

Memote: A community driven effort towards a standardized genome-scale metabolic model test suite

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Abstract:

Several studies have shown that neither the formal representation nor the functional requirements of genome-scale metabolic models (GEMs) are precisely defined. Without a consistent standard, comparability, reproducibility, and interoperability of models across groups and software tools cannot be guaranteed.

Here, we present memote (<https://github.com/opencobra/memote>) an open-source software containing a community-maintained, standardized set of metabolic model tests. The tests cover a range of aspects from annotations to conceptual integrity and can be extended to include experimental datasets for automatic model validation. In addition to testing a model once, memote can be configured to do so automatically, i.e., while building a GEM. A comprehensive report displays the model's performance parameters, which supports informed model development and facilitates error detection.

Memote provides a measure for model quality that is consistent across reconstruction platforms and analysis software and simplifies collaboration within the community by establishing workflows for publicly hosted and version controlled models.

Introduction

The reconstruction and analysis of metabolic reaction networks provide mechanistic, testable hypotheses for an organism's metabolism under a wide range of empirical conditions¹. At the current state of the art, genome-scale metabolic models (GEMs) can include thousands of metabolites and reactions assigned to subcellular locations,

gene-protein-reaction rules (GPR), and annotations, which provide meta-information by referencing large biochemical databases. This development has been facilitated by standard protocols for reconstruction² and guidelines for provenance-tracking and interoperability³⁻⁵. However, the quality control of GEMs remains a formidable challenge that must be solved to enable confident use, reuse, and improvement.

Both Ravikrishnan and Raman⁶ and Ebrahim et al.⁷ lamented the lack of an agreed-upon description format as they found that GEMs can be published as SBML⁸, MATLAB files, spreadsheets, and PDF. While the former noted that incompatible formats limit the scientific exchange and, thus, the ability to reproduce calculations on different setups, the latter elaborated how formatting errors can directly cause inconsistent results when parsed and evaluated with various software packages.

When comparing four previously published models for *Pseudomonas putida* KT2440, Yuan et al. discovered that in identical simulation conditions the predicted growth rate of one model was almost twice as high as that of another⁹. Moreover, one of the examined models could generate ATP without needing to consume any substrate, rendering some model predictions useless.

This behavior occurs when a model's reaction directions are not checked for thermodynamic feasibility, leading to the formation of flux cycles which provide reduced metabolites to the model without requiring nutrient uptake. Fritzemeier et al.¹⁰ detected such erroneous energy-generating cycles (EGCs) in the majority of GEMs specifically in the MetaNetX^{11,12} (~66%) and ModelSEED¹³ (~95%) databases, which mostly contain automatically-generated, non-curated metabolic models. Although the authors found that EGCs are rare in manually-curated GEMs from the BiGG Models database (~4%), their effect on the predicted growth rate in FBA may account for an increase of up to 25%. This

makes studies involving the growth rates predicted from such models unreliable. It is possible to identify and correct these issues either with functions included in the COBRA Toolbox¹⁴, or the modified GlobalFit algorithm¹⁵ presented by Fritzemeier et al.¹⁰. Yet, as the models of *P. putida* analyzed by Yuan et al. show, this is not done consistently⁹.

Investigating the biomass compositions (BCs) of 71 manually-curated prokaryotic GEMs, Xavier et al. found that organic cofactors (e.g., Coenzyme A, pyridoxal 5-phosphate, and S-adenosyl-methionine) are missing even though their inclusion is vital to a model's performance in gene-essentiality studies¹⁶.

Chan et al. highlighted deviations in molecular weight as another problem with the formulation of BCs¹⁷. Conforming to the defined molecular weight of 1 g/mmol is essential to reliably calculate growth yields, cross-compare models, and obtain valid predictions when simulating microbial consortia. Half of the 64 tested models deviated from the defined 1 g by up to 5%, with the other half differing even more strongly. Any discrepancy, however, should be avoided as the smallest error affects the predicted biomass yield, favoring models containing BCs which sum to lower molecular weight.

In addition to discussing encoding related problems, Ravikrishnan and Raman stressed that missing metabolite and reaction annotations are further fundamental issues when trying to exchange GEMs which have been generated from different platforms, or when attempting to integrate them into existing computational workflows⁶. Mapping annotations between biochemical databases is not trivial but semi-automatic approaches help to reduce the required manual effort¹⁸. Nonetheless, they reported the absence of metabolite annotations (i.e., metabolite formula, database-dependent (e.g., ChEBI ID), and database-independent

i.e. derived from the properties of the object itself (e.g., SMILES, InCHI) references) in almost 60% of the 99 models they examined.

