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Efficient measurement and factorization of high-order drug interactions in Mycobacterium tuberculosis

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Combinations of three or more drugs are used to treat many diseases, including tuberculosis. Thus, it is important to understand how synergistic or antagonistic drug interactions affect the efficacy of combination therapies. However, our understanding of high-order drug interactions is limited because of the lack of both efficient measurement methods and theoretical frameworks for analysis and interpretation. We developed an efficient experimental sampling and scoring method [diagonal measurement of \( n \)-way drug interactions (DiaMOND)] to measure drug interactions for combinations of any number of drugs. DiaMOND provides an efficient alternative to checkerboard assays, which are commonly used to measure drug interactions. We established a geometric framework to factorize high-order drug interactions into lower-order components, thereby establishing a road map of how to use lower-order measurements to predict high-order interactions. Our framework is a generalized Loewe additivity model for high-order drug interactions. Using DiaMOND, we identified and analyzed synergistic and antagonistic antibiotic combinations against Mycobacterium tuberculosis. Efficient measurement and factorization of high-order drug interactions by DiaMOND are broadly applicable to other cell types and disease models.

INTRODUCTION

Combinations with three or more drugs are the cornerstone of treatment for many diseases including cancer, HIV, and tuberculosis (TB). Therefore, it is important to understand how combining drugs changes treatment efficacy. In synergistic or antagonistic drug interactions, drugs in combination may be, respectively, more or less effective than what is expected from the individual drugs. Treatments that include a synergistic combination achieve the same efficacy with a relatively lower total dose, potentially improving efficacy, alleviating side effects, and shortening treatment (1, 2).

TB treatment remains difficult, and combinations of three or more drugs (high-order combinations) are the standard of care for active TB. The current treatment regimen for drug-sensitive TB uses a four-drug combination for 2 months, followed by a two-drug combination for at least 4 months (3). Drug-resistant TB therapy is more arduous, involving long-term, high-order combinations of antibiotics with severe side effects (4). Despite the critical importance of this information for treatment, high-order interactions are rarely measured and analyzed in any disease. For example, because of the high number of measurements required to measure three-way interactions by the standard checkerboard assay, there is a relative scarcity of these measurements. Targeted studies that have sought to predict three-way interactions from pairwise measurements in Escherichia coli have found that pairwise and three-way interaction measurements are correlated (5, 6). These prediction methods are likely to be useful in prioritizing combinations for further study, but cannot fill the need for experimental measurement of high-order interactions. In the case of TB, we lack measurement of high-order drug interactions and reliable, systematic measurement of pairwise interactions against Mycobacterium tuberculosis (Mtb), the causative agent of TB. Therefore, drug interactions are not routinely considered in drug regimen design (7–11). Thus, a more efficient methodology to measure drug interactions would enable us not only to comprehensively measure pairwise interactions but also to interrogate high-order drug combinations and establish tools to predict high-order synergistic combinations for further testing early in preclinical regimen design.

The absence of systematic studies of high-order drug interactions is due to three significant challenges. The first challenge is the experimental measurement of high-order drug interactions. Pairwise drug interactions are traditionally determined by a two-dimensional (2D) checkerboard assay (Fig. 1A). To evaluate \( D \) different doses for each drug in a pairwise combination, the response to \( D^2 \) = 100 dose combinations must be measured, which requires the preparation and measurement of cell growth in 100 microplates. A high-order drug interaction among \( n \) drugs can be determined using a high-dimension checkerboard assay (12). However, this measurement technique is very expensive and can be logistically prohibitive (13): If \( D \) doses are used for each drug, then \( D^n \) dose combinations need to be assessed. If, for example, we were to measure a five-way drug interaction \( (n=5) \) at a resolution of 10 doses for each drug \( (D=10) \), a 5D checkerboard assay necessitates the measurement of cell response in \( 10^5 \) combinations, requiring 1000 microplates. Another serious limitation is that these tests would require vast amounts of drugs, and the cost and availability of drugs, especially natural products, preclude such experiments. Because we lack efficient measurement procedures, drug interactions of four or more drugs remain uncharacterized, and there are few examples in literature that report three-way drug interactions.

