic differences between even the most closely related strains
266 used in this study may have hindered our ability to detect more subtle genomic effects on
267 evolvability. While it is clear that inducible *ampC* β -lactamase expression is an important driver
268 of evolvability in this genus, it is clear that other genes must influence the ability to evolve
269 ceftazidime resistance. For example, *P. stutzeri* ATCC17588 and *P. putida* KT2440, both of
270 which lack *ampR*, have similar evolvability to *P. aeruginosa* PAO1. We are currently extending
271 this research program by focusing on studying variation in evolvability between clones from the
272 same species, and we hope that this approach will enable us to identify genetic drivers of
273 evolvability in greater depth.

274 The differing modes of *ampC* expression among the pseudomonads affect their ability to evolve
275 resistance to β -lactams by interacting with genes in the peptidoglycan biosynthesis pathway. In
276 strains possessing *ampR*, the intracellular accumulation of peptidoglycan catabolites converts the
277 AmpR transcription factor into an activator of *ampC* expression in response to peptidoglycan
278 damage. Mechanistically, *ampR* increases evolvability by allowing mutations in peptidoglycan
279 biosynthesis genes to induce high levels of β -lactamase expression, which effectively amplifies
280 the *ampC* expression plasticity that occurs when cell wall synthesis is compromised by β -
281 lactams²⁰. From a more conceptual perspective, *ampR* can be thought of as a conduit that
282 translates genetic variation in the peptidoglycan biosynthesis gene network into phenotypic
283 variation in *ampC* expression. This suggests that response pathways that are involved in sensing
284 environmental change may have a general role as evolutionary catalysts, linking plastic and
285 mutational responses to environmental change. Intriguingly, these alternative expression modes
286 are disseminated among the enterobacteria; however, insertion of the *ampR* gene into
287 constitutive producers is not sufficient to restore inducible expression, suggesting a distinct
288 regulatory mechanism in constitutive producers²⁹. To avoid the evolution of high levels of
289 ceftazidime resistance, and subsequent treatment failure, treatment with ceftazidime should be
290 avoided in infections caused by strains with inducible *ampC* expression.

291 Understanding the evolutionary trajectory to high levels of ceftazidime resistance makes it
292 possible to design a simple two-drug mixture consisting of ceftazidime and avibactam that can be
293 used to effectively eliminate populations of the pathogen *P. aeruginosa*. We argue that this
294 strategy is successful because avibactam effectively prevents mutations in peptidoglycan
295 biosynthesis genes and *dacB* from increasing ceftazidime resistance, eliminating their fitness
296 benefit. One possible solution to this evolutionary challenge would be to first evolve avibactam
297 resistance, and then evolve ceftazidime resistance. However, avibactam does not have any
298 detectable toxic effects on *Pseudomonas* at concentrations where it is able to effectively inhibit
299 AmpC, rendering this evolutionary pathway to combined avibactam/ceftazidime resistance
300 inaccessible. Although these results are encouraging, we emphasize that there are a number of
301 confounding factors that may affect the efficacy of this combination of drugs *in vivo*. For
302 example, the pharmacokinetic properties of the two drugs may make it difficult to effectively
303 maintain the drug mixtures at the site of bacterial infections, and it is also possible that

304 avibactam resistant alleles of *ampC* or other β -lactamases capable of hydrolyzing ceftazidime are
305 already present in pathogen populations.

306 Predicting the evolution of antibiotic resistance is a challenging and important objective. Here
307 we show that comparative experimental evolution can be used to identify genes and pathways
308 that make some bacterial strains prone to evolving resistance, and to exploit this to design
309 treatment strategies for preventing resistance evolution. High throughput sequencing is
310 revolutionizing clinical microbiology^{30,31}, and it may be possible to identify such potentiator
311 genes in clinical pathogen populations and to use this information to optimize antimicrobial
312 treatment strategies.

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317 **Statement of author contributions:** This study was designed by RCM. Experiments were
318 carried out by VF, AP and TV. Bioinformatics were done by DRG. VF, AP, DRG and RCM
319 analyzed data. AO contributed reagents and expertise. The manuscript was written by RCM,
320 DRG and VF.

