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Species-specific activity of antibacterial drug combinations

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Abstract

The spread of antimicrobial resistance has become a serious public health concern, making once treatable diseases deadly again and undermining breakthrough achievements of modern medicine^{1,2}. Drug combinations can aid in fighting multi-drug resistant (MDR) bacterial infections, yet, are largely unexplored and rarely used in clinics. To identify general principles for antibacterial drug combinations and understand their potential, we profiled ~3,000 dose-resolved combinations of antibiotics, human-targeted drugs and food additives in 6 strains from three Gram-negative pathogens, *Escherichia coli*, *Salmonella* Typhimurium and *Pseudomonas aeruginosa*. Despite their phylogenetic relatedness, more than 70% of the detected drug-drug interactions are species-specific and 20% display strain specificity, revealing a large potential for narrow-spectrum therapies. Overall, antagonisms are more common than synergies and occur almost exclusively between drugs targeting different cellular processes, whereas synergies are more conserved and enriched in drugs targeting the same process. We elucidate mechanisms underlying this dichotomy and further use our resource to dissect the interactions of the food additive, vanillin. Finally, we demonstrate that several synergies are effective against MDR clinical isolates *in vitro* and during *Galleria mellonella* infections with one reverting resistance to the last-resort antibiotic, colistin.

Main text

To study the characteristics and conservation of drug-drug interactions in bacteria, we selected three γ -proteobacterial species, *E. coli*, *Salmonella enterica* serogroup Typhimurium, and *P. aeruginosa*, all belonging to the highest risk groups for antibiotic resistance³. We used model lab strains rather than MDR isolates to derive general principles behind drug-drug interactions without being confounded by horizontally transferred antibiotic resistance elements, and to facilitate follow-up experiments and comparisons with results from others. To further assess whether drug responses vary between strains of the same species, we included two strains per species (ED Fig. 1a), probing each in up to 79 compounds alone and in pairwise combinations. The compounds comprised 59% antibiotics (all major drug classes), 23% human-targeted drugs and food additives, most with reported antibacterial/adjuvant activity^{4,5}, and 18% of other compounds with known bacterial targets or genotoxic effects – e.g. proton motive force (PMF) inhibitors or oxidative damage agents, due to their potential relevance for antibiotic activity and/or uptake^{6,7} (ED Fig. 1a; Supplementary Table 1). Altogether, we profiled up to 2,883 pairwise drug combinations in each of the 6 strains (17,050 total). We assessed each drug combination in a 4x4 tailored dose matrix (Methods, Supplementary Table 1), using optical density as growth readout, and calculated fitness as the growth ratio between drug treated and untreated cells (ED Fig. 1-2, Methods). All experiments were done at least twice and on average 4x, with good replicate correlation (average Pearson Correlation =0.93; ED Fig. 3a-b).

We quantified all drug-drug interactions using the Bliss independence model (ED Fig. 1b, Methods). Consistent with its null hypothesis, interaction scores are zero-centered for all species (ED Fig. 3c). From all the scores (ε) obtained per combination (4x4 dose matrix), we derived a single interaction score $\tilde{\varepsilon}$ ranging from -1 to 1 (Methods). Synergies and antagonisms were considered significant if p-value < 0.05 (Benjamini-Hochberg corrected, 10,000 repetitions of a two-sided Wilcoxon rank-sum test). Strong interactions had an additional effect size requirement for $|\tilde{\varepsilon}| > 0.1$, whereas weak interactions could satisfy the effect size threshold for one of the two strains of the same species, but be just below for the other ($|\tilde{\varepsilon}| > 0.06$; Methods). In total we detected ~19% interactions for *E. coli*, ~16% for *S. Typhimurium*, and ~11% for *P. aeruginosa* (Supplementary Table 2). This is in between the >70% hit rate for a limited set of antibiotics tested in *E. coli*⁸ and the <2% for a larger set of antifungals tested in different fungi⁹. Discrepancies are likely due to: (i) drug selection biases, (ii) single drug concentrations used in previous studies (which increases false negative and positive rates), and (iii) different data analysis. For example, we observed that drugs lacking antibacterial activity engage in fewer interactions (ED Fig. 3e). Robbins *et al.*

screened pairwise combinations of 6 antifungals with 3,600 drugs, most of which had no antifungal activity⁹, thus explaining the low number of interactions detected, whereas Yeh *et al.* profiled only bioactive antibiotics⁸. Out of 79 drugs tested here, 69 had at least one interaction and a median of 5-13 interactions per drug in the different strains (ED Fig. 3f).

Since drug combinations have not been systematically probed in bacteria before, we lacked a ground truth for benchmarking our dataset. To overcome this limitation, we selected 242 combinations across the 6 strains, and created a validation set using higher-precision 8x8 checkerboard assays (ED Fig. 4a-b, Supplementary Table 3, Methods). We used this validation set to both assess the performance of our interaction identification approach and to benchmark our screen (ED Fig. 4c-d). Overall, we had precision and recall of 91% and 74%, respectively. The slightly lower recall can be partially explained by the larger coverage of drug concentration range in the validation experiments, which increases our ability to detect interactions (ED Fig. 5). We further confirmed 90% of all weak interactions we tested (n=46; Supplementary Table 3, ED Fig. 6), validating the rationale of our interaction identification approach. Indeed, including weak interactions in our hits contributes to higher recall (ED Fig. 4d). For a handful of the synergies observed between antibiotics of the same class (β -lactams), we confirmed the interactions using the Loewe additivity model (ED Fig. 4e), which is more suitable for assessing interactions between drugs with the same target.

Overall, we detected 1354 antagonistic and 1230 synergistic interactions. Although this suggests that the two occur with similar frequencies, antagonisms are nearly 50% more prevalent than synergies, when correcting for the ability to detect both types of interactions (Fig. 1a). This is because we can detect antagonisms only for 75% of combinations (when at least one individual drug inhibits growth; ED Fig. 3d, Methods), whereas synergies are detectable for nearly all combinations. Higher prevalence of antagonisms has also been reported for antifungals¹⁰.

Strikingly, antagonisms and synergies exhibited a clear dichotomy in our data. Antagonism occurred almost exclusively between drugs targeting different cellular processes, while synergies were also abundant for drugs of the same class or targeting the same process (Fig. 1b-e, ED Fig. 7). Mechanistically, antagonism could be explained by interactions at the drug target level, with the two inhibitors helping the cell to buffer the distinct processes perturbed. DNA and protein synthesis inhibitors act this way in bacteria (Fig. 1b)¹¹. Consistent with this being a broader phenomenon, in genome-wide genetic interactions studies in yeast, alleviating interactions (antagonisms) are enriched between essential genes (the targets of anti-infectives), which are part of different functional processes¹².

However, antagonism can also occur at the level of intracellular drug concentrations (ED Fig. 8a). We tested 16 antagonistic interactions of different drugs with gentamicin or ciprofloxacin in *E. coli* to investigate to what extent this occurs. Although initially detected at a growth inhibition level, all antagonisms held true at a killing level, with 14/16 decreasing the intracellular gentamicin or ciprofloxacin concentrations (ED Fig. 8b). In several cases tested, this likely occurred because the second drug either decreased the PMF-energized uptake of gentamicin or increased AcrAB-TolC-dependent efflux of ciprofloxacin, as antagonisms were neutralized in the respective mutant backgrounds (ED Fig. 8c). Overall, our results suggest that a large fraction of antagonisms is due to modulation of intracellular drug concentrations, rather than due to direct interactions of the primary drug targets (ED Fig. 8d-e).

Unlike antagonistic interactions, synergies often occurred between drugs targeting the same cellular process (Fig. 1b-e, ED Fig. 7). In fact, synergies are significantly enriched within drugs of the same category across all three species (p-value < 10^{-16} , Fischer's exact test), given that there are ~15-fold more possible drug combinations across drug categories in our dataset. Mechanistically, targeting the same functional process at different steps could tease apart its redundancy. For example, β -lactams have different affinities to the numerous and often redundant penicillin-binding-proteins (PBPs), likely explaining the many synergies between them (Fig. 1b, ED Fig. 4e & 7a-b).

Like antagonisms, synergies can also occur due to modulation of intracellular drug concentrations. Consistent with a general permeabilization role of membrane-targeting compounds in many organisms^{9,13,14}, and with drug uptake being a major bottleneck for Gram-negative pathogens, one fourth of all detected synergies contain at least one out of eight membrane-targeting drugs in our screen (Wilcoxon rank-sum test, p-value=0.044). However, membrane-targeting compounds account also for ~16% of antagonisms, suggesting that perturbations in membrane integrity can also decrease intracellular drug concentrations. Consistently, benzalkonium decreases the intracellular concentration of both gentamicin and ciprofloxacin, likely because it interferes with their active import in the cell (ED Fig. 8b-c).

We next examined the conservation of drug-drug interactions. Interactions within species were highly correlated (Fig. 2a, ED Fig. 9a-b): 53-76%, depending on the species (Fig. 2b). Conservation is actually higher (68-87%, and on average 80%), if we disregard the non-comparable interactions for which the concentration range tested preclude us from detecting synergy or antagonism for both strains (Fig. 2b, ED Fig. 3d). High conservation of drug-drug

interactions within species is consistent with the finding that such interactions are generally robust to simple genetic perturbations ¹⁵. Despite this overall high-degree conservation within species, 13-32% of the interactions were strain-specific, with the majority being neutral in the second strain. Very few drug combinations synergized for one strain and antagonized for the other (16 interactions), but such strain differences held in our validation set (Supplementary Table 2).

While conservation is relatively high within species, it is very low across species (Fig. 2c, ED Fig. 9c). The majority (70%) of interactions occurred in one species, and only 5% were conserved in all three phylogenetically close-related species. Since conservation is much higher at the single-drug level for the three species (sharing resistance/sensitivity to 73% of the drugs; Supplementary Table 1, Methods), this indicates that drug combinations can impart species specificity to the drug action. Such specificities can be beneficial for creating narrow spectrum therapies with low collateral damage, by using synergies specific for pathogens and antagonisms specific for abundant commensals.

Moreover, we found that synergies are significantly more conserved than antagonisms (Fig. 2d), despite being less prevalent (Fig. 1a). This is presumably because: i) synergies are enriched between drugs of the same category, and interactions within functional processes are conserved across evolution ¹⁶; ii) membrane-targeting drugs have a general potentiation effect in Gram-negative bacteria; and iii) antagonisms often depend on drug import/uptake (ED Fig. 8), which are controlled by less conserved envelope machineries.

