

1 Non-antibiotic pharmaceuticals exhibit toxicity
2 against *Escherichia coli* at environmentally
3 relevant concentrations with no evolution of
4 cross-resistance to antibiotics

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15

16 **Abstract**

17 Antimicrobial resistance can arise in the natural environment via prolonged exposure
18 to the effluent surrounding manufacturing facilities. These facilities also produce
19 non-antibiotic pharmaceuticals, and the effect of these on the surrounding microbial
20 communities is less clear; whether they have inherent toxicity, or whether long-term
21 exposure might select for cross-resistance to antibiotics. To this end, we screened
22 four non-antibiotic pharmaceuticals (acetaminophen, ibuprofen, propranolol, met-
23 formin) and titanium dioxide for toxicity against *Escherichia coli* K-12 MG1655

24 and conducted a 30 day selection experiment to assess the effect of long-term expo-
25 sure. All compounds reduced the maximum optical density reached by *E. coli* at
26 a range of concentrations including one of environmental relevance, with transcrip-
27 tome analysis identifying upregulated genes related to stress response and multidrug
28 efflux in response to ibuprofen treatment. The non-antibiotic pharmaceuticals did not
29 select for significant genetic changes following a 30 day exposure, and no evidence
30 of selection for cross-resistance to antibiotics was observed for population evolved in
31 the presence of ibuprofen in spite of the differential gene expression after exposure
32 to this compound. This work suggests that these non-antibiotic pharmaceuticals, at
33 environmental concentrations, do not select for cross-resistance to antibiotics in *E.*
34 *coli*.

35 Introduction

36 Antimicrobial resistance (AMR) is a leading public health concern with global but
37 unequal causes and consequences [1; 2; 3]. There is concern about the extent to
38 which AMR is driven by effluent from pharmaceutical production and wastewater
39 treatment facilities that can enter local ground and surface water [4; 5; 6; 7; 8]. This
40 long-term supply of wastewater is known to have considerable effects on the local mi-
41 crobial community [8; 9; 10]. Metagenomic studies have for example identified AMR
42 genes in water bodies close to pharmaceutical plants in countries including India and
43 Croatia [6; 11; 12]. High levels of antibiotics and antibiotic resistance genes have
44 also been measured around production facilities in Lahore, Pakistan [13]. Whilst the
45 evolution of bacteria in response to antibiotics is well understood [14; 15; 16], less
46 is known about the possible impact of long-term exposure to non-antibiotic phar-
47 maceuticals. Importantly, manufacturing facilities typically produce more than one
48 chemical entity, meaning waste from these sites can contain a range of biologically
49 active chemicals including both antibiotics and non-antibiotic compounds [17].

50 Like antibiotics, non-antibiotic pharmaceuticals are ubiquitous contaminants that
51 have been detected in water bodies globally following inadvertent release into the
52 environment [5; 18; 19], and efforts have been made to begin characterising their ef-
53 fects on different bacterial species. Compounds including vasodilators and selective
54 norepinephrine re-uptake inhibitors have, at various concentrations, antimicrobial
55 activity against *Escherichia coli*, *Staphylococcus aureus*, *Pseudomonas aeruginosa*,
56 and *Candida albicans* [20]. A screen of non-antibiotic pharmaceuticals against 40
57 species of gut bacteria found over 200 human-targeted drugs that had a negative ef-
58 fect on the growth of at least one of the species tested [21]. Interestingly, the majority

59 of these compounds were found to be active against only a few strains, suggest-
60 ing that non-antibiotic pharmaceuticals may have strain-specific effects. Titanium
61 dioxide (TiO₂), found in a wide range of products from cosmetics to paints [22], has
62 been shown, at various concentrations, to have antibacterial effects on organisms
63 including *E. coli* [23; 24; 25], *P. aeruginosa* [23], *S. aureus* [23; 24; 25; 26], and
64 *Enterococcus faecalis*, amongst others [27; 28]. In many instances, the antimicrobial
65 properties of TiO₂ have been evaluated as a surface coating [29; 24] rather than a
66 suspension, the latter representing the form in which microbial communities would
67 be exposed to TiO₂ in waste effluent. The anti-inflammatory drug ibuprofen has
68 been demonstrated, using disc diffusion methods, to exhibit antimicrobial activ-
69 ity against species including *S. aureus* and *Bacillus subtilis*, whilst not against *E.*
70 *coli* or *P. aeruginosa* [30], further highlighting the species-specific activity of these
71 compounds.

72 There has also been recent interest in the potential of non-antibiotic pharmaceuti-
73 cals to influence the susceptibility of bacterial species to antibiotics. Compounds
74 including ibuprofen, diclofenac, and propranolol have been linked to enhanced up-
75 take of antibiotic resistance genes, possibly due to an increase in cell competency and
76 membrane permeability, and the promotion of conjugative plasmid transfer [31; 32].
77 The antiepileptic drug carbamazepine has also been shown to promote the transfer
78 of plasmid-encoded resistance genes via conjugation [33]. In contrast, experiments
79 in clinically relevant species suggest that metformin, used in the treatment of type
80 2 diabetes, and the non-steroidal anti-inflammatory drug benzydamine can promote
81 uptake of tetracyclines, thereby reversing a resistance phenotype in multidrug resis-
82 tant pathogens [34; 35], and antibiotic-non-antibiotic drug combinations have been
83 suggested as possible routes for treating infections caused by such species [36]. The
84 short- and long-term effects of exposure to these compounds on bacterial popula-
85 tions therefore warrant further study; whether they have intrinsic toxicity and, if so,
86 the mechanism of action and the likelihood that they could act as selection pressures
87 resulting in significant genetic changes. Importantly, discovery of the latter might
88 indicate a possible mechanism by which exposure to non-antibiotic pharmaceuticals
89 could select for cross-resistance to antibiotics. This is therefore an area of research
90 with important clinical ramifications.

91 Here, we undertook an investigation of the short- and long-term effects of a panel
92 of non-antibiotic compounds on *Escherichia coli* K-12 MG1655, with an emphasis
93 on testing at environmentally-relevant concentrations [37; 38; 39; 40; 41; 42; 43; 44]
94 to examine the potential for selecting for cross-resistance to antibiotics. The com-
95 pounds selected were TiO₂, acetaminophen (the active ingredient in the painkiller
96 paracetamol), ibuprofen (an anti-inflammatory), propranolol (a beta-blocker used

97 to treat heart conditions), and metformin (a medication for type 2 diabetes). These
98 compounds were all found to have a degree of toxicity against this strain of *E.*
99 *coli* at a range of concentrations including those of environmental relevance, with
100 transcriptome analysis identifying upregulated genes involved in stress response and
101 multidrug efflux during ibuprofen treatment. Through experimental evolution in the
102 presence of environmentally relevant concentrations of these compounds we found
103 evolved populations displayed decreased fitness relative to the ancestral lineage when
104 grown in the presence of the selection compound. However, analysis of hybrid as-
105 semblies of the evolved isolates found no single nucleotide polymorphisms (SNPs)
106 between independently evolved populations, and there was no change in minimum
107 inhibitory concentration (MIC) for a panel of antibiotics against isolates evolved in
108 the presence of ibuprofen compared to the ancestor. Together, this suggests that
109 the toxicity of the non-antibiotic pharmaceuticals does not exert a selection pressure
110 sufficiently strong enough to lead to the fixation of mutations under the conditions
111 tested, and with no observed selection for cross-resistance to antibiotics.

