Comparative analyses of parasites with a comprehensive database of genome-scale metabolic models

Maureen A. Carey ID1,2*, Gregory L. Medlock ID3, Michał Stolarczyk ID4,5, William A. Petri, Jr. ID2, Jennifer L. Guler ID2,4, Jason A. Papin ID2,3,6*

1 Department of Microbiology, Immunology, and Cancer Biology, University of Virginia School of Medicine, Charlottesville, Virginia, United States of America, 2 Division of Infectious Diseases and International Health, Department of Medicine, University of Virginia School of Medicine, Charlottesville, Virginia, United States of America, 3 Department of Biomedical Engineering, University of Virginia School of Medicine, Charlottesville, Virginia, United States of America, 4 Department of Biology, University of Virginia, Charlottesville, Virginia, United States of America, 5 Center for Public Health Genomics, University of Virginia School of Medicine, Charlottesville, Virginia, United States of America, 6 Department of Biochemistry & Molecular Genetics, University of Virginia School of Medicine, Charlottesville, Virginia, United States of America

* Current address: Vedanta Biosciences, Inc., Cambridge, Massachusetts, United States of America

* maureen.carey@mac.com (MAC); papin@virginia.edu (JP)

Abstract

Protozoan parasites cause diverse diseases with large global impacts. Research on the pathogenesis and biology of these organisms is limited by economic and experimental constraints. Accordingly, studies of one parasite are frequently extrapolated to infer knowledge about another parasite, across and within genera. Model in vitro or in vivo systems are frequently used to enhance experimental manipulability, but these systems generally use species related to, yet distinct from, the clinically relevant causal pathogen. Characterization of functional differences among parasite species is confined to post hoc or single target studies, limiting the utility of this extrapolation approach. To address this challenge and to accelerate parasitology research broadly, we present a functional comparative analysis of 192 genomes, representing every high-quality, publicly-available protozoan parasite genome including Plasmodium, Toxoplasma, Cryptosporidium, Entamoeba, Trypanosoma, Leishmania, Giardia, and other species. We generated an automated metabolic network reconstruction pipeline optimized for eukaryotic organisms. These metabolic network reconstructions serve as biochemical knowledgebases for each parasite, enabling qualitative and quantitative comparisons of metabolic behavior across parasites. We identified putative differences in gene essentiality and pathway utilization to facilitate the comparison of experimental findings and discovered that phylogeny is not the sole predictor of metabolic similarity. This knowledgebase represents the largest collection of genome-scale metabolic models for both pathogens and eukaryotes; with this resource, we can predict species-specific functions, contextualize experimental results, and optimize selection of experimental systems for fastidious species.
Author summary

Comparative genomics and phylogeny-based assumptions are useful approaches to generate predictions about cellular behavior for data-poor organisms, such as unculturable but clinically-relevant pathogens. Computational approaches, including metabolic modeling, can accelerate such comparisons. Genome-scale metabolic network models serve as a knowledgebase for an organism and enable rigorous and quantitative comparisons of disparate and sparse data, such as genomics and biochemical data, within and across species. Here, we generated a pipeline to create metabolic network models for 192 genomes from protozoan parasites, including the malaria parasite and organisms that cause diarrhea, African sleeping sickness, and leishmaniasis. Importantly, this pipeline was developed to propagate manual curation efforts from one model to others as manual curation remains the field’s ‘gold standard’ for high-quality networks. We compare metabolic behavior across parasites to contextualize experimental results and compare metabolism. We identify which organisms are metabolically similar for the purpose of identifying experimental model systems and find that both metabolic niche and phylogeny influence metabolic similarity.

Introduction

Malaria, African sleeping sickness, many diarrheal diseases, and leishmaniasis are all caused by eukaryotic single-celled parasites; these infections result in over one million deaths annually and contribute significantly to disability-adjusted life years [1–3]. In addition, human infectious and related parasites infect domestic and wild animals, resulting in a large reservoir of human pathogens and diseased animal population [4]. This combined global health burden makes parasitic diseases a top priority of many economic development and health advocacy groups [5–7]. However, effective prevention and treatment strategies are lacking. No widely-used, efficacious vaccine exists for any parasitic disease (e.g. [8–12]). Patients have limited treatment options because few drugs exist for many of these diseases, drug resistance is common, and many drugs have stage specificity (e.g. [13–15]). Thus, there is a pressing need for novel, effective therapeutics. Beyond the economic constraints associated with antimicrobial development [16,17], antiparasitic drug development is technically challenging for two primary reasons: these parasites are eukaryotes and they are challenging to manipulate in vitro.

As protozoa, these parasites share many more features with their eukaryotic host than prokaryotic pathogens do. Thus, antiparasitics must target the parasite while minimizing the effect on potentially similar host targets, similar to cancer therapeutics. Enzyme kinetics can be leveraged such that the drug targets the pathogen’s weak points while remaining below the lethal dose for host [18] or drugs can synergize with the host immune response (e.g. [19,20]). Unique parasite features (i.e. signalling cascades as in [21] or plastid organelles as in [22]) can also be targeted once identified.

Drug target identification and validation are further complicated by experimental challenges associated with these parasites. Many of these organisms have no in vitro culture systems, such as Plasmodium vivax (malaria) and Cryptosporidium hominis (diarrheal disease), or in vivo model system, such as Cryptosporidium meleagridis (diarrheal disease). Some parasite species have additional unique biology and resultant experimental challenges hindering drug development, such as resistance to genetic modification. For example, Plasmodium falciparum (malaria) was considered refractory to genetic modification until recently [23,24]. Entamoeba...
histolytica (diarrheal disease) has also been refractory to efficient genetic manipulation, and the genomes of Leishmania develop significant aneuploidy under selective pressure [25,26]. Although these challenges may be circumvented with new technology, the use of clinical samples, and reductionist approaches, little data exist relative to that which is available for most bacterial pathogens. Without adequate profiling data (genome-wide essentiality, growth profiling in diverse environmental conditions, etc.), we do not have the knowledge to rationally identify novel drug targets. Untargeted and unbiased screens of chemical compounds for antiparasitic effects have proven useful (if the parasite can be cultured, e.g. [27–32]), but this approach provides little information about mechanism of action or mechanisms of resistance development. Typical approaches to study drug resistance, such as evolving resistance to identify mutations in a drug’s putative target, are not possible without a long-term culture system and a relatively well-annotated genome.

As a result of these difficulties (Table 1), data collected in one organism are frequently extrapolated to infer knowledge about another parasite, across and within genera (Fig 1). Toxoplasma gondii is frequently used as a model organism for other apicomplexa due to its genetic and biochemical manipulability [33–36]. Mouse models of malaria [37,38] and cryptosporidiosis [39,40] imperfectly represent the disease and/or use different species than the human pathogen. However, the modest characterization of functional differences among parasite species, beyond comparative genomics (e.g. [41–44]), limits the utility of this extrapolation-based approach, especially broadly among protozoa. Systematic assembly of existing knowledge about parasites and their predicted capabilities could greatly improve the extrapolation-based knowledge transfer by facilitating rigorous in silico comparison. Such systems biology approaches (e.g. genome-scale metabolic modeling) provide a framework to understand parasite genomes, highlight knowledge gaps, and generate data-driven hypotheses about parasite metabolism.

Genome-scale metabolic models are built from genomic data and by inferring function to complete or connect metabolic pathways; these models are supplemented with data from functional genetic and biochemical studies, representing our best understanding of an organism’s

| Species               | Disease            | Treatable? | Drug Resistance? | Culturable? | Genetically tractable? | Percent of genome is 'hypothetical'? |
|-----------------------|--------------------|------------|-----------------|-------------|------------------------|------------------------------------|
| Trypanosoma brucei    | African sleeping sickness | yes        | yes             | yes         | yes                    | 76.40%                             |
| Babesia bovis         | babesiosis         | yes        | no              | yes         | yes                    | 72.00%                             |
| Trypanosoma cruzi     | Chagas disease     | yes        | yes             | yes         | yes                    | 52.90%                             |
| Cryptosporidium hominis| diarrhea          | no         | -               | no          | no                     | 54.10%                             |
| Cryptosporidium parvum| diarrhea          | no         | -               | yes         | yes                    | 4.1%*                              |
| Entamoeba histolytica | diarrhea          | yes        | yes             | yes         | yes                    | 79.80%                             |
| Giardia intestinalis  | diarrhea          | yes        | yes             | yes         | yes                    | 39.20%                             |
| Naegleria fowleri     | encephalitis       | yes        | yes             | yes         | no                     | 31.70%                             |
| Leishmania major      | leishmaniasis      | yes        | yes             | yes         | yes                    | 76.60%                             |
| Plasmodium falciparum | malaria           | yes        | yes             | yes         | yes                    | 37.60%                             |
| Plasmodium vivax      | malaria           | yes        | yes             | yes         | no                     | 43.50%                             |
| Toxoplasma gondii     | toxoplasmosis      | yes        | yes             | yes         | yes                    | 56.20%                             |
| Trichomonas vaginalis | trichomoniasis    | yes        | yes             | yes         | yes                    | 94.00%                             |

Table 1. Summary of select parasitic diseases and their causal organism. Parasites cause important human and animal diseases and have unique biological and experimental challenges that have made interpretation of in vivo and in vitro data challenging. Several examples are shown. Current treatments and associated observed drug resistance are noted. Many well-studied parasites remain refractory to genetic modification and/or still have poor genome annotation. ‘Uncharacterized’ genes were identified via EuPathDB searches for terms such as ‘uncharacterized’, ‘putative’, ‘hypothetical’, etc., for a representative strain. Because each database is heavily influenced by the respective scientific community, some databases such as CryptoDB do not use these terms because the function of so few genes have been validated in the Cryptosporidium parasites. Thus, the genomes of the Cryptosporidium parasites are mostly hypothetical and proposed functions are only putative; the reported percent of genome that is hypothetical is low for this reason (highlighted by an asterisk).

https://doi.org/10.1371/journal.pcbi.1009870.t001
biochemistry and cellular biology. Unfortunately, existing approaches for the construction of metabolic network models are lacking in standardization and scalability and/or biological relevance for eukaryotes. While there are pipelines that include compartmentalization (i.e. RAVEN [45] and merlin [46]), individual high-quality parasite reconstructions (e.g. [47–51]), and scalable pipelines for the construction of many networks (i.e. CarveMe [52], ModelSEED [53]), we sought to build on these tools and the Eukaryotic Pathogens Database (EuPathDB [54]) to leverage genomic information on the EuPathDB database and existing effort towards manual curation of individual reconstructions.

