A metabolic network approach for the identification and prioritization of antimicrobial drug targets

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**Abstract**

For many infectious diseases, novel treatment options are needed in order to address problems with cost, toxicity and resistance to current drugs. Systems biology tools can be used to gain valuable insight into pathogenic processes and aid in expediting drug discovery. In the past decade, constraint-based modeling of genome scale metabolic networks has become widely used. Focusing on pathogen metabolic networks, we review in silico strategies used to identify effective drug targets and highlight recent successes as well as limitations associated with such computational analyses. We further discuss how accounting for the host environment and even targeting the host may offer new therapeutic options. These systems-level approaches are beginning to provide novel avenues for drug targeting against infectious agents.

**Systems biology and pathogen metabolism**

Systems biology methods have been applied extensively to the study of infectious diseases across multiple scales of biological organization to generate predictions ranging from pathogen gene lethality in particular microenvironments [1,2] to dynamics involved in the host immune response to infection [3]. Utilizing the predictive power of computational modeling and systems analysis, wideranging questions related to pathogen virulence, disease progression and host response can be explored to generate hypotheses for more thorough experimental investigation.

The increasing availability of high-quality genome-scale metabolic reconstructions [4] presents an opportunity for the rational and systematic identification of metabolic drug targets in a pathogen of interest. Built bottom-up from functional genome annotations (and a variety of other data sources) and analyzed with computational methods such as flux balance analysis (FBA; see Glossary), these biochemical networks can account for hundreds to thousands of metabolites participating in enzymatic reactions across a range of metabolic subsystems (e.g. carbohydrate, amino acid, lipid, nucleotide and energy metabolism) and cellular localizations (e.g. extracellular space, cytosol and compartments specific to particular organisms) (Figure 1a) [5]. Since the initial genome-scale reconstructions of Escherichia coli [6,7] and Haemophilus influenzae [8,9], the metabolic networks of over 50 organisms (bacteria, archaea and eukaryotes) have been reconstructed (reviewed in [4]). Elements of this network reconstruction process have been automated, allowing the preliminary analysis of hundreds of draft network reconstructions [10]. Among these, metabolic networks have been reconstructed for several pathogenic organisms (Table 1). Indeed, the study of pathogen metabolism–for the elucidation of highpriority drug targets and metabolic factors contributing to pathogenicity—is an exciting application for metabolic network modeling and systems biology.

In this review, we explore several techniques and approaches used to predict antimicrobial drug targets from metabolic network modeling using FBA. Where possible we present examples that have led to novel data, drug targets, or drugs. Metabolic network modeling is still in its infancy, but has allowed for predictions that align with previous data and has provided many hypotheses that continue to be developed. We first discuss the fundamental aspects of network analysis and FBA in particular. Subsequently, we delve into how computational metabolic reconstructions can be used to prioritize drug target predictions. Furthermore, we review recent developments on model-guided pipelines for drug target discovery against pathogens. Finally, we extend the discussion to include host cell metabolism and propose directions for future modeling efforts in infectious disease.

**Reconstructing the metabolic network and defining an objective**

A metabolic network reconstruction is assembled piece-bypiece by compiling data on known enzymes, genes encoding these enzymes, and the stoichiometry of the reactions catalyzed by these enzymes ([11] for a list of databases containing such data). Gene–protein–reaction (GPR) relationships, in the form of Boolean logic statements, define which genes are necessary for each enzyme and which enzymes are necessary for each reaction [5] (Figure 1b). The information for all the reactions in a network reconstruction with m metabolites and n reactions can be stored in an m by n table or matrix, the stoichiometric or S matrix. Each element or cell in this matrix corresponds to the stoichiometric coefficient of one particular metabolite in one particular reaction [5,12]. The S matrix enables strict accounting for the underlying biochemistry and allows a quantitative description of complex interactions between metabolites that are responsible for driving a cellular phenotype. This matrix formalism facilitates interrogation of the structural and functional properties of the network.

The application of FBA to a network reconstruction results in the identification of combinations of reaction fluxes that correspond to a maximum flux through a targeted reaction (an objective) while requiring that constraints are satisfied, for example that the mass entering the network is equal to the mass exiting the network. In more mathematical terms, FBA involves the use of a linear programming formulation wherein an objective is optimized subject to a set of governing constraints (Box 1) [4,12,13]. In addition to requiring mass balance for every reaction, thermodynamic, topological, environmental, and regulatory data may provide additional constraints that dictate the feasible flux space [14]. An objective often used with FBA is biomass production, which is represented in silico as a drain capturing crucial metabolites necessary for growth of the organism [15,16]. With the ultimate goal of identifying antimicrobial drug targets (and associated drugs) to slow or stop the growth of a pathogen, a biomass objective is often very appropriate for computational modeling efforts. Network reconstructions of most pathogenic organisms have incorporated biomass reactions as their objectives (Table 1).