321 **Competing interests:** The authors declare the absence of any competing interests.

322

323 **Materials and Methods**

324 *MIC Determination for parental strains*

325 Three independent estimations of the MIC for each parental strain were determined in 96-well
326 plates using the broth microdilution method. Briefly, 5-10 morphologically similar colonies of
327 each strain were resuspended in sterile saline solution (NaCl 0.9%). The solution was adjusted to
328 the adequate optical density so that it would contain approximately 1.5×10^8 cells/mL. This
329 standardized inoculum and was diluted a further 200-fold in Mueller-Hinton 2 (MH2, Sigma-
330 Aldrich, United Kingdom) broth containing ceftazidime (Sigma-Aldrich) at a concentration
331 between 64 mg/L and 0.0625 mg/L. After 24h of incubation at 30 °C with shaking at 250 rpm,
332 optical density at 595nm was determined for each well with a Synergy 2 plate reader (Biotek,
333 Winooski, USA). We considered that bacterial growth had been inhibited if the optical density
334 was less than 25% of that of antibiotic-free cultures. The lowest antibiotic concentration at which
335 growth had been inhibited was considered the MIC. The measured MIC was used to calculate the
336 ramping ceftazidime concentration regime in the selection experiment (see "Experimental
337 evolution").

338 *Effect of avibactam on MIC*

339 The effect of avibactam on MIC was evaluated by measuring growth inhibition by ceftazidime at
340 the presence/absence of avibactam. The procedure was identical to MIC determination described
341 above, except that one group of replicates was supplemented with 4 mg/L of avibactam
342 (BioVision Inc. USA). No avibactam was added to a control group. The avibactam treatment and
343 control groups were tested at concentrations ranging from 0.1 mg/L to 3.8 mg/L of ceftazidime
344 with 4 replicates each.

345 *Mutation rate estimation*

346 Mutation rates were estimated by fluctuation assays, with the antibiotic rifampicin as the
347 selection agent, using the method of Luria and Delbruck³². 480 replicate cultures were inoculated
348 with approximately 50 cells from an overnight culture of each parental strain and incubated for
349 48 hours in 200 µl of KB media at 30 °C with constant shaking at 200 rpm. Approximately 10⁷
350 cells from each culture were then plated onto KB-agar containing rifampicin at the appropriate
351 MIC (minimum inhibitory concentration) for each strain (60 mg/L for *P. aeruginosa* PAO1, 30
352 mg/L for all other strains). For each strain the proportion of cultures yielding no mutants was
353 scored, from which the mutation per culture was calculated using the negative natural logarithm.
354 This value was then divided by the number of cells plated, which provides an estimate of the
355 mutation rate per cell division.

356 *Experimental evolution*

357 To initiate the selection experiment, all parental strains were recovered from -80 °C stocks and
358 cultured overnight in MH2 broth at 30 °C for 24 h with shaking at 250 rpm. Next, the cultures
359 were diluted by 10⁻⁶ in MH2 broth and distributed on 96-well plates (200 µL per well). After 48 h
360 of incubation at 30°C, we initiated the first transfer by diluting these cultures 1:100 in MH2
361 broth containing 1/8 MIC of ceftazidime, relative to the measured MIC of each strain. Bacterial
362 populations were incubated for 24 h at 30°C with shaking at 250 rpm and diluted 1:100 for the
363 next transfer. Every transfer ceftazidime concentration was doubled, reaching 8× MIC in the
364 final transfer. Population survival was monitored during the course of the selection experiment
365 by measuring optical density at 595 nm using a Synergy 2 microtiter plate reader (BioTek,
366 Winooski, VT, USA). We additionally confirmed population survival after the last transfer by
367 plating a 1 uL sample of each population on antibiotic-free MH2 agar plates that were scored for
368 growth after overnight incubation at 30 °C. We performed the evolution experiment in two
369 independent blocks. In each block we propagated 60 replicates populations of each strain that
370 were challenged with increasing doses of ceftazidime and 12 replicate controls populations of
371 each strain that were allowed to evolve in antibiotic-free MH2. At the end of the experiment, a
372 maximum of 20 population per strain were streaked in MH2 agar plates and a clone was picked
373 for each population was picked and amplified for further analyses. To avoid bias by conducting
374 the experiment at different temperatures and incubators, a common growth environment (i.e. 30
375 °C, MH2) and growth medium (Mueller-Hinton 2) that supports the growth of all strains was
376 chosen for all strains. Although the strains have different optimal growth temperatures (28 °C for
377 *P. protegens*, *P. fluorescens*, and *P. fulva*; 30 °C for *P. putida* and *P. mendocina*, 35 °C for *P.*
378 *stutzeri* and 37°C for *P. aeruginosa*), all were capable of vigorous growth in this environment,
379 hence the number of generations per day (6-7) is instead dictated by the dilution factor (1/100).