Exploring the network of conserved drug-drug interactions across the three species (ED Fig. 9d) exposed potential Achilles heels of Gram-negative bacteria, such as the strong synergy of colistin with macrolides ¹⁷, but also revealed that known antibiotic classes often behave non-uniformly. For example, the well-known synergy between β -lactams and aminoglycosides is confined to potent aminoglycosides used in our screen (amikacin and tobramycin) and β -lactams that target specifically the cell-division related PBPs (piperacillin, aztreonam, cefotaxime), in agreement with previous reports ¹⁸. To address whether pairwise drug interactions are Mode of action (MoA)-driven (i.e. drug classes interacting purely synergistic or antagonistic with each other) ⁸, we calculated a monochromaticity index (MI) for all drug category pairs, across all species (Methods). For highly monochromatic category pairs, MI approaches 1 and -1 for antagonism and synergy, respectively. MI is overall high, especially between well-defined antibiotic classes. Yet, a number of them, including β -lactams, tetracyclines and macrolides, have mixed antagonisms and synergies with other

antibiotic classes (ED Fig. 9e). While β -lactams have diverse affinities to their multiple PBP targets (potentially explaining the mixed interactions with other classes), the same does not apply to protein synthesis inhibitors, which have unique targets. In this case, non-uniform class behavior may be due to different chemical properties of the class members, and thus different dependencies on uptake and efflux systems. Aggregating the MI per drug category reinforced the view that broader categories exhibit less concordant interactions (ED Fig. 9f). Interestingly, human-targeted drugs were the category exhibiting more synergies, supporting the hypothesis that many human-targeted drugs may act as adjuvants.

Since antibiotic classes interacted largely monochromatically, as expected, clustering drugs according to their interactions recapitulated the class groupings (ED Fig. 10). For example, cell-wall inhibitors grouped together, with further subdivisions being reflective of target specificity. Yet, exceptions were also evident, such as the macrolides, which split, with azithromycin, the only dibasic macrolide separating from its class co-members and clustering with two other basic antibiotics, bleomycin and phleomycin. Azithromycin interacts with and crosses the OM of Gram-negative bacteria distinctly to other macrolides^{17,19}, and has also different binding kinetics to the peptide exit tunnel of the 50S ribosomal subunit²⁰. For drugs with unknown or less-well defined targets, clustering hinted towards possible MoA's. Among them, we selected the flavoring compound vanillin, which clusters together with the structurally related acetylsalicylic acid (aspirin). Salicylate and aspirin induce the expression of the major efflux pump in enterobacteria, AcrAB-TolC via binding and inactivating the transcriptional repressor MarR²¹ (Fig. 3a). Consistent with a similar action, vanillin treatment increased AcrA protein levels in *E. coli*, due to *marA* overexpression (Fig. 3b-c). Higher AcrA levels upon vanillin or aspirin treatment led to higher chloramphenicol and ciprofloxacin MICs (Fig. 3d-e). As previously reported for salicylate²², vanillin exerts an additional minor effect on drug resistance in a MarR/A-independent manner, presumably via the MarA homologue, Rob (Fig. 3c-e).

To test whether detected interactions are relevant for resistant isolates, we selected seven strong and conserved synergies, comprising antibiotics, human-targeted drugs or food additives, and assessed their efficacy against six MDR and XDR *E. coli* and *Klebsiella pneumoniae* clinical isolates. All strains were recovered from infected patients, belonging to successfully spread clonal lineages harboring extended spectrum β -lactamase (ESBL) resistance and various highly prevalent carbapenemases^{23,24}. One *K. pneumoniae* strain (929) is also resistant to the last-resort antibiotic, colistin, due to a chromosomal mutation (Supplementary Table 4). All drug pairs acted synergistically in most of the strains tested

(Fig. 4a, ED Fig. 11a). We further tested colistin-clarithromycin and spectinomycin-vanillin, with an established infection model for evaluating antibacterial activity, that of the greater wax moth, *Galleria mellonella*. Both combinations acted also synergistically *in vivo* by increasing *Galleria mellonella* survival during infection (Fig. 4b & ED Fig. 11b).

The strongest of these synergies is between colistin and different macrolides (Fig. 4, ED Fig. 11). Although other polymyxins are known to help macrolides cross the OM of Gram-negative bacteria¹⁷, this particular synergy occurred at low colistin concentration (< 0.3 µg/ml) and was active even for the intrinsically colistin-resistant strain (Fig. 4, *K. pneumoniae* 929), implying that macrolides may also potentiate colistin's action via a yet unknown mechanism. Similar resensitization of colistin-resistant pathogens to colistin by macrolides was recently reported for plasmid-borne colistin resistance²⁵, indicating that this synergy is independent of the resistance mechanism. In addition to antibiotic pairs, combinations of human-targeted drugs or food additives with antibiotics were also effective against MDR isolates, even if former did not have antibacterial activity on their own (ED Fig. 11).

Finally, vanillin potentiated the activity of spectinomycin in *E. coli* MDR isolates. This was intriguing, since vanillin antagonizes many other drugs, including other aminoglycosides (Supplementary Table 2). We confirmed that this interaction is specific to spectinomycin and vanillin, and not to other aminoglycosides or aspirin, and thus independent of the vanillin effect on AcrAB-TolC (ED Fig. 12a-c). We then probed a genome-wide *E. coli* gene knockout library²⁶ to identify mutants that abrogate the vanillin-spectinomycin interaction, but do not influence the amikacin (another aminoglycoside)-vanillin interaction. One of the top hits was *mdfA*, which encodes for a Major Facilitator Superfamily (MFS) transporter, exporting both electrogenic and electroneutral compounds²⁷ (ED Fig. 12c). Consistent with MdfA modulating spectinomycin uptake, $\Delta mdfA$ cells were more resistant to spectinomycin and not responsive to vanillin (ED Fig. 12d), whereas cells overexpressing *mdfA* were more sensitive to spectinomycin (ED Fig. 12e, not visible at the MIC level in ED Fig. 12d), as previously reported²⁸, with vanillin further exacerbating this effect (ED Fig. 12d). Vanillin addition also increased the intracellular spectinomycin concentration in an *mdfA*-dependent manner (ED Fig. 12e). At this point, it is unclear how MdfA, which is known to export compounds out of the cell, facilitates spectinomycin import in the cell. However, the phylogenetic occurrence of *mdfA* is concordant with the species-specificity of this interaction, as we detected the synergy in *E. coli* and *S. Typhimurium*, but not in the phylogenetically more distant, *P. aeruginosa* and *K. pneumoniae* isolates, which lack *mdfA*. This synergy underlines the importance of exploring the role of food additives in combinatorial therapies⁵.

In summary, we generated a comprehensive resource of pairwise drug combinations in Gram-negative bacteria, illuminating key principles of drug-drug interactions and providing a framework for assessing their conservation across organisms or individuals (see Supplementary Discussion). Such information can nucleate equivalent screens in other microbes, studies investigating the underlying mechanism of pairwise drug combinations^{11,15,29} and computational predictions of their outcomes^{30,31}. Moreover, some of the general principles of drug-drug interactions may hold true beyond anti-infectives and microbes³². For antibacterial drug therapies, our study highlights the promise that non-antibiotic drugs hold as adjuvants, offers a new path for narrow spectrum therapies and effective synergies against MDR clinical isolates (see Supplementary Discussion). Further experimentation is required to address whether such synergies have clinical relevance.

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Author contributions

ARB and ATy conceived and designed the study. ARB, ATe and JB performed the screen; ARB, ATe and NN the validation screen; and ARB, MB, AM, JS, SB, MZ and JZB the mechanistic follow-up work. SG characterized the clinical isolates. ARB, ATe and SF performed the clinical isolate checkerboards, and EH and SG the *G. mellonella* infection experiments. ARB analyzed all data. BP, FB, SG and ATy supervised different parts of this study; BE, MS and PB provided advice. ARB and ATy wrote the paper with input from MS, PB and SG. All authors approved the final version.

Figure legends

Figure 1: Principles of drug-drug interaction networks. **a)** Antagonism is more prevalent than synergy. Fraction of observed over detectable interactions in the 6 strains. We detect more antagonistic (1354) than synergistic (1230) interactions, although our ability to detect antagonisms is lower than synergy: 12,778 versus 16,920 combinations. **b & d)** Drug-drug interaction networks in *E. coli*. Nodes represent either drug categories (**b**) or drugs grouped according to the general cellular process they target (**d**). Node color is as ED Fig. 1a and node size reflects the number of drugs within category. Edges represent synergy (blue) and antagonism (orange); thickness reflects number of interactions. Interactions between drugs of the same category/general cellular target are represented by self-interacting edges. Conserved interactions, including weak, are presented. **c & e)** Antagonisms occur almost exclusively between drugs belonging to different categories (**c**) or targeting different cellular processes (**e**), whereas synergies are also abundant between drugs within the same category (**c**) or targeting the same process (**e**). Quantification and Chi-squared test p-values from *E. coli* drug-drug interactions are shown in **b** and **d**, respectively.

Figure 2: Drug-drug interaction conservation. **a)** Drug-drug interactions are conserved in *E. coli*. Scatter plot of interaction scores from the two *E. coli* strains; significant interactions for at least one of the strains are shown. Dark blue: strong and conserved interactions in both strains; light blue: strong interactions in one strain and concordant behavior in other (weak and conserved); grey: interactions occurring exclusively in one strain or conflicting between strains (non-conserved). R denotes the Pearson correlation, n the number interactions plotted. **b)** Drug-drug interactions are highly conserved within all three species. Colors as in **a**; non-comparable refers to combinations that have significantly different single drug dose responses between strains (Methods). **c)** Drug-drug interactions are largely species-specific; n = total number of interactions; n_c = conflicting interactions between species (synergy in one species, antagonism in other), not accounted for in Venn diagram. **d)** Synergies are more conserved than antagonisms. Mosaic plots and Chi-squared test p-values show the quantification of synergy and antagonism among conserved (fully and partially) and non-conserved interactions between species.