The inability of the metabolic network model to synthesize even one metabolite of the biomass reaction will result in a predicted value of zero for the objective (biomass) flux, and analogously no growth. Therefore, growth predictions are sensitive to metabolites that are placed in the biomass reaction. FBA can be used to investigate the ability of the model to produce each metabolite within biomass. In the Porphyromonas gingivalis metabolic network, the ability of the model to produce each of the 52 metabolites within the biomass objective was evaluated after systematic reaction deletions [17]. Crucial groups of reactions were identified that were responsible for lipopolysaccharide (LPS) production, coenzyme A production, glycolysis, or purine and pyrimidine biosynthesis [17]. Corresponding enzymes that are essential for growth, as well as for the production of important bacterial components such as LPS, could serve as potential drug targets. In addition, in a study of Leishmania major metabolism, the contribution of minimal media components to the synthesis of individual biomass constituents was analyzed [18]. The study found that the absence of cysteine and oxygen in the minimal media had a drastic impact on the overall metabolic network, limiting the generation of 30 of the 40 biomass constituents [18]. Network analyses as delineated in these examples of P. gingivalis and L. major may permit the formulation of a hypothesis for the role of specific metabolites and their influence on growth. Therefore, defining an appropriate biomass reaction (Boxes 2 and 3) is crucial for useful predictions and identification of vulnerable parts of the metabolism of a pathogen.

**The quest for drug targets in metabolic networks of pathogens**

*Gene essentiality analysis*

The most common method to identify potential drug targets has been through the prediction of essential genes (Table 1). The enzymes encoded by essential genes are typically hypothesized as drug targets. Gene knockouts might lead to a redistribution of flux through the network if the perturbed gene or gene product affects the removal of a particular flux-carrying reaction [4]. A GPR aids in mapping the effects of a genetic (or pharmacological) perturbation on the associated reactions, and thus the network. Gene-level perturbations that result in reduced or zero flux through a biomass reaction correspond to growth-reducing or lethal gene knockouts, respectively (Figure 2). For example, in a reconstruction of Mycobacterium tuberculosis, five previously known drug targets were encoded by genes predicted to be essential from computational analysis [19]. For metabolic reconstructions of pathogens, enzymes that are predicted to be essential will offer new experimental hypotheses and avenues for drug discovery. In the following subsections we discuss several other approaches for drug targeting using metabolic network analysis. These other approaches may provide a separate list of potential targets; however, they can also be used in tandem with gene essentiality analysis for step-by-step prioritization of drug targets.

*Enzyme robustness and flux variability*

In addition to an analysis of gene essentiality, assessing enzyme robustness could identify vulnerable or sensitive portions of a metabolic network suitable for drug targeting. To determine the robustness of a metabolic network to the inhibition of an enzyme-catalyzed reaction, the flux of a reaction may be constrained to a fraction of its wild-type flux (simulating partial to complete inhibition), and the effects on an objective flux (e.g. biomass production) can be evaluated [20] (Figure 2). Such an approach was used in analyzing the network reconstruction of Francisella tularensis, which revealed that the growth rate was sensitive to changes in H+ and NH4 flux in a simulated in vitro medium but not in a simulated in vivo medium that mimicked the environment during infection [21]. Analyzing enzyme robustness with different constraints provides a detailed view of possibly non-lethal reactions whose change in flux has a strong effect on the objective of a network under different environmental conditions, therefore suggesting important reactions during an infection or perturbations for drug targeting.

Alternatively, the objective function could be constrained to a fixed percentage of its wild-type flux and the allowable range of flux for each reaction can be determined using an approach termed flux variability analysis (FVA; Box 4) [22]. A recognized shortcoming of FBA is that the typical implementation calculates only one of many possible solutions that optimize flux through the objective reaction. Consequently, there may be many possible routes through the network that achieve the same optimal flux for a given objective [14,22]. In an effort to circumvent this shortcoming, FVA was developed to determine the range of fluxes over which a particular reaction operates, while still allowing for optimal, or near-optimal, objective flux [22]. FVA also identifies blocked reactions (reactions incapable of carrying any flux in a given model under specified constraints) or reactions with different flux ranges in various media. Enzymes that catalyze reactions with little to no variability in flux for a given objective could be selected as potential drug targets given that the network may be sensitive to even modest inhibition of its activity. In analyzing the reconstruction of M. tuberculosis, the average of the highest and lowest allowable flux for each reaction was used as a point of comparison for the model under constraints for two growth rates [19]. In the study, the reaction catalyzed by isocitrate lyase – an enzyme important for persistence in a host – was predicted to have increased flux during slow growth, and the activity of the enzyme was experimentally confirmed to be greater in slow-growing cells of the closely related Mycobacterium bovis [19]. Hence, flux variability and enzyme robustness aid in further prioritizing drug targets identified from other methods such as gene essentiality.

