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Exploring the Mode of Action of Bioactive Compounds by Microfluidic Transcriptional Profiling in Mycobacteria

Paul Murima1, Paola Florez de Sessions2*, Vivian Lim1, Ahmad Nazri Mohamed Naim2, Pablo Bifani1, Helena I. M. Boshoff3, Vasan K. Sambandamurthy1*, Thomas Dick1,4, Martin L. Hibberd2, Mark Schreiber1,5*, Srinivasa P. S. Rao1*

1 Novartis Institute for Tropical Diseases, Singapore, Singapore, 2 Genome Institute of Singapore, A*STAR, Singapore, Singapore, 3 Tuberculosis Research Section, Laboratory of Clinical Infectious Diseases, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Bethesda, Maryland, United States, 4 Department of Microbiology, National University of Singapore, Singapore, Singapore, 5 Novartis Institutes For Biomedical Research, Cambridge, Massachusetts, United States of America

Abstract

Most candidate anti-bacterials are identified on the basis of their whole cell anti-bacterial activity. A critical bottleneck in the early discovery of novel anti-bacterials is tracking the structure activity relationship (SAR) of the novel compounds synthesized during the hit to lead and lead optimization stage. It is often very difficult for medicinal chemists to visualize if the novel compounds synthesized for understanding SAR of a particular scaffold have similar molecular mechanism of action (MoA) as that of the initial hit. The elucidation of the molecular MoA of bioactive inhibitors is critical. Here, a new strategy and routine assay for MoA de-convolution, using a microfluidic platform for transcriptional profiling of bacterial response to inhibitors with whole cell activity has been presented. First a reference transcriptome compendium of Mycobacterial response to various clinical and investigational drugs was built. Using feature reduction, it was demonstrated that subsets of biomarker genes representative of the whole genome are sufficient for MoA classification and deconvolution in a medium-throughput microfluidic format ultimately leading to a cost effective and rapid tool for routine antibacterial drug-discovery programs.

Introduction

Since the early 20th century, bioactive inhibitors used for anti-infective chemotherapy have been identified by phenotypic screens and further examined in complex biological systems [1]. Advances in genome sequencing, molecular biology and biochemistry led to an evolution from the traditional phenotypic screens to a more ‘reductionist’ target-based approach, which was thought to be more rational and efficient [2]. Despite the rapid identification of diverse, novel drug targets characterized by genetic tools [3], target-based anti-bacterial lead discovery has been less successful [4–6]. In many cases, these target-based screens reveal small molecules with potent activity against the purified target in vitro but fail to render anti-bacterial activity in both in vitro and in vivo models [4,7].

The large-scale failure of genomics driven anti-bacterial lead discovery programs has led to the renaissance of empirical phenotypic screens for the identification of new chemotypes [6,8,9]. In contrast to target-based screening, molecules identified using this approach have the advantage of not only possessing desirable physicochemical properties from the beginning (such as cell penetration), but are also active against the relevant target in its intracellular context, under physiological conditions. Despite this key advantage, success in defining the target, mechanism of action (MoA), and the final lead optimization of hits derived from phenotypic screens has been low [4,6].

One of the daunting tasks for medicinal chemists during hit to lead and lead optimization of hits, and scaffolds derived from whole cell screen, is to make sure that the compounds they are synthesizing also have similar MoA as that of the parent molecule. In order to understand the structure activity and property relationship (SAR and SPR) medicinal chemists synthesize multiple compounds in and around the parent molecule. It is very critical that the new molecules are acting in a similar way as that of the parent in order to get desired final effect. Currently, lead optimization of hits from phenotypic screens can only be best done with a known target. Although various approaches for MoA and target deconvolution have been established, including...
characterization of resistant mutants, biochemical affinity-based methods, genetic complementation, protein and DNA microarrays [10], target identification is still a challenging and inefficient task to support the early discovery process [6].

Until the last decade, MoA deconvolution was largely limited to model organisms whose metabolic pathways have been well characterized. Transcriptional profiling by microarray analysis has been used to analyze the MoA of early anti-bacterial [11,12], anti-fungal [13], and anti-malarial compounds [14]. Despite the elegance of this approach for MoA deconvolution, it is not practical for use as a routine assay [15–17]. To benefit from the transcriptional profiling body of evidence we have established a miniaturized gene expression assay for efficient MoA deconvolution and discovery chemistry based on microfluidics. The microfluidic integrated fluidic circuits (IFC) contain tens of thousands of microfluidic-controlled valves and interconnected channels for transporting and combining cDNA molecules and qPCR reagents in complex patterns [18]. As a result of the miniaturization inherent in this approach, a single assay is capable of increasing the throughput of traditional qPCR by 2 orders magnitude using nanolitre reaction volumes compared to the standard techniques [19]. Here we report the application of this tool as a routine assay for MoA deconvolution, and its help in hit to lead and lead optimization of novel compounds obtained by phenotypic screens. We demonstrate that a minimal number of differentially expressed genes are sufficient to classify the MoA of novel chemical entities (NCE).