Figure 3: Vanillin induces a multi-antibiotic-resistance (*mar*) phenotype. **a)** Vanillin and aspirin (acetylsalicylic acid) have similar drug-drug interaction profiles (see ED Fig. 10), suggesting similar MoA's. A schematic representation of the *mar* response induction via deactivation of the MarR repressor by salicylate/aspirin²¹ is illustrated. **b)** Vanillin increases AcrA levels in a *marA*-dependent manner. A representative immunoblot of exponentially

growing cells, untreated or after treatment with vanillin (150µg/ml) or aspirin (500µg/ml) is shown - loading controlled by cell density and constitutively expressed RecA. Barplots depict AcrA protein level quantification (all western blots available in Supplementary Fig. 1); c) *marA* expression levels upon vanillin (150µg/ml) or aspirin (500µg/ml) treatment are stronger in wildtype than in $\Delta marR$ mutant. Expression is measured by RT-qPCR and normalized to no-drug treatment in wildtype; **d & e**) Vanillin (150 µg/ml) and aspirin (500µg/ml) increase the MIC of chloramphenicol (**d**) or ciprofloxacin (**e**). Antagonism is weaker and abolished in $\Delta marA$ and $\Delta acrA$ mutants, respectively. n = number of independent biological replicates and error bars depict standard deviation (**b-e**)

Figure 4: Potent synergistic combinations against Gram-negative MDR clinical isolates. **a)** *In vitro* synergies, shown as 8x8 checkerboards, for 3 MDR stains (more strains and synergies in ED Fig. 11). One of two biological replicates is shown. **b)** Drug synergies against the same MDR strains in the *Galleria mellonella* infection model (see also ED Fig. 11). Larvae were infected by *E. coli* and *K. pneumoniae* MDR isolates (10^6 and 10^4 CFU, respectively) and left untreated, or treated with single drugs or combination. % larvae survival was monitored at indicated intervals after infection – n=10 larvae per treatment. Here is shown the average of 3 biological replicates; error bars depict standard deviation.

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Methods

Strains, plasmids and drugs

For each of the three Gram-negative species profiled in this study, we used two broadly-used and sequenced strains: *Escherichia coli* K-12 BW25113 and O8 IA11, *Salmonella enterica* serovar Typhimurium LT2 and 14028s, *Pseudomonas aeruginosa* PAO1 and PA14. To validate selected synergies, we profiled 6 MDR clinical *Enterobacteriaceae* isolates recovered from human patient specimens: *E. coli* 124, 1027, 1334 and *Klebsiella pneumoniae* 718, 929 and 980 (see Supplementary Table 4 for details on antibiotic resistance determinants).

All mutants used in this study were made using the *E. coli* Keio Knockout Collection²⁶ - after PCR-confirming and retransducing the mutation to wildtype BW25113 with the P1 phage (Supplementary Table 5). The kanamycin resistance cassettes was excised when necessary using the plasmid pCP20³³. The plasmid used for *mdfA* overexpression was obtained from the mobile *E. coli* ORF library³⁴.

Drugs used in this study were purchased from Sigma Aldrich, except for metformin hydrochloride (TCI Chemicals), clindamycin and bleomycin (Applichem), CHIR-090 (MedChemtronica) and vanillin (Roth). Stocks were prepared according to supplier recommendations (preferably dissolved in water) and kept in the dark at -30°C until arrayed into the plates.

Minimal Inhibitory Concentration (MIC) calculation

We defined MIC as the lowest concentration required to inhibit growth of a microorganism after 8 hours of incubation in Lysogeny Broth (LB) at 37°C with shaking (384 wells plates, starting OD_{595nm} 0.01). MICs of all drugs were computed using a logistic fit of growth (OD_{595nm} for 8h) over 2-fold serial dilutions of the antibiotic concentrations for all strains used for the high-throughput screening and follow-up experiments.

High-throughput screening of pairwise drug interactions

For all drug combination experiments, drugs were diluted in LB to the appropriate working concentrations in transparent 384-well plates (Greiner BioOne GmbH), with each well containing 30µl in total. After the addition of drugs, cells were inoculated at initial OD_{595nm} ~0.01 from an overnight culture. The same inoculum size was used for all strains. All liquid handling (drug addition, cell mixing) was done with a Biomek FX liquid handler (Beckman Coulter). Plates were sealed with breathable membranes (Breathe-Easy®) and incubated at

37°C in a humidity-saturated incubator (Cytomat 2, Thermo Scientific) with continuous shaking and without lids to avoid condensation. OD_{595nm} was measured every 40 min for 12 hours in a Filtermax F5 multimode plate reader (Molecular Devices).

A flowchart of the experimental and analytical pipeline is shown in ED Fig. 2a. Data analysis was implemented with R and networks were created with Cytoscape³⁵.

Experimental Pipeline

The drug-drug interaction screen was performed using 4x4 checkerboards. 62 drugs were arrayed in 384 well plates with the different concentrations in duplicates (array drugs). Each plate contained 12 randomly distributed wells without arrayed drug: 9 wells containing only the query drug, and 3 wells without any drug. One query drug at a single concentration was added in all wells of the 384-well plate, except for the 3 control wells. All drugs were queried once per concentration, occasionally twice. We used 78 drugs as query in *E. coli* and *S. Typhimurium*, and 76 in *P. aeruginosa*. In total 79 query drugs were screened, out of which 75 were common for all three species (Supplementary Table 1). The 62 array drugs were a subset of the 79 query drugs. The same drug concentrations were used in both query and array drugs (Supplementary Table 1). Three drug concentrations (2-fold dilution series) were selected based on the MIC curves, tailored to the strain and drug. We targeted for nearly full, moderate, and mild/no growth inhibition –on average, corresponding to 50-100%, 25-50% and 0-25% of the MIC, respectively. The highest drug concentration and the lowest fitness obtained per single drug, are listed in Supplementary Table 1. For drugs that do not inhibit growth on their own, we selected concentrations according to sensitivity of other strains/species or to their use in clinics or for research. *E. coli* and *S. Typhimurium* exhibited largely similar single drug dose responses within species, thus the same drug concentrations were used for both strains of each species. For *P. aeruginosa*, MICs often differed by several fold, thus drug concentrations were adjusted between the two strains (Supplementary Table 1).

Growth curves analysis

The Gompertz model was fitted to all growth curves (when growth was observed) by using the R package *grofit* version 1.1.1-1 for noise reduction. Quality of fit was assessed by Pearson correlation (R), which was > 0.95 for ~95% of all growth curves. R < 0.95 was indicative of either non-sigmoidal-shaped growth curves, typical of some drugs such as fosfomycin, or of highly noisy data. In the first case, the original data was kept for further analysis. In the second case, noisy data was removed from further analysis. Plate effects were corrected by fitting a polynomial to the median growth of each row and column.

Background signal from LB was removed by subtracting the median curve of the non-growing wells from the same plate. These were wells in which either the single or the double drug treatments fully inhibited growth; each plate contained at least three such wells. Data was processed per strain and per batch to correct for systematic effects.

Fitness estimation

We used a single time-point OD_{595nm} measurement (growth) for assessing fitness. This corresponded to the transition to stationary phase for cells grown without perturbation, as this allows us to capture the effect of drugs on lag-phase, growth rate or maximum growth. Thus, we used OD_{595nm} at 8 hours for *E. coli* BW25113 and both *P. aeruginosa* strains, at 7 hours for the fast-growers *E. coli* iAi1 and *S. Typhimurium* 14028s, and at 9 hours for the slower growing *S. Typhimurium* LT2.

We used the Bliss model to assess interactions, as it can accommodate drugs that have no effect alone, but potentiate the activity of others (adjuvants)³⁶. This feature is especially relevant here, since we probed intrinsically antibiotic-resistant microbes (*P. aeruginosa* and MDR clinical isolates), and human-targeted drugs or food additives lacking antibacterial activity. According to the Bliss independence model³⁷ and assuming that drug-drug interactions are rare, for most drug combinations the fitness of arrayed drugs (f_a) equals the fitness in the presence of both drugs (f_{aq}) divided by the fitness of the query drug alone (f_q):

$$\varepsilon = f_{aq} - f_a * f_q \quad (\text{Eq. 1})$$

$$\text{if } \varepsilon = 0$$

$$f_a = \frac{f_{aq}}{f_q} \Leftrightarrow f_a = \frac{g_{aq}/g_0}{g_q/g_0} \Leftrightarrow f_a = \frac{g_{aq}}{g_q} \quad (\text{Eq. 2})$$

where ε denotes the Bliss score, f denotes fitness, g denotes growth, a denotes an arrayed drug, q denotes a query drug and 0 denotes no drug. The fitness in the presence of both drugs (f_{aq}) was calculated by dividing the growth in the presence of both drugs (g_{aq}) by the median of the growth of drug-free wells from the same plate (g_0). The fitness of the single query drugs (f_q) was obtained by dividing the top 5% growing wells across each batch by the median of the growth of drug-free wells of each plate (g_0). This metric is more robust to experimental errors than using only the 9 wells containing the query drug alone. Nevertheless, both estimators for f_q yield very similar results (Pearson correlation = 0.98). In line with Eq. 2, the fitness of arrayed drugs (f_a) was estimated by the slope of the line of best fit between g_{aq} and g_q across all plates (query drugs) within a batch:

$$\begin{bmatrix} g_{q1} \\ \vdots \\ g_{qn} \end{bmatrix}_{n \times 1} \cdot f_{am} = \begin{bmatrix} g_{amq1} \\ \vdots \\ g_{amqn} \end{bmatrix}_{n \times 1}, \quad 1 \leq m \leq nr \text{ arrayed drugs} \quad (\text{Eq. 3})$$

for given array drug m (a_m) across n query drugs q within a batch (ED Fig. 2b).

For array drugs with Pearson correlation between g_{aq} and g_q below 0.7 f_a was estimated using only the query drugs corresponding to the interquartile range of g_{aq}/g_q (minimum $n = 18$ query drugs, ED Fig. 2b). Wells where r was still below 0.7, even after restricting the number of plates, were removed from further analysis due to high noise (~2%). For wells exhibiting no growth for > 75% of the plates within a batch f_a was deemed as zero.