Here, we present a parasite knowledgebase, Parasite Database Including Genome-scale metabolic Models (ParaDIGM), for this purpose. ParaDIGM is a collection of publicly available genome-scale metabolic models, and the computational tools needed to generate and regenerate these models iteratively as new data becomes available. Importantly, these tools also enable the propagation of experimental data collected in a manual curation to closely related organisms. The integration of this genomic and experimental evidence into genome-scale metabolic models enables direct comparison of predicted metabolic capabilities in specific
contexts, rather than the purely qualitative comparisons that can be performed with traditional genomic approaches. We demonstrate the utility of ParaDIGM by comparing metabolic capacity, gene essentiality, and pathway utilization. Ultimately, ParaDIGM can be used to better leverage experimentally tractable model systems for the study of eukaryotic parasites and antiparasitic drug development.

Results

Building ParaDIGM, a parasite knowledgebase

To build a comprehensive collection of genome-scale network reconstructions representing parasite metabolism, we designed a novel network reconstruction pipeline optimized for eukaryotic organisms (Fig 2A). Our pipeline builds on publicly available, open source software and resources [52,54–56] and focuses on the compartmentalization of biochemical reactions (Fig 2A). We applied this pipeline to assemble networks for all publicly available reference genomes from parasite isolates representing 119 species (see Data Availability for link to code and reconstructions). In brief, we obtained 192 high-quality genomes from the parasite genome resource, EuPathDB [54], to generate a de novo reconstruction for each genome (Fig 2A, step 1). We mapped the protein sequence of all open reading frames against a biochemical database [56] to identify putative metabolic functions via gene-protein-reaction mappings. Reaction compartmentalization was adjusted to maintain each gene-protein-reaction mapping but only with the subcellular compartments relevant for each organism. A large proportion of parasite gene-reaction pairs would otherwise be misassigned or removed from the network due to assignment to an incorrect compartment, due to lack of orthologous and compartmentalized reactions in biochemical databases; our pipeline reassigns these reactions to the cytosol or extracellular space (Fig 2B). Although not all functions annotated on EuPathDB are integrated into our de novo reconstructions using this approach, well studied enzymes and pathways are well represented (S1 Fig); discrepancies between EuPathDB and de novo reconstructions can be prioritized in future curation efforts. We also identify metabolic functions not currently annotated on EuPathDB (S1A and S1B Fig).

We next leveraged the manual curation in one parasite reconstruction, P. falciparum (Fig 2A, step 2, curation from [47,57] and in S1 Table), to generate a semi-curated reconstruction for a subset of phylogenetically-related organisms, specifically all Plasmodium sp.. To build these semi-curated reconstructions, we transformed the manually-curated reconstruction using genetic orthology (Fig 2A, step 3) and added all transformed reactions to the recipient de novo reconstruction (Fig 2A, step 4), improving the overlap between our curated and draft networks for Plasmodium reconstructions (S2 Fig). Lastly, all draft and semi-curated reconstructions were gapfilled using parsimonious flux balance analysis (pFBA)-based gapfilling [60,61] to complete biochemical requirements identified in the experimental literature (Fig 2A, step 5) and to produce biomass (see the Additional Information: Online Methods). Gapfilling too adds to the metabolic scope of all reconstructions (S3 Fig). As a result, when compared to manually-curated parasite reconstructions [47–50], semi-curated reconstructions are larger in scope than de novo reconstructions (Fig 2C) and generate predictions with comparable or improved accuracy (Fig 2D). These reconstructions are also more compliant with community standards [62,63] than previous reconstructions for parasites (representative MEMOTE examples shown at https://github.com/maureencarey/paradigm/tree/master/memote_reports).

Our de novo draft reconstructions contain only genetically supported information (prior to gapfilling) and, unsurprisingly, reconstruction size is correlated with genome size (Fig 3A and 3B). The large genome of Chromera velia CCMP2878 (a non-parasitic organism on CryptoDB with 31,799 ORFs and 3,064 reactions) corresponds to a reconstruction with the second most

PLOS Computational Biology | https://doi.org/10.1371/journal.pcbi.1009870 February 23, 2022
unique reactions with 58. Unique reactions are defined here as reactions found in only one reconstruction and no other reconstructions. However, even small reconstructions contain unique reactions prior to gapfilling (Figs 3A and S4A) and the vast majority of these unique reactions are well connected within the network (S4B and S4C Fig). In fact, 39 reconstructions are well connected within the network.
Fig 3. Reconstructions for all eukaryotic organisms with published genomes. (A): Model summary. Genome size is measured here by the number of amino acid sequences encoded by the genome (triangle) and model size is measured by the number of reactions present in the network (square points). Grey rings highlight 100, 500, 1000, 5000, and 10,000 ORFs moving from the center outwards. Genomes are grouped by database, a rough phylogenetic grouping (see Fig 1). Note: T. gondii RH is excluded from all future analyses given only a subset of the genome is available from EuPathDB. (B): Model size is correlated with genome size. Larger genomes tend to generate larger models. Line is fit to a linear regression with R^2 noted (p-value < 0.001); the standard error is not shown. Points are color-coded by database. (C): Unique reactions by database. Number of unique metabolic reactions per database. Unique reactions are defined here as reactions found in every reconstruction within a database and in no other reconstructions outside of that database grouping. Reactions found in different cellular compartments are considered distinct reactions.

https://doi.org/10.1371/journal.pcbi.1009870.g003

contain at least one unique reaction (S4A Fig) and every database has unique functions, or functions that are found in every reconstruction within a database but not other reconstructions (Fig 3C). For example, the group of *Plasmodium* networks share over 200 reactions that are only found in *Plasmodium* reconstructions; among these reactions include hemoglobin breakdown (Fig 3C). The number of unique reactions is correlated with genome size, both before and after gapfilling (S4F and S4G Fig).

In sum, 34% of reactions are in fewer than 10% of models (Fig 4, light grey) and 352 reactions are unique to just a single model (examples in Fig 4). Importantly, these unique reactions are typically well-connected within the network and rarely represent blocked or unconnected reactions (S4B and S4C Fig). A core set of 45 reactions are contained in all 192 reconstructions (Fig 4). Just 3% of reactions are in at least 90% of models (dark grey in Fig 4); reactions shared

Unique functions (in <10 models): 343 reactions, including:
- Molybdopterin synthase (*C. veia*, CryptoDB)
- Gluconokinase (*T. vaginalis*, TrichDB)
- Butyrate transport (*E. histolytica* HMLIMSS,B, AmoebaDB)
- Urate oxidase (*A. castellanii*, AmoebaDB)
- Thiaminase (*V. brassicaformis*, CryptoDB)
- Glutamate-1-semialdehyde aminotransferase (*N. fowleri*, AmoebaDB)
- Dihydropyrimidinase (*T. cruzi* Sylvio X10.1, TriTrypDB)

Core metabolism (in >166 models): 45 reactions
- Trp-tRNA synthetase
- Fatty acid-CoA ligases
- ABC transporters
- Mg2, Ca2, phospholipids

Fig 4. Reaction frequency ranges from unique to core metabolism. Reconstructions help identify rare metabolic functions (light grey box and on histogram, in fewer than 10 reconstructions) and core parasite metabolism (dark grey box and on histogram, in more than 166 reconstructions). Example rare reactions include seven metabolic reactions that are found in only one reconstruction. Of the 45 reactions found in all reconstructions (core metabolism), most reactions correspond to ABC transporter functions for ions or phospholipids. One reaction corresponds with a tRNA synthetase and the remaining correspond to fatty acid-CoA ligases for various fatty acids.

https://doi.org/10.1371/journal.pcbi.1009870.g004
by many models include functions such as glycolytic enzymes. The relationship between genome size and model size is weakened following gapfilling (S4D Fig), likely due to the same biomass formulation for all reconstructions, and the frequency of rare reactions (light grey reactions in Fig 4) increases. ParaDIGM can be used to tease apart the difference between unique, species-specific functions and poorly annotated functions to illuminate the uncharacterized fraction of parasite genomes. To illustrate additional examples of using this resource, we identified niche-specific functions, predicted fluxomics studies to identify divergent enzymes, and identified representative model systems for drug development.

**Niche-specific metabolic functions**

To identify niche-specific functions, we used ParaDIGM to compare the enzymatic capacity of each organism. Specifically, we compared which enzymes are genetically supported and, therefore, present in each reconstruction prior to gapfilling. We performed classical multidimensional scaling using the Euclidean distance between reaction presence for each reconstruction (Fig 5A and 5B). We observe that phylogenetically-related parasites tend to contain similar reactions (Fig 5A). However, while networks generated from genomes within a common genus or species cluster together, models also cluster within environmental niche rather than broader phylogenetic grouping such as phylum. Apicomplexan parasites cluster tightly within genus but not across genera (Fig 5A, Apicomplexa colored by database). Cryptosporidium parasites cluster with other gut pathogens (Fig 5A, gut pathogens in black) rather than other Apicomplexa. Thus, phylogeny is not the sole predictor of model similarity (permutational multivariate analysis of variance, \( p = 0.001 \) using groups of Cryptosporidium, Toxoplasma, Plasmodium, and all other, and homogeneity of dispersion, \( p < 0.001 \)).

Next, we performed random forest classification using reconstruction reaction content to identify the specific metabolic reactions associated with the metabolic niche of the gut environment. The classifier performed well with an AUC of 0.98 and an out-of-bag error rate of less than 8%, supporting our observation that gut parasites contain distinct metabolic reactions. Most important variables (reactions) were associated with being more frequently observed in non-gut pathogens, including gamma-glutamylcysteine synthetase (GLUCYS), glycerol-3-phosphate dehydrogenase (G3PD and G3PD4), an extracellular membrane proton pump (PPA_1), the glycine cleavage system (GCCb, GLYCL_2), phosphogluconate dehydrogenase (GND), and a pyruvate dehydrogenase using lipoamide (PDHa; Fig 5C). Reactions associated with gut pathogenicity included Butanal:NAD+ oxidoreductase (BNORhc), glucan 1-4-alpha-glucosidase (GLCGSD), and starch synthase (STARCH300Sc; Fig 5C).

Similarly, parasites that invade red blood cells, including Plasmodium and Babesia, are dissimilar when comparing their full reaction content (Fig 5B, triangles); however, the same analysis limited to each organism’s genetically encoded transporters reveals that these parasites have relatively similar transporter capabilities (Fig 5C, triangles). This result indicates that these red blood cell-invading parasites rely on similar nutrients from their host red blood cell. On the contrary, the broad metabolic niche of extracellular growth yielded some outliers regarding enzymatic capacity and transporter profile (Fig 5B and 5C, circles), likely due to the range of environments that parasites capable of extracellular growth encounter.