**Materials and Methods**

**Bacterial cultures and RNA extraction**

*Mycobacterium tuberculosis* (ATCC 27294) and *M. bovis* BCG (Pasteur) were grown with aeration at 37°C in Middlebrook 7H9 (Difco) liquid culture medium supplemented with 0.5% (w/v) bovine serum albumin fraction V, 0.2% dextrose, 0.08% sodium chloride, 0.5% glycerol, 0.5% glycerol, and 0.05%, tween 80 to mid-log phase. The mid log phase culture was concentrated and re-suspended to A_{600} nm of 0.3 (Amersham Ultrospec 3300). Antimicrobial compounds were added at either 0.5×, 1×, or 5× minimum inhibitory concentration (MIC) determined using a turbidimetric microplate assay. For microarray experiments *M. tuberculosis* H37Rv strain was anti-tubercular compounds were treated at different concentrations for 6 hours prior to isolation of RNA for analysis. All the details are provided in GEO accession # GSE46212. For real time qPCR experiments, *M. bovis* BCG cells were treated at 0.5× MIC_{50} of various anti-tubercular agents for 3 hours prior to isolation of RNA. Dimethyl sulfoxide (DMSO, drug vehicle) treated mycobacterial cultures were used as untreated controls in all experiments. The culture of cells treated/untreated with the inhibitor were pelleted in RNAlater (Qiagen) and bacterial RNA were extracted using the GTC/Trizol® method [20].
Table 1. List of investigational and FDA approved anti-tubercular drugs and their mode of action.

| Compounds       | Mode of action | BCG MIC<sub>50</sub> (µM) |
|-----------------|----------------|-----------------------------|
| NITD101         | Unknown        | -                           |
| NITD102         | Unknown        | -                           |
| NCE1a (AD841)   | Unknown        | 0.02                        |
| NCE1b (AD1015)  | Unknown        | 0.04                        |
| NCE1c (AD842)   | Unknown        | 0.2                         |
| NCE2a (AD1158)  | Unknown        | 0.38                        |
| NCE2b (AD1441)  | Unknown        | 1.28                        |
| NCE2c (AD1259)  | Unknown        | 2.06                        |
| NCE2d (AD1260)  | Unknown        | 2.04                        |
| NCE2e (AD1534)  | Unknown        | 7.2                         |
| NCE2f (AD1540)  | Unknown        | 0.44                        |
| NCE3a (NITD_TB034) | Unknown | 10                          |
| NCE3b (ATP425)  | Unknown        | 1.6                         |
| NCE3c (ATP359)  | Unknown        | 6                           |
| NCE3d (ATP435)  | Unknown        | 2                           |
| NCE3e (ATP355)  | Unknown        | 10                          |
| Cyclomarin-     | Targets the ClpC1 subunit of the Caseinolytic Protease |
| Series          |                |                             |
| Liparmycin-     | DNA-dependent RNA polymerase inhibitor |
| Series          |                |                             |
| PA-824          | Possibly acts via generation of radicals having nonspecific toxic effects. Might possibly target enzymes in mycolic acid & protein biosynthesis. | 0.44 |
| Linezolid       | Binds to the A site of the 50S subunit, and prevents the assembly of the ribosome initiation complex. | 1.52 |
| Ethambutol      | Cell wall inhibitor, inhibits arabinosyl transferases involved in cell wall biosynthesis. | 2.96 |
| Isoniazid       | Cell wall inhibitor. Isoniazid inhibits inhA, a NADH-specific enoyl-ACP coA reductase involved in mycolic acid synthesis. | 0.28 |
| Ethionamide     | Cell wall inhibitor, disrupts of mycolic acid biosynthesis | 2.04 |
| Prothionamide   | Cell wall inhibitor | 0.9 |
| Amikacin        | **Aminoglycosides**: Inhibit translocation of the peptidyl-tRNA from the conserved A-site to the P-site in the 30S ribosomal Subunit. | 0.38 |
| Capreomycin     | 1              |                             |
| Streptomycin    | 0.16           |                             |
| Kanamycin       | 1.56           |                             |
| Clofazimine     | Binds to guanine bases of bacterial DNA, causing steric hindrance in the template function of the DNA. Causes the accumulation of toxic lysosphospholipids which inhibit bacterial growth. | 0.26 |
| Ampicillin      | **Penicillin**: Beta lactam antibiotic. Penicillins bind to transpeptidase enzymes (which are involved in the continuous remodelling of the peptidoglycan layer by cross linking peptides chains). [Note transpeptidase catalyses the terminal step of cell wall biosynthesis by cross linking the peptidoglycan]. By inhibiting this enzyme, penicillin prevents the formation of peptide bonds, weakens the cell wall, which subsequently cause lysis of the cell membrane. | 6.76 |
| Moxifloxacin    | **Quinolones**: Inhibit the bacterial ATP-dependent enzyme DNA gyrase and topoisomerase IV, preventing DNA unwinding during bacterial replication, transcription and bacterial DNA repair. | 0.28 |
| Ofloxacin       | 1.56           |                             |
| Levofloxacin    | 0.84           |                             |
| Gatifloxacin    | 0.2            |                             |
| Sparfloxacin    | 0.22           |                             |
| Valnemulin      | **Pleuromutilin**: Binds to the peptidyl transferase component of the 50S subunit of ribosomes. | 6.76 |
| Vancomycin      | **Glycopeptide**: Vancomycin binds with high affinity to the D-Ala-D-Ala C-terminus of the pentapeptide, thereby blocking the addition of late precursors by transglycosylation to the nascent peptidoglycan chain and preventing subsequent cross-linking by transpeptidation | 14.08 |
| Rifampicin      | Inhibits the essential rpoB gene product a subunit of DNA-dependent RNA polymerase activity | 0.02 |
| Rifabutin       | 0.02           |                             |
| Rifapentene     | 0.02           |                             |
| Thioridazine    | A phenothiazine interfering with NADH dehydrogenase | 14.08 |
| TMC207          | **Diarlyquinoline** which inhibits the proton pump of ATP synthase. | 0.32 |
Table 1. Cont.