Interaction scores

Bliss independence

Bliss scores (ε) were calculated for each well as described above (Eq. 1). At least 3 x 3 drug concentrations x 2 (duplicates) x 2 (query and array drugs) = 36, or 18 (drugs used only as query) scores were obtained per drug pair. Drug-drug interactions were inferred based on the Bliss independence model in three steps: a) strong interactions based on complete ε distributions, b) strong interactions based on ε distributions restricted to relevant drug concentrations and c) weak and conserved interactions within species. Cross-species comparison, drug-drug interaction networks and monochromaticity analysis shown in this study include all drug-drug interactions.

a) Strong drug-drug interactions based on complete ε distributions

Strong drug-drug interactions were statistically assigned using a re-sampling approach. 10,000 repetitions of a two-sided Wilcoxon rank-sum test (per drug pair, per strain) were performed, in order sample a representative set of ε for a given strain. For every repetition, the ε distribution of a given combination was compared to a ε distribution of the same size randomly sampled from the complete ε set for a given strain. Permutation p-values were calculated as follows:

$$p = \frac{\sum_{n=1}^N (p_n > 0.1) + 1}{N + 1} \quad (\text{Eq. 4})$$

where N is the total number of repetitions (10,000) and p_n is the p-value of the Wilcoxon rank-sum test obtained for the n^{th} repetitions. Strong drug-drug interactions were assigned to those drug pairs simultaneously satisfying two criteria: i) 1st or 3rd quartile of the ε distribution below -0.1 or higher than 0.1, for synergies or antagonisms respectively, and ii) $p < 0.05$ (after correcting for multiple testing, Benjamini-Hochberg). Only one-sided drug interactions were taken into account, thus those few interactions satisfying the criteria concurrently for synergy and antagonism were re-assigned as neutral (only $n=1$ for $\tilde{\varepsilon} > |0.1|$). The highest

absolute ε value between 1st and 3rd quartile was used as single interaction score ($\tilde{\varepsilon}$) to reflect the strength of the drug-drug interactions.

b) Strong drug-drug interactions based on ε distributions restricted to relevant drug concentrations

Because drug interactions are concentration dependent, the same statistical procedure was repeated after restricting the drug concentration ratios to those relevant for either synergy or antagonism. This constraint was added by excluding ε values corresponding to concentration ratios where the expected fitness (product of the fitness on single drugs, $f_a \cdot f_b$) was below 0.2 for synergy and above 0.8 for antagonism – blind spots for either interaction type (ED Fig. 3d). These interactions are described by their p-value and $\tilde{\varepsilon}$ obtained with restricted drug concentration ratios. Although most interactions were detected based on both full and restricted ε distributions, each of the different methods had uniquely identified interactions (ED Fig. 4c). With the expected fitness cutoff of 0.2, we identified the highest number of strong interactions (1950) with 90 uniquely identified interactions from full ε distributions and 379 from restricted (see also sensitivity analysis).

Restricting ε values based on expected fitness also allows defining whether synergy or antagonism is detectable for any given drug pair. No significant p-value was found for drug pairs with less than 5 ε scores within the relevant expected fitness space, as their sample size is insufficient. Synergy and antagonism could not be detected for 1% and 25% of all drug combinations, respectively.

c) Weak and conserved drug-drug interactions within species

For drug pairs with a strong drug-drug interaction in only one of the two strains per species, the criteria for assigning interactions for the second strain was relaxed to $|\tilde{\varepsilon}_{second\ strain}| > 0.06$, provided that the interaction sign was the same. Interactions assigned with this approach are termed weak and conserved.

Loewe Additivity

For combinations between β -lactams for which high-resolution 8x8 checkerboards with sufficient growth inhibition was available in the validation dataset, Loewe additivity³⁸ was used to confirm the interactions. Drug-drug interactions were inferred by the shape of the isoboles (lines of equal growth) in two-dimensional drug concentration plots. Unless stated otherwise, all isoboles correspond to 50% growth inhibition (IC_{50}) and were obtained by fitting a logistic model – with lines representing isoboles and dots IC_{50} interpolated

concentrations. To interpolate IC_{50} concentrations (or other $IC_{n\%}$), a logistic model was used to fit the growth for each concentration of the first drug across different concentrations of the second drug. The null-hypothesis of this model is represented by the additivity line: a linear isobole connecting equal individual IC's by the two drugs.

Sensitivity analysis

We confirmed the adequacy of the main statistical parameters used to assign interactions by conducting a sensitivity analysis. Several expected fitness ($f_a * f_b$) cutoffs were tested, while keeping the other parameters constant (ED Fig. 4c). The added value of restricting the ε distributions to relevant drug concentrations (based on expected fitness) was strongly supported by the proportion of strong drug-drug interactions found exclusively using this criterion (~19% with our selected cutoff). The selected cutoff (0.2; disregarding wells with $f_a * f_b < 0.2$ for synergies and with $f_a * f_b > 0.8$ for antagonisms) resulted in the largest number of total interactions assigned, and the highest precision (91%) and recall (74%) after benchmarking against the validation dataset (ED Fig. 4c).

The suitability of the thresholds applied to define strong ($|\tilde{\varepsilon}| > 0.1$) and weak ($|\tilde{\varepsilon}| > 0.06$) interactions was assessed by their impact on the true and false positive rates (TPR and FPR respectively, ED Fig. 4d). A threshold of $|\tilde{\varepsilon}| > 0.1$ is beneficial, as it imposes a minimum strength to assign interactions. 0.1 corresponds to ~3 times the median of the 1st and 3rd quartiles across all ε distributions (ED Fig. 2c). Lowering this threshold results in lower TPR, because several drug pairs are reassigned to neutral due to ambiguity in calling interaction (we do not allow interactions being both a synergy and an antagonism). Increasing this threshold lowers the TPR, because only very strong interactions will be assigned (ED Fig. 4d). Drug-drug interactions are highly conserved within species, exhibiting high correlation of $\tilde{\varepsilon}$ observed for all species (Fig. 2a and ED Fig. 9a-b). This motivated us to relax the interaction strength threshold for the second strain if interaction score $|\tilde{\varepsilon}|$ was above 0.1 in first, dubbing these interactions weak and conserved. Including weak and conserved interactions in our analysis increased the TPR by 15%. Adding a threshold for weak interactions of $|\tilde{\varepsilon}| > 0.06$ (~2 times the median of the 1st and 3rd quartiles of all $\tilde{\varepsilon}$ distributions) is key for maintaining a suitable FPR (ED Fig. 4d).

Benchmarking & clinical isolates checkerboard assays

8x8 checkerboard assays were performed for validating our screen (242 drug combinations - benchmarking dataset, Supplementary Table 3), as well as to test 7 selected synergies against 6 MDR clinical isolates (Fig. 4 & ED Fig. 11). As in the screen, growth was assessed

based on OD_{595nm} at the transition to stationary phase for the no drug controls. The timepoints used in the screen were used again for the validation set, whereas 8 hours were used for all *E. coli* and *K. pneumoniae* MDR isolates. Fitness was calculated by dividing OD_{595nm} after single or double drug treatment by no drug treatment for each individual checkerboard. Bliss scores (ε) were calculated as before, resulting in 49 ε values per drug pair. Drug combinations were analyzed based on ε distributions, after removing wells in which one of the drugs alone and its subsequent combinations with the second drug completely inhibit growth. Antagonism was assigned when the median of the ε distribution was above 0.1 or the 3rd quartile was above 0.15. Similarly, synergies were assigned when the median of the ε distribution was below -0.1 or the 1st quartile was below -0.15. All experiments were done in biological duplicates, and interactions were considered effective when duplicates agreed.

Assessing conservation of drug-drug interactions

Conservation of drug-drug interactions between strains of the same species was assessed by Pearson correlation of the interactions scores, $\tilde{\varepsilon}$. For potentially non-conserved drug-drug interactions, the expected fitness distributions of the two strains were taken into account. When the two distributions were significantly different according to a two-sided Wilcoxon rank-sum test (p-value < 0.05 after BH correction for multiple testing), the drug pairs were deemed as non-comparable between the two strains.

To assess the cross-species conservation of drug-drug interactions, we took into account only drug pairs that were probed in all three species. Drug-drug interactions were defined as being detected within a species, when detected in at least one of the two strains and no change of interaction sign was observed for the other strain. Interactions were then compared across the three species. Cases in which an interaction between drugs changed sign across species (conflicting interactions; ~7% of all interactions -Supplementary Table 2) were excluded from the comparative “across-species” Venn diagram (Fig. 2c). Note that with current analysis a given drug-drug interaction may be conserved across species, but not conserved within the species.

Conservation at the single drug level was defined based on shared resistance and sensitivity (Supplementary Table 1). A strain was considered sensitive to a given drug if one of the drug concentrations resulted in at least 30% growth inhibition. In line with conservation of drug-drug interactions across species, single drug responses are conserved across species when at least one strain of both species has the same sign (sensitive or resistant).

Monochromaticity index

The monochromaticity index (MI) between drug pairs was defined as in Szappanos *et. al*³⁹:

$$\begin{aligned} \text{if } r_{ij} > b, MI_{ij} &= \frac{(r_{ij} - b)}{1 - b} \\ \text{if } r_{ij} = b, MI_{ij} &= 0 \\ \text{if } r_{ij} < b, MI_{ij} &= \frac{(r_{ij} - b)}{b} \end{aligned} \quad (\text{Eq. 5})$$

where r_{ij} denotes the ratio of antagonism to all interactions between drugs from classes i and j , and b denotes the ratio of antagonism to all interactions. We set a minimum of 2 interactions between drugs from classes i and j in order to calculate the MI. MI equals 1 if only antagonisms occur between drugs from classes i and j , and -1 if only synergies occur. MI equals zero if the fraction of antagonism reflects the background ratio b . Both strong and weak drug interactions were taken into account across all species, in order to obtain one MI index per drug category pair.

Assessment of drug combinations in the *Galleria mellonella* infection model

Larvae of the greater wax moth (*Galleria mellonella*) at their final instar larval stage were used as an *in vivo* model to assess efficacy of drug combinations. Larvae were purchased from UK Waxworms (Sheffield, UK) and TZ-Terraristik (Cloppenburg, Germany). Stock solutions of vanillin (in 20% DMSO), spectinomycin (Aqua dest.), colistin (Aqua dest.) and clarithromycin (20% DMSO/0.01% glacial acetic acid) were freshly prepared and diluted in PBS to the required concentration. Drugs and bacterial suspensions were administered by injection of 10 μ L aliquots into the hemocoel via the last left (drugs) and right (antibiotic) proleg using Hamilton precision syringes. Controls included both uninfected larvae, and larvae which were injected into both last prolegs with the solvent used for the drugs. Drug toxicity was pre-evaluated by injection of serial dilutions of either single drugs or drug combination, and drugs were used at amounts that caused little/no toxicity. Similarly, time-kill curves were generated by inoculating the larvae with 10 μ L of serial diluted bacterial suspensions (1×10^2 to 1×10^7 colony forming units [CFU]) to identify an optimal inoculum. For final experiments, groups of ten larvae were injected per strain/drug combination and placed into Petri dishes and incubated at 37 °C. Larvae were infected with a (sub)lethal dose of 10^6 and 10^4 CFU for *E. coli* and *K. pneumoniae* isolates, respectively, and subsequently injected with the drugs, 1-hour post infection. Larvae survival was monitored at the indicated time points by two observers independently. Each strain/drug combination was evaluated in 3-4 independent experiments.

