

1 **Environmental fluctuations do not select for increased**  
2 **variation or population-based resistance in *Escherichia coli***

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14 Running title: **Fluctuations do not increase variation**

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16 Submitted for review.

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28 **Abstract**

29  
30 In nature, organisms often face unpredictably fluctuating environments. However, little is  
31 understood about the mechanisms that allow organisms to cope with such unpredictability. To  
32 address this issue, we used replicate populations of *Escherichia coli* selected under complex,  
33 randomly changing environments. We assayed growth at the level of single cells under four  
34 different novel stresses that had no known correlation with the selection environments. Under  
35 such conditions, the individuals of the selected populations had significantly lower lag and  
36 greater yield compared to the controls. More importantly, there were no outliers in terms of  
37 growth, thus ruling out the evolution of population-based resistance. We also assayed the  
38 standing phenotypic variation of the selected populations, in terms of their growth on 94  
39 different substrates. Contrary to extant theoretical predictions, there was no increase in the  
40 standing variation of the selected populations, nor was there any significant divergence from the  
41 ancestors. This suggested that the greater fitness in novel environments is brought about by  
42 selection at the level of the individuals, which restricts the suite of traits that can potentially  
43 evolve through this mechanism. Given that day-to-day climatic variability of the world is rising,  
44 these results have potential public health implications. Our results also underline the need for a  
45 very different kind of theoretical approach to study the effects of fluctuating environments.

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**Keywords:** evolvability, standing variation, experimental evolution, antibiotic resistance, neutral space

## 57 INTRODUCTION

58 The last few decades have witnessed a global increase in day-to-day climatic variability  
59 (Medvigy & Beaulieu, 2012). As a result of this, many organisms are now subjected to  
60 environmental changes at much shorter time scales than what they would have probably  
61 experienced for much of their evolutionary history. This has led to a number of empirical  
62 (reviewed in Kassen, 2002; Hedrick, 2006) and theoretical (Levins, 1968; Ishii *et al.*, 1989;  
63 Taddei *et al.*, 1997) studies, that seek to investigate the effects of environmental variability on  
64 the physiology (Hagemann, 2011) and evolution (Coffey & Vignuzzi, 2011; Ketola *et al.*, 2013)  
65 of organisms. The primary insight that has emerged from these studies is that various aspects of  
66 the environmental heterogeneity, e.g. the number of components that constitute the environment  
67 (Barrett *et al.*, 2005; Cooper & Lenski, 2010), the speed with which the environment changes  
68 (Ancel, 1999; Cohan, 2005; Meyers *et al.*, 2005) or the predictability of environmental changes  
69 (Hughes *et al.*, 2007; Alto *et al.*, 2013) - can act singly, or in combinations with each other, to  
70 affect the evolutionary trajectory of populations. More interestingly, such fluctuations can lead  
71 to very different patterns of fitness in different test environments. For instance, in a recent study,  
72 when replicate populations of *E. coli* were subjected to fluctuating complex environments  
73 (random, stressful combinations of pH, salt and H<sub>2</sub>O<sub>2</sub>), the selected populations had no fitness  
74 advantage over the controls in stresses in which they were selected (i.e. pH or salt or H<sub>2</sub>O<sub>2</sub> or  
75 combinations thereof) (Karve *et al.*, 2015). Yet, the same selected populations had significantly  
76 greater fitness in completely novel environments that had never been encountered by the bacteria  
77 before and had no known correlation with the stresses under which they had been selected  
78 (Karve *et al.*, 2015). Similar patterns of advantage under novel environments have been observed  
79 in other bacteria (Ketola *et al.*, 2013) and viruses (Turner & Elena, 2000), when subjected to

80 fluctuating selection pressure (although see Coffey & Vignuzzi, 2011). These results are  
81 consistent with the general observation that disturbed habitats give rise to a large number of  
82 invasive species which, by definition, have fitness advantages in novel environments (Lee &  
83 Gelembiuk, 2008 and references therein).

84

85 Unfortunately, in spite of a considerable corpus of theoretical predictions (Levins, 1968; Meyers  
86 *et al.*, 2005), there is little empirical work on the mechanisms that allow organisms to adapt to  
87 novel environments. Two major ways by which organisms can have greater fitness in novel  
88 environments are through an enhanced capacity to generate adaptive variations or by possessing  
89 larger amount of standing genetic variation. Although several organisms are known to respond to  
90 stress through increased mutation rate (Bjedov *et al.*, 2003) or enhanced phenotypic variation  
91 (Rohner *et al.*, 2013), it is not clear whether such traits can evolve due to exposure to  
92 environmental fluctuations. Recently, it has been shown that exposure to complex fluctuating  
93 environments do not lead to a significant change in mutation rates in *E. coli* (Karve *et al.*, 2015).  
94 However, it is hard to generalize on the issue as empirical studies on evolutionary effects of  
95 environmental fluctuations often do not investigate changes in mutation rates. The situation is  
96 not much different *w.r.t* the evolution of standing genetic variation under fluctuating  
97 environments. It has been shown *in vitro* that accumulated cryptic genetic variation in ribozymes  
98 can increase fitness in novel environments (Hayden *et al.*, 2011). However, chikungunya virus  
99 populations selected under fluctuating environments, show much less increase in genetic  
100 diversity compared to those raised in constant environments (Coffey & Vignuzzi, 2011). Again,  
101 generalization of any kind is difficult, since we could not locate any other study that reports the  
102 changes in genetic variation in response to selection in fluctuating environments.

