Dynamic Flux Balance Analysis Models in SBML

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2 ABSTRACT

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- Computational models in systems biology and systems medicine are typically simulated using a 3 single formalism such as ordinary differential equations (ODE). However, more complex models require the coupling of multiple formalisms since different biological phenomena are better described by different methods. For example, metabolism in steady state is often modeled using 7 flux-balance analysis (FBA) whereas dynamic changes of model components are better described via ODEs. The coupling of FBA and ODE modeling formalisms results in dynamic FBA models. A major challenge is how to describe such hybrid models that couple multiple formalisms in a 10 standardized way so that they can be exchanged between tools and simulated consistently in a reproducible manner. This paper presents a scheme for encoding and implementation of dynamic FBA models in the Systems Biology Markup Language (SBML), thereby enabling the exchange of 12 multi-framework computational models between software tools. We demonstrate the feasibility of the approach using various example models and show that different tools are able to simulate the hybrid models and agree on the results. As part of this work, two independent implementations of a multi-framework simulation method for dynamic FBA have been developed supporting such 16 models: iBioSim and sbmlutils. 17
- Keywords: dynamic flux balance analysis, DFBA, flux balance analysis (FBA), ordinary differential equations (ODE), static optimization approach (SOA)

1 INTRODUCTION

- 21 In systems biology, mathematical modeling is used to investigate biological systems (Kitano, 2002). The
- 22 resulting computational models enable researchers to make predictions in silico which can be validated
- 23 experimentally. However, the process of model building is time-consuming and error-prone. Model
- 24 reproducibility and exchangeability are of major importance for independent validation of results and
- 25 model reuse, especially in the case of more complex models.
- To achieve reproducibility, interoperability, and consistent model interpretation, a well-defined modeling
- 27 representation with unambiguous syntax is crucial. To this end, standard model representation formats

- exist that enable model exchange, such as the Systems Biology Markup Language (SBML) (Hucka et al.,
- 29 2003; Keating et al., 2020; Hucka et al., 2019).
- 30 SBML has been successfully applied to the encoding of single formalism models, but the encoding of
- 31 hybrid models using SBML has yet to be explored. Some tools have implemented hybrid simulation, such
- 32 as COPASI (Hoops et al., 2006) and E-CELL (Tomita et al., 1999), nonetheless, they lack reproducibility.
- 33 In COPASI, the models fall short of necessary pieces of information for model exchange. Namely, these
- 34 models lack the information about their own model formalism which results in hybrid models being only
- 35 specific to COPASI. In E-CELL, most models are encoded in C++ and only few in SBML. Even though
- 36 the C++ models are repeatable, they are not reproducible because other tools cannot use these files and
- 37 even models encoded in SBML are incomplete and lack the integration of different formalisms.
- 38 The support of hybrid modeling adds new challenges. The present work addresses this problem by
- 39 developing a methodology in conjunction with implementations to support such hybrid modeling efforts.
- 40 We demonstrate the usefulness of our approach by exchanging two models between two distinct simulation
- 41 tools with both implementations leading to similar simulation results.

42 1.1 Coupling multiple modeling formalisms

- Various simulation and analysis methods have been developed in systems biology. Depending on the
- 44 biological question, different methods are preferred. Kinetic time-course simulations based on ordinary
- 45 differential equations (ODE) are often employed to study the dynamics of entities in a model over time.
- 46 Depending on the research question and biological system, such simulations can be non-deterministic
- 47 (stochastic). Other popular simulation methods are Boolean (Thomas, 1973; Kauffman, 1969) models,
- 48 logical models (Morris et al., 2010), and constraint-based approaches (Bordbar et al., 2014).
- 49 Dynamical modeling of metabolic networks by ODE approaches is particularly challenging since kinetic
- 50 parameters needed for ODE models are often unobtainable (Varma and Palsson, 1994). Hence, steady-state
- 51 approaches that do not need kinetic information are employed to model metabolism such as flux balance
- 52 analysis (FBA) (Savinell and Palsson, 1992; Varma et al., 1993) which is based on constraint-based
- 53 optimization. This method only requires the connectivity of the reactions and metabolites along with the
- 54 respective stoichiometry, an objective function, such as cell growth, and additional constraints like flux
- 55 bounds. The idea is to constrain the model based on the stoichiometry of the reactions and optimize the
- 56 objective function while satisfying the flux constraints. These models do not require kinetic information
- 57 and can be simulated efficiently even in case of very large systems.
- 58 Biological research questions often require the coupling of different model formalisms. One such recent
- 59 example is the whole-cell model for the *Mycoplasma genitalium* (Karr et al., 2012) that is encoded using a
- 60 mixture of Boolean networks, stochastic processes, differential equations, and FBA.