Cell viability assays and intracellular antibiotic concentration

Ciprofloxacin

Overnight cultures of *E. coli* BW25113 were diluted 1:1,000 into 50 ml LB and grown at 37°C to OD_{595nm} ~0.5. Paraquat (50 µg/ml), Vanillin (150 µg/ml), Benzalkonium (5 µg/ml), Caffeine (200 µg/ml), Doxycycline (0.5 µg/ml), Rifampicin (5 µg/ml), Trimethoprim (5 µg/ml) or Curcumin (100 µg/ml), were added to the cultures and incubated at 37°C for 30 minutes prior to the addition of 2.5 µg/ml final concentration ciprofloxacin. The cultures were incubated at 37°C for 1 hour in the presence of both drugs. Cell viability was determined by counting CFUs after 16 hours incubation of washed cell pellets plated onto drug-free agar petri dishes. Intracellular ciprofloxacin was quantified using liquid chromatography coupled to tandem mass spectrometry (LC-MS/MS), as previously described^{40,41}. Non-washed cell pellets⁴² were directly frozen and lysed with 350 µl of acetonitrile, followed by three freeze-thaw cycles (thawing was performed in an ultrasonic bath for 5 min). Cell debris was pelleted at 16,000 g and the supernatant was filtered through a 0.22 µm syringe filter prior to injection. Chromatographic separation was achieved on a Waters BEH C18 column (2.1 × 50 mm; 1.7 µm) at 40 °C, with a 2 min gradient with flow rate of 0.5 mL/min: (i) 0–0.5 min, 1% mobile phase B; (ii) 0.5–1.2 min, linear gradient from 1 to 95% mobile phase B; (iii) 1.2–1.6 min, 95% mobile phase B; and (iv) 1.6–1.7 min, return to initial conditions (mobile phase A consisted of 0.1% formic acid in water, and mobile phase B consisted of 0.1% formic acid in acetonitrile). Samples were kept at 4 °C until analysis. Sample injection volume was 5 µL. Detection of ciprofloxacin was performed on a Waters Q-ToF premier instrument with electrospray ionization in positive mode. The transition 332>314 was monitored, with cone voltage set at 8 and collision energy set at 20. Intracellular ciprofloxacin was normalized to CFU at the time of ciprofloxacin addition.

Gentamicin

Intracellular gentamicin was quantified by measuring [3H]-gentamicin (1 mCi/ml; Hartmann Analytic Corp.), as previously described⁷. Overnight cultures of *E. coli* MG1655 (the parental strain of BW25113) were diluted 1:1,000 into 5 ml LB and grown to OD_{595nm} ~0.1. [3H]-gentamicin was diluted in cold gentamicin to get a 5 mg/ml (0,1 mCi/ml) stock solution, which was then added to the culture at a final concentration of 5 µg/ml (0,1 µCi/ml), simultaneously with the second drug: Berberine (200 µg/ml), Erythromycin (15 µg/ml), Metformin (13000 µg/ml), Procaine (6000 µg/ml), Loperamide (400 µg/ml), Benzalkonium (5 µg/ml), Rifampicin (5 µg/ml) or Clindamycin (200 µg/ml). Cultures were then incubated at 37°C on a rotary shaker. At 0, 0.5, 1, 1.5 and 2h time-points, 500 µl aliquots were removed

and applied to a 0.45 µm-pore-size HAWP membrane filter (Millipore) pretreated with 1 ml of unlabeled gentamicin (250 µg/ml). Filters were washed with 10 ml of 1.5% NaCl, placed into counting vials, and dried for 30 min at 52°C. 8 ml of liquid scintillation were then added to the dried filters and vials were incubated overnight at room temperature before being counted for 5 min. Gentamicin uptake efficiency is expressed as total accumulation of gentamicin (ng) per 10⁸ cells, and plotted here for the final timepoint (2h). Cell viability was determined by CFUs.

Spectinomycin

Intracellular spectinomycin was quantified by measuring [3H]-spectinomycin (1 µCi/mg; Hartmann Analytic Corp.). Overnight cultures of *E. coli* BW25113 were diluted 1:1,000 into 1 ml LB with and without vanillin (150 µg/ml) and grown to OD_{595nm} ~0.5. 50 µg/ml [3H]-spectinomycin:spectinomycin 1:100 was added and the cultures were incubated for 1 h. Cultures were pelleted, washed twice with PBS with 50 µg/ml non-labeled spectinomycin, re-suspended in 1% SDS and incubated for 20 min at 85°C. The lysate was mixed with 8 ml liquid scintillation (Perkin Elmer ULTIMA Gold) and counted for 1 min using a Perkin Elmer Tri-Carb 2800TR. Measured radioactivity was normalized to cell number as measured by OD_{595nm}.

RNA isolation, cDNA preparation and Quantitative RT-PCR

Overnight cultures of *E. coli* BW25113 and the *marR* deletion mutant were diluted 1:2,000 into 20 ml LB and grown at 37°C to OD_{595nm} ~0.2. Aspirin or vanillin were added to the cultures to 500 and 150 µg/ml final concentration respectively (DMSO was added in the control), followed by a 30 min incubation period at 37°C with agitation. Cells were harvested and RNA extraction was done with the RNeasy Protect Bacteria Mini Kit (Qiagen) according to manufacturer's directions. cDNA was prepared for qRT-PCR using SuperScript™ III Reverse Transcriptase (Thermo Fisher Scientific). *marA* expression levels were estimated by quantitative RT-PCR using SYBR™ Green PCR master mix following the manufacturer's instructions (Thermo Fisher Scientific). Primer sequences for *marA* and *recA* are previously described²⁹. All experiments were conducted in at least three biological replicates, and relative expression levels were estimated according to Livak *et al.*⁴³, using *recA* expression as reference.

Immunoblot analysis for protein quantification

Overnight cultures of *E. coli* BW25113 and the *marA* deletion mutant ($\Delta marA$) were diluted 1:1,000 into 50 ml LB containing 500 µg/ml aspirin, 150 µg/ml vanillin or DMSO (drugs

solvent control), followed by growth with agitation at 37°C to OD_{595nm} ~0.5. Cells were washed in PBS containing corresponding drugs or DMSO, then resuspended to match OD_{595nm} = 1. Cell pellets were resuspended in Laemmli buffer and heated to 95°C for 3 minutes followed by immunoblot analysis with α -AcrA polyclonal antiserum (gift from K.M. Pos) at 1:200,000 dilution. Primary antiserum was detected using anti-rabbit HRP (A0545 Sigma) at 1:5,000 dilution. An anti-RecA antibody (rabbit, ab63797 Abcam) was used for loading control. Pixel densities of bands were quantified using ImageJ. At least four different biological replicates were blotted and summarized by their mean and standard deviation. Each biological replicate was run and blotted at least twice (technical replicates). Relative AcrA levels per biological replicate correspond to the average intensities of the technical replicates.

Screening the *E. coli* Keio Knockout Collection for identifying MoA of drug interactions

The *E. coli* Keio Knockout Collection ²⁶ (two independent clones per mutant) was arrayed in 1536-format in LB agar plates using a Rotor HDA (Singer Instruments) as previously described ²⁹. The growth of each mutant was estimated by colony opacity ⁴⁴ after 13 hours incubation at 37°C in the absence and presence of vanillin (200 μ g/ml), spectinomycin (4 μ g/ml), and their combination. All plates were imaged under controlled lighting conditions (splmager S&P Robotics) using an 18-megapixel Canon Rebel T3i (Canon). Experiments were done in triplicates. Fitness of each mutant was calculating by dividing the growth in condition (vanillin, spectinomycin or both) by the growth in LB, after correcting for outer-frame plate effects ⁴⁴. Bliss scores were calculated as per Eq. 1 per replicate and then averaged (Supplementary Table 7)

Data availability statement

Data supporting the findings of this study are included in this article as supplementary files.

Code availability

The code used for data analysis is available from the corresponding author upon reasonable request.

Extended Data Figures Legends

Extended Data Figure 1: High-throughput profiling of pairwise drug combinations in Gram-negative bacteria. **a)** Drug and species selection for screen. The 79 drugs used in the combinatorial screen are grouped to categories (Supplementary Table 1). Antibacterials are grouped by target with the exception of antibiotic classes for which enough representatives were screened (>2) to form a separate category: β -lactams, macrolides, tetracyclines, fluoroquinolones and aminoglycosides. Classification of human-targeted drugs and food additives is not further refined, because the MoA is unclear for most. A subset of 62 arrayed drugs was profiled against the complete set of 79 drugs in 6 strains. Strains are color coded according to species. Strain colors and abbreviations are used in main and ED figures. **b)** Quantification of drug-drug interactions. Growth was profiled by measuring optical density (OD_{595nm}) over time in the presence of no, single and both drugs. Interactions were defined according to Bliss independence. Significantly lower or higher fitness than expectation ($f_a \cdot f_q$) indicates synergy or antagonism, respectively. Synergy and antagonism were assessed by growth in 4x4 checkerboards (Methods).

Extended Data Figure 2: Data analysis pipeline. **a)** Flowchart of the data analysis pipeline. **b)** Estimating single drug fitness of arrayed drugs. As drug-drug interactions are rare, the slope of the line of best fit between g_{aq} (growth with double drug) and g_q (growth with query drug alone – deduced from average of the top 5% growing wells across plates) across n_p query drugs (plates) corresponds to a proxy of the fitness of the arrayed drug alone, f_a (Methods, Eq 3). R denotes the Pearson correlation coefficient between g_{aq} and g_q across n_p plates. Well A9 from *E. coli* BW25113 containing 3 μ g/ml spectinomycin is shown as an example of arrayed drugs with several interactions; several query drugs (plates) deviate from the expected fitness (light grey points), therefore only half of the plates corresponding to the interquartile range of f_a were used to estimate f_a . **c)** Density distributions of quartiles 1, 2 and 3 of Bliss scores (ε) distributions for *E. coli*. Q1, Q2 and Q3 denote the median of quartiles 1, 2 and 3 of ε distributions, respectively. n denotes the number of drug combinations used.