| Compounds | Mode of action | BCG MIC\textsubscript{50} (\textgreek{M}) |
|------------|----------------|----------------------------------|
| Tetracycline | Bind to the 30S ribosomal subunit in the mRNA translation complex thus, inhibiting binding of aminoacyl-tRNA to the mRNA-ribosome complex. | |
| p-aminosalicylic acid (PAS) | Inhibitor of folic acid biosynthesis pathway | 0.56 |

MIC\textsubscript{50} listed for anti-tubercular investigational and SAR compounds, when available.
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Quantitative reverse transcription qPCR assay

The expression of genes selected as probes to represent the whole genome was quantified using real time qPCR, after normalizing the RNA expression levels to SigA as previously described [11].

Microarray gene expression

Microarray analysis was performed as described [11]. The microarray data obtained for the compounds has been submitted to NCBI (GEO accession # GSE46212 and GPL1343). Expression ratios were calculated as the feature pixel median minus background pixel median for one color channel divided by the same for the other channel. In cases where more than 10% of the feature pixels were saturated, the feature pixel mean was used instead of the median. When the feature pixel mean did not exceed the background pixel mean by more than two standard deviation values (calculated from the background pixel distribution), the feature pixel median was used in the ratio without background subtraction. In cases where both color channels were near background (same criterion), the ratio value was set to “missing.” Expression ratios were transformed to the log base 2 for all further calculations.

Microfluidic gene expression analysis

Using the previously described BioMark Fluidigm 96.96 microfluidic chip (Fluidigm, South San Francisco, Calif., USA) [19], we analyzed bacterial transcriptional response in the antibiotic or vehicle treated cells. Total RNA was isolated using Trizol and RNA was quantified using a Nanodrop and adjusted to 150 ng for reverse transcription using High-Capacity RT Kit (Applied Biosystems, 4368813), as per the manufacture’s recommendations. Pre-amplification of the cDNA was performed using a Taqman PreAmp Master Mix Kit (Applied Biosystems, 4391128) as per the manufacture’s recommendations, and the preamplified product was diluted at 1:20. The pre-amplified product was used for the BioMark 96.96 Real-time PCR assay (Fluidigm) according to the manufacture’s recommendations using Eva green DNA binding dye (Biotium). Fluidigm uses integrated fluidic circuits to generate a medium throughput for real-time PCR. In the BioMark platform 96 primer sets and 96 samples can be loaded and via the on-chip network of microfluidic channels, chambers and valves individual PCR reactions will be assembled simultaneously (a maximum of 9,216 reactions can be evaluated at one time). The complete mechanism of how microfluidic circuit and valve system technology work have been well described elsewhere (http://www.fluidigm.com/technology.html).

Data processing and analysis

To identify differentially regulated genes, a measure of significance t-test was applied to the normalized data set. To discriminate genes that significantly deviated from the 1:1 ratio (treated:vehicle), a minimum p-value of 0.05 was used. Differentially expressed genes were subsequently subjected to Benjamini and Hochberg correction to account for multiple experimental testing.

Feature reduction

Feature reduction is the process of reducing the number of variables (in this case genes) defining a condition. Feature reduction often attempts to identify a minimum set of non-redundant features that are useful for classification. Microarray data of mycobacteria treated with a variety of antibacterial compounds were hierarchically clustered using Euclidian distance as a distance matrix and a minimal variance method (Ward linkage) to join adjacent clusters [21]. It can be summarized by the following five steps: a) Distances between all gene pairs are calculated, using Euclidean distance, b) The resulting distance matrix is thoroughly inspected to find the smallest distance between expression profiles, c) The corresponding genes are joined together in the tree and form a new cluster, d) The distances between the newly formed cluster and the other genes are recalculated, e) Steps b, c and d are iteratively repeated until all genes and clusters are linked in a final tree.

