

# Network analysis links genome-wide phenotypic and transcriptional stress responses in a bacterial pathogen with a large pan-genome.

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## 29 ABSTRACT

30 **Background:** Bacteria modulate subcellular processes to handle stressful environments. Genome-wide  
 31 profiling of gene expression (RNA-Seq) and fitness (Tn-Seq) allows two views of the same genetic  
 32 network underlying these responses. However, it remains unclear how they combine, enabling a  
 33 bacterium to overcome a perturbation.

34 **Results:** Here we generate RNA-Seq and Tn-Seq profiles in three strains of *S. pneumoniae* in response  
 35 to stress defined by different levels of nutrient depletion. These profiles show that genes that change  
 36 their expression and/or become phenotypically important come from a diverse set of functional  
 37 categories, and genes that are phenotypically important tend to be highly expressed. Surprisingly, we  
 38 find that expression and fitness changes rarely occur on the same gene, which we confirmed by over 140  
 39 validation experiments. To rationalize these unexpected results we built the first genome-scale metabolic  
 40 model of *S. pneumoniae* showing that differential expression and phenotypic importance actually  
 41 correlate between nearest neighbors, although they are distinctly partitioned into small subnetworks.  
 42 Moreover, a meta-analysis of 234 *S. pneumoniae* gene expression studies reveals that essential genes and  
 43 phenotypically important subnetworks rarely change expression, indicating that they are shielded from  
 44 transcriptional fluctuations and that a clear distinction exists between transcriptional and phenotypic  
 45 response networks.

46 **Conclusions:** We present a genome-wide computational/experimental approach that contextualizes  
 47 changes that occur on transcriptomic and phenomic levels in response to stress. Importantly, this  
 48 highlights the need to connect disparate response networks, for instance in antibiotic target identification,  
 49 where preferred targets are phenotypically important genes that would be overlooked by transcriptomic  
 50 analyses alone.

## INTRODUCTION

How organisms handle and overcome stress in their environment is a central question in biology, with applications in fields ranging from bioengineering to drug(-target) discovery. With the advent of genome-wide approaches, it has become clear that even relatively simple perturbations, such as a change in extracellular pH or the presence of an antibiotic, require complex physiologic responses affecting multiple subcellular processes [1-7]. A complete cellular stress response can be separated into at least two organization levels. The *transcriptional response* describes the change in gene expression following stress, while the *phenotypic response* describes how the importance of each gene (fitness) changes due to stress. Although the transcriptional and phenotypic stress responses are inter-dependent, they are measured experimentally by two disparate technologies. Transcriptomic profiling, as measured by cDNA microarrays or RNA-Seq, has been particularly popular in deciphering the complex bacteria-environment interaction to identify genes that change in their transcript abundance upon various environmental perturbations, such as exposure to antibiotics [8,9], interaction with host niches [10], and disruption of iron homeostasis [11]. Although such profiles provide a detailed picture of the transcriptional landscape, it remains to be determined whether transcript abundance alone is predictive of the phenotypic importance of a gene [12,13]. Alternatively, genome-wide mutant fitness profiling approaches, such as transposon insertion sequencing (Tn-Seq), have been developed to directly link genotypes to phenotypes and thereby measure the phenotypic stress response on a genome-wide scale [14-18]. This means that Tn-Seq determines the phenotypic importance of each gene in the genome in a specific environment (referred to as “gene fitness” in this paper) by measuring the effect each gene knockout in the genome has on fitness. For example, the lower the fitness, the more important a gene is for maintaining survival under a specific (e.g. stressful) condition [2,14,18]. Although both transcriptomic and phenotypic fitness profiling interrogate the same gene network, little is known about how these two data types correlate, i.e. are differentially expressed genes also phenotypically important during stress? Indeed, it is generally assumed that differentially expressed genes also represent phenotypically important genes. However, this assumption has not been thoroughly investigated, raising the possibility that if transcriptional and phenotypic stress responses are not correlated, then measuring either one alone offers an incomplete and incorrect picture of the cellular response. Here we determine

whether a correlation between transcription and phenotype exists and whether these stress responses can be combined to obtain a complete physiologic response.

As our model system we employ the human pathogenic bacterium *Streptococcus pneumoniae*, a major respiratory pathogen and source of morbidity and mortality. *S. pneumoniae* colonizes the nasopharynx asymptotically, but by disseminating to other tissues it can trigger disease, including pneumonia, meningitis, sepsis, and otitis media, which results in ~1 million deaths annually among children <5 years of age and ~0.5 million among groups including the immunocompromised and the elderly (>65 years) [19-21]. One possible challenge in exploring the physiologic response in a species such as *S. pneumoniae* is the genomic variation among strains. The increasing availability of fully sequenced genomes for this and other species has demonstrated a distinction between the species' core genome (the set of genes shared by all strains) and its pan-genome (the species' global genetic repertoire) [22-25]. Because no gene or pathway functions in a vacuum, rather they are connected by complex genetic networks [26,27], the presence or absence of genes among different pneumococcal strains suggests that each strain's genomic network may be differently wired, which could potentially make phenotypes strain-dependent. Indeed, we have recently shown that strain-specific phenotypic stress responses in *S. pneumoniae*, for instance in response to antibiotics, may be common [1]. Thus, when working with a bacterium with a large pan-genome, an ideal approach should obtain a species-wide, generalizable view of how the bacterium overcomes a stressful environment.

In this study, we investigate whether differentially expressed genes are also phenotypically important during stress in *S. pneumoniae*. We generate an unbiased, high-quality, and extensively validated profile of the bacterial stress response by employing two genome-wide approaches, RNA-Seq and Tn-Seq, to measure both the transcriptional and phenotypic stress response for three pneumococcal strains under three different levels of nutrient depletion. Surprisingly, there is little correlation between differential expression and phenotypic importance across the entire genome. To contextualize the transcriptional and phenotypic profiles, we built and curated the first genome-scale metabolic model in the genus *Streptococcus*. By integrating all our data into this model we show that the disparate genes with expression and fitness changes are actually closely connected in the genomic network. However, phenotypically important and essential genes seem to be transcriptionally shielded from large

107    fluctuations in expression and therefore organizationally separated from transcriptionally plastic, but  
 108    phenotypically unimportant, genes. Importantly, we provide a detailed roadmap to develop similar  
 109    systems-level approaches in other microorganisms. Our approach facilitates profiling and reconciling  
 110    transcriptomic and mutant fitness datasets and enables mapping of an organism's full physiologic  
 111    response to an environmental disturbance. Moreover, this study provides a clear rationale that  
 112    emphasizes the importance of targeting phenotypically important genes rather than differentially  
 113    expressed genes in applications such as in drug target discovery.

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# RESULTS AND DISCUSSION

**Designing a robust nutrient depletion assay for *S. pneumoniae*.** To avoid bias in the bacterial response to nutrient depletion that might result from genomic variation in a particular strain, we selected three phylogenetically distant strains to represent *S. pneumoniae*: TIGR4 (T4), Taiwan-19F (19F) and D39 (**Additional File 1**). Both T4 and 19F can cause invasive pneumococcal disease (IPD): T4 is a serotype 4 strain that was originally isolated from a Norwegian patient with IPD [28,29]; 19F is a multi-drug resistant strain isolated from a patient with IPD in Taiwan [30,31] (**Table 1**). D39 is a historically important and commonly used serotype 2 strain that was originally isolated from a patient about 90 years ago [32] (**Table 1**). Concerning their genomic content, the three strains share 1647 genes, while T4 has 217, 19F has 140 and D39 has 93 strain-specific genes (**Additional Files 2 and 3**). To simulate systemic nutrient depletion, we designed three increasingly restrictive media to cultivate *S. pneumoniae*. The first, semi-defined minimal media (SDMM), is a relatively rich media we have used previously [1,2] that contains a single carbon source (glucose), yeast extract, casein hydrolysate (digested amino acids), salts, trace metals, and vitamins (**Table 2**). The second, a chemically defined media (CDM), is based on a previously described recipe [33], however the original composition did not allow each strain to grow robustly. By adjusting the recipe, mainly by taking SDMM as the basis and replacing the yeast extract and casein hydrolysate by an equimolar mixture of the 20 amino acids, comparable growth rates to SDMM are achieved. CDM is thus completely defined, is less nutrient rich than SDMM, but still contains several non-essential components. By iteratively removing components of CDM, we created a minimal CDM, or MCDM, that still enables each strain to grow. In contrast to SDMM and CDM, each component in MCDM is essential for growth in at least one of the three strains, and removing any component of MCDM triggers severe growth defects in at least one strain. Nutrient availability therefore decreases from SDMM to CDM, and further to MCDM (**Additional File 4**), which is illustrated by a decrease in the growth rate of *S. pneumoniae* (**Table 1, Additional File 5**). By using multiple strains and three media conditions, generalizable -- instead of strain and/or environment-specific -- profiles are assembled that map the phenotypic and transcriptomic stress responses.

**Genome-wide fitness and expression profiling reveal the phenotypic and transcriptional importance of cellular processes upon nutrient depletion.** We applied two high-throughput, genome-wide methods to determine on two different levels how *S. pneumoniae* deals with nutrient depletion stress. The first level is measured by Tn-Seq, which quantifies a gene's effect on the growth rate (fitness) resulting from a genetic disruption by transposon insertions [18]. The second level is determined by RNA-Seq, which measures gene expression by quantifying transcript abundance. Tn-Seq fitness data thus provides insight into the phenotypic importance of each gene, while RNA-Seq provides insight into which genes respond to stress on a transcriptional level.