1.2 Dynamic flux balance analysis

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- One disadvantage of FBA is that it cannot express the dynamics of the metabolites since it does not change
- 64 amounts or concentrations of species, but only provides information about the optimal flux distribution
- 65 for the given optimization problem. Due to this limitation, the field of dynamic FBA (DFBA) (Varma and
- 66 Palsson, 1994) has emerged, which couples the stationary flux distribution resulting from FBA with the
- 67 kinetic update of the metabolites taken up or consumed by the FBA network. For DFBA models, a FBA
- 68 submodel is coupled to a kinetic model (ODE).

- Besides the whole-cell model, which uses DFBA as a core module, various metabolic pathways have
- 70 been modeled using DFBA. DFBA has been applied in small-scale examples (Varma and Palsson, 1994;
- 71 Mahadevan et al., 2002; Luo et al., 2006), over medium-size models (Pizarro et al., 2007; Lequeux et al.,
- 72 2010; Meadows et al., 2010), and up to genome-scale DFBA applications (Hanly and Henson, 2011;
- 73 Hjersted et al., 2007). For an overview, see Table 1 in (Höffner et al., 2013).
- The coupling between FBA and kinetic model parts can be implemented via three main approaches: static optimization approach (SOA), dynamic optimization approach (DOA), and direct approach (DA) (Gomez et al., 2014). The SOA approach solves the linear programming (LP) problem of FBA at each time step using an Euler forward method assuming constant fluxes over the time step (Gomez et al., 2014). DOA approaches optimize simultaneously over the entire time period by solving a nonlinear programming problem (NLP). The DA approach directly includes the LP solver on the right-hand side of the ordinary
- The advantage of the SOA is its relatively simple implementation, which is why most of the published DFBA models use the SOA approach. However, SOA is often less accurate compared to other computational more demanding methods such as DOA. The DA method exhibits the best trade-off between accuracy and runtime performance but has its downsides in terms of implementation difficulty. For this work, we use the SOA method. Its simplicity makes it a good candidate to use as proof of concept for this work.

1.3 Exchangeability & reproducibility of models

differential equations (ODEs).

- Despite the wide range of published DFBA models no standard for the exchange of such models exists. Existing models are hard-coded, such as the whole-cell model which is implemented in MATLAB. Hereby, the mathematical model is separated in the respective kinetic and FBA formalisms in a script along with the connections between the kinetic and flux balance parts of the models. As a consequence, it is not possible to exchange existing DFBA models between different software tools. Thus, they cannot be reproduced or validated. This is especially problematic in the case of DFBA models because often multiple optima can exist for the FBA model part (and the various time steps). The resulting DFBA results are not unique since they depend on the analysis implementation (how a solver selects one of the possible FBA solutions). In addition, the simulation results may depend on the selected step size of the SOA algorithm, in particular, if the step size is not small enough.
- While it is possible to replicate the same scripts in different programming languages, it is unpractical, error-prone, unnecessary, often leads to data loss, and most importantly does not solve the underlying problem of non-exchangeability of such models. For these reasons, script replication makes achieving reproducibility difficult and often infeasible. The necessity of an exchange format for DFBA emerged from efforts trying to encode and reproduce the DFBA submodel of the whole-cell model using standards during the whole-cell workshop (Waltemath et al., 2016).

1.4 Model standards

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To achieve exchangeability and reproducibility of models, standards for the encoding of models have been created. The de-facto standard for systems biology models is SBML (Hucka et al., 2003; Keating et al., 2020). SBML core elements are used to describe mathematical models of reaction-based networks and provide the means to encode computational models based on reaction networks that can be represented both deterministically and stochastically. SBML uses packages for extending the functionality of core elements. While SBML is used to encode mathematical models of biological networks, there are different standards for other purposes: the Simulation Experiment Description Markup Language (SED-ML) is used for describing simulations (Waltemath et al., 2016; Bergmann et al., 2018), the Systems Biology Graphical

- 112 Notation (SBGN) is used for describing visualizations (Le Novère et al., 2009), and COMBINE Archives
- 113 are used for grouping files in a single archive necessary to reproduce a modeling experiment (Bergmann
- et al., 2014). The main advantage of using these standards over hard-coding models in code is the ability to
- 115 exchange models between research groups and reproduce results using various tools that support these
- 116 standards. In addition, these standards enable the use of semantic annotations to document the model and
- model components which enhances the reusability and interoperability (Neal et al., 2019, 2020).
- One of the challenges in SBML models is the limitation of models to a single formalism lacking support
- 119 for the expression of models using multiple formalisms. Although there are several tools that support ODE
- 120 simulation and FBA, they all support them independently. In order to overcome this challenge, this paper
- introduces a scheme that enables the coupling of ODE and FBA models. This paper demonstrates that this
- 122 scheme facilitates exchangeability and reproducibility by encoding and simulating DFBA models in both
- 123 iBioSim (Watanabe et al., 2019) and sbmlutils (König, 2022).