Figure 2. Transcriptional profile of gene clusters in response to antibiotics. X-axis denotes different drugs, and the y-axis denotes relative expression of the gene in response to the particular drug on the x-axis. Whole genome microarray data were used for this analysis. (A) A cluster replete with genes in respiratory metabolic pathway as well as subunits of the secretion system. The ESX section associated protein EspC was selected as the gene best representing the cluster transcription profile. (B) Genes of a related metabolic pathway cluster together in response to antibiotics. All three genes are from one metabolic pathway. NadB involved in quinolone biosynthesis served as the representative of the cluster.
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A hierarchical cut off was selected that gave 90 gene clusters. A single biomarker gene to represent each cluster was identified by finding the gene in the cluster with the best Pearson correlation against the cluster mean. In total, 90 genes were selected optimizing the number of biomarkers measured in the 96 probe fluidigm format while allowing for up to 6 control probes. The reduction method included following steps; a) computation of the mean expression of genes in a cluster in all conditions, b) computation of Pearson product moment correlation coefficient to the cluster mean for each gene, and c) the selection of gene with the best correlation co-efficient in each cluster as the feature to represent the cluster.

Figure 3. Transcriptional responses to drug treatment reveal that a single gene representing logically associated gene clusters is sufficient for MoA diagnosis in *Mycobacterium tuberculosis*. Groups of drugs clustered separately based on the known mechanism of action, while SAR series of compounds with novel MoA (cyclomarin) showed a fingerprint distinct from any of the compounds tested. Periods following drug names represent duplicates. X-axis represents the 90 genes that were chosen from the microarray as representative of 90 clusters, y-axis lists the different drug treatments. i, ii, and iii, represent 3 clusters namely, cell wall inhibitor, RNA polymerase inhibitor and Cyclomarin series, respectively. doi:10.1371/journal.pone.0069191.g003

| Class ID | Functional Class                          | PCR array | Genome (H37Rv) |
|----------|------------------------------------------|-----------|----------------|
|          | Virulence, detoxification & adaptation    | 10.59%    | 5.22%          |
| 1        | Lipid metabolism                         | 5.88%     | 5.84%          |
| 2        | Information pathways                     | 7.06%     | 5.17%          |
| 3        | Cell wall and cell processes             | 20%       | 18.49%         |
| 5        | Insertion sequences & phages             | 2.35%     | 3.62%          |
| 6        | PPE/PE                                   | 3.53%     | 4.14%          |
| 7        | Intermediary metabolism & respiration    | 15.29%    | 22.11%         |
| 9        | Regulatory proteins                      | 4.08%     | 4.78%          |
| 10       | Conserved hypotheticals                  | 15.29%    | 22.06%         |
| 16       | Conserved hypotheticals with an orthologue in *M.bovis* (BCG) | 8.24% | 6.43% |
| -        | Not annotated on H37Rv                   | 7.06%     | 7.06%          |

Table 2. Pathway enrichment of functional classes of genes on PCR array.

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The robustness of experimental set up was evaluated using Pearson correlation coefficient of expression values between two independent biological duplicates. Similarities in the expression profiles were analyzed by generating heat maps using R [22]. Correlation similarity matrix comparing different compounds, or SAR analogues was evaluated using both R and the Pearson product moment correlation coefficient.

Results

A feature reduced compendium of chemical-genetic profiles

We established the transcriptional response of \textit{M. tuberculosis} to 12 different inhibitors. Included in this collection were 5 synthetic compounds and natural products derived from phenotypic screens, the rest were FDA approved drugs with anti-tubercular activity. The list of inhibitors used for perturbing bacteria and MoA diagnosis are included in Table 1. Microarray results showed differential expression for 1860 genes out of 4734 genes that were used in the array. (GEO accession # GSE46212 and GPL1343). The hierarchical heat map of bacterial transcriptional response of 1860 genes to the list of antibacterials showed that groups of drugs clustered separately based on their known mechanism of action (Fig. 1). Thus, protein synthesis inhibitors, transcriptional inhibitors, cell wall synthesis inhibitors and novel antibacterials fell into distinct groups. Boshoff and co-workers had previously described the application of transcriptional profiling using microarrays for MoA diagnosis of a natural product extract with no known mode of action [11]. However, the approach is not easily scalable and suitable for routine MoA diagnosis for hundreds of compounds from phenotypic screening. To exploit this property amenable for phenotypic screening, we limited the number of genes defining a condition to 90 using feature reduction.