Extended Data Figure 3: Data quality control. **a)** High replicate correlation for single and double drug treatments. Transparent boxplots contain Pearson correlation coefficients between plates of the same batch containing arrayed drugs only (LB was used instead of the second drug). n represents the total number of correlations. Full boxplots contain Pearson correlation coefficients between double drug replicate wells within the same plate, across all plates. n represents the number of wells used for correlation, $n_{max} = (62 \text{ drugs} + 1 \text{ LB}) \times 3$

concentrations = 189. Only wells with median growth above 0.2 were taken into account (see panel **b**). For all box plots the center line, limits, whiskers and points correspond to the median, upper and lower quartiles, 1.5x IQR and outliers, respectively. **b**) Wells with lower median growth have lower replicate correlation. The double drug correlation coefficients used to generate the boxplot from **a** are plotted as function of the median growth of all wells across all plates for *E. coli* iAi1. Wells with overall lower growth (due to strong inhibition of arrayed drug) are less reproducible due to a combination of the lower spread of growth values and the sigmoidal nature of the drug dose response curves. **c**) Drug-drug interactions are rare. Density distributions of all Bliss scores (ε) obtained per strain. **d**) The ability to detect synergies and antagonisms depends on the effects of single drug treatments. Bliss scores (ε) are plotted as function of expected fitness ($f_a * f_b$) for all drug concentration ratios for all combinations in *E. coli* BW (example). Boxplots summarizing both variables are shown besides the axes ($n=99,907$ Bliss scores, center line, limits, whiskers and points correspond to the median, upper and lower quartiles, 1.5x IQR and outliers, respectively.). Blind spots for detecting antagonism and synergy are indicated; they are both based on the expected fitness (see also ED Fig. 4c-d) and thus dependent on the growth of the strain with the single drugs. The number of drug combinations falling in the blind spot for antagonism is larger, due to the number of drugs used in the screen that do not inhibit *E. coli* on their own. **e**) Scatter plot of number of interactions per drug versus the minimum fitness of the drug alone (as obtained in screen, Supplementary Table 1). Strong and weak interactions are represented. n denotes the total number of interactions and R is the Pearson correlation coefficient. Strains are color coded as panels **a** & **c**. **f**) Density distributions of the number of interactions per drug for all strains.

Extended Data Figure 4: Benchmarking. **a**) Validation set is enriched in synergies and antagonisms to assess better true and false positives. Comparison of the interaction fractions between the screen and validation set. Both strong and weak interactions (Fig. 2b) are accounted for the screen tally. **b**) Number of benchmarked interactions per strain. **c & d**) Sensitivity analysis of the statistical thresholds for calling interactions. **c**) Total amount of interactions as function of the expected fitness ($f_a * f_b$) cutoff used for restricting the ε distributions to relevant drug concentrations. Strong drug-drug interactions are classified according to the ε distribution where they were significant: complete distribution only (i.e. all expected fitness wells), relevant wells only (i.e. all wells with $f_a * f_b >$ cutoff for synergies and all wells with $f_a * f_b <$ (1-cutoff) for antagonisms), or in both. Weak drug-drug interactions are independently assigned, and represented in white for completeness. We selected an expected fitness cutoff of 0.2, as it resulted in the largest number of total interactions detected, with the highest precision and recall (91 and 74% respectively) after benchmarking

against the validation dataset. **d)** Receiver operating characteristic (ROC) curve for the screen across different p-value thresholds (two-sided permutation test of Wilcoxon rank-sum) as a unique criterion for assigning interactions. The selected p-value (0.05) for screen threshold is indicated by a grey cross. Sensitivity to additional parameters for calling hits is shown: allowing interactions to be either antagonisms or synergies but not both (1-sided); strong and weak interaction thresholds. True and false positive rates were estimated based on the validation dataset. Precision and recall for the final and best performing set of parameters, are shown: one-sided interactions, $p < 0.05$, $f_a * f_b$ cutoff = 0.2 and $|\varepsilon| > 0.1$ for strong interactions, $|\varepsilon| > 0.06$ for weak interactions. TP, TN, FP and FN stand for True Positives, True Negatives, False Positives and False Negatives, respectively. n indicates the total number of benchmarked drug combinations (Supplementary Table 3). **e)** Synergies between β -lactams according to Loewe additivity interaction model. The results of 8x8 checkerboards for 3 combinations between β -lactams in 4 strains are shown. The grey line in each plot represents null hypothesis in the Loewe additivity model, whereas the black line corresponds to the IC_{50} isobole, estimated by fitting a logistic curve to the interpolated drug concentrations (colored dots, Methods). Piperacillin did not reach 50% growth inhibition in *E. coli*, thus IC_{20} and IC_{40} isoboles were used for the amoxicillin + piperacillin combination in *E. coli* BW and *E. coli* iAi1, respectively.

Extended Data Figure 5: Benchmarking of non-comparable drug-drug interactions. a)

The barplot illustrates the division of benchmarked drug combinations according to their degree of conservation within species. The pie chart shows the proportion of False & True Positive (FP & TP) and False and True Negatives (FN & TN) within non-comparable drug-drug interactions. **b)** Combination of amoxicillin with cefotaxime in *P. aeruginosa*: an example of a non-comparable drug-drug interaction. The results of the screen are presented on the upper box. Bliss scores as function of expected fitness for both strains are presented on the left hand side, while a density distribution of the Bliss scores is shown on the right hand side. n denotes the total number of Bliss scores, Q1 and Q3 indicate the Bliss score for quartiles 1 and 3, respectively. Antagonism was detected only for PAO1 (Q3 > 0.1). PA14 was resistant to both drugs in concentrations screened (upper left panel), rendering detection of antagonism impossible. The benchmarking results indicate that interaction is antagonistic in both strains (lower box), albeit weaker at PA14 and visible mostly at higher concentrations. Color on checkerboard reflects fitness and black dots correspond to drug-ratios where the Bliss score is above 0.1.

Extended Data Figure 6: Benchmarking of weak conserved drug-drug interactions. a)

The barplot illustrates the division of benchmarked drug combinations as in ED Fig. 5a. The pie chart shows the proportion False Positives (FP) and True Positives (TP) within weak conserved interactions. **b)** Combination of doxycycline with amikacin in *S. Typhimurium*: an example of a weak conserved drug-drug interaction. The results of the screen are presented on the upper box. Bliss scores as function of expected fitness for both strains are presented on the left hand side, while a density distribution of the Bliss scores is shown on the right hand side. n denotes the total number of Bliss scores, $Q1$ and $Q3$ indicate the Bliss score for quartiles 1 and 3, respectively. A strong synergy was detected only for ST14028 ($Q1 < -0.1$), and then a weak conserved synergy was assigned afterwards to ST LT2 ($Q1 < -0.06$). The benchmarking results, presented on the box below, confirm that the interaction is synergistic in both strains. Color on checkerboard reflects fitness and black dots correspond to drug-ratios where the Bliss score is below -0.1 .

Extended Data Figure 7: *Salmonella* and *Pseudomonas* drug-drug interaction networks. a & b)

Drug category interaction networks. Nodes represent drug categories according to ED Fig. 1a. Node color/size and edge color/thickness are plotted as in Fig. 1b. Conserved interactions, including weak conserved, are shown here. One of the most well-known and broadly used synergies is that of aminoglycosides and β -lactams⁴⁵. Consistent with its use against *P. aeruginosa* in clinics, we detected multiple strong synergies between specific members of the two antibiotic classes in *P. aeruginosa*, but fewer interactions in the other two species (see also Fig 1b). **c & d)** Drug-drug interactions across cellular processes. Representation as in **a & b**, but drug categories targeting the same general cellular process are grouped here. **e)** Quantification of synergy and antagonism in the networks from **a & b**. Corresponding Chi-squared test p-value shown. As in *E. coli*, antagonism occurs more frequently than synergy and almost exclusively between drugs belonging to different categories in *S. Typhimurium* and *P. aeruginosa*. In *P. aeruginosa*, there are very few interactions occurring between drugs of the same category.

Extended Data Figure 8: Drug antagonisms are often due to decrease in intracellular drug concentrations. a)

Cartoon of possible MoAs for drug-drug interactions that workin via modulation of the intracellular drug concentration. A drug (antagonist; blue) inhibits the uptake or promoting the efflux of another one (black), and thus decreases the intracellular concentration. **b)** Different antagonists (see methods for concentrations) of gentamicin (red – 5 $\mu\text{g/ml}$) and ciprofloxacin (gold – 2.5 $\mu\text{g/ml}$) identified in our screen for *E. coli* BW also rescue the killing effect of the two bactericidal drugs in the same strain or its parental MG1655 (top right and left panels, respectively). With the exception of clindamycin (for

gentamicin) and curcumin (for ciprofloxacin) all other antagonists decrease the intracellular concentration of their interacting drug (bottom panels) – gentamicin detected by using radiolabeled compound and ciprofloxacin with LC-MS/MS (see Methods). The degree of rescue (upper panel) in many cases follows the decrease of intracellular concentration (lower panel), implying that most of these interactions depend to a large extent on modulating the intracellular concentration of the antagonized drug. **c)** Antagonisms are resolved in *E. coli* BW mutants lacking key components controlling the intracellular concentration of the antagonized drug. Aminoglycosides depend on PMF-energized uptake and thus respiratory complexes^{7,46}; ciprofloxacin is effluxed by AcrAB-TolC^{29,47}. For gentamicin, most interactions are resolved when respiration is defected, even the one with clindamycin (not modulating intracellular gentamicin concentration- see panel **(b)**) presumably because MoA and import of aminoglycosides are linked in a positive feedback loop^{7,48}. For ciprofloxacin, antagonisms with paraquat and caffeine are resolved in the Δ *acrA* mutant, implying that both compounds induce the AcrAB-TolC pump (known for paraquat). In contrast, interactions with curcumin, benzalkonium and doxycycline remain largely intact in the Δ *acrA* mutant. First is expected as curcumin does not modulate intracellular ciprofloxacin concentration (see panel **(b)**). In other two cases, other component(s) besides AcrAB-TolC are likely responsible for the altered ciprofloxacin import/export; ciprofloxacin is well known to largely use OmpF to enter the cell⁴⁹. Ciprofloxacin and gentamicin concentrations were adjusted in all strains according to MIC (70% and 100% MIC for ciprofloxacin and gentamicin, respectively; all drug concentrations are listed in Supplementary Table 6). Bliss interaction scores (ϵ) were calculated as in screen. Barplots and error bars in **c** & **d** represent the average and standard deviation, respectively, across *n* independent biological replicates. **d)** Gentamicin and ciprofloxacin antagonism networks for *E. coli* BW. Nodes represent drugs colored according to targeted cellular process (as ED Fig. 1a). Full and dashed edges represent antagonistic drug-drug interactions for which intracellular antibiotic concentration was and was not measured, respectively. Drug interactions that result in decreased intracellular concentration of the antagonized drug are represented by black edges. **e)** Quantification of antagonistic drug-drug interactions from the networks in **(d)**. The bars for fluoroquinolones and aminoglycosides account for an extrapolation of antagonistic interactions to all other members of the two classes, assuming they behave same as ciprofloxacin and gentamicin, respectively.

