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## 1 Selective conditions for a multidrug resistance plasmid depend on the

- 2 sociality of antibiotic resistance
- 3
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- 9 Running heading: Social selection of a MDR plasmid
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- 13

## 14 ABSTRACT

15 Multidrug resistance (MDR) plasmids frequently encode antibiotic resistance

- 16 genes conferring qualitatively different mechanisms of resistance. We show that
- 17 the antibiotic concentrations selecting for the RK2 plasmid in Escherichia coli
- 18 depend upon the sociality of the drug resistance: Selection for a selfish drug
- 19 resistance (efflux-pump) occurred at very low drug concentrations, just 1.3% of
- 20 the sensitive's MIC, whereas selection for a cooperative drug resistance
- 21 (modifying-enzyme) occurred at drug concentrations exceeding the MIC of the
- 22 plasmid-free strain.
- 23

25 Antibiotics are critical to modern medicine, but their widespread use and misuse 26 has lead to the evolution of resistant strains to most commonly used antibiotics 27 (1, 2). Antibiotic resistance has become a major threat to global health, with 28 multi-drug resistant (MDR) bacteria observed globally (3). Environmental 29 antibiotic resistance genes (ARGs) are a major source of clinical resistance (4). 30 ARGs can be selected for at very low concentrations of antibiotic, far below the 31 minimum inhibitory concentration (MIC) of sensitive cells (5, 6), with antibiotic 32 contamination at sub-MIC concentrations being proposed as the main driving 33 force behind environmental selection for resistance (7–9). However, ARGs can 34 encode gualitatively different forms of resistance ranging from selfish to 35 cooperative. Selfish drug resistances only confer a benefit to the individual cell 36 harbouring it, for example efflux pumps, reduced membrane permeability and 37 alteration of antibiotic targets (10, 11). By contrast cooperative antibiotic 38 resistances benefit both the resistant cell and surrounding cells whether they are 39 resistant or not. For example, modifying enzymes such as  $\beta$ -lactamase inactivate 40 the antibiotic through hydrolysis, decreasing its environmental concentration. 41 Localisation of the  $\beta$ -lactamase enzyme in the periplasmic space may enhance 42 the share of the benefit for the resistant cell, but nevertheless, the decrease in 43 the overall environmental concentration of antibiotic will benefit both resistant and 44 sensitive cells (12). We hypothesised that the sociality of drug resistance could 45 alter the selective conditions for the spread of ARGs (13, 14). Specifically, 46 because the benefits of selfish drug resistance are directed solely to the resistant

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47 cell, whereas the benefits of cooperative drug resistance are shared between 48 resistant and sensitive cells, we predict that selfish drug resistance should be 49 selected at lower relative drug concentrations (i.e. % of the sensitive MIC) than 50 cooperative resistance.

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52 Multiple ARGs are frequently clustered together onto conjugative plasmids 53 including combinations of selfish and cooperative drug resistances (15). How 54 combinatorial antibiotic usage selects for MDR plasmids is not clear, especially 55 for combinations of antibiotics requiring qualitatively different modes of drug 56 resistance, such as selfish or cooperative drug resistances. Here we tested how 57 the sociality of drug resistance, and single versus combined antibiotic treatment, 58 altered the selective conditions for the MDR plasmid RK2 (16) in Escherichia coli 59 MG1655. RK2 encodes both cooperative ampicillin resistance, mediated by a β-60 lactamase, and selfish tetracycline resistance, mediated by an efflux pump. We 61 report that the selfish drug resistance is selected for at far lower relative antibiotic 62 concentrations than the cooperative drug resistance, and that combined antibiotic 63 selection is additive, showing no interaction.