Ideally, drug interactions are scored from a checkerboard assay relative to the simple null model, indicating that the interaction of a drug with itself is additive (14, 15). Thus, with the Loewe additivity model, if a combination’s effect is higher or lower than additive, the combination is synergistic or antagonistic, respectively (Fig. 1A). We may visualize this intuitive interpretation of the checkerboard by evaluating the contour of a common phenotype (isobole). A straight contour indicates additivity, whereas concave and convex contours are observed in synergistic and antagonistic interactions, respectively (Fig. 1A). The shape of this curve is quantified using the fractional inhibitory concentration (FIC), the standard metric of interaction scores. In the
Fig. 1. DiaMOND for measuring pairwise drug interactions. (A) Simulated 2D checkerboard assay as demonstration of additive (left), synergistic (middle), and antagonistic (right) pairwise drug combinations. In each assay, one drug is linearly increased on each axis, and growth inhibition (white to blue color bar) is recorded for each concentration combination. A combination of a drug with itself will yield linear isophenotypic contours (isoboles). Drug combinations with linear, concave, or convex isoboles are classified as additive, synergistic, or antagonistic, respectively. To efficiently approximate the isoboles, DiaMOND samples the most information-rich dose responses of the checkerboard: the single-drug dose responses (white rectangles) and the 1:1 two-drug combination dose response (yellow rectangle; along the diagonal of the checkerboard). The dashed blue line connecting the observed single-drug dose responses approximates the additivity isobole for a given phenotype (for example, IC50), and the intersection of this line with the two-drug dose response is the expected dose in the null model (€1; blue diamond). The circles indicate the drug doses that result in the particular phenotype for each dose response. In this setting, the blue diamond (€1) will be the same distance as, closer to, or further from the origin in additive, synergistic, or antagonistic combinations. The circles are colored by drug interaction type throughout the manuscript: white, additivity; green, synergy, red, antagonism. (B) DiaMOND experimental setup to measure pairwise drug interactions requires three dose-response experiments, where drug dose is linearly increased. Two dose responses are single-drug dose responses. In the third dose-response experiment (yellow rectangle), a 1:1 mixture of drug X and drug Y is linearly increased. This mixture has half the concentration of both drug X and drug Y, and its dose response is the dose response of the diagonal sampling shown in (A). The expected dose (€1), shown by a blue diamond, is calculated using the geometric argument described in (A). When the observed dose is the same as, less than, or more than the expected dose, additivity, synergy, or antagonism is concluded, respectively. The brackets illustrate the expected (€1) and observed (o2) doses for the two-drug dose response (see also fig. S1). (C) All pairwise interactions among nine antibiotics tested in Mtb. The dose response of nine antibiotics and all 36 pairwise combinations is shown. Circles or blue diamonds mark the observed or expected IC50, respectively. Circles smaller or larger than expectation are marked by green or red, indicating synergy or antagonism. FIC2 scores obtained by dividing observed with expected IC50 are shown on the right of each two-drug dose response. (D) Hierarchical clustering of the pairwise interactions among nine antibiotics. The geometric mean of two biological replicates is represented. (E) Validation of two synergies and one antagonism via the traditional checkerboard method in Mtb. Simulated dose-response data are shown in blue gradient, and experimentally determined dose-response data are shown in grayscale gradient throughout.
absence of an efficient checkerboard assay, the Bliss multiplicative model has become a commonly used scoring system for determining drug interactions. The Bliss multiplicative model defines the expected phenotype [such as median inhibitory concentration (IC50) for 50% growth inhibition] as the multiplication of the phenotype induced by each drug, and therefore, Bliss scoring enables an efficient experimental setup where measurement at only four conditions is adequate to score a pairwise interaction. Hence, Bliss scoring is useful for understanding the interaction of drugs at therapeutically relevant concentrations (16). Despite this advantage in efficiency, the Bliss model lacks a null model; therefore, one cannot do a control drug interaction experiment that, by definition, will result in noninteraction (17). The same drug’s combination with itself might result in synergy, independence, or antagonism, depending on the concentration chosen to sample the drug’s dose response (18). In contrast, the Loewe additivity model produces a single interaction score, which is an integration of the whole range of concentrations that may be combined. Thus, a measurement methodology that is more efficient than a traditional checkerboard with Loewe additivity scoring would empower systematic measurement of pairwise and high-order drug interactions.

The second challenge is the evaluation and interpretation of high-order drug interactions. When two drugs are combined, the measured interaction is simply their pairwise interaction. However, when three drugs are combined, the measured interaction encapsulates three pairwise interactions and one three-way interaction. A three-way interaction, therefore, may refer to either (i) a total observed interaction, which compares the three-way combination effect with single drugs, or (ii) an emergent interaction, which compares the three-way combination effect with the expectation from the underlying pairwise interactions. For example, a three-way combination would have emergent synergy if the total interaction is more synergistic than the expectation based on the three pairwise interactions alone. In this setting, if pairwise interactions were antagonistic and total interaction was additive, then the emergent three-way interaction must be synergistic. Understanding the emergent effect of a high-order interaction that is not predicted by lower-order combinations requires a scoring framework to factorize high-order interaction scores into meaningful components. Score factorization was recently developed for three-way interactions scored with the Bliss independence model (19), but no such framework exists for the factorization of high-order drug interactions using intuitive Loewe additivity scoring.

The third challenge is predicting the strongest synergies among the large number of possible high-order drug combinations to facilitate efficient prioritization of optimal drug regimens. As noted above, the experimental measurement of even one five-way drug interaction is prohibitively expensive. However, there are, for example, 15,504 different five-way combinations possible among 20 drugs. An experimental screen of high-order drug interactions is therefore not possible, and finding synergistic high-order combinations requires the development of tools to predict synergistic combinations. Two recent studies used Bliss scoring to analyze three-way drug interactions in E. coli and in cancer cells and found that they are largely determined by the pairwise interactions among the constituent drugs (5, 6). Neither of these studies, however, systematically evaluated or analyzed the emergent interactions of high-order combinations. Several pairwise drug interaction screens have reported that synergy is rare, whereas antagonism is more common than additivity and synergy (20–23). Because any experimental search for synergy will necessarily screen a very small percentage for high-order combinations, the ability to predict high-order interactions is essential to prioritize candidate synergistic high-order combinations.

The absence of an efficient measurement technique for drug interactions has limited the ability to fully pursue these challenges to systematically measure drug interactions, understand central questions about high-order drug interactions, and prioritize synergistic combinations in preclinical testing. Here, we present a methodology that uses a diagonal sampling of the checkerboard assay to efficiently measure and score pairwise or high-order drug interactions, which extends on the theoretical work of Berenbaum (24). Our methodology, DiaMOND (diagonal measurement of n-way drug interactions), requires only n + 1 dose responses and produces the FIC of the n-way drug interaction. We used DiaMOND to measure all pairwise interactions among nine TB antibiotics and all possible high-order combinations among five antibiotics in Mtb, revealing synergies in three pairwise, three three-way, and two four-way combinations. With these drug interaction data, we generalized the Loewe additivity model to systematically explore high-order drug combinations and develop a framework that factorizes the components of a high-order drug interaction to resolve contributions from each emergent drug interaction. We demonstrate that high-order interactions can often be predicted by their lower-order combinations, but we also identified one three-way drug combination with strong emergent antagonism, highlighting the importance of measuring high-order interactions. Our study provides a unifying framework to measure, score, classify, and predict high-order drug interactions. We applied DiaMOND to identify and analyze synergistic drug combinations against Mtb; however, DiaMOND’s application is not limited to drug studies in Mtb and may also be used to systematically and efficiently measure and analyze the interactions of multiple drugs, cytokines, or other molecules on cellular behavior.