2 MATERIAL AND METHODS

124 2.1 Model encoding

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- The DFBA models presented in this paper were created in the proposed scheme either using a graphical
- 126 user interface in iBioSim or a script-based approach in sbmlutils. For a given model, the four
- submodels (TOP, FBA, BOUNDS, and UPDATE) were packaged with the corresponding simulation files
- 128 using SED-ML in COMBINE archives in order to facilitate the exchange between tools. All models and
- 129 simulation results are available from https://github.com/matthiaskoenig/dfba.

2.2 Stationary optimization approach (SOA)

- A stationary optimization approach for DFBA was implemented as a simulation algorithm in iBioSim and sbmlutils following the simulation scheme depicted in Figure 1.
- The following paragraph assumes familiarity with SBML and we refer to the SBML specification for
- more information (Hucka et al., 2019). As the first step, all of the species and parameters in the model are
- initialized and each variable is assigned an initial value. After the initialization step, the FBA submodel is
- 136 executed. During the FBA step, reaction fluxes are computed using the initial flux bound values where
- 137 the flux bounds for the reactions come from the top-level using replacements from the SBML comp
- 138 package (Smith et al., 2015). In SBML, replacements of parameters and species indicate the top-level
- 139 entities are the same entity as the one being replaced. Once the fluxes are computed, they are assigned on
- 140 the top-level to parameters using assignment rules. These parameters represent reaction rates.
- 141 After computing reaction fluxes, the update step is performed concurrently with a dynamic step by
- 142 computing the time-evolution of every species in the UPDATE and KINETIC submodels. Species that
- 143 affect any flux bound in the FBA submodel are updated on the top-level. The new bounds are used in the
- 144 FBA submodel for the next time step. Simulation time is incremented at the end. If the time limit is reached,
- then the simulation is complete. Otherwise, all of the steps are repeated.
- 146 The SOA simulation algorithm has been implemented in iBioSim and sbmlutils. The iBioSim
- 147 tool uses the structure of (Watanabe and Myers, 2014) for simulation. The sbmlutils tool uses
- 148 roadrunner (Somogyi et al., 2015) for the kinetic simulation and cobrapy (Ebrahim et al., 2013) to solve
- 149 the FBA problem. Both iBioSim and sbmlutils take an SBML file that describes a DFBA model and
- 150 a SED-ML file that describes the simulation experiment. In the proposed approach, SED-ML is mainly
- 151 used to indicate which simulation algorithm to use, the time points in which tools should print out the
- 152 values of the variables, the initial time, and the time limit. The SED-ML files provide a minimal simulation

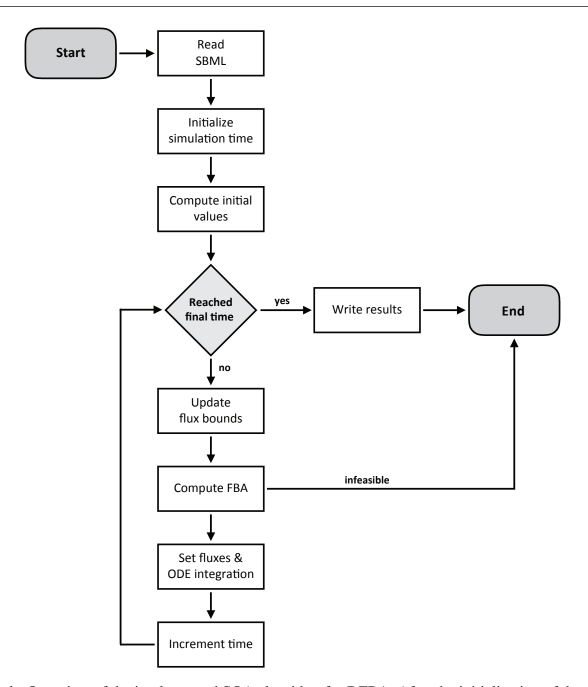


Figure 1. Overview of the implemented SOA algorithm for DFBA. After the initialization of the model, the FBA and kinetic simulations are run iteratively until the simulation endpoint. In every step, FBA is used to compute the reaction rates of the FBA network. Subsequently, based on the computed FBA rates, the values of the species are updated dynamically. In the SOA approach, FBA fluxes are assumed to be constant within a time step. For a detailed description see the Material And Methods Section.

experiment to check reproducibility between implementations. The value of each time increment for SOA is defined by a parameter with id dt in the SBML model, which can be overwritten by the SED-ML file for the actual simulation. An ontology term for the description of DFBA simulation algorithms has been introduced in the Kinetic Simulation Algorithm Ontology (KISAO) (Zhukova et al., 2011), term KISAO: 0000500 corresponding to the DFBA-SOA method, and is used in the SED-ML descriptions.