*Metabolite essentiality*

Many current drugs have high similarity to natural metabolites and compete for and/or inhibit normal enzymatic activity [23]. Therefore, another very interesting avenue towards identifying drug targets in metabolic networks is via the prediction of essential metabolites. In contrast to the more traditional individual gene knockouts mentioned previously, metabolites in the S matrix can also be systematically removed. Consequently, all reactions in which a given metabolite participates are removed, and the resultant effects on the objective are assessed (Figure 2). A total of 211 essential metabolites in the Acinetobacter baumanii reconstruction were narrowed to 9 following removal of (i) currency metabolites (ATP, NADH, H2O, etc.); (ii) metabolites present in the human metabolic network; and (iii) metabolites participating in reactions catalyzed by enzymes with human homologs [24]. Enzymes that catalyze reactions involved in the consumption or production of these essential metabolites may be considered as drug targets. Moreover, structural analogs of essential metabolites may be considered as test compounds for experimental evaluation, thus sidestepping extensive screening or computational predictions [25].

*Combination gene and reaction perturbations*

Many anti-infectives on the market act on multiple targets. This multiplicity of targets was exemplified in a network analysis of 890 FDA-approved drugs targeting 394 human proteins (derived from the DrugBank database), where approximately 38% of the drugs were associated with more than one protein target, and a few drugs were associated with as many as 14 targets [26]. In that regard, a drugdiscovery strategy incorporating compounds known to act simultaneously on multiple targets can be adopted for microbial pathogens. FBA allows predictions involving the perturbation of multiple genes or reactions in a rapid timeframe (minutes or hours)—a single in silico combination taking only a fraction of a second [27]. In a reconstruction of M. tuberculosis that accounted for features specific to an in vivo environment, all non-trivial double-deletion mutants (synthetic lethal pairs) were tested for in silico growth using FBA [2]. Of two experimentally-characterized double gene deletions, the model accurately predicted reduced in silico growth in both in vitro and in vivo environmental conditions [2]. A combination of drugs that affect synthetic lethal targets may act synergistically to inhibit growth of a pathogen, thereby paving the way for model-guided predictions of drug synergy. Experimentally screening for all possible drug combinations against a particular pathogen is costly and is often not feasible. Therefore, predicted combinations of drugs associated with synthetic lethal targets can direct more specific experiments and perhaps reveal entirely novel treatment strategies.

*Groups of targets and network topology*

Other approaches characterizing the structure of a genomescale network reconstruction identify sets of reactions that act together and therefore may be targeted as an entire pathway. Sets of correlated reactions (or Co-sets) consist of groups of reactions whose fluxes are linked, and which represent functional modules within a biochemical network [28]. Co-sets, which can aid in suggesting alternative drug targets by identifying reactions that are functionally related to each other, can be divided into several categories. A perfect Co-set consists of a group of reactions such that, for any given pair in the group, a non-zero flux in one reaction implies a non-zero flux in the other, with a fixed ratio [29]. Other categories include partial Co-sets (pairs of linked reactions, but with a variable flux ratio) and directional Co-sets (a non-zero flux in one reaction implies a nonzero flux in the other, but the converse is not true) [29]. By calculating hard-coupled reaction (HCR) sets – a subgroup of perfect Co-sets where sets of reactions are defined by participating metabolites sharing a consumption to production connectivity of 1:1 (i.e. two reactions are linked by a metabolite that is connected to no other reaction) – one study found 25 of 147 HCR sets contained previously identified drug targets in M. tuberculosis [1]. Because an altered flux in one reaction results in an altered flux of all reactions within a HCR set, only one enzyme needs to be targeted. This approach aids in prioritizing the list of potential drug targets by identifying linked enzymatic reactions. Hence, analyses of the topology of a metabolic network can reveal key local and structural features that may be important drug targets when data related to environment fluxes or appropriate objectives are not necessarily available.

*Environment and conditional essentiality*

Finally, the metabolic phenotype is dependent on the media environment and the exchange of metabolites into and out of the system (see Box 3 for discussion of knowledge gaps associated with nutrient availability). By constraining uptake or secretion fluxes, a minimal set of metabolites that allows flux through the objective can be computed [18]. Moreover, enzymes or metabolites that are necessary for growth in various environments (e.g. minimal media, defined media, or rich media with an abundance of nutrients and carbon sources) can be predicted. In the reconstruction of F. tularensis, genes that were essential in a simulated macrophage environment and five other environmental conditions were considered to be unconditionally essential genes (in other words, they represented a core set of genes that were essential regardless of the medium). Of the 17 virulence factors cataloged from previous literature, eight were unconditionally essential genes [21]. By contrast, enzymes that may be necessary for growth under one condition, but not another, are conditionally essential (Figure 2). With a careful consideration of an objective and appropriate nutrient uptake, gene essentiality analysis may reveal new drug targets specific to particular growth conditions and environments in which a pathogen must survive. Such an analysis can also inform strategies for manipulating the environment of the pathogen that could be effective as a treatment option.