**Predicting metabolic function**

Beyond the direct comparison of enzyme presence, we can use ParaDIGM to predict metabolic functions and the functional consequences of reaction presence and network connectivity. This approach augments the analysis beyond mere genetic comparisons: some enzymes may not be discovered in the genome despite being necessary to perform biochemical function
Fig 5. Identifying metabolic niches. (A): Reaction content. Classical multidimensional scaling was performed on the reaction content of all de novo reconstructions; each reconstruction is represented by a point (grey/black or colored by database for emphasis). Thus, this analysis focuses exclusively on the genetically supported features of each reconstruction. Apicomplexan parasites (colored by database) and all other gut pathogens (black points) are highlighted. (B): Reaction content with alternative color scheme. Parasites that invade red blood cells (triangles, *Plasmodium* and *Babesia*) or can replicate extracellularly (circles) are highlighted; all other parasites are in lighter grey squares. (C): Important variables for the classification of gut pathogens. We performed a random forest classification to distinguish organisms that are considered gut pathogens from other organisms in ParaDIGM (AUC = 0.98 and an out-of-bag error rate of less than 8%). Important variables with a difference in occurrence score of 1 were present in 100% of gut pathogens and 0% of other organism’s reconstructions and those with a score of -1 were present in 100% of non-gut pathogens and 0% of gut pathogen’s reconstructions. (D): Transporter profile. Again, parasites that invade red blood cells (triangles) or can replicate extracellularly (circles, like the kinetoplastids and *Giardia*, among others) are highlighted, with all other parasites are in lighter grey squares. Red blood cell-invading parasites cluster.
Fig 6. Predicting metabolic function. (A): Advantage of network-based approaches. Metabolic models include hypothetical functions (i.e. the enzyme encoded by gene2) that are unsupported by direct genetic evidence but may be indirectly required based on biochemical evidence. These functions are added through gapfilling. Using models augments our analysis beyond mere genetic comparisons: some enzymes may not be discovered in the genome despite being
necessary for biochemical observations made and are included in these models. **(B): Defining metabolic capacities.** With our gapfilled models, we can identify if metabolites are consumed and/or produced. **(C): Experimentally-derived metabolic functions.** We compiled data providing evidence for consumption or production of select metabolites from the literature (S1 Table). Consumed metabolites are imported by the parasite from the extracellular environment (e.g. the in vitro growth medium). Produced metabolites are synthesized by the parasite even when the metabolite is not in the extracellular environment. See Additional Information: Online Methods for more detail. Data are sparse. **(D): Analogous in silico metabolic capacity.** Inferred metabolic capacity of each organism from Panel C for every metabolite from panel C. Data from panel C was used to gapfill reconstructions to generate data presented in Panel D (see Fig 2A for methods). See Panel B for definitions. Metabolites that are neither produced nor consumed are consumed intracellularly but are not taken up from the extracellular environment. Metabolites noted as ‘complex or unknown’ here are represented by multiple metabolite identifiers in the reconstructions (e.g., lactate is measured experimentally, but could represent both D-lactate and L-lactate within the reconstructions). **(E-G): Example gapfilled functions in the Vitamin B6 pathway.** These reactions were added to support the observed metabolic functions in Panel C or to support in silico growth. Panel E shows L-alanine-alpha-keto acid aminotransferase (ASPTA6, added to 58 reconstructions), Panel F shows pyridoxamine-pyruvic transaminase (PDYXPT_c, added to 64 reconstructions), and Panel G shows pyridoxamine oxidase (PYDHO, named pyridoxal oxidase in BiGG, added to 90 reconstructions). Note, a deaminating pyridoxaminoxygen oxidoreductase (PYDHO_1) is also added to 12 reactions to interconvert pyridoxal and pyridoxamine.

https://doi.org/10.1371/journal.pcbi.1009870.g006

which there is a genetically-encoded transporter (S2 Table) or gapfilled transport reaction (S3 Table). A schematic for each metabolite categorization is shown in Fig 6B with experimental data shown in Fig 6C, with untested or unknown results in white, and analogous in silico results in Fig 6D. All models except for one (Chromera velia CCMP2878 with the largest genome) required gapfilling to synthesize one or more metabolites and/or biomass. We can expand the in silico predictions to all metabolites in all models (a total of 5,141 metabolites by 192 models, S5 Fig) to generate hypotheses about understudied metabolites and enzymes.

Interestingly, several metabolic enzymes were consistently predicted to be necessary for observed metabolic capabilities (metabolic tasks in Fig 6C) or growth across all parasites (gapfilled reactions in Table 2); three common examples are shown in Fig 6E and 6G including three steps in Vitamin B6 metabolism. Pyridoxamine oxidase (Fig 6G) is an understudied enzyme involved in Vitamin B6 metabolism; fewer than 300 articles on PubMed describe the enzyme. Not surprisingly given the lack of literature, the reaction associated with this enzyme is in just seven reconstructions in the BiGG database, including two iterations of the S. cervisiae S288C model [56]. The deaminating version of this reaction is in only 10 reconstructions in the BiGG database; all ten of these reconstructions are for eukaryotes including five Plasmodium genomes. Pyridoxamine oxidase was only added to the V. brassicaformis CCMP3155 and G. niphandrosodes reconstructions in the bioinformatics-driven model construction steps; however, this enzyme was added in 90 gapfilling solutions to satisfy experimentally-derived functions. Thus, we predict that it is important for parasite growth. We also predict that the unidentified sequences for pyridoxal oxidase are highly divergent from known sequences because they were not identified using bioinformatic annotation methods. By comparing the reconstructions within ParaDIGM, we can identify high-confidence reactions that are encoded by divergent genetic sequences and missed by purely bioinformatic approaches.

Selecting the most representative model system for an experiment

Genome-wide essentiality screens are available for Plasmodium falciparum [64] and berghei [58,65], Toxoplasma gondii [35], and Trypanosoma brucei. Using the models generated with ParaDIGM, we can perform the equivalent in silico simulations (Fig 2D and S3 Table) regardless of experimental genetic tractability (Table 1). These analyses can be used to identify drugs for repurposing or the best model system for testing a novel drug target. To do this, we sequentially removed each reaction from the reconstruction to identify which reactions are necessary for growth (i.e. production of biomass). These simulations are performed in an unconstrained model (i.e. all metabolites with a transporter can be imported, all enzymes can be used) to simulate the parasite’s growth intracellularly in the nutrient-rich host cell. Dissimilarity of reaction essentiality for all Toxoplasma, Plasmodium, and Cryptosporidium reconstructions was calculated using the Euclidean distance (Fig 7).
Reaction essentiality is generally more similar for closely related organisms (i.e. within genera). However, some genera generate more similar predictions than others. Essentiality predictions were more similar when comparing *Plasmodium* genomes to one another than between *Toxoplasma*, despite all genomes being of the same species, or *Cryptosporidium* genomes. *Cryptosporidium* genomes generate predictions that are significantly less similar than *Plasmodium* genomes. Essentiality predictions in *T. gondii* are less similar to *Cryptosporidium* parasites than to *Plasmodium* (*Fig 7*). As *T. gondii* is a popular model system for other parasites, this result supports the use of *T. gondii* to test hypotheses about *Plasmodium* over *Cryptosporidium*. Moreover, we can identify organisms that are particularly unique within a genus. For example, *C. parvum* is a poor representative of *C. ubiquitum* whereas *C. muris* and *C. andersoni* are quite similar. Despite being distinct immunotypes, *T. gondii* VEG and GT1 are the most similar *Toxoplasma*. *P. vivax* Sal-1, an unculturable and clinically relevant *Plasmodium* species, is more similar to *P. knowlesi* H than the average two *Plasmodium* genomes, whereas *P. falciparum* 3D7 and *P. berghei* ANKA are among the most dissimilar *Plasmodium* genomes. Importantly, more complete models generate less similar predictions indicating differences in essentiality reflects functional differences, not merely incomplete genome annotation resulting in incomplete reconstructions (*S6 Fig*). These results highlight how ParaDIGM can be used to identify functional similarities and differences between parasites that directly inform experiments for developing and studying new drugs.

### Table 2. Most frequently gapfilled reactions.

These reactions (in the BiGG namespace) were the most commonly added reactions as a result of all gapfilling steps.

| Reaction       | Gapfilled N times | Reaction Name                                                                 |
|----------------|-------------------|-------------------------------------------------------------------------------|
| NADPPPS        | 96                | NADP phosphatase                                                             |
| PYDXO          | 90                | Pyridoxal oxidase                                                            |
| IMPtr           | 86                | Transport of IMP                                                             |
| SO4HCOtex       | 84                | Sulfate transport via bicarbonate countertransport                            |
| EX_lyslyls_e   | 81                | LysLysLys exchange                                                           |
| LYSLYSLYSr      | 81                | Metabolism (Formation/Degradation) of LysLysLys                              |
| LYSLYSLYSt      | 81                | LysLysLys transport                                                           |
| PSERT           | 80                | Phosphoserine transaminase                                                   |
| PGCD            | 75                | Phosphoglycerate dehydrogenase                                               |
| GTHOXii         | 74                | Glutathione transport                                                        |
| CYSLY3         | 65                | Cysteine lyase                                                                |
| NNDPR           | 65                | Nicotinate-nucleotide diphosphorylase (carboxylating)                         |
| PDYXPT_c        | 64                | Pyridoxamine-pyruvic transaminase                                             |
| H2O2t           | 63                | Hydrogen peroxide transport                                                  |
| PSP_L           | 60                | Phosphoserine phosphatase (L-serine)                                          |
| EX_ileargle_e  | 59                | IleArgle exchange                                                            |
| ILEARGILEr      | 59                | Metabolism (Formation/Degradation) of IleArgle                               |
| ILEARGILEt      | 59                | IleArgle transport                                                            |
| lipid2          | 59                | aggregation of all fatty acyl-CoAs                                            |
| ASPTA6          | 58                | L-alanine-alpha-keto acid aminotransfer                                       |
| GMPR            | 56                | GMP reductase                                                                 |
| GTHRDH_syn      | 55                | Glutathione hydralase                                                        |
| GTHPe           | 53                | Glutathione peroxidase                                                        |
| H2Ot            | 51                | Water transport                                                               |
| HISD_c          | 48                | Histidine degradation to glutamate                                            |

https://doi.org/10.1371/journal.pcbi.1009870.t002
Here, we presented a novel pipeline for generating metabolic network reconstructions from eukaryotic genomes and applied it to create 192 reconstructions for parasites, expanding the scope of parasite modeling. These reconstructions represent the first genome-scale metabolic network reconstructions for all but nine of these organisms, making ParaDIGM the broadest computational biochemical resource for eukaryotes to date. ParaDIGM uses reaction and metabolite nomenclature from the Biochemical, Genetic and Genomic knowledge base (BiGG, which includes both microbial and mammalian genome-scale metabolic network reconstructions) [56], facilitating future work involving host-pathogen interaction modeling. Gene nomenclature used in ParaDIGM is from the Eukaryotic Pathogens Database (EuPathDB) [54], consistent with the parasitology field standards and 'omics data collection. Reproducible data integration approaches are used to curate each reconstruction, making this the first fully automated reconstruction pipeline for eukaryotes; code and data are available in the Data Availability section for iterative improvements by ourselves and others.