2.3 Reproducibility between tools

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In order to test interoperability based on the proposed scheme, models were built in both the iBioSim and the sbmlutils tools. Models built in iBioSim were then imported into sbmlutils and viceversa to check whether models could be interpreted by both tools consistently. This was done in an iterative manner and issues were solved by clarifying the encoding scheme by adding additional rules which resolved ambiguities. Ensuring reproducibility for DFBA models is challenging because there may exist several possible outcomes that satisfy the objective function and constraints of the FBA models. Different trajectories can result from the DFBA simulation depending on how a solver and implementation selects one of the multiple optima. The issue of multiple optima was solved by guaranteeing uniqueness of the solution in every time step based on Flux Variability Analysis (FVA) (Mahadevan and Schilling, 2003). FVA gives the possible minimal and maximal fluxes for each reaction in each step of the simulation. If all minimal fluxes are equal to all maximal fluxes for a time point a solution is unique in the time point. If all time points are unique the solution is unique. As a practical note: If the solution is not unique, the addition of additional constraints to the FBA problem allows to make the solution unique. Reproducibility of the model simulations was tested by comparing the numerical SOA results between the two tools for models with unique solutions (see Supplementary Material S2). Results were assumed as numerically identical if the absolute difference for every time point t_k for all dynamical FBA species in the model c_k was smaller than the tolerance $\epsilon = 10^{-5}$. The difference is computed as follows:

$$abs(c_i(t_k)_{sbmlutils} - c_i(t_k)_{ibiosim}) \le \epsilon \ \forall c_i, t_k$$

In SOA-DFBA, it is important that the time steps dt are small enough so that the solution converges against the optimal solution. Solutions vary if selected step sizes are too large. To highlight this fact, changing the step size in the toy_wholecell model from 1.0 to 0.1 resulted in differences in steady-state concentrations of up to 10%. Consequently, the step size was reduced until the changes did not affect the simulation results.

3 RESULTS

- 181 The major result of this work is the creation of the first schema for a DFBA encoding in SBML,
- 182 demonstrating hybrid computational models to be exchangeable and reproducible between tools. In
- the following, the schema and its application to multiple DFBA models is presented.

184 3.1 Schema for dynamic flux balance analysis

This paper proposes for the first time a schema to encode hybrid models, such as DFBA model, in SBML. The developed schema consists of rules, guidelines, and additional information and is available in the Supplementary Material S1. The latest version of the document is available from https://github.com/matthiaskoenig/dfba/. Proposals, errata, and updates to the schema are managed via the respective issue tracker and releases.

In this Section, we provide a high-level overview of the underlying concepts used in the schema, followed by an application of the schema to encode DFBA models.

The DFBA model is constructed hierarchically using the SBML comp package, separating the hybrid model into different building blocks based on the respective functionality and modeling frameworks (Figure 2). The top-level model is hereby composed of four submodels: (i) a kinetic submodel that computes flux bounds based on the dynamic metabolite availability and ensures that the FBA problem is

196 constrained by the available metabolite resources (BOUNDS submodel); (ii) a FBA submodel that encodes

