Antibiotics are the most effective tool to treat bacterial infections, but bacteria are rapidly developing both resistance and persistence to these drugs. The use of antibiotics has increased steadily over the past 15 years and is now further increasing due to the ongoing coronavirus pandemic; nearly all patients with severe COVID-19 are receiving antibiotics to prevent and treat secondary bacterial infections. Meanwhile, bacterial pathogens—most prominently, the widespread opportunistic pathogen Staphylococcus aureus—are evolving ways to evade antibiotic killing through various mechanisms of resistance as well as persistence, whereby a small isogenic subpopulation of bacteria undergoes a phenotypic change to temporarily avoid the cidal effect of the antibiotic. Antibiotic persistence leads to a biphasic killing curve, beginning with an early phase of rapid killing of the normal cells and continuing with a later phase of a much slower killing of the persistent cells. Persistence can therefore reduce the ultimate infection clearance efficacy of antibiotic treatments.

As pathogens evolve resistance and persistence to antibiotics, drug combinations are gaining increased attention. Multidrug therapies can have potential benefits in increasing the spectrum of targeted pathogens, preventing the emergence of antibiotic resistance and improving clinical efficacy. Considering the effects of subinhibitory drug combination on bacterial growth (growth-inhibition interactions), laboratory studies have found that the potency of drug mixes increases with the number of co-mixed drugs, yet specific drug pairs and multidrug combinations can substantially deviate from this general trend due to drug synergy and antagonism. Considering the bactericidal effects of drug combination, drug antagonism is commonly revealed, especially between static and cidal drugs. In some cases, drug antagonism can be so strong that the combined effect of co-mixed drugs is weaker not only compared with their expected additive effects but also compared with the effect of one of the drugs alone. Such suppressive, or hyperantagonistic, interactions are typically non-reciprocal—the combined effect of the drugs is weaker than one of the mixed components but not the other (one drug suppresses the effect of the other, but not vice versa). In principle, drug combinations of which the effect is weaker than the effects of either one of the individual drug components are also possible (reciprocal suppression). However, despite the potential clinical implications of such reciprocal suppression—which, while jeopardizing drug efficacy, may counteract resistance—such extreme interactions are not observed in drug combination effects on bacterial growth inhibition or on the early-killing phase.
Although drug combination effects on growth inhibition have been systematically mapped\(^{2-11}\), much less is known about how drugs combine to affect longer-term population clearance and persistence (clearance interactions)\(^{12-14}\). Measuring bacterial survival after prolonged exposures to individual drugs and drug combinations is inherently difficult as it requires the quantification of low densities of surviving cells, and the use of conventional plating techniques limit the number of antimicrobial combinations that can be tested\(^{12}\). Initial studies mapping the effect of specific drug pairs on intermediate (2–4 h) or long-term clearance (8 h) identified synergistic, antagonistic as well as non-reciprocal suppressive interactions\(^{12-14}\). At longer-term survival (1 day exposure), there is even an example of reciprocal suppression\(^{13}\). Owing to the lack of a high-throughput approach to quantify cell viability en masse, a systematic study of such clearance interactions is lacking. It is therefore unclear to what extent these clearance interactions correlate with early-killing and growth-inhibition interactions, how the overall clearance efficacy varies with the number of co-mixed drugs, and whether reciprocal suppression could be identified and used to counteract resistance to the single drugs.

### High-throughput cell-viability assay

To systematically quantify bacterial survival under a range of drug combinations, we adapted a high-throughput cell-viability assay based on miniaturized plating and automated image analysis\(^{14}\). In brief, growing \(S.\) \(aureus\) bacterial cultures were treated with individual drugs or drug mixes. Then, at several time points after drug exposure, the cultures were microplated at a range of dilutions. Automated imaging and image analysis enabled the quantification of the number of microcolonies grown at each microplating spot (up to 100 colonies were detected in each microplating spot in a 96-well format) (Methods and Fig. 1a). Detection sensitivity and reliable colony counting was facilitated by differential labelling and co-culturing of the same \(S.\) \(aureus\) strain with two different fluorescent markers (DsRed and GFP). Using this assay, we comprehensively quantified the effect of pairwise and multidrug mixes on the early- and late-killing phases of a drug-sensitive \(S.\) \(aureus\) strain as well as on the competition among drug-sensitive and drug-resistant strains (Methods; a total of over 25,000 microplating images).

### Drug interactions in early killing

Mapping the growth inhibition and early-killing interactions among all pairwise combinations within a set of 14 diverse antibiotics (Supplementary Table 1) revealed two distinct networks of synergistic, antagonistic and non-reciprocal suppressive interactions (Extended Data Fig. 1). Inspired by previous studies\(^{13}\), we defined for each pair of drugs A and B two directional interaction scores signifying the effect of drug A on drug B and the effect of drug B on drug A. For early killing, directional interactions were defined on the basis of measurements of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (each drug A is applied at a concentration of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (each drug A is applied at a concentration of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (each drug A is applied at a concentration of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (each drug A is applied at a concentration of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (each drug A is applied at a concentration of the decrease in cell viability during 90 min after exposure to each of the 14 drugs when applied individually (Methods and Extended Data Fig. 1c). Measuring bacterial survival during 8-h exposure to cidal concentrations of individual drugs and drug pair combinations and calculating their directional interaction scores (as defined above) (Fig. 1b and Methods). These clearance interactions were not correlated with the growth inhibition interactions but were strongly correlated with the early-killing interactions, indicating that the long-term effect of the drug combinations can be predicted, to some extent, from the early-killing efficacy (Extended Data Fig. 2c,d). However, we also discovered some substantial differences between the early-killing and late-clearance interactions. Indeed, some strong non-reciprocal suppressive early-killing interactions, such as the suppression of the early-killing activity of the cidal drugs by the classically static drugs, are weakened or completely diminished at longer time scales (such as the effect of protein-synthesis inhibitors on the early-killing efficacy of tobramycin and the effect of the trimethoprim on the early-killing efficacy of cell-wall inhibitors or ciprofloxacin) (Fig. 1c and Extended Data Figs. 1f, 4 and 5). By contrast, many other known early-killing non-reciprocal suppressive interactions were strongly enhanced at longer time scales, often developing into reciprocal suppression. In particular, we found that, when cell-wall and classically static protein-synthesis inhibitors are combined, the persisting bacterial population is larger than when any of the drugs are administered alone, even at the same dose and despite using both drugs at cidal concentrations (Fig. 1c and Extended Data Fig. 5). These reciprocal suppressive clearance interactions were robust to changes in drug concentrations, growth environments, incubation time and bacterial physiological state, and were also present in an unrelated \(S.\) \(aureus\) strain with a different genetic background (Methods and Extended Data Fig. 6a–e).