RESULTS

Diagonal sampling method to measure pairwise drug interactions

Pairwise drug interactions are traditionally measured by a 2D checkerboard assay. Examples of checkerboards for additive, synergistic, and antagonistic pairs are shown in Fig. 1A with simulated values. The synergy or antagonism of a drug pair is assessed by the shape of the isobole (or contour of the chosen phenotype, such as IC50 for 50% growth inhibition). For two drugs, X and Y, Loewe additivity is defined as $\text{FIC}_2 = \frac{\text{IC}_{50X} + \text{IC}_{50Y}}{2}$. On a checkerboard assay, an FIC2 = 1 corresponds to the straight line connecting IC50X and IC50Y (Fig. 1A). Synergistic pairs achieve the same phenotype with lower concentrations of the combined drugs and their checkerboards have a concave contour, giving FIC2 scores between 0 and 1 (Fig. 1A, middle), and checkerboards of antagonistic pairs have convex contours with FIC2 scores > 1 (Fig. 1A, right).

We sought to minimize the number of measurements required by this assay and reasoned that the entirety of the checkerboard was not necessary to quantify the curvature of the contour. Instead, we aimed to sample the checkerboard in regions that provide the most information about the shape of the contour. To do so, we measure three dose responses for each pairwise interaction: (i) the dose response for drug X, (ii) the dose response for drug Y, and (iii) the dose response for an equipotent (1:1) mixture of drug X and drug Y (referred to hereafter as a two-drug dose response). These dose responses correspond to the horizontal, vertical, and diagonal (yellow) rectangles drawn on each checkerboard in Fig. 1A. For each dose response, one can identify the dose that produces a specified phenotype response, such as IC50. In Fig. 1A, we show how a line connecting the IC50 values for drug X, drug Y, and
the two-drug combination approximates the contour. To quantify the curvature from these three dose responses, the expected IC$_{50}$ ($e_1$; blue diamonds in Fig. 1A) of the two-combination dose response is computed on the basis of the null hypothesis that the expected interaction is additive. Thus, $e_1$ is the distance from the origin to the intersection of the straight line connecting IC$_{50}$ values of single drugs with the two-drug dose response (bracketed in Fig. 1B and see fig. S1 and method S1 for calculation of $e_1$ given two IC$_{50}$ values).

The curvature of the contour can be approximated by the ratio of the observed IC$_{50}$ of the two-drug combination ($o_2$; circles in the two-drug dose responses in Fig. 1A) to the expected IC$_{50}$ in additivity ($e_1$; blue diamonds in Fig. 1A). We therefore define the FIC of a pairwise combination (FIC$_2$) as

$$FIC_2 = \frac{o_2}{e_1}$$

(1)

The FIC, in both traditional checkerboard assays and with DiaMOND, is not only a measure of contour curvature but also a quantification of the relative efficacy of the combination compared to single drugs. For example, an FIC$_2$ score of 1 (Fig. 1B, left) corresponds to an additive interaction, whereas 0.5 (synergy) or 2 (antagonism) indicate that half or double the dose leads to the same effect, respectively (see fig. S1 for examples of step-by-step calculations of FIC$_2$ given two single-drug and one two-drug dose responses).

The experimental setup for DiaMOND is shown in Fig. 1B, with simulated examples of additivity, synergy, and antagonism. The assay requires two single-drug dose responses and just one two-drug dose response, where the two-drug dose response is a 1:1 mixture of drug X and drug Y concentrations used for the single dose responses (Fig. 1B and fig. S1). This mixture will have half the concentration of both drug X and drug Y, and its dose response will be the dose response of the diagonal sampling shown in Fig. 1A (yellow outline).

**Experimental measurement of pairwise drug interactions in Mtb**

We used DiaMOND to measure all the pairwise interactions among nine first- and second-line antibiotics (Table 1) against Mtb. We generated the dose response for nine antibiotics individually and the two-drug combination dose response for the 36 pairwise combinations, each in biological duplicate. For each two-drug dose response, the expected IC$_{50}$ ($e_1$) was computed using single-drug IC$_{50}$ values. Figure 1C shows one of our experimental screens, where the expected IC$_{50}$ for each combination is indicated with a blue diamond. To compute FIC$_2$ (Eq. 1), the observed IC$_{50}$ of the combinations ($o_2$; shown with a circle) was divided by the expected IC$_{50}$ ($e_1$). The FIC$_2$ scores were highly correlated between biological replicates (Spearman’s correlation $r = 0.86$, $P < 3.7 \times 10^{-9}$; fig. S2). The expected and observed drug concentrations used for FIC calculations are not limited to IC$_{50}$ values, and any measurable parameter from a dose response may be used. Because IC$_{50}$ is broadly used to measure drug sensitivity in microbes and in cancer cell lines, we chose IC$_{50}$ for this study. However, dose responses are usually non-linear, and therefore the FICs may depend on the choice of what level of growth inhibition is used in the FIC calculations (for example, IC$_{50}$ versus IC$_{30}$ versus IC$_{70}$). Taking advantage of the single-drug and two-drug dose responses measured with DiaMOND, we reanalyzed these data and computed FIC$_2$ scores using IC$_{30}$, IC$_{40}$, IC$_{60}$, and IC$_{70}$. We found that the FIC scores correlated significantly in this range (fig. S3), suggesting that the FIC is not highly dependent on the choice of inhibition level. We conclude that DiaMOND’s measurement layout and scoring (FIC$_2$) are reliable and robust.