380 *Experimental evolution with ΔampC and ΔampR mutants*

381 We obtained ΔampC and ΔampR mutants of *P. aeruginosa* PAO1 that were constructed
382 following well-established procedures based on the *cre-lox* system for gene deletion and
383 antibiotic resistance marker recycling³³. We determined the MIC of these mutants using the
384 microbroth dilution method, as above. To test evolvability of ΔampC and ΔampR mutants, we
385 followed the same protocol as the main selection experiment, as outlined above. We challenged
386 90 replicate populations of each deletion mutant and 30 replicate populations of PAO1 wild-type

387 with increasing doses of ceftazidime. In addition, we evolved 18 control populations per strain in
388 antibiotic-free culture medium. This experiment was carried out in a single block.

389 *Experimental evolution to test the effect of avibactam*

390 The effect of avibactam on evolvability was tested for $\Delta ampC$ and $\Delta ampR$ mutants and for wild-
391 type PAO1. 120 replicate populations of each strain were passaged following exactly the same
392 procedure as in the two previous experiments. The ceftazidime concentration was doubled every
393 transfer from 1/8 to 8× MICs. For each strain, half of the populations (60 replicates) were
394 additionally challenged with avibactam (always 4 mg/L, BioVision Inc. USA). Population
395 survival was monitored for 7 serial transfers by measuring optical density. We also included 20
396 control populations evolving at the presence of avibactam but without the antibiotic. There was
397 no extinction observed in the control treatment.

398 *Inhibition zone assays*

399 Evolved clones were cultured in MH2 broth overnight (30 °C, 250 rpm). A sterile swab was
400 dipped then into a 10⁻³ dilution of this overnight culture to and the swab was used to inoculate
401 the surface of three MH2 agar plates. Then we placed four different antibiotic susceptibility
402 testing discs (Oxoid) on each plate, testing a total of 12 antibiotics: ceftazidime, piperacillin,
403 meropenem, imipenem, aztreonam, cloramphenicol, tetracycline, rifampicin, amikacin,
404 tobramycin, ciprofloxacin and levofloxacin. After 24 h of incubation at 30 °C, the diameter of
405 the different inhibition zones was measured with a ruler taking the average of three
406 measurements in different axis. Assays were performed in 4 randomized blocks containing a
407 similar number of evolved clones for each strain, and all ancestral strains were tested in each
408 block as a control. Change in antibiotic sensitivity was estimated as the difference in diameter of
409 the inhibition zone of each clone compared to its ancestor for each antibiotic.

410 *DNA extraction and sequencing*

411 DNA from the evolved clones surviving the duration of the experiment was extracted using the
412 Wizard Genomic DNA Purification Kit (Promega, UK) as per the manufacturer's instructions.
413 To maximize phylogenomic coverage and reduce bias toward *P. fluorescens* strains, *P.*
414 *fluorescens* SBW25 was excluded from sequencing due to being highly similar to *P. fluorescens*
415 Pf0-1. We assessed the purity of DNA extractions by measuring absorbance at 230, 260, and 280
416 nm and by visualizing migration on a 0.7% agarose gel. The concentration of each genomic
417 DNA in each sample was then accurately determined using QuantiFluor^{ds}DNA System
418 (Promega, UK) and samples were diluted to 30 ng/μL in TE Buffer before sequencing.