**Discussion**

Here, we presented a novel pipeline for generating metabolic network reconstructions from eukaryotic genomes and applied it to create 192 reconstructions for parasites, expanding the scope of parasite modeling. These reconstructions represent the first genome-scale metabolic network reconstructions for all but nine of these organisms, making ParaDIGM the broadest computational biochemical resource for eukaryotes to date. ParaDIGM uses reaction and metabolite nomenclature from the Biochemical, Genetic and Genomic knowledge base (BiGG, which includes both microbial and mammalian genome-scale metabolic network reconstructions) [56], facilitating future work involving host-pathogen interaction modeling. Gene nomenclature used in ParaDIGM is from the Eukaryotic Pathogens Database (EuPathDB) [54], consistent with the parasitology field standards and 'omics data collection. Reproducible data integration approaches are used to curate each reconstruction, making this the first fully automated reconstruction pipeline for eukaryotes; code and data are available in the Data Availability section for iterative improvements by ourselves and others.

ParaDIGM or individual reconstructions can be used for comparative analyses or applied to interrogate clinically- and biologically-relevant phenotypes. The adherence to community standards for metabolic modeling throughout ParaDIGM enables easier manual curation for users interested in studying a specific parasite in more detail. Together, this adherence to standards and the automated approach for integration of experimental data, will accelerate further curation of ParaDIGM itself as genome annotation improves, more experiments are
performed with individual parasites, and ParaDIGM users provide feedback on reconstruction usage and performance.

This eukaryote-specific reconstruction process (Fig 2A) generates comprehensive networks of comparable quality to manually curated parasite reconstructions (Fig 2B-2D). However, manual network curation and adding condition-specific constraints remain the gold standard approaches to maximize the accuracy of network predictions, especially for modeling stage-specific metabolism (i.e. [58]). Even so, our semi-automated curation approach enhances the genome-wide coverage of each reconstruction (Fig 2C) and generates models with comparable accuracy to previously published manually curated reconstructions (Fig 2D); maximum model accuracy is dependent on including compartmentalized reactions in the reconstruction process (S7 Fig).

To evaluate these networks, we compare in silico predictions to experimental results; all have imperfect accuracy regarding gene essentiality (Fig 2D), emphasizing how challenging it is to make a truly predictive model without integrating extensive stage-specific experimental data. High rates of false positives (when the model incorrectly identifies a gene as essential) are a product of the model building process; these reconstructions are built to summarize all metabolic capabilities of the organism, not the specific stage-dependent phenotype of an organism in the experimental system. Thus, constraining a reconstruction with in vitro expression data will reduce the false positive rate (e.g. [47,58]). We also compared our manually curated P. falciparum 3D7 reconstruction to our new semi-curated reconstruction for the same species. Differences in the two iterations fall into three groups: (1) non-specific genes that map to multiple reactions, (2) non-enzymatic genes (specifically, tRNAs), and (3) metabolic functions not yet encoded in the BiGG database for which reaction objects were created in our manual curation efforts. These differences can inform the first round of curation for semi-curated reconstructions.

However, this pipeline offers a few key limitations. First, simulation accuracy remains low (Fig 2D), largely because annotation pipelines may over-annotate function and (importantly) these models represent the metabolic capacity of many life stages, whereas experimental data is derived from a single timepoint. Thus, without constraining these models with stage-specific data, models will under predict essential genes and have poor accuracy. Furthermore, all reconstructions are limited by the data used for their construction; for example, we used the reactions already documented in the BiGG database as a universal set of reactions. Thus, reactions not contained in BiGG will not be included in ParaDIGM and only the reactions with specific cofactor utilization or directionality documented in BiGG will be included. Similarly, limited experimental data are available for the localization of specific enzymes and transporters and there has been limited successful experimental validation of computational predictions. Transporters in particular will influence pathway usage and a large proportion of transporters were added in the gapfilling process (S8 Fig). Lastly, the data incorporated in objective reactions and extracellular environment (i.e. media formulation) heavily influences which reactions are considered essential and which non-genetically supported reactions are added via gapfilling. Currently, these constraints within ParaDIGM are not experimentally-derived for each organism.

The metabolic capacity represented in ParaDIGM will be expanded and the accuracy of each reconstruction and associated simulations will be improved as (1) BiGG is further expanded, (2) more confidence is gained regarding protein localization, and (3) metabolomics analyses improve biomass and media formulation. Additionally, we used our orthology-based semi-curation approach for only Plasmodium models; however, this approach can be used for other organisms to propagate manual curation efforts (from our group and others, e.g. [48–50]) from one species to closely-related organisms as well. Finally, these reconstructions have
not been manually curated and require such attention for improved accuracy, especially in well-studied pathways and transporters and to represent stage-specific phenotypes. Despite these limitations, the models within ParaDIGM perform similarly to manually curated models (Fig 2D) and so we highlight example use cases. First, we used ParaDIGM to better leverage model systems for drug development by identifying divergent or conserved metabolic pathways between select human pathogens. Network structures were quite unique with only 25.8% of all reactions in more than 50% of the reconstructions (Fig 4); network topology did however cluster by genus, and transport ability is associated with specific host environments (Fig 5). Despite these structural similarities, minor topological differences in networks (and unique functions, Figs 3C and 4) confer key metabolic strengths or weaknesses (Figs 6 and 7). We compare metabolic reaction (or enzyme) essentiality to identify the best in vitro system or non-primate infection model of disease for drug development (Fig 7). For example, enzyme essentiality is broadly more consistent between Toxoplasma gondii and Cryptosporidium parasites than between T. gondii and the malaria parasites. By leveraging network context (Fig 6A and 6B), we can impute fluxomic studies in all 192 parasites (Figs 6D and S5) to contextualize the variable results across species in relatively few in vitro fluxomics studies (Fig 6C) and to expand these observations to untested organisms and metabolites (S5 Fig).

Beyond our use cases of ParaDIGM, the pipeline and reconstructions presented here can be used broadly by the field. The study of microbial pathogens generated paradigm-shifting results in biology. The study of viral pathogens revealed basic cellular machinery present nearly ubiquitously in eukaryotic cells, such as the discovery of alternative RNA splicing in adenovirus [66]. The study of bacteria has provided a nearly real-time observation of evolution, allowing researchers to perform hypothesis-driven evolutionary biology experiments in addition to observational research [67]. These microorganisms have shed light on cell biology and the history of life in impactful yet highly unanticipated ways; experimental challenges associated with parasites have slowed their utility in this regard. However, both the genetic ‘dark matter’ of eukaryotic parasites and known parasite-specific functions are abundant (Fig 1); thus, parasites too have the capacity to inform our understanding of life. The reconstructions in ParaDIGM can be used broadly to contextualize existing experimental data and generate novel hypotheses about eukaryotic parasite biochemistry as it relates to the rest of the tree of life.

ParaDIGM provides a framework for organizing and interpreting knowledge about eukaryotic parasites. The reconstruction pipeline designed for ParaDIGM implements and builds on field-accepted standards for genome-scale metabolic modeling and the latest genome annotations in the parasitology field; moreover, it is uniquely tailored to eukaryotic cells by recognizing the importance of compartmentalization and the design of the objective function. The pipeline can be implemented with other organisms and re-implemented iteratively to incorporate novel genome sequences, biochemical datasets, genome annotations, and reconstruction curation efforts. The genome-scale metabolic network reconstructions organized in ParaDIGM also can be used broadly by the scientific community, using the reconstructions as-is as biochemical and genetic knowledgebases or as draft reconstructions for further manual curation to maximize the utility and predictive accuracy of the models. These reconstructions can be used to generate targeted experimental hypotheses for exploring parasite phenotypes, ultimately improving the accessibility of modeling approaches, increasing the utility of parasites as model systems, and accelerating clinically-motivated research in parasitology.

Supporting information
S1 Table. Automated curation tasks. All reconstructions were gapfilled to ensure the network could consume or produce all relevant metabolites outlined below. Data from multiple strains
of one species were aggregated. The first two columns describe each metabolite with a sub-
ystem and name. The first two columns represent the genus and species for which literature evi-
dence was compiled. Each \( \{i,j\} \) position in the matrix represents whether there is experimental
evidence for a species’ consumption or production of the metabolite. Blank cells indicate no
literature evidence was found for that metabolite in that species.

**S2 Table. Genetically-encoded transporters.** Transporters in each reconstruction prior to
gapfilling. These transporters are annotated into each genome.

**S3 Table. Gapfilled transporters.** Transporters added to each reconstruction in the gapfiling
process. These transporters are necessary to generate flux through the biomass reaction using
parsimony-based gapfilling.

**S4 Table. Available essentiality datasets.** Experimental genome-wide essentiality datasets
that are available in the literature. These data for *Toxoplasma* and *Plasmodium* were used to
evaluate model performance and specifically simulated gene essentiality.

**S5 Table. Compartmentalization.** Subcellular compartments used for reconstructions in each
genus.

**S6 Table. Blocked reactions.** Each row represents a reconstruction (named in column 1). All
following columns list out the BiGG identifiers for the blocked/unconnected reactions in that
reconstruction. The *products* of blocked reactions are not used in any other reaction, whereas
the *reactants* of unconnected reactions are not generated by any other reaction. Row headings
are arbitrarily counting the number of reactions in each reaction in that category.

**S1 Fig. Comparison of ParaDIGM gene coverage to genes annotated on EuPathDB with
GO terms relating to metabolism.** (A): Schematic for gene count comparisons. Genes
found on EuPathDB with a GO term related to ‘metabolic processes’, genes incorporated into
de *novo* reconstructions (\( N = 192 \) reconstructions), and genes in both of these categories are
described. (B): **Number of genes per category.** Boxplots represent the total number of genes
per category across all reconstructions. (C): **Number of genes EuPathDB associated with
three example pathways.** These genes represent the intersection of EuPathDB genes and
reconstruction genes for three select pathways. For all panels, the box extends from the lower
to upper quartile values of the data; the center line marks the median and whiskers shows the
range of the data. Outliers are not shown. (D): **Percent of genes on EuPathDB associated
that are represented in reconstructions.** Again, these genes represent the intersection of
EuPathDB genes and reconstruction genes for three select pathways. For each example path-
way in Panel C, the percent of total genes from EuPathDB that are represented in the *de novo*
reconstructions are shown.

**S2 Fig. Reaction content overlap between Plasmodium reconstructions in ParaDIGM and
a manually-curated reconstruction, iPfal22.** Semi-curation improves reaction content over-
lap between *Plasmodium* reconstructions in ParaDIGM and a well-curated reconstruction.
Venn diagram of reaction content between three *Plasmodium* species or strains (*falciparum*,
3D7 in A, falciparum Dd2 in B, and berghei ANKA in C) for the draft reconstruction including only genes identified by Diamond and the semi-curated reconstruction (see Additional Information: Online Methods), and compared to iPfal22.