Feature reduction and real-time qPCR for MoA deconvolution

Feature reduction attempts to identify a set of non-redundant genes that are useful for MoA classification. To reduce the features amenable to a robust 96 well format, a total of 90 genes from 90 clusters derived from agglomerative hierarchical clustering were selected. Briefly, the expression profiles of single genes was successfully joined to form nodes, which in turn were joined further until a total of ninety clusters were obtained. At the finest level, redundant gene representations across all perturbations clustered together with genes in each cluster showing similar expression profile. Two representative pattern of transcriptional profile of gene clusters in response to various antibiotics are shown in Fig. 2A & B. However despite the similar expression patterns, individual genes could be distinguished from each other with subtle differences in their expression profiles (Fig. 2A). To reduce the number of features a single gene was selected from each cluster that best represented the cluster expression profile by finding the gene most correlated with the mean expression profile of the cluster. The 90 genes selected with their known function are listed in Table S1.

To validate the functionality of the selected gene probes for MoA diagnosis, we analyzed bacterial transcriptional response to the chemical perturbation from microarrays. Hierarchical clustering analysis of the transcription reference compendium revealed that 90 genes are sufficient to define the MoA of hits derived from phenotypic screens (Fig. 3). We found that compounds with similar cellular effects showed similar transcription fingerprints and thereby cluster together on the vertical axis in revealing both anticipated and novel insights into their mode-of-action. In particular there were a couple of examples where the cluster analysis grouped inhibitors targeting the same pathway or target (Fig. 3). Individual clusters are annotated by roman numbers: (i) Cell wall inhibitors: Ampicillin (peptidoglycan) and Isoniazid.
(Mycolic acids); (ii) RNA polymerase inhibitors: Rifampicin and Lipiarmycin [23], and (iii) a series of ClpC1 caseinolytic proteasome inhibitors cyclomarin whose mode of action is novel [24]. The quality of the data was tested using Pearson rank correlation and Euclidean Distance, which verified that the expression fingerprint within each compound perturbation was tightly correlated (Fig. S1).

MoA deconvolution of a natural product with antibacterial activity identified by chemical genetics

Given that, transcriptional profiling is applicable to any molecule that impairs bacterial growth, we evaluated the MoA of the natural product cyclomarin. To test whether chemical-genetic profiling may be particularly useful in driving the SAR for a molecule whose MoA was distinct from the compendium, we analyzed the transcriptional fingerprint of different derivatives of the natural product. All derivatives had a tight correlation, showing a similar fingerprint suggesting that they all had the same mode of action, and were probably targeting the same target (Fig. 3 & S1, Fig. 1). A detailed study on the elucidation of the MoA of the natural product cyclomarin is described elsewhere [24].

To validate the functional representation of the selected biomarker genes using qPCR, we evaluated the functional enrichment of genes. Table 2 summarizes the gene function enrichment of biomarker genes. Functional classes were defined according to Cole et al [25]. Gene function enrichment revealed that the dominant feature was a large percentage of virulence, detoxification and adaptation genes more than twice the expected number by chance. Genes from this functional class include; a heat shock protein groES, some conserved hypothetical proteins with PIN domains, a sub unit of the alkyl-hydrogen peroxide reductase ahpC involved in the oxidative stress response and genes in toxin-anti toxin operons. This demonstrates that the feature reduction can successfully enrich for genes involved in stress responses [26] as well as central pathways targeted by antibiotics [27].

One of the challenges of working with microarrays for MoA deconvolution is large sample volume, the need for multiple replicates, further compounded by the noise inherent in these systems. As a proof of concept to circumvent these properties, application of using oligos to PCR amplify the 90 selected genes as biomarkers to represent the whole genome was evaluated. A small collection of anti-tubercular drugs were analyzed by qPCR arrays using M. bovis BCG bacterial RNA. Analysis of the qPCR arrays showed that expression profiles from this platform were able to distinguish the MoA between anti-bacterial drugs (Fig. 4A & B). The minimal gene list to decipher the antibiotic mechanism of action on a PCR array has been summarized in Table S1.

Figure 5. Hierarchical clustering and correlations of anti-tubercular drugs. Transcriptional responses data obtained from microfluidic experiment are used for the analysis. The detailed descriptions for genes represented in this figure are provided in Table S1. (A) Hierarchical clustering via average linkage of Pearson correlations for FDA approved drugs. The individual genes are represented in y-axis and the compound treatment is in the x-axis. (B) Pearson correlations for 23 anti-tubercular drugs. Correlations were calculated between 2 independent microfluidic experiments, following median normalization to account for plate effects.
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Despite the excellent dynamic range, sensitivity and reproducibility of data using real-time qPCR, this technique is a low-throughput and expensive method to analyze a limited number of genes. Microfluidic digital PCR is able to generate higher-throughput data technically identical in quality to the standard qPCR [19]. We explored the use of microfluidic chips to study the bacterial transcriptional response to a collection of anti-tubercular agents. Microfluidics as Tool to Study Antimycobacterials

Figure 6. Transcriptional response of selected 90 genes to novel chemical entities. The heatmap was drawn using Pearson correlations through single linkage for both genes and compound treatments. The individual genes are represented in y-axis and the compound treatment is in the x-axis. Compounds from the same SAR library (designated as NCE1 and NCE2) show a similar expression fingerprint, while NCE3 SAR compound series displays more variability in expression. The detailed descriptions for genes represented in this figure are provided in Table S1.