419 Resequencing was done using Illumina HiSeq2000 with 100bp paired-end reads (Wellcome
420 Trust Centre for Human Genetics, Oxford, UK). Sequencing analysis was performed using the
421 pipeline first described in San Millan et al.³⁴. Read filtering was done using the NIH-
422 QCToolkit³⁵. Read ends were trimmed if the Phred quality score was less than 20. We discarded
423 reads <50bp after trimming, with >2% ambiguous bases, or with >20% bases of Phred score <20.
424 BWA was used to map reads to the reference genome of each strain. Mapped reads were
425 processed to increase the quality of the variant calling: 1) reads with multiple best hits were
426 discarded; 2) duplicated reads were discarded using MarkDuplicates from the Picard package
427 (<http://picard.sourceforge.net>); 3) reads around indels were locally realigned using
428 RealignerTargetCreator and IndelRealigner from the GATK package to correct for

429 misalignment; and 4) mate pairs were sorted using FixMateInformation in the Picard package.
430 Variant calling was performed with GATK UnifiedGenotyper³⁶ and Samtools mpileup³⁷.
431 VCFtools vcf-annotate³⁸, and GATK toolkit VariantFiltration³⁹, were used to filter the raw
432 variants for strand bias, end distance bias, base quality bias, SNPs around gaps, low coverage
433 and erroneously high coverage. Variants were combined using GATK's CombineVariants
434 (keeping any unfiltered). High quality variants not filtered were annotated using SnpEff⁴⁰. Three
435 approaches were used to detect structural variants: BreakDancer⁴¹ (indels, inversions and
436 translocations), Pindel⁴² (indels, inversions, tandem duplications and breakpoints), and
437 ControlFREEC (copy number variants⁴³ with mappability tracks generated by gem-mappability
438 (GEM library⁴⁴).

439 *Comparative genomics of resistance pathways*

440 Using pairwise reciprocal BLAST between the reference sequences of the sequenced strains, we
441 determined their similarity in genome content. This approach was taken because the strains differ
442 in the extent to which their genomes are annotated. Using the KEGG database⁴⁵, we compared
443 the genes in the β -lactam resistance and peptidoglycan recycling pathways (irrespective of
444 whether they had mutated during selection).

445 *Competition experiment with $\Delta ampC$ and $\Delta ampR$ mutants*

446 To measure relative fitness of the deletion mutants, we performed a competition experiment.
447 $\Delta ampC$, $\Delta ampR$ and their isogenic PAO1 wild-type were competed against a YFP-marked tester
448 strain PAO1 strain that carries a constitutively expressed YFP integrated at the mini-Tn7
449 insertion site¹⁴. Competition experiments were carried out in MH2 broth containing ceftazidime
450 at a concentration of 0, 0.25 or 0.5 mg/L. All competition experiments were replicated 9 times.
451 First, the strains were recovered from -80 °C stock and cultured overnight in MH2 broth medium
452 at 30 °C with shaking at 250 rpm. The overnight cultures were diluted 1:50 in MH2 broth and
453 used to prepare 1:1 mixtures of PAO1-YFP with each of the 3 strains to be tested. Before starting
454 competition, we first estimated the exact starting proportion of strains using flow cytometry (for
455 details see below). Next, we combined 10 μ L of these mixtures and 190 μ L of MH2 with a
456 corresponding ceftazidime concentration (0, 1/4 and 1/2 MICs). This resulted in an additional
457 1:20 dilution. The bacterial strains were let to compete in 96-well plates for 24 h at 30 °C. The
458 next day, the cultures were diluted 1:50 in saline solution (0.9% NaCl) and analyzed on a flow
459 cytometer in order to estimate the resulting proportion of the YFP-labeled versus unlabelled cells
460 after competition (see below).

461 Flow cytometry was performed on Accuri C6 (BD Biosciences, UK). The cell densities were
462 adjusted to give around 1000 events per second. During data acquisition, a lower cut off was set
463 at 10,000 for FSC-H and at 8000 for SSC-H. The data were exported as FCS-files and processed
464 in R using a custom pipeline based on flowCore and flowViz packages⁴⁶⁻⁴⁸. In the pipeline, the
465 events were automatically gated on size by retaining the cells within 2 standard deviations
466 around the median in the bivariate normal distribution of FSC-A and SSC-A. Then, k-mean
467 clustering algorithm was applied on fluorescence intensity FL1-H to differentiate fluorescent
468 versus non-fluorescent cells. For each antibiotic concentration, we ensured that YFP-expressing
469 strain can be well separated from non-fluorescent strains by overlaying non-mixed controls
470 (overlap is usually less than 2% of the cells). Figure S3 shows a representative plot of the gating
471 strategy.