**From target to drug and the development of model guided pipelines for drug discovery**

Computational analysis of metabolic processes in pathogens can yield a ranking of predicted drug targets. Bioinformatics and network analyses were performed to yield a high-confidence list of targets against M. tuberculosis [30]. By implementing a multilayered approach, targets that did not pass sequential cut-off values were removed (e.g. elimination of enzymes with human homologs or targets with no computationally predicted binding pocket) [30]. In another proof-of-concept study it was noted that essential type II fatty acid biosynthesis (FAS II) reactions in the E. coli MG1655 metabolic network were also essential in several Staphylococcus aureus strains [31]. Following network analysis, a virtual screening strategy was employed whereby small molecules from a library of approximately 106 compounds were docked to enzymes catalyzing essential reactions, and 41 inhibitors of FAS II enzymes were selected for experimental validation [31]. In cell viability assays, six of the inhibitors had growth-retarding effects against E. coli and S. aureus strains in standard LB agar plates [31]. Finally, following the identification of 163 essential metabolites, a third study used a layered approach to prioritize five essential metabolites in the metabolic network of the opportunistic pathogen Vibrio vulnificus [25]. Currency metabolites, metabolites consumed by a single reaction, metabolites present in the human metabolic network, and metabolites associated with enzymes with human homologs were removed. The study screened 352 compounds found to be structurally similar to one of the five essential metabolites and identified one compound that most potently inhibited growth, more so than a currently used drug [25]. These studies provide various examples of model-guided pipelines to drug discovery by primarily using network analyses to identify and prioritize drug targets. Additional constraints such as enzyme druggability and elevated gene expression can also be used to prioritize drug targets, which can then guide the screening and selection of compounds.

A common approach in proposing drug targets using metabolic networks has been to rule out targets that overlap with host cell metabolism – the idea being that offtargets can be minimized and drug interference plus subsequent complications with the host can be avoided. However, there are arguments to be made in favor of retaining targets that overlap with human metabolism. First, accounting for the drug selectivity between host and pathogen targets at the respective binding sites may preclude off-target influences [32]. Second, if the goal is to discover drugs against infectious diseases quickly, then the best option may be to focus on finding new clinical indications for existing FDA-approved drugs (i.e. pursuing drug repurposing strategies) instead of developing new investigational compounds that are subject to regulatory hurdles [33]. Also, the majority of FDA-approved drugs target human proteins. Hence, eliminating pathogen targets that overlap with human proteins reduces the number of potential drugs that could be evaluated experimentally.

**A host cell perspective**

Interaction with a host cell is often crucial to the metabolism and survival of a pathogen. For instance, the kinetoplastid parasite L. major is unable to synthesize several essential amino acids and therefore obtains them from the host macrophage [34]. As another example, Legionella pneumophila, the bacterium responsible for Legionnaire’s disease, ceases to replicate inside a host macrophage when it cannot access or process threonine [35]. Consequently, identifying the particular niche of nutrients and resources in the host cell required by a pathogen is vital to discovering treatment options that specifically target host–pathogen interactions [36].

A systems-level analysis of pathogen metabolism interfaced with cell type-specific host metabolic networks can also be conducted. Because P. falciparum invades mature erythrocytes to establish infection in its human host, a metabolic model of the human erythrocyte was built in conjunction with the P. falciparum reconstruction to make predictions which aligned closer to known conditions in the infected erythrocyte [37]. This model, modifying an existing approach from Shlomi et al. [38], integrated previous gene expression data where several enzymes were constrained to be ‘on’ and ‘off’ during specific life-cycle stages. The combined erythrocyte-blood stage P. falciparum network correctly predicted metabolite exchanges between the microbe and host [37]. A recently active area of research has been the development of algorithms to create cell type-specific metabolic networks by integrating gene and protein expression data with existing human metabolic reconstructions [38–40]. Inclusion of host-specific factors into pathogen metabolic-network reconstructions or developing systems-level models of host and pathogen networks will continue to enable investigations into the complexities of the host–pathogen interplay.

Finally, inhibiting host pathways and perturbing the flux of metabolites in the host cell may alter the ambient environment and require pathogens to adapt their metabolic needs. Therefore, targeting the machinery of a host cell at the host–pathogen interface can provide new therapeutic approaches [41]. For example, host proteins hijacked for viral replication are potentially important drug targets. Recently, a high-throughput screening assay identified a lipophilic compound – NA255 – that inhibits the host serine palmitoyltransferase, an enzyme needed for association of hepatitis C virus (HCV) with host lipid rafts [42]. Moreover, to characterize transformations in host functions, data specific to the host cell preand post-infection must be obtained. The analysis of transcription profiles is one approach that has been successfully implemented to identify genes in the host cell that are differentially regulated due to pathogenic infection [43,44]. Similarly, profiling the proteome and lipidome of a hepatocyte over the time-course of infection and integrating these data with protein–protein interaction networks revealed multiple lipids and enzymes differentially regulated in HCV-infected cells [45]. A third approach for identifying factors in the host necessary for establishing infection involves the use of genetic screens in which largescale insertional mutagenesis is performed to develop null mutants in a human cell line [46]. Ultimately, the use of new experimental technologies along with metabolic modeling will be vital to discovering host components crucial to the survival of a pathogen.