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104 Apart from these variation-based mechanisms, there are a few other ways in which organisms  
105 can potentially deal with novel environments. The most well investigated of these seems to be  
106 phenotypic plasticity (Fusco & Minelli, 2010) which is expected to evolve when the environment  
107 changes faster than the life span of the organisms (Ancel, 1999; Meyers *et al.*, 2005). Another  
108 potential mechanism in this context might be an increase in broad-spectrum stress tolerance  
109 which is consistent with a recent finding that enhanced efflux activity evolves in *E. coli* in  
110 response to selection in fluctuating environments (Karve *et al.*, 2015). The third possible way to  
111 have greater fitness in novel environments is the evolution of population-based resistance,  
112 wherein a small fraction of individuals in the population synthesize some chemicals into the  
113 environment, which allows the entire population to become stress resistant (Lee *et al.*, 2010;  
114 Vega *et al.*, 2012). This kind of division of labor, in principle, can allow the population to  
115 become resistant to a wider spectrum of environments, thus enabling them to have greater fitness  
116 in a multitude of novel environments.

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118 Here we investigate two of the above mentioned mechanisms for improved fitness in novel  
119 environments. We use replicate *E. coli* populations previously selected under unpredictable,  
120 complex environmental fluctuations for ~170 generations (Karve *et al.*, 2015). We test whether  
121 the phenotypic variation of the selected populations, in terms of usage of 94 substrates, have  
122 sufficiently diverged from the controls or not. We also count the number of progenies produced  
123 by individual bacterial cells, to ascertain whether population-based resistance has evolved in our  
124 selected populations. We find that our selected populations retain the fitness advantage even at

125 the level of individual cells. However, there was no evidence of evolution of either increased  
126 phenotypic diversity or population-based resistance. Thus we can say that environmental  
127 fluctuations do not lead to increased variation, at least in the short time scale.

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## 163 **MATERIALS AND METHODS**

### 164 **Selection under constant and fluctuating environments**

165 In this study, we used three replicate populations (henceforth F populations) of *E. coli* (strain  
166 NCIM 5547) that had been previously selected in unpredictably fluctuating, complex stressful  
167 environments. During the process of selection, the F populations were subjected to stressful  
168 combinations of salt, hydrogen peroxide and pH that changed unpredictably every 24 hrs. We  
169 also maintained corresponding controls (henceforth S populations) in the form of three replicate  
170 *E. coli* populations that were passaged in Nutrient Broth (see S1 for composition). After 30 days  
171 of selection (~170 generations) these S and F populations were stored as glycerol stocks at -  
172 80°C. The details of the maintenance regime for both the F and the S populations have been  
173 mentioned elsewhere (Karve *et al.*, 2015).

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### 175 **Fitness of the individual bacteria in novel environments**

176 To estimate the fitness of individual bacterium and characterize the possible heterogeneity within  
177 a population, we employed a slide-based observation technique (Lele *et al.*, 2011). Pilot studies  
178 were conducted to determine the sub-lethal concentrations for the four novel environments when  
179 the bacteria were grown on slides (see Table S2 for concentrations). The identity of these novel  
180 environments were chosen such that there are no known correlations between the mechanism of  
181 stress resistance to them and the three stresses used in the fluctuating selection (Karve *et al.*,  
182 2015). Glycerol stock for S or F population was revived overnight in 50 ml Nutrient Broth. This  
183 revived culture was used to flood the slide layered with nutrient agar (see S1 for composition)  
184 containing one of the novel environment. After the broth had dried off (~ 30 minutes at room

185 temperature under aseptic conditions), the agar surface was covered with a cover slip, excess  
186 agar outside the cover slip was removed with the help of a scalpel, and the sides were sealed with  
187 the mounting medium DPX (Di-n-butyl phthalate in xylene). The slide was then placed on the  
188 stage of a microscope (Primo Star<sup>TM</sup>, Zeiss, Jena, Germany) which in turn was placed at 37<sup>0</sup> C  
189 throughout the observation time.

190 A suitable field containing 6 to 20 single, well-spaced cells was focused under 100X  
191 magnification. For each cell in the field of view, we manually scored the time taken by the cell  
192 and its progenies to divide over a period of 240 minutes from the preparation of the slide i.e.  
193 from the time when broth was poured on the agar slide. Two trials were conducted for every  
194 replicate population of S and F in every novel environment ( $2 \times 6 \times 4 = 48$  trials). The yield of  
195 each cell was estimated as the number of progenies produced by the cell at the end of 240  
196 minutes. We also measured the ‘lag’ as the time taken for the first division. Since the cells were  
197 not synchronized, the lag estimate is likely to be associated with some amount of error. However,  
198 there is no reason to believe that this would affect S and F populations differentially. Moreover,  
199 since we measured substantial number of cells per population, such errors arising due to lack of  
200 synchronicity should be further ameliorated.

201 The yield and lag data were analyzed separately using mixed model ANOVA with novel assay  
202 environment (4 levels: Cobalt, Zinc, Norfloxacin and Streptomycin) and selection (2 levels: S  
203 and F) as fixed factors and replication (3 levels, nested within selection) and trial (2 levels,  
204 nested in assay environment  $\times$  selection  $\times$  replication) as random factors.

205 We also performed the individual mixed model ANOVAs for each of the novel assay  
206 environments. For this set of analysis, selection (2 levels: S and F) was treated as a fixed factor

207 and replication (3 levels, nested within selection) and trial (2 levels, nested in selection ×  
208 replication) as random factors. For the control of family-wise error rate, we used sequential  
209 Holm-Šidák correction of the  $p$  values (Abdi, 2010). All ANOVAs in this study were performed  
210 on STATISTICA v5 (Statsoft Inc., Tulsa, OK, USA).

211 To estimate the effect size of the differences between the means, we computed Cohen's  $d$   
212 (Cohen, 1988) using the freeware Effect Size Generator (Deville, 2004). The effect sizes were  
213 interpreted as small, medium and large for  $0.2 < d < 0.5$ ,  $0.5 < d < 0.8$  and  $d > 0.8$  respectively  
214 (Cohen, 1988).