### Multidrug-reduced long-term clearance

Not only was reciprocal suppression common among drug pairs, increasing the number of drugs beyond two further reduced the efficacy of long-term clearance. Following the same procedure used for the drug pairs, we measured the population survival in the presence of all possible multidrug mixes of six commonly used cidal and classically static drugs (clindamycin (CLI), tetracycline (TET), fusidic acid (FUS), meropenem (MER), ciprofloxacin (CPR), oxacillin (OXA); 63 combinations) (Fig. 2a, Extended Data Fig. 7a and Supplementary Table 1). In contrast to the growth inhibitory effects of multidrug combinations, for which potency increases with the number of drugs\(^{15}\), focusing on long-term clearance, we found the opposite behaviour—increasing the number of drugs reduced the efficacy of cell-wall inhibitors or ciprofloxacin) (Fig. 1c and Extended Data Figs. 1f, 4 and 5). Although non-reciprocal suppression was common, reciprocal suppression was not detected in the inhibitory or the early-killing networks (Extended Data Figs. 1e,f and 3–4).

### Drug interactions in long-term clearance

In contrast to early-killing and growth-inhibition interactions, quantifying the effects of antibiotic combinations on long-term clearance revealed pronounced and widespread reciprocal suppression, especially among combinations of cell-wall and classically static protein-synthesis inhibitors (Fig. 1c). To quantify clearance interactions, we measured cell viability during 8-h exposure to cidal concentrations of individual drugs and drug pair combinations and calculated their directional interaction scores (as defined above) (Fig. 1b and Methods). These clearance interactions were not correlated with the growth inhibition interactions but were strongly correlated with the early-killing interactions, indicating that the long-term effect of the drug combinations can be predicted, to some extent, from the early-killing efficacy (Extended Data Fig. 2c,d). However, we also discovered some substantial differences between the early-killing and late-clearance interactions. Indeed, some strong non-reciprocal suppressive early-killing interactions, such as the suppression of the early-killing activity of the cidal drugs by the classically static drugs, are weakened or completely diminished at longer time scales (such as the effect of protein-synthesis inhibitors on the early-killing efficacy of tobramycin and the effect of the trimethoprim on the early-killing efficacy of cell-wall inhibitors or ciprofloxacin) (Fig. 1c and Extended Data Figs. 1f, 4 and 5). By contrast, many other known early-killing non-reciprocal suppressive interactions were strongly enhanced at longer time scales, often developing into reciprocal suppression. In particular, we found that, when cell-wall and classically static protein-synthesis inhibitors are combined, the persisting bacterial population is larger than when any of the drugs are administered alone, even at the same dose and despite using both drugs at cidal concentrations (Fig. 1c and Extended Data Fig. 5). These reciprocal suppressive clearance interactions were robust to changes in drug concentrations, growth environments, incubation time and bacterial physiological state, and were also present in an unrelated \(S.\) \(aureus\) strain with a different genetic background (Methods and Extended Data Fig. 6a–e).

Furthmore, such suppression also appeared in human serum and in an in vivo infection model of the larva of the great wax moth (\(G.\) mel-
number of drugs reduces the overall clearance efficacy of the combinations (Fig. 2a and Extended Data Fig. 7a, b). This phenomenon is unique to long-term clearance interactions; whereas long-term efficacy decreased with the number of drugs, early-killing efficacy did not change across the different multidrug combinations (Extended Data Fig. 7c). The same trend was also observed in an independent set of five additional drugs of diverse classes (cefoxitin (FOX), linezolid (LIN), cefazolin (CEF), minocycline (MIN) and pristinamycin (PRI); Fig. 2b and Extended Data Fig. 8). Furthermore, considering each multidrug set as a combination of two drug mixes (for example, mix ABC is decomposed to AB + C, A + BC, AC + B), we found enrichment of reciprocal suppressions between drug mixes (Fig. 2c and Extended Data Fig. 7d), which was not observed in early-killing interactions (Extended Data Fig. 7e). Separating drug sets by all-cidal and all-static or combined drug mixes...
highlights that this reciprocal suppression is mostly driven by combinations of cidal and classically static drugs (Fig. 2b,c). However, including specific drugs of which the cidal activity is only weakly dependent on cell metabolism strongly synergized the clearance efficacy of all of the other drugs and drug mixes, efficiently eradicating the long-term surviving cell population (persistence-targeting: daptomycin (DAP), mitomycin (MIT)) (Extended Data Fig. 8). The dot colour represents drug combinations that are only CS (cyan), only cidal (purple) or a mix (grey, black cubic fit), and sets including persistence-targeting drugs (green shades; green linear fit). The distribution of the interaction scores among all tested non-overlapping pairs of drug sets (for example, drug mix ABC is decomposed into AB + C, A + BC, AC + B) without (c) and with (d) persistence-targeting drugs. n = 391 (c) and n = 422 (d). The combined effect of the drug mix is shown relative to the minimum (min) and maximum (max) of the effect of the two drug sets (Methods). Drug mix pairs are classified as only cidal (purple), only CS (cyan), mix of all CS and all cidal (pink), pairs of mixed drug sets (mix, grey) or sets including DAP or MIT (green shades).

Selection against antibiotic resistance

Next, we competed drug-resistant and drug-sensitive strains in mixtures of reciprocal suppressive drug pairs, and we found that these interactions can select against strains that are resistant to either of the single drugs. It has been shown that non-reciprocal suppressive interactions can reverse the selective advantage of a strain that is resistant to one of the drugs (if drug A suppresses drug B, the combination of the two drugs selects against resistance to drug A, but not against resistance to drug B)\(^{12,13}\). Extrapolating from these results, it has been hypothesized that, with reciprocal suppression, it might be possible to select against any of the single-drug-resistant mutants\(^ {31} \).
Focusing on the combination of TET and ampicillin (AMP) (a reciprocal suppressive pair for long-term clearance; Fig. 1c), we competed our sensitive ancestral strain with either a strain that is resistant to TET (TET evolved) (Methods, Fig. 3a and Extended Data Fig. 9c,d) or a strain that is resistant to cell-wall inhibitors (methillin-resistant Staphylococcus aureus (MRSA) (Methods, Fig. 3b and Extended Data Fig. 9e–g), or we compared the sensitive strain with and without an inducible β-lactamase (Methods and Extended Data Fig. 9a,b). By contrasting the survival of the drug-sensitive and drug-resistant strains after 8 h of exposure to a two-dimensional concentration gradient of TET and AMP, we found that, under an extended region of the two-drug concentration space, the long-term survival of the resistant strains was much lower compared with the sensitive strain (10-fold to 100-fold) (Extended Data Fig. 9a–g). These selection inversions resulted from the interaction itself; there was no selection against the resistant strains under any of the single treatments. Selection against each of the single-drug-resistant mutants also occurred in competition within human serum (Fig. 3c,e and Extended Data Fig. 9h) and selection against MRSA was even observed within the G. mellonella infection model (in TET–AMP (Fig. 3f) as well as in MIN–AMP (Methods and Extended Data Fig. 9i). However, note that in this in vivo environment, even the single drugs, TET and MIN, showed selection against the MRSA strain (Fig. 3f and Extended Data Fig. 9i; perhaps reflecting suppressive interactions of these drugs with natural antimicrobial peptides in vivo)40. Selection against TET-resistant mutants did not occur within larvae, possibly due to the effect of oxygen depletion on the action of β-lactams39 and due to competition for resources within larvae (Fig. 3d).