**Next steps**

Advanced meta-network analyses, such as comparative modeling of metabolic reconstructions across multiple strains and species or community-based modeling of metabolism across differing pathogenic organisms, have broad implications for understanding and investigating infectious diseases. Below, we highlight several future directions in this realm and provide a few examples of efforts already underway.

The recent completion of metabolic reconstructions of the pathogen Pseudomonas aeruginosa [47] and the related nonpathogen Pseudomonas putida [48] creates new opportunities for investigating species-specific differences in metabolism and the metabolic basis for virulence of P. aeruginosa. Towards that end, a reconciliation of the two reconstructions was completed such that any differences in the metabolic networks of P. aeruginosa and P. putida would be indicative of true biological variations as opposed to artifacts of the reconstruction and modeling process [49]. In the reconciliation study, the model for each organism was analyzed to characterize the tradeoffs of producing biomass versus the production of individual metabolites. Compared to P. putida, P. aeruginosa was able to produce a small proportion of the shared virulence factor precursors with only a slight decrease in biomass production. In general, the metabolic flexibility analysis suggested that the virulence of P. aeruginosa is complex and highly multifactorial, and has more flexibility than P. putida in many metabolic pathways. This computational analysis paves the way for future modeling efforts of other infectious disease-causing agents and their basis for establishing virulence.

As another example, syntrophic mutualism between a sulfate-reducing bacterium, Desulfovibrio vulgaris, and a methanogen, Methanococcus maripaludis, was investigated by performing FBA on a compartment-based model involving the metabolic reconstructions of both organisms and a culture medium [50]. In another study, gene expression data were integrated with the genome-scale metabolic reconstruction of P. aeruginosa in the context of a chronic cystic fibrosis lung infection over a 44-month time-course [51]. This analysis provided a systems-level view of bacterial adaptations in a cystic fibrotic lung environment over time. Subsequent studies can shed light on the interactions between multiple organisms over the time course of an infection.

Finally, automated reconstruction platforms such as ModelSEED permit the rapid reconstruction of hundreds of draft bacterial metabolic networks [52,53]. Integration of many such reconstructed networks may help elucidate interactions within the host microbiome and partially explain the development of opportunistic infections that occur primarily because of an altered bacterial flora and environment. For example, an integrative metabolic analysis of organisms in the human gut microbiome could aid understanding of the intricate balance between non-pathogenic and potentially pathogenic organisms during healthy and infectious states in the gastrointestinal tract. An expected outcome of such analyses could result in the selection of drugs or drug cocktails that specifically target pathogens without eliminating non-pathogenic members.

**Concluding remarks**

As reconstructions of metabolic networks become more standard and automated [5], the need for computational tools to characterize these networks becomes more apparent. In addition, the generation and management of large datasets pertaining to both host cell and pathogen intracellular processes of metabolism, signal transduction or regulation has necessitated a systems approach and, therefore, the computational methods used to analyze these data are becoming increasingly important. Experimental methods will continue to improve, thereby generating data that have so far been either impossible or prohibitively laborious to obtain, and which have constrained the value of some model predictions. For example, TraDIS (a new experimental method used to identify all essential genes simultaneously) directly measures gene essentiality that the model could only predict [54]. However, the iterative relationship between modeling and experiment will always permit the generation of novel hypotheses and the contextualization of large datasets, often in a quicker and more cost-efficient manner (e.g. rapid essentiality prediction of all double gene knockouts). Network-based approaches such as genome-scale metabolic reconstructions have been effective in drug target prediction and will continue to expand in scope and applicability. In addition, integration of networks and data into more standard pipelines that traverse the spectrum from computational prediction to experimental evaluation and back again will speed the process dramatically. By including many types of data sources that have yet to be coupled, entirely new classes of drug targets or treatment strategies may be found. The already enormous amount of data is only increasing, and the use of systems biology approaches will be vital to driving future research of drug and drug target discovery against infectious diseases.

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**Glossary**

Biomass: an objective reaction consisting of important metabolites that the cell needs in order to grow (e.g. amino acids, lipids and carbohydrates). Correlated reaction sets: groups of reactions whose fluxes always change in relation to one another.

Exchange reaction: a reaction that transports metabolites into or out of the system.

Feasible flux space: the possible combination of allowable fluxes given a defined set of constraints which bound reaction fluxes, for example reversible and irreversible reactions.

Flux balance analysis (FBA): a linear programming problem formulated to maximize the flux through an objective reaction, which the analyst defines, under steady-state flux constraints that are derived from the stoichiometric matrix of the metabolic network.

Flux distribution: the set of all reaction fluxes within a metabolic network.

Flux variability analysis (FVA): a linear programming-based method which determines the minimum and maximum reaction fluxes that allow for optimal or near-optimal flux through the objective reaction.

Gene–protein–reaction (GPR) relationship: the combination of proteins or protein components sufficient to carry out an enzymatic reaction and the combination of genes sufficient to express each of the protein components.