Six transposon insertion libraries for each strain were constructed and grown in each media to generate Tn-Seq profiles and a comprehensive genotype-phenotype map for each strain/environment pair (**Figure 1a**). As expected, and as we have shown before [1,2,14], the majority of insertions do not produce a significant fitness change, and more fitness defects (fitness < 1) were observed than increases in fitness (fitness > 1) (**Additional File 6**). To complement the Tn-Seq fitness data, genome-wide gene expression was measured for each strain/media pair by RNA-Seq. Four replicates from mid-exponential phase cells were prepared for each strain/media pair and sequenced at high depth (3.5-16 million reads/sample) [34]. Each sample contained reads mapping to 88-96% of all annotated genes in the corresponding genome, and transcript abundance between genes varied by nearly  $10^5$  (**Figure 1b**, **Additional File 6**). Comparing the raw expression value (i.e. transcript abundance, not fold change) with fitness revealed that most genes with a fitness defect are highly expressed (**Figure 1c**), which is a pattern that holds across all strains and all three media. However, this pattern is of little predictive value, since the majority of highly expressed genes do not have a fitness defect or increase, and thus transcript abundance alone does not predict fitness.

**Validation experiments confirm that high-throughput profiles consist of high confidence data.** The high-throughput genome-wide approaches Tn-Seq and RNA-Seq provide comprehensive profiles of cell physiology at two different levels. However, as with any high-throughput experiment the resulting datasets require careful validation to assess their accuracy. Tn-Seq data were validated by constructing 31 individual gene deletion mutants across the three strains. The mutants were used in monoculture growth

assays and 1x1 competition assays in which the wild type strain is competed against the mutant to obtain fitness. In total 122 genotype-phenotype relationships were validated across the three strains and three media conditions, which to our knowledge is the largest validation set generated to date for different strains of a bacterial pathogen for which no ordered knockout arrays exist. This resulted in a strong correlation ( $R^2 = 0.82$ ), which is similar to correlations we achieved previously [1,2] and confirms high-confidence Tn-Seq fitness data (**Figure 2a; Additional File 7**). Importantly, these data also give detailed information on what type of stress is experienced by the bacterium. For example, Tn-Seq data shows that in rich media (SDMM)  $\Delta aroE$  (*SP1376*), which catalyzes the conversion of 3-dehydroshikimate to shikimate and is associated with an upstream reaction in the *de novo* synthesis pathway of aromatic amino acids, has a growth defect (SDMM:  $W_{aroE} = 0.75$ ), which we indeed validated (**Figure 2b**). However, a more severe defect in growth is measured when  $\Delta aroE$  is grown in CDM and thus under stress by limited availability of amino acids (CDM:  $W_{aroE} = 0.26$ ). Moreover, the gene becomes almost conditionally lethal in MCDM in the absence of aromatic amino acids (MCDM:  $W_{aroE} < 0.10$ ). These data not only show that the Tn-Seq data consist of high confidence fitness values, but it also pinpoints environment-dependent weaknesses in the genomic network, highlighting how different genes become important in an environment-dependent manner. Moreover, it shows what type of stress is experienced in the environment; the conditional importance of *aroE* suggests an increasing lack of aromatic amino acids in CDM and MCDM.

Additionally, the RNA-Seq data were validated by qPCR by measuring the expression of nine genes in the three media conditions (**Figure 2c**). The changes in expression measured by qPCR match the RNA-Seq differential expression data across all strains and media (**Additional File 8**), confirming that the generated genome-wide expression profiles represent real changes in transcription.

**Identifying genetic drivers of the nutrient stress response.** The validated high-throughput data were used to identify genes responsible for the phenotypic (Tn-Seq) and transcriptomic (RNA-Seq) stress responses. By comparing fitness across environments we determined which genes changed their fitness ( $\Delta$ fitness, or  $\Delta W$ ) as *S. pneumoniae* transitions from rich media (SDMM) to defined (CDM) or minimal (MCDM) media (**Figure 3a**). Thus genes whose importance increases upon nutrient depletion will have

a decreased fitness (*i.e.* a negative  $\Delta$ fitness) in the more restrictive media, while those whose importance does not change will have a  $\Delta$ fitness of 0. Those genes whose importance decreases will have an increased fitness (*i.e.* a positive  $\Delta$ fitness). Following a change from SDMM to either CDM or MCDM, each strain had an average of 12 genes increase in fitness and 29 genes decrease in fitness. For example, gene *SP1555* (dihydrodipicolinate reductase) is a key enzyme for lysine biosynthesis. A deletion of *SP1555* thus blocks *de novo* lysine synthesis and should hamper *S. pneumoniae*'s ability to overcome the depletion of extracellular lysine in CDM and MCDM. Indeed, Tn-Seq data shows that *SP1555* is not important in rich media (SDMM:  $W_{SP1555} = 0.95$ ), but shows decreasing fitness, and thus increasing importance, in the more stringent media (CDM:  $W_{SP1555} = 0.78$ ,  $\Delta W_{SP155} = -0.17$ ; MCDM:  $W_{SP155} = 0.07$ ,  $\Delta W_{SP155} = -0.71$ ).

Besides genes that change in fitness, genes that change in expression were identified by comparing transcript abundances between SDMM and either CDM or MCDM (**Figure 3b**). On average, the media shift caused 101 genes to significantly increase expression and 125 genes to decrease expression. Overall, 5.4 times more genes showed significant expression changes compared to significant fitness changes. Importantly, in both the Tn-Seq and RNA-Seq datasets, the significant changes were distributed across a variety of cellular subsystems, indicating that the nutrient depletion environments trigger stress that is experienced network-wide (**Figure 3c**). Among metabolic subsystems, amino acid pathways are especially well represented, with 17% of genes showing a fitness change and 23% of genes differentially expressed (**Additional File 6**).

**Genome-wide data visualization with a whole cell model reveals that expression profiles are poor predictors of phenotypic importance.** *S. pneumoniae* designates a large fraction of its genome to metabolism, and a number of metabolic enzymes have been linked to the bacterium's phenotypic stress response to nutrient and antibiotic perturbations [1,2]. Given the large number of metabolic genes with fitness or expression changes during nutrient depletion (**Figure 3c**), we focused on metabolic pathways to identify patterns in how *S. pneumoniae* handles and overcomes stress. To analyze metabolism in a systematic and comprehensive way, a genome-scale metabolic model of *S. pneumoniae* was assembled. A draft metabolic model was derived from the KBase system [35] (<http://kbase.us>) by collecting metabolic

227 reactions associated with the annotated genomes for T4, 19F, and D39. To account for non-enzymatic  
 228 reactions and reactions with misannotations, a gap-filling algorithm [36] added reactions to ensure growth  
 229 of the model on SDMM, CDM, and MCDM. Pathways in the model were manually curated by comparing  
 230 reactions and gene associations to KEGG [37] and BioCyc [38], with a particular emphasis on amino acid  
 231 and nucleotide metabolism (**Additional Files 9 and 10**). The final model, called iSP16, details the  
 232 interconversion of 866 metabolites by 928 reactions, catalyzed by 463 genes (43.9% of all ORFs in T4).  
 233 To our knowledge, iSP16 is the first curated, genome-scale metabolic model in the genus *Streptococcus*.

234 Using iSP16, we searched for patterns in the location of fitness and expression changes during  
 235 nutrient stress. Initially, we expected the Tn-Seq and RNA-Seq data to align, as increasing expression of a  
 236 metabolic enzyme can increase flux through an important pathway [39]. However, overlaying the datasets  
 237 onto the network shows that, when transitioning from SDMM to CDM, metabolic pathways either change  
 238 on a transcriptional level or their fitness (phenotypic importance) changes, but they almost never change  
 239 in the same location (**Figure 4a**). Moreover, upon transitioning to MCDM, the clusters of fitness or  
 240 expression changes expand but they almost never merge (**Figure 4b**). This means that, contrary to our  
 241 expectations (and popular belief), overlap between fitness and expression changes are incredibly rare  
 242 (**Figure 4**).

243 A clear example of the disconnect between fitness and expression changes is the shikimate pathway,  
 244 the biosynthetic route for the aromatic amino acids (**Figure 5a**). Beginning with the glycolytic and  
 245 pentose phosphate pathway intermediates phosphoenol-pyruvate (PEP) and erythrose-4-phosphate (E4P),  
 246 the shikimate pathway uses seven enzymatic reactions before branching in sub-pathways specific for  
 247 tryptophan (Trp), phenylalanine, and tyrosine. The branchpoint occurs immediately after gene *SP1374*,  
 248 with gene *SP1816* catalyzing the first reaction of the Trp-specific branch (**Figure 5a**). Compared to  
 249 SDMM, CDM contains reduced tryptophan, and MCDM contains no tryptophan (**Table 2, Additional**  
 250 **File 4**). We expected that the removal of Trp would cause increased expression of the shikimate pathway  
 251 to compensate for the absence of Trp and decreased fitness when any of the pathway's genes are  
 252 interrupted. Although this pathway shows both fitness and expression changes in our data, the fitness  
 253 changes are restricted to the genes above the branch into tryptophan synthesis (*SP1700* - *SP1374*), while  
 254 the expression changes are below the branch point (*SP1817* - *SP1812*) (**Figure 5a**). The sole biosynthetic

route to tryptophan thus contains disjoint sets of genes with fitness and expression changes, indicating that a single pathway can be split between the phenotypic and transcriptional stress responses.

