Abstract: Genome-scale metabolic models (GEMs) are essential tools for in silico phenotype prediction and strain optimisation. The most straightforward GEMs reconstruction approach uses published models as templates to generate the initial draft, requiring further curation. Such an approach is used by BiGG Integration Tool (BIT), available for merlin users. This tool uses models from BiGG Models database as templates for the draft models. Moreover, BIT allows the selection between different template combinations. The main objective of this study is to assess the draft models generated using this tool and compare them BIT, comparing these to CarveMe models, both of which use the BiGG database, and curated models. For this, three organisms were selected, namely Streptococcus thermophilus, Xylella fastidiosa and Mycobacterium tuberculosis. The models’ variability was assessed using reactions and genes’ metabolic functions. This study concluded that models generated with BIT for each organism were differentiated, despite sharing a significant portion of metabolic functions. Furthermore, the template seems to influence the content of the models, though to a lower extent. When comparing each draft with curated models, BIT had better performances than CarveMe in all metrics. Hence, BIT can be considered a fast and reliable alternative for draft reconstruction for bacteria models.

Keywords: BiGG Integration Tool; BiGG models; genome-scale metabolic models; merlin.

1 Introduction

The reconstruction of metabolic networks in Systems Biology has peaked at an unprecedented scale, with over 6000 Genome-Scale Metabolic Models (GEMs) representing the metabolism of 434, 40 and 117 taxonomic families of bacteria, archaea and eukarya, respectively [1]. GEMs can predict the metabolic behaviour of an organism in several genetic and environmental conditions using methods, such as Flux Balance Analysis (FBA) [2]. Therefore, GEMs are relevant computational biology tools due to their many metabolic engineering applications, such as in silico strain optimisation [3].

Despite the comprehensive collection of GEMs, only a small share of these organisms (113 bacteria, ten archaea, and 60 eukaryotes) have manually and high-quality GEM reconstructions [1]. As of 2010, the

*Corresponding author: Oscar Dias, Centre of Biological Engineering, University of Minho, 4710-057 Braga, Portugal; and LABBELS – Associate Laboratory, Braga, Guimarães, Portugal, E-mail: odias@ceb.uminho.pt. https://orcid.org/0000-0002-1765-7178

Alexandre Oliveira, Emanuel Cunha, Fernando Cruz, João Capela, João C. Sequeira, Marta Sampaio and Cláudia Sampaio, Centre of Biological Engineering, University of Minho, 4710-057 Braga, Portugal; and LABBELS – Associate Laboratory, Braga, Guimarães, Portugal. https://orcid.org/0000-0002-2255-8983 (A. Oliveira)
Biochemical, Genetic and Genomic (BioGG) Models database has collected a wide variety of these high-quality and manually curated GEMs [4]. Up to the present date, this knowledgebase integrates 108 GEM reconstructions from bacteria, archaea and eukarya organisms. Moreover, BioGG Models provides the standardisation and unification of the reactions and metabolites available in these GEMs using universal non-redundant identifiers called BioGG identifiers. Then, reactions, metabolites and genes are mapped to external databases such as Kyoto Encyclopedia of Genes and Genomes (KEGG) [5], BioCyc [6], National Center for Biotechnology Information (NCBI) protein [7], among others.

The reconstruction of GEMs can usually follow two diverse approaches: bottom-up [8] and top-down [9].

The fast and automated reconstruction of metabolic networks consists of reconstructing a universal simulation-ready model that can be carved into an organism-specific model by pruning metabolic reactions for which genomic evidence is missing. CarveMe computational tool is an example of this top-down approach, used to create microbial community models by merging fast and automated single-species GEMs [9].

On the other hand, the widely-used bottom-up approach comprehends four main steps for generating high-quality metabolic networks [8]. This approach includes the genome functional annotation, network refinement and gap-filling, conversion to a stoichiometric model, and validation with experimental data. Although the bottom-up paradigm provides a protocol for reconstructing high-quality and relevant GEMs, the process tends to be time-consuming. Thus, fast and automated reconstructions have gained more interest in recent years.

With this in mind, we propose a novel computational tool for the fast reconstruction of GEMs using the BioGG Models database. The BioGG Integration Tool (BIT) has been implemented in the Metabolic Models Reconstruction Using Genome-Scale Information (merlin) software [10, 11]. This tool only requires the organism’s genome sequence to perform homology searches against the BioGG Models database and assemble a draft metabolic network. Thus, one can easily generate draft metabolic networks using only the Graphical User Interface (GUI).