Increasing numbers of manually-curated and automatically-generated GEMs are published each year, growing both in scale and scope; from models on single cells to multi-organism communities¹⁹ to multi-compartmental plant²⁰, human and cancer tissue models²¹.

Especially when considering the growing application of models to human health and disease, it becomes essential to address any remaining issues concerning reproducibility and interoperability to pave the way for reliable systems medicine²².

Thus, we need to establish a standard framework which ensures that:

- Models are formulated consistently in a software agnostic manner.
- Components of GEMs are uniquely identifiable using standardized database-independent identifiers which can be converted easily using cross-references.
- Default conditions and mathematically specified modeling formulations are precisely defined to allow the reproduction of the original model predictions.
- Models yield biologically feasible phenotypes when analyzed under alternating conditions.
- Data that has been used to curate/parametrize the model are adequately documented to precisely understand the model refinement process.

Here, we argue for a two-pronged approach in creating this framework: 1) We advocate the use of the latest version of the *SBML Level 3 Flux Balance Constraints (FBC) Package*²³ as the agreed-upon description format, which renders GEMs to be independent through a

unified formulation. 2) Borrowing tools and best practices from software development ^{24,25}, we present *memote* as a unified approach for benchmarking metabolic models.

Results

SBML: Tool-independent model formulation

Historically, GEMs have been structured and stored in many non-standard ways, for example, tool specific formats or language dialects ⁶. This prevented the accurate exchange of models between various software tools and the unambiguous, machine-readable description of all model elements such as chemical reactions, metabolites, gene associations, annotations, objective functions and flux capacity constraints. While a widely used model description standard, such as the Systems Biology Markup Language (SBML) Level 3 Core⁸, can describe some of these components, e.g., reactions, metabolites, or annotations, it cannot present other model components needed to describe a parameterised GEM or FBA model in a structured and semantic way.

Consequently, an adequate model description format is needed that allows for the unambiguous definition and annotation of such a model's components and underlying mathematics.

With the release of SBML Level 3 it has become possible to load specific modeling packages that extend the core format with additional features. The *SBML Level 3 Flux Balance Constraints (FBC) Package* (SBML3FBC) has been specifically designed to address the problems described above. Such extensions allow users to take advantage of infrastructure built around SBML, while also providing a smaller set of specifications that can be adjusted to cater to the quickly changing needs of a specific research area. The FBC

package allows for the unambiguous, tool neutral and validatable SBML description of domain-specific model components such as flux bounds, multiple linear objective functions, gene-protein-reaction associations, metabolite chemical formulas, charge and related annotation²³. The SBML and constraint-based modeling communities collaboratively develop this package and update it based on user input. As a result, FBC Version 2 is the *de facto* standard for encoding GEMs. Critical to this process is its implementation in a wide range of constraint-based modeling software and adoption by public model repositories^{22,26–34}. We believe these factors make SBML3FBC the optimal format for sharing and representing GEMs, thus models encoded in SBML3FBC serve as the input to memote.

Memote: Community-driven quality control

In software engineering, test-driven development ensures that in response to a defined input a piece of code generates the expected output³⁵. Distributed version control represents an efficient way of tracking and merging changes made by a group of people working on the same project³⁶. Finally, continuous integration ties these two principles together by automatically triggering tests to be executed after each change that is introduced to the project³⁷. Memote (/ˈmiːmoʊt/ (IPA)), short for metabolic model tests, is an open-source python software that applies these engineering principles to genome-scale metabolic models.

Memote accepts stoichiometric models encoded in SBML3FBC as input, allowing users to benchmark them against a set of consensus tests. By enabling researchers to quickly interrogate the quality of GEMs, problems can be addressed before they affect reproducibility and scientific discourse, or increase the amount of time spent troubleshooting

³⁸

Memote supports two basic workflows (Figure 1a). First, by running the test suite on a model once, memote generates a comprehensive, human-readable report, which quantifies the model's performance. By this information, a definitive assessment of model quality can be made, i.e., by editors or reviewers. This workflow is accessible through a web interface (<https://memote.dd-decaf.eu>), analogous to the SBML validator ²⁷, or locally through the command line.