Strikingly, the lack of correlation between fitness and expression changes is not limited to metabolic genes. When plotting genome-wide expression changes against fitness changes (**Figure 5b**), no clear relationship is observed between fitness and expression changes; almost all genes appear on either the horizontal or vertical axes of **Figure 5b**, indicating either a fitness change with no expression change, or vice versa. Therefore, even though fitness and expression changes occur across the genome, their disjointedness would go unnoticed if only Tn-Seq or RNA-Seq experiments were performed, suggesting that upon exposure to stress, expression profiles are not good predictors of what is actually phenotypically important in a bacterium.

**Cellular networks link changes in gene expression and fitness.** In the metabolic network, fitness and expression changes rarely overlap, but they often seem to be located near one another (**Figure 4**). Thus even though expression changes are not indicative of genes that are phenotypically important, expression changes suggest that phenotypically important genes are often very close neighbors. To test this hypothesis a nearest neighbor analysis was performed in which each gene with a fitness change is paired with the closest gene in the network with an expression change (**Figure 5c**). The same procedure was repeated, starting with the genes with expression changes and searching for nearby fitness changes. Importantly, the nearest neighbor transformation restores the expected relationship between increased expression and decreased fitness. Prior to considering the nearest neighbors, 33% of the genes in **Figure 5a** appeared in the upper left quadrant (decreased fitness and increased expression) (**Figure 5d**). After pairing with nearby genes, 59% of the genes moved into this quadrant, the largest change for any quadrant of the graph.

This raises at least two questions: 1.) How close, on average, are paired neighbors; and 2.) Is there a neighbor-to-neighbor relationship between the size of the fitness change and the change in expression? For instance, are genes with the largest fitness changes neighbors with genes with the largest expression changes? To answer these questions distances between genes in the metabolic network were quantified by counting the number of reactions between enzymes (**Figure 6a**). For instance, genes associated with the

283 same reaction have a distance of zero, while genes associated with reactions that share a common  
 284 metabolite have a distance of one. To quantify the magnitude of both the fitness and expression changes,  
 285 we multiply the two changes. This product corresponds to the area of a rectangle drawn between a point in  
 286 **Figure 5c** and the origin (**Figure 6b**). This “area off axes” is maximized when both the fitness and  
 287 expression changes are large, and it is near zero when either the fitness or expression change is small.  
 288 Finally, by plotting the fitness-expression product against the distance between the corresponding genes  
 289 (**Figure 6c**), two conclusions can be drawn. First, the majority of fitness and expression changes pair at  
 290 short distances, with 80% of all pairs within a distance of two, and 93% within a distance of three. Second,  
 291 the size of the fitness and expression changes in each pair decreases with distance ( $p < 0.005$ , ANOVA).  
 292 This analysis (**Figure 5**) thus shows quantitatively what is visually suggested in **Figure 4**; fitness and  
 293 expression changes occur in distinct, but co-localized genes. Moreover, the largest changes in fitness are  
 294 close to the largest expression changes, which means that fitness and expression changes are not only  
 295 co-located, but have a magnitude that matches their neighbors'.

296 Further visual inspection of the metabolic network maps (**Figure 4**) suggests that genes with fitness  
 297 changes and genes with expression changes are not distributed evenly throughout the network. Instead, it  
 298 appears that small clusters of fitness and expression changes are present, suggesting small sub-networks  
 299 that either change transcriptionally or that are conditionally important. To quantify the existence of these  
 300 clusters, we compared the distribution of distances among all genes, genes with fitness changes, and genes  
 301 with expression changes (**Figure 6d**). Genes with an expression change ( $\Delta\text{expression} \rightarrow \Delta\text{expression}$ ) are  
 302 indeed clustered at shorter distances to other differentially expressed genes, and the same is true for genes  
 303 that change in fitness ( $\Delta\text{fitness} \rightarrow \Delta\text{fitness}$ ) ( $p < 0.005$ ,  $t$ -test of Poisson fit to distributions of distances)  
 304 (**Figure 6d**). In contrast, essential genes, derived from these and previous Tn-Seq data [2,14] (**Additional**  
 305 **File 11**) are not closely clustered (**Figure 6e**, essential  $\rightarrow$  essential). Instead, essential genes are closer to  
 306 genes with fitness changes than they are to either genes with expression changes or other essential genes  
 307 (**Figure 6e**, essential  $\rightarrow \Delta\text{essential}$  and essential  $\rightarrow \Delta\text{fitness}$ ,  $p < 0.05$ ). This distribution shows that: 1.)  
 308 Phenotypic stress response genes lie closer to essential genes than transcriptional stress response clusters;  
 309 and 2.) Both transcriptional control and important functions are intertwined, but separated into small  
 310 clusters that correspond to either the phenotypic or transcriptional stress responses. Therefore, it appears

that genes that become phenotypically important in a new environment are shielded from highly fluctuating changes in expression, while nearby genes that are less important on a phenotypic level may fluctuate to a much larger extent, suggesting these different sets of genes are part of differently organized regulatory modules.

**Meta-analysis reveals partitioning of phenotypic and transcriptomic stress responses across multiple conditions.** If these distinct sets of genes are indeed organized in different regulatory modules, this would suggest that essential genes and the phenotypically important genes we identified would be universally shielded from expression changes across any condition. To test this hypothesis, we performed a meta-analysis using all publicly available *S. pneumoniae* datasets in the Gene Expression Omnibus (GEO) database. Using microarray data from 234 experiments (**Additional File 9**), we calculated the “expression plasticity” across a wide range of genetic and environmental perturbations (**Figure 7a**). The plasticity is the relative variance in expression of the gene (and its homologs) across all datasets in the GEO database. This means that genes with low plasticity rarely vary in their expression levels, while high plasticity genes vary widely in their expression across conditions. The GEO meta-analysis shows that both essential genes (which are phenotypically important in all environments) as well as genes with fitness changes in our dataset have low expression plasticity across the GEO datasets (**Figure 7b**). Furthermore, plotting the GEO expression plasticity against the size of the fitness change in our experiment reveals that genes with the largest fitness changes have the lowest plasticity (**Figure 7c**). And thus, not only do the phenotypically important genes have no corresponding expression change in our experiments, they also hold their expression relatively constant across all the experiments in GEO. This meta-analysis suggests that the partitioning of the phenotypic and transcriptional stress responses extends beyond the nutrient depletion stresses analyzed in this study and further suggests that the phenotypic and the transcriptional response networks are composed of highly dissimilar regulatory modules.

## CONCLUSIONS

A popular assumption is that differentially expressed genes are good proxies for genes that are important for maintaining the survival of a microbe under stressful conditions [8-11]. Our validated genome-wide profiles for three strains of *S. pneumoniae* show in detail that genes and pathways that become important upon nutrient depletion are generally not differentially expressed. It is possible that the products of the genes that change their fitness are characterized by post-transcriptional or post-translational, rather than transcriptional, changes [40,41]. For instance, enzymes involved in protein and carbohydrate biosynthesis are heavily modified by lysine succinylation in multi-drug resistant *Mycobacterium tuberculosis* strains [42], while S-glutathionylation serves as a major regulation mechanism when *Cyanobacteria* are under oxidative stress conditions [43]. However, in general these mechanisms only affect a relatively small number of genes, and overall there seems to be a strong correlation between transcription and protein levels, especially during mid-exponential growth [44]. We thus do not believe that our findings are coincidental and related to our organism or experimental setup, also because several other studies on bacterial and fungal species have implied similar patterns in certain metabolic pathways and cellular functions, even though not all of them were validated or conducted on a genome-wide level [40,45,46]. It seems that a poor correlation between transcriptional change and functional importance is, at least for bacteria, universal. This means that transcriptomic profiling studies should be assessed carefully when they are used as a predictor of genes that matter phenotypically, e.g. in genetic engineering or drug target identification studies. Drug target identification across different domains of life has heavily relied on transcriptomic data as a surrogate for functional importance [47-53]. For instance, inhibitors of streptokinase gene expression have been proposed as novel antimicrobials for group A streptococcus [52], and gene expression datasets were used as a source for co-target identification using a random-walk model based on *M. tuberculosis* [51]. Our results provide an important argument to search for phenotypically important genes instead of differentially expressed genes in future antimicrobial discovery -- blocking differentially expressed genes may fail to cause a growth defect while phenotypically important genes directly affect an organism's fitness.

In our attempts to unify fitness and expression changes, we showed that leveraging genome-scale metabolic modeling and topological analyses links both sets of genes in a cellular metabolic network.

Although changes in transcription status and changes in functional importance occur on separate sets of genes, our result show that these sets of genes are either in the same pathways or closely related pathways that share intermediates. This pattern is consistent with the ideas of metabolic control analysis (MCA), where pathway flux can be controlled by changes in either enzyme abundance or substrate concentrations [39,54]. A central result of MCA is that the relative importance of enzyme or substrate changes can vary along a pathway. In our example of the shikimate pathway (**Figure 5a**), the upper half of the pathway (SP1371-SP1377) may be controlled by substrate abundance, while flux through the amino acid-specific branches (e.g. SP1811-SP1818) may be controlled by enzyme abundance. It is important to note that all of the reversible reactions in the shikimate pathway occur before the branch point (**Figure 5a**), and the (reversible) enzymes above the branch point are not differentially expressed. Reversibility allows pathway intermediates to control flux through feedback mechanisms, possibly lessening the importance of transcriptional control in the upper branch. By deferring most of the transcriptional control until the lower branches of the shikimate pathway, *S. pneumoniae* may be able to control the production of Trp separately from the other aromatic amino acids while still maintaining adequate flux through the upper branch via substrate-level feedback.