Reverse effect of resistance inhibition

Finally, given the ability of suppressive drug pairs to select against AMP resistance, we examined the effect of supplementing these drug combinations with a β-lactamase inhibitor. β-Lactamase inhibitors are commonly used together with β-lactams to increase treatment efficacy against strains carrying the β-lactamase enzyme. We measured the long-term survival of a β-lactamase-mediated AMP-resistant strain (Methods) as a function of increasing doses of avibactam, a known β-lactamase inhibitor, in the presence of three different clear ance-suppressive drug combinations (AMP combined with TET, CLI or FUS; Fig. 4). Consistent with these combinations selecting against AMP resistance, we found that adding avibactam to these clearance-suppressive combinations decreased rather than increased the long-term clearance efficacy against the β-lactam-resistant strain (Fig. 4 and Extended Data Fig. 10a–d). Thus, in the presence of clearance-suppressive drug combinations, a β-lactamase inhibitor jeopardizes, rather than facilitates, treatment efficacy.

Discussion

By systematically quantifying the growth inhibition and killing dynamic of individual drugs and drug combinations at cidal, clinically relevant concentrations, we found that reciprocal suppressive interactions emerge in prolonged treatment efficacy. In contrast to shorter-term drug effects and conventional wisdom, drug efficacy is reduced rather than increased as more drugs are combined. These strong suppressive clearance interactions are suggestive of induced
Fig. 4 | Supplementing clearance suppressive combinations with avibactam, a β-lactamase inhibitor, decreased rather than increased the long-term clearance efficacy against a β-lactamase-carrying strain. a. As expected, the killing efficacy of AMP at fixed dosage (0.5 μg ml⁻¹) is enhanced by the addition of an increasing dosage of avibactam (AVI). b–d. By contrast, the treatment efficacy is reduced rather than enhanced when avibactam is added to a combination of AMP (0.5 μg ml⁻¹) with a fixed dosage of TET (b; 20 μg μl⁻¹), CLI (c: 10 μg ml⁻¹) or AMP (d: 0.4 μg ml⁻¹) with FUS (d; 20 μg ml⁻¹). No AMP controls show that this inverted effect of AVI is mediated by the interaction among AMP and these protein-synthesis inhibitors (grey dashed lines). The error bars represent 95% confidence intervals calculated from the colony counts of one sample by the Poisson's model. Statistical analysis was performed using two tests based on the Poisson-derived confidence intervals; P < 10⁻⁶.

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### Online content

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### PERSPECTIVE

As expected, the killing efficacy of AMP at fixed dosage (0.5 μg ml⁻¹) is enhanced by the addition of an increasing dosage of avibactam (AVI). By contrast, the treatment efficacy is reduced rather than enhanced when avibactam is added to a combination of AMP (0.5 μg ml⁻¹) with a fixed dosage of TET (b; 20 μg μl⁻¹), CLI (c: 10 μg ml⁻¹) or AMP (d: 0.4 μg ml⁻¹) with FUS (d; 20 μg ml⁻¹). No AMP controls show that this inverted effect of AVI is mediated by the interaction among AMP and these protein-synthesis inhibitors (grey dashed lines). The error bars represent 95% confidence intervals calculated from the colony counts of one sample by the Poisson's model. Statistical analysis was performed using two tests based on the Poisson-derived confidence intervals; P < 10⁻⁶.
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Strains and plasmids

The *S. aureus* sp. Rosenbach (ATCC 29213) strain was transformed with either a DsRed (pHC4S, sarAP1_DsRed, CHLR) or a sGFP fluorescent reporter gene (pCM29, sarAP1_sGFP, CHLR) under a constitutive promoter (antibiotic-sensitive *S. aureus*). To test for generality and for counter-selection experiments, the MW2 *S. aureus* strain was similarly transformed with these two plasmids (MRS). A β-lactamase-inducible plasmid was constructed by amplifying the blaZ gene from *Enterococcus faecalis* (ATCC 49757) and cloning by Gibson assembly under the TET-inducible promoter of the *S. aureus* gene.

Data

**Methods**

**Strains and plasmids**

The *S. aureus* sp. Rosenbach (ATCC 29213) strain was transformed with either a DsRed (pHC4S, sarAP1_DsRed, CHLR) or a sGFP fluorescent reporter gene (pCM29, sarAP1_sGFP, CHLR) under a constitutive promoter (antibiotic-sensitive *S. aureus*). To test for generality and for counter-selection experiments, the MW2 *S. aureus* strain was similarly transformed with these two plasmids (MRS). A β-lactamase-inducible plasmid was constructed by amplifying the blaZ gene from *Enterococcus faecalis* (ATCC 49757) and cloning by Gibson assembly under the TET-inducible promoter of the pG+off plasmid. This plasmid was transformed into the sensitive *S. aureus* strain to form the AMP-resistant strain in the presence of 0.05 μM anhydrotetracycline (Fig. 4 and Extended Data Figs. 9a,b and 10a–d). Laboratory evolution was performed according to a previously established protocol to evolve the antibiotic-sensitive *S. aureus* strain to TET.

A resistant clone with the highest TET minimal inhibitory concentration (25-fold increase compared with the ancestral strain) was transformed with either the DsRed or the GFP plasmid and used in the experiments of Fig. 3a,c,d and Extended Data Fig. 10a–d. Fresh antibiotic solutions were prepared from powder stocks (Sigma-Aldrich, Cayman chemical, AdooQ Bioscience) on a monthly basis, filter-sterilized and kept at −20 °C until use.