Second, for model maintenance and reconstruction, memote coordinates version control and continuous integration, such that each tracked-edit in the reconstruction process can progressively be tested. Users edit the model with their preferred reconstruction tool, and export to SBML afterward. This way, each incremental change can be tested with the suite. Then, a report on the entire history of results serves as a guide towards a functional, high-quality GEM. This workflow is accessible through the command line and may be extended to include custom tests against experimental data. Memote allows researchers to test a model repository offline, but we encourage and support community collaboration in reconstruction via distributed version control development platforms such as GitHub (<https://github.com/>), GitLab (<https://gitlab.com/>) or BioModels ³⁹ (<http://wwwdev.ebi.ac.uk/biomodels/>).

Either development platform supports a branching strategy (Figure 1b), which model builders could use to curate different parts of the model simultaneously or to invite external experts to improve specific model features. Memote further enables model authors to act as gatekeepers, choosing to accept only high-quality contributions. Identification of functional differences happens in the form of a comparative 'diff' report, while for the file-based discrepancies memote capitalizes on the platform's ability to show the line-by-line changes between different versions of a model. For this purpose, the model is written in a sorted YAML format ⁴⁰ after every change.

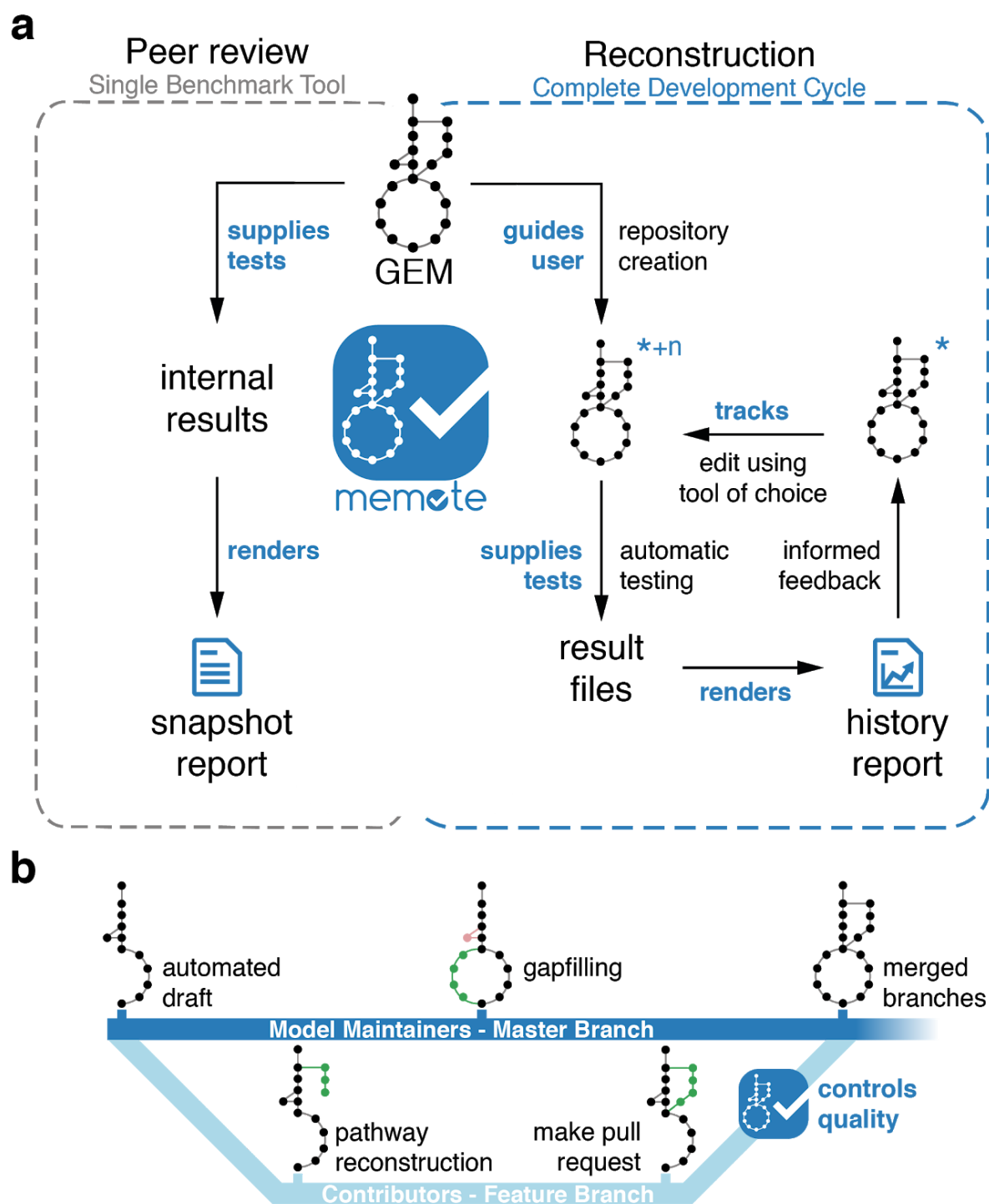


Figure 1: **Functionality offered by memote.** (a) Graphical representation of the two principal workflows in detail. For peer review, memote serves as a benchmark tool offering a quick snapshot report. For model reconstruction, memote helps the user to create a

version-controlled repository for the model (indicated by the *), and to activate continuous integration. The model is tested using memote's library of tests, the results are saved, and an initial report of the model is generated. This constitutes the first iteration of the development cycle. Now, the user may edit the model using a tool of their choice creating a new version (indicated by the +n). This will restart the cycle by running the tests automatically, saving the results for each version and including them incrementally in a history report. (b) An example of a potential branching strategy employing memote as a benchmark of external contributions. **Bold blue text** denotes actions performed by memote.

Description of the test library