In silico: in contrast to in vivo or in vitro, the term indicates a computational process in a simulated environment.

Linear programming: also termed linear optimization, an area of mathematics developed for maximizing a linear combination of variables (e.g. A v1 + B v2 + C v3), such that the variables are constrained by many linear equalities and inequalities (e.g. the constraint v1 - v2 = 0 implies that flux v1 is constrained to be twice flux v2).

Mass balance: a requirement that the mass entering the system or any pathway within the system equals the mass exiting the system or pathway; a crucial characteristic of metabolic reconstructions from which functional mathematical models can be derived.

Metabolic network reconstruction: a manually-curated computational network of the metabolism of an organism with all the GPRs assembled from a functionally annotated genome, biochemical data, and literature, that are compiled into a stoichiometric matrix, which serves as the framework for further computational analysis.

Objective flux: the flux through the objective reaction

Objective reaction: a reaction which sets a demand for particular metabolites in the network. The typical goal in formulating this reaction is to simulate a biological objective, whether it be growth, energy, virulence, or a combination of other factors.

Reaction flux: moles consumed in a reaction per unit time.

Steady state: with respect to flux, a key assumption in FBA that the reaction fluxes within the system, and therefore the amounts of each metabolite, do not change over time, an assumption often justified by the very short time-scale of metabolic reactions compared to the time necessary for changes in cell phenotype.

Stoichiometric matrix (S matrix): a mathematical formalization of a metabolic reconstruction; a matrix in which each element contains the stoichiometric coefficient for a metabolite (row) participating in the corresponding reaction (column).

**Box 1. A brief primer on FBA**

Nutrient availability, restrictions on surrounding environmental pH, and temperature are examples of basic constraints that, when imposed upon cells, can affect the resulting phenotypes [12]. Such constraints can be mathematically described, and they serve to narrow the operating range of the cell and yield a set of feasible reaction fluxes for a metabolic network. In other words, the constraints (which can be physicochemical, topological, environmental and regulatory) restrict the number of possible phenotypes, which then allows the function of the biochemical network to be characterized [12,14].

Flux balance analysis (FBA) is one constraint-based method that has been extensively applied in the study of prokaryotic and eukaryotic metabolic networks [47,70]. First, reactions of a metabolic network are assembled into a stoichiometric or S matrix, whose elements correspond to the stoichiometric coefficients describing the conversions from reactants to products (Figure I) [5,12]. This simple matrix formalism permits quantitative description of the complex interactions between metabolites. Following network reconstruction, the concentrations of metabolites and fluxes through reactions can be represented as follows (Equation I):

dC/dt =*Sv*

Here, C is a vector of concentrations of metabolites, t is time, S is the stoichiometric matrix consisting of m rows of metabolites and n columns of reactions, and v is a vector of fluxes through the corresponding reactions. Invoking the steady-state assumption so that the rate of production of every metabolite equals its rate of consumption yields the following (Equation II):

*Sv = 0*

Limits can be applied to individual fluxes as follows:

*vmin ≤ v ≤ vmax*

Particular reactions can have set upper limits (vmax) that may align closely with experimental enzyme capacity measurements, whereas other irreversible reactions will have vmin set to 0.

The principle physicochemical constraint in Equation II represents a set of linear equations. The number of equations is usually less than the number of unknown fluxes [12], and a linear programming optimization can be used to generate a single flux distribution for the entire network by choosing an appropriate objective (or goal):

Max *vbiomass*

Traditionally, maximization of biomass has been chosen as the objective of choice in FBA [12]. A set of metabolites (e.g. amino acids, lipids, nucleotides and carbohydrates) that are necessary for the cell or organism to grow are typically included in the biomass reaction. Therefore, the optimization problem can be summarized as simply:

Max *v*biomass

Subject to *Sv = 0*

*vmin ≤ vi ≤ vmax*  for all *i*

**Box 2. Cellular objectives of pathogenic organisms**

A biomass objective reaction is not applicable under all conditions, and very often growth alone may not be a realistic objective. Other objectives such as maximizing or minimizing ATP or maximizing the production of particular cellular by-products (e.g. lactate or pyruvate) can also be used.

The metabolism of an organism may be adapted for increased virulence or pathogenicity. Pathogens and host cells may also temporarily opt for alternative objectives while under selective pressures (e.g. changes in nutrients or environmental influences of secreted toxins). In addition, different morphological stages of a pathogen (e.g. the sporozoite stage of Plasmodium falciparum in mosquitoes vs the merozoite stage in humans) may be characterized by varying metabolic requirements. Consequently, the objective function must be appropriately defined to find relevant enzyme targets crucial in particular stages of infection and/or environmental conditions.