Extended Data Figure 9: Drug-drug interactions are largely conserved within species and only partially MoA-driven. a & b) Drug-drug interactions are conserved in *S. Typhimurium* (**a**) and *P. aeruginosa* (**b**). Scatter plot of interaction scores in the two strains of

each species; only significant interactions for at least one strain are shown. Colors and grouping as in Fig. 3a. R denotes the Pearson correlation and n the total number interactions plotted. Lower correlation in *P. aeruginosa* is presumably due to fewer and weaker interactions in total. **c)** Drug interaction profiles are phylogeny driven. Clustering of strains based on Pearson correlation of their drug interaction profiles (taking into account all pairwise drug combinations; $n=2759-2883$, depending on the drug). Strains of the same species cluster together, with the two enterobacterial species, *E. coli* and *S. Typhimurium*, behaving more similar to each other than to the phylogenetically more distant *P. aeruginosa*. **d)** Conserved drug-drug interaction network. Nodes represent individual drugs grouped and colored by targeted cellular process (as in ED Fig. 1a). Drug names are represented by 3 letter codes (Supplementary Table 1). Dashed and full edges correspond to conserved interactions between two or three species, respectively. Many of the human-targeted drugs, such as loperamide, verapamil and procaine exhibit a general potentiating effect, similar to that of membrane-targeting drugs, suggesting that they may also facilitate drug uptake or impair efflux, consistent with previous reports on the role of loperamide in *E. coli* and verapamil in *Mycobacterium tuberculosis*^{4,50}. **e)** Monochromaticity between all drug categories. The monochromaticity index (MI) reflects whether interactions between drugs of two categories are more synergistic (MI=-1) or antagonistic (MI=1) than the background proportion of synergy and antagonism. MI equals zero when interactions between two drug categories have the same proportion of synergy and antagonism as all interactions together. (Methods). MI was calculated using all interactions from the 6 strains for all category pairs that had at least 2 interactions. White cells in the heat map correspond to category pairs for which no (or insufficient number of) interactions were observed. **f)** Human-targeted drugs, and LPS or PMF inhibitors are strong and promiscuous adjuvants. Density distributions of the MIs per drug category from panel **e** are shown. n denotes the amount of drugs in category involved in i interactions.

Extended Data Figure 10: Hierarchical clustering of drugs according to their interaction profiles. Rows depict the 75 drugs common to all strains (colored according to drug category – ED Fig. 1a), and columns account for their interactions with other drugs in all six strains tested. The median of the ε distributions, uncentered correlation and average linkage were used for clustering.

Extended Data Figure 11: Active synergies against Gram-negative MDR clinical isolates *in vitro* and in *G. mellonella* infection model. Both human-targeted drugs (lately found to have an extended impact on bacteria⁵¹) and food additives can promote the action of antibiotics in MDR strains, indicating that should be explored more in the future. **a)** Drug

combinations active against MDR *E. coli* and *K. pneumoniae* clinical isolates (related to Fig. 4). Interactions are shown as 8x8 checkerboards and synergies have a black bold border. Drug pairs are the same per line and indicated at the first checkerboard. The species in which interaction was detected in screen are indicated after the last checkerboard. Concentrations increase on equal steps per drug (see legend); only minimal and maximal concentrations are shown for first strain of each species. Apart from colistin, the same concentration ranges were used for all *E. coli* and *K. pneumoniae* MDR strains. One of two replicates is shown. b) Drug synergies against the same MDR strains in the *Galleria mellonella* infection model. Larvae were infected by *E. coli* and *K. pneumoniae* MDR isolates (10^6 and 10^4 CFU, respectively) and left untreated, or treated with single drugs or combination. % larvae survival was monitored at indicated intervals after infection – n=10 larvae per treatment. Here is shown the average of 3-4 biological replicates; error bars depict standard deviation.

Extended Data Figure 12: Mode of Action for the vanillin-spectinomycin synergy. a) Spectinomycin MIC decreases upon addition of 100 $\mu\text{g/ml}$ vanillin in the wildtype *E. coli* BW, as well as single-gene knockouts of members of the AcrAB-TolC efflux pump or its MarA regulator. Thus, the vanillin-spectinomycin synergy is independent of the effect of vanillin on AcrAB-TolC (Fig. 3). **b)** Synergy is specific to vanillin-spectinomycin, as 500 $\mu\text{g/ml}$ of the vanillin-related compound, aspirin antagonizes spectinomycin, increasing the MIC ~ 3 -fold. **c)** Profiling the vanillin-spectinomycin combination in the *E. coli* BW Keio collection ²⁶ to deconvolute its MoA. Violin plots of the drug-drug interaction scores ε of all mutants (n=9216; Methods) are presented for the vanillin-spectinomycin combination (synergy) and as control, for the combination of vanillin with another aminoglycoside, amikacin (antagonism). The interaction scores of the two *mdfA* deletion clones present in the Keio library are indicated by red dots. The vanillin-spectinomycin synergy is lost in the absence of *mdfA*, whereas the vanillin-amikacin antagonism remains unaffected, indicating that the vanillin-spectinomycin synergy depends specifically on MdfA. **d)** Deletion of *mdfA* leads to increased spectinomycin MIC and abolishes the synergy with vanillin, independent of the presence or absence of AcrAB-TolC. Mild overexpression of *mdfA* from a plasmid (*pmdfA* - methods) further enhances the synergy with Vanillin, further decreasing the spectinomycin MIC by ~ 2 -fold (comparing to MIC of combination in wildtype). Thus, MdfA levels are directly correlated to the degree of the spectinomycin-vanillin synergy. **e)** Overexpression of *mdfA* leads to increased spectinomycin sensitivity, even though MIC does not change. The growth of *E. coli* BW and *pmdfA* was measured ($\text{OD}_{595\text{nm}}$ after 8h) over 2-fold serial dilutions of spectinomycin and normalized to the no-drug growth of the corresponding strain (white and

black dots represent the average of n=3 independent biological replicates, error bars represent standard deviation.). Spectinomycin dose response was computed using a logistic fit of the averaged data points (MICs are calculated by fitting individual replicates first and then averaging). Fitted curves are represented by full and dashed lines for *pmdfA* and *E. coli* BW respectively. **f)** Vanillin leads to accumulation of spectinomycin in the cell in an *mdfA*-dependent manner. Intracellular spectinomycin is measured with the tritiated compound (Methods). Barplots and error bars in a, b, d & f represent the average and standard deviation, respectively, across n independent biological replicates.

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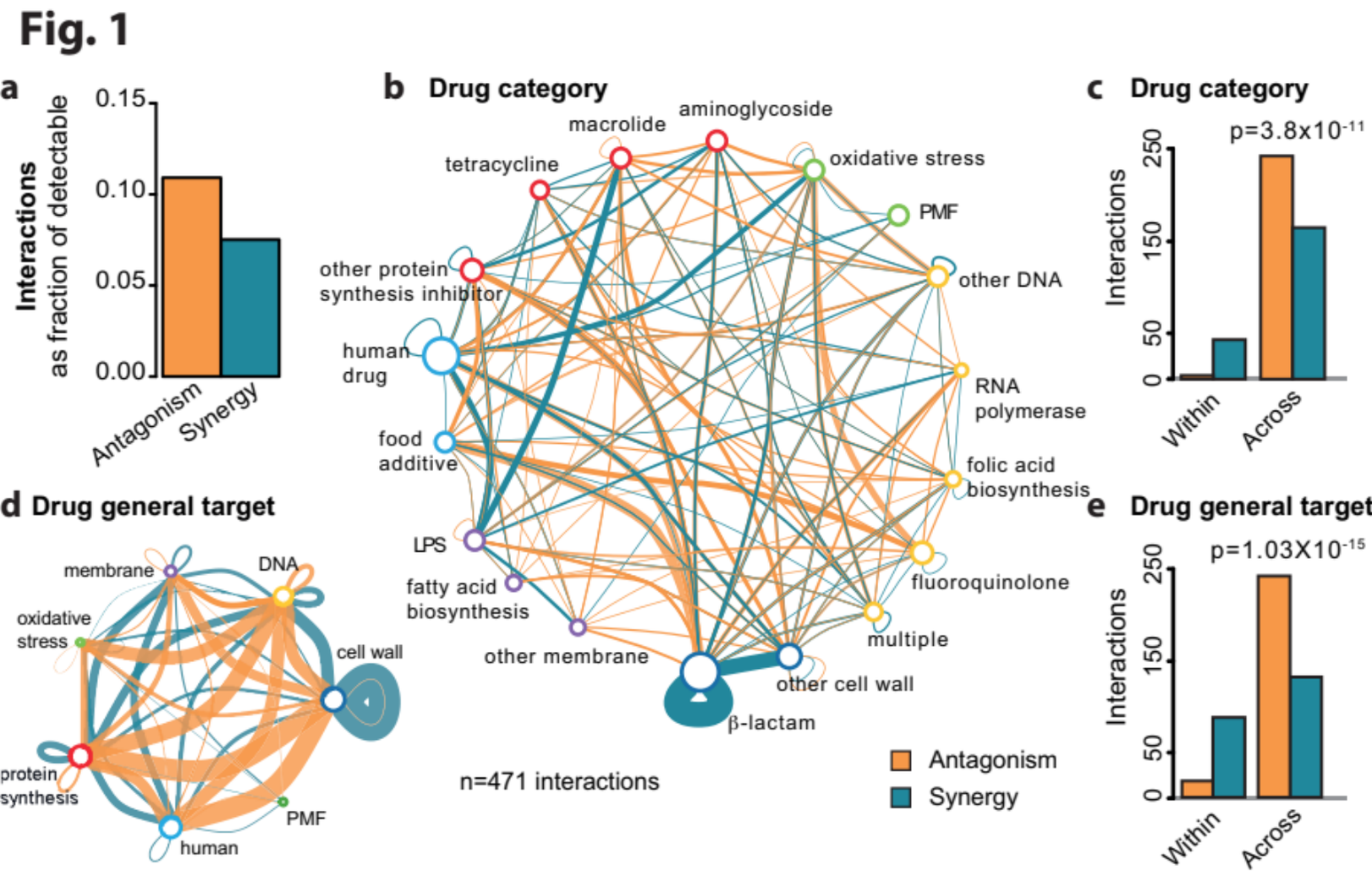