Quantifying bacterial survival under antibiotic exposure

We adapted a high-throughput cell-viability assay in which bacterial cultures were exposed to single or multiple drugs in 96-well assay plates and their viability across time was quantified by microplating and colony counting. Specifically, these experiments were performed in 6 steps. (1) Preparation of bacterial stocks. Single colonies of the GFP and DsRed fluorescently labelled sensitive *S. aureus* strains were isolated and grown overnight in LB supplemented with 10 μg ml⁻¹ CHL (to maintain the plasmid) at 37 °C, 250 rpm. Overnight cultures were diluted 1:5,000 and grown to an optical density at 600 nm (OD₆₀₀) of 0.25. These cultures were concentrated by centrifugation and small aliquots of 50 μl containing ~2.2 × 10⁶ cells were frozen in glycerol (stored at −80 °C). (2) Preparing assay plates with antibiotics. Deep 96-well plates were filled with 600 μl LB and supplemented with single or multiple antibiotics using an automated digital dispenser (D300e, Tecan). At least two replicate wells were randomly distributed for each condition. (3) Antibiotic exposure. For each biological replicate, frozen cultures of the GFP and DsRed strains were thawed and mixed at a 1:1 ratio, diluted 1:250 in 25 ml LB and incubated for 40 min at 37 °C and 250 rpm. These co-growing bacterial cultures were then aliquoted into the 96-well deep-well antibiotic plates (150 μl culture; total volume of 750 μl) at a low inoculum density of about 10⁻⁴ CFU per ml to minimize the risk of pre-existing resistant mutants and prevent nutrient depletion (Extended Data Fig. 10–m; the growth of the resistant strain was not limited in the supernatant of the FOX regimes that kill the sensitive strain (~1 μg ml⁻¹) but not the resistant strain (~3.8 μg ml⁻¹), especially in the wells in which the starting density of the sensitive strain was initially low (~10⁻¹); this result indicates that nutrient depletion is negligible during our killing assay starting with a 10⁻¹₀ initial population size). This inoculum density follows recommended clinical testing standards for killing assays. No-cell and no-antibiotic wells were designated on each plate to control for contamination (2–5 of the wells in each plate). (4) Microplating assay for quantification of bacterial viability (colony-forming units (CFU)). At fixed time points after antibiotic exposure at 37 °C and 250 rpm (0, 0.25, 0.5, 1 and 1.5 h for the early-killing assay; and 0, 0.5, 1, 2, 4 and 8 h for the long-term clearance assay), small aliquots (10–150 μl) were taken from each well, and fivefold serial diluted for plating. The 8 h time point was the last time point for sampling as the clearance efficacy of many of the single drugs (Extended Data Fig. 5) already reached the detection limit of our assay (1 colony in 7 μl volume plated; ~140 CFU per ml; extending beyond 8 h requires larger plating volumes, which cannot be done in microplates). Large-volume experiments for the two chosen drug pairs showed robustness of the clearance interactions even at 24 h (Extended Data Fig. 6c). For the 8 h clearance assay, in which cell viability declined substantially, we also plated the undiluted culture (dilutions of 5ⁿ, where n = 0–4) and therefore added a wash step by centrifugation and resuspension to remove the drugs (three washes with 800 μl PBS at 4,000 rpm for 10 min). For the killing-assay, in which cell viability was higher, an initial dilution of at least 5₀ was used, precluding the washing step (final dilutions of 5ⁿ × 5ⁿ, where n = 0–2). Small microdrops (7 μl) from each dilution were then carefully microplated (Gilson 96-channel plate master) in one to four technical replicates onto omni single-well (Greiner) plates filled with 45 ml of agar with 0.2× diluted LB and incubated for 24 h at 37 °C. The diluted LB was used to limit the substrate availability on the agar thereby allowing the growth of many small colonies on a relatively small surface area. (5) Imaging. After incubation, the agar plates were imaged for GFP (excitation:470/30; emission: 540/50) and DsRed (excitation: 590/50; emission: 641/75) with a custom-made automated macroscope device. (6) Detection of resistance. At the last time point, 15 μl aliquots from each well of the assay plates were also inoculated into a 96-well plate with 200 μl liquid LB containing, in each well, the same antibiotic concentrations of the assay plates and incubated overnight at 37 °C for 250 rpm. After incubation, the growing cultures (OD₆₀₀ > 0.2) indicating resistant cells, were excluded from further analysis. Overall, our data included ~15,000 microdrops: n = 2 biologically independent strains (DsRed, GFP) measured in 585 drug conditions (25 singles (1×, 2× dose) and 91 pairs), 5 × 36 singles (1×, 2×, 3×, 4×, 5× and 6× dose), 5 × 57 multidrug mixes; 1–5 timepoints; 2 time ranges (short-term and long-term killing), at least 2 independent replicates and 4 to 8 platings (dilutions + technical replicates).

**Automated image analysis**

The number of red and green colonies in each microplating spot was counted using a custom Python script, implementing the following steps. (1) The GFP and DsRed microplating images (one for each microplating spot) were contrast-enhanced and noise-reduced (mean shift pyramid). (2) These enhanced GFP and DsRed images were thresholded (mahotas.otsu), yielding two corresponding binary images. (3) An exact Euclidean distance transform was used to yield a distance matrix of each pixel to its nearest zero pixel (scipy.ndimage.distance_transform_log). (4) Local peaks in the distance matrix (skimage.feature.peak_local_max) were clustered into connected components (nearest neighbours, scipy.ndimage.label) to yield labelled feature arrays. (5) These feature arrays were used as seeds to the watershed algorithm to segment the colonies (skimage.morphology.watershed). (6) Thresholding on the basis of the size and circularity of the colonies was applied to remove very tiny colonies or merged spots. (7) Microplating spots where the colonies merged extensively or with colonies that were too dense to reliably count were designated uncountable (over 100 colonies per spot). (8) For each given culture and time point, the colony counts at dilutions at which there was no uncountable plating were used to evaluate the CFU per ml (total number of colonies in all of the countable dilutions divided by total culture volume plated). Our high-throughput viability assay was highly reproducible, showing good agreement among biological replicates (Extended Data Fig. 10–h). Co-cultures of GFP/DsRed strains enabled us to: (1) increase the sensitivity of the colony-detection algorithm—different colour colonies can be more easily distinguished even if they grow at close proximity; (2) spot low-frequency antibiotic-resistant mutants that manifest as red-only or green-only colonies on the agar spot where two identical strains labelled with different fluorescent tags; (3) conduct competition experiments in which the sensitive and resistant strains were labelled differentially (see below).
Calculating early-killing and clearance interaction scores between drug pairs

Early-killing and long-term clearance directional interaction scores (DIS) among pairwise combinations of the 14 antibiotics were quantified based on the fraction of surviving cells after short (average of the 1 and 1.5 h time points; the killing delay was almost always shorter than half an hour across the different conditions indicating that the average cell survival after 1 and 1.5 h is an effective killing score that combines both the killing rate and drug action delay; Extended Data Fig. 10) and long (8 h time point; the choice of 8 h incubation time is based on the two hour marking ~2 × 10^9 cells were frozen in glycerol (−80 °C). For each independent replicate, frozen aliquots of these cultures were tested for Extended Data Fig. 6). Our high-throughput cell-viability assay was applied with minor modifications: for each biological replicate, the frozen cultures of Gfp and DsRed were thawed and mixed at a 1:1 ratio, diluted 1:250 dilution in 5 ml of the appropriate medium (LB, BHI, TSB, in case of the serum experiment LB was used) in culture tubes and incubated for 20 min at 37 °C and 250 rpm. The cell-viability assay, microplating and automated image analyses were then performed as described above.