To explore the effects of targeted perturbations (pharmacological or genetic), different objectives can be explored. It may be that the evolutionary pressures that dictate wild-type cells are different for knockout mutants. Additionally, mutants may not have the ability for immediate regulation of fluxes that allow for optimal growth. Based on these ideas, an approach termed minimization of metabolic adjustment (MOMA) was developed. The requirement for optimal growth is relaxed for gene deletions in MOMA. Instead, MOMA assumes that the overall flux distribution of a gene-knockout mutant will probably not change significantly from that of the corresponding wild type [71]. In terms of flux values, the gene-knockout mutant will remain as close as possible (in Euclidean distance) to the wild-type optimal flux state. MOMA aided in correcting gene essentiality predictions associated with knockouts of fructose-1,6-bisphosphatate aldolase, triosephosphate isomerase and phosphofructokinase in the E. coli central metabolic model [71]. These genes were predicted to be nonessential when biomass was used as the objective for E. coli growth on glucose, which was inconsistent with supporting literature evidence. MOMA yielded a suboptimal flux distribution for a knockout mutant that would not necessarily equal the optimum as dictated by traditional FBA [71].

Approaches such as MOMA that consider alternate hypotheses for the objective of metabolic networks provide a basis for understanding a potential biological goal for pathogenic organisms, especially considering the complexity of the environment surrounding the pathogen of interest.

**Box 3. Knowledge gaps and caveats to metabolic network analysis**

Multiple steps in the model-building process and subsequent analyses are prone to errors that may greatly affect flux and growth predictions and, consequently, predicted drug targets. FBA provides a quality-assurance check that ensures mass balance. Growth rates and gene essentiality predictions are validated against experimental data to ensure the model truly reflects biological processes. Gene essentiality predictions (commonly between 55% and 90%) provide confidence that downstream analyses are based on a high-quality model [1,72]. To ensure the usefulness of any computational pipeline, drug target predictions should be compared to known targets from the literature. Below we discuss more of the particular difficulties and limitations encountered when using metabolic models to identify drug targets.

*Genome annotation*

In any metabolic reconstruction, there may be hundreds of putative metabolic enzymes with no experimentally identified function. These reactions and associated GPR relationships may be assembled strictly based on existing functional annotations of the genome or based on evidence from related organisms. Even some very well characterized enzymes may have other unexpected activities. In such cases, misannotated enzymes may yield incorrect model predictions leading to errors in drug targeting. For instance, an enzyme may be incorrectly predicted to be essential if the activity of another enzyme, which is not included in the network, can account for the same function. Therefore, this represents one important knowledge gap in the assembly of metabolic networks, and the inclusion of more refined enzyme annotations will directly improve drug target predictions [73,74].

*Nutrient availability*

An important challenge in reconstructing and modeling metabolic networks is determining the composition of in silico media. Nutrients available in host environments are poorly characterized. In addition, for any nutrients that are identified, quantitative data on uptake rates are unavailable. Knowledge of the transporters of an organism elucidates which metabolites are transported into the cell. However, information on transporters is particularly limited and, in general, transport reactions lack any experimental evidence or gene associations supporting their presence. Instead, transport reactions are added for proper functioning of the computational model. Because model predictions are dependent on the media environment, nutrients and transporters must be carefully defined.

*Objective function*

As stated previously, in FBA-based modeling of microbial organisms, a biomass objective reaction has often been used. The purpose of the reaction is to ensure a drain of metabolites that are deemed essential to support the growth of the organism. Starting with the estimated weight fraction of important macromolecular components of the cell (e.g. protein, lipid, RNA, DNA and carbohydrate), the relative abundance of metabolites comprised in each group (e.g. amino acids, phospholipids, nucleotides) can be computed [15]. Among several available, a few experimental methods to measure biomass components include chloroform–methanol extraction (lipids), colorimetric protein assays, and gas chromatography–mass spectrometry (protein content) [75]. Fluxes measured directly by metabolic flux analysis, which tracks the movement of 13C from an initial 13C-labeled substrate, may also help to define the objective function [76,77].

An objective function is most likely to cause errors if metabolites are entirely missing or incorrectly included; the relative amounts of each metabolite in the objective reaction (i.e. the stoichiometric coefficients) do not greatly affect FBA results [16,78]. Therefore, logically deducing a biomass objective is often adequate to estimate the growth of an organism. However, additional experimental data can reveal interesting peculiarities that may be used to design a specific biomass objective for a particular organism. Data on growth- and non-growth-associated ATP maintenance can be included in the biomass reaction [79]. Under in vivo conditions when a pathogen is interacting with its host, an aspect that is often unclear is which biomass component(s) to include in the reaction. The composition of the biomass reaction is likely to vary under different physiological conditions, and the choice of metabolites can directly influence model predictions regarding drug targets. For example, failure to include a particular cell-wall component will not necessarily direct flux through reactions that may be crucial in vivo, and therefore the associated enzymes will not be targeted.

**Box 4. A brief primer on FVA**

After the application of various constraints on the biochemical network, the number of allowable network states (or possible phenotypes) is typically large. Depending on the size and interconnectedness of the cellular network, there may be several alternative optimal phenotypes [14]. FBA calculates one of many feasible solutions that result in the same optimal value of the cellular objective.