In this study, we have used BIT to create several draft metabolic networks for three bacteria, namely Streptococcus thermophilus, Xylella fastidiosa and Mycobacterium tuberculosis. Gram negative and Gram positive bacteria available in the BioGG database were selected as case studies for this work [12–14]. Then, the metabolic content of the draft models was compared with reference GEMs [15–17]. Moreover, CarveMe’s top-down approach [9] has also been used to generate draft GEMs that were equally assessed against the BIT and reference reconstructions.

## 2 Workflow

BIT is a fully automated tool to generate draft GEMs using the BioGG Models database. This computational tool has been implemented in merlin, which is a graphical and user-oriented solution for the reconstruction of GEMs [10, 11].

BIT takes the genome sequence of the organism of interest and parameters related to the BLAST homology search (scoring matrix, e-value, and bit score) and gene-protein-reaction (GPR) associations.

As shown in the first step of Figure 1, BIT performs bidirectional BLAST similarity searches between the organism’s genome sequences and protein sequences available at the BioGG Models database. As a result, reciprocal matches between the direct and reverse homology searches allow identifying homologous genes between the organism of interest and the BioGG Models database.

BIT performs the association between the homologous genes found in the database and corresponding BioGG reactions in the next step. As described in stage 2 of Figure 1, GPR associations are used to find the BioGG reactions to be included in the draft metabolic network.

For that, GPR associations are decomposed in a set of Boolean logic rules using the common operators “AND” and “OR”. Then, the integration of BioGG reactions in the draft reconstruction is based on the evaluation of the Boolean logic expression using the homologous genes and similarity searches. The assessment of each Boolean logic rule is performed according to the following principles:
Figure 1: BiGG integration tool workflow implemented in merlin. Users have access to BIT within merlin’s graphical interface. The genome sequence fasta file is submitted to the BIT webservice for an organism of interest. A bidirectional BLAST similarity search is performed against the protein sequences available at the BiGG models database (1). This process allows the identification of homologous genes in the database. Then, Gene-Protein-Reaction (GPR) associations are elaborated based on the homologous genes and reactions available in the BiGG models database (2). Identified reactions are further integrated and validated based on the associated metabolites, namely reactants and products (3). Finally, the model can be exported or integrated into merlin’s workspace.

(i) When the GPR association contains a single gene, the corresponding homologous gene must be present in the similarity searches to consider the BiGG reaction in the draft model.

(ii) When the GPR association only contains “OR” operators, a single homologous gene present in the similarity searches is enough to assemble a partial GPR association and consider the BiGG reaction into the draft model.

(iii) When the GPR association only contains “AND” operators, all homologous genes must be present in the similarity searches to assemble a complete GPR association and consider the BiGG reaction into the draft model.

(iv) When the GPR association contains both “OR” and “AND” operators, the Boolean logic rule is decomposed by the “OR” operator and evaluated as in the principle (ii). As demonstrated in the example of Figure 2, the GPR association of the PI45P3K_cho reaction available in the BiGG Models database can be simplified by the “OR” operators into a single Boolean logic expression.

(100689251 OR 100689252 OR 100750761 OR 100774067)

Figure 2: Decomposition of the GPR association assigned to the PI45P3K_cho reaction in BiGG models database.
Once all GPR associations have been evaluated using the homologous genes, all identified reactions are integrated and validated to avoid redundancy (stage 3 in Figure 1). As the BiGG Models database contains duplicated reactions due to the reactions’ reversibility, models’ compartmentalisation or compounds’ protonation state, BIT would automatically integrate all reactions in the draft metabolic network. Despite increasing the redundancy of reactions in the draft model, these cases are resolved by verifying that the relevant intervenient metabolites are the same in the redundant reactions.

After integrating the BiGG reactions, BIT assembles a draft metabolic network that can be exported as an SBML model [18]. Alternatively, the draft model can be integrated into the merlin workspace to be curated and refined.

3 Methods

3.1 Genomes’ Comparative Functional Analysis

The protein sequences of the organisms present in BiGG models and three case study bacteria were retrieved from NCBI’s RefSeq database (S. thermophilus – NZ_LR822015; X. fastidiosa – NC_004556.1; M. tuberculosis – NC_000962.3). Recogniser (version 1.4.6) [19] was run on the protein sequences to compare the functional similarity between the organisms with the following parameters: threads set to 14, maximum e-value set to 1e-10 and Clusters of Orthologous Genes (COG) [20] set as the reference database. Only COG identifiers belonging to the “METABOLISM” general functional category were selected for further analyses.