- 197 metabolism as a FBA problem (FBA submodel); and (iii) a kinetic submodel that updates the amounts and
- 198 concentrations of the dynamic metabolites changed via the FBA submodel via consumption or production
- 199 (UPDATE submodel); (iv) an optional kinetic submodel that represents a dynamic part with all kinetics
- 200 other than the metabolic pathway, such as DNA transcription, DNA translation, and protein degradation,
- among others (KINETIC submodel). Alternatively, arbitrary kinetics can be part of the top model.
- The top-level model couples the three different submodels using SBML comp replacedElement
- and replacedBy constructs with the interface between the submodels defined via comp ports (which
- 204 define which model components of the submodels can be connected, i.e, are exposed).
- In order to describe the different formalisms of each submodel, the Systems Biology Ontology (SBO)
- 206 is used (Courtot et al., 2011). The SBO defines controlled vocabulary terms used in the systems biology
- 207 field. The SBO terms are arranged in a taxonomic hierarchy using a tree structure. This allows the
- 208 grouping of terms that are related to one another. The modeling formalisms of the individual submodels
- 209 are described using terms on the modeling framework branch, where FBA models are described using the
- 210 flux balance framework term, stochastic processes are described using the non-spatial discrete framework
- 211 term, and differential equations are described using the non-spatial continuous framework term. Although
- 212 the terms for stochastic processes and differential equations can be used for describing either stochastic or
- 213 deterministic simulation methods, these terms were selected because they are the ones that best describe
- 214 these two formalisms.
- In addition to the modeling formalism, other key components are annotated in the submodels via SBO
- 216 terms in the schema, like the upper and lower flux bounds and the exchange reactions in the FBA submodel
- 217 defining which metabolites can be consumed or produced in the FBA part of the DFBA, or the dynamic
- 218 species in the top model changed by the FBA submodel. By the means of these annotations, the interface
- 219 between the hybrid submodels can be clearly defined.
- 220 All of the interconnections between the submodels are encoded in SBML rather than using an external
- 221 approach like for instance via SED-ML. The connections between model components are crucial
- 222 information of the model and should be part of the model encoding. SED-ML is only used to encode which
- 223 simulation to run with the model. As a consequence, this schema requires only a single hierarchical SBML
- 224 model and a single SED-ML file.

225 3.2 Minimal Example (toy_wholecell)

- In order to illustrate the proposed schema, a simplified example of a whole-cell model was created
- 227 and visualized. The corresponding files (i.e. COMBINE archive and Cytoscape visualization) are in
- 228 Supplementary Material S3. The visualization shows how the different submodels connect with each other
- 229 in a flat form.
- 230 This model is constructed hierarchically where a top-level model is created to instantiate different
- 231 submodels (BOUNDS, UPDATE, and FBA) and make the necessary connections between them. The figure
- 232 illustrates the structure of each submodel and how each submodel ties in with each other in a flat version of
- 233 the model once all of the connections are established.
- In the example, the FBA submodel imports species A and convert it via a linear chain of reactions to
- 235 species C. The exchange reactions EX_A and EX_C contain the rate of consumption and production of
- 236 the respective species. The TOP model contains assignment rules that assign the fluxes to the parameters
- 237 pEX_A and pEX_C. The pEX_A and pEX_C parameters are used by the UPDATE model to compute the new

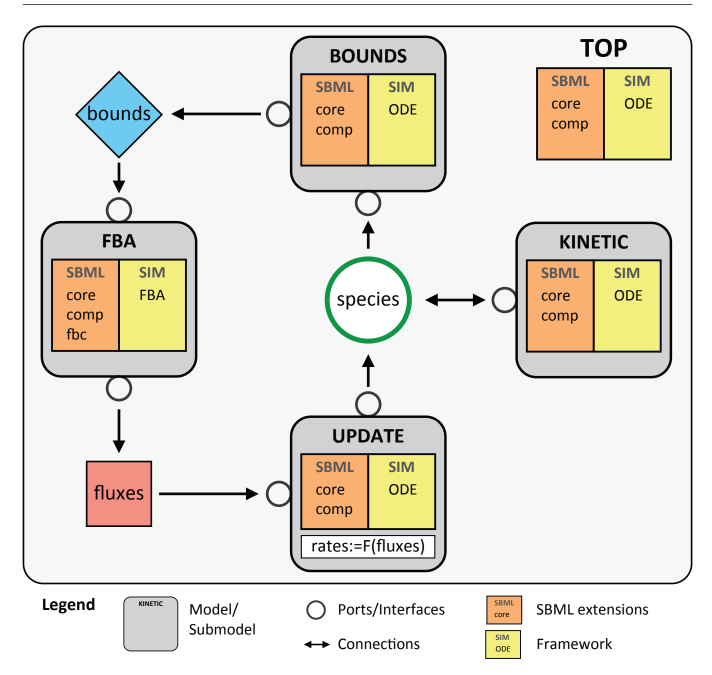


Figure 2. Overview of the schema for DFBA model encoding in SBML. The hierarchical SBML model is composed of a top-level model with four submodels: FBA, BOUNDS, UPDATE, and KINETIC. The individual submodels are connected via ports. The respective SBML packages used are listed in the models, as well as the employed simulation method. The BOUNDS submodel calculates the upper and lower flux bounds based on metabolite availability. The FBA submodel computes the reaction fluxes of the metabolic model encoded via the SBML fbc package using the bounds as constraints. The UPDATE submodel calculates the dynamic update of the dynamic metabolites affected by the FBA model. The rates of change are hereby functions of the FBA fluxes. The KINETIC submodel includes all of the other processes in the model, which may affect or be affected by entities in metabolism. The top-level model ties together the different submodels using SBML comp replacedElement and replacedBy constructs.