To visualize the overall pattern of synergy and antagonism in the 36 pairwise combinations and the similarities in overall interaction patterns among the nine antibiotics, we show the geometric mean of interaction scores (FIC$_2$) from both replicate experiments in Fig. 1D. We used a geometric mean rather than an arithmetic mean because the FIC is a ratio and therefore must be averaged by a geometric mean. For example, an FIC score of 0.5 means half the dose is required for the same effect and a score of 2 means that double the dose is required for the same effect. If we imagine two (noisy) experiments of the same pair that gives scores of 0.5 and 2, the conclusion would be that these drugs are additive. The arithmetic mean of 0.5 and 2 is 1.25, erroneously suggesting antagonism. However, the geometric mean of 0.5 and 2 is 1, and is scored correctly as additive.

In agreement with previous studies, pairwise synergy was rare and antagonism was common in our pairwise measurement set (20–23). However, we identified three pairs with strong and reproducible synergy: clofazimine + bedaquiline, clofazimine + isoniazid, and pretomanid + rifampicin. Of these, synergy between clofazimine and isoniazid and that between clofazimine and bedaquiline have been previously reported (25, 26), but the synergy between pretomanid and rifampicin against Mtb is novel. Other interactions such as the antagonism between moxifloxacin and rifampicin and that between moxifloxacin and linezolid have been reported previously (16, 27). We note the tendency of moxifloxacin to exhibit antagonism with nearly all antibiotics tested. Rifampicin + isoniazid was found to be mildly antagonistic, consistent with patient studies that showed antagonism or additivity depending on treatment dose (28–30). The lower bound and upper bound of FIC$_2$ in this set were 0.45 (bedaquiline + clofazimine and clofazimine + isoniazid) and 1.90 (bedaquiline + moxifloxacin), respectively. These FIC$_2$ values correspond to combinations with approximately double or half the potency of the single drugs. We validated the clofazimine + bedaquiline and pretomanid + rifampicin synergy and the moxifloxacin + pretomanid antagonism using traditional checkerboard assays (Fig. 1E). We conclude that DiaMOND is an efficient and robust way to measure pairwise drug interactions, and we report a novel synergistic antibiotic pairing that had not yet been reported among TB drugs.

| Drug | Abbreviation | Target process | MIC (µg/ml) |
|------|--------------|----------------|------------|
| Bedaquiline | BDQ | ATP synthase | 0.6 |
| Clofazimine | CLZ | DNA replication | 2.8 |
| Ethionamide | ETA | Mycolic acid | 3 |
| Ethambutol | ETH | Mycolic acid | 1.5 |
| Isoniazid | INH | Mycolic acid | 0.18 |
| Linezolid | LIN | Protein synthesis | 3 |
| Moxifloxacin | MOX | DNA gyrase | 0.35 |
| Pretomanid | PRE | Protein synthesis | 0.8 |
| Rifampicin | RIF | RNA polymerase | 0.06 |

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Diagonal sampling method to measure high-order drug interactions

In both DiaMOND and the traditional checkerboard assay, the experimental setup of a three-way drug interaction is an extension of the pairwise measurement technique. With a checkerboard assay, a 3D dose space is sampled exhaustively and the curvature of the contour as a plane (as opposed to a line for the pairwise assay) is measured (Fig. 2A). With DiaMOND, we instead sample the most information-rich areas of the 3D dose space with the goal of measuring the contour curvature. Like pairwise drug combinations, for three-way drug interactions, we measure the single-drug dose responses (X, Y, and Z) and the combination dose response (a 1:1:1 mixture of the three drugs). DiaMOND defines an additivity plane using the IC50s for individual drugs, as shown in Fig. 2A, with simulated examples. The distance from the origin to the intersection of this plane with the three-drug dose response is defined as the expected IC50 of the combination, given single drugs (e1; see method S1 for calculation of e1 given three IC50 values). The comparison of the observed IC50 (o3) and e1 is then used to obtain a three-way interaction FIC3 score

\[
FIC_3 = \frac{o_3}{e_1}
\]  

(2)

DiaMOND can be directly scaled to measure the interaction score for combinations with any number of drugs. To produce an FICn score for an n-way combination, one needs n single-drug dose responses and one n-drug combination mixture dose response, totaling n + 1 dose-response experiments. Using the IC50 for each of the n single drugs, DiaMOND defines a hyperplane (the contour of additivity). The distance from the origin to the intersection of this hyperplane with the n-drug dose response is defined as the expected IC50 of the combination, given single drugs (e1). Calculation of e1 given IC50 measurements of n drugs is given in method S1. For example, measuring a five-way interaction would only take six dose-response experiments and, hence, can fit into one microplate, making DiaMOND 1000 times more efficient than a 5D checkerboard assay.

\[
FIC_n = \frac{o_n}{e_1}
\]  

(3)

Experimental measurement of high-order interactions in Mtb

Thus, having developed an efficient method to measure high-order drug interactions, we next sought to use this method to find high-order synergistic combinations in Mtb. We applied DiaMOND to five anti-
biotics that showed pairwise synergy in the pairwise screen: bedaquiline, clofazimine, isoniazid, pretomanid, and rifampicin. We used DiaMOND to measure the interaction of a five-way combination of these antibiotics. This interaction was close to additivity (FIC$_5$ = 0.93). To explore this interaction further, we measured all the pairwise, three-way, and four-way combination dose responses for these drugs in biological duplicate (Fig. 2B). Using DiaMOND, we then computed the interaction scores among all combinations of these five antibiotics (10 FIC$_2$ values, 10 FIC$_3$ values, 5 FIC$_4$ values, and 1 FIC$_5$ value). Our results were reproducible among duplicates ($r = 0.56$, $P < 3 	imes 10^{-3}$, fig. S4).