A binary matrix was generated to perform a Principal Components Analysis (PCA) of functional characterisations, with rows representing organisms, columns representing the COG identifiers, and 0 and 1 representing the absence or presence of COGs, respectively. Variables with low variance were removed, and standardisation was applied using the VarianceThreshold and StandardScaler methods of the scikit-learn package [21]. Plots were generated with the PCA method of the scikit-learn package, comparing functional characterisation at the Domain and Phylum levels (Figure 1).

3.2 Draft models

S. thermophilus, X. fastidiosa, and M. tuberculosis were the selected organisms for draft reconstruction with BIT and CarveMe [9]. For each organism, seven draft models were generated with BIT. Three different approaches were used, namely:

1. all models templates, which selects all the models from BiGG to build with available genome sequences;
2. selected models template, with models of the three metabolically closest organisms;
3. random models templates, in which five drafts were assembled, using three random models as templates in each model.

For estimating the closest organisms to S. thermophilus, X. fastidiosa, and M. tuberculosis, metabolic functions wise, results from Genomes’ Comparative Functional Analysis were used to create a distance matrix, where the presence/absence of COGs was quantified between all organisms and the top three selected.

Parameters for GEMs reconstruction with BIT were as follows: e-value, bit score and query coverage were set to 1e-20, 50 and 0.75, respectively, and BIT did not include BiGG reactions with missing or with incomplete GPRs. CarveMe was run with default parameters [9], without gapfilling.

3.3 Quantitative analysis of draft models

The diversity of reactions and functional annotation was analysed for the draft models. The COBRApy package [22] was used to read the draft models in SBML [18] format. The draft models were compared concerning their BiGG reaction and metabolite identifiers, and their COG annotations. To determine COG identifiers of each model, the RefSeq identifiers of the enzymes indicated in the GPRs associations of the model were retrieved, and cross-referenced with the annotation obtained in Section 3.1, in order to obtain the collection of COG identifiers of each model.

The first approach to compare reactions, metabolites and COG identifiers was Venn diagrams. This analysis aimed to assess how the different tools and templates affected the variability of the drafts GEM. The Venn diagrams were plotted using the matplotlib-venn package (available at https://github.com/konstantint/matplotlib-venn) for Python 3.8.

The draft models’ reactions and metabolites were further compared using PCA. A binary matrix was constructed, in which each row represented a draft model, and each column represented either a BiGG reaction identifiers, a BiGG metabolite identifiers or a COG identifiers, depending on the functional information being analysed. Matrix cells were filled with presence (1) or absence (0) of each identifier. Low-variance features (columns $s^2 < 2 = 0$) were removed using scikit-learn [21]. In addition, standardisation
was also applied to each binary matrix, using the standard score (z-score: \( Z = (x - \mu)/\sigma \)). A single PCA analysis was performed for each binary matrix using scikit-learn. Principal Components (PC) 1, and 2 were selected for each analysis. PCA scores were plotted and coloured according to the set of template models. Each dot was annotated with a model-specific identifier, using the first letter of the genus and the first three letters of the species’ specific name. As five random-template models were created using a different set of template models each time, these models are also numbered accordingly.

### 3.4 Model assessment

The draft models were assessed to published manually curated GEMs [15–17]. The comparison was applied to all models’ reactions, excluding exchange, sink, and demand reactions. A confusion matrix was generated for each model and used to calculate the evaluation metrics. Precision (\( P \)), recall (\( R \)), and F1 score (\( F1 \)) were used for this evaluation and calculated as demonstrated in the following equations:

\[
P = \frac{TP}{TP + FP} \tag{1}
\]

\[
R = \frac{TP}{TP + FN} \tag{2}
\]

\[
F1 = 2 \cdot \frac{P \cdot R}{P + R} \tag{3}
\]

where \( P \) is the precision, \( R \) is the recall, \( F1 \) is the F1 score, TP refers to the true positives, FP to the false positives, and FN to the false negatives. Reactions present in both the reference model and the generated drafts were deemed TP. Moreover, reactions present in the drafts but missing in the reference model were regarded as FP. Conversely, the reactions of the reference model that were not in the drafts were considered FN.