Testing the robustness of reciprocal suppression clearance interactions to different media

The clearance interactions between two representative reciprocal suppressive drug combinations (CLI–CLO and FOX–TET) (Fig. 1c and Extended Data Fig. 5) were measured in different media including Luria–Bertani (LB), brain–heart infusion (BHI), tryptic soy broth (TSB) and human serum (Extended Data Fig. 6b). Antibiotic concentrations were adjusted to give a similar clearance efficacy in each condition (source data for Extended Data Fig. 6). Our high-throughput cell-viability assay was applied with minor modifications: for each biological replicate, the frozen cultures of Gfp and DsRed were thawed and mixed at a 1:1 ratio, diluted 1:250 dilution in 5 ml of the appropriate medium (LB, BHI, TSB, in case of the serum experiment LB was used) in culture tubes and incubated for 20 min at 37 °C and 250 rpm. The cell-viability assay, microplating and automated image analyses were then performed as described above.

Testing the robustness of reciprocal suppression clearance interactions to longer incubation times

To compare drug interactions after 8 h and 24 h incubation time, we repeated the cell-viability assay of two representative reciprocal suppressive drug combinations (CLI–CLO; TET–FOX) (Fig. 1c and Extended Data Fig. 5) using a standard plating technique on regular Petri dishes (Extended Data Fig. 6c). Culture tubes were filled with 2 ml LB medium supplemented with single or multiple antibiotics. For each independent replicate, frozen cultures of Gfp and DsRed were thawed and mixed and incubated at a 1:1 ratio, diluted 1:250 in 5 ml LB in a culture tubes and incubated for 15 min at 37 °C and 250 rpm. These co-growing bacterial cultures were then aliquoted into each tube (135 µl culture) at a low inoculum density of about 10^5–10^7. After incubation for 8 h and 24 h at 37 °C and 250 rpm, 400 µl samples were taken, washed three times with 1 ml PBS and resuspended in 200 µl PBS. Then, 100 µl samples were spread from each dilution onto LB agar Petri dishes and incubated for 24 h at 37 °C. The plate imaging and automated colony counting were performed according to the same steps as described above.

Testing the robustness of reciprocal suppression clearance interactions to bacterial physiological state in a plasmid-less strain

The clearance interaction was measured between CLI and CLO starting from two different biologically independent S. aureus cultures in different physiological conditions (Extended Data Fig. 6d; exponential growing and late growing phase). Frozen aliquots of these cultures were prepared by taking two single colonies of the plasmid-less sensitive S. aureus strain, growing overnight in LB, then diluted 1:5,000 and grown to either OD_600 = 0.1 or OD = 0.8. Then, both of these cultures were concentrated by centrifugation and small aliquots of 30 µl containing ~2 × 10^10 cells were frozen in glycerol (~80 °C). For each independent replicate, these frozen cultures were thawed, diluted 1:250 in 2 ml LB and incubated for 15 min at 37 °C and 250 rpm. A standard plating technique was applied (as described above) to measure cell viability after incubation for 24 h in human serum supplemented with the single drugs or the combination. Plates were imaged using the Canon EOS T3i camera followed by manual colony counting.

Clearance efficacy and competition in the G. mellonella infection model

Larvae of the G. mellonella at their final instar stage were used to assess the clearance efficacy of drug combinations. Larvae were either purchased from UK BioSystem Technology (TRUELarvae) or from the Israel Agricultural Research Organization, Volcani Institute (AROLarvae). Antibiotic stock solutions were freshly prepared and diluted in PBS to the required concentration. Drugs and fluorescently tagged S. aureus (~10^7) bacterial suspensions (with chloramphenicol resistance) were
Measuring growth inhibition under 2D concentration gradients

Growth inhibition of all the pairwise combinations of the 14 antibiotics (n = 91; Extended Data Figs. 1a–e and 3) was measured on the sensitive S. aureus (ATCC 29213) according to a previously established experimental protocol. First, for each drug A, a concentration gradient covering its effective range was defined as $D_A(k) = IC_{90A} \times w^{-k}$, where $D_A(k)$ is the dosage of the drug at dilution number $w = (IC_{90A}/IC_{10})^{1/3}$ is the dilution factor, $IC_{10}$ and $IC_{90A}$ are the concentrations of A drug resulting in 10% and 90% inhibition, respectively, and $k$ ranges from 1 to 8 (such that the lowest concentration is $D_A(1) = IC_{10}$ and the highest concentration is $D_A(8) = IC_{90A}$). Then, for each pair of drugs A and B, a 96-well microplate was filled with 150 μL LB, supplemented (D300e concentration is w) resulting in 10% and 90% inhibition, respectively, and $k$ varies from 3 to 8, total of 24 wells), (3) bacteria-free control (2 wells), (4) no-drug control (6 wells). These assay plates were then inoculated with the sensitive S. aureus at 10^6 cells per mL and incubated for 18 h in 37 °C, shaken at 250 rpm. After incubation, the OD600 (BioTek Synergy 2) was measured as the basis for calculating the interaction scores between the two drugs.

Calculating growth-inhibition interaction score

For each pair of the 14 antibiotics, the OD of their 2D drug assay plates was used to quantify directional growth-inhibition interaction scores (DGIS). First, the OD was background-subtracted and normalized to the no-drug control wells. Second, a polynomial surface model was fitted (Loess fit) to these normalized OD values to determine the concentration of each drug leading to 90% growth inhibition ($IC_{90A}$, $IC_{90B}$) as well as the 90% growth inhibition isobole ($IC_{90i}$). To quantify directional growth-inhibition interaction scores, we considered the distance from the origin to the IC90 isobole along two straight lines representing fixed A:B ratios (in IC90 units) of 1:3 for effect of drug A on drug B (A→B) and 3:1 for effect of drug B on drug A (B→A). These two distances were then interpolated relative to the corresponding distances associated with the intersection of these fixed-ratio lines with 4 reference isobole lines representing extreme synergy (no drug, DGIS = 1), Loewe’s dosage additivity (DGIS = 2), antagonism (highest single agent (HSA), DGIS = 3) and strong suppression (2× HSA, DGIS = 4) (Extended Data Figs. 1a–e and 3).

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

All data supporting the findings of this study are provided as Source Data. The raw data files, including the microplating images with the bacterial colonies, are available in a public data repository (https://osf.io/wh62f/). Source data are provided with this paper.

Code availability

Custom script developed for the study was deposited in a public data repository (https://osf.io/wh62f/).

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Author contributions

V.L. and R.K. conceived and designed the study. V.L. performed all of the in vitro experiments. V.L., O.S., R.K. and D.B. designed the experiments in the larvae infection model. V.L. and R.K. performed the experiments in the larvae infection model. V.L. and D.B. analyzed the data and wrote the paper.

Competing interests

The authors declare no competing interests.