A common explanation for the lack of an expected fitness defect is redundancy in the surrounding network. Since all of the fitness defects in the shikimate pathway occur before it branches into pathways for individual amino acids, it is possible that redundancies exist that can overcome the loss of a biosynthetic route for a single amino acid, but not for all three. Although there is no known alternative route for aromatic amino acid synthesis outside of the shikimate pathway, the Trp-specific genes could be redundant and thus some genes in the pathway could alleviate the absence of other genes and provide functional redundancy. Identifying these “hidden” redundancies is a powerful benefit of overlaying genome-scale phenotypic data onto mathematical models.

In addition to allowing more parsimonious gene regulatory networks, separating transcriptional control from phenotypic importance may allow bacteria more flexibility to respond to new environments without incurring a fitness cost. Genes that fluctuate transcriptionally are not important for sustaining growth and hence have more flexibility in their expression. Phenotypically important genes seem to be shielded from large, fluctuating expression changes and are possibly controlled by feedback loops, which

is a mechanism adept at retaining expression levels within tight boundaries [55]. Taken together, these features allow a bacterium to maintain a tightly controlled, robust core of essential genes while simultaneously preserving metabolic flexibility.

In this report, we present a transferable, systems-level approach to reconcile transcription and fitness changes within a network, which serves as an important attempt to achieve a systems-level understanding of how a bacterium deals with environmental perturbations. Although metabolism represents the majority of genes that change in either expression or fitness during nutrient depletion (**Figure 3c**), we are striving to achieve a truly holistic view by integrating additional parts of the genomic network, including energy generation, cell division, transport, and formation and turnover of the cell wall and membrane. Lastly, applying network topological analyses to contextualize high-throughput experiments has the potential to provide value in genetic engineering, predicting drug target candidates, and re-evaluating current drug targets with the goal to achieve a higher success rate in developing novel strategies to eradicate microbial pathogens.

## 407 **METHODS**

### 408 **Bacterial strains, growth and media**

409 Experiments were performed with *S. pneumoniae* strains TIGR4 (T4; NCBI Reference Sequence:  
 410 NC\_003028.3) and Taiwan-19F (19F; NC\_012469.1), and D39 (NC\_008533). All gene numbers are  
 411 according to the TIGR4 genome, except the unique genes, which are preceded by SP, SPT, and SPD for  
 412 TIGR4, Taiwan-19F, and D39, respectively. A "correspondence table" that matches homologous genes in  
 413 the three strains can be found in **Additional File 2**. Single gene knock-out strains were constructed by  
 414 replacing the coding regions with a chloramphenicol or spectinomycin resistance cassette as described  
 415 previously [2,14,56]. Except for Tn-Seq experiments, RNA-Seq experiments, and specific growth  
 416 conditions, *S. pneumoniae* was cultivated statically in Todd Hewitt broth with 5% yeast extract and 5  
 417  $\mu$ L/mL of Oxyrase (Oxyrase, Inc), or on sheep's blood agar plates at 37°C in a 5% CO<sub>2</sub> atmosphere.  
 418 When appropriate, liquid culture and blood agar plates contained 4  $\mu$ g/mL of chloramphenicol (Cm) or  
 419 200  $\mu$ g/mL of spectinomycin (Spec) for selecting strains or mutant libraries that contain drug markers.  
 420 Tn-Seq and RNA-Seq experiments were performed in three growth media that contain gradually  
 421 decreasing nutrient levels, namely, semi-defined minimal medium (SDMM) [2], chemically defined  
 422 medium (CDM; **Additional File 4**) and minimal chemically defined medium (MCDM; **Additional File**  
 423 **4**).

### 424 425 **Tn-Seq library construction and selection experiments**

426 Transposon insertion mutant libraries were constructed as previously described with the transposon  
 427 Magellan6, which lacks transcriptional terminators therefore allows for read-through transcription and  
 428 diminishing polar effects [2,14,57]. Additionally, the mini-transposon contains stop codons in all three  
 429 frames in either orientation when inserted into a coding sequence. Six independent transposon libraries  
 430 were constructed in T4, 19F, and D39. Tn-Seq experiments were performed with each transposon library  
 431 for each of the three strains under the three media conditions (pH 7.3 with 20mM of supplemental  
 432 glucose in SDMM and MCDM and 28 mM glucose in CDM). Mutant libraries were grown to mid- to  
 433 late-log phase and harvested for genomic DNA isolation.

434

## **Tn-Seq sample preparation, sequencing and fitness calculation**

Sample preparation, Illumina sequencing, and fitness calculations were performed as previously described [2,14,18,58]. For each insertion, fitness ( $W_i$ ) representing the growth rate is calculated by determining the change in frequency for the mutant in the population [58]. Fitness for single genes is calculated by averaging  $W_i$  over all the insertions in the same gene. To determine whether fitness effects significantly differ between conditions, three requirements must be fulfilled: 1.)  $W_i$  is calculated from at least three data points (insertions), 2.) the difference in fitness between conditions has to be larger than 15% ( $|W_i - W_j| > 0.15$ ), and 3.) the difference in fitness has to be significantly different in a one sample  $t$ -test with Bonferroni correction for multiple testing.

## **Competition and single strain growth assays**

1x1 competition experiments were performed by mixing a single gene knock-out strain with the corresponding wild type strain in a 1:1 ratio and growing for approximately eight generations to late exponential phase in a particular growth medium [2]. A sample was taken at the beginning and the end of a competition experiment for CFU counts and plated on blood agar plates (for a total CFU count) and on blood agar plates with selective antibiotics (for a CFU count of the knock-out strain). Fitness was then calculated using the same approach as Tn-Seq by determining the ratios of the competing strains at the start and end of the competition and determining the expansion of the population using CFU counts. Single-strain growth assays were performed in 96-well plates by taking OD<sub>600</sub> measurements on a Tecan Infinite 200 PRO plate reader. Both competition and single-strain growth assays were performed no fewer than three times.

## **RNA-Seq sample collection**

The three strains were grown under each of the three growth media conditions (SDMM, CDM, and MCDM) in four biological replicates. 5 mL of early mid-log phase liquid culture was harvested by centrifugation at 4°C, snap frozen, and stored at -80°C for RNA isolation. Total RNA was isolated using the RNeasy Mini kit (Qiagen).

## **RNA-Seq sample preparation, sequencing and expression level calculations**

RNA-Seq cDNA libraries were generated following the RNAtag-Seq protocol as previously described [59]. Briefly, 400ng of RNA was fragmented, depleted of genomic DNA using TURBO DNA-free kit (Ambion), 5'-dephosphorylated, and subsequently ligated to barcoded RNA adapters at the 3'-terminus. Barcoded RNA samples were pooled and purified with RiboZero (Illumina). The ribosomal RNA-depleted samples were converted to Illumina cDNA sequencing libraries in three key steps: 1.) First strand cDNA synthesis with AffinityScript Multiple Temperature cDNA Synthesis kit (Agilent) and RNA degradation, 2.) ligation to a 3'-linker, and 3.) PCR amplification using primers that target the 3'-linker and a constant region of the RNA barcodes and contain the Illumina flow cell sequences. The cDNA libraries were sequenced on an Illumina NextSeq500 platform (single read, 50 base pair).

Raw reads were demultiplexed, trimmed to 40 base pairs, and quality filtered using custom R scripts and the ShortRead package. Reads were mapped to the corresponding *S. pneumoniae* genome using Bowtie [60] with settings "-n 2 -l 60 -m 1 -B 1". Reads were aggregated to genes using the GenomicRanges R package and differential expression was calculated using DESeq2 [61].

## **qPCR expression analysis**

The three wildtype strains were grown in SDMM, CDM, and MCDM to early mid-log phase. Sample collection and total RNA isolation were performed following the same procedure as RNA-Seq. 4 ug of RNA from each sample was treated with the TURBO DNA-free kit, after which 400 ng of cleaned-up RNA was subjected to first strand cDNA synthesis using iScript reverse transcription Supermix (BioRad). Quantitative PCR was performed using a BioRad MyiQ; each sample was measured in two biological replicates and three technical replicates. No-reverse transcriptase and no-template controls were included for each sample. Expression levels from all samples were normalized against the 50S ribosomal gene SP2204.

## **Statistical analysis**

Statistical analyses were performed in R (<http://www.r-project.org>). Gene distributions were fit to either Poisson (gene distance distributions) or Gamma (GEO plasticity) distributions using the fitdistr function

491 in the MASS toolbox. Expected values of the distributions were compared by a *t*-test.

492

## 493 **DECLARATIONS**

494 **Ethics approval and consent to participate.** Not applicable.

495 **Consent for publication.** Not applicable.

496 **Availability of data and materials.** The datasets generated during the current study are available as  
497 Additional Files and in the Sequence Read Archive (SRP082544).

498 **Competing interests.** The authors declare that they have no competing interests.

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500 **Authors' contributions.** PAJ led the computational experiments. ZZ led the wet-lab experiments. PAJ  
501 and ZZ performed computational and wet-lab experiments. All authors analyzed data, interpreted results,  
502 and wrote and approved the final manuscript.

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504

## 505 **ADDITIONAL FILES**

506 **Additional File 1:** *S. pneumoniae* phylogenetic tree.

507 **Additional File 2:** Gene correspondence table.

508 **Additional File 3:** Breakdown of *S. pneumoniae* genes by strain and category.

509 **Additional File 4:** Media composition for SDMM, CDM, and MCDM.

510 **Additional File 5:** Growth curves for T4, 19F, and D39 in SDMM, CDM, and MCDM.