According to these metrics, models with high \( R \) have an increased number of reactions correctly included (TP) and a low number of reactions missing (FN); thus, the efforts of manually curating the model are mostly associated with removing eventual FPs and adding few missing reactions (FN). On the other hand, models with low \( R \) have a high number of missing reactions (FN), requiring a more time-consuming refinement focused on adding the absent information.

Models with high \( P \) have an increased number of reactions correctly included (TP) when assessed to the number of reactions incorrectly added (FP). Proportionally, models with low \( P \) have a high number of reactions incorrectly assigned compared with reactions correctly inserted. In this regard, low \( P \) models are more suitable than low \( R \) from a curator’s point of view, as it is less tiresome to remove additional information than thoroughly searching for the missing information.

The F1 score combines the precision and recall of the classifier into a single metric, allowing a more balanced and direct evaluation of the draft models.

### 4 Results and discussion

BIT was used to reconstruct seven draft metabolic models for each bacterium: \( M. \) tuberculosis, \( S. \) thermophilus and \( X. \) fastidiosa. The models were created using three different approaches, namely using all models, selected models and random models as templates. CarveMe [9] was also used to develop a draft model for each bacterium. Then, all models were analysed and assessed regarding the variability of reactions and compared with curated models.

#### 4.1 Genomes’ comparative functional analysis

The first step of this work was to analyse the metabolic variability of all species present in the BiGG database and \( S. \) thermophilus and \( X. \) fastidiosa, unavailable in BiGG. For this purpose, the recogniser tool [19] was used to retrieve the COG identifiers for all organisms and PCA was applied to check the variability of the COG-annotated metabolic functions. Figure 3 shows the contribution of principal components (PC) 1 and 2 to the variability of metabolic functions of all organisms. These components group BiGG organisms by phylum, meaning that organisms in the same phylum share more metabolic functions. In addition, Figure 3 shows an evident separation between eukaryotes and prokaryotes, which confirms the results published by the database authors in their latest publication [4]. Therefore, these results validate the use of metabolic COG identifiers to study the metabolic capacities of the organisms.
The metabolic functions of the three case study bacteria were quantitatively compared using the metabolic COG identifiers and a Venn diagram (Figure 4). Analysing Figure 4, 77% of all COGs identified for the three models are present in *M. tuberculosis*, of which 544 are unique for this organism, indicating that *M. tuberculosis* has more and more distinct metabolic functions. On the other hand, *S. thermophilus* and *X. fastidiosa* share most of their metabolic functions, having only 126 and 155 unique metabolic functions, respectively. Overall, the three bacteria only share 226 metabolic COGs, indicating that the respective models should reflect these metabolic differences. In addition, these microorganisms exhibit different features: *S. thermophilus* is a lactic...
acid Gram-positive bacterium; \textit{X. fastidiosa} is a plant-pathogen Gram-negative; and \textit{M. tuberculosis} is a well-studied bacterium and already has a GEM available on BiGG.

4.2 Model analysis

Draft metabolic models for the three case study bacteria were generated using BIT and CarveMe. The models used for the \textit{selected models’ template}, were determined through similarity of metabolic functions, as shown in SI Table 1. The details of the 24 models generated for this study are summarised in SI Table 2.

The generated drafts were then compared to assess the effect of the template on the content of the models. For that, the reaction content was analysed in a Venn diagram (Figure 5), representing the number of unique and shared reactions between the different models of each bacterium.

As expected, \textit{all models’ template} generated models with the highest number of reactions for all organisms. For \textit{M. tuberculosis}, the model includes 75.4\% of all reactions, of which 1093 are not shared with the models from other templates. This value is even greater for the other two bacteria: 1391 and 1580 unique reactions for \textit{S. thermophilus} and \textit{X. fastidiosa} models, respectively. Unsurprisingly, the \textit{all models’ template} comprises most reactions available in the other templates. This is not the case of \textit{M. tuberculosis}, as the BiGG database includes models for \textit{M. tuberculosis}, which were used to reconstruct the model with the \textit{all models’ template}, while in the other templates, homology searches may lead to different matches according to the models selected for the reconstruction. Therefore, these results suggest that the template used in BIT for reconstructing draft metabolic models impacts the reaction content of the generated models.

The draft models generated from BIT using the \textit{selected template} and the models reconstructed by CarveMe were jointly analysed to investigate the differences in the models obtained with these two approaches. For this purpose, a Venn diagram was created for each method (Figure 6), representing the number of unique and shared reactions between the models of the three bacteria.