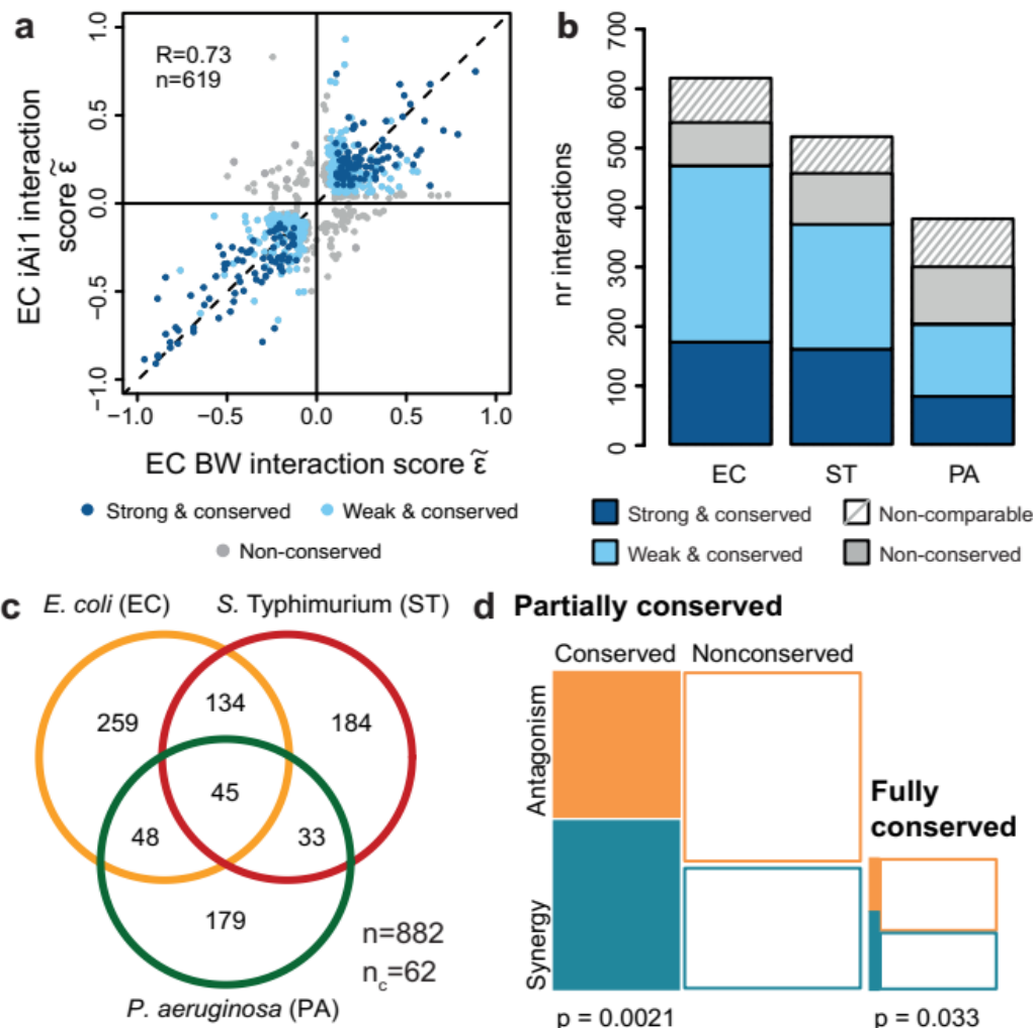
Fig. 2

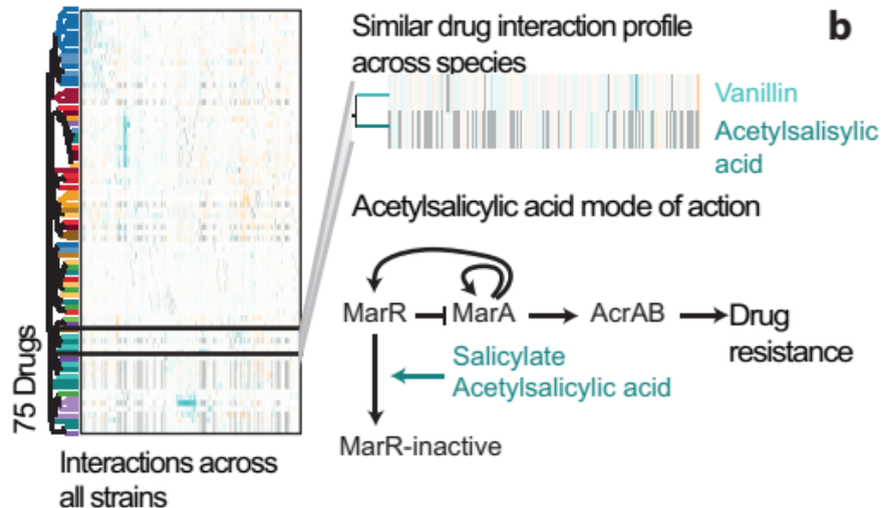
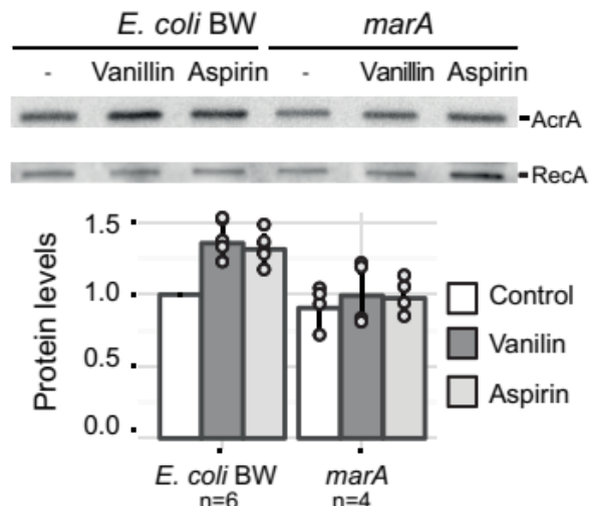
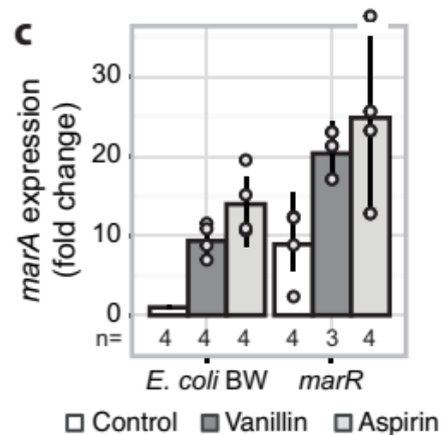
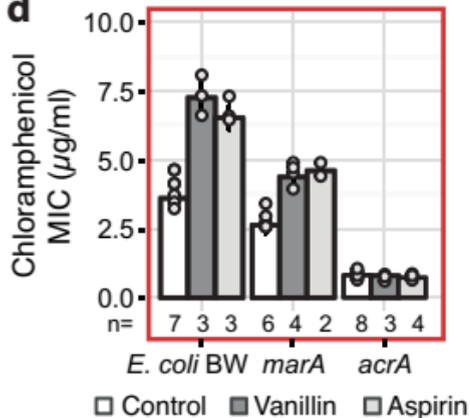
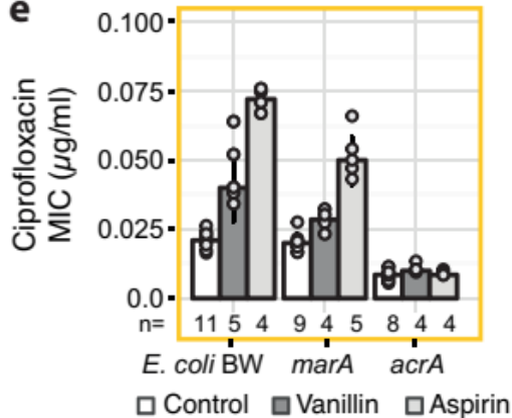
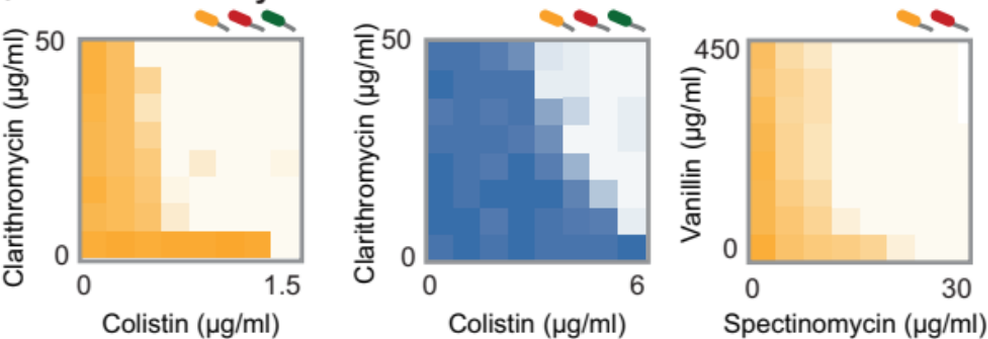
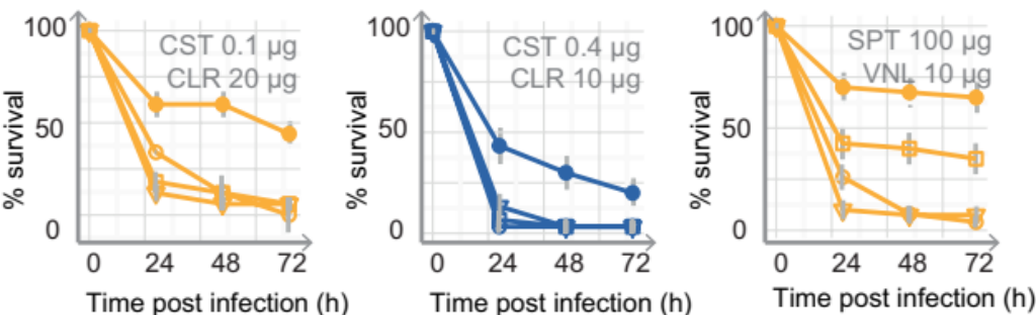
Fig. 3**a****b****c****d****e**

Fig. 4**a In vitro activity****b G. mellonella infections****Legend**

Synergy in screen

 *E. coli*
 *S. typhimurium*
 *P. aeruginosa*

In vitro activity*E. coli* 1334

fitness

0.0 1.0

K. pneumoniae 929*G. mellonella* infections

○ No drug

□ Drug A

▽ Drug B

● Drug A + Drug B