112 **Methods**

113 **Strains and growth conditions**

114 To measure potential toxicity of compounds, *E. coli* K-12 MG1655 was streaked from
115 a glycerol stock on to a Luria Bertani (LB) agar plate (E & O Laboratories Ltd),
116 incubated overnight at 37°C. A single colony was then used to inoculate 5 mL LB
117 (E & O Laboratories Ltd) in a 30 mL universal before overnight incubation at 37°C
118 with agitation. The overnight cultures were diluted to an optical density at 600 nm
119 (OD₆₀₀) of approximately 0.5 in LB. Serial dilutions of acetaminophen, ibuprofen,
120 titanium dioxide (TiO₂, in the form of nanoparticles), propranolol, and metformin
121 were prepared as per Table S1. These compounds were selected as both published
122 literature and preliminary investigations identified their presence in wastewater and
123 receiving water environments, and they are commonly used non-antibiotic phar-
124 maceuticals [5]. Additionally, TiO₂ is found in a wide range of products and has
125 suggested applications in water treatment [27; 45; 46]. Environmentally relevant con-
126 centrations were identified following a literature search and are provided in Table
127 S1. A solution of 99 μ L of LB + compound was added to each test well of a 96-well
128 plate, including an LB-only control, with 1 μ L of the dilute cell suspension then
129 added. Plates were incubated for 24 hours in a microplate reader (Tecan) at 37°C
130 with continuous double orbital shaking, with absorbance measurements (OD₆₀₀)

131 taken every 30 minutes in triplicate. To assess the growth kinetics of evolved popu-
132 lations, ten colonies were selected for incubation as representative of the population
133 and the kinetics monitored in a microplate reader as described previously, in the
134 presence of the compound to which they were exposure during the selection exper-
135 iment. Compounds were tested at 1x and 100x selection concentrations (Table S1)
136 to measure whether the evolved isolates would show improved growth compared to
137 the ancestral lineage when stressed with a higher concentration of the compound to
138 which they had been exposed during the selection experiment.

139 **Genome sequencing and bioinformatics**

140 Illumina short read sequencing of the ancestral and evolved isolates was performed
141 by MicrobesNG (UK). Long-read sequencing of the same isolates was performed us-
142 ing MinION sequencing (Oxford Nanopore Technologies, UK). Briefly, genomic DNA
143 was extracted from overnight cultures using the Monarch Genomic DNA Purifica-
144 tion Kit (New England Biolabs). DNA was quantified using a Qubit 4 fluorometer
145 (Invitrogen) and accompanying broad-range double stranded DNA assay kit (Invit-
146 rogen). Sequencing libraries were prepared using SQK-LSK109 ligation sequencing
147 kit and EXP-NBD114 native barcode expansion (Oxford Nanopore Technologies,
148 UK), as per manufacturer instructions. Long-read sequencing was performed on a
149 MinION sequencer using an R9.4.1 flow cell (Oxford Nanopore Technologies, UK).
150 Base calling was conducted using Guppy (v6.0.1). Reads were filtered using Filt-
151 long (v0.2.1) using a cut-off of 600000000 target bases and demultiplexed using qcat
152 (v1.1.0). Hybrid assemblies were then generated using Unicycler (v0.4.8-beta) in
153 bold mode. Panaroo (v1.2.10) was used to generate gene presence/absence and core
154 gene alignment files with the latter used to construct a maximum likelihood tree
155 with IQ-TREE (v2.2.0.3). The tree and gene presence/absence data were visualised
156 in Phandango [47] to look for differential gene presence patterns across the evolved
157 isolates. A custom ABRicate (v0.8) database was used to investigate the presence
158 and identity of the *ldrA* gene across the evolved isolates. The presence of SNPs
159 was analysed using snippy (v4.3.6). Potential movement of insertion sequence (IS)
160 elements was investigated using ISEScan (v1.7.2.3), ISFinder [48], and a custom
161 ABRicate (v0.8) database, with sequences interrogated in Unipro UGENE (v47.0).

162 Transcriptome sequencing

163 RNA sequencing was performed on *E. coli* grown in the presence and absence of
164 50 $\mu\text{g}/\text{mL}$ ibuprofen in triplicate. For the control (absence) replicates, an equiv-
165 alent volume of the ibuprofen solvent (ethanol) was added. For sample prepa-
166 ration, a single colony for each replicate was picked following overnight growth
167 on LB agar and added to 5 mL of LB broth (Sigma-Aldrich, UK). A 100 μL
168 suspension of each overnight culture was then transferred into 10 mL fresh LB
169 in the presence or absence of 50 $\mu\text{g}/\text{mL}$ ibuprofen, with cultures then incubated
170 at 37°C with agitation until an optical density at 600 nm (OD600) of approxi-
171 mately 0.9. A 1 mL sample was centrifuged for five minutes at 10,000 rpm (Eppen-
172 dorf MiniSpin F-45-12-11), resuspended in 1 mL phosphate buffered saline (PBS,
173 VWR), and this wash step repeated. The supernatant was aspirated and the pellet
174 frozen prior to processing and RNA sequencing by GENEWIZ from Azenta Life
175 Sciences (Frankfurt, Germany) using their standard RNA sequencing service. Dif-
176 ferential gene expression was quantified using Kallisto (v0.48.0) A long-read as-
177 sembly of the ancestral *E. coli*, annotated using Prokka (v1.14.6), was used as a
178 reference. The annotated assembly was processed using `genbank_to_kallisto.py`
179 (https://github.com/AnnaSyme/genbank_to_kallisto.py). GNU parallel [49] was
180 used for job parallelization. Differential gene expression was analyzed in Degust
181 (v4.1.1) with a false discovery rate threshold of $p < 0.05$ and an absolute log fold
182 change of at least 1.