Regarding the models generated by BIT with the \textit{selected models’ template}, 180 reactions are shared between the three models. Furthermore, the most distinct \textit{selected} model belongs to \textit{S. thermophilus}, with 556 unique reactions. In contrast, the model of \textit{X. fastidiosa} is the one that presents more overlap with the other models, sharing 351 reactions, which represents 60\% of its reaction content. However, the results do not agree with the inferences obtained from the metabolic annotation of the organisms (4.1), in which \textit{M. tuberculosis} was the one with more unique COG-annotated metabolic functions (Figure 3).

On the other hand, the three models generated by CarveMe share 384 reactions, representing a slightly higher overlap than the models from BIT. In this case, the model with more unique reactions (643 reactions)

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{venn_diagrams.png}
\caption{Venn diagrams for the reaction content of the different draft models for each organism. Draft models have been generated from BiGG using BIT with different approaches to building templates: \textit{all models as templates}, \textit{random models as templates} and \textit{selected models as templates}. Venn diagrams show the number of reactions shared between the different models for \textit{M. tuberculosis} (A), \textit{S. thermophilus} (B), and \textit{X. fastidiosa} (C).}
\end{figure}
belongsto X. fastidiosa, while the S. thermophilus’ model is the most indistinctive, sharing 72.6% of its reaction content (725 reactions). These results are different from those obtained with BIT and from the results of the metabolic annotation (4.1).

CarveMe generates simulation-ready models, thus adding more reactions, such as sink and demand, to fill the model gaps. In contrast, BIT models require curation and gap-filling to be ready for simulation [9]. Therefore, although both approaches use BiGG to reconstruct draft metabolic models, the resulting drafts are considerably different.

The final step of the model analysis was to assess the metabolic functions included in the draft models. For that, genes included in the draft models were cross-referenced to the COG annotation of the genomes.

Hence, PCA was used to assess the variability of the COG metabolic functions present in the models for the three case study bacteria. The resulting PCA scores and the contribution of PC 1 and PC 2 to the variability of the data are shown in Figure 7. Analysing this figure, PC 1 unquestionably differentiates the metabolic functions of S. thermophilus models from those of X. fastidiosa and M. tuberculosis models. Similarly, PC 2 clearly separates the functional features of M. tuberculosis models from those of X. fastidiosa models.

These results show that the metabolic COG annotation of each model seems to be independent of the template or method used in the model reconstruction. All draft models from the same organism have a similar set of metabolic functions, different from other organisms’ models. Therefore, this suggests that the models’ metabolic functions are not influenced by the approach or the template, in contrast with the models’ reaction content analysis.

4.3 Models assessment

Twenty-four draft metabolic models were reconstructed from the BiGG database using BIT and CarveMe, and compared with manually curated models of the respective organisms under study. Herein, Tables 1–3 showcase the results of the mentioned comparison regarding P, R, and F1.

Table 1 shows the results of M. tuberculosis draft models. The best model in terms of P, R and F1 score was the all models’ template by a large margin. Accordingly, this draft model delivered a P of 0.537, an R of 0.821, and an F1 of 0.649, whereas the second-best model (CarveMe) presented 0.513, 0.617, and 0.560, respectively. Notably, the two best models significantly outperformed all random and selected models templates, especially considering R and F1. In this case, the second-best model (CarveMe) outperformed the best random and selected model templates by 29% and 21%, respectively.
Figure 7: PCA plot of the COG-annotated metabolic functions of all draft models generated by BIT, using different templates, and by CarveMe, for each bacterium. The plot shows the influence of principal components (PC) 1 and 2 on the variability of metabolic features, including the percentage of variance explained by each PC. The dots represent each draft model, and their colour corresponds to the approach used in its reconstruction. Each dot is annotated with the organism-specific identifier, consisting of the first letter of the genus and the first three letters of the species' second name. The set of models for each organism is surrounded by ellipses that are only used for illustration purposes and do not represent real clusters.

Table 1: Comparison of the evaluation metrics between draft models generated with CarveMe and with BIT for M. tuberculosis.

| Model       | Precision score | Recall score | F1 score |
|-------------|-----------------|--------------|----------|
| CarveMe     | 0.513           | 0.617        | 0.560    |
| BIT all     | **0.537**       | **0.821**    | **0.649**|
| BIT selected| 0.396           | 0.269        | 0.320    |
| BIT random 1| 0.438           | 0.281        | 0.343    |
| BIT random 2| 0.401           | 0.259        | 0.314    |
| BIT random 3| 0.383           | 0.277        | 0.322    |
| BIT random 4| 0.383           | 0.323        | 0.350    |
| BIT random 5| 0.356           | 0.237        | 0.285    |

These bold values were included to highlight the model with the highest metrics.