The tests within memote are divided into independent core tests and tests that depend on user-supplied experimental data. Core tests are further divided into a scored and an unscored section (Figure 2).

The tests in the scored section are independent of the type of the modeled organism, the complexity of the model itself or the types of identifiers that are used to describe the model components. Calculating a score for these tests allows for the quick comparison of any two given models at a glance. The unscored section provides general statistics and covers specific aspects of a model that are not universally applicable. For instance, dedicated quality control of the biomass equation only applies to metabolic models which are used to investigate cell growth. Tests in either section belong to one of four general areas:

- 1) Basic tests give an insight into the formal correctness of a model, verifying the existence of the main model components such as metabolites, compartments, reactions, and genes. These tests also check for the presence of formula and charge

information of metabolites, and for the presence of gene-protein-reaction rules of reactions. General quality metrics such as the degree of metabolic coverage representing the ratio of reactions and genes ⁴¹ are also covered here.

- 2) Some tests are dedicated to testing the biomass reaction. This includes testing the model's ability to produce all biomass precursors in different conditions, the biomass consistency, a non-zero growth rate and direct precursors. The biomass reaction is based on the biomass composition of the modeled organism and expresses its ability to produce the necessary precursors for *in silico* cell growth and maintenance. Hence, an extensive, well-formed biomass reaction is crucial for accurate predictions with a GEM ¹⁶.
- 3) Stoichiometric inconsistency, erroneously produced energy metabolites ¹⁰ and permanently blocked reactions, are identified by testing the model's consistency. Errors here may lead to the production of ATP or redox cofactors from nothing ² and are detrimental to the performance of the model when using FBA ⁶.
- 4) Annotation tests maintain that a model is equipped according to the community standards with MIRIAM-compliant cross-references ⁴², that all primary IDs belong to the same namespace as opposed to being fractured across several namespaces, and that components are described semantically with Systems Biology Ontology terms ⁴³. A lack of explicit, standardized annotations complicates the use, comparison, and extension of GEMs, and thus strongly hampers collaboration ^{3,6,44}.

A detailed list of all the test in memote is available at

<https://github.com/opencobra/memote/wiki>.