64

65 Conventionally, ARGs are thought to be positively selected at antibiotic

66 concentrations exceeding the MIC of sensitive cells in monoculture (17) (i.e. the

67 conventional selective window, Fig 1). To determine whether the sociality of

68 resistance affected the selection window for the RK2 MDR plasmid, we estimated

69 the relative fitness of plasmid bearing versus isogenic plasmid free cells by direct

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72 decreasing the fitness of *E. coli* by 19% (Fig. 1A/B, t test, p < 0.001, t = -9.8674, 73 df = 23). An intrinsic cost is often associated with plasmid carriage when 74 accessory traits are not under positive selection due to cellular disruption and 75 increase transcriptional load (18). Cooperative ampicillin resistance was 76 positively selected at ampicillin concentrations exceeding the MIC of sensitive E. 77 coli (Fig. 2A). Importantly, sensitive cells were able to maintain positive growth in 78 mixed cultures at ampicillin concentrations that completely inhibited their growth 79 in monoculture (>8µg/ml; cf. Fig. 1A & Fig. S4), justifying the assignment of 80 ampicillin resistance as cooperative. Thus cooperative resistance permits 81 persistence of a sensitive subpopulation beyond the sensitive MIC due to the 82 inactivation of the antibiotic, potentially allowing reinvasion by sensitive cells 83 once the antibiotic concentration is sufficiently reduced by the action of resistant 84 cells. 85 86 In contrast, selfish tetracycline resistance was positively selected at tetracycline 87 concentrations of just 1.3% of the MIC of sensitive E. coli (Fig. 2B). Indeed, at

competition following standard methodology (see supplementary material). In the

absence of antibiotics the plasmid imposed a significant cost of carriage,

88 concentrations of tetracycline above 10% of the MIC of sensitive E. coli, the 89 resistant plasmid bearers competitively excluded the plasmid-free bacteria, with 90 no plasmid-free cells observable (Fig. S1). This is despite the fact that plasmid-

91 free E. coli could survive at these tetracycline concentrations when grown alone

92 (Fig. 1B). Our data suggest that selfish tetracycline resistance is positively

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93 selected in the sub-MIC selective window at very low tetracycline concentrations,

94 similar to those observed in the natural environment (19).

95

96 When ampicillin and tetracycline were applied in combination there was no 97 significant interaction ( $F_{1.68}$  = 0.2395, p = 0.6261) indicating that when these two 98 antibiotics were used in combination their selective effects were independent and 99 additive (Fig. 2C). This means that very low concentrations of tetracycline were 100 sufficient to completely mask the population-level effects of cooperative ampicillin 101 resistance. With increasing tetracycline concentrations, the ampicillin 102 concentration positively selecting for the MDR plasmid shifted to lower and lower 103 sub MIC levels, reducing the window of selective conditions where sensitive cells 104 could persist (Fig. 2D). 105 106 Residues of multiple antibiotics are commonly found contaminating the same 107 environments at low concentrations (19, 20). These combinations, and 108 particularly the presence in the environment of antibiotics like tetracycline 109 targeted by selfish efflux-mediated resistance, will select for the spread of MDR 110 plasmids and competitive exclusion of sensitive cells. This is despite being 111 present at concentrations far below the level required to positively select

112 resistance individually. This adds further evidence that ARGs, whether

113 chromosomal or plasmid encoded, can be positively selected at antibiotic

114 concentrations far below the MIC of sensitive strains (5, 6, 9).

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116 Our study has a number of possible limitations: First, it is possible that other 117 factors, in addition to sociality, may have contributed to differences in the fitness 118 reaction norms of the antibiotics, including the contrasting effects of sub-MIC 119 concentrations on monoculture densities and the fact that ampicillin is 120 bacteriocidal whereas tetracycline is bacteriostatic. Second, we use exemplars of 121 cooperative and selfish resistance but more research will be required to test the 122 importance of sociality on the selective conditions for other resistance 123 mechanisms. 124 125

Here we show that the extent to which an ARG is positively selected at sub-MIC 126 antibiotic concentrations depends upon the sociality of the mechanism of drug 127 resistance. Cooperative ampicillin resistance is positively selected at ampicillin 128 concentrations exceeding the MIC, whereas selfish tetracycline resistance is 129 positively selected at 100-fold lower relative drug concentrations. This striking 130 difference in the selective window for ARGs co-located on the same MDR 131 plasmid probably arises because of the population-level effects of the ARGS: 132 Cooperative ampicillin resistance allowed sensitive bacteria to survive past their 133 MIC by reducing the ampicillin concentration and sharing the benefits of 134 resistance, whereas, selfish tetracycline resistance drove complete competitive 135 exclusion of sensitive cells at >10% MIC due to the exclusively individual benefits 136 of efflux-mediated resistance. Combining the two antibiotics – at concentrations 137 that would not normally select for resistance individually - selects for both 138 resistances and spread of the MDR plasmid. Taken together these findings