511 **Additional File 6:** Tn-Seq and RNA-seq data

512 **Additional File 7:** Tn-Seq validation

513 **Additional File 8:** RNA-seq validation

514 **Additional File 9:** Supplementary Methods: Model assembly and curation; minor metabolite listing;  
515 GEO datasets and analysis pipeline.

516 **Additional File 10:** iSP16 model in SBML format.

517 **Additional File 11:** Essential genes for T4, 19F, and D39.

518

# REFERENCES

1. van Opijnen T, Dedrick S, Bento J. Strain-specific antibiotic sensitivity in *S. pneumoniae*. PLoS Pathog. 2016. In Press.
2. van Opijnen T, Camilli A. A fine scale phenotype-genotype virulence map of a bacterial pathogen. Genome Research. 2012;22:2541–51.
3. Drlica K, Malik M, Kerns RJ, Zhao X. Quinolone-Mediated Bacterial Death. Antimicrob. Agent Chemother. 2008;52:385–92.
4. Tomasz A. The Mechanism of The Irreversible Antimicrobial Effects of Penicillins: How the Beta-Lactam Antibiotics Kill and Lyse Bacteria. Annu Rev Microbiol. 1979;:1–25.
5. Vakulenko SB, Mobashery S. Versatility of Aminoglycosides and Prospects for Their Future. Clin Microbiol Rev. 2003;16:430–50.
6. Floss HG, Yu T-W. Rifamycin Mode of Action, Resistance, and Biosynthesis. Chem. Rev. 2005;105:621–32.
7. Rajagopal M, Martin MJ, Santiago M, Lee W, Kos VN, Meredith T, et al. Multidrug Intrinsic Resistance Factors in Staphylococcus aureus Identified by Profiling Fitness within High-Diversity Transposon Libraries. mBio. 2016;7:e00950–16–11.
8. Chatterjee A, Saranath D, Bhattar P, Mistry N. Global Transcriptional Profiling of Longitudinal Clinical Isolates of Mycobacterium tuberculosis Exhibiting Rapid Accumulation of Drug Resistance. Dheda K, editor. PLoS ONE. 2013;8:e54717–8.
9. Nielsen PK, Andersen AZ, Mols M, van der Veen S, Abee T, Kallipolitis BH. Genome-wide transcriptional profiling of the cell envelope stress response and the role of LisRK and CesRK in Listeria monocytogenes. Microbiology. 2012;158:963–74.
10. Bao H, Kommadath A, Liang G, Sun X, Arantes AS, Tuggle CK, et al. Genome-wide whole blood microRNAome and transcriptome analyses reveal miRNA-mRNA regulated host response to foodborne pathogen Salmonella infection in swine. Sci Rep. 2015;:1–12.
11. Butcher J, Stintzi A. The Transcriptional Landscape of Campylobacter jejuni under Iron Replete and Iron Limited Growth Conditions. Zhang Q, editor. PLoS ONE. 2013;8:e79475–16.

- 546 12. Guimaraes JC, Rocha M, Arkin AP. Transcript level and sequence determinants of protein  
547 abundance and noise in Escherichia coli. *Nucleic Acids Research*. 2014;42:4791–9.
- 548 13. Feder ME, Walser JC. The biological limitations of transcriptomics in elucidating stress and  
549 stress responses. *J Evolution Biol*. 2005;18:901–10.
- 550 14. van Opijnen T, Bodi KL, Camilli A. Tn-seq: high-throughput parallel sequencing for fitness and  
551 genetic interaction studies in microorganisms. *Nat Meth*; 2009;;1–9.
- 552 15. Langridge GC, Phan MD, Turner DJ, Perkins TT, Parts L, Haase J, et al. Simultaneous assay of  
553 every Salmonella Typhi gene using one million transposon mutants. *Genome Research*.  
554 2009;19:2308–16.
- 555 16. Gawronski JD, Wong SM, Giannoukos G, Ward DV, Akerley BJ. Tracking insertion mutants  
556 within libraries by deep sequencing and a genome-wide screen for Haemophilus genes required in  
557 the lung. *PNAS*. 2009;1–6.
- 558 17. Goodman AL, McNulty NP, Zhao Y, Leip D, Mitra RD, Lozupone CA, et al. Identifying Genetic  
559 Determinants Needed to Establish a Human Gut Symbiont in Its Habitat. *Cell Host & Microbe*.  
560 2009;6:279–89.
- 561 18. van Opijnen T, Camilli A. Transposon insertion sequencing: a new tool for systems-level  
562 analysis of microorganisms. *Nat Rev Microbiol*. 2013;11:435–42.
- 563 19. Tuomanen EI, Mitchell TJ, Morrison BG. *The Pneumococcus*. Washington: ASM Press.
- 564 20. CDC. Antibiotic Resistance Threats in the United States. 2013;1–114.
- 565 21. WHO. Pneumococcal conjugate vaccine for childhood immunization--WHO position paper.  
566 Relevé épidémiologique hebdomadaire Section d'hygiène du Secrétariat de la Société des Nations  
567 Weekly epidemiological record Health Section of the Secretariat of the League of Nations.  
568 2007;93–104.
- 569 22. Henriques-Normark B, Tuomanen EI. *The Pneumococcus: Epidemiology, Microbiology, and*  
570 *Pathogenesis*. Cold Spring Harbor Perspectives in Medicine. 2013;3:a010215–5.

- 571 23. Medini D, Serruto D, Parkhill J, Relman DA, Donati C, Moxon R, et al. Microbiology in the  
572 post-genomic era. *Nat Rev Microbiol.* 2008;1–12.
- 573 24. Medini D, Donati C, Tettelin H, Masignani V, Rappuoli R. The microbial pan-genome. *Curr Opin*  
574 *Genet Dev.* 2005;15:589–94.
- 575 25. Tenaillon O, Skurnik D, Picard B, Denamur E. The population genetics of commensal  
576 *Escherichia coli*. *Nat Rev Microbiol.* 2010;8:207–17.
- 577 26. Boone C, Bussey H, Andrews BJ. Exploring genetic interactions and networks with yeast. *Nat.*  
578 *Rev. Genet.* 2007;8:437–49.
- 579 27. Dixon SJ, Costanzo M, Baryshnikova A, Andrews B, Boone C. Systematic mapping of genetic  
580 interaction networks. *Annu. Rev. Genet.* 2009;43:601–25.
- 581 28. Aaberge IS, Eng J, Lermak G, Lovik M. Virulence of *Streptococcus pneumoniae* in mice: a  
582 standardized method for preparation and frozen storage of the experimental bacterial inoculum.  
583 *Microbial Pathogenesis.* 1995;1–12.
- 584 29. Tettelin H, Ciammola A, Rigamonti D, Leavitt BR, Goffredo D, Conti L, et al. Complete Genome  
585 Sequence of a Virulent Isolate of *Streptococcus pneumoniae*. *Science.* 2001;293:493–8.
- 586 30. McGee L, McDougal L, Zhou J, Spratt BG, Tenover FC, George R, et al. Nomenclature of major  
587 antimicrobial-resistant clones of *Streptococcus pneumoniae* defined by the pneumococcal  
588 molecular epidemiology network. *Journal of Clinical Microbiology.* 2001;39:2565–71.
- 589 31. Shi Z-Y, Enright MC, Wilkinson P, Griffiths D, Spratt BG. Identification of Three Major Clones of  
590 Multiply Antibiotic- Resistant. *Journal of Clinical Microbiology.* 1998;:1–6.
- 591 32. Lanie JA, Ng WL, Kazmierczak KM, Andrzejewski TM, Davidsen TM, Wayne KJ, et al. Genome  
592 Sequence of Avery's Virulent Serotype 2 Strain D39 of *Streptococcus pneumoniae* and Comparison  
593 with That of Unencapsulated Laboratory Strain R6. *J. Bacteriol.* 2006;189:38–51.
- 594 33. Härtel T, Eylert E, Schulz C, Petruschka L, Gierok P, Grubmüller S, et al. Characterization of  
595 central carbon metabolism of *Streptococcus pneumoniae* by isotopologue profiling. *J. Biol. Chem.*  
596 2012;287:4260–74.

597 34. Haas BJ, Chin M, Nusbaum C, Birren BW, Livny J. How deep is deep enough for RNA-Seq  
598 profiling of bacterial transcriptomes? BMC Genomics. 2012;1–11.

599 35. Department of Energy Systems Biology Knowledgebase (KBase), <http://kbase.us>.

600 36. Satish Kumar V, Dasika MS, Maranas CD. Optimization based automated curation of metabolic  
601 reconstructions. BMC Bioinformatics. 2007;8:212–6.

602 37. Kanehisa M, Goto S, Sato Y, Furumichi M, Tanabe M. KEGG for integration and interpretation of  
603 large-scale molecular data sets. Nucleic Acids Research. 2011;40:D109–14.

604 38. Karp P, Billington R, Holland T, Kothari A, Krummenacker M, Weaver D, et al. Computational  
605 Metabolomics Operations at BioCyc.org. Metabolites. 2015;5:291–310.

606 39. Fell D. Understanding The Control of Metabolism. London: Portland Press; 1997.

607 40. Giaever G, Chu AM, Ni L, Connelly C, Riles L, Veronneau S, et al. Functional profiling of the  
608 *Saccharomyces cerevisiae* genome. Nature. 2002;1–5.

609 41. Grangeasse C, StÅ¼lke JR, Mijakovic I. Regulatory potential of post-translational modifications  
610 in bacteria. Front. Microbiol. 2015;6:1–2.