472 Relative fitness was calculated according to the formula

$$473 w = \log_2[p_1/(p_0/1000)] / \log_2[(1-p_1)/[(1-p_0)/1000]],$$

474 where p_0 is an initial proportion of an unlabelled stain, and p_1 is a final proportion of an
 475 unlabelled stain after competition. 1000 is a dilution factor, which reflects a difference in cell
 476 density at the beginning and at the end of the competition.

477 **Data availability**

478 Data generated or analysed during this study are included in this published article (and its
 479 supplementary information files), with the exception of sequence data, which are deposited in
 480 European Nucleotide Archive (PRJEB20060).

481

482 **Figure legends**

483

484 **Figure 1:** Responses of *Pseudomonas* to ceftazidime. **a** Phylogeny of the strains used in this
 485 study, all nodes were supported with >99% confidence and the scale bar shows genetic distance
 486 (adapted from ref. 11 and 49 with permission under Creative Commons licence CC-BY-4.0). **b**
 487 The proportion of populations ($n = 120$ populations/strain) of each strain that survived exposure
 488 to increasing doses of ceftazidime. Doses were standardized relative to the MIC of the ancestral
 489 clone of each strain, and doses increased 2 fold daily up to $8\times$ MIC. Evolvability differs between
 490 strains that are not connected by red lines (Post-hoc test on Cox's proportional hazard, $P<0.05$).

491

492 **Figure 2: Resistance in evolved clones.** Each column in this figure represents a single,
 493 randomly chosen clone from a population that survived until the end of the selection experiment
 494 ($8\times$ MIC). **a** Black boxes show the presence of mutations in known ceftazidime resistance genes,
 495 as determined by whole genome resequencing. Note that some clones carry mutations in multiple
 496 resistance genes, and that some clones lack mutations in known resistance genes (online
 497 supplementary data table S2). **b** Coloured boxes show the change in ceftazidime MIC of evolved
 498 clones (mean of $n=3$ replicates), and **c** changes in the zone of inhibition for a large panel of
 499 antibiotics, as determined by disc diffusion assay (mean of $n=3$ replicates).

500

501 **Figure 3: The AmpR transcription factor potentiates the evolution of ceftazidime resistance**
 502 **in *P. aeruginosa* PAO1.** **a** The survival of populations of an *ampR* deletion strain
 503 (PAO1:: $\Delta ampR$; $n = 90$) relative to an isogenic PAO1 control ($n = 30$) under increasing doses of
 504 ceftazidime. The *ampR* deletion reduces evolvability to levels comparable to those observed in a
 505 mutant lacking the *ampC* β -lactamase (PAO1:: $\Delta ampC$; $n=90$). **b** Relative fitness (mean +/- s.e; n
 506 = 9) of the PAO1:: $\Delta ampR$ mutant (grey triangles) and the PAO1:: $\Delta ampC$ mutant (blue circles) in
 507 direct competition with a PAO1 reference strain carrying a neutral YFP marker. Symbols denote
 508 statistical significance, as determined by a Bonferroni-corrected Wilcoxon rank sum test (N.S. =
 509 $P>0.05$, * = $P<0.05$, *** = $P<0.001$).

510

511 **Figure 4: Blocking the evolution of ceftazidime resistance.** **a** Survival of populations of *P.*
 512 *aeruginosa* PAO1 that were challenged with increasing doses of ceftazidime in either the
 513 presence or absence of the AmpC-inhibitor avibactam ($n = 60$ populations/treatment). Avibactam
 514 was administered at a constant, non-inhibitory dose (4 mg/L). Avibactam increases the rate of
 515 population extinction in the presence of increasing doses of ceftazidime. **b** and **c** The survival of
 516 *ampR* or *ampC* deletion strains (PAO1:: $\Delta ampR$ and PAO1:: $\Delta ampC$) under the same
 517 experimental conditions as for the isogenic wild-type POA1 ($n = 60$ populations/treatment for
 518 each strain). Avibactam had no effect on the survival of *ampR* or *ampC* deletion mutants.