To interpret FIC scores as synergistic, additive, or antagonistic, we developed a model of experimental error. Because a drug is, by definition, additive with itself, the FIC$_n$ values obtained from interaction measurements of a drug against itself are expected to be 1, and any deviation from this is due to experimental error. We measured the FIC$_n$ of isoniazid against itself for pairwise, three-way, four-way, and five-way combinations (fig. S5). As expected, the FIC$_n$ scores from these experiments centered around 1 (mean = 1.00) with an SD of 0.09. We calculated a 95% confidence interval for additivity as 0.82 to 1.18; therefore, combinations with scores smaller or larger than this range are significantly synergistic or antagonistic.

To identify synergistic (and antagonistic) high-order drug combinations against Mtb, we evaluated the geometric mean of two replicates. We discovered one three-way combination (bedaquiline + clofazimine + isoniazid) with particularly strong three-way synergy (FIC$_3$ = 0.62). In addition, we identified the three-way combinations bedaquiline + clofazimine + rifampicin and bedaquiline + pretomanid + rifampicin and four-way combinations bedaquiline + clofazimine + isoniazid + rifampicin and clofazimine + isoniazid + pretomanid + rifampicin as synergistic high-order combinations. We validated the three-way synergy of bedaquiline + clofazimine + isoniazid using a traditional 3D checkerboard of drug concentration combinations where each drug was increased linearly in one axis in a $6 \times 6 \times 6$ matrix (fig. S6). We conclude that the DiaMOND approach can efficiently and reliably measure high-order drug interactions.

**Factorization method of high-order drug interaction scores**

We observed many high-order synergistic combinations from our exhaustive search among five Mtb drugs. This was not unexpected, because these five drugs were selected for having synergistic pairwise combinations. We next asked whether the high-order interaction scores (FICs) were due entirely to the strength of the underlying lower-order interactions. Thus, we aimed to quantify distinct contributions of the FIC$_n$: (i) the emergent interactions of the $n$-way interactions and (ii) the lower-order interactions that underlie the $n$-way interaction. DiaMOND can precisely break down an FIC$_n$ score into these two components (fig. S7). We will first demonstrate this factorization for a three-way combination and then describe a general model for $n$-way combinations.

The expected value $e_1$ used for FIC$_3$ scoring above was calculated using the IC$_{50}$ of three individual drugs. Alternatively, we can define expectation based on the strengths of the pairwise interactions. To do so, we construct an expected surface (magenta plane) with the IC$_{50}$ of the three two-drug dose responses (Fig. 3A). This plane is then the expected contour of additivity accounting for the underlying interactions of the three pairwise interactions that comprise the three-drug combination. We define $e_2$ as the distance from the origin to the intersection of this surface with the three-drug dose response. $e_2$ corresponds to the expected concentration to give IC$_{50}$ if there is additivity among three pairwise combinations. Notably, calculation of $e_1$ and $e_2$ requires three IC$_{50}$ values, supplied from single-drug or two-drug dose responses, respectively. $e_1 = e_2$ if pairwise combinations are overall additive. If the $e_2$ plane is closer to the origin than the $e_1$ plane ($e_2 < e_1$), then pairwise combinations are overall synergistic, whereas if $e_2 > e_1$, then pairwise combinations are overall antagonistic.

We observe that

$$
FIC_3 = \frac{o_3}{e_1} = \frac{o_3}{e_2} \cdot \frac{e_2}{e_1}
$$

This operation factorizes a three-way drug interaction into two components: (i) $o_3/e_2$ is a measure of how different the three-way effect is from what would be expected given the two-drug dose-response results. We define $o_3/e_2$ as emergent FIC$_3$ ($e$FIC$_3$), using the same term used by Deshpande et al. (16) in their analysis with the Bliss multiplicative model. (ii) $e_2/e_1$ is a measure of the total effect of three pairwise interactions among three drugs. We define $e_2/e_1$ as lower-order FIC$_2$ ($\lambda$FIC$_2$). Therefore, the total FIC$_3$ score may be factored directly into emergent and lower-order components (see also fig. S7).

Emergent and lower-order scores ($e$FIC$_3$ and $\lambda$FIC$_2$, respectively) are interpreted like the FIC, with 1 for additivity, <1 for synergy, or >1 for antagonism. For example, when each of the drug pairs in a three-way combination is additive, there is no net effect of drug interaction coming from pairwise interactions ($\lambda$FIC$_2 = 1$; Fig. 3B, left). In this case, the interaction score is due to the emergent three-way interaction; hence, FIC$_3 = e$FIC$_3$. When pairs in a three-way combination are synergistic ($\lambda$FIC$_2 < 1$) or antagonistic ($\lambda$FIC$_2 > 1$), this agreement is not necessarily true. For example, if the observed IC$_{50}$ falls between $e_1$ and $e_2$, FIC$_3$ and eFIC$_3$ will have opposite interaction types (indicated by "*" in fig. 3B). We note that both eFIC$_3$ and $\lambda$FIC$_2$ scores for a particular combination are expected to correlate with FIC$_3$, explaining the previous success in predicting three-way synergy using pairwise interactions (17, 18).