611 42. Xie L, Liu W, Li Q, Chen S, Xu M, Huang Q, et al. First Succinyl-Proteome Profiling of Extensively  
612 Drug-Resistant Mycobacterium tuberculosis Revealed Involvement of Succinylation in Cellular  
613 Physiology. J. Proteome Res. 2015;14:107–19.

614 43. Chardonnet S, Sakr S, Cassier-Chauvat C, Le Maréchal P, Chauvat F, Lemaire SD, et al. First  
615 Proteomic Study of S-Glutathionylation in Cyanobacteria. J. Proteome Res. 2015;14:59–71.

616 44. Taylor RC, Webb Robertson B-JM, Markillie LM, Serres MH, Linggi BE, Aldrich JT, et al. Changes  
617 in translational efficiency is a dominant regulatory mechanism in the environmental response of  
618 bacteria. Integr. Biol. 2013;5:1393–14.

619 45. Price MN, Deutschbauer AM, Skerker JM, Wetmore KM, Ruths T, Mar JS, et al. Indirect and  
620 suboptimal control of gene expression is widespread in bacteria. Mol Syst Biol. Nature Publishing  
621 Group; 2013;9:1–18.

622 46. Turner KH, Everett J, Trivedi U, Rumbaugh KP, Whiteley M. Requirements for *Pseudomonas*  
623 *aeruginosa* Acute Burn and Chronic Surgical Wound Infection. Garsin DA, editor. PLoS Genet.  
624 2014;10:e1004518–12.

625 47. Patil SD, Sharma R, Srivastava S, Navani NK, Pathania R. Downregulation of *yidC* in *Escherichia*  
626 *coli* by Antisense RNA Expression Results in Sensitization to Antibacterial Essential Oils Eugenol  
627 and Carvacrol. van Veen HW, editor. PLoS ONE. 2013;8:e57370–9.

628 48. Fadhal E, Mwambene E, Gamiedien J. Modelling human protein interaction networks as metric  
629 spaces has potential in disease research and drug target discovery. BMC Syst Biol. 2014;:1–12.

630 49. Mizuarai S, Yamanaka K, Itadani H, Arai T, Nishibata T, Hirai H, et al. Discovery of gene  
631 expression-based pharmacodynamic biomarker for a p53 context-specific anti-tumor drug Wee1  
632 inhibitor. Mol Cancer. 2009;8:34–12.

633 50. Liu M, Healy MD, Dougherty BA, Esposito KM, Maurice TC, Mazzucco CE, et al. Conserved  
634 Fungal Genes as Potential Targets for Broad-Spectrum Antifungal Drug Discovery. Eukaryotic Cell.  
635 2006;5:638–49.

636 51. Chen L-C, Yeh H-Y, Yeh C-Y, Arias CR, Soo V-W. Identifying co-targets to fight drug resistance  
637 based on a random walk model. BMC Syst Biol. 2012;:1–14.

638 52. Sun H, Xu Y, Sitkiewicz I, Ma Y, Wang X, Yestrepesky BD, et al. Inhibitor of streptokinase gene  
639 expression improves survival after group A streptococcus infection in mice. PNAS. 2012;:1–6.

640 53. Mukhopadhyay S, Nair S, Ghosh S. Pathogenesis in tuberculosis: transcriptomic approaches to  
641 unraveling virulence mechanisms and finding new drug targets. FEMS Microbiol Rev.  
642 2012;36:463–85.

643 54. Heinrich R, Schuster D. The Regulation of Cellular Systems. Boston: Springer; 1996.

644 55. Shah NA, Sarkar CA. Robust Network Topologies for Generating Switch-Like Cellular  
645 Responses. PLoS Comp Biol. 2011;1–12.

646 56. Iyer R, Baliga NS, Camilli A. Catabolite Control Protein A (CcpA) Contributes to Virulence and  
647 Regulation of Sugar Metabolism in *Streptococcus pneumoniae*. J. Bacteriol. 2005;187:8340–9.

648 57. van Opijnen T, Lazinski DW, Camilli A. Genome-Wide Fitness and Genetic Interactions  
649 Determined by Tn-seq, a High-Throughput Massively Parallel Sequencing Method for  
650 Microorganisms. *Curr Protoc Mol Biol*. 2015;36:1E.3.1–1E.3.24.

651 58. van Opijnen T, Boerlijst MC, Berkhout B. Effects of Random Mutations in the Human  
652 Immunodeficiency Virus Type 1 Transcriptional Promoter on Viral Fitness in Different Host Cell  
653 Environments. *J.Virol*. 2006;80:6678–85.

654 59. Shishkin AA, Giannoukos G, Kucukural A, Ciulla D, Busby M, Surka C, et al. Simultaneous  
655 generation of many RNA-seq libraries in a single reaction. *Nat Meth*. 2015;12:323–5.

656 60. Langmead, B., Trapnell, C., Pop, M., & Salzberg, S. L. (2009). Ultrafast and memory-efficient  
657 alignment of short DNA sequences to the human genome. *Genome Biology*, 10(3), R25.  
658 <http://doi.org/10.1186/gb-2009-10-3-r25>

659 61. Love MI, Huber W, Anders S. Moderated estimation of fold change and dispersion for RNA-seq  
660 data with DESeq2. *Genome Biology*. 2014;15:31–21.

661

662

**Table 1** Summary of the three *S. pneumoniae* strains in this study

| Strains             | TIGR4                                                                        | Taiwan-19F                                             | D39                        |
|---------------------|------------------------------------------------------------------------------|--------------------------------------------------------|----------------------------|
| RefSeq              | NC_003028                                                                    | NC_012469                                              | NC_008533                  |
| Genome size (bp)    | 2,160,842                                                                    | 2,112,148                                              | 2,046,115                  |
| Number of genes     | 2287                                                                         | 2119                                                   | 2165                       |
| Origin              | Isolated from the blood of a 30-year-old male patient in Kongsvinger, Norway | A patient with invasive pneumococcal disease in Taiwan | Clinical isolate, 1916     |
| Reference           | Aaberge <i>et al.</i> (1995)<br>Tettelin <i>et al.</i> (2001)                | Shi <i>et al.</i> (1995)<br>McGee <i>et al.</i> (2001) | Lanie <i>et al.</i> (2006) |
| Doubling time (min) |                                                                              |                                                        |                            |
| SDMM                | 51 ± 5                                                                       | 74 ± 2                                                 | 57 ± 4                     |
| CDM                 | 80 ± 7                                                                       | 110 ± 10                                               | 74 ± 4                     |
| MMCDM               | 108 ± 9                                                                      | 118 ± 13                                               | 100 ± 8                    |

666 **Table 2.** Comparison of nutrient availability in SDMM, CDM and MCDM.

| Components         | SDMM | CDM | MCDM |
|--------------------|------|-----|------|
| Casein Hydrolysate | X    |     |      |
| Yeast Extract      | X    |     |      |
| L-Ala              |      | X   |      |
| L-Arg              |      | X   | X    |
| L-Asn              | X    | X   |      |
| L-Asp              |      | X   |      |
| L-Cys              | X    | X   | X    |
| L-Gln              | X    | X   | X    |
| L-Glu              |      | X   |      |
| Gly                |      | X   | X    |
| L-His              |      | X   | X    |
| L-Ile              |      | X   | X    |
| L-Leu              |      | X   | X    |
| L-Lys              |      | X   |      |
| L-Met              |      | X   | X    |
| L-Phe              |      | X   |      |
| L-Pro              |      | X   | X    |
| L-Ser              |      | X   | X    |
| L-Thr              |      | X   | X    |
| L-Trp              | X    | X   |      |
| L-Tyr              |      | X   |      |
| L-Val              |      | X   | X    |
| Adenine            | X    | X   | X    |
| Uracil             | X    | X   | X*   |
| Ca-Pantothenate    | X    | X   | X    |
| Nicotinic Acid     | X    | X   | X    |
| Pyridoxine.HCL     | X    | X   |      |
| Thiamine.HCL       | X    | X   |      |
| Riboflavine        | X    | X   |      |
| Biotin             | X    | X   |      |

|                                      |   |   |   |
|--------------------------------------|---|---|---|
| K <sub>2</sub> HPO <sub>4</sub>      | X | X | X |
| NaOAc                                | X | X | X |
| NaHCO <sub>3</sub>                   | X | X | X |
| MgCl <sub>2</sub> ·6H <sub>2</sub> O | X | X | X |
| CaCl <sub>2</sub>                    | X | X | X |
| CuSO <sub>4</sub> ·5H <sub>2</sub> O | X | X | X |
| ZnSO <sub>4</sub> ·7H <sub>2</sub> O | X | X | X |
| MnSO <sub>4</sub> ·4H <sub>2</sub> O | X | X | X |
| D-Glucose                            | X | X | X |

667 X: present in a medium.

668 \* In MCDM uracil was supplied at half the concentration compared to SDMM (**Additional File 4**).

669

670

## FIGURE LEGENDS

**Figure 1:** High-resolution profiles of phenotype and gene expression during stress incurred by nutrient depletion. **A.** Fitness values from Tn-Seq. **B.** Transcript abundance from RNA-Seq. Both high-throughput methods were performed on three *S. pneumoniae* strains (19F, T4, and D39) and in three media conditions (SDMM, CDM, and MCDM). **C.** By plotting Tn-Seq and RNA-Seq data on the same graph it becomes clear that genes with fitness defects ( $W < 0.85$ ) are highly expressed.