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investigational and 23 FDA approved drugs (Table 1). As expected, drugs with similar mode of action, targeting the same cellular organelles clustered together (Fig. 5A). For example the fluoroquinolones clustered together, which inhibit the bacterial ATP-dependent enzyme DNA gyrase and topoisomerase IV, preventing DNA unwinding during bacterial replication, transcription and bacterial DNA repair. To validate the dynamic microfluidic PCR as a platform for routine gene expression analysis, the variation between and within measurement platforms for all genes under investigation using a subset of drugs was analyzed. The correlation between independent biological assays was as good as qPCR readouts (Fig. 5B).

Driving SAR of a chemical genetics hits using microfluidic arrays

The principal advantage of a cell-based screen is that it represents an unbiased picture of both known and unknown cellular pathways that a molecule can modulate. The disadvantage of this empirical approach is that the target remains unknown. Hit to lead and lead optimization is generally more efficient when the target is known, thus target deconvolution is essential despite being time consuming and tedious. During SAR expansion, derivative molecules of the primary compound are made with a greater potency than the starting hit molecule (parent hit). It is imperative to monitor that these derivatives inhibit the same target or similar MoA, meaning that they have a similar transcription fingerprint as the parent hit molecule. In order to evaluate if SAR analyses can be improved for a compound with anti-tubercular activity, we evaluated the transcription fingerprint of a series of derivatives around particular pharmacophores. We hypothesized that a change in the transcription profile would indicate a change of target or MoA. To investigate this, a subset of an SAR expansion of NCE1, NCE2, and NCE3 were analyzed. SAR groups NCE1 and NCE2 showed a similar transcription fingerprint, indicating that they probably were still “on target” (Fig. 6 & S2). The NCE3 series displayed more variability in expression in our feature reduction microfluidic experiment; this is evident in both the heatmap and the correlation matrix (Figure 6 & S2). All derivatives had anti-bacterial activity (Table 1).

Discussion

There is an urgent need for new therapeutic approaches to antibiotic discovery as a result of the rapid emergence of multi-resistance to existing drugs, which is further exacerbated by the current gap in the development of new molecules. Phenotypic screening has re-emerged as the standard anti-bacterial drug discovery approach to avoid some of the attrition in target-based lead identification. When successful, it provides lead compounds with proven in vivo activity against the organism of interest [8,9]. This approach has produced a large set of starting chemical points for anti-bacterial & anti-parasitic lead discovery [14,28–32]. However, the cost of not having a biochemical assay for whole cell activity is that hit to lead and lead optimization becomes more challenging.

It is more difficult to establish consistent structure-activity relationships from phenotypic screens because whole-cell potency is a composite result of several compound properties, such as target affinity, cell permeability, efflux, and or intracellular modifications. Nonetheless, evidence supports the effectiveness of this approach in the identification of novel chemical classes with anti-bacterial or anti-malarial activity [23,24,28–32]. The key to success is a good chemical starting point and an adequate progression path to identify the potential to become a drug as quickly as possible.

The current study describes a more efficient tool for MoA deconvolution by transcriptional profiling of compounds derived from phenotypic screens using a microfluidic platform with integrated fluidic circuits. We show that the approach is indeed feasible from both a conceptual and practical point of view. A major goal was to identify a minimal set of genes that can be used as biomarkers for MoA deconvolution on an amenable miniaturized assay. Analysis of the whole genome bacterial transcriptional responses to these compounds helped in selecting a set of biomarker genes important in the general bacterial response to antibiotics, which was utilized in the development of a routine assay. Unsupervised grouping by hierarchical clustering revealed gene clusters whose expression profile for these inhibitors was consistent with previous studies on the MoA of these agents. More importantly, the clustering grouped together inhibitors with a similar mode of action.

A common theme, in all compound expression fingerprints was the remarkable over representation of the genes in bacterial defense, detoxification adaptation and virulence systems. Interestingly, a majority of the functionally characterized biomarkers appeared to be only indirectly related to the primary molecular targets of most of the inhibitors used. These were largely identified owing to their consistent expression pattern when exposed to antibiotic stress. Some of the biomarkers included heat shock proteins, toxin anti-toxin regulons, and detoxification systems against reactive oxygen species. These findings are in line with an increasingly more acknowledged mechanism of anti-bacterial action where the accumulation of broadly toxic intermediates and a suicidal derailing of central homeostatic system result in bacterial death [4,7,8,27].