215

## 216 **Population based resistance in novel environments**

217 Most formal tests of outlier detection assume the underlying data to be normally distributed  
218 (Barnett & Lewis, 1978). However, since our yield data did not meet this assumption, we used  
219 plots of the cumulative yield percentage to check for outliers. For this, we computed the  
220 percentage contribution of each parental bacteria to the final yield, arranged the values from both  
221 trials in ascending order and plotted the cumulative percentage yield against the percentage of  
222 the parental cells. In this plot, any cell(s) with disproportionate contribution to the overall yield  
223 can be easily identified by a sharp upward inflection towards the right of the graph.

224

## 225 **Assay for phenotypic variation**

226 We assayed the phenotypic variation in the population using GEN III MicroPlate™ (Catalog no.  
227 1030 Biolog, Hayward, CA, USA). Each of these plates contains 94 separate substrates of which

228 71 can be utilized as carbon sources while 23 can act as growth inhibitors. The presence or  
229 absence of growth is indicated with the help of tetrazolium redox dye where intensity of purple  
230 color is proportional to the amount of growth.

231 From each of the F and S populations, we obtained 8 clones by streaking the glycerol stock on a  
232 Nutrient Agar plate and incubating overnight at 37<sup>0</sup>C. Thus a total of 48 clones were isolated  
233 over the three S and three F populations. Every clone was then characterized for the 94 different  
234 phenotypes on the Biolog plate using standard protocol (for detailed methods, see S3). An  
235 ancestral clone was processed in the same way to obtain the ancestral phenotypic profile.

236 Following a previous study (Cooper & Lenski, 2000), we measured absorbance of the plates at  
237 590 nm using a microplate reader (SynergyHT BioTek, Winooski, VT, USA). For the 23 wells  
238 with inhibitory compounds, considering the recommendations of the product manual, we scored  
239 optical densities that were 50% or more of the corresponding positive control as 1 (i.e. no  
240 inhibition) and others as 0 (inhibition). Similarly, for the 71 wells with substrate utilization test,  
241 optical density that was  $\geq 200\%$  of the corresponding negative control was scored as 1 (i.e.  
242 utilized) while the others were scored as 0 (i.e. not-utilized). These binary scores were then used  
243 to determine standing phenotypic variation as well as the differences from the ancestral  
244 phenotypic profile. 33 phenotypes showed no variation in S and F (i.e. all individuals in S and F  
245 were either 0 or 1) and were ignored. For estimating standing phenotypic variation over the  
246 remaining 61 phenotypes, we computed the sum score of every replicate population over the  
247 eight clones. These values, ranging from 0 to 8, denote the variation within every population for  
248 that phenotype. It should be noted here that in some of the 94 substrates, absence of growth (i.e.  
249 0) was the dominant phenotype while for the other substrates, the presence of growth (i.e. 1) was  
250 the dominant one. We were not interested in the qualitative nature of the phenotype (1 or 0) and

251 wanted to analyze the variation over the entire set of 94 phenotypes. Therefore, we mapped  
252 phenotypic variation values of 5, 6, 7 and 8 to 3, 2, 1 and 0 respectively. In other words, a  
253 population in which three clones showed no growth (i.e. 3 zero values) and five clones showed  
254 growth (i.e. 5 values of 1), was deemed to have the same phenotypic variation for a given  
255 phenotype as a population which had five non-growers and three growers for a different  
256 phenotype. These mappings work only across phenotypes and fail if there are differences  
257 between the three replicates of S or F for the same phenotype. However, only three such cases  
258 were found in S populations and none at all in the F populations. The interpretations of our  
259 statistical analysis did not change with or without these points and hence we have retained these  
260 three data points. The phenotypic variations were then analyzed by a two way ANOVA with  
261 phenotype (61 levels) and selection (2 levels: S and F) as fixed factors.

262 For estimating the phenotypic divergence from the ancestor, we recorded the number of clones  
263 displaying phenotype that was different from the ancestral one, for all the F and S populations.  
264 The number of differences for each phenotype was then analyzed using a two way ANOVA with  
265 phenotype (61 levels for Phenotypes) and selection (2 levels: S and F) as fixed factors.

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273 **RESULTS**

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275 **Fitness of individual bacterial cells**

276 When pooled over all the novel environments, individuals from F populations displayed  
277 significantly lower lag time (Fig 1A) and higher yield (Fig 1B) than individuals from S  
278 populations, with medium and high effect sizes respectively (Table 1). There was a significant  
279 effect of the novel environment in both cases ( $F_{3, 12} = 66.75$ ,  $p < 0.001$  for lag and  $F_{3, 12} = 88.93$ ,  
280  $p < 0.001$  for yield) indicating that the difference in the fitness varies across different novel  
281 environments. This is intuitive as all the environments are not expected to affect fitness similarly.  
282 When analyzed separately for each novel environment, F populations had significantly and  
283 marginally significantly lower lag time in cobalt and streptomycin respectively (Table S4, Fig  
284 1A) and significantly higher yield in cobalt, streptomycin and norfloxacin (Table S4, Fig 1B).  
285 The effect sizes were large in all these cases (Table S4). It is important to note that in all the four  
286 novel environments, F populations showed lower lag time and higher yield compared to S  
287 populations. Overall, these results demonstrate the growth advantage for individuals of F  
288 populations in the four novel environments, corroborating the population level outcomes  
289 observed in an earlier study (Karve *et al.*, 2015).

290

291 **Population-based resistance**

292 Inspection of the data suggested that there were no individual cells whose progeny contributed  
293 disproportionately to the final population size. This can also be seen from the plot of the  
294 cumulative percentage yield of the cells, where the F populations showed a linear trend in three  
295 out of the four novel environments (Fig 2). Only in zinc, there was a small departure from the

296 linearity (Fig 2D). However, even then ~20% of the cells contributing to ~40-60% of the  
297 observed yield and hence, there was nothing to suggest the presence of a small number of  
298 outliers that contributed disproportionately to the growth. Interestingly, zinc was the only novel  
299 environment where F populations did not display a fitness advantage in terms of yield or lag (see  
300 Discussion), thus ruling out the possibility of a few individuals conferring fitness advantage to  
301 the entire population.