183 Selection experiment

184 The ancestral *E. coli* isolate was streaked from a glycerol stock on to an LB plate
185 and incubated overnight at 37°C. A single colony was used to inoculate 5 mL nu-
186 trient broth (NB) (Sigma) in a 30 mL universal, with six independent biological
187 replicates per condition. Acetaminophen (5 ng/mL), ibuprofen (2 $\mu\text{g}/\text{mL}$), TiO_2
188 (1 $\mu\text{g}/\text{mL}$), propranolol (0.5 ng/mL), and metformin (0.5 ng/mL) were tested in-
189 dividually, including a NB-only control. Microcosms were incubated for 24 hours
190 at 37°C with agitation, before a 1% transfer of cell suspension into fresh media.
191 This 1% transfer was repeated every 24 hours for 30 days. After 30 days, the whole
192 population was centrifuged at 3600 rpm (Thermo Scientific Megafuge 40R TX-1000)
193 for five minutes, resuspended in 1 mL 50% glycerol, and stored at -80°C. To assess
194 whether the populations were experiencing short-term, reversible toxicity as a result
195 of compound exposure, ten colonies from each end-point population were selected
196 from a UTI ChromoSelect agar plate (Millipore) and used to inoculate 5 mL NB

197 only. The microcosms were incubated for 24 hours at 37°C with agitation, before a
198 1% transfer of cell suspension into fresh media every 24 hours for seven days. After
199 seven days, the whole population was centrifuged at 3600 rpm (Thermo Scientific
200 Megafuge 40R TX-1000) for five minutes, resuspended in 1 mL 50% glycerol, and
201 stored at -80°C.

202 **Minimum inhibitory concentration assay**

203 Stocks of ethidium bromide, ampicillin, ciprofloxacin, chloramphenicol, trimetho-
204 prim, and colistin were prepared to 1000 $\mu\text{g}/\text{mL}$. Cultures of the *E. coli* MG1655
205 ancestor and an *E. coli* ATCC 25922 control strain were prepared by inoculating 5
206 mL of LB with a single colony of bacteria and incubating with agitation at 37° for 18
207 hours. Iso-Sensitest broth (ISB) (Thermo Fisher Scientific) was then used through-
208 out the assay. The overnight culture was then diluted 1:100 and working stocks
209 of antibiotics prepared, both in ISB. U-bottom 96-well plates were set up so that
210 the cultures were incubated with no antibiotic and with 11 different concentrations
211 of antibiotic ranging from 0.008 to 8 $\mu\text{g}/\text{mL}$. The first column of the 96-well plate
212 contains the highest concentration of antibiotic and the 11th column contains the
213 lowest concentration, with the 12th column containing no antibiotic, and 50 μL of
214 the diluted cell suspension was added to all wells. Plates were incubated at 37° for
215 18 hours and examined for growth the next day. Results were only accepted if the
216 observed MIC for the ATCC 25922 strain was within one doubling dilution of the
217 expected result.

218 **Checkerboard minimum inhibitory concentration assay**

219 Cultures of the *E. coli* MG1655 ancestral lineage and one of the six end-point isolates
220 evolved in the presence of ibuprofen were prepared using a single colony inoculated
221 into LB broth before incubation overnight at 37° with agitation. Working stocks
222 of ethidium bromide, ampicillin, ciprofloxacin, chloramphenicol, trimethoprim, col-
223 istin, and ibuprofen were prepared at four times the highest final concentration
224 required by diluting in ISB. Overnight cultures were diluted 1:2000 in ISB. A 50
225 μL aliquot of ISB was added to all columns of a 96-well plate, and 50 μL working
226 ibuprofen stock added to all wells of columns one and two. Starting with column
227 two, the ibuprofen was serially diluted 1:2 across the plate up to and including col-
228 umn 11. A 50 μL aliquot of one antibiotic working stock was then added to all
229 wells of row A, the 1:2 dilution repeated down the plate up to and including row

230 G, and 50 μ L removed from column 11 and row G before adding cells to keep the
231 volume consistent. A 50 μ L sample of diluted overnight culture was then added to
232 each well and mixed gently before the plate was covered and incubated for 18 hours
233 at 37°static. Plates were read following incubation and the presence or absence of
234 growth noted.

235 **Statistical analyses**

236 Area under the curve measurements were calculated using `numpy.trapz` in Python
237 (v3.9.10). Significance testing was conducted using a one-way analysis of variance
238 (ANOVA).

239 **Results**

240 **Observed toxicity from pharmaceutical compounds against** 241 ***E. coli***

242 To first establish whether non-antibiotic pharmaceuticals can have observable toxi-
243 city, we screened a panel of compounds at a range of different concentrations (Table
244 S1) against *E. coli* K-12 MG1655 as a model organism. Acetaminophen, ibuprofen,
245 TiO₂, propranolol, and metformin were all found to have significant negative effects
246 on *E. coli* growth over a 24 hour incubation in comparison to the no-compound con-
247 trol at all concentrations tested ($p < 0.05$, one-way ANOVA, Fig. 1, Fig S1). The
248 effect was predominantly noted as a reduction in the maximum OD reached. With
249 the exception of TiO₂ (where the highest concentration of 100 μ g/ml had a larger
250 effect than other concentrations), altering the concentration of the compounds had
251 little effect on the resulting growth kinetics. The non-antibiotic pharmaceuticals
252 tested can therefore negatively impact growth of *E. coli* MG1655.

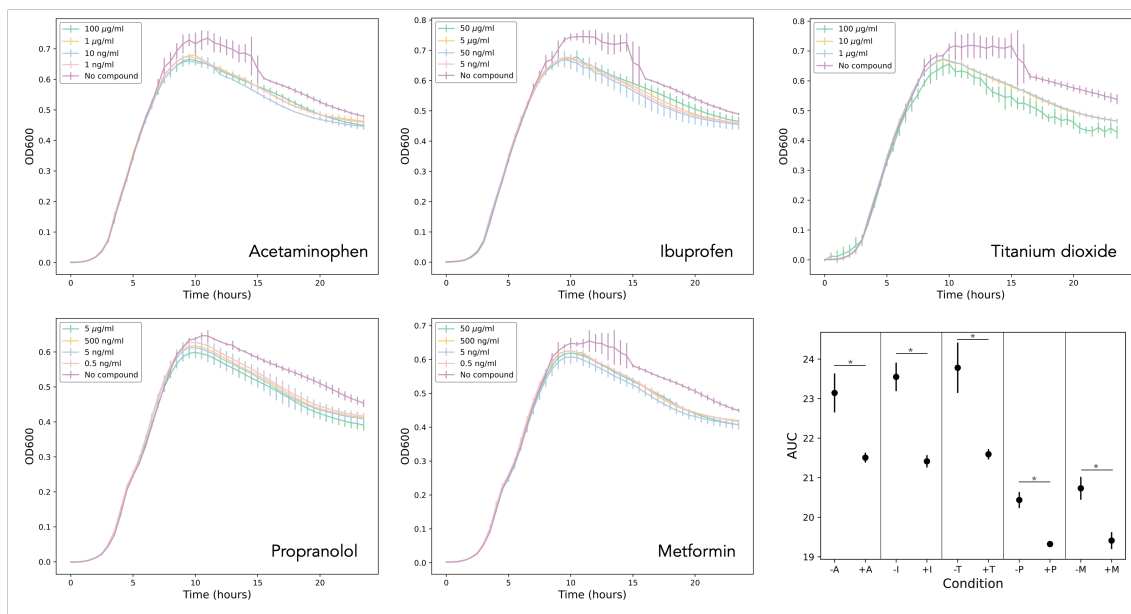


Fig. 1 Toxicity screen of acetaminophen (+A), ibuprofen (+I), TiO₂ (+T), metformin (+M), and propranolol (+P) at various concentrations against *E. coli* over a 24 incubation in a 96-well plate in a microplate reader. A no-compound control ('No compound', purple) was included for all screens. AUC values for the following concentrations are given against their representative no-compound control (-); 1 ng/mL acetaminophen, 5 ng/mL ibuprofen, 1 µg/mL TiO₂, 0.5 ng/mL propranolol, 0.5 ng/mL metformin. * p < 0.05, one-way ANOVA. All AUC values are shown in Fig. S1. Measurements in triplicate, error bars depict standard deviation.