values of the dynamic species A and C via the update reactions update_A and update_C. The BOUNDS model calculates the bounds of all FBA exchange reactions (constraining the availability of the dynamic species). In the example, the upper bound ub_R1 of reaction R1 is changed via a rate rule. Additional

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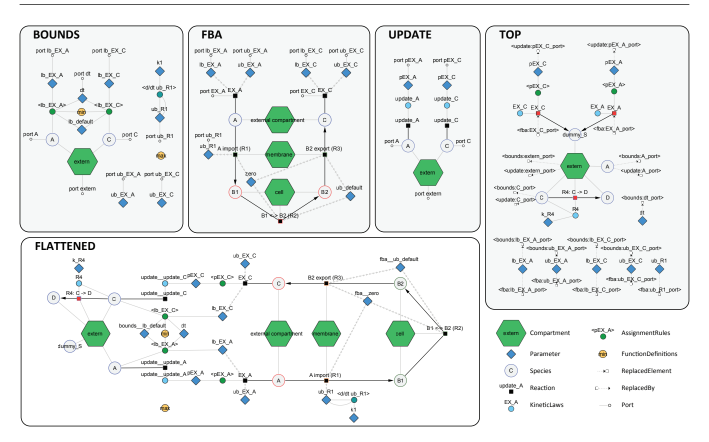


Figure 3. Detailed schema of the minimal example model (toy_wholecell). The figure shows the components in the BOUNDS, FBA and UPDATE submodels. Links between submodel components are based on ports which are connected elements via TOP model replacements (replacedElements and replacedBy). The flattened SBML comp model (FLATTENED) shows the resolved connections between the different submodels after these replacements have been performed. The flattened model can not be simulated because the separation of the modeling formalisms is lost in the flattening process. The network visualization are available as interactive graphs in Cytoscape as Supplementary Material, which provide additional information and annotation of the components. The figure was created with cy3sbml using the SBML models (König et al., 2012).

- kinetics are encoded in the TOP model, such as the kinetic conversion of C to D (these could also be in a separate KINETIC submodel).
- In order to validate the exchangeability and reproducibility of the model, simulations were performed using the simulation algorithm described in Figure 1 with results depicted in Figure 4. Both implementations resulted in numerically identical results (see Section 2.3). Importantly, our encoding schema allowed to reproduce the numerical results even if the step sizes were not yet small enough to have converged against the correct solution, thereby allowing to test the effects of varying step sizes in a reproducible manner.
- In addition to the presented minimal model, a second model and its corresponding Cytoscape visualization of a simplified DFBA glycolysis (toy_atp) is available in the supplement (COMBINE archive in Supplementary Material S3)

3.3 Diauxic growth in Escherichia coli (diauxic_growth)

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The next example is an encoding and reproduction of results from a published DFBA model of diauxic growth of the *Escherichia coli* (Mahadevan et al., 2002) consisting of four reactions between four

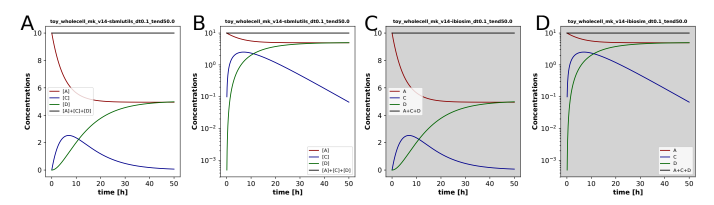


Figure 4. DFBA simulation results for the toy_wholecell model in two different tools (sbmlutils: A, B; iBioSim: C, D). This demonstrates that models can be exchanged by different tools using standards and the results can be reproduced when using the same simulation algorithm. Species A is converted to C via the FBA subnetwork over time. Species C is converted to D via the kinetic parts in the top model. Species A is not consumed completely because of the import of A in the FBA subnetwork via R1 which is shut down over time via a rate rule for the upper flux bound. The model was simulated for 50[h] with a time step dt of 0.1[h].

metabolites: glucose (Glext), oxygen (O_2), acetate (A_c), and biomass (X). The model can grow either aerobically on acetate (v_1), aerobically on glucose (v_2 or v_3), or anaerobically convert glucose to acetate:

$$v1: 39.43A_c + 35O_2 \rightarrow X$$

 $v2: 9.46Glcxt + 12.92O_2 \rightarrow X$
 $v3: 9.84Glcxt + 12.73O_2 \rightarrow 1.24A_c + X$
 $v4: 19.23Glcxt \rightarrow 12.12A_c + X$