302

### 303 **Phenotypic Variation**

304 For 33 out of the 94 substrates tested, no variation was found i.e. all the 48 clones of S and F  
305 gave the same phenotype. In the remaining 61 substrates, at least 1 out of the 48 clones (8 clones  
306 each for three S and three F populations) gave a different phenotype. ANOVA on the phenotypic  
307 distances showed a significant main effect of phenotype ( $F_{60, 244} = 3.69, p \ll 0.001$ ) suggesting  
308 some phenotypes harbored more variation than others. This is intuitive as one does not expect  
309 similar number of variation for 61 traits over six populations. However, more crucially, there  
310 was no significant difference for the phenotypic variation across S and F populations, with a low  
311 effect size for the difference (Table 1 Fig 3A). Thus, we conclude that there was no evidence of  
312 an increased phenotypic variation in the F populations.

313 61 out of 94 phenotypes (not the same 61 as above though) showed at least one clone that was  
314 phenotypically different from the ancestor. Although, averaged over the 61 phenotypes, the S  
315 populations showed greater divergence which was marginally statistically significant (Fig 3B)  
316 the corresponding effect size of the difference was low (Table 1). More crucially, there was no  
317 phenotype for which all S or F populations were different from the ancestor. Barring two cases,

318 no consistent pattern was observed in terms of acquiring or losing a phenotypic trait. 43 out of all  
319 the 48 clones tested acquired the ability to utilize methyl pyruvate while 39 became capable of  
320 utilizing  $\beta$ -methyl- D-glucoside. Although prior studies indicate that the ability to catabolize  
321 methyl pyruvate (Timonen *et al.*, 1998) and  $\beta$ -methyl- D-glucoside (Perkins & Nicholson, 2008)  
322 often evolves under different kinds of stresses, the reason for the same remains unknown. Since,  
323 both S and F populations acquired the ability to utilize these compounds it is possible that there  
324 is some fitness advantage of these two phenotypes in nutrient broth. Crucially, there were no  
325 clear patterns in terms of phenotypic divergence from the ancestor, indicating that the variation  
326 accumulated is likely to be either neutral or have very weak effect on fitness. Apart from one  
327 replicate population of S in which all the individuals tested had lost the ability to utilize D-  
328 raffinose and pectin as a carbon source, there was not a single population in S or F in which all  
329 eight individuals had diverged from the ancestor. This suggests that the observed phenotypic  
330 variation is unlikely to be a result of a strong and /or directional selection pressure on one of the  
331 phenotypes. The divergence from ancestral phenotype varied significantly across different  
332 phenotypes ( $F_{60, 244} = 18.82, p \ll 0.001$ ) with a significant interaction with selection ( $F_{60, 244} =$   
333  $2.98, p \ll 0.001$ ). Both these results are intuitive since one neither expects similar levels of  
334 divergence over 61 substrates nor similar patterns of divergence in S and F populations.

335 **DISCUSSION**

336 **Higher fitness of individuals in novel environments**

337 For all the four novel environments, the lag times were lower for F populations and yields were  
338 higher (Fig 1) although the differences were not statistically significant for each comparison  
339 (Table S4). This corroborates similar observations at the population level in a previous study  
340 (Karve *et al.*, 2015).

341 Increased fitness in multiple novel environments can come about in at least two major ways: an  
342 increased rate of generating new variation or the existence of larger amount of standing  
343 variation. If the first case were true, then one would not expect the progenies of all individuals of  
344 the F populations to acquire the favorable mutations at the same time in a novel environment. If  
345 the F populations had increased standing variation which was contributing to their enhanced  
346 fitness under novel environments, then again one would expect that most individuals would fail  
347 to grow and the progenies of only few individuals would primarily contribute to the final  
348 population size. However, we found no outliers in terms of contributions to the final size of the  
349 population (Fig 2) which suggests that whatever the mechanism that had evolved, was  
350 benefitting all the existing F individuals similarly. This observation does not fit with either  
351 increased rate of generation of variation or increased standing genetic variation.

352

353 **No evidence for population-based resistance**

354 When the magnitude and direction of selection fluctuates continuously, traits that are favorable  
355 under one set of conditions, might become neutral or even deleterious when the environment  
356 changes. This can lead to a scenario where a population is continuously changing with each shift

357 in environment, without really evolving to greater fitness. One way by which a population can  
358 escape such a stasis is through the evolution of cooperation which allows subsets of the  
359 population to specialize in countering particular stresses and then confer resistance to the  
360 population as a whole (West *et al.*, 2007). For example, it has been shown that in populations of  
361 the bacteria *Pseudomonas aeruginosa*, the proportion of individuals that synthesize the iron-  
362 scavenging siderophore pyoverdinin, changes based on the kind of competition and genetic  
363 relatedness (Griffin *et al.*, 2004). Similarly, when *E. coli* populations are challenged with  
364 antibiotics, a very small percentage (0.1 - 1%) of the individuals secrete excess amounts of indole  
365 to the external environment, which then allows the entire population to become antibiotic  
366 resistant (Lee *et al.*, 2010). Since only a small fraction of the population needs to evolve the  
367 resistant mechanism for a given stress, in principle, this mechanism allows different subsets of  
368 the population to evolve resistance to different stresses. This should increase the population level  
369 variation in terms of the ability to resist diverse stresses, and hence increase fitness in different  
370 kinds of novel environments. Given that antibiotics were among the novel environments that we  
371 studied, population-based resistance was a possible explanation for the fitness advantages of F  
372 populations. However, we did not find any outliers in terms of the yield and, except in the case  
373 of zinc, all the plots of cumulative yield were linear (Fig 2). Even in the case of zinc, where there  
374 was a slight departure from linearity, at the point of inflection, ~20% of the parents contribute to  
375 ~40-60% of the yield. Overall, the conclusions are unambiguous, the observed increase in yield  
376 of the F populations were not attributable to a small fraction of the population.