**Figure 2:** Tn-Seq and RNA-Seq data correlate strongly with validation experiments. **A.** A strong correlation between Tn-Seq fitness and fitness calculated from individual mutant growth curves or 1x1 competitions ( $W_{validation}$ ;  $n = 122$ ; shown are mean  $\pm$  SEM; linear fit yields  $R^2 > 0.82$ ) emphasizes that the strain-dependent sensitivity profiles are composed of high-confidence Tn-Seq data. **B.** Representative growth curves for comparisons between wild type (WT) and mutant  $\Delta SP1376$  in three different conditions, showing the dependence of the mutant on the presence of aromatic amino acids in the environment. **C.** Fold change in transcript abundance measured by qPCR and RNA-Seq. Vertical bars indicate SEM for both experiments. No qPCR and RNA-Seq pairs differed significantly ( $p < 0.0042$ ,  $t$ -test with Bonferroni correction) indicating RNA-Seq data are composed of high-confidence data as well.

**Figure 3:** Changes in gene fitness and expression occur across all strains, media, and cellular subsystems. **A.**  $\Delta fitness$  ( $\Delta W$ ) depicts how genes change their fitness as a strain transitions from rich media (SDMM) to defined (CDM) or minimal (MCDM) media. **B.** Shown are how genes change their expression as a strain transitions from rich media (SDMM) to defined (CDM) or minimal (MCDM) media. Expression is  $\log_2$  fold change in transcript abundance from RNA-Seq. In both figures statistically significant changes are colored and both assays were performed on three *S. pneumoniae* strains (T4, 19F, and D39) and two media transitions (SDMM→CDM, SDMM→MCDM). **C.** Percentage of genes in each category with significant changes in fitness (red) and expression (green). For total number of genes in each category (by strain), see **Additional File 3**.

699

700 Figure 4: Genes with significant changes in expression (green), fitness (red), or both (blue) are  
 701 distributed throughout the iSP16 metabolic model. Lines indicate reactions connecting metabolites  
 702 (circles). Minor and currency metabolites are not shown (see **Additional File 9**). Reactions are colored  
 703 based on gene associations in the iSP16 model.

704

705 Figure 5: Fitness and expression changes do not occur on the same gene but appear to be related. **A.**  
 706 Changes in gene expression and fitness are separated in the tryptophan biosynthesis branch of the  
 707 shikimate pathway. Single and double-headed arrows indicate reversibility or non-reversibility of  
 708 individual chemical reactions, respectively, while arrows spanning two genes indicate enzymatic  
 709 subunits catalyzing the same reaction. The dashed blue line (between SP1374 and SP1816) indicates the  
 710 branch point into tryptophan biosynthesis. PEP=phosphoenol-pyruvate, E4P=D-erythrose-4-phosphate,  
 711 Trp=L-tryptophan. **B.** Changes in gene expression (RNA-Seq:  $\Delta expression$ ) vs. changes in fitness  
 712 (Tn-Seq:  $\Delta fitness$  ( $\Delta W$ )) are not correlated. For each data point (gene)  $\Delta fitness$  ( $\Delta W$ ) and  $\Delta expression$  are  
 713 plotted depicting how a gene's fitness and expression change as a strain transitions from rich media  
 714 (SDMM) to defined (CDM) or minimal (MCDM) media. **C.** Genes with significant  $\Delta expression$  or  
 715  $\Delta fitness$  ( $\Delta W$ ) (black) migrate off the horizontal and vertical axes when paired with the nearest neighbor  
 716 that changes in fitness or expression (red) (NB. If multiple genes appear at the same distance from the  
 717 gene with a fitness change, the expression changes are averaged.). **D.** The number of genes in each  
 718 quadrant of the  $\Delta expression/\Delta fitness$  plot shifts before (black) and after (red) nearest-neighbor pairing,  
 719 showing that fitness and expression changes are somehow linked.

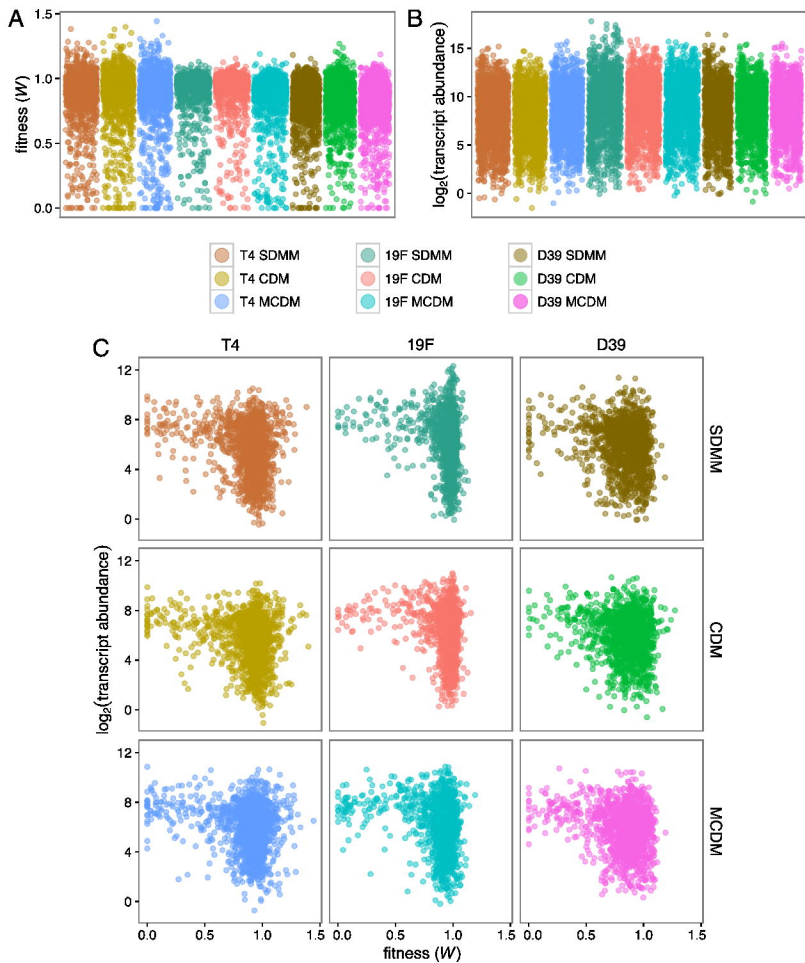
720

721 Figure 6: Metabolic models can quantify distances between reactions and genes. **A.** Shown is how  
 722 distances are calculated. For instance, the distance between the subunits *pgmA* and *pgmB* in glycolysis is  
 723 zero, while either enzyme has a distance one to *eno* and a distance two to *pyk*. **B.** The area off axes is the  
 724 product of  $\Delta expression$  and  $\Delta fitness$  ( $\Delta W$ ) and is maximized when both the fitness and expression  
 725 changes are large. The area off axes is near zero when either the fitness or expression change is small. **C.**  
 726 Extent of off-axis migration after nearest-neighbor averaging decays with distance between the gene and

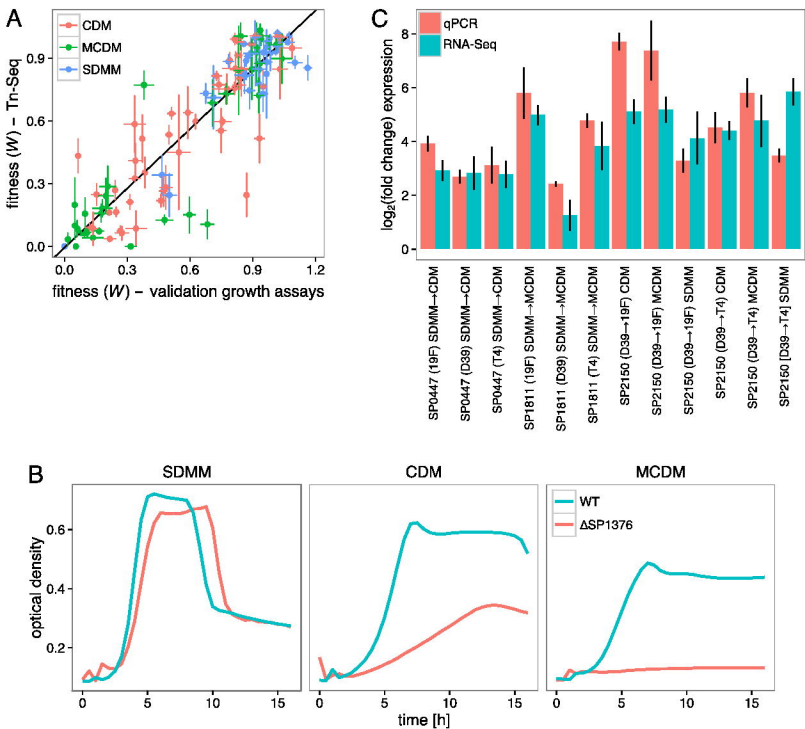
its nearest neighbor, indicating that changes are closely located within the network and that the largest changes in fitness are close to the largest expression changes. **D.** Distribution of network distances varies between genes with fitness and expression changes. Distances between all genes (*any*), genes with significant changes in expression ( $\Delta expression$ ) or fitness ( $\Delta fitness$ ). Unconnected genes (distance =  $\infty$ ) are not shown. The short distances between two genes with expression changes or two genes with fitness changes indicates small subnetworks of either fitness or expression changes. **E.** Essential genes and genes with fitness or expression changes are not evenly distributed. On average, essential genes (*essential*) are closest to genes with fitness changes ( $\Delta fitness$ ) and farthest from genes with expression changes ( $\Delta expression$ ).