There is a clear difference in performance between the draft models generated with the all models’ template or and CarveMe, which encompass a large part of the BiGG models’ database, and others that used a small part of it (random and selected models’ templates). This discrepancy might have occurred due to the utilisation of M. tuberculosis GEM to generate CarveMe and the all model, as it was stored in BiGG’s database. Such fact affected all other models’ performance, as they did not consider the mentioned GEM during reconstruction.

Nevertheless, the model generated with BIT’s all models’ template surpassed the model generated with CarveMe in all metrics. This result suggests that BIT can generate draft models with higher similarity to curated GEMs than CarveMe, particularly when the reconstruction is based on a curated GEM of the case study. Table 2 describes the assessment of all generated models of S. thermophilus.
Table 2: Comparison of the evaluation metrics between draft models generated with CarveMe and with BIT for *S. thermophilus*.

| Model         | Precision score | Recall score | F1 score |
|---------------|-----------------|--------------|----------|
| CarveMe       | 0.303           | 0.651        | 0.414    |
| BIT all       | 0.155           | **0.812**    | 0.261    |
| BIT selected  | 0.406           | 0.724        | 0.520    |
| BIT random 1  | **0.508**       | 0.679        | **0.581**|
| BIT random 2  | 0.413           | 0.478        | 0.443    |
| BIT random 3  | 0.360           | 0.541        | 0.432    |
| BIT random 4  | 0.466           | 0.703        | 0.560    |
| BIT random 5  | 0.399           | 0.420        | 0.409    |

These bold values were included to highlight the model with the highest metrics.

Table 3: Comparison of the evaluation metrics between draft models generated with CarveMe and with BIT for *X. fastidiosa*.

| Model         | Precision score | Recall score | F1 score |
|---------------|-----------------|--------------|----------|
| CarveMe       | 0.378           | 0.615        | 0.468    |
| BIT all       | 0.261           | **0.672**    | 0.376    |
| BIT selected  | 0.549           | 0.331        | 0.413    |
| BIT random 1  | 0.590           | 0.570        | 0.580    |
| BIT random 2  | **0.652**       | 0.529        | **0.584**|
| BIT random 3  | 0.585           | 0.387        | 0.466    |
| BIT random 4  | 0.454           | 0.052        | 0.093    |
| BIT random 5  | 0.500           | 0.381        | 0.432    |

These bold values were included to highlight the model with the highest metrics.

The best-performance model regarding R was generated using the *all models’ template* (0.812), followed by the model built using the *selected models’ template* with 0.724. The model generated with CarveMe was only the fifth-best model regarding R, having been outperformed by models generated with the *random* method of BIT. Models obtained with CarveMe and BIT *all models’ template* delivered a low number of FN and consequently a high R. This was most expected, as they were constructed considering a higher number of reactions (nearly all BiGG reactions), thus having lower chances of leaving reactions out, though with a higher chance of adding FPs to the model.

As for P, the top-performance model was the *random1* with 0.508, followed by *random4* (0.466) and *random2* (0.417). The models generated with BIT’s *all models’ template* and CarveMe obtained a P of 0.155 and 0.303, respectively. As expected, the number of FP in these models was considerably higher than the others, as these considered a higher number of reactions, having higher chances of adding “noise” to the model.

The models with the best F1 score were the *random1* (0.581), *random4* (0.560) and the *selected* (0.520), outperforming CarveMe, *random5* and *all models*.

Table 3 showcases the performance of draft models of *X. fastidiosa*. In terms of R, the best models were the *all* with a performance of 0.672 and the one of CarveMe with 0.615. The former model surpasses the *random* and *selected* models by at least 10%, whereas the latter only performs better by a difference of 4.5%. As expected and similar to the models of *S. thermophilus*, the R values were higher for the models that made use of a broader collection of template models.