S3 Fig. Benchmarking of gapfilling approach on Plasmodium reconstructions. Only Plasmodium reconstructions were assessed here because we can compare the resultant reconstructions to a manually-curated reconstruction, iPfal22. (A): Number of reactions added at each step of the reconstruction pipeline per reconstruction. For each boxplot, the box extends from the lower to upper quartile values of the data; the center line marks the median and whiskers shows the range of the data. Outliers are not shown. (B-D): Venn diagrams highlighting the shared reaction content of three Plasmodium reconstructions with and without gapfilling. Gapfilling increases the coverage of functions contained in the well-curated reconstruction, iPfal22.

S4 Fig. Further characterization of ParaDIGM reconstructions. (A): Unique reactions by reconstruction 39 reconstructions contain at least one unique metabolic reaction, or a reaction not found in any of the other 191 models. Reconstructions are colored by EuPathDB grouping, like in panel A, and the bar represents the number of unique reactions in that reconstruction. (B-C): Unique reactions are well connected. Percent (B) and total number (C) of all unique reactions per reconstruction that are blocked, unconnected, or both blocked and unconnected. Blocked reactions are defined as those whose products are not utilized by any other reactions (including transport and exchange reactions), whereas unconnected reactions are those whose reactants are not produced by other reactions. For all panel B, the box extends from the lower to upper quartile values of the data; the center line marks the median and whiskers shows the range of the data. Outliers are shown as points. For panel C, each column or bar represents an individual reconstruction. Most unique reactions are connected and unblocked. (D-E): Gapfilled model size remains correlated with genome size. Following gapfilling, the relationship between genome size (as measured by open reading frames or genes with metabolic GO terms, D and E respectively) and model size remains; however, the correlation is weak due to an increase in the number of reactions for reconstructions built from medium-sized genomes. (F-G): Larger genomes have more unique reactions before (F) and after (G) gapfilling. Genome size is measured by open reading frames. For panels D-G, line is fit to a linear regression with R² noted (p-value < 0.001); the standard error is not shown. Points in D-G are color coded by database.

S5 Fig. Complete in silico metabolic capacity. Inferred metabolic capacity of each organism (rows) for metabolites (columns) for every reconstruction and metabolite in ParaDIGM (5,141 metabolites by 192 models). See Fig 6B for definitions. Note the sheer volume of data acquired from ParaDIGM.

S6 Fig. Evaluation of prediction dissimilarity and reconstruction coverage. (A-C): Distribution of reconstruction completeness scores for Cryptosporidium, Plasmodium, and Toxoplasma reconstructions. Completeness scores were calculated by identifying the ratio of metabolic functions (reactions) in a single reconstruction compared to the sum of metabolic functions covered by all reconstructions from the respective genus. (D): Prediction dissimilarity is correlated to model completeness. For each pair of models, reaction essentiality predictions were compared to generate a dissimilarity score (Fig 7A). Identical predictions have a
dissimilarity score of 0 whereas a high dissimilarity score indicates divergent predictions.

**S7 Fig. Compartmentalization improves prediction accuracy and coverage of the genome.**

(A): Sensitivity and specificity of *in silico* gene essentiality predictions when compared to experimental data. Reconstructions generated using our pipeline with and without compartmentalization are connected with a line; other points represent published models for reference. Our compartmentalization approach improves the sensitivity and or specificity of gene essentiality predictions for *P. berghei* ANKA, (*P. falciparum*) 3D7, and *T. gondii* GT1. (B): Venn diagram of gene content for the ParaDIGM *P. falciparum* 3D7 reconstruction (with and without compartmentalization) and our manually curated *P. falciparum* 3D7 reconstruction, iPfal22. Incorporating compartmentalization improves genome coverage by adding 12 genes also found in iPfal22 and 17 genes not found in iPfal22.

**S8 Fig. Only half of transporters are genetically supported.** (A): Percent of all transporters with gene annotations. (B): Percent of intracellular transporters with gene annotations. Red dotted line indicates mean.

**S9 Fig. Distribution of blocked and unconnected reactions.** The distribution of poorly connected reactions (A: blocked, B: unconnected, C: both) was similar before and after gapfilling.

**Acknowledgments**

The authors would like to acknowledge the helpful discussion and feedback from members of the Petri, Mann, Guler, and Papin labs, as well as Drs. Alison Criss, Norbert Leitinger, Herve Agaisse, and Young Hahn. The authors would also like to thank the University of Virginia ARCS staff for support regarding UVA’s High Performance Computing cluster, especially Karsten Siller and Katherine Holcomb. Lastly, the authors would like to thank the EuPathDB, BiGG, and CobraPy communities for providing essential tools (software and database infrastructure) and inspiration.

**Additional information**

**Online methods**

**Code dependencies**

R [68] and R packages tidyverse, ggpubr, ggdendro, seqinr, Biostrings, msa, reshape2, UpSetR, cluster, ade4, RColorBrewer, readxl, dplyr, and ggdendro were used for analysis or visualization [69–80]. Python 3.6.4, pandas [81], and CobraPy 0.14.1 [55], as well as code to implement Diamond-based annotation scoring from CarveMe [52], were used for the reconstruction and modeling. Memote [62] was used to evaluate all reconstructions.

**Genomic analyses**

Sequences were obtained from EuPathDB release 44 [54]. EuPathDB curates and compiles genome annotation for all genomes hosted by the database. We used open reading frames identified on EuPathDB and annotated the sequences with Diamond, described below. EuPathDB’s OrthoMCL was used for mapping orthology between *Plasmodium* species. In brief, orthology was mapped within each EuPathDB database by the ‘map by orthology’ tool from the genome of each organism with a curated reconstruction to all other genomes within that database. The search protocol was ‘new search > genes > taxonomy > organism {pick} >
transform by orthology. We mapped each organism’s amino acid sequences using Diamond annotation against proteins referenced in the BiGG databases. Diamond is a similar approach to BLAST, with sensitive and fast performance on protein annotations.

**Model generation**

We generated draft reconstructions by first annotating each organism’s amino acid sequences, obtained from EuPathDB, using Diamond annotation against enzymes and transporters referenced in the BiGG databases. We next mapped all functional annotations to reactions contained in the BiGG database inspired by the approach conducted with the reconstruction pipeline CarveMe. Next, for all reconstructions, we added any KEGG reaction-associated genes from EuPathDB if the KEGG identifier could be converted into a BiGG identifier, using KEGG-BiGG mappings found at http://bigg.ucsd.edu/static/namespace/bigg_models_metabolites.txt. Importantly, EuPathDB annotations used for this KEGG-BiGG mapping are derived from a mix of variable automated and manual approaches and include user contributions. Methods are included in the analytic code hosted on Github, at https://github.com/maureencarey/paradigm.

Unlike the CarveMe approach, we included all high-scoring reactions. CarveMe maximizes the number of high-scoring hits while building a functional network. Our conservative approach generates broadly inclusive but incomplete reconstructions (i.e. that are not able to produce biomass until gapfilled). This approach added redundant reactions from multiple different compartments (e.g. peroxisome, mitochondria, and cytosol) so all reaction versions other than the cytosolic version were pruned unless localized to a relevant compartment (Table); for genera not included in Table, only the cytosol and extracellular space were used to avoid inclusion of erroneous compartments. For example, if a Plasmodium reconstruction contained a reaction in the cytosol, mitochondria, and chloroplast, only the cytoplasmic and mitochondrial reaction versions would be kept. Following this step, a large percentage of each reconstruction’s reactions remained in unsupported compartments because there was no analogous cytosolic (or extracellular) reaction (Fig 2B). Next, reactions only found in an unsupported compartment were moved to the extracellular space or cytosol; specifically, periplasmic metabolites were moved to the extracellular space and all internal subcompartment metabolites were moved to the cytosol. However, this step removed all reactions that summarized a transport reaction from the extracellular space to periplasm or from the cytosol to an unsupported organelle. The extracellular compartment corresponds to the parasitophorous vacuole space contained within the host cell for intracellular parasites (i.e. Plasmodium, Toxoplasma, Cryptosporidium) and the host serum for extracellular parasites (i.e. Trypanosoma).

**Manual curation**

We performed brief manual curation from literature sources, building on our curation conducted in [47,57]. Manual curation included updating reaction identifier and notes fields for consistency to CobraPY standards, updating metabolite identifiers for consistency to BiGG standards, replacing old PlasmoDB gene identifiers with updated identifiers, mass balancing seven reactions (with phosphate or hydrogen), removing duplicate reactions and metabolites (12 modifications), pruning unused metabolites, ensuring all metabolites have associated compartments, adding annotations to reactions and metabolites if missing (i.e. KEGG identifiers, EC codes, InChI strings), and adding SBO terms to all objects. Our previous asexual blood-stage Plasmodium falciparum 3D7 reconstruction ([57] adapted from [47]) was manually curated generating iPfal22. See Data Availability for code documenting all for modifications and implementation of manual curation.

Additional manual curation was performed on lipid metabolism of the asexual blood-stage Pl. falciparum using the lipidomics study presented in [83] adding over 700 reactions, 400 metabolites, and 18 genes. This curation removed aggregate reactions representing lipid
metabolism and replaced them with individual reactions for individual lipid species, as supported by the lipidomics study. This model is available, but was not used for the analyses presented here as the metabolic demands for lipids in our biomass reaction are also aggregated. Inclusion of these reactions is appropriate for understanding lipid metabolism but would create random distributions of flux through the individual reactions that may distract from meaningful changes in flux distributions. All code for this curation is available at https://github.com/gulermalaria/iPfal17_curation.

Automated orthology-driven curation

We developed a novel automated curation approach using orthologous transformation, similar to the approach taken by [48]. Our approach leverages the curation conducted in one organism for closely-related organisms and applied it to all draft Plasmodium reconstructions using iPfal22 (Fig 2A). We first mapped orthology of P. falciparum to each other Plasmodium species to build an orthology thesaurus (Fig 2A). We then added genes and associated reactions from iPfal22 if there was an orthologous gene in the target species’ reconstruction and the reaction was not already present (Fig 2A) resulting in a significant increase in the number of genes and reactions in each reconstruction (Fig 2C). Notably, this approach facilitates the compartmentalization of these reconstructions, a function most automated pipelines except RAVEN [45] and merlin [46] fail to include. This step is particularly important for parasite-specific compartments such as the apicoplast, which is not included in any database.

Gapfilling (part 1)—Automated data-driven curation

Gapfilling is an analytic process used to bridge or complete genetically-supported metabolic pathways to permit the network to fulfill metabolic functions, and was used to generate functional models. To increase the scope of a reconstruction (i.e. to add reactions), we performed gapfilling to fill in gaps in pathways to ensure that the reconstruction can complete a particular task. This optimization process adds reactions to allow the reconstruction to carry flux under given constraints.