$\lambda$FIC$_3$ can be approximated by the geometric mean of the FIC$_2$ scores of the three pairwise combinations (fig. S7). Then, FIC$_3$ can be factorized as a product of eFIC$_3$ and the geometric mean of three FIC$_2$ scores (FIC$_2$)

$$
FIC_3 = \frac{o_3}{e_1} = \frac{o_3}{e_2} \cdot \frac{e_2}{e_1} = eFIC_3 \cdot \lambdaFIC_2 \approx eFIC_3 \cdot FIC_2
$$

The recursive factorization of a four-way drug interaction is shown in method S2. The factorization of an $n$-order drug interaction is generalized from these examples. For $n$-way interactions, the expected plane that accounts for the lower-order interactions ($e_{n-1}$) is defined as the multidimensional plane defined by all the $(n-1)$-way drug combinations. Therefore, we may write the following to factorize an $n$-way drug interaction into two components

$$
FIC_n = \frac{o_n}{e_1} = \frac{o_n}{e_{n-1}} \cdot \frac{e_{n-1}}{e_1} = eFIC_n \cdot \lambdaFIC_{n-1} \approx eFIC_n \cdot FIC_{n-1}
$$

Each of the FIC$_{n-1}$ may also be represented as $(eFIC_{n-1} \cdot FIC_{n-2})$, and the equation above takes a recursive form. It is solved as

$$
FIC_n \approx \frac{FIC_2 \cdot \prod_{k=3}^{n} eFIC_k}{FIC_{n-1}}
$$
Fig. 3. DiaMOND for measuring emergent high-order drug interactions. (A) Each circle indicates an observed IC50 in either single-drug or two-drug dose responses for a three-drug combination. The expected surfaces obtained by connecting the IC50 values observed in two-drug combinations are shown in magenta. For clarity, only one two-drug axis \([X + Y]/2\) is shown as a magenta line. If the observed IC50 of the equipotent mixture of three drugs is smaller than, equal to, or larger than the pairwise expectation, emergent synergy, additivity, or antagonism is concluded, respectively. The blue triangle represents the expected surface given single-drug IC50 values. When pairwise interactions are additive, synergistic, or antagonistic, the expectation given pairs (magenta triangle) will be equal to, smaller than, or larger than the expectation given singles, respectively (blue triangles). (B) Simulated dose responses are given for single drugs, pairwise drugs, and possible outcomes for three-drug dose responses. The blue and magenta triangles mark the additive expectation given single drugs or pairwise combinations, respectively. The top and bottom of the circles representing the observed IC50 values are colored according to the interaction given singles (FICn) and emergent interaction (eFICn) that would be concluded. While two interaction types are the same when drug pairs are additive, a three-drug pair may have opposite interaction types when pairs are synergistic or antagonistic (indicated by **). Simulated dose-response data are shown in blue gradient. (C) All interaction scores for two-, three-, four-, and five-way combinations among five antibiotics are shown as a network representation. Nodes are specific combinations, whose contents are indicated by the barcode. The color of a combination shows the FICn or eFICn (geometric mean of two replicates) of a combination, whereas the size corresponds to deviation from additivity (FICn = 1). Edges are drawn between each combination that differs by one drug.
FIC\textsubscript{2} can also be represented as \((\varepsilon \text{FIC}\textsubscript{2} - \text{FIC}\textsubscript{1})\). Drugs are additive with themselves; therefore, FIC\textsubscript{1} is 1, which makes FIC\textsubscript{2} = \varepsilon \text{FIC}\textsubscript{2}. By substituting this equivalency, Eq. 7 is transformed into the following equation that allows the quantification of each lower-order emergent interaction’s effect on the final high-order drug interaction observed.

\[
\text{FIC}_n \approx \prod_{k=2}^{n} \varepsilon \text{FIC}_k
\]  

(8)

Thus, DiaMOND’s measurements may be simply analyzed to parse total interaction scores into the product of the geometric mean of emergent interaction scores for each lower-interaction group (fig. S7). From this factorization, we can quantify the extent to which each lower-order component influences the \(\text{FIC}_n\) score of a high-order drug interaction.

**Factorization of high-order drug interaction measurements in Mtb**

To understand the relative contribution of the lower-order combinations and the emergent interaction of the high-order combinations to the overall \(\text{FIC}_n\), we used this framework to factorize each of the high-order \(\text{FIC}_n\), we measured above into emergent (\(\varepsilon \text{FIC}_n\)) and lower-order interactions (\(\lambda \text{FIC}_{n-1}\)) (fig. S7). Using the two-, three-, and four-drug dose responses among five TB antibiotics, we computed the \(\varepsilon \text{FIC}_n\) score for each high-order interaction experiment (20 three-way, 10 four-way, and 2 five-way). We used the geometric mean of two replicates as \(\varepsilon \text{FIC}_n\) of a combination to evaluate the interactions (see the Supplementary Materials). The \(\text{FIC}_n\) and \(\varepsilon \text{FIC}_n\) scores for all combinations are shown as nodes on a tree in Fig. 3C, where addition or subtraction of a drug is indicated with an edge. The most striking \(\varepsilon \text{FIC}_n\) score was 1.3 for clofazimine + pretomanid + rifampicin. This emergent antagonism makes this three-way combination additive, despite the expected synergy given pairwise combinations. To a weaker extent, the same explanation applies for the five-way combination, for which synergy would be the expectation given lower-order interactions. An antagonistic \(\varepsilon \text{FIC}_5\) score of 1.14 pushes the \(\text{FIC}_5\) score to 0.93, close to an additive interaction. \(\text{FIC}_4\) and \(\varepsilon \text{FIC}_4\) scores for clofazimine + isoniazid + pretomanid + rifampicin were 0.78 and 0.9, respectively, indicating that this four-way combination is synergistic and also has emergent synergy. This contrasts with the three-way synergy of bedaquiline + clofazimine + rifampicin, which is synergistic \((\text{FIC}_3 = 0.78)\), but has no emergent synergy \((\varepsilon \text{FIC}_3 = 1.06)\), suggesting that this three-way synergy is primarily due to pairwise synergies.