The kinetic part of the model is described by the following differential equations:

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$$\frac{dGlcxt}{dt} = A^{Glcxt}\nu X$$

$$\frac{dA_c}{dt} = A^{A_c}\nu X$$

$$\frac{dO_2}{dt} = A^{O_2}\nu X + k_L a(0.21 - O_2)$$

$$\frac{dX}{dt} = (v1 + v2 + v3 + v4)X$$

where A^{Glext} , A^{A_c} , A^{O_2} are the respective rows of each variable in the stoichiometry matrix and $k_L a$ is the mass transfer coefficient of oxygen. For a detailed description see (Mahadevan et al., 2002).

The model and its corresponding Cytoscape visualization is available in Supplementary Material S3.

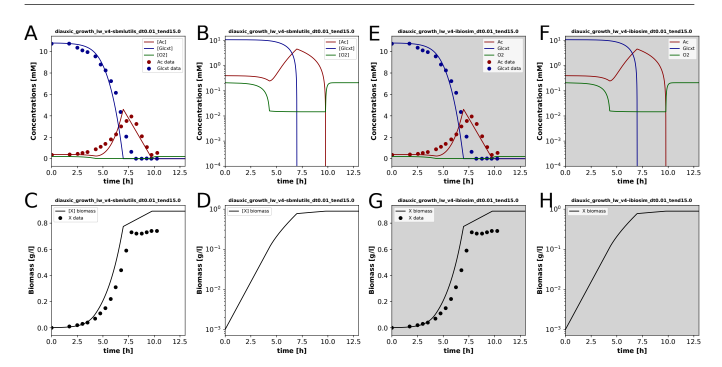


Figure 5. Simulation results for the diauxic growth of *Escherichia coli* in sbmlutils (**A**, **B**, **C**, **D**) and iBioSim (**E**, **F**, **G**, **H**). The model is able to reproduce the general behavior from experimental data. The cell is growing exponentially while glucose is present, but when the cell runs out of glucose, growth slows down and is limited mainly by oxygen. However, when the cell runs out of glucose and oxygen, growth diminishes significantly. The model was simulated for 15[h] with a time step dt of 0.01[h].

The results in Figure 5 depict an exponential growth phase using glucose aerobically until running out of glucose, which at this point the cell grows linearly due to oxygen. When both oxygen and glucose run out, the cell growth stagnates. Experimental data from (Varma and Palsson, 1994) is plotted alongside the simulation results. The model is able to capture the behavior observed in the experimental data. The results are equivalent to the models in (Mahadevan et al., 2002).

We hereby showed that our schema is able to encode published DFBA models, resulting in a reproducible and exchangeable model representation between tools.

3.4 Escherichia coli Core Metabolism (ecoli)

To demonstrate the feasibility of the proposed schema and method for real-world examples of DFBAs, a larger metabolic network for the core metabolism of *Escherichia coli* (Orth et al., 2010) was encoded in the proposed schema and simulated as shown in Figure 6. The model is available as COMBINE archive in Supplementary Material S3. The FBA submodel was downloaded from BiGG (King et al., 2016) (core metabolism of *Escherichia coli* str. K-12 substr. MG1655) and transformed to an DFBA model in an automatic fashion using sbmlutils. BiGG models encode the exchangeable species via annotated exchange reactions which allows an automatic inference of the dynamic species. Only additional information required to run a DFBA simulations are initial concentrations for the species. The automatic encoding of larger scale examples demonstrates the scalability of the proposed encoding approach.

While sbmlutils is able to find a solution for the model, iBioSim cannot as it runs into an unfeasible solution in the middle of simulation. This captures the well-known problem of DFBA with multiple solutions. The FBA problem is not constrained enough to result in a unique solution and depending on which solution the simulator picks, different solutions and thereby trajectories arise. Despite the existence