Flux variability analysis (FVA) calculates the range of flux in each reaction that allows for the same optimal flux through the objective reaction. The objective flux of the reaction is specified as an additional constraint and multiple optimizations are performed to compute the maximum and minimum flux for every reaction in the network:

Max/Min *v*i

Subject to *Sv = 0*

*vbiomass = Zobj*

*vmin ≤ vi ≤ vmax for i = 1…n*

Here, n refers to the number of reactions in the biochemical network, and Zobj is the optimal value of the cellular objective (vbiomass) as obtained by FBA [22].

**Figure captions**

**Figure 1.** The iterative process of model building and refinement. (a) A functionally annotated genome together with data from the biochemical literature are used to assemble a network reconstruction. Flux balance analysis (FBA) allows for modeling and simulation of the reconstructed network. Advanced network analyses (such as gene essentiality or flux variability) allow for identifying potential anti-microbial drug targets. These targets can then be associated with drugs using bioinformatics approaches and obtaining target drug information from a variety of publicly available databases (e.g. STITCH or DrugBank). Predictions involving targets and drugs can be experimentally validated. Any discrepancies between computational predictions and experimental validation can be informative to improving upon and refining the original reconstruction and modeling platform. (b) Gene–protein–reaction (GPR) relationships, central to the assembly of a metabolic reconstruction, define the genes and gene products needed for each enzymatic reaction. Isozymes can be represented with ‘OR’ statements, whereas enzyme subunits required to function together to catalyze a particular reaction can be represented with ‘AND’ statements.

**Figure 2.** Drug targeting in metabolic networks. Various strategies are illustrated for identifying drug targets by performing FBA on metabolic reconstructions. The sample network shows an input media that represents the environment and exchange reactions, intracellular reactions, and an objective reaction that drains metabolites out of the system. An essential reaction and metabolite that, when removed, block any flux through the objective are highlighted in red. In the conditionally essential panel, the absence of the metabolite highlighted in blue causes the highlighted reaction to become essential in the selected media. One of the synthetic lethal pairs of the network is denoted by ‘SL’. The dashed line in the flux variability illustration may represent ‘near-optimal’ objective flux. A robust reaction maintains near-optimal objective flux over a larger range of reaction fluxes.

Table 1. Drug targeting-related analysis of pathogen metabolic networks

Review

|  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| Pathogen | Disease | Drug targeting-related in silico analysis | | | | | | | | Validation | | Novel compounds identified | Refs |
| Gene/reaction essentiality | Minimal media prediction | Conditional essentiality | Synthetic lethality | Flux variability analysis | Enzyme robustness | Metabolite essentiality | Correlated reaction sets | Literature- derived | Novel experimental validation |
| Acinetobacter baumannii | Opportunistic; nosocomial infection | X |  |  |  |  |  | X |  | X |  |  | [24] |
| Burkholderia cenocepacia | Opportunistic; cepacia syndrome | X |  | X |  |  |  |  |  | X |  |  | [55] |
| Francisella tularensis | Tularemia | X |  | X |  | X | X |  |  | X | X |  | [21] |
| Haemophilus influenzae | Otitis media and respiratory infections | X | X |  | X |  |  |  |  | X |  |  | [8,9] |
| Helicobacter pylori | Gastritis; peptic ulceration; gastric cancer | X | X | X | X |  |  |  |  | X |  |  | [56,57] |
| Klebsiella pneumoniae | Klebsiella pneumonia; urinary tract infection |  |  |  |  |  |  |  |  |  | X |  | [58] |
| Mycobacterium tuberculosis | Tuberculosis | X |  | X | X | X | X |  | X | X | X |  | [1,2,19] |
| Neisseria meningitidis | Meningitis; meningococcal septicemia |  | X |  |  |  |  |  |  |  | X |  | [59] |
| Porphyromonas gingivalis | Periodontal disease | X | X |  |  |  |  |  |  | X |  |  | [17] |
| Pseudomonas aeruginosa | Opportunistic; nosocomial infection | X |  |  |  |  |  |  |  | X |  |  | [47] |
| Salmonella  Typhimurium | Gastroenteritis;  diarrhea | X |  |  | X | X |  |  |  | X | X |  | [60–62] |
| Staphylococcus aureus | Opportunistic; nosocomial infection | X | X |  | X |  |  |  |  | X | X |  | [63–65] |
| Vibrio vulnificus | Cellulitis;  septicemia |  |  |  |  |  |  | X |  | X | X | X | [25] |
| Yersinia pestis | Bubonic, pneumonic, and septicemic plague | X | X |  | X |  |  |  |  | X |  |  | [66] |
| Cryptosporidium hominis | Cryptosporidiosis | X |  |  |  |  |  |  |  | X |  |  | [67] |
| Leishmania major | Leishmaniasis | X | X |  | X |  | X |  |  | X |  |  | [18] |
| Plasmodium falciparum | Malaria | X |  |  | X |  |  |  |  | X | X | X | [37,68] |
| Trypanosoma cruzi Chagas disease X X X [69] | | | | | | | | | | | | | |

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