253 Upregulation in stress response and multidrug efflux genes in 254 response to ibuprofen exposure

255 We noted a reduction in maximum OD following exposure to the compounds. To
256 investigate the cause of this further, we conducted transcriptomic analysis on *E. coli*
257 populations grown in the presence and absence of 50 µg/mL ibuprofen. Ibuprofen
258 was selected as exposure to this compound resulted in one of the larger reductions
259 in maximum OD over a 24 hour time course, and it has been linked previously to a
260 resistance phenotype by enhancing the transfer of resistance genes [31]. We found
261 16 genes were significantly upregulated in the presence of ibuprofen relative to the
262 untreated control (Fig. 2). Those with the largest log fold change that could be in-
263 fluencing phenotype include *insC* (4.247), *nikA* (2.539), *yhcN* (1.396), *yhiM* (1.340),
264 *lit* (1.289), and *mdtE* (1.285) (Table 1). NikA is a periplasmic binding protein for a
265 nickel ATP-binding cassette (ABC) transporter. The *mdtE* gene encodes the mem-
266 brane fusion protein component of a multidrug efflux system. Genes involved in a
267 second multidrug efflux transporter, *emrA* and *emrD*, are also significantly upregu-
268 lated, albeit to a lesser extent. The *yhcN* gene is linked to a response to stress, and

269 *yhiM* to acid resistance. These data suggest that the observed reduction in maxi-
 270 mum OD following ibuprofen exposure could be attributed to the cells undergoing
 271 a stress response or actively exporting the compound.

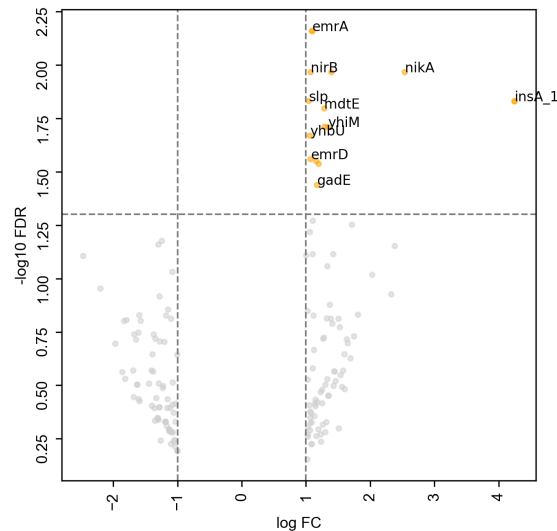


Fig. 2 Genes significantly upregulated (yellow) (false discovery rate [FDR] threshold of $p < 0.05$ and an absolute log fold change [FC] of at least one) in the presence of ibuprofen. Genes which had an absolute log FC of at least one but did not reach the FDR threshold are shown in grey and are considered to not be significantly differentially expressed. Selected genes are labelled.

Table 1 *E. coli* genes upregulated significantly in the presence of ibuprofen, their function as assigned by Prokka, and the average (of biological triplicate) log fold change (FC) when normalised against *E. coli* grown in the absence of ibuprofen.

Gene	Function	log FC
<i>insA</i>	IS2 element protein	4.247
<i>insA</i>	IS2 element protein	4.247
<i>nikA</i>	Nickel ABC transporter - periplasmic binding protein	2.539
<i>yhcN</i>	Stress-induced protein	1.396
<i>yhiM</i>	Inner membrane protein with a role in acid resistance	1.340
<i>lit</i>	Cell death peptidase; phage exclusion; e14 prophage	1.289
<i>mdtE</i>	MdtEF-TolC multidrug efflux transport system - membrane fusion protein	1.285
<i>yhiD</i>	Putative Mg(2+) transport ATPase	1.199
<i>gadE</i>	DNA-binding transcriptional activator	1.172
<i>bhsA</i>	Outer membrane protein involved in copper permeability, stress resistance and biofilm formation	1.161
<i>aegA</i>	Putative oxidoreductase, Fe-S subunit	1.099
<i>emrA</i>	EmrAB-TolC multidrug efflux transport system - membrane fusion protein	1.093
<i>nirB</i>	Nitrite reductase, large subunit	1.073
<i>emrD</i>	Multidrug efflux transporter	1.072
<i>yhbU</i>	Putative peptidase (collagenase-like)	1.059
<i>slp</i>	Starvation lipoprotein	1.044

272 **Co-exposure to ibuprofen and antibiotics does not alter**
273 **MICs for *E. coli* MG1655**

274 Microbial communities residing in or near industrial wastewater will be exposed
275 to a cocktail of antibiotic and non-antibiotic compounds. The presence of a non-
276 antibiotic pharmaceutical may induce a response that could alter the MIC of an
277 antibiotic during co-exposure. To assess this, *E. coli* MG1655 and ATCC 25922 (as
278 a control strain) were co-exposed to ibuprofen plus one of the following antimicrobial
279 agents; ethidium bromide, ampicillin, ciprofloxacin, chloramphenicol, trimethoprim,
280 and colistin. No change in MIC was observed for *E. coli* MG1655 for any ibuprofen-
281 antimicrobial pair, with FIC scores indicating no synergy or antagonism (Table 2).
282 This suggests that in laboratory strains of *E. coli*, co-exposure to ibuprofen alongside
283 an antibiotic does not alter the resistance profile.

Table 2 MICs (mg/L) for six antimicrobial agents against the MG1655 ancestral lineage, the MG1655 strain evolved in the presence of ibuprofen, and an ATCC 25922 control strain (n=3, one biological replicate each, modal MIC value given with the exception of ciprofloxacin against MG1655 evolved where the median value is given). Checkerboard (CB) MICs (mg/L) for *E. coli* MG1655 and ATCC 25922 co-exposed to ibuprofen plus an antimicrobial agent, one of; ethidium bromide, ampicillin, ciprofloxacin, chloramphenicol, trimethoprim, or colistin (n=3, one biological replicate each, modal MIC value given). Fractional Inhibitory Concentration (FIC) scores calculated whereby 0.5 – 4 indicates no synergy or antagonism. - indicates not applicable/not tested.