**Figure 7:** Meta-analysis shows that essential and phenotypically important genes are shielded from large changes in expression. **A.** Quantifying gene expression plasticity using meta-analysis of GEO expression studies. Plasticity is the normalized variance in gene expression across all *S. pneumoniae* data in GEO (see **Additional File 9**). **B.** Gene expression plasticity is significantly lower for essential genes ( $p < 10^{-14}$ ) and conditionally essential genes (genes with a significant fitness change) ( $p < 10^{-34}$ , both comparisons *t*-test on means of fitted Gamma distributions). Both the essential and conditionally essential genes appear to be shielded from transcriptional changes not only in our RNA-Seq data, but across all the expression datasets in GEO. **C.** Gene expression plasticity decreases with increasing magnitude of the fitness change (relative to SDMM). Thus the amount of shielding is proportional to a gene's phenotypic importance, and genes with the largest fitness changes show the smallest variation in expression across the experiments in GEO.

Figure 1



**Figure 2**



**Figure 3**

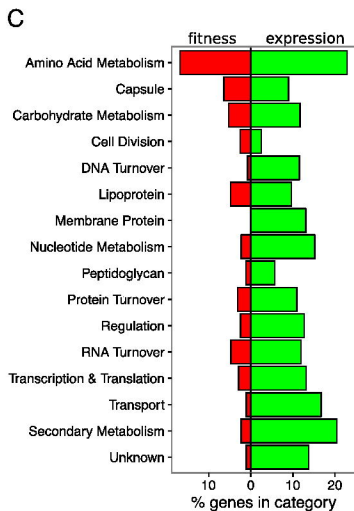
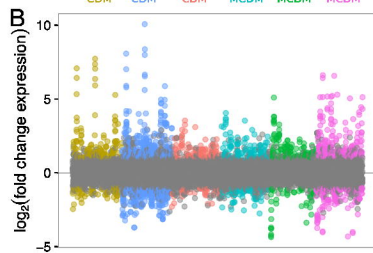
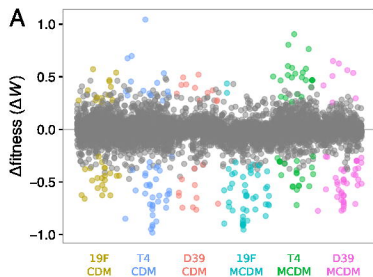


Figure 4

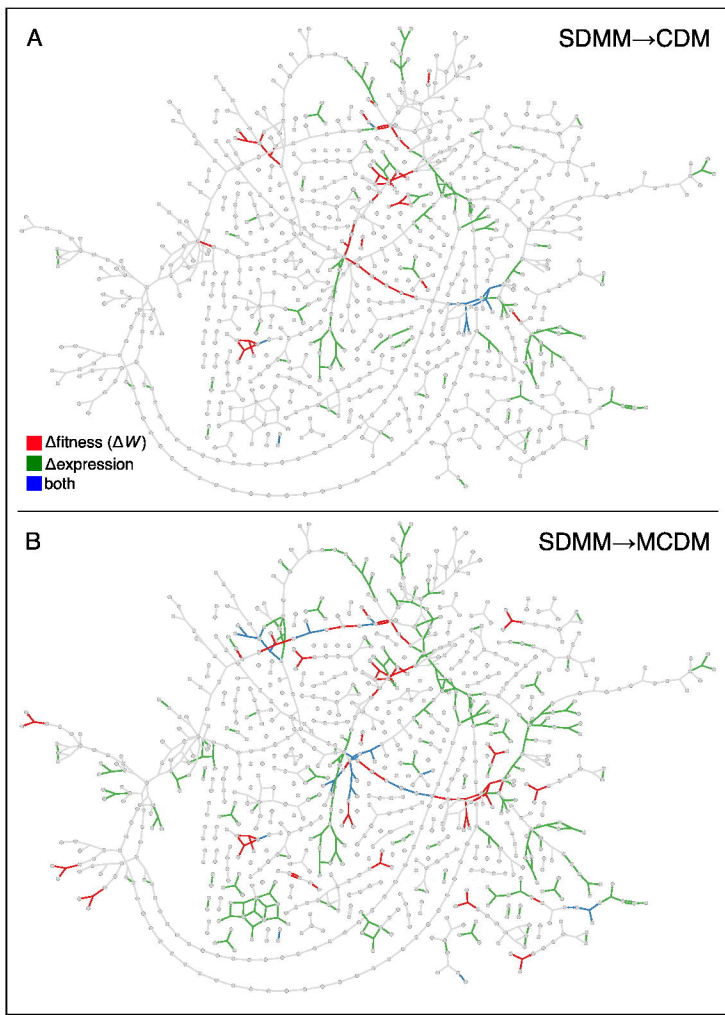


Figure 5

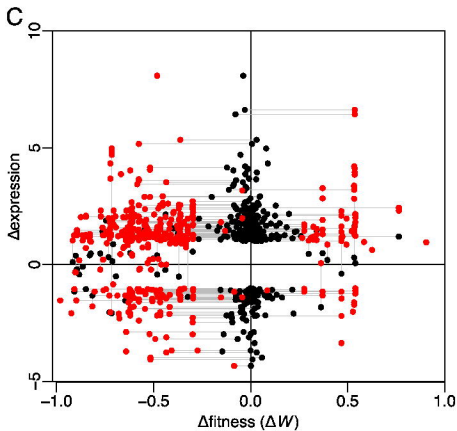
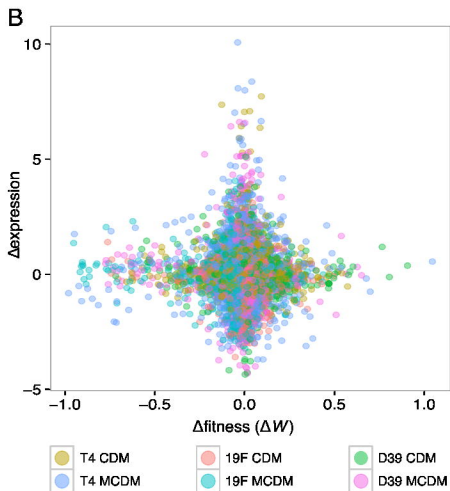
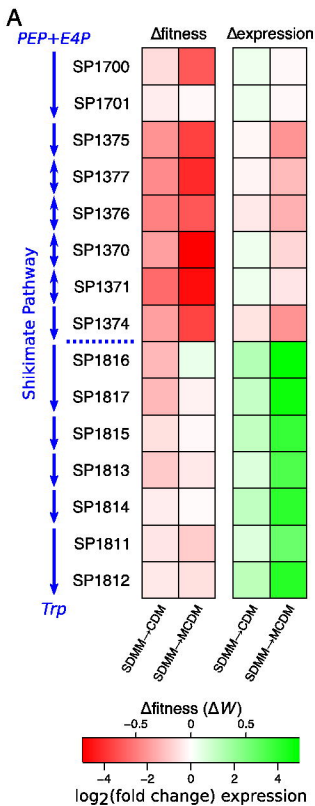


Figure 6

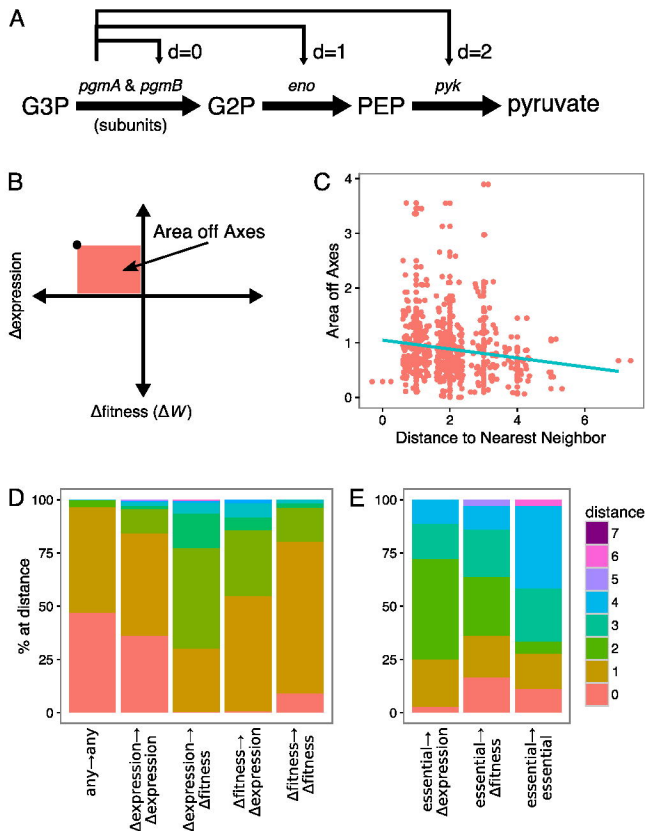
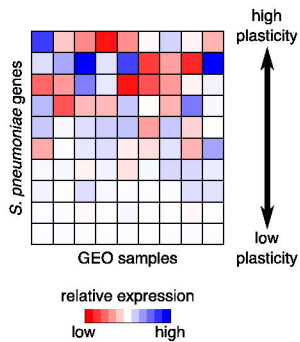
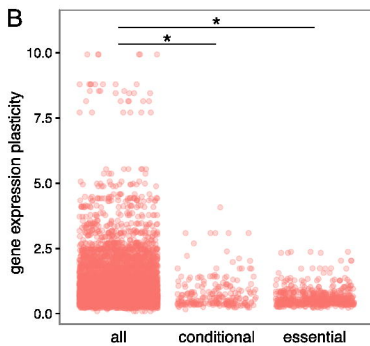


Figure 7

A



B



C