To understand how much the underlying components contribute to the overall higher-order interactions, we next evaluated the correlations of emergent and lower-order interaction scores to overall \(\text{FIC}_n\), in our full experimental set (fig. S7). Emergent interaction scores \(\varepsilon \text{FIC}_n\) significantly correlated with total interaction scores \(\text{FIC}_n\) (Spearman’s \(r = 0.45, P = 9.6 \times 10^{-3}\); Fig. 4A). Therefore, emergent interactions, which are independent of low-order interactions, are an important component of a high-order drug interaction. We next calculated a lower-order interaction score (\(\lambda \text{FIC}_{n-1}\)) for each combination by dividing \(\text{FIC}_n\) with \(\varepsilon \text{FIC}_n\). \(\lambda \text{FIC}_{n-1}\) scores correlated with \(\text{FIC}_n\), indicating that lower-order interactions also correlated with high-order interaction scores \((r = 0.55, P < 1.0 \times 10^{-3})\). However, \(\varepsilon \text{FIC}_n\) and \(\lambda \text{FIC}_{n-1}\) scores did not have a significant correlation \((r = -0.35, P = 5.2 \times 10^{-2})\). Together, our analysis suggests that both lower-order interactions and emergent interactions correlate with the overall higher-order interaction, but that the lower-order and emergent interactions are independent of each other.

Finally, we used our measured and calculated component scores to test the theoretical predictions made by our factorization methodology. \(\lambda \text{FIC}_{n-1}\) significantly correlated with the geometric mean of \(\text{FIC}_{n-1}\) \((\text{FIC}_{n-1}; \quad r = 0.97, P = 2.8 \times 10^{-23}\); figs. S7 and S8), verifying the approximation used in Eq. 6. The multiplication of \(\varepsilon \text{FIC}_n\) and \(\text{FIC}_{n-1}\) also significantly correlated with \(\text{FIC}_n\) \((r = 0.97, P = 3.1 \times 10^{-20}\); Fig. 4B), in agreement with Eq. 8.

**Prediction of high-order interactions using lower-order emergent drug interactions**

With DiaMOND, we have established the methodology for efficient measurement and analysis of pairwise and high-order drug combinations. However, systematic searches for synergistic high-order combinations are impeded by the large number of combinations, and even the improved efficiency of DiaMOND’s measurement procedure cannot fully enable systematic experiments. Previous work has shown that pairwise interactions could largely predict three-way drug interactions (5, 6). These studies suggest that prediction of three-way interactions from the nature of the underlying pairwise interactions, if quantitatively tractable, may enable us to enrich candidates for synergistic three-way combinations for further measurement.

We next aimed to use our high-order drug interaction scores and their factorization into emergent and lower-order contributions to test
the hypothesis that lower-order interactions could be used for data-driven prediction of high-order interactions. The factorization of high-order interactions provides an immediate way to predict $FIC_n$ (see above). Experimentally determined emergent interaction scores ($\epsilon FIC_n$) can be used in Eq. 8 to produce an approximation for $FIC_n$ using the definition of $\epsilon FIC_2 = FIC_2$ to simplify the recursive formula. As we have already determined all $\epsilon FIC_2$, $\epsilon FIC_3$, $\epsilon FIC_4$, and $\epsilon FIC_5$ scores for all combinations in our screen, we next calculated the $\epsilon FIC_n$ scores for all three-way, four-way, and five-way interactions; the $\epsilon FIC_3$ scores for all four-way and five-way interactions; and the $\epsilon FIC_5$ score for the five-way interaction. According to Eq. 8, each of these $\epsilon FIC_n$ score sets equally influences a high-order drug interaction score $FIC_n$. The multiplication of each of these $\epsilon FIC_n$ score sets and $\epsilon FIC_n$ of a high-order combination should approximate the $FIC_n$ of the combination (see fig. S7 for examples of $\epsilon FIC_3$ and $FIC_3$ calculation). The final approximation of $FIC_n$ depends on the data quality of experimental measurements of lower-order interactions, highlighting the importance of reproducibility in the prediction of high-order drug interactions.

When all combinations were considered, pairwise scores ($\epsilon FIC_2$) did not significantly correlate with high-order interaction scores ($FIC_2$, $r = 0.33$, $P = 0.06$; Fig. 4C). However, when only three-way interactions were considered (triangles), pairwise interaction scores ($\epsilon FIC_2$) significantly correlated with the interactions at high order ($FIC_2$, $r = 0.55$, $P = 1.2 \times 10^{-2}$). These observations suggest that a simple geometric mean of all $FIC_2$ scores is a good predictor of three-way interaction scores, but that pairwise scores could not predict interactions of more than three drugs. In contrast, we find a significant but weaker correlation when $\epsilon FIC_3$ and $\epsilon FIC_4$ scores are included to predict $FIC_n$ for four-way and five-way interactions, respectively ($r = 0.45$, $P = 9.9 \times 10^{-3}$), suggesting that lower-order interactions influence but do not strongly predict overall $FIC$. This suggests that the knowledge of all lower-order emergent interactions has some predictive potential, even in the absence of $\epsilon FIC_n$. When the high-order combination’s $\epsilon FIC_n$ is also considered, the agreement between prediction and experiments was very strong ($r = 0.90$, $P = 1.6 \times 10^{-12}$) (Fig. 4D). Therefore, we conclude that the prediction of high-order drug interactions can be achieved by systematically measuring emergent interaction scores. These scores can be combined in Eq. 8 to make interaction predictions on high-order combinations and rank these combinations by likelihood of synergy.