Additional information

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Extended Data Fig. 1 | Systematic mapping of pairwise growth-inhibition and early-killing interactions. a–d. Definition of directional growth-inhibition interaction scores. Measuring the growth as the final optical density (OD) in a 2-D gradient of drugs A and B after 18-hour of incubation, we define the concentration of each drug leading to 90% growth inhibition (IC90\textsubscript{A}, IC90\textsubscript{B}) as well as the 90% growth inhibition isobole (IC90 isobole, black line). To quantify bi-directional interactions, we consider the distance from the origin to the IC90 isobole along two straight lines representing fixed A:B ratios (in IC90 units) of 1:3 for effect of drug A on drug B (A→B, dashed orange) and 3:1 for effect of drug B on drug A (B→A, dashed magenta). These two distances are then interpolated relative to the corresponding distances associated with the intersection of these fixed-ratio lines with 4 reference isobole lines representing extreme synergy (red), Loewe’s dosage additivity (grey), antagonism (Highest Single Agent, HSA, green) and strong suppression (blue). The four panels illustrate four representative examples of synergism (a), antagonism (b), non-reciprocal suppression (c) and reciprocal suppression (d).

e. Measurements of the directional growth-inhibition interaction scores for pairwise combinations of 14 antibiotics. Upper (lower) triangles show the interaction score for the effect of the drugs in the row (column) on the drugs in the column (row).
f. Measurements of directional interaction scores for pairwise combinations of 14 antibiotics in early-killing (2 biologically independent strains measured in \(A\)\textsubscript{1×}, \(A\)\textsubscript{2×}, and \(A\)\textsubscript{1×}+\(B\)\textsubscript{1×} at 6, 4, 2 experiments, respectively). Upper (lower) triangles show the interaction score for the effect of the drugs in the row (column) on the drugs in the column (row). Insets show the killing curves and the average survival fractions following short (1 and 1.5-hour) treatment with a representative suppressive drug combination. For each treatment, representative micro-plating images of one of the replicates are shown. Error bars represent 95% confidence intervals calculated from the colony counts of the experiments by the Poisson’s model (f, middle panel) or 2x s.e.m. (f, right panel).
Extended Data Fig. 2 | a, The dose response curves of 19 drugs. Cell viability (CFU/ml) measured following 8-hour exposure to different concentrations of 19 drugs. Hill function fit indicated (note though a slight non monotonic behaviour for some of the antibiotics). The fixed cidal concentration chosen for each drug is indicated as a red line (“x1”, Methods, Fig. 1b, Supplementary Table 1). The cell viability of 2 biologically independent strains were measured over each drug. Error bars represent 95% confidence intervals calculated from the colony counts by the Poisson’s model. b–d, Lack of correlation between directional interactions of growth-inhibition and early-killing or long-term clearance. The directional interaction scores for all the pairwise combination of 14 antibiotics (n = 182, coloured by three different drug-pair groups: (1) black, pairs of classically static (CS) drug pairs; (2) purple, pairs of cidal drug pairs; (3) cyan, combining a cidal and CS drugs ranged from synergistic to antagonistic and suppression. There is a lack of correlation between growth-inhibition interactions (Extended Data Fig. 1e) and early-killing interactions (Extended Data Fig. 1f) seen for all drug pairs (b; Rho:0.04) and within each of the three drug-pair groups (CS pairs, Rho:0.01; cidal pairs, Rho:0.01; CS-cidal pairs, Rho:−0.13); as well as between growth-inhibition and long-term killing interactions (Fig. 1c) between all drug pairs (c, Rho:−0.02) and within each of the three drug-pair groups (CS pairs, Rho:−0.1; cidal pairs, Rho:−0.13; CS-cidal pairs, Rho:−0.18). While there is a significant strong correlation (d, Rho:0.73, P-value:4.5 × 10−32) between early-killing and long-term clearance interactions seen for all drug pairs and within each of the three drug-pair groups (CS pairs, Rho:0.69, P-value:10−3; cidal pairs, Rho:0.47, P-value:3.4 × 10−5; CS-cidal pairs, Rho:0.61, P-value:1.9 × 10−10) indicating that the long-term effect of the drug combinations can be predicted, to a good extent, from the early-killing efficacy. Spearman’s test was applied.
Extended Data Fig. 3 | Pairwise interactions in growth inhibition. For each pair of the 14 antibiotics (91 pairs), *S. aureus* (ATCC 29213) growth was measured following 18-hour exposure to 2-D dosage gradients of two drugs (normalized final OD, size of black dots). A polynomial surface model was fitted (Loess fit; R stats) to these normalized OD values represented by the isoboles for growth inhibitions of 30%, 60% and 90%. The 90% inhibition isobole was contrasted with four reference isobole lines representing extreme synergy (red), Loewe’s dosage additivity (grey), antagonism (green) and strong suppression (blue). The distance of the 90% inhibition line from these reference isoboles was measured along two linear lines of fixed A:B drug dosage ratios of 1:3 and 3:1 (in IC90 units, dashed magenta, and orange lines), representing the effect of drug A on drug B and the effect of drug B on drug A, respectively. The unit of the A and B axis is μg/ml.
Extended Data Fig. 4 | Pairwise interactions in early-killing. Decline in cell viability is measured as colony forming unit per ml (CFU/ml) along 90 min following exposure to each of the 14 drugs when applied individually (orange and magenta) and when applied, at the same doses, in pairwise combinations (black). The line represents the least square fit of the viability measurement over time to $\exp(-L(\mu+\mu)(e^{-t/L}-1)-kt)$ where $t$ is time, $\mu$ is the measured growth rate in absence of drug (=1.4/hour), and $L$ and $k$ are two fitted parameters representing an effective time delay for drug action, and the killing rate, respectively (2 biologically independent strains measured in $A_1\times$, $A_2\times$, and $A_1\times+B_1\times$ at 6, 4, 2 experiments, respectively). Error bars represent 95% confidence intervals of the experiments calculated from the colony counts by the Poisson’s model.
Extended Data Fig. 5 | Pairwise interactions in long-term killing. Decline in cell viability is measured as colony forming unit per ml (CFU/ml) along 8 h following exposure to each of the 14 drugs when applied individually (orange and magenta) and when applied, at the same doses, in pairwise combinations (black). The line represent the least square fit of the CFU measurement over time to $(1-f_p)\exp\left((-1+\mu)\left(e^{-\frac{t}{L}}\right)\right) + f_p \exp\left(-k_p t\right)$ where $t$ is time, $\mu$ is the measured growth rate in absence of drug (=1.4/hour), and $L$, $k_n$, $k_p$, and $f_p$ are four fitted parameters representing an effective time delay for drug action, the fraction of persistent cells, and the killing rates of the normal and the persistent cells, respectively (2 biologically independent strains measured in $A_{1×}$, $A_{2×}$, and $A_{1×} + B_{1×}$ at 6, 4, 2 experiments, respectively). Error bars represent 95% confidence intervals of the experiments calculated from the colony counts by the Poisson’s model.
Extended Data Fig. 6 | Reciprocal suppressive interactions in long-term clearance efficacy are robust to changes in drug concentrations, growth environments, incubation time, bacterial physiological state, and strain backgrounds. a, Survival of the S. aureus (ATCC 29213) following 8-hour of exposure to cidal concentration of tetracycline and cloxacillin (TET-CLO) indicating robust reciprocal suppression over a wide range of drug concentrations (no bacteria well at 0,0 concentration point). b, Bacteria survival measured following 8-hour exposure to clindamycin-cloxacillin (CLI-CLO, left) or tetracycline-cefoxitin (TET-FOX, right) demonstrating strong reciprocal suppression in different media: Lauria Delbruck (LB), Brain-Heart Infusion (BHI), Tryptic Soy Broth (TSB), and human serum (2 biologically independent strains measured at 2 experiments; 4, for TET-FOX in TSB). c, Survival following exposure to CLI-CLO (10 μg ml⁻¹, 2 μg ml⁻¹) or TET-FOX (8 μg ml⁻¹, 4 μg ml⁻¹) in LB demonstrate reciprocal suppression both after 8-hour incubation (top, control) as well as after 24-hour (bottom; 2 strains measured at 2 replicates). d, Viability of 2 biologically independent strains in different physiological state (left, exponential growing state, OD = 0.1; right, late growing phase, OD = 0.8) was measured in 2 experiments following 8-hour exposure in human serum to CLI-CLO (50 μg ml⁻¹, 10 μg ml⁻¹; b-d **** P values < 10⁻¹⁰; two tailed z-test). e, Reciprocal suppression is demonstrated in an unrelated, methicillin resistant S. aureus (MRSA, Methods). f, Viability of the sensitive strain measured 8-hour post inoculation into larvae treated with CLI-CLO (15 μg ml⁻¹, 40 μg ml⁻¹; n = 5 TRUE larvae per condition; left to right P values: 0.008, 0.103; Mann-Whitney test). g-i, Viability of the sensitive strain (GFP) measured 8-hour competition with the MRSA strain (g,f,i; DsRed) or with the TETR strain (h; DsRed) within larvae treated with TET-AMP (g, 50 μg ml⁻¹, 100 μg ml⁻¹; h, 1.5 μg ml⁻¹, 38 μg ml⁻¹) or with CLI-AMP (i, 6 μg ml⁻¹, 6 μg ml⁻¹; g, n = 8 AROL larvae per condition, left to right P values: 1.5 × 10⁻⁴, 1.5 × 10⁻⁴; Mann-Whitney test, same experiment as in Fig. 3f; h, n = 10 TRUE larvae per condition, same experiment as in Fig. 3d; i, n = 10 TRUE larvae per condition, same experiment as in Extended Data Fig. 9i). Data are presented as mean +/- 95% confidence intervals (points on the axis are below the detection limit).