377 The above result could have arisen in at least two other ways. It was possible that the F  
378 populations do exhibit population-based resistance, but we had managed to sample only those  
379 bacteria that conferred resistance to the population. The chances of such an event happening are

380 probably negligible since, as stated already, the fraction of bacteria that confer the population-  
381 wide resistance is typically very low (Lee *et al.*, 2010). As we had sampled around 12- 40  
382 bacteria out of  $\sim 2 \times 10^8$  (over two trials) for each F population, it is highly unlikely that only  
383 individuals with altruistic capacities were sampled. In fact, the second possibility was far more  
384 likely, namely that we had sampled only those bacteria that did not confer any resistance to the  
385 population. In principle, this could also explain the absence of outliers in the F populations in  
386 terms of overall yield. However, in that case, we could not have observed an increase in the yield  
387 when compared to the S populations. Since the F populations did show a significantly larger  
388 yield compared to S populations (Fig 1B), we conclude that whatever mechanism was  
389 responsible for it, was not present only in a small number of individuals.

390 There can be multiple, non-exclusive reasons for which population-based resistance failed to  
391 evolve in our F populations. Our F populations were sub-cultured every 24 hours with 1/50 of  
392 the existing population forming an inoculum for the next generation (Karve *et al.*, 2015). It is  
393 difficult for population-based resistance to evolve in such a system due to a high chance of losing  
394 the resistant cells (which are in very low frequency) during each sub-culture. Moreover, it is  
395 known that when the environment changes, the production of the chemical that benefits the  
396 whole population can be costly for the producer cell (Lee *et al.*, 2010). Thus, in our F  
397 populations, there could have been a strong selection against the resistant cells, each time the  
398 environment changed. Taken together, perhaps it is not surprising that population-based  
399 resistance did not evolve in our F populations.

400

401

## 402 **Fluctuating selection does not increase standing variation**

403 Populations with greater standing variation are expected to respond faster to selection pressures  
404 compared to those with increased mutation rates. This is because with standing variation, the  
405 population need not wait for a beneficial mutation and such mutations are typically at a slightly  
406 higher frequency than those that arise *de novo* after exposure to the selection pressure (Barrett &  
407 Schluter, 2008). Furthermore, theoretical studies show that fluctuating environments are  
408 expected to promote standing variation in the populations (Gillespie & Turelli, 1989; Turelli &  
409 Barton, 2004). Taken together, the greater fitness of the F populations in novel environments can  
410 be potentially explained if such populations have greater standing variation. However, it is  
411 difficult to visualize how large standing variation can be maintained when the direction of  
412 selection is changing very often (Via & Lande, 1987). One way out of this problem is contextual  
413 neutrality, i.e. the assumption that at least some genetic changes are neutral in some  
414 environments (thus escaping selection) but affect fitness in other environments (thus contributing  
415 to standing genetic variance)(Wagner, 2005a). Thus, a population with a greater “neutral space”  
416 (i.e. contextually neutral variation) would be expected to have greater fitness across novel  
417 environments (Wagner, 2005b). Although some studies have directly measured genetic diversity  
418 through quantification of the number of mutations present (Coffey & Vignuzzi, 2011), it is hard  
419 to determine how much of that diversity is functionally relevant. This is because, practically  
420 speaking, it is difficult to ascertain from the sequence data, whether a particular genomic  
421 mutation is deleterious, neutral or contextually neutral. Therefore, we favored a direct  
422 measurement of the phenotypic variation in the populations, through their ability to grow on 94  
423 different conditions on the Biolog GEN III MicroPlate™ plate (Cooper, 2002). This way, we

424 quantify those variations that can cause an observable change at the phenotypic level and hence,  
425 are functionally important.

426

427 Our results suggest that selection for unpredictable fluctuations did not increase the phenotypic  
428 variation in F populations. If anything, the mean phenotypic variation was slightly larger for the  
429 S populations (Fig 3A), although the difference was not statistically significant. This is  
430 consistent with a previous study on viruses demonstrating that genetic diversity (as measured by  
431 genomic mutations) is larger in populations that experience a steady environment as opposed to  
432 those facing fluctuating ones (Coffey & Vignuzzi, 2011). Our results also agree with a previous  
433 observation that constant selection environments lead to increase in the genetic variance for  
434 fitness in novel environments (Travisano *et al.*, 1995). In terms of the phenotypic divergence  
435 from the ancestors, we found no consistent differences or reversal of phenotypes that were  
436 specific to the F or S populations (Fig 3B).

437 There might be several reasons for which phenotypic variation did not increase in the F  
438 populations. The duration of selection (~170 generations) might have been too less to lead to a  
439 significant divergence in terms of phenotypic variation. Moreover, the fact that the environment  
440 (and hence the selection pressure) changed every ~6 generations, might have caused a much  
441 stronger selection pressure that prevented maintenance of phenotypic variation. One way by  
442 which standing variation can be increased even in the face of changing environments, is through  
443 increased mutation rates (Ishii *et al.*, 1989). However, since the mutation rates of the F  
444 populations did not evolve to be significantly larger than the S populations (Karve *et al.*, 2015),  
445 this route was closed to the selected populations. It is important to note here that we only scored

446 the presence or absence of phenotypes, a process that is biased towards catching large  
447 phenotypic differences. In principle, one can also think of variations which affect the rate at  
448 which the substrates are metabolized or the intensity of the effect of stress substrates on the  
449 bacterial cells. However, quantifying such effects would require replicate measurements at the  
450 level of single clones and increased number of replicate clones due to the inherent variation in  
451 the metabolic rates of the cells, and hence was beyond the scope of this work.