Thus, automated curation of all reconstructions was performed by gapfilling for metabolites measured to be consumed in fluxomic or select media formulation studies. Detailed analysis is provided in our code. In brief, following an extensive literature review, we compiled data providing evidence for consumption or production of select metabolites (S1 Table). Metabolites were defined as consumed by the parasite if: (1) the metabolite was radiolabeled, added to media, incorporated into the parasite or converted by the parasite, and this was not seen to the same degree in uninfected host cells; (2) the metabolite rescued inhibitor treatment of a metabolically upstream parasite enzyme; or (3) the metabolite is an essential media component for parasite culture. Metabolites were defined to be produced by the parasite if the metabolite was radiolabeled following growth in a media containing a radiolabeled precursor metabolite, and this was not seen to the same degree in uninfected host cells.

To gapfill for these tasks, import or excretion of these metabolites were added to the reconstruction. Next, the model objective was changed to an internal demand reaction for the metabolite or excretion reactions, respectively, and was gapfilled sequentially; this ensures import or synthesis of each of these measured metabolites. We then used a parsimonious flux balance analysis (pFBA)-based approach as originally used in [60] and further developed in [61] to add the minimal number of reactions required to carry flux through the internal demand or excretion reaction. Code is linked in the Data Availability section. In essence, this pFBA approach minimizes the flux through genetically unsupported reactions from a biochemical database such that the network can carry flux through the objective reaction (i.e. metabolite production or consumption or biomass synthesis). Any reaction from the database that carries flux during this problem was added to the network.

Gapfilling (part 2)—Biomass as the objective function
After compartmentalization, manual curation, and gapfilling for individual metabolites, we used pFBA-based gapfilling to ensure each network was capable of generating biomass. We use two classes of biomass functions here to robustly evaluate model performance, specifically a genus-specific curated biomass reaction (for *Plasmodium* reconstructions) and a parasite-specific generic biomass reaction. The genus-level curated biomass reaction was developed and curated for our manually curated *Plasmodium falciparum* model; see [47] for justification of this biomass reaction. Our generic biomass contains the consensus set of metabolites from several manually curated reconstructions for *Plasmodium falciparum, Leishmania major,* and *Cryptosporidium hominis.* The stoichiometry for this generic biomass is from the *iPfal22* biomass reaction [47]. Unfortunately, variability in reconstruction namespace (i.e. the database used for metabolite and reaction naming conventions) make it difficult to reconcile data compiled for some parasite reconstructions, such as the *Toxoplama* and *Plasmodium* reconstructions *ToxoNet1* in the KEGG namespace and *iPfal22* in the BiGG namespace, as there are not always one-to-one mappings of variables across databases. This generic biomass was used to capture the most conservatively defined required parasite biosynthetic capacity. All reconstructions were gapfilled to ensure biomass could be synthesized via the generic parasite biomass reaction; all *Plasmodium* reconstructions were also gapfilled to ensure biomass could be synthesized via the genus-specific biomass reaction. For all biomass gapfilling steps and simulations, all exchange reactions were kept open to simulate a nutrient rich extraparasitic environment. Blocked and unconnected reactions were kept in the reconstructions (S6 Table). The distribution of blocked and unconnected reactions were similar in *de novo* and gapfilled reconstructions (S9 Fig).

**Model performance evaluation**

Network accuracy was evaluated against gene essentiality data (S4 Table) if available. Gene essentiality was simulated by performing single gene deletion studies in our models. All simulations were performed in a model state without additional experimental constraints; specifically, all exchange reactions were permitted to carry flux reversibly, simulating a nutrient rich environment such as intracellular growth. Of note, these ‘unconstrained’ models do not necessarily represent the *in vitro* or *in vivo* environment in which all experiments were conducted, merely the metabolic capacity an organism encodes. If this extracellular environment was constrained to represent the specific host environment, we would see further separation of model predictions.

Gene deletions were simulated by removing the gene of interest from the model using CobraPy’s ‘single_gene_deletion’ function. This change results in the inhibition of flux through all reactions that require that gene to function. If the model could not produce biomass with these constraints, the gene was deemed essential. Specifically, we defined an essential gene as a knockout that resulted in 10% or less of the maximal biomass (measured in mmol/(gDW•hr)). Knockout accuracy was defined as the sum of true positives (refractory to knockout or lethal genes) and true negatives (nonlethal genes) divided by the total number of predictions. As targeted metabolomics data were used for model generation (by gapfilling), we excluded these data from the evaluation data.

Models were tested for thermodynamically-infeasible loops and energy-generating cycles; the approach outlined in [84] was used with minor modifications for eukaryotic cells.

**Memote evaluation**

Models were evaluated using Memote [62]; example outputs are presented on [https://github.com/maureencarey/paradigm/tree/master/memote_reports](https://github.com/maureencarey/paradigm/tree/master/memote_reports). Additionally, Memote was used to quality control the reconstructions throughout the development of the ParaDIGM pipeline to improve standard compliance (especially annotation coverage) and biological relevance (e.g. network connectivity and topology).
Reconstruction pipeline for eukaryotes

Our pipeline itself (Fig 2A) is uniquely tailored to eukaryotic cells by recognizing the importance of compartmentalization and the design of the objective function. Compartmentalization biases predictions [47]; both our de novo reconstruction and orthology-driven approaches addresses this important step. Compartmentalization was incorporated into our de novo reconstruction pipeline and implemented for several genera (S5 Table). We adjusted the localization of reactions if inappropriately compartmentalized and added unique parasite-specific compartments (e.g. the apicoplast in Plasmodium). This addition was done by adding genetically supported reactions to all feasible compartments. If a gene-encoded enzyme corresponds to both mitochondrial and cytoplasmic reactions, both versions will be included: adding network redundancy that may not be biologically accurate. Alternatively, if an enzyme maps to a chloroplast reaction, our approach moved the reaction to the cytosol. It is plausible that chloroplast reactions like this example are not catalyzed by the parasite. These modifications are encoded in our analytic pipeline for future reference (see code, Data Availability).

Additionally, we used a curated model to inform the compartmentalization of each semi-curated model (Fig 2A); genes associated with compartmentalized reactions were mapped via orthology, assuming orthologous genes has comparable localization across species. Similarly, the objective function (often representing biomass synthesis [85]) influences network simulations [86] and the assumptions used in formulating a biomass reaction for prokaryotes may not apply to eukaryotes [87]. For example, in the first genome-scale metabolic model of any Cryptosporidium species, C. hominis [88], 30 reactions involved in lipid synthesis were unsupported by genetic evidence and manually added to the network. The selection of lipid species as biomass precursors may have impacted the addition of these unsupported reactions and resultant simulations with the model; for example, the 30 gapfilled reactions might not be added if alternative or fewer lipid species were included in the biomass reaction.

Thus, we use a consensus biomass reaction derived from multiple curated parasite reconstructions, as well as a genus-specific biomass for Plasmodium reconstructions. These objective functions influence reactions added via gapfilling, adding uncertainty in network structure [60,61]; thus, we emphasize which reactions were gapfilled frequently (i.e. for many species within a genera or ParaDIGM broadly) to increase our confidence in these predicted functions. Targeted investigation of these reactions (such as pyridoxal oxidase and shikimate dehydrogenase) will increase our confidence in all of our parasite network reconstructions.

Furthermore, initial de novo reconstructions were supplemented with curated information from the EuPathDB databases. All genes associated with KEGG reaction identifiers were collected from EuPathDB. These KEGG identifiers were then converted to the BiGG namespace, whenever possible, and added these reactions to each genome reconstruction. If the reaction was already present but the gene was not, the gene was added to the reaction’s gene-protein-reaction rule (using an ‘OR’ relationship if another gene already was included). This process improved the scope of the network reconstruction for each species/genome and leverages one of the best features of the EuPathDB family of databases: user contributions and literature support. This builds on the bioinformatic approach (i.e. Diamond annotation) to build de novo reconstructions by integrating curated gene annotations that have been documented on EuPathDB.

Even with this addition, it is important to note that not every gene with a metabolic annotation on EuPathDB will be included in the reconstructions for three main reasons: (1) some metabolic processes are not included in genome-scale metabolic network reconstructions (group 1), (2) some functions are insufficiently characterized to include as a reaction (group 2), and (3) unfortunately not all KEGG IDs can be converted to the BiGG namespace (group...
3). For example, 1032 genes are on EuPathDB for *T. gondii* ME49, associated with a metabolic GO term, and not found via our Diamond annotation of metabolic genes. This list of genes includes TGME49_201130, TGME49_321660, and TGME49_326200. TGME49_201130 is annotated as a rhoptry kinase family protein ROP33a with GO terms including ‘ATP binding’ and ‘protein kinase activity’; rhoptry biology is not traditionally the kind of function that would be included in genome-scale metabolic network reconstructions because it is not a biosynthesis or degradation. Thus, this gene is associated with group 1 above. On the other hand, it may be appropriate to include the function associated with TGME49_321660 in a reconstruction as its associated GO terms include ‘transferase activity’ and ‘transferring glycosyl groups’. However, the annotation for TGME49_321660 is not detailed enough to include in a reconstruction (i.e. there is no EC number or KEGG ID), so this gene belongs in group 2. Lastly, TGME49_326200 encodes a type I fatty acid synthase (EC 1.6.5.5) is associated with the KEGG ID (R02364) but no BiGG ID (group 3). The groups 2 and 3 (2: incompletely annotated genes and 3: those associated with only KEGG IDs) are excellent starting points for manual curation.

In addition, to curate these reconstructions, we recommend readers focus on false positives, or enzymic functions that were identified but do not occur *in vivo* or *in vitro*. For example, when the model incorrectly predicts a function or gene is not essential, there is inaccurate redundancy upstream of the prediction; thus, a curator can use these observations to identify and remove false positive gene annotations.

**Author Contributions**

**Conceptualization:** Maureen A. Carey, Gregory L. Medlock, Jason A. Papin.

**Data curation:** Maureen A. Carey, Michał Stolarczyk.

**Formal analysis:** Maureen A. Carey.

**Funding acquisition:** Maureen A. Carey, Gregory L. Medlock, William A. Petri, Jr., Jennifer L. Guler, Jason A. Papin.

**Investigation:** Maureen A. Carey, Gregory L. Medlock.

**Methodology:** Maureen A. Carey.

**Project administration:** William A. Petri, Jr., Jason A. Papin.

**Software:** Maureen A. Carey, Gregory L. Medlock.

**Supervision:** Jennifer L. Guler, Jason A. Papin.

**Validation:** Maureen A. Carey.

**Visualization:** Maureen A. Carey.

**Writing – original draft:** Maureen A. Carey.

**Writing – review & editing:** Maureen A. Carey, Gregory L. Medlock, Michał Stolarczyk, William A. Petri, Jr., Jennifer L. Guler, Jason A. Papin.

**References**

1. World Health Organization. World Malaria Report 2018. World Health Organization; 2019.

2. World Health Organization. Research priorities for Chagas disease, human African trypanosomiasis and leishmaniasis. World Health Organ Tech Rep Ser. 2012; v–xii, 1–100. PMID: 23484340

3. World Health Organization. Weekly epidemiological record Relevé épidémiologique hebdomadaire. Weekly Epidemiological Report. 2018; 20.
4. Elsheikha HM, Khan NA. Essentials of Veterinary Parasitology. Horizon Scientific Press; 2011.