- 139 suggest that selfish efflux-mediated drug resistances are likely to be especially
- 140 important for the selective maintenance and spread of MDR plasmids.
- 141
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- 154

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- 219
- 220 FIG 1
- 221 Cell density (OD<sub>600</sub>) of sensitive plasmid free bacteria (green line) and resistant
- 222 plasmid containing bacteria (blue line) as a function of A ampicillin concentration,
- 223 B tetracycline concentration after 24 hours growth in monoculture. Error bars

show SEM (n=6). Area shaded in green shows the sub-MIC selective window,
and the area shaded in blue shows the selective window conventionally thought
to select for resistance.

227

228 FIG 2

229 Fitness reaction norms as a function of antibiotic concentration during

230 competition experiments between *E. coli* harboring the RK2 plasmid and isogenic

231 plasmid free sensitive strains. Competitions in the presence of A ampicillin, B

232 tetracycline, red lines show fitted regression. C/D Fitness reaction norms of

233 combination treatments with both ampicillin and tetracycline during competition

234 experiments between RK2 harboring and plasmid free strains. There is no

235 significant interaction of antibiotic treatments upon the relative fitness (F<sub>1,68</sub> =

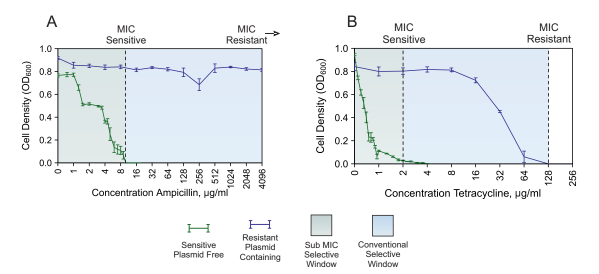
236 0.2395, p = 0.6261) indicating treatments were non-interacting and additive. Error

237 bars show SEM (n=6), Antibiotic concentrations shown as percentages of

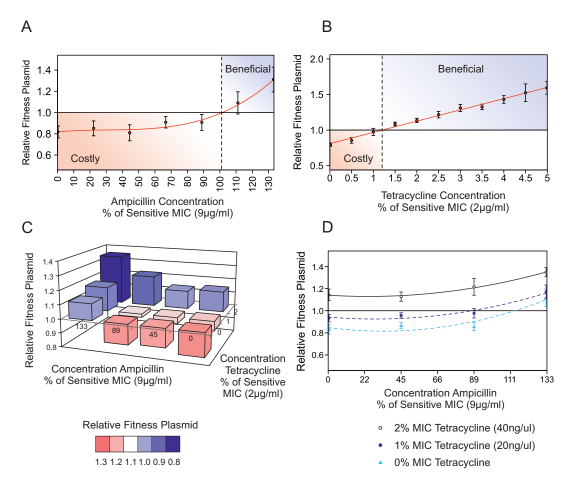
238 sensitive MIC.

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**FIG 1** Cell density (OD600) of sensitive plasmid free bacteria (green line) and resistant plasmid containing bacteria (blue line) as a function of A ampicillin concentration, B tetracycline concentration after 24 hours growth in monoculture. Error bars show SEM (n=6). Area shaded in green shows the sub-MIC selective window, and the area shaded in blue shows the selective window conventionally thought to select for resistance.



**FIG2** Fitness reaction norms as a function of antibiotic concentration during competition experiments between E. coli harboring the RK2 plasmid and isogenic plasmid free sensitive strains. Competitions in the presence of A ampicillin, B tetracycline, red lines show fitted regression. C/D Fitness reaction norms of combination treatments with both ampicillin and tetracycline during competition experiments between RK2 harboring and plasmid free strains. There is no significant interaction of antibiotic treatments upon the relative fitness (F1,68 = 0.2395, p = 0.6261) indicating treatments were non-interacting and additive. Error bars show SEM (n=6), Antibiotic concentrations shown as percentages of sensitive MIC.