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454 **CONCLUSIONS**

455 Bacterial populations exposed to randomly fluctuating environments evolve to have greater  
456 fitness in novel stresses (Ketola *et al.*, 2013; Karve *et al.*, 2015). However, this is not attributable  
457 to an increase in standing variation, nor evolution of population-based resistance, nor an increase  
458 in the rate of generation of variation through mutations (Karve *et al.*, 2015). This suggests that  
459 the greater fitness in novel environments is perhaps due to direct individual-level selection on  
460 broad-spectrum stress resistance traits like change in membrane structure (Viveiros *et al.*, 2007),  
461 multi-drug efflux pumps (Nikaido & Pagès, 2012) etc. This observation is consistent with a  
462 previous result that the efflux activity of the F populations had increased significantly (Karve *et*  
463 *al.*, 2015). If the evolved increase in fitness were due to mutations or standing genetic variations,  
464 then there are a large number of ways available for the bacteria to evolve. However, the number  
465 of individual-level broad-spectrum resistance mechanisms is relatively small and typically well-  
466 studied, which at least gives some hopes in terms of developing containment strategies against  
467 such mechanisms. Moreover, most theoretical studies on evolutionary effects of fluctuating  
468 environments seek to model changes in mutation rates and standing variation (Leigh, 1970; Ishii  
469 *et al.*, 1989; Taddei *et al.*, 1997). Our results suggest that such studies have perhaps failed to  
470 consider the critical mechanism that enables organisms to adapt to such situations in nature and a  
471 new class of theoretical modeling is needed to investigate this issue.

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474 **ACKNOWLEDGEMENTS**

475 We thank Milind Watve for helpful discussions and Madhur Mangalam for help with the single-  
476 cell assay. SK was supported by a Senior Research Fellowship from Council of Scientific and  
477 Industrial Research, Govt. of India. This study was supported by a research grant from  
478 Department of Biotechnology, Government of India and internal funding from Indian Institute of  
479 Science Education and Research, Pune.

Assay	Means		ANOVA F (df effect, df error)	ANOVA p	Effect size $\pm 95\%CI$	Interpretation
	S	F				
Lag	147.66	118.69	19.20 (1,4)	0.012	0.50 $\pm$ 0.18	Medium
Yield	3.89	6.26	92.94 (1,4)	0.0006	0.94 $\pm$ 0.19	Large
Phenotypic variation	0.61	0.51	1.51 (1,244)	0.22	0.11 $\pm$ 0.2	Small
Phenotypic divergence from ancestor	0.97	0.79	3.98 (1,244)	0.047	0.1 $\pm$ 0.2	Small

480

481 **Table 1: Summary of the main effects of selection in the pooled ANOVAs.**

482 Effect size was measured as Cohen's  $d$  statistic and interpreted as small, medium and large for

483  $0.2 < d < 0.5$ ,  $0.5 < d < 0.8$  and  $d > 0.8$  respectively.

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485 **SUPPORTING INFORMATION**

486 **S1** – Composition of Nutrient Agar and Nutrient Broth

487 **Table S2** – Novel Environments used for estimating fitness at the individual level

488 **S2** – Protocol for using the Biolog plates

489 **Table S4** – Summary of the main effects of selection in the ANOVAs under individual

490 environments

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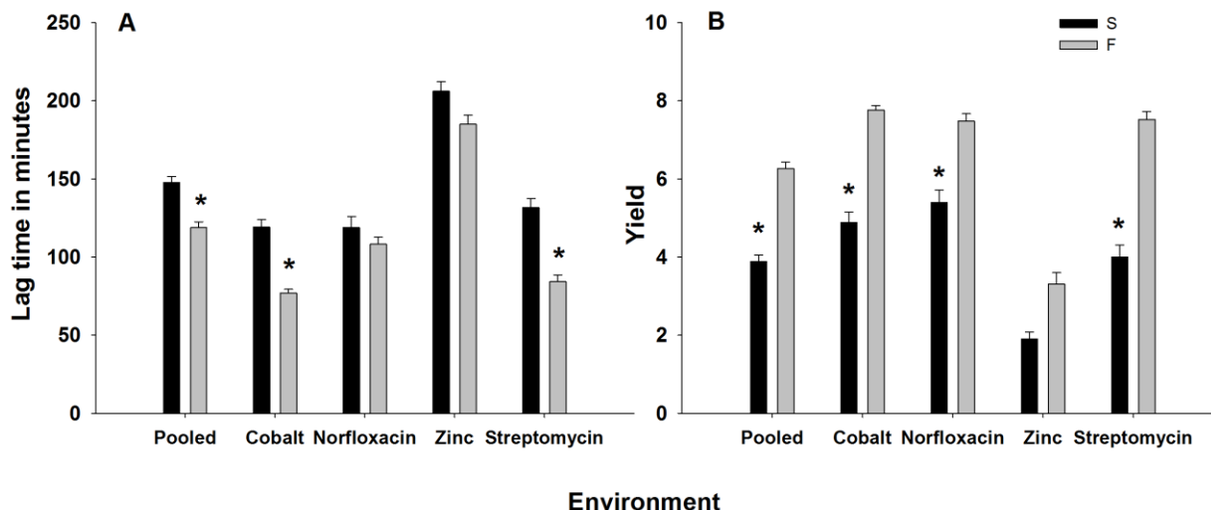
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618 **FIGURES**

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621 **Fig 1: Fitness of individual bacterial cells. A.** Mean ( $\pm$ SE) lag time is significantly lower for F

622 populations than S populations when pooled over four novel environments. When compared

623 separately for each novel environment, F populations show significantly lower lag time in cobalt

624 and streptomycin and similar lag time in norfloxacin and zinc. **B.** Mean ( $\pm$ SE) yield is

625 significantly higher for F populations than S populations when pooled over four novel

626 environments. When compared separately for each novel environment, F populations show

627 significantly higher yield for cobalt, norfloxacin and streptomycin and similar yield for zinc.