As before, models created with BIT’s *selected* and *random models templates* beat CarveMe and the *all models’ template* models’ performances in P. Accordingly, *random2* obtained the best P result (0.652), outperforming *random1* (0.590), *random3* (0.585), *selected* (0.549), *random5* (0.500), and *random4* (0.454).
Finally, the F1 top-performing models were unequivocally the random2 models’ template (0.584) and random1 models’ template (0.581). Interestingly, these two were also the best P and showed R values closer to CarveMe.

4.3.1 General considerations

Eight models for each of the three organisms were reconstructed and compared against curated GEMs. The results obtained with these comparisons allowed us to reach a few conclusions. As expected, generating models from template GEMs of the organisms under study is beneficial compared with not using those templates. Accordingly, the models of M. tuberculosis generated with CarveMe and the all models’ template method performed better for all metrics. Nevertheless, BIT’s all models’ template method surpassed CarveMe in all metrics.

R allows us to define how many TPs were predicted compared with the total number of real positives. In short, it evaluates how many reactions are left behind during reconstruction, as lower R represents that there are few TP compared with FN. From a curator’s point of view, it is easier to remove reactions than search and add new reactions to the model. Hence, a high recall indicates that the model’s curation process will be less tiresome and time-consuming. BIT’s all and selected models’ template, and CarveMe models were the best for this indicator. Notably, the two best BIT models obtained higher R than CarveMe.

High P translates in a low number of FP compared with all cases predicted as positive. Hence, high values of this metric indicate that a model has less “noise” and has more reactions in a potential curated model. On the other hand, a model with a high P in detriment of R would require more effort in the curation process, as there would be several reactions missing. Therefore, the best models of S. thermophilus and X. fastidiosa were generated with the random models’ template methods of BIT.

Finally, F1 combines P and R into a single metric, directly evaluating the performance of draft models’ construction in terms of balance between FP and FN. High F1 indicates that the reconstruction method is not leveraging errors, both adding and discarding reactions. The best models were generated using BIT random models’ template methods according to this definition.

5 Conclusions

This study concludes that BIT and CarveMe reconstruct different drafts for specific organisms using BiGG as a source of template models. Analysis of the present reactions and metabolic functions of the models’ genes revealed a clear distinction between the drafts reconstructed for each studied organism.

Even though the distribution of genes’ metabolic functions revealed small variability between the different templates used, the comparison of draft reactions found in each template showed that a careful selection of the template models is required as it impacts the reactions added to the draft.

Moreover, when comparing the reactions of the generated drafts to curated and validated models, BIT’s models provided higher metrics’ than CarveMe models. As for the best templates to use, it depends on the end goal. Suppose the aim is to have the lowest number possible of FN. In that case, even if that means having a high number of FP, the best templates seem to be the all models’ template, as removing reactions is less time consuming than adding missing reactions, and the user may not mind the extra “noise” in the draft. However, if the goal is to have the lowest number of FP and FN combined, the F1 metric suggests that the random template leads to better results.

Nevertheless, it is worth considering that BIT only retrieves an initial draft model for the organisms under study. Further curation and validations of the model are still required to obtain a final model that can be used for phenotype prediction. In addition, the conclusions reached by this work can only be stated when considering simple bacterial organisms. Due to the distribution of studied organisms, the applicability of
BIT to eukaryotes or bacteria with a different metabolic profile was not assessed and cannot be guaranteed. Hence, expected future work includes extending this analysis to more organisms.

**Author contribution:** All authors have accepted responsibility for the entire content of this manuscript and approved its submission.

**Research funding:** This study was supported by the Portuguese Foundation for Science and Technology (FCT) under the scope of the strategic funding of UIDB/04469/2020 unit. A. Oliveira (DFA/BD/10205/2020), E. Cunha (DFA/BD/8076/2020), F. Cruz (SFRH/BD/139198/2018), J. Sequeira (SFRH/BD/147271/2019), and M. Sampaio (SFRH/BD/144643/2019) hold a doctoral fellowship provided by the FCT. J. Ribeiro hold a fellowship provided by the University of Minho (UMI-NHO/BIM/2020/68). Oscar Dias acknowledge FCT for the Assistant Research contract obtained under CEEC Individual 2018.