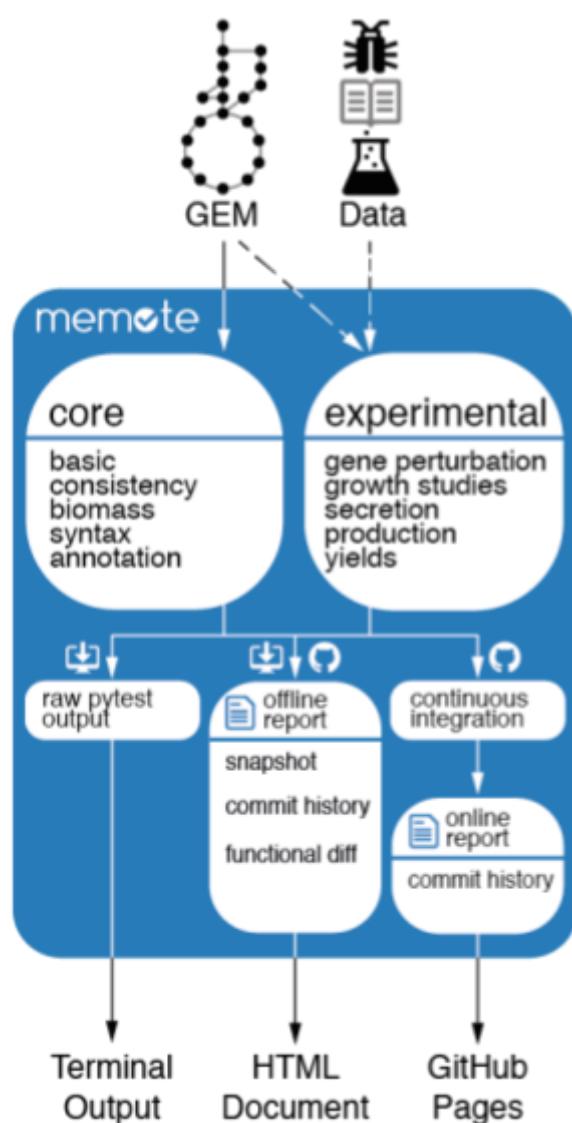


Figure 2: Functional overview of the Metabolic Model Tests (memote) package: A genome-scale metabolic model (GEM) is supplied by the user and tested in all core test categories. Optionally, if the user supplies experimental data, the model will also be subjected to the corresponding experimental tests. After testing, user input through the command line interface determines how the results are displayed. In addition to a high-level output in the terminal, the user can generate a variety of HTML-formatted reports. “Snapshot” will provide a performance benchmark of a single specified model; with a functional diff the user can benchmark two models side-by-side; and the commit-history will show the development of a model’s performance over the course of changes to a version controlled model. The latter is the type of report that is generated automatically when continuous integration is enabled. Then the results are displayed online on the project’s GitHub pages.

In addition to the core tests, researchers may supply experimental data from gene perturbation studies from a range of input formats (CSV, TSV, XLS or XLSX). Gene perturbation studies, especially gene essentiality studies are useful to refine GEM reconstructions by allowing researchers to identify network gaps and by providing a basis for model validation ⁴⁵, as well as providing grounds for a hypothesis about an organism’s physiology ⁴⁶.

To constrain the model concerning the experimental conditions underlying the supplied data, researchers may optionally define a configuration file (.yml) in which they can set the medium, FBA objective, and known regulatory effects. Without memote, this would typically be done through the use of custom scripts, which can vary significantly depending on the researcher writing them. Moreover, scripts tend to suffer from software rot if they are not actively maintained after publication²⁵. The use of configuration files instead of scripts avoids software rot since the configuration files do not require dependencies other than memote, which is likely to be maintained in the future. In conjunction, setting up a version-controlled model repository not only allows researchers to publish a ‘default’ unspecific GEM of the investigated organism, but also reproducible instructions on how to obtain a model that is specific to the organism in a defined experimental context including, and validated against the data supporting this context. This formulaic approach of deriving a GEM into a condition-specific form supports Heavner and Price’s³ call for more transparency and reproducibility in metabolic network reconstruction.