628 \* denotes  $p < 0.05$  (after Holm-Šidák correction in the case of comparisons under individual

629 environments).

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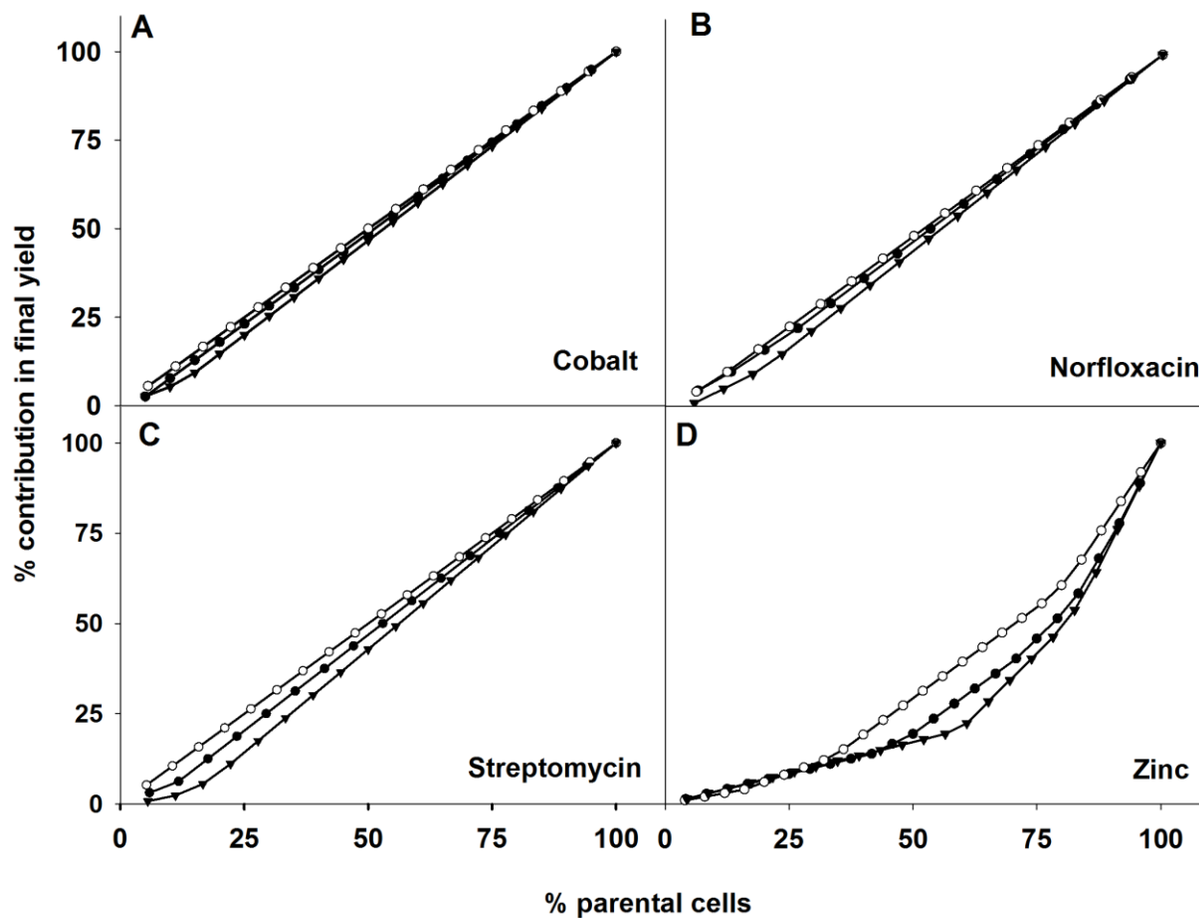
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639 **Fig 2: Population-based resistance in F populations.** Contribution of each parent cell to the

640 total yield is plotted for three replicate F populations in four novel environments. Each line in a

641 figure stands for a replicate population of F. **A.** Cobalt, **B.** Norfloxacin, **C.** Streptomycin, **D.**