Agent	MIC ancestor	MIC evolved	MIC ATCC 25922	CB MIC ancestor	FIC ancestor	CB MIC ATCC 25922	FIC ATCC 25922
Ethidium bromide	5	512	256	512	2	256	2
Ampicillin	4	8	8	8	3	8	2
Ciprofloxacin	0.0156	0.0156	0.0156	0.0156	2	0.0156	2
Chloramphenicol	8	8	4	8	2	4	2
Trimethoprim	0.25	0.25	0.25	0.5	3	0.5	3
Colistin	4	4	4	2	1.5	4	2
Ibuprofen	>200	-	>200	>200	-	-	-

284 **Long-term exposure to non-antibiotic pharmaceuticals**
285 **impacts *E. coli* growth but does not select for**
286 **cross-resistance to antibiotics**

287 After establishing the negative impact on *E. coli* growth in the presence of selected
288 non-antibiotic pharmaceuticals, we investigated whether this would be sufficient to
289 act as a selective pressure during long-term exposure. We therefore propagated
290 populations in the presence and absence of acetaminophen, ibuprofen, TiO₂, met-
291 formin, and propranolol individually, passaging cells every 24 hours for 30 days.
292 The evolved populations were then screened in the presence and absence of their

293 selection compound to assess growth in comparison to the ancestral lineage. We
294 observed a decrease in the maximum OD readings reached by the evolved popula-
295 tions in their selection media in comparison to the ancestral lineage, suggesting that
296 prolonged exposure to the compounds did not select for improved growth (Fig. 3).
297 This difference was statistically significant in the majority of populations ($p < 0.05$,
298 one-way ANOVA, Fig. S2), and was most notable for the populations exposed to
299 ibuprofen and TiO_2 (Fig. 3). The reduction in OD observed in the no compound
300 control was calculated to be not significant (Fig. S2). The difference remained when
301 the populations were exposed to 100x concentration of the compounds (Fig. S3),
302 and when all populations were passaged through a seven-day ‘recovery’ experiment
303 in NB with no added pharmaceutical (Fig. S4). This suggests therefore that the
304 growth patterns observed in the evolved populations was not a transient, reversible
305 effect as a result of long-term toxicity.

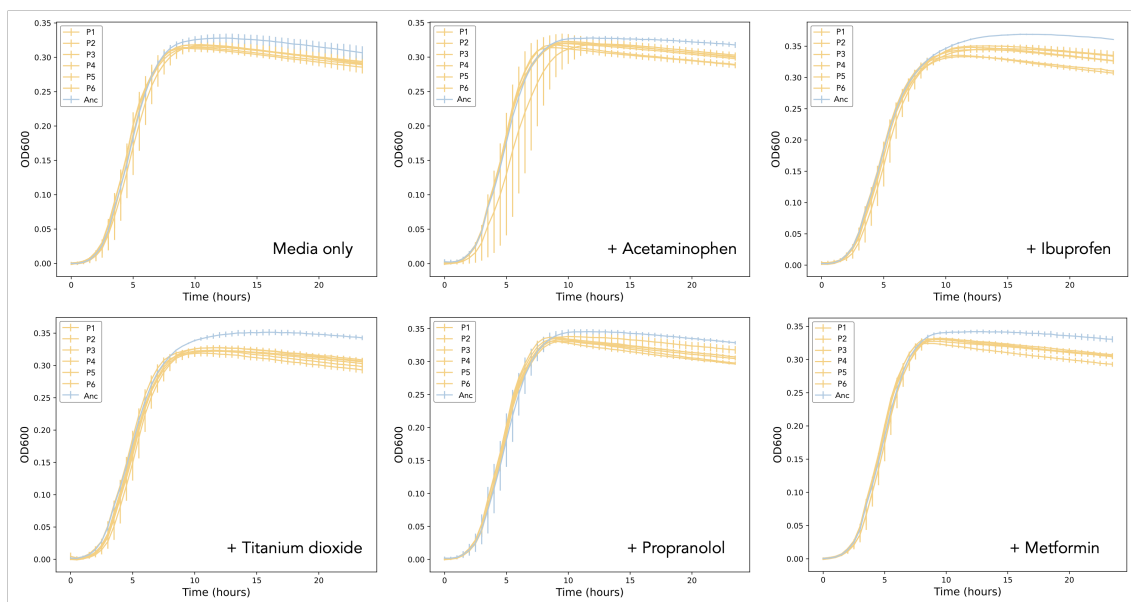


Fig. 3 Growth of evolved populations (yellow, six independent biological replicates P1-P6) in the presence of the compound in which their selection experiment was conducted in comparison to growth of the ancestral lineage (blue, Anc); (clockwise from top left) media-only control, acetaminophen, ibuprofen, metformin, propranolol, and TiO_2 . Measurements in triplicate, error bars depict standard deviation.

306 To establish whether there were any significant single nucleotide polymorphisms
307 (SNPs) arising as a result of the selection experiments, short-read assemblies were
308 generated for three replicates from each of the control and test conditions and the
309 sequences analysed using Snippy. No mutations parallel between independent evolu-
310 ting populations were found within treatments (Table S3). SNPs in *recQ* (ATP-
311 dependent DNA helicase) and *ygeA* (a putative racemase) were observed in single
312 replicates of evolved populations exposed to acetaminophen. The singular occur-

313 rence of each suggests they arose as a result of drift rather than selection, or that
314 selection was not strong enough for them to reach fixation within the other popula-
315 tions. An analysis of the gene presence/absence patterns across the hybrid assem-
316 bled evolved isolate genomes highlighted sequence variation in *ldrA* gene in three
317 sequences; one control, and one each evolved in acetaminophen and TiO₂. All had
318 three SNPs compared to the ancestral MG1655 (Table S2). The TiO₂ and the control
319 had identical SNPs, whereas the three SNPs in the acetaminophen-exposed isolate
320 were different. Again, these variations occurred in a single replicate per condition
321 only.

322 The IS2 element *insA* was shown to be upregulated in the presence of ibuprofen. IS
323 element transposition could be a cause of the differences in growth patterns between
324 the ancestral and evolved isolates. To assess this, hybrid assemblies were generated
325 and the distribution of IS elements then established using ISEScan and ISFinder.
326 All IS elements were present in equal numbers between the ancestor and all evolved
327 lineages. IS2, IS30, and IS1 elements were interrogated in depth and showed no
328 evidence of movement between any evolved isolate and the ancestor. Overall, these
329 data suggest that although the presence of the non-antibiotic pharmaceuticals has
330 a negative effect on the growth on *E. coli*, they do not exert a selection pressure
331 during prolonged exposure at the timescale and concentrations tested.