519

520 **References**

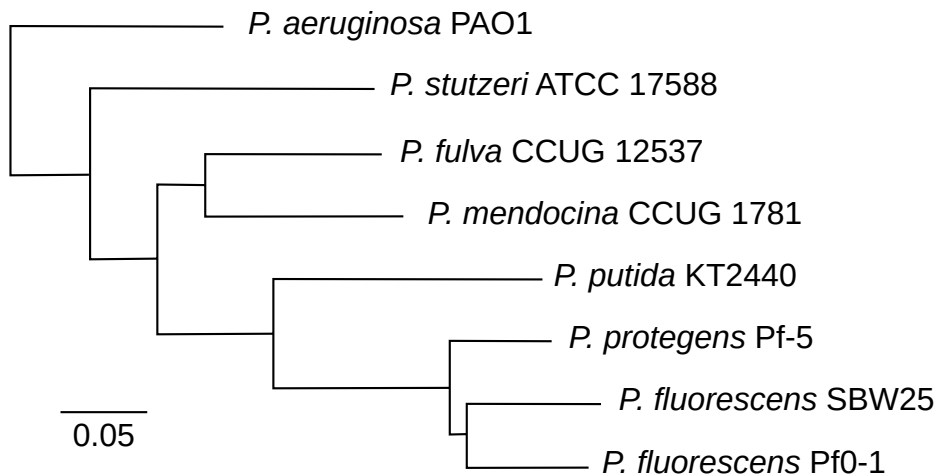
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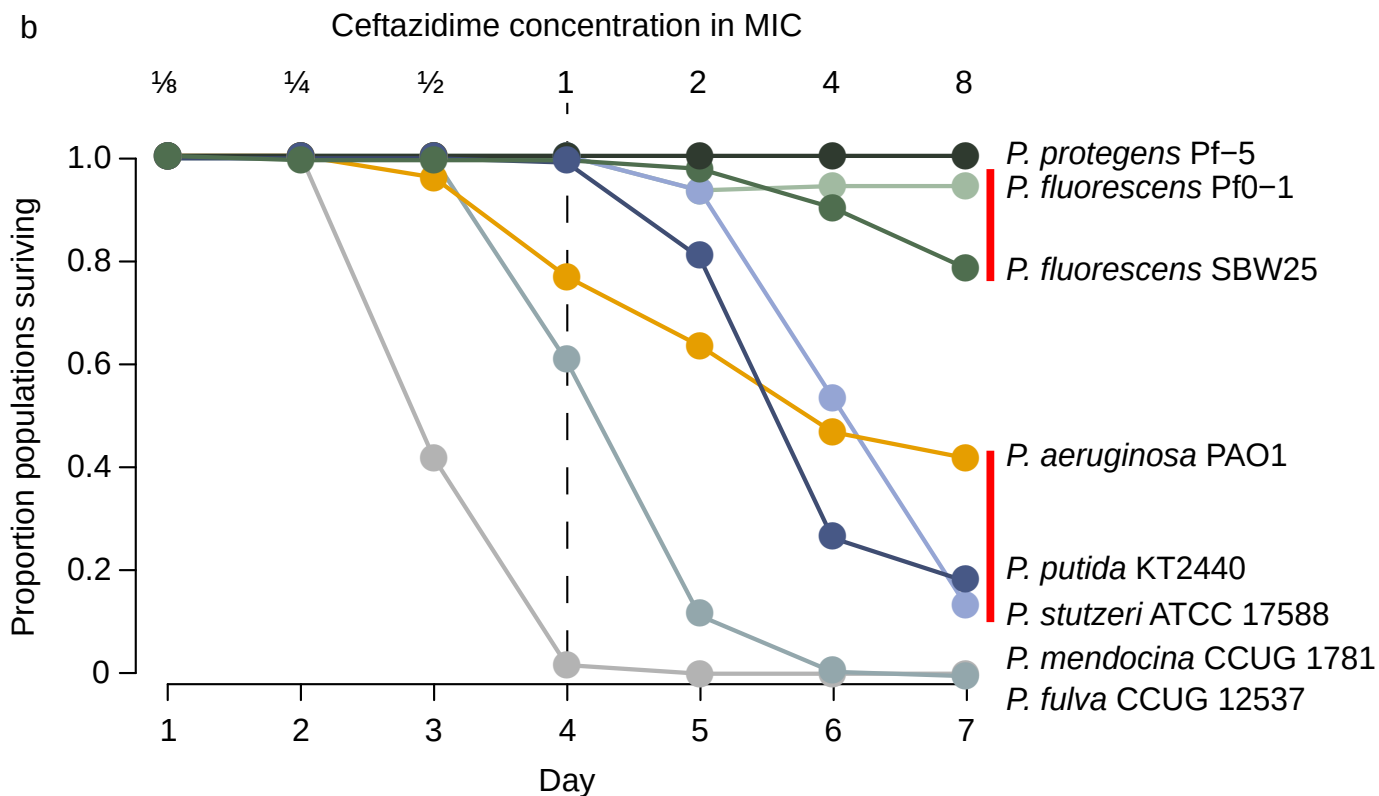
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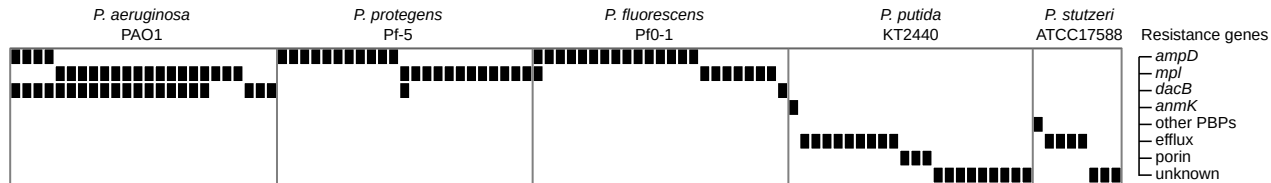