5. Day MJ. One health: the importance of companion animal vector-borne diseases. Parasit Vectors. 2011; 4: 49. https://doi.org/10.1186/1756-3305-4-49 PMID: 21489237

6. May RM. Parasites, people and policy: infectious diseases and the Millennium Development Goals. Trends Ecol Evol. 2007; 22: 497–503. https://doi.org/10.1016/j.tree.2007.08.009 PMID: 17904247

7. McCoy D, Kembhavi A, Patel J, Luintel A. The Bill & Melinda Gates Foundation’s grant-making programme for global health. Lancet. 2009; 373: 1645–1653. https://doi.org/10.1016/S0140-6736(09)60571-7 PMID: 19427959

8. Ghorbani M, Farhoud R. Leishmaniasis in humans: drug or vaccine therapy? Drug Des Devel Ther. 2018; 12: 25–40. https://doi.org/10.2147/DDDT.S146521 PMID: 29317800

9. Sacks DL. Vaccines against tropical parasitic diseases: a persisting answer to a persisting problem. Nat Immunol. 2014; 15: 403–405. https://doi.org/10.1038/ni.2853 PMID: 24747701

10. Dumonteil E, Herrera C, Buekens P. A therapeutic preconceptional vaccine against Chagas disease: A novel indication that could reduce congenital transmission and accelerate vaccine development. PLoS Negl Trop Dis. 2019; 13: e0006985. https://doi.org/10.1371/journal.pntd.0006985 PMID: 30703092

11. Pance A. How elusive can a malaria vaccine be? Nat Rev Microbiol. 2019; 17: 129. https://doi.org/10.1038/s41579-018-0148-3 PMID: 30626910

12. Checkley W, White AC Jr, Jaganath D, Arrowood MJ, Chalmers RM, Chen X-M, et al. A review of the global burden, novel diagnostics, therapeutics, and vaccine targets for cryptosporidium. Lancet Infect Dis. 2015; 15: 85–94. https://doi.org/10.1016/S1473-3099(14)7772-6 PMID: 25278220

13. Sparks H, Nair G, Castellanos-Gonzalez A, White AC Jr. Treatment of Cryptosporidium: What We Know, Gaps, and the Way Forward. Curr Trop Med Rep. 2015; 2: 181–187. https://doi.org/10.1007/s40475-015-0056-9 PMID: 26568906

14. Menard D, Dondorp A. Antimalarial Drug Resistance: A Threat to Malaria Elimination. Cold Spring Harb Perspect Med. 2017; 7. https://doi.org/10.1101/cshperspect.a025619 PMID: 28289248

15. Delves M, Plouffe D, Scheurer C, Meister S, Wittlin S, Winzeler EA, et al. The activities of current antimalarial drugs on the life cycle stages of Plasmodium: a comparative study with human and rodent parasites. PLoS Med. 2012; 9: e1001169. https://doi.org/10.1371/journal.pmed.1001169 PMID: 22363211

16. Simpkin VL, Renwick MJ, Kelly R, Mossialos E. Incentivising innovation in antibiotic drug discovery and development: progress, challenges and next steps. J Antibiot. 2017; 70: 1087–1096. https://doi.org/10.1038/ja.2017.124 PMID: 29089600

17. DiMasi JA, Hansen RW, Grabowski HG. The price of innovation: new estimates of drug development costs. J Health Econ. 2003; 22: 151–185. https://doi.org/10.1016/S0167-6296(02)00126-1 PMID: 12606142

18. Haanstra JR, Gerdin A, Dolga AM, Sorgdrager FJH, Buist-Homan M, du Toit F, et al. Targeting pathogen metabolism without collateral damage to the host. Sci Rep. 2017; 7: 40406. https://doi.org/10.1038/srep40406 PMID: 28084422

19. Bogdan C, Stenger S, Rollenhoff M, Solbach W. Cytokine interactions in experimental cutaneous leishmaniasis. Interleukin 4 synergizes with interferon-γ to activate murine macrophages for killing of Leishmania major amastigotes. Eur J Immunol. 1991; 21: 327–333. https://doi.org/10.1002/eji.1830210213 PMID: 1900240

20. Kumaratiilake LM, Ferrante A, Robinson BS, Jaeger T, Poulos A. Enhancement of neutrophil-mediated killing of Plasmodium falciparum asexual blood forms by fatty acids: importance of fatty acid structure. Infect Immun. 1997; 65: 4152–4157. https://doi.org/10.1128/iai.65.10.4152-4157.1997 PMID: 9317021

21. Zheng W. Sirtuins as emerging anti-parasitic targets. Eur J Med Chem. 2013; 59: 132–140. https://doi.org/10.1016/j.ejmech.2012.11.014 PMID: 23206411

22. Dahl EL, Shock JL, Shenai BR, Gut J, DeRisi JL, Rosenthal PJ. Tetracyclines specifically target the apicoplast of the malaria parasite Plasmodium falciparum. Antimicrob Agents Chemother. 2006; 50: 3124–3131. https://doi.org/10.1128/AAC.00394-06 PMID: 16940111

23. Ghorbal M, Gorman M, Macpherson CR, Martins RM, Scherf A, Lopez-Rubio J-J. Genome editing in the human malaria parasite Plasmodium falciparum using the CRISPR-Cas9 system. Nat Biotechnol. 2014; 32: 819–821. https://doi.org/10.1038/nbt.2925 PMID: 24880488

24. Lee MC, Fidock DA. CRISPR-mediated genome editing of Plasmodium falciparum malaria parasites. Genome Med. 2014; 6: 63. https://doi.org/10.1186/s13073-014-0063-9 PMID: 25473431

25. Downing T, Imamura H, Decuyper S, Clark TG, Coombs GH, Cotton JA, et al. Whole genome sequencing of multiple Leishmania donovani clinical isolates provides insights into population structure and mechanisms of drug resistance. Genome Res. 2011; 21: 2143–2156. https://doi.org/10.1101/gr.123430.111 PMID: 22038251
26. Sterkers Y, Lachaud L, Bourgeois N, Crobu L, Bastien P, Pagès M. Novel insights into genome plasticity in Eukaryotes: mosaic aneuploidy in Leishmania. Mol Microbiol. 2012; 86: 15–23. https://doi.org/10.1111/j.1365-2958.2012.08185.x PMID: 22857263

27. Jumani RS, Bessoff K, Love MS, Miller P, Stebbins EE, Teixeira JE, et al. A Novel Piperazine-Based Drug Lead for Cryptosporidiosis from the Medicines for Malaria Venture Open-Access Malaria Box. Antimicrob Agents Chemother. 2018; 62. https://doi.org/10.1128/AAC.01505-17 PMID: 29339392

28. Chao AT, Lee BH, Wan KF, Selva J, Zou B, Gedeck P, et al. Development of a Cytopathic Effect-Based Phenotypic Screening Assay against Cryptosporidium. ACS Infect Dis. 2018; 4: 635–645. https://doi.org/10.1021/acsinfecdis.7b00247 PMID: 29341586

29. Love MS, Beasley FC, Jumani RS, Wright TM, Chatterjee AK, Huston CD, et al. A high-throughput phenotypic screen identifies clofazimine as a potential treatment for cryptosporidiosis. PLoS Negl Trop Dis. 2017; 11: e0005373. https://doi.org/10.1371/journal.pntd.0005373 PMID: 28158186

30. Subramanian G, Belekar MA, Shukla A, Tong JX, Sinha A, Chu TTT, et al. Targeted Phenotypic Screening in Plasmodium falciparum and Toxoplasma gondii Reveals Novel Modes of Action of Medicines for Malaria Venture Malaria Box Molecules. mSphere. 2018; 3. https://doi.org/10.1128/mSphere.00534-17 PMID: 29359192

31. Lucantoni L, Duffy S, Adjalley SH, Fidock DA, Avery VM. Identification of MMV malaria box inhibitors of plasmodium falciparum early-stage gametocytes using a luciferase-based high-throughput assay. Antimicrob Agents Chemother. 2013; 57: 6050–6062. https://doi.org/10.1128/AAC.00670-13 PMID: 24060871

32. Antonova-Koch Y, Meister S, Abraham M, Luth MR, Ottilie S, Lukens AK, et al. Open-source discovery of chemical leads for next-generation chemoprotective antimalarials. Science. 2018;362. https://doi.org/10.1126/science.aat9446 PMID: 30523084

33. Li F-J, He CY. Autophagy in protozoan parasites: Trypanosoma brucei as a model. Future Microbiol. 2017; 12: 1337–1340. https://doi.org/10.2217/fmb-2017-0158 PMID: 28972410

34. Meyer KJ, Caton E, Shapiro TA. Model System Identifies Kinetic Driver of Hsp90 Inhibitor Activity against African Trypanosomes and Plasmodium falciparum. Antimicrob Agents Chemother. 2018;62. https://doi.org/10.1128/AAC.00056-18 PMID: 29866861

35. Sidik SM, Huet D, Ganesan SM, Huynh M-H, Wang T, Nasamu AS, et al. A Genome-wide CRISPR Screen in Toxoplasma Identifies Essential Apicomplexan Genes. Cell. 2016; 166: 1423–1435.e12. https://doi.org/10.1016/j.cell.2016.08.019 PMID: 27594426

36. Kim K, Weiss LM. Toxoplasma gondii: the model apicomplexan. Int J Parasitol. 2004; 34; 423–432. https://doi.org/10.1016/j.ijpara.2003.12.009 PMID: 15003501

37. Huang BW, Pearman E, Kim CC. Mouse Models of Uncomplicated and Fatal Malaria. Bio Protoc. 2015; 5. Available: https://www.ncbi.nlm.nih.gov/pubmed/26236758 https://doi.org/10.1126/science.aat9446 PMID: 30523084

38. Huang BW, Pearman E, Kim CC. Mouse Models of Uncomplicated and Fatal Malaria. Bio Protoc. 2015; 5. Available: https://www.ncbi.nlm.nih.gov/pubmed/26236758 https://doi.org/10.1126/science.aat9446 PMID: 30523084

39. Galinski MR. Functional genomics of simian malaria parasites and host-parasite interactions. Brief Funct Genomics. 2019; 18: 270–280. https://doi.org/10.1093/bfgp/elz013 PMID: 32141151

40. Carlton JM, Adams JH, Silva JC, Bidwell SL, Lorenzi H, Caler E, et al. Comparative genomics of the neglected human malaria parasite Plasmodium vivax. Nature. 2008; 455: 757–763. https://doi.org/10.1038/nature07327 PMID: 18843361

41. El-Sayed NM, Myler PJ, Blandin G, Berriman M, Crabtree J, Aggarwal G, et al. Comparative genomics of trypanosomatid parasitic protozoa. Science. 2005; 309: 404–409. https://doi.org/10.1126/science.1112181 PMID: 16020724