332 Given the observed upregulation in known efflux systems following ibuprofen treat-
333 ment, we examined whether prolonged exposure to ibuprofen may select for cross-
334 resistance to antibiotics. The MICs of several antimicrobials were analysed for a
335 strain evolved in the presence of ibuprofen compared to the ancestral isolate. Ethid-
336 ium bromide, ampicillin, ciprofloxacin, chloramphenicol, trimethoprim, and colistin
337 were chosen based on the previously mentioned transcriptomic data as compounds
338 which might logically have altered MICs as a result of ibuprofen exposure. The
339 MICs of the evolved isolate for all antibiotics tested were either the same or within
340 one doubling dilution of the ancestor (Table 2). This indicates the long-term expo-
341 sure to ibuprofen, regardless of the differential gene expression, does not co-select
342 for resistance to the antibiotics tested.

343 Discussion

344 The evolution of bacteria, including *E. coli*, in response to exposure to antibiotics
345 is well understood. The effects of short- and long-term exposure to non-antibiotic
346 pharmaceuticals is however less clear. This includes their potential toxicity and the

347 likelihood that their presence might select for cross-resistance to antibiotics. With
348 increasing evidence for the presence of non-antibiotic pharmaceuticals in proximity
349 to production plants [5; 18], it is becoming apparent that these are compounds that
350 require further investigation into their potential to impact local microbial popula-
351 tions. To begin to address this, we screened a panel of five non-antibiotic phar-
352 maceuticals against a laboratory strain of *E. coli* to assess their potential toxicity
353 and found that all five, to various degrees, reduced the maximum OD reached by
354 the population over a 24 hour incubation. This therefore suggests that these com-
355 pounds may, even at low concentrations, have a negative effect on members of the
356 microbial communities surrounding production plants. Our results contribute to
357 published work on ibuprofen toxicity, whereby it was demonstrated through disc
358 diffusion assays to not have antimicrobial activity against *E. coli* [30].

359 We then uncovered 16 genes that are upregulated significantly when *E. coli* was
360 grown in the presence of ibuprofen. Notable amongst these genes were two periplas-
361 mic adaptor proteins from different multidrug efflux systems; *mdtE* and *emrA*.
362 MdtEF-TolC is a resistance nodulation division (RND) family pump with beta-
363 lactams, benzalkonium chloride, macrolides, and oxazolidinones as known substrates,
364 and EmrAB-TolC is a major facilitator superfamily (MFS) efflux pump that confers
365 resistance to compounds including fluoroquinolones in *E. coli* [50]. Efflux pump
366 expression is often upregulated in the presence of toxic compounds to prevent their
367 accumulation inside the cell [51; 52; 53; 54]. There is some existing evidence linking
368 efflux pumps to a response to non-antibiotic pharmaceuticals. Exposing *S. aureus*
369 to diclofenac has been shown to downregulate a putative *emrAB*-family pump [55],
370 which contrasts the upregulation we observed in *E. coli* following ibuprofen expo-
371 sure. The response could therefore be specific to the species, compound, or pump,
372 and more work is needed to unravel this potential interaction.

373 Ibuprofen is an example of a partial proton motive force (PMF) uncoupler that
374 can inhibit the function of RND and MFS pumps [56]. The nitrite reductase *nirB*
375 was identified as another significantly upregulated gene. Nitrate reduction has been
376 shown to enhance bacterial survival in the presence of agents that dissipate PMF
377 [55]. This therefore provides tentative support to a hypothesis that *E. coli* may
378 be using nitrate reduction to ameliorate the dissipation of PMF in the presence of
379 ibuprofen, enabling the function of the RND and MFS pumps.

380 Ibuprofen exposure also resulted in the upregulation of several genes linked to a
381 response to stress, including *yhcN*, the inner membrane protein *yhiM*, and the outer
382 membrane protein *bhsA*. Existing research has shown a *bhsA* mutant of *E. coli* to be
383 more sensitive to a variety of stressors including acid [57], and *yhcN* has been linked

384 to cytoplasm pH stress [58]. Additionally, the gene observed to be upregulated to
385 the greatest extent in our data was *insA*, and IS2 and other IS elements are known
386 to be upregulated in response to stress [59; 60]. An upregulation of stress response
387 genes in response to non-antibiotic pharmaceuticals has been noted previously in *A.*
388 *baylyi*, where the uptake of antibiotic resistance genes was shown to be facilitated
389 by the presence of compounds including ibuprofen and propranolol [31]. There,
390 analysis including transcriptomics linked the observation to increased stress and the
391 over-production of reactive oxygen species, amongst other characteristics.

392 Our data suggest that the non-antibiotic components within pharmaceutical pro-
393 duction waste may affect the local microbial communities, as over time the toxicity
394 observed here may deplete species or genera within the communities, altering their
395 composition. Despite the observed reduction in maximum OD, we found that pro-
396 longed exposure to this panel of non-antibiotic pharmaceuticals did not result in
397 significant genetic changes across multiple independent populations. It is possible
398 that the stress induced by the compounds was dealt with sufficiently by, for example,
399 the upregulation of efflux pumps, thereby reducing the selection pressure, or that the
400 reduced OD was not due to changes in carrying capacity but rather due to morpho-
401 logical changes following induction of stress responses. We also found no evidence
402 of synergy when the ancestral strain was co-exposed to ibuprofen and one of a panel
403 of antibiotics. Additionally, when the same panel of antibiotics were tested against
404 the ancestor and the strain evolved in the presence of ibuprofen, there was no evi-
405 dence of altered MICs in the latter that would indicate selection for cross-resistance
406 to antibiotics. This is a reassuring initial investigation given the large quantities
407 of pharmaceutical production waste entering local ecosystems. Whilst the panel of
408 non-antibiotic pharmaceuticals tested here is small, they are commonly used and
409 consistently found in water bodies, and therefore can be considered a representa-
410 tive sample [61; 62; 63; 64]. Acetaminophen, ibuprofen, metformin, and propranolol
411 have also been identified as priority pharmaceuticals in India [65]. The use of a
412 standard laboratory strain of *E. coli* is a useful starting point, but previous work
413 suggesting that the activity of non-antibiotic pharmaceuticals may be strain-specific
414 [21] underscores the need for broad spectrum testing before definitive conclusions
415 can be drawn. Variations in concentrations over time as a result of effluent changes,
416 dry seasons, and climate change should also be considered, and there is a need
417 to extend this research to encompass production waste as a holistic entity against
418 environmentally relevant populations.

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424 Competing interests

425 The authors declare no competing financial interests in relation to the work de-
426 scribed.

427 Data availability statement

428 The datasets generated and analysed during the current study are available from
429 NCBI BioProject with accession PRJNA1005239.

430 Author contributions

431 RJH: methodology, formal analysis, investigation, data curation, writing - original
432 draft, visualization. AES: investigation, data curation. SJE: methodology, inves-
433 tigation, vizualisation. RAM: methodology. HS: validation, investigation. EAC:
434 methodology. MJB: methodology. JMAB: conceptualization, resources, supervision.
435 IA: funding acquisition. LJC: conceptualization, funding acquisition. AM: concep-
436 tualization, methodology, supervision, funding acquisition. All authors: writing -
437 review & editing.