Extended Data Fig. 7 | Long-term clearance efficacy of multidrug combinations is reduced as more drugs are added. We repeated the Fig 2a experiment with newly prepared antibiotic and bacterial culture stocks and the results were consistent with our prior experiment (Fig. 2a). a, Bacterial survival following 8-hour of exposure to all possible combinations of 6 drugs (drug combinations are indicated by the colours at the top of the hexagonal columns, legend, Supplementary Table 1; 2 strains measured in 2 independent experiments) b, c, The survival fractions following long (8-hour, b) or short (1-hour, c) drug exposures as a function of the number of co-mixed drugs for all 6-drug combinations (average survival fraction of the two replicates of the 63 combinations; black circles; box-plots; centre line, median; box limits, upper and lower quartiles; whiskers, maximum and minimum values). Grey line shows quadratic fit. The coloured dots and coloured linear fit lines represent the efficacy of the single treatments as a function of increasing doses (from 1× to 6× of the dosage used in the combination; antibiotic color codes indicated in panel a). Data are presented as mean +/- 95% confidence intervals. d, e, The distribution of the clearance (d) and early-killing (e) interaction scores among all tested non-overlapping pairs of drug sets (for example, drug mix ABC is decomposed into AB+C, A+BC, AC+B; n = 301). The combined effect of the drug mix is shown relative to the minimum (min) and maximum (max) of the effect of the two drug sets (Methods). Drug mix pairs are classified as only cidal (purple), only classically static (CS; cyan), mix of all CS and all cidal (pink), pairs of mixed drug sets (mix, grey).
Extended Data Fig. 8 | The reduced clearance efficacy of multidrug mixes can be circumvented by specific persistence-targeting drugs. a, c. Bacterial survival following 8-hour of exposure to all possible combinations of 6 drugs including two persistence-targeting drugs (daptomycin, DAP; mitomycin C, MIT; drug combinations are indicated by the colours at the top of the hexagonal columns, legend, Supplementary Table 1; 2 biologically independent strains measured in 2 experiments). Inset: The survival fractions following long (8-hour) drug exposures as a function of the number of co-mixed drugs for all 6-drug combinations (n = 63 combinations; black circles). The coloured dots and coloured linear fit lines represent the efficacy of the single treatments as a function of increasing doses (from 1× to 6× of the dosage used in the combination; antibiotic color codes indicated in panel a; note though a slight non monotonic behaviour of the minocycline (MIN) and pristinamycine (PIR), possibly due to Eagle effect of the drugs). Data are presented as mean +/- 95% confidence intervals. b, d. The distribution of the clearance interaction scores among all tested non-overlapping pairs of drug sets (for example, drug mix ABC is decomposed into AB+C, A+BC, AC+B; n = 301). The combined effect of the drug mix is shown relative to the minimum (min) and maximum (max) of the effect of the two drug sets (Methods). Drug mix pairs are classified as only cidal (purple), only classically-static (CS; cyan), mix of all CS and all cidal (pink), pairs of mixed drug sets (mix, grey), or sets including DAP or MIT (green shades, legend). Persistence-targeting drugs strongly synergize the clearance efficacy of other drug mixes (Source Data for Fig. 2).
Extended Data Fig. 9 | Reciprocal suppressive drug combination selects against antibiotic resistance to either one of the single drugs. a, b, The survival fraction of the sensitive strain with and without an inducible β-lactamase (sensitive and AMP′, respectively) was measured separately in two parallel experiments following 8-hour of exposure to a two-dimensional gradient of tetracycline (TET) and ampicillin (AMP) (e) or clindamycin (CLI) and AMP (f). c–f, Survival fraction measured after 8-hour competition of equally mixed sensitive (ATCC 29213, Methods) versus evolved tetracycline resistant strains (c, Sensitive, red; TETR, green; d, Sensitive, green; TET R, red) or MRSA strain (e, Sensitive, red; MRSA, green; f, Sensitive, green; MRSA, red) in two-dimensional gradient of tetracycline (TET) and ampicillin (AMP). The log₁₀ of the ratio of the survival of the sensitive and resistant strains are shown in red and green color scale (at the bottom of each panel). g, Survival fraction of the sensitive (grey) and MRSA (black) strain following an 8-hour treatment with TET alone (11 μg ml⁻¹; blue box in panel e,f) and with TET-AMP mix (11 μg ml⁻¹, 0.4 μg ml⁻¹; red box in panel e,f; n = 24 replicates of sensitive-DsRed versus MRSA-GFP and n = 24 replicates of a corresponding marker-swap competition (P value = 3 × 10⁻¹⁷, two sided Mann Whitney test). h–i, The log₂ of the ratio of the survival of the sensitive (GFP) and MRSA (DsRed) resistant strains after competing them for 8-hour within human serum (h, 2 biologically independent strains measured in 5 independent experiments per condition; two sided Mann-Whitney test; from left to right P values: 0.0072, 0.65, 0.0002) or within larvae (i, n = 10 TRUELarvae per condition, two sided Mann-Whitney test; from left to right P values: 0.0004, 0.0017, 0.0002) with MIN (h, 40 μg ml⁻¹; i, 6 μg ml⁻¹), AMP (h, 35 μg ml⁻¹; i, 6 μg ml⁻¹) or MIN+AMP. Box-plots: centre line, median; box limits, upper and lower quartiles; whiskers, maximum and minimum value.
Extended Data Fig. 10 | a–d, Supplementing persistence-suppressive drug combinations with avibactam, a β-lactamase inhibitor, decreased rather than increased clearance efficacy against a β-lactam resistant strain. The clearance efficacy of ampicillin (AMP) is enhanced with the addition of increasing dosage of avibactam (AVI, a). In contrast, the treatment efficacy is reduced rather than enhanced when avibactam is added to the reciprocal suppressive combination of AMP and TET (b, fixed dosage of TET, 20 μg ml−1), AMP and CLI (c, fixed dosage of CLI, 10 μg ml −1) and AMP and FUS (d, fixed dosage of FUS, 20 μg ml −1). e–h, Reproducibility of the microplating assay between two biological replicates. e,f, Correlation between the colony forming units (CFU per ml) of two replicates measured during the pairwise drug combination screens (e, Extended Data Fig. 5; f, Extended Data Fig. 4). g–h, Correlation between the colony forming units of the two replicates of the multidrug screens (g, left panel, Fig. 2a; g, right panel, Extended Data Fig. 7a; h, Extended Data Fig. 7c). Single, two-drug treatments and high-order drug combinations were differentially coloured as red, light blue and dark blue, respectively. i, The distribution of the killing delays across all single-drug and two-drug treatments. After taking the best fit of the killing curves to the killing equation (see the killing equation on the caption of Extended Data Fig. 4) we plot the distribution of delay (L) across all conditions. j–m, Nutrient depletion is negligible during the killing assay. j–k, The optical density (OD, j) and the green fluorescence signal (GFP, k) of the sensitive strain following 8-hour incubation with increasing concentration of cefoxitin. After incubation the supernatant of each well was transferred to a new microplate and was re-inoculated with 2 × 10^6 GFP tagged cefoxitin resistant strain (MRSA) and incubated for another 6-hour. l–m, The OD (l) and GFP signal (m) of the cefoxitin resistant culture after 6-hour incubation.
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Our web collection on statistics for biologists contains articles on many of the points above.