642 Zinc. No individual cells contribute disproportionately to the total yield.

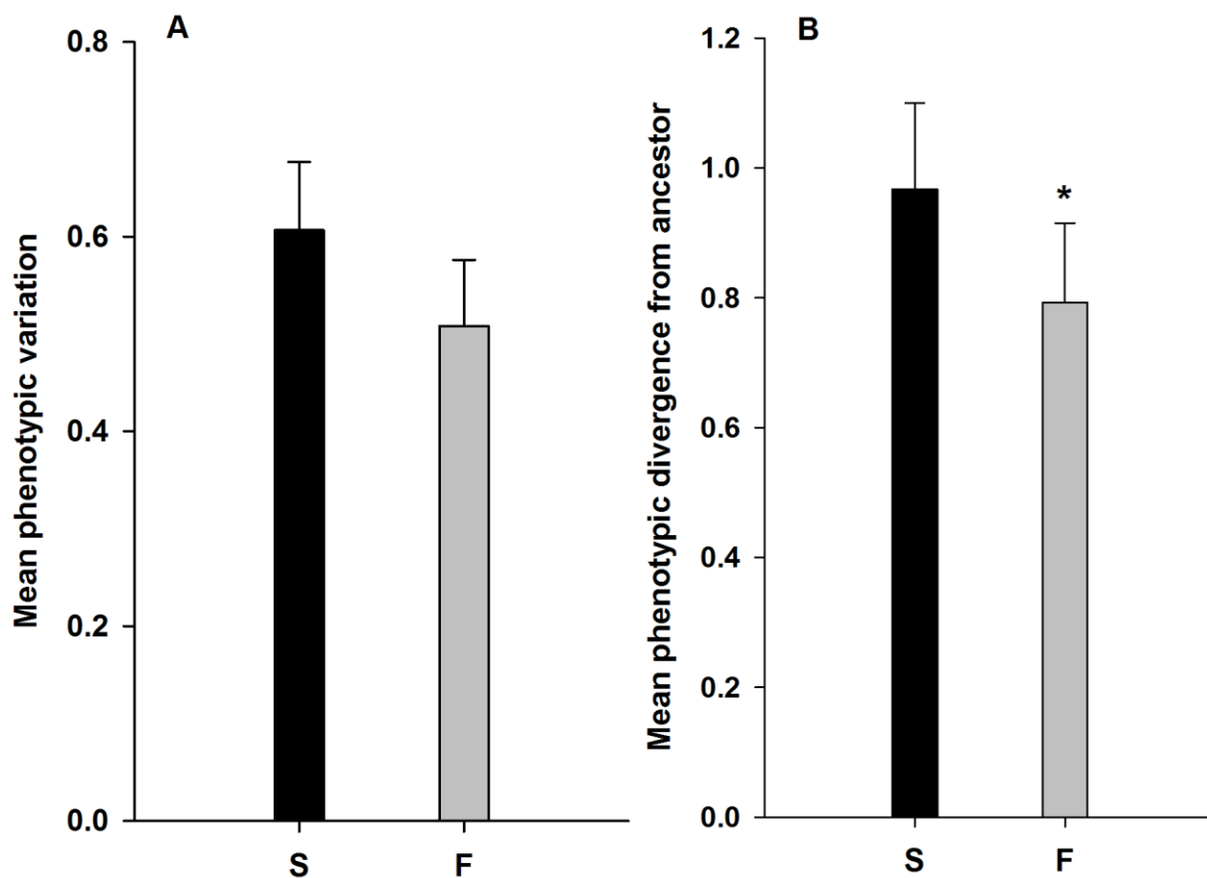
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650 **Fig 3: Phenotypic variation and divergence from ancestors.** A. Mean ( $\pm$ SE) phenotypic  
651 variation for S and F populations. B. Mean ( $\pm$ SE) phenotypic divergence from the ancestors. The  
652 S populations show slightly higher variation and divergence albeit with small effect sizes. \*  
653 denotes  $p < 0.05$

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## **SUPPLEMENTARY ONLINE MATERIAL**

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675 **S1 – Composition of Nutrient Agar and Nutrient Broth**

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677 Composition of Nutrient Agar (NA):

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Ingredients	g/L
Peptic digest of animal tissue	5.00
Sodium chloride	5.00
Beef extract	1.50
Yeast extract	1.50
Agar	20.00

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680 Final pH (at 25<sup>0</sup>C) 7.4 ± 0.2

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682 The composition for Nutrient Broth (NB) is the same as above except the absence of agar.

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684 For the slide based technique, we used 12 g / L of Agar.

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691 **Table S2 – Novel Environments used for estimating fitness at the individual level**

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<b>Assay Environment</b>	<b>Concentration</b>
Cobalt chloride	28.5 mg%
Zinc sulfate	120 mg%
Streptomycin	0.0065mg%
Norfloxacin	0.0032 mg%

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710 **S3 – Protocol for using the Biolog plates**

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712 We used GEN III MicroPlate™ along with inoculating fluid A (IF-A) for estimating the  
713 phenotypic variation. Both plates and inoculating fluid were stored at 4<sup>0</sup>C and thawed at room  
714 temperature before use.

715 A part of glycerol stock was streaked on nutrient agar plate for every replicate population of S  
716 and F. The plates were incubated at 37<sup>0</sup>C overnight. 8 isolated clones of comparable sizes were  
717 selected for every population and inoculated into the separate inoculation fluid tube. The  
718 transmittance was in the range of 95% to 98% for the selected clones. 100 µl of this well mixed  
719 inoculation fluid was used to inoculate the GEN III plate. The plates were incubated at 37<sup>0</sup>C for  
720 24 hours after which they were stored at 4<sup>0</sup>C for another day, during which time we measured  
721 optical density for all the 48 plates at 590 nm (Cooper & Lenski, 2000) using a microplate reader  
722 (SynergyHT Biotek, Winooski, VT, USA).

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733 **Table S4 – Summary of the main effects of selection in the ANOVAs under individual**  
 734 **environments.**

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Fitness Measure	Environment	Means		ANOVA F(1,4)	Holm- Šidák corrected p values	Effect Size±95% CI	Interpretation
		S	F				
Yield	Cobalt	4.877	7.759	43.33	0.008	1.79±0.43	Large
	Norfloxacin	5.396	7.479	40.77	0.006	1.15±0.42	Large
	Zinc	1.897	3.306	6.47	0.064	0.69±0.34	Medium
	Streptomycin	4.000	7.519	99.53	0.002	1.77±0.42	Large
Lag	Cobalt	119.070	77.000	34.53	0.017	1.46±0.36	Large
	Norfloxacin	118.854	108.208	0.99	0.375	0.26±0.4	Small
	Zinc	206.059	185.056	1.62	0.47	0.42±0.33	Small
	Streptomycin	131.667	84.278	14.93	0.053	1.22±0.4	Large

736

737 This table shows yield and lag measurements for individual cells under four novel environments.

738 Effect size was measured as Cohen's *d* statistic and interpreted as small, medium and large for

739  $0.2 < d < 0.5$ ,  $0.5 < d < 0.8$  and  $d > 0.8$  respectively.

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741 **REFERENCE**

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