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647 Supplementary

Table S1 Concentrations used to screen the non-antibiotic pharmaceutical compounds for toxicity against *E. coli*, and the final concentration chosen for selection experiments. No-compound controls were also used as comparisons for all screens. Relevant references are also included.

Compound	Conc. (screen)	Conc. (selection)	Manufacturer	Ref
Acetaminophen	100 µg/mL, 1 µg/mL, 10 ng/mL, 1 ng/mL	5 ng/mL	Merck Life Science UK Ltd	[37]
Ibuprofen	50 µg/mL, 5 µg/mL, 50 ng/mL, 5 ng/mL	2 µg/mL	Merck Life Science UK Ltd	[38; 39]
Titanium dioxide	100 µg/mL, 10 µg/mL, 1 µg/mL	1 µg/mL	Merck Life Science UK Ltd	[40; 41]
Propranolol	5 µg/mL, 500 ng/mL, 5 ng/mL, 0.5 ng/mL	0.5 ng/mL	VWR International Ltd	[42]
Metformin	50 µg/mL, 500 ng/mL, 5 ng/mL, 0.5 ng/mL	0.5 ng/mL	VWR International Ltd	[43; 44]

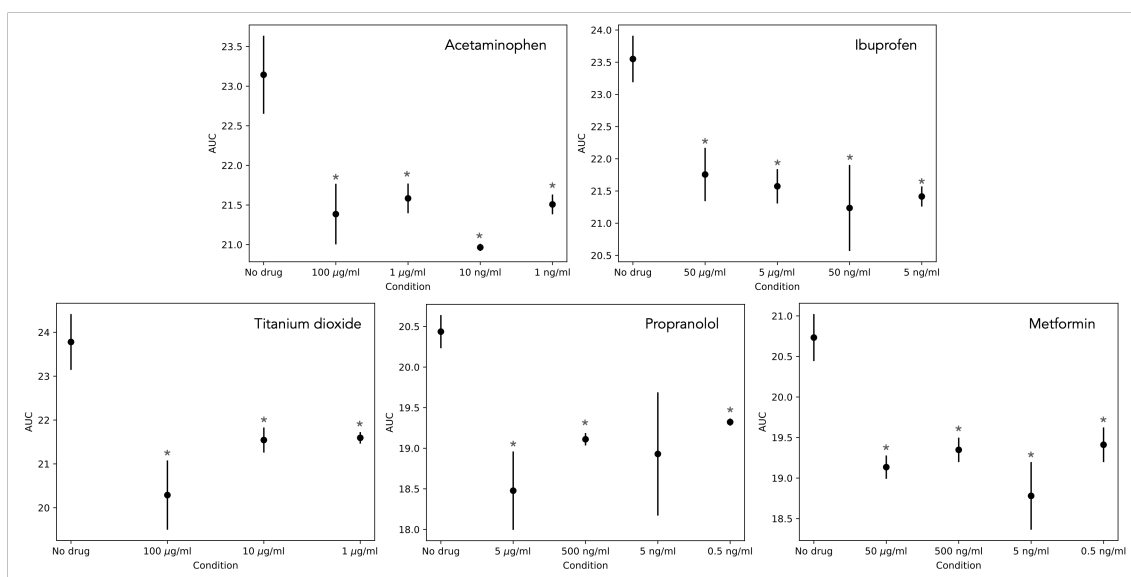


Fig. S1 AUC values for a toxicity screen of (clockwise from top left) acetaminophen, ibuprofen, metformin, propranolol, and TiO₂ at various concentrations against *E. coli*, relating to Fig. 1. A no-compound control ('No drug') was included for each screen. * p < 0.05, one-way ANOVA. Measurements in triplicate, error bars depict standard deviation.

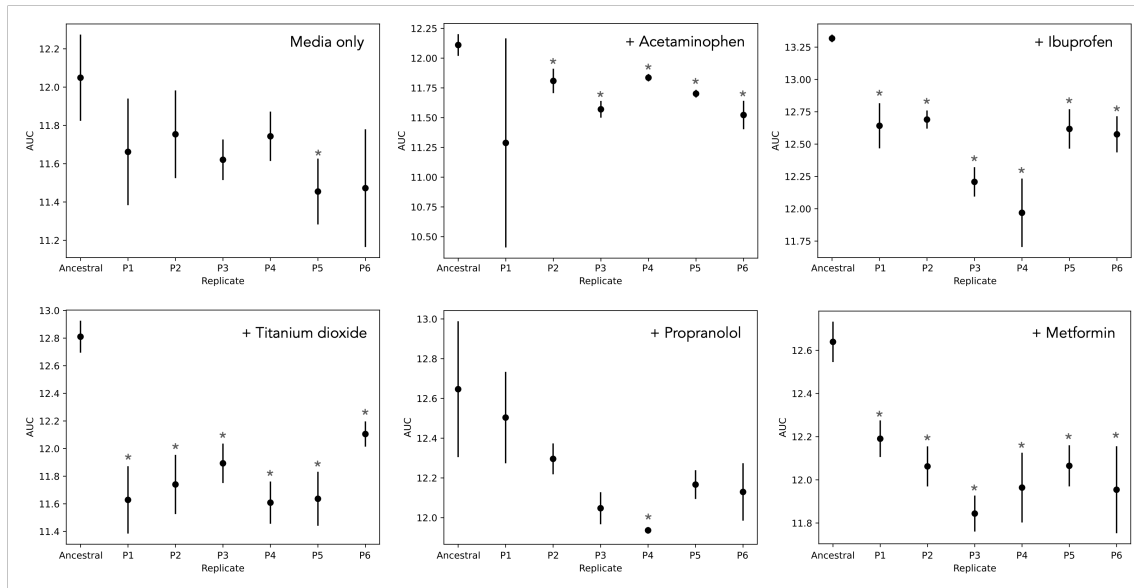


Fig. S2 AUC values for evolved populations (six independent biological replicates P1-P6) in a fresh sample of their respective evolution media in comparison to the ancestral lineage, relating to Fig. 3. * $p < 0.05$, one-way ANOVA. Measurements in triplicate, error bars depict standard deviation.