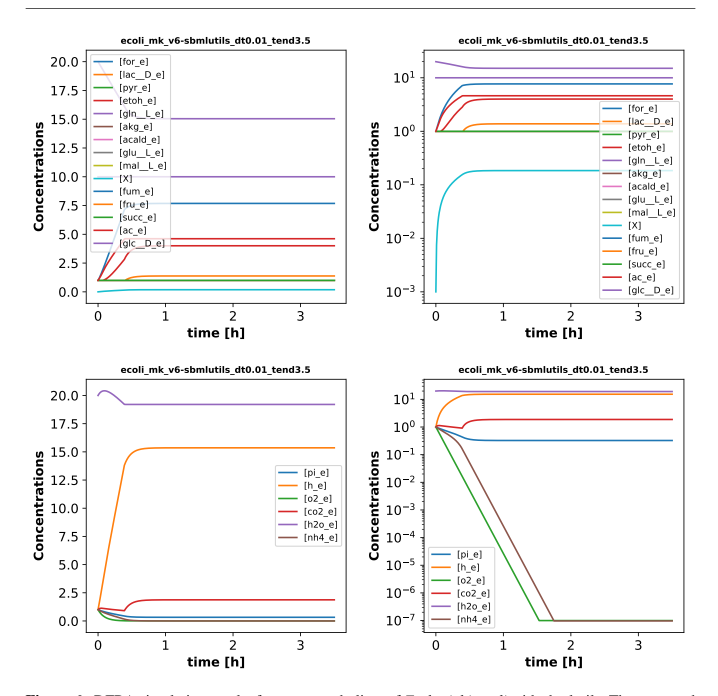


Figure 6. DFBA simulation results for core metabolism of *Escherichia coli* with sbmlutils. The proposed approach can be used in larger models, such as the *Escherichia coli* model described in the paper. The model is growing aerobically on glucose in the initial phase and reaches a steady state after oxygen is consumed. The model was simulated for 3.5[h] with a time step dt of 0.01[h].

of multiple solutions, tools and LP solvers typically pick solutions deterministically. Hence, single tools can reproduce their own results, but results are irreproducible between different implementations. Without the use of standards, this could never be demonstrated because variations in results could be due to discrepancies in the model, and not in the tool.

4 DISCUSSION

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The ability to encode hybrid models, modularity and reproducibility of models are indispensable for encodings of complex models in computational biology. In this work, we presented such an approach,

- 286 which allows a clear separation of the different modeling formalisms via hierarchical models and defining
- 287 the interfaces between the submodels. Here we propose and implement an exchangeable and reproducible
- 288 hybrid modeling scheme. This scheme for encoding DFBA models has been implemented in two different
- 289 tools, demonstrating the exchangeability and reproducibility of our approach on various examples models.
- 290 iBioSim and sbmlutils are freely available for download and offer the necessary infrastructure for
- 291 anyone to develop DFBA models using the proposed scheme. Currently, the proposed approach supports
- 292 the modeling of DFBA models based on the SOA simulation algorithm. Hence, our approach only covers a
- 293 subset of DFBA algorithms and a subset of possible frameworks.
- 294 Most DFBA models are stiff. Hence, short time steps are required for stability and for accurate results.
- 295 Due to the need for short time steps, the SOA approach is computationally expensive. Future directions
- 296 include the exploration of adaptive time steps for executing the DFBA with SOA, alternative DFBA
- 297 methods, such as DOA or DA, and extending our scheme to encode such models.
- 298 Our current approach is limited to the coupling of ODEs to FBA models. Different hybrid modeling
- 299 types, such as any mixtures of differential equations, stochastic processes, or Boolean models may yield
- 300 promising results in the future. The proposed approach of decoupling different modeling formalisms via
- 301 the comp package could work similarly for other modeling frameworks like Boolean models.
- 302 So far, only small to medium-size DFBA models have been encoded in our proposed approach. For future
- 303 work, we will encode genome-scale metabolic models such as HepatoNet1 (Gille et al., 2010) which will
- 304 allow us to assess the scalability and performance of the proposed approach.

CONFLICT OF INTEREST STATEMENT

- 305 All authors declare that the research was conducted in the absence of any commercial or financial
- 306 relationships that could be construed as a potential conflict of interest.

AUTHOR CONTRIBUTIONS

- 307 MK and LHW designed the study, developed the computational models, implemented and performed
- 308 the analysis, and wrote the initial draft of the manuscript. All authors discussed the results. All authors
- 309 contributed to and revised the manuscript critically.

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DATA AVAILABILITY STATEMENT

- 320 Availability: All materials and models are available from https://github.com/matthiaskoenig/dfba. The
- 321 tools used in this project are freely available: iBioSim at https://www.async.ece.utah.edu/ibiosim and

322 sbmlutils at https://github.com/matthiaskoenig/sbmlutils/.

5 SUPPLEMENTARY MATERIAL

- 324 Supplementary Material is available online.
- 325 S1 Schema for encoding DFBA in SBML.
- 326 S2 Reproducibility results between sbmlutils and iBioSim.
- 327 S3 texttttoy_wholecell COMBINE archive and Cytoscape figure for DFBA models (minimal model,
- minimal glycolysis model, diauxic model, and Escherichia coli core model).

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