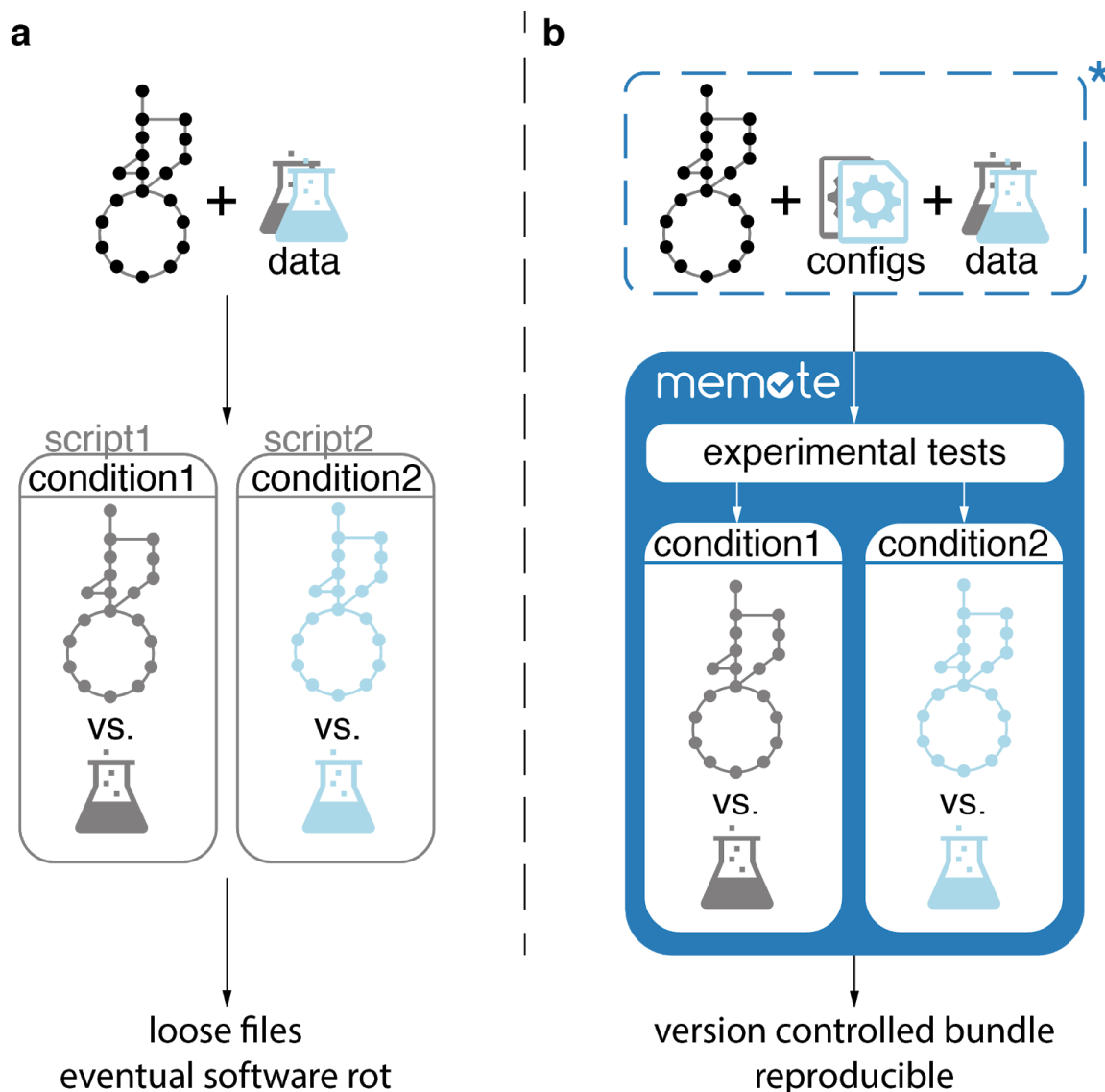


Figure 3: **Experimental tests can be tailored to a specific condition through the use of one or several configuration files (configs).** (a) To validate GEMs against experimental data measured in specific conditions, researchers usually write their scripts which constrain the model. This is problematic as scripts can vary a lot and they are, unless actively maintained, susceptible to software rot. (b) With memote, user-defined configuration files replace scripts, which allows the experimental validation of GEMs to be unified and formalized. Bundling the model, configuration files, and experimental data within a version-controlled repository (indicated by the *) supports cohesive releases.

Discussion

By providing a performance benchmark based on community guidelines and commonly-referenced SOPs, memote facilitates informed model reconstruction and quality control. The tests within memote cover semantic and conceptual requirements which are fundamental to SBML3FBC and constraint-based modeling, respectively. They are extensible to allow the validation of a model's performance against experimental data and can be executed as a stand-alone tool or integrated into existing reconstruction pipelines. Capitalizing on robust workflows established in modern software development, memote promotes openness and collaboration by granting the community tangible metrics to support their research and to discuss assumptions or limitations openly.