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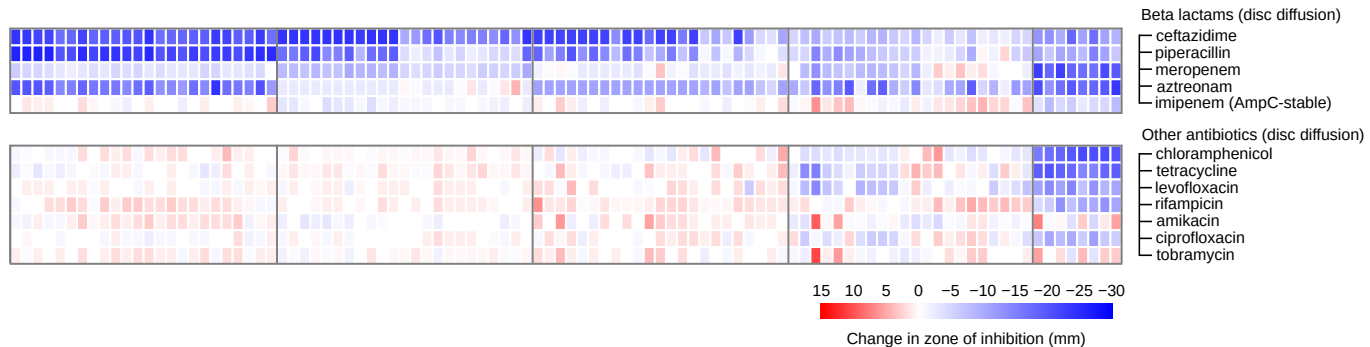
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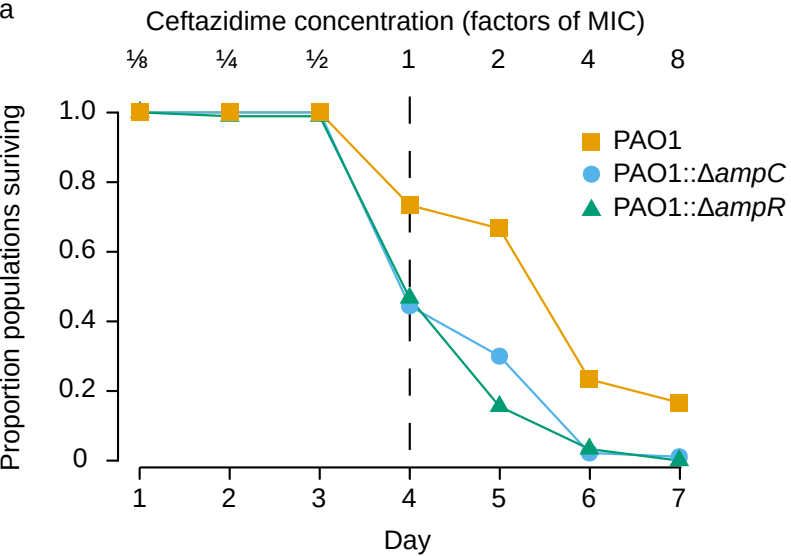
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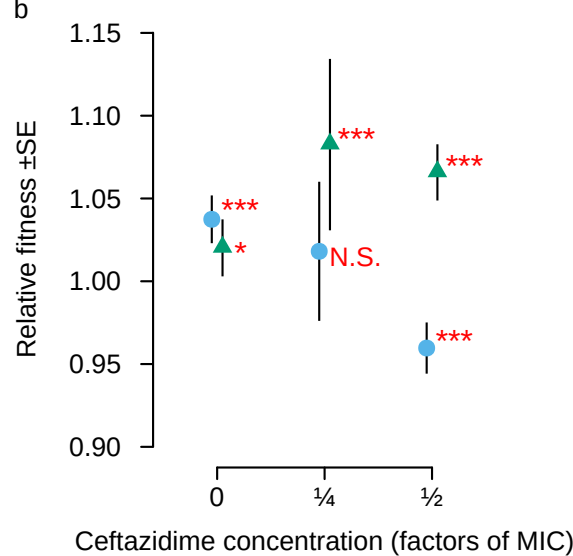