42. Reid AJ, Vermont SJ, Cotton JA, Harris D, Hill-Cawthorne GA, Koen-Waisman S, et al. Comparative genomics of the apicomplexan parasites Toxoplasma gondii and Neospora caninum: Coccidia differing in host range and transmission strategy. PLoS Pathog. 2012; 8: e1002567. https://doi.org/10.1371/journal.ppat.1002567 PMID: 22457617

43. Agren R, Liu L, Shoae S, Vongsangnak W, Nookaew I, Nielsen J. The RAVEN toolbox and its use for generating a genome-scale metabolic model for Penicillium chrysogenum. PLoS Comput Biol. 2013; 9: e1002980. https://doi.org/10.1371/journal.pcbi.1002980 PMID: 23555215
46. Dias O, Rocha M, Ferreira EC, Rocha I. Reconstructing genome-scale metabolic models with merlin. Nucleic Acids Res. 2015; 43: 3899–3910. https://doi.org/10.1093/nar/gkv294 PMID: 25845595

47. Carey MA, Papin JA, Guler JL. Novel Plasmodium falciparum metabolic network reconstruction identifies shifts associated with clinical antimalarial resistance. BMC Genomics. 2017; 18: 543. https://doi.org/10.1186/s12864-017-3905-1 PMID: 28724354

48. Abdel-Haleem AM, Hefzi H, Mineta K, Gao X, Gojobori T, Palsson BO, et al. Functional interrogation of Plasmodium genus metabolism identifies species- and stage-specific differences in nutrient essentiality and drug targeting. PLoS Comput Biol. 2018; 14: e1005895. https://doi.org/10.1371/journal.pcbi.1005895 PMID: 29300748

49. Chiappin o-Pepe A, Tymoshenko S, Ataman M, Soldati-Favre D, Hatzimanikatis V. Bioenergetics-based modeling of Plasmodium falciparum metabolism reveals its essential genes, nutritional requirements, and thermodynamic bottlenecks. PLoS Comput Biol. 2017; 13: e1005397. https://doi.org/10.1371/journal.pcbi.1005397 PMID: 28333921

50. Tymoshenko S, Oppenheim RD, Agren R, Nielsen J, Soldati-Favre D, Hatzimanikatis V. Metabolic Needs and Capabilities of Toxoplasma gondii through Combined Computational and Experimental Analysis. PLoS Comput Biol. 2015; 11: e1004261. https://doi.org/10.1371/journal.pcbi.1004261 PMID: 26001086

51. Caspi R, Billington R, Fulcher CA, Keseler IM, Kothari A, Krummenacker M, et al. The MetaCyc database of metabolic pathways and enzymes. Nucleic Acids Res. 2018; 46: D633–D639. https://doi.org/10.1093/nar/gkx935 PMID: 29059334

52. Machado D, Andrejev S, Tramontano M, Patil KR. Fast automated reconstruction of genome-scale metabolic models for microbial species and communities. Nucleic Acids Res. 2018; 46: 7542–7553. https://doi.org/10.1093/nar/gky337 PMID: 30192979

53. Seaver SMD, Liu F, Zhang Q, Jeffryes J, Faria JP, Edirisinghe JN, et al. The ModelSEED Biochemistry Database for the integration of metabolic annotations and the reconstruction, comparison and analysis of metabolic models for plants, fungi and microbes. Nucleic Acids Res. 2021; 49: D575–D588. https://doi.org/10.1093/nar/gkaa746 PMID: 32996334

54. Amos B, Aurrecoechea C, Barba M, Barreto A, Basenko EY, Bažant W, et al. VEuPathDB: the eukaryotic pathogen, vector and host bioinformatics resource center. Nucleic Acids Res. 2022; 50: D98–D111. https://doi.org/10.1093/nar/gkaa929 PMID: 34718728

55. Ebrahim A, Lerman JA, Palsson BO, Hyduke DR. COBRApy: COnstrain ts-Based Reconstruction and Analysis for Python. BMC Syst Biol. 2013; 7: 74. https://doi.org/10.1186/1752-0509-7-74 PMID: 23927696

56. King ZA, Lu J, Draeger A, Miller P, Federowicz S, Lerman JA, et al. BiGG Models: A platform for integrating, standardizing and sharing genome-scale models. Nucleic Acids Res. 2016; 44: D515–D22. https://doi.org/10.1093/nar/gkv1049 PMID: 26476456

57. Untaroiu AM, Carey MA, Guler JL, Papin JA. Leveraging the effects of chloroquine on resistant malaria parasites for combination therapies. BMC Bioinformatics. 2019; 20: 186. https://doi.org/10.1186/s12859-019-2756-y PMID: 30987583

58. Stanway RR, Bushell E, Chiappino-Pepe A, Roques M, Sanderson T, Franke-Fayard B, et al. Genome-Scale Identification of Essential Metabolic Processes for Targeting the Plasmodium Liver Stage. Cell. 2019; 179: 1112–1126. https://doi.org/10.1016/j.cell.2019.10.030 PMID: 31730853

59. Krishnan A, Kloehn J, Lunghi M, Chiappino-Pepe A, Waldman BS, Nicolas D, et al. Functional and Computational Genomics Reveal Unprecedented Flexibility in Stage-Specific Toxoplasma Metabolism. Cell Host Microbe. 2020; 27: 290–306.e11. https://doi.org/10.1016/j.chom.2020.01.002 PMID: 31991093

60. Biggs MB, Papin JA. Managing uncertainty in metabolic network structure and improving predictions using EnsembleFBA. PLoS Comput Biol. 2017; 13: e1005413. https://doi.org/10.1371/journal.pcbi.1005413 PMID: 28263984

61. Medlock GL, Papin JA. Guiding the Refinement of Biochemical Knowledgebases with Ensembles of Metabolic Networks and Machine Learning. Cell Systems. 2020. https://doi.org/10.1016/j.cels.2019.11.006 PMID: 31926940

62. Lieven C, Beber ME, Olivier BG, Bergmann FT, Ataman M, Babaei P, et al. MEMOTE for standardized genome-scale metabolic model testing. Nat Biotechnol. 2020; 38: 272–276. https://doi.org/10.1038/s41587-020-0446-y PMID: 32123384

63. Carey MA, Dräger A, Beber ME, Papin JA, Yurkovich JT. Community standards to facilitate development and address challenges in metabolic modeling. Mol Syst Biol. 2020; 16: e8235. https://doi.org/10.1525/msb.20199235 PMID: 32845080
64. Zhang M, Wang C, Otto TD, Oberstaller J, Liao X, Adapa SR, et al. Uncovering the essential genes of the human malaria parasite Plasmodium falciparum by saturation mutagenesis. Science. 2018;360. https://doi.org/10.1126/science.aap7847 PMID: 29724925

65. Bushell E, Gomes AR, Sanderson T, Anar B, Girling G, Herd C, et al. Functional Profiling of a Plasmodium Genome Reveals an Abundance of Essential Genes. Cell. 2017; 170: 260–272.e8. https://doi.org/10.1016/j.cell.2017.06.030 PMID: 28708996

66. Berk AJ. Discovery of RNA splicing and genes in pieces. Proc Natl Acad Sci U S A. 2016; 113: 801. https://doi.org/10.1073/pnas.1525084113 PMID: 26787897

67. Baym M, Lieberman TD, Kelsic ED, Chait R, Gross R, Yelin I, et al. Spatiotemporal microbial evolution on antibiotic landscapes. Science. 2016; 353: 1147–1151. https://doi.org/10.1126/science.aag0822 PMID: 27609891

68. R Core Team. R: A Language and Environment for Statistical Computing. Vienna, Austria: R Foundation for Statistical Computing; 2017. Available: https://www.R-project.org/

69. Wickham H. Tidyverse: Easily install and load‘tidyverse’packages. 2017.

70. Kassambara A. ggpubr:‘ggplot2’ based publication ready plots. R package version 0.1. 2017; 6.

71. de Vries A, Ripley BD. Ggdendro: tools for extracting dendrogram and tree diagram plot data for use with ggplot. 2013.

72. Charif D, Lobry JR. SeqinR 1.0–2: A Contributed Package to the R Project for Statistical Computing Devoted to Biological Sequences Retrieval and Analysis. In: Bastolla U, Porto M, Roman HE, Vendruscolo M, editors. Structural Approaches to Sequence Evolution: Molecules, Networks, Populations. Berlin, Heidelberg: Springer Berlin Heidelberg; 2007. pp. 207–232.

73. Pagès H, Aboyoun P, Gentleman R, DebRoy S. Biostings: Efficient manipulation of biological strings. 2017.

74. Bodenhofer U, Bonatesta E, Horejš-Kainrath C, Hochreiter S. msa: an R package for multiple sequence alignment. Bioinformatics. 2015; 31: 3997–3999. https://doi.org/10.1093/bioinformatics/btv494 PMID: 26315911

75. Wickham H. reshape2: Flexible reshape data: a reboot of the reshape package. R package version. 2012; 1. Available: http://cran.ms.unimelb.edu.au/web/packages/reshape2/

76. Gehlenborg N. UpSetR: A More Scalable Alternative to Venn and Euler Diagrams for Visualizing Intersecting Sets. 2017. Available: https://CRAN.R-project.org/package=UpSetR

77. Dray S, Dufour A-B, Others. The ade4 package: implementing the duality diagram for ecologists. J Stat Softw. 2007; 22: 1–20.

78. Neuwirth E, Neuwirth ME. The RColorBrewer Package. 2007. Available: http://ftp.uni-bayreuth.de/math/statlib/R/CRAN/packages/RColorBrewer.pdf

79. Wickham H, Bryan J. readxl: Read Excel Files. R package version 1.0. 0. URL https://CRAN.R-project.org/package=readxl; 2017.

80. Feist AM, Palsson BO. The biomass objective function. Curr Opin Microbiol. 2010; 13: 344–349. https://doi.org/10.1016/j.mib.2010.03.003 PMID: 20430689

81. Xavier JC, Patil KR, Rocha I. Integration of Biomass Formulations of Genome-Scale Metabolic Models with Experimental Data Reveals Universally Essential Cofactors in Prokaryotes. Metab Eng. 2017; 39: 200–208. https://doi.org/10.1016/j.ymben.2016.12.002 PMID: 27939572

82. Liebermeister W, Noer E, Flamholz A, Davidi D, Bernhardt J, Milo R. Visual account of protein investment in cellular functions. Proc Natl Acad Sci U S A. 2014; 111: 8486–8493. https://doi.org/10.1073/pnas.1314810111 PMID: 24889604
88. Vanee N, Roberts SB, Fong SS, Manque P, Buck GA. A Genome-Scale Metabolic Model of Cryptosporidium hominis. Chem Biodivers. 2010; 7: 1026–1039. https://doi.org/10.1002/cbdv.200900323 PMID: 20491062