Software and code

Policy information about: availability of computer code

Data collection

The optical density were read by Biotek Synergy 2 plate reader, software version GeneS 1.115. The agar plates were imaged for GFP (excitation:470/30; emission: 540/50) and RFP (excitation: 590/30; emission 643/75) with a custom-made automated imaging device (microscope).

Data analysis

Code availability statement is provided as a separate section after the data availability statement. The custom python script developed for image analysis is deposited in a public DOI-minting repository (Open Science Framework) and was cited in the appropriate Methods section. The detailed algorithm for calculating the drug interaction scores is available in the Methods section.

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The raw data including all the micro plating images with the bacterial colonies that support the findings of this study are available as Source Data files in a public DOI-minting repository [https://osf.io/ypl5/].
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Life sciences study design

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Sample size

We screened the cell viability of 2 biologically independent strains (DsRed, GFP) measured in 585 drug conditions (28 singles [1X, 2X dose; at 6 and 4 independent experiments] and 91 pairs in 2 independent experiments; 5X 36 singles [1X, 2X, 3X, 4X, 5X and 6X dose], 3X52 multigene mixtures at 2 independent experiments; 15 timepoints; 2-time ranges (short and long-term killing) and 4 to 8 plating (dilutions-technical replicates). The sample size was chosen to allow detection of possible outliers while still allowing a valid systematic quantification of pairwise and multi-drug interactions. Sample size for the larval experiments was large enough to allow statistically significant detection of variation assuming the Mann Whitney test. Growth inhibition interactions between all the pairwise combinations of the 14 antibiotics were measured as the final optical density of the cultures under a 2-D concentration gradients (8x12 different concentrations). Then a polynomial surface model (Loess-fit, degree 2) was fitted to the normalized optical density values.

Data exclusions

At the last time point of the early killing and the long-term clearance assay, 15μl aliquots from each well of the assay plates were also inoculated into a 96-well plate with 150μl liquid LB containing, in each well, the same antibiotic concentrations of the assay plates and incubated overnight at 37°C, 250rpm. Following incubation, OD was measured and growing cultures (OD600>0.2), indicating the presence of resistant cells, were excluded from further analysis. We excluded these wells to specifically focus on the survival fraction of the sensitive cell population. Micro-plating spots where the colonies merged extensively or with too dense colonies to reliably count were designated "uncountable" (over 100 colonies per spot).

Replication

Our high-throughput cell viability assay was highly reproducible, showing good agreement among the independent experiments (Extended Data Fig. 12a-d). The number of biologically independent replicates and experiments for each experiment are listed in the appropriate figure legends.

Randomization

The allocation of the single and multi-drug conditions were randomized on the 96-well plates during the high-throughput screen to measure early and long-term killing interactions.

Blinding

The surviving colonies were counted in the presence of different compounds in the same way, while blind to the compound identity of each well.

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Animals and other organisms

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Laboratory animals

Larvae of the Galleria mellonella at their final instar stage were used to assess the clearance efficacy of drug combinations. Larvae were either purchased from UK BioSystem Technology (TrueLarv) or from Israeli Agricultural Research Organization, Volcani Institute (ARCLarv).

Wild animals

Provide details on animals observed in or captured in the field; report species, sex and age where possible. Describe how animals were caught and transported and what happened to captive animals after the study (if killed, explain why and describe method; if released, say where and when). OR state that the study did not involve wild animals.
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Ethical approval is not required.

Note that full information on the approval of the study protocol must also be provided in the manuscript.