The concept of having a set of defined metabolic model tests is not dependent on the implementation in memote presented herein. In fact, for some platforms, it may be more desirable to implement these tests separately as this could streamline the user experience. However, an independent, central, community-maintained library of tests and a tool to run them offers 1) an unbiased approach to quality control as the tests are continuously reviewed by the community, 2) a long-lived resource as the project is independent of individual funding sources, 3) flexibility as updates can be propagated rapidly and 4) consistent results as the codebase is unified. To encourage integration as opposed to duplication, memote provides a python API as well as being available as a web-service. We plan to make memote available in the Department of Energy's Knowledge Base ⁴⁷ as an app and integrate it with the BiGG Database ³³, BioModels ³¹, and the RAVEN toolbox ⁴⁸. The memote test suite plug-in for OptFlux ⁴⁹ will approximately be released with version 3.4 scheduled for mid July 2018.

The variety of constraint-based modeling approaches and the fundamental differences between various organisms compound the assessment of GEMs. For instance, authors may publish metabolic networks, which are constrained to reflect one experimental condition or publish unconstrained metabolic databases, which need to be initialized before simulation. Both can be encoded in SBML. With having a scored test section, we attempt to normalize each type of model such that they become comparable. Despite memote's code itself being unit tested, it is difficult to anticipate all edge cases *a priori*. Also, memote depends on external resources such as MetaNetX¹² and identifiers.org⁵⁰ that are likely to change over time. Subsequently, individual users may identify potential false-positive and false-negative results. Hence, we recommend to approach the report with scrutiny and encourage users to reach out to the authors to report any errors.

The tests that memote offers only apply to stoichiometric models. However, the underlying principles and individual tests behind memote may apply to models of metabolism and expression (ME-models)⁵¹, kinetic⁵², or even systems pharmacological models⁵³.

The cloud-based distributed version control for GEMs encoded as single SBML files supported by memote is only one possible implementation approach for version control and collaboration on stoichiometric models. For instance, the reconstruction and modeling software Pathway Tools internally stores organism data in the form of a database, which can be queried and altered through the provided guided user interface and access forms⁵⁴. AuReMe, follows a similar approach, by allowing users to interact with a database through automatically generated wikis⁵⁵. While databases offer greater capacity and speed than single, large data files, the programmatic or form-based interaction required for databases may not be most immediately accessible to a broad community.

In the future, with respect to rising big data streams, memote ought to be extended to provide support for tests based on multi-omics data. Moreover, to distribute all files of a model repository together, i.e., the model, supporting data and scripts, these could be automatically bundled into one COMBINE archive file ⁵⁶, additionally including SED-ML documents which further describe relevant simulation experiments ⁵⁷.

The greater flexibility and awareness of community-driven, open-source development and the trend towards modular approaches exhibited by the solutions that were put forth in the field of systems biology ⁴⁴, motivate us to keep the development of memote open. We believe that a robust benchmark can only come to fruition when actively supported by the whole community and thus call for interested experts to involve themselves, be it through testing our tool, discussing its content or improving its implementation. We intend to keep extending memote with additional tests and functionality.

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Code availability

Memote source code is available at <https://github.com/opencobra/memote> under the Apache License, Version 2.0.

The supporting documentation is available at <https://memote.readthedocs.io/en/latest/>.

The memote web-interface is hosted at <https://memote.dd-decaf.eu>.

Author Contributions

CL, MEB and NS conceived the study. MEB, CL, SC and NS wrote the software memote.

WvH and JOV alpha-tested memote and provided early ideas for the memote report interface. CL drafted all parts of the manuscript except for the section “SBML:

Tool-independent model formulation” which was drafted by BGO and FTB. JOV helped shape the “Introduction”. PM and PV created a memote plug-in for OptFlux. JJK provided a configuration for continuous integration with Gitlab. CL, MEB, BEG, FTB, PB, JAB, LMB, SC, KC, CD, AD, BEE, JNE, AF, RMTF, BGJ, WvH, CSH, HH, MJH, HUK, ZK, JJK, SK, EK, ML, NL, DYL, SYL, SL, NEL, HM, DM, PM, AM, GLM, JMM, RM, JN, LKN, JN, IN, ORA, BØP,

JAP, KRP, NDP, AR, IR, PJS, RSMS, SS, NS, BT, PV, JOV, JAW, JCX, QY, MZ, and CZ

beta-tested memote and provided feedback and suggestions which shaped the software. All authors read, corrected and approved the manuscript.

Competing Interests

The authors declare no conflict of interest.

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