**DISCUSSION**

Here, we present DiaMOND, a methodology to make robust measurements for pairwise or higher-order drug interactions. DiaMOND relies on the construction of dose responses for drug mixtures and comparison with the dose response of component drugs under the Loewe additivity model for drug interactions. Sampling a small part of the traditionally used experimental setup allows efficient measurements for pairwise or high-order drug interactions. DiaMOND’s efficiency also enables the practical use of Loewe additivity in producing immediately interpretable and widely used $FIC$ interaction scores for combinations of many drugs. Drug interaction measurement by DiaMOND is broadly applicable for any agent that results in a specified phenotype. Here, we successfully applied DiaMOND to measure pairwise and high-order interactions among antibiotics effective against Mtb. We propose DiaMOND as an alternative to checkerboard assays for drug interaction measurements, especially for high-order interactions. The robustness of DiaMOND, despite the difficulty of working with slow-growing mycobacteria, suggests that DiaMOND may be applied to other challenging experimental systems.

With precise measurement of interactions among many drugs, we can evaluate the current guidelines and consider alternative regimens in search of better drug combination options for latent, drug-sensitive, and drug-resistant TB. Here, we uncovered surprisingly strong drug synergies including pairwise (bedaquiline + clofazimine, pretomanid + rifampicin), three-way (bedaquiline + clofazimine + isoniazid), and four-way (bedaquiline + clofazimine + isoniazid + rifampicin) combinations. Interestingly, none of the high-order interactions were as synergistic as pairwise synergies. However, note that multidrug regimens are often used to block resistance evolution paths (31). Although the identified three-way and four-way synergistic combinations have weaker synergy than pairwise combinations, they may be more effective against resistance. Of particular interest is the tendency of moxifloxacin to show frequent antagonism, including with rifampicin (32) or pretomanid, because moxifloxacin is used in combination with these antibiotics in clinical trials (33, 34). These observations are a reminder that synergy that is observed in vitro is not necessarily associated with a better treatment, because resistance evolution, adverse drug interactions, combinatorial pharmacokinetics/pharmacodynamics, and other considerations may influence treatment success. Given the toxicity and adverse side effects of many second-line antibiotics and drugs in the development pipeline of TB therapeutics, a detailed, systematic, and quantitatively rigorous understanding of drug synergies and antagonisms may help accelerate the design of multidrug regimens for testing in animals and in adaptive clinical trials (3).

Apart from measuring a high-order interaction, DiaMOND can also factorize a high-order drug interaction into components. Here, we characterize a straightforward relationship between the three basic types of interaction scores: The total score is a product of the emergent action of the high-order interaction and the geometric means of each lower-order interaction group. This breakdown of interaction scores, applied to our compendium of drug interaction scores in Mtb, enabled us to address several important questions. First, we established that lower-order interactions are moderate predictors of high-order interactions. This may help us use easy-to-measure pairwise interactions to apply data-driven design of screens for high-order drug interactions instead of exhaustive searches. Even with the efficacy of DiaMOND, an exhaustive search of all high-order combinations among 20 drugs is not practical because this would take 100,000 microplates, whereas a comprehensive pairwise interaction screen requires many fewer plates (by four orders of magnitude). Second, despite the good predictive value of low-order interactions to higher orders, there were strong emergent interaction scores, highlighting the importance of experimental verification of high-order drug interaction measurements in the design of multidrug regimens.

DiaMOND’s conceptual framework to study specific drug interactions may be adapted to study interactions in a more nuanced way. For example, in TB, some interactions can be synergistic or antagonistic in animal models or during the course of clinical treatment depending on treatment timing and dose (16, 28–30). The dose dependency may be due to scoring by the Bliss methodology, which can yield fundamentally different interaction scores depending on drug concentration, or the dose dependencies may be due to contours of the checkerboard that are convex in one combination-dose region and concave in another. The DiaMOND methodology can be customized to identify interactions that are inherently dose dependent by expanding the regions of geometric sampling.
Combination therapy is used not only to treat TB and other infectious diseases but also for many cancers. With combination therapies, we have the opportunity to combine synergistic drugs but have been limited by a practical inability to systematically measure and prioritize drug combinations for testing. The DiaMOND methodology is efficient, inexpensive, and intuitive, and therefore may be used to develop a more balanced emphasis on data-driven modeling and experimental verification in preclinical drug regimen design for a broad range of disease models.

**METHODS**

**Strains and culturing**

A panthoanen and leucine auxotrophic strain of Mtb H37Rv was used for experiments (29). Bacteria were grown to mid-log phase with shaking at 37°C, in 7H9 medium supplemented with 0.05% Tween 80, 0.2% glycerol, 10% BBL Middlebrook OADC enrichment (VWR), leucine (50 μg/ml), and panthoanen (24 μg/ml).

**Growth inhibition assays**

Drugs were dissolved in dimethyl sulfoxide and stored at −20°C. The minimum inhibitory concentration for each drug is given in Table 1. Assays were performed in clear, flat-bottom 384-well microplates by dispensing nanoliter volumes of drugs using a digital drug dispenser (D300e Digital Dispenser, HP). Dispense locations were randomized within each plate using the software to minimize the plate position effect. Mid-log phase cultures were diluted to an optical density (OD) of 0.05 in fresh supplemented 7H9 medium, and 50-μl diluted cells were added to each well. Plates were sealed with aluminum plate seals and incubated without shaking at 37°C. After 5 days, aluminum seals were replaced with optically clear plate seals, and growth measurements (OD600) were made (Synergy HT, BioTek).

**Data analysis**

Using the randomization map from the digital dispense software, plate measurement data were reconstituted and analyzed (MATLAB, Mathworks), as described in Results and methods S1 and S2.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/10/e1701881/DC1

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