Hit to lead and lead optimization programs characteristically result in the generation of multiple compound series. Whilst it is important that cellular penetration, physiochemical profiles, and target inhibition are maintained or improved during SAR expansion, the gathering of this information is critical for good lead selection. Biomarker gene expression fingerprinting revealed that hit to lead and lead optimization could be guided in part by transcriptional profiling of the SAR series using the biomarker genes (Fig. 6). Derivatives of NCE1 and NCE2 showed similar fingerprints to the primary hit molecules, suggesting that they retained the same MoA. The tools described enable MoA diagnosis whilst prioritizing compounds from an SAR library of phenotypic screen derived leads. This will help in distinguishing the new series from other compound classes based on the transcriptional outcomes reflected in the gene expression by the chemical perturbations. Although the microfluidics, helped in tracking whether the novel compounds perturbed similar genes, it fails to give more details regarding the exact mode of action. Once the final compound has been identified, other techniques such as spontaneous mutant generation followed by whole genome sequencing and or affinity based target pull down studies need to be employed to identify the actual target.

The low number of biomarker genes required for MoA deconvolution and the establishment of a simple, robust and routine assay highlights the value of this technology in anti-infective drug discovery programs. As evidence to the potential application of this diagnostic platform, Barczak and co-workers recently underscored the application of drug RNA signatures for rapid clinical diagnosis of antibiotic susceptibilities in a broad range of bacterial pathogens [33]. The continued expansion of the drug transcriptome (collection of transcriptional fingerprints profiles generated by different chemical entities) using microfluidic
Supporting Information

Figure S1 A prediction correlation matrix using feature-reduced data to validate the classification of individual inhibitors. A heat map representation of gene expression similarity matrix of chemical inhibitors using selected features. The blue-red color scale shows the degree of similarity of drugs expression profiles ranging from -1 to 1 respectively. All expression values were transformed to log base 2. Periods following drug names represent duplicates.

(TIF)

Figure S2 A correlation matrix based on Pearson correlations of compounds from the same SAR library.