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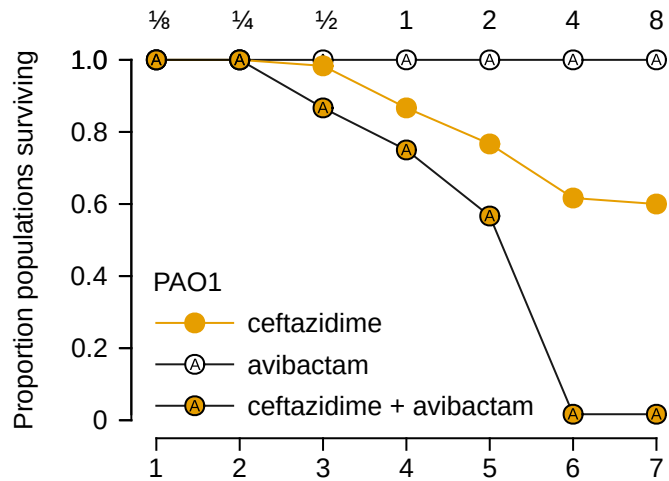
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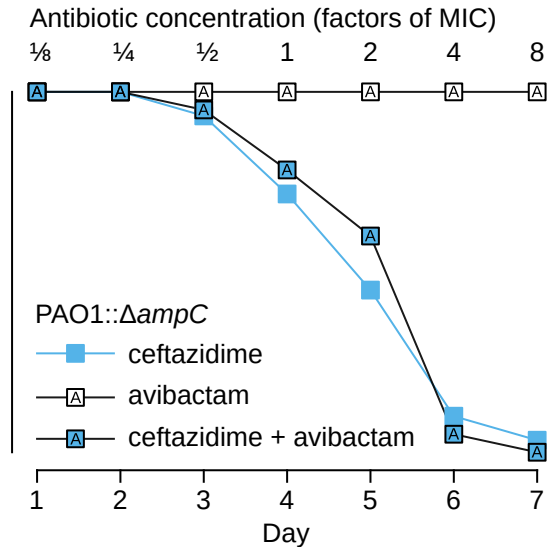
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