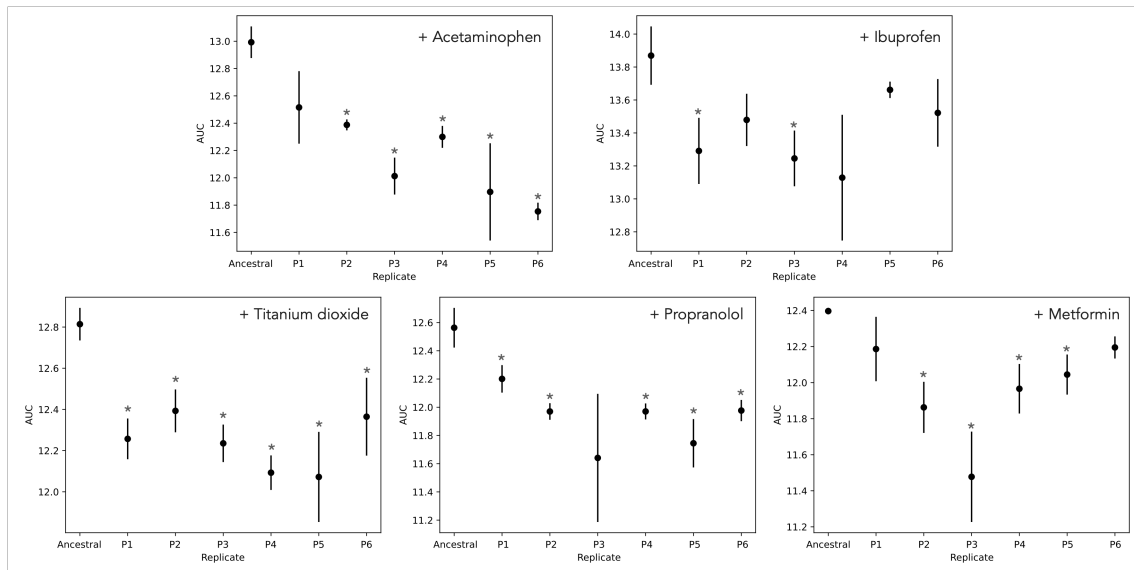


Fig. S3 AUC values for evolved populations (six independent biological replicates P1-P6) in the presence of 100x concentration of the compound in which their selection experiment was conducted in comparison to growth of the ancestral lineage. * $p < 0.05$, one-way ANOVA. Measurements in triplicate, error bars depict standard deviation.

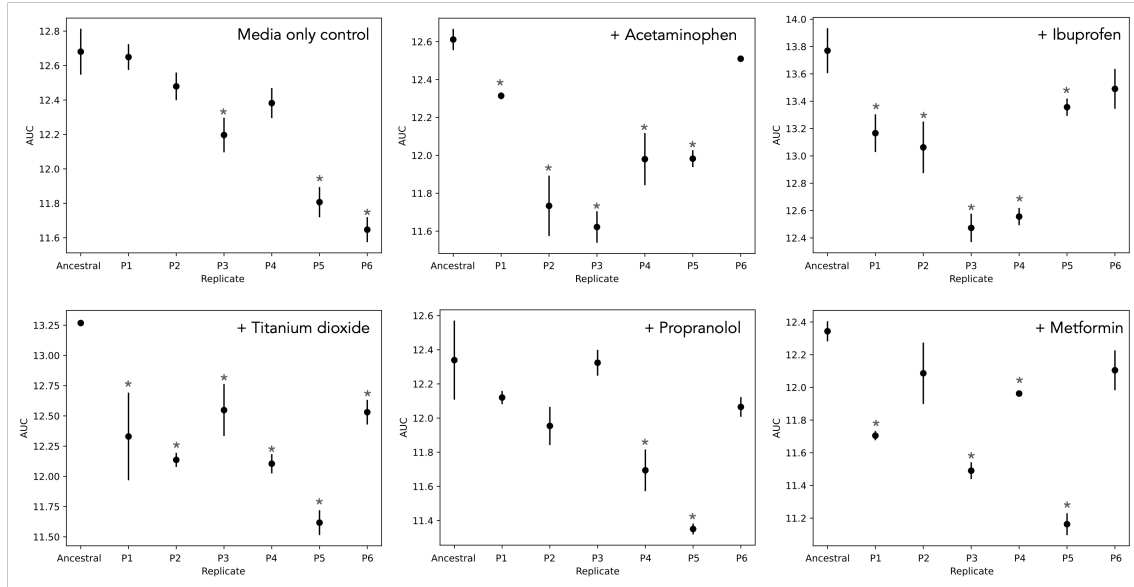


Fig. S4 AUC values for evolved populations (six independent biological replicates P1-P6) following serial passaging for seven days in the absence of pharmaceuticals in comparison to the ancestral lineage. * $p < 0.05$, one-way ANOVA. Measurements in triplicate, error bars depict standard deviation.

Table S2 SNP differences observed in the *ldrA* gene in the hybrid assemblies of the evolved isolates in comparison to the ancestor. R = test replicate, SNP = single nucleotide polymorphism.

Condition	SNP	Position
Control R3	T→C	28
	A→G	47
	A→G	87
+ Acetaminophen R3	T→C	24
	T→C	64
	A→G	83
+ Titanium dioxide R3	T→C	28
	A→G	47
	A→G	87

Table S3 SNPs identified in each sequence (R = test replicate) as predicted by Snippy. The nucleotide(s) in the reference (ref) and comparison (alt) sequences, the locus tag of the gene, and its product are given.

Condition	Ref	Alt	Locus tag	Gene	Product
Ancestor	A	C	00551	<i>rhcC</i>	Protein RhsC
Media only R1	A	G	00431		Uncharacterized protein HI_1672
	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
	T	G			
Media only R2	A	C	00551	<i>rhcC</i>	Protein RhsC
Media only R3	A	C	00551	<i>rhcC</i>	Protein RhsC
	A	C	03981	<i>sspA</i>	Stringent starvation protein A
+Acetaminophen R1	A	C	00551	<i>rhcC</i>	Protein RhsC
	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
	C	A			
+Acetaminophen R2	T	G	03610	<i>recQ</i>	ATP-dependent DNA helicase
	A	C	00551	<i>rhcC</i>	Protein RhsC
	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
+Acetaminophen R3	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
	GCGCC	G	01387	<i>ygeA</i>	Putative racemase
	A	C	00551	<i>rhcC</i>	Protein RhsC
+Ibuprofen R1	A	C	00551	<i>rhcC</i>	Protein RhsC
+Ibuprofen R2	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
+Ibuprofen R3	A	C	00551	<i>rhcC</i>	Protein RhsC
	A	G	00553		Hypothetical protein
+Titanium dioxide R1	A	C	00551	<i>rhcC</i>	Protein RhsC
	C	G	00553		Hypothetical protein
	A	G	00553		Hypothetical protein
+Titanium dioxide R2	A	C	00551	<i>rhcC</i>	Protein RhsC
	A	G	00553		Hypothetical protein
+Titanium dioxide R3	A	C	00551	<i>rhcC</i>	Protein RhsC
	C	G	00553		Hypothetical protein
+Propranolol R1	A	G	00553		Hypothetical protein
	A	C	00551	<i>rhcC</i>	Protein RhsC
	T	G			
+Propranolol R2	A	C	00551	<i>rhcC</i>	Protein RhsC
+Propranolol R3	A	C	00551	<i>rhcC</i>	Protein RhsC
	A	G	00553		Hypothetical protein
+Metformin R1	G	A			
	A	C	00551	<i>rhcC</i>	Protein RhsC
	C	G	00553		Hypothetical protein
+Metformin R2	A	G	00553		Hypothetical protein
	A	C	00551	<i>rhcC</i>	Protein RhsC
	A	G	00553		Hypothetical protein
+Metformin R3	G	T			
	None				