References

1. Monaghan RL, Barrett JF (2006) Antimicrobial drug discovery – Then, now and the genomics future. Biochemical Pharmacology 71: 901–909. doi:10.1016/j.bcp.2005.11.023.
2. Payne DJ, Holmes DJ, Rosenberg M (2001) Delivering novel targets and antibiotics from genomics. Curr Opin Investig Drugs 2: 1028–1034.
3. Chan PF, Macarren R, Payne DJ, Zalacain M, Holmes DJ (2002) Novel antibacterials: a genomics approach to drug discovery. curr drug targets infect disord 2: 291–308.
4. Payne DJ, Gwynn MN, Holmes DJ, Pompliano DL (2006) Drugs for bad bugs: confronting the challenges of antibacterial discovery. Nat Rev Drug Discov 6: 29–49. doi:10.1038/nrd2001.
5. Sams-Dodd F (2005) Target-based drug discovery: is something wrong? Drug Discov Today 10: 139–147. doi:10.1016/S1359-6464(03)03161-6.
6. Dick T, Young D (2011) How antibacterials really work: impact on drug discovery. Future Microbiology 6: 603–604. doi:10.2217/fmb.11.26.
7. Pethe K, Sequeira PC, Agarwalla S, Rhee K, Kuhen K, et al. (2010) A chemical genetic screen in Mycobacterium tuberculosis identifies carbon-source-dependent growth inhibitors devoid of in vivo efficacy. Nature Communications 1: 1–10. doi:10.1038/ncomms1060.
8. Butcher EC (2005) Can cell systems biology rescue drug discovery? Nat Rev Drug Discov 4: 362–363.
9. Keller TH, Shi PY, Wang QY (2011) Anti-infectives: Can cellular screening deliver? Current Opinion in Chemical Biology 15: 529–533.
10. Testuppen GC, Schlopen C, Raggia R, Gavraviga G (2007) Target deconvolution strategies in drug discovery. Nat Rev Drug Discov 6: 891–903. doi:10.1038/nrd2410.
11. Boshoff HM (2004) The Transcripational Responses of Mycobacterium tuberculosis to Inhibitors of Metabolism: NOVEL INSIGHTS INTO DRUG MECHANISMS OF ACTION. Journal of Biological Chemistry 279: 40174–40184. doi:10.1074/jbc.M406796200.
12. Hutter B, Schaub C, Albrecht S, Borgmann M, Brunner NA, et al. (2004) Prediction of mechanisms of action of antibacterial compounds by gene expression profiling. Antimicrobial Agents and Chemotherapy 48: 2538–2544. doi:10.1128/AAC.48.6.2538–2544.2004.
13. Parsons AB, Lopez A, Givoni IE, Williams DE, Gray CA, et al. (2006) Exploring the Mode-of-Action of Bioactive Compounds by Chemical-Genetic Profiling in Yeast. Cell 136: 611–623. doi:10.1016/j.cell.2006.06.049.
14. Plouffe D, Beinker A, McNamara C, Henson K, Kato N, et al. (2008) In silico activity profiling reveals the mechanism of action of antimicrobials discovered in a high-throughput screen. Proc Natl Acad Sci USA 105: 9039.
15. Frances S (2005) An array of problems. Nat Rev Drug Discov 4: 362–363. Available: http://www.nature.com/nrd/journal/v4/n5/abs/nrd1743.html.
16. Ioannidis JP (2005) Microarrays and molecular research: noise discovery? The Lancet 365: 454–455. doi:10.1016/S0140-6736(05)17678-7.
17. Michiels S, Koscielny S, Hill C (2005) Prediction of cancer outcome with microarrays: a multiple random validation strategy. The Lancet 365: 486–492. doi:10.1016/S0140-6736(05)17866-0.
18. Melin J, Qake SR (2007) Microfluidic large-scale integration: the evolution of design rules for biological automation. Annu Rev Biophys Biomol Struct 36: 213–231. doi:10.1146/annurev.biophys.36.040406.132646.
19. Spurgeon SL, Jones RC, Ramkrishnan R (2008) High throughput gene expression measurement with real time PCR in a microfluidic dynamic array. PLoS ONE 3: e1662. doi:10.1371/journal.pone.0001662.
20. Mangan JA, Sole KM, Minchison DA, Butcher PD (1997) An effective method of RNA extraction from bacteria refractory to disruption, including mycobacteria. Nucleic Acids Research 25: 675–676.
21. Stekel D (2003) Analysis of Relationships Between Genes, Tissues or Treatments. Microarray Bioinformatics. Cambridge University Press. 139–182.
22. R Development Core Team (2008) R: A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria. ISBN: 3-900051-09-6. URL: http://www.R-project.org.
23. Kurabachew M, Lu SJH, Krastel P, Schmitt EK, Suresh BL, et al. (2008) Lipoyamycin targets RNA polymerase and has good activity against multidrug-resistant strains of Mycobacterium tuberculosis. Journal of Antimicrobial Chemotherapy 62: 713–719. doi:10.1093/jac/dkn269.
24. Schmitt EK, Rivanto M, Sambhandamurthy V, Roggo S, Mauel C, et al. (2011) The Natural Product Cyclomarin Kills Mycobacterium Tuberculosis by Targeting the ClpC1 Subunit of the Caspase-like Protease. Angew Chem Int Ed 50: 3859–3861. doi:10.1002/anie.201101740.
25. Cole ST, Brosch R, Parkhill J, Garnier T, Churcher C, et al. (1998) Deciphering the biology of Mycobacterium tuberculosis from the complete genome sequence. Nature 393: 537–544. doi:10.1038/31159.
26. Arcus VL, Rainey PB, Turner SJ (2005) The PIN-domain toxin–antitoxin array in mycobacteria. Trends in Microbiology 13: 360–365.
27. Kohanski MA, Dwyer DJ, Hayete B, Lawrence CA, Collins JJ (2007) A common mechanism of cellular death induced by bacterial antibiotics. Cell 130: 797–810. doi:10.1016/j.cell.2007.06.049.
28. Rottmann M, McNamara C, Yeung BKS, Lee MCS, Zou B, et al. (2010) Spironolodones, a Potent Compound Class for the Treatment of Malaria. Science 329: 1173–1180. doi:10.1126/science.1152253.
29. Makarov V, Manina G, Mikouva K, Meimonn U, Ryaboova O, et al. (2009) Benzothiazinones kill Mycobacterium tuberculosis by blocking arabian synthesis. Science 324: 801–804. doi:10.1126/science.1171583.
30. AUDREY K (2005) A Diaryquinolin Drug Active on the ATP Synthase of Mycobacterium tuberculosis. Science 307: 221–227. doi:10.1126/science.1106753.
31. Bax BD, Chan PF, Eggleston DS, Fosbery A, Gentry DR, et al. (2010) Type IIa topoisoermerase inhibition by a new class of antibacterial agents. Nature 466: 935–940. doi:10.1038/nature09197.
32. Sanz LM, Jimenez-Diaz MB, Crespo R, De-Cozar C, Almela MJ, et al. (2011) Cyclopropyl Carboxamides, a Chemically Novel Class of Antimalarial Agents Identified in a Phenotypic Screen. Antimicrobial Agents and Chemotherapy 55: 5740–5745. doi:10.1128/AAC.05188-11.

33. Barczak AK, Gomez JE, Kaufmann BB, Hinson ER, Cosimi L, et al. (2012) RNA signatures allow rapid identification of pathogens and antibiotic susceptibilities. Proceedings of the National Academy of Sciences 109: 6217–6222. doi:10.1073/pnas.1119340109.