**Conflict of interest statement:** Authors state no conflict of interest.

**References**

1. Gu C, Kim GB, Kim WJ, Kim HU, Lee SY. Current status and applications of genome-scale metabolic models. Genome Biol 2019;20:1–18.
2. Orth JD, Thiele I, Palsson BO. What is flux balance analysis? Nat Biotechnol 2010;28:245–8.
3. Lu H, Li F, Sánchez BJ, Zhu Z, Li G, Domenzain I, et al. A consensus S. cerevisiae metabolic model Yeast8 and its ecosystem for comprehensively probing cellular metabolism. Nat Commun 2019;10:1–13.
4. Norsigian CJ, Pusarla N, McConnell JL, Yurkovich JT, Dräger A, Palsson BO, et al. BiGG Models 2020: multi-strain genome-scale models and expansion across the phylogenetic tree. Nucleic Acids Res 2020;48:D402–6.
5. Kanehisa M, Furumichi M, Tanabe M, Sato Y, Morishima K. KEGG: new perspectives on genomes, pathways, diseases and drugs. Nucleic Acids Res 2017;45:D353–61.
6. Karp PD, Billington R, Caspi R, Fulcher CA, Latendresse M, Kothari A, et al. The BioCyc collection of microbial genomes and metabolic pathways. Briefings Bioinf 2019;20:1085–93.
7. Sayers EW, Bolton EE, Brister JR, Canese K, Chan J, Comeau DC, et al. Database resources of the national center for biotechnology information. Nucleic Acids Res 2022;50:D20–6.
8. Thiele I, Palsson B. A protocol for generating a high-quality genome-scale metabolic reconstruction. Nat Protoc 2010;5:93–121.
9. 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;46:7542–53.
10. Dias O, Rocha M, Ferreira EC, Rocha I. Reconstructing genome-scale metabolic models with merlin. Nucleic Acids Res 2015;43:3899.
11. Capela J, Lagoa D, Rodrigues R, Cunha E, Cruz F, Barbosa A, et al. merlin v4.0: an updated platform for the reconstruction of high-quality genome-scale metabolic models. bioRxiv 2021.02.24.432752 2021.
12. Cui Y, Xu T, Qu X, Hu T, Jiang X, Zhao C. New insights into various production characteristics of Streptococcus thermophilus strains. Int J Mol Sci 2016;17:1701.
13. Baldi P, la Porta N. Xylella fastidiosa: host range and advance in molecular identification techniques. Front Plant Sci 2017;8:944.
14. Russell DG. Mycobacterium tuberculosis: here today, and here tomorrow. Nat Rev Mol Cell Biol 2001;2:569–78.
15. Gerlin L, Cottret L, Cesbron S, Taghouti G, Jacques MA, Genin S, et al. Genome-scale investigation of the metabolic determinants generating bacterial fastidious growth. mSystems 2020;5. https://doi.org/10.1128/mSystems.00698-19.
16. Pastink MI, Teusink B, Bols P, Visser S, de Vos WM, Hugenholtz J. Genome-scale model of Streptococcus thermophilus LMG18311 for metabolic comparison of lactic acid bacteria. Appl Environ Microbiol 2009;75:3627–33.
17. Kavvas ES, Seif Y, Yurkovich JT, Norsigian C, Poudel S, Greenwald WW, et al. Updated and standardized genome-scale reconstruction of Mycobacterium tuberculosis H37Rv, iEK1011, simulates flux states indicative of physiological conditions. BMC Syst Biol 2018;12. https://doi.org/10.1186/s12918-018-0557-y.
18. Hucka M, Bergmann FT, Chaouiya C, Dräger A, Hoops S, Keating SM, et al. The Systems biology markup language (SBML): language specification for level 3 version 2 core release 2. J Integr Bioinform 2019;16. https://doi.org/10.1515/jib-2019-0021.
19. Sequeira JC, Rocha M, Alves MM, Salvador AF. UPIMAPI, reCOGnizer and KEGGCharter: bioinformatics tools for functional annotation and visualization of (meta)-omics datasets. Comput Struct Biotechnol J 2022;20:1798–810.
20. Galperin MY, Kristensen DM, Makarova KS, Wolf YI, Koonin E. Microbial genome analysis: the COG approach. Briefings Bioinf 2019;20:1063–70.
21. Pedregosa F, Michel V, Grisel O, Blondel M, Prettenhofer P, Weiss R, et al. Scikit-learn: machine learning in Python. J Mach Learn Res 2011;12:2825–30.
22. Ebrahim A, Lerman JA, Palsson BO, Hyduke DR. COBRApy: COnstraints-based reconstruction and analysis for Python. BMC Syst Biol 2013;7:1–6.

Supplementary Material: The online version of this article offers supplementary material (https://doi.org/10.1515/jib-2022-0014).