

IO Tutorial

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Reviewers:

INTRODUCTION:

This tutorial aims at providing information on how to load models into the COBRA toolbox and export them to other formats.

Available input formats

The COBRA Toolbox supports the use of models in multiple formats. Internally, it uses a simple MATLAB struct with fields defined in the [Documentation](#). These models are commonly provided in a .mat file.

There are additional model formats for which io functions exist in the Toolbox. These include

- Models in SBML Format
- Models in SimPheny format
- Models in an Excel Format

Matlab save files:

The format of a model struct provided in a .mat file has to stick to the rules defined in the [Documentation](#).

SBML Format

The COBRA Toolbox currently supports SBML models provided as Level 3 version 1 (as defined [here](#)) and has legacy support for older versions of SBML. It also supports the Level 3 FBC package (both in version 1 and version 2).

The Toolbox will use the provided SBML IDs as IDs for the respective elements of the model structure, and use the name fields as names. It is assumed (but not necessary), that metabolite IDs start with a "M_", reaction ids start with a "R_" , gene ids start with a "G_" and compartment ids start with a "C_". This is due to the limitation on Identifiers in SBML and those starting sequences will be removed if they are consistently present in the model. Since metabolites in the COBRA Toolbox model struct are commonly provided with a metabolite ID followed by a compartment identifier e.g. ('ala_L[c]'), and brackets are illegal characters for an SBML id, it is assumed, that if all non boundary species have a trailing compartment identifier preceded by an underscore (e.g. SBML ID: M_ala_L_c) those identifiers should be converted to model compartment identifiers.

The Toolbox has a legacy support for the NOTE Fields defined in [Schellenberger et al, Nature Protocols, 2011](#), but it is suggested to instead use annotations whenever possible. In general, if a fbc-package field and a NOTES field is present, the fbc-package value will be used (e.g. CHARGE for metabolites, or GENE_ASSOCIATION for reactions). The same applies to annotations, i.e. if there is an annotation for an EC number, the Notes field EC Number will be ignored. However, the charge field in SBML Level 2 will be overwritten by the Notes field definitions.

SimPheny Format

SimPheny models provide their models in 3 or 4 files (4 if GPR rules are provided). The model identifiers will be used as presented in the SimPheny files.

Excel Format

Excel files adhering to the COBRA xls specifications listed in the [Documentation](#). Parseable excel models contain two sheets for reactions and metabolites respectively.

Available output formats

The toolbox also allows storage in multiple file types as detailed below:

Matlab .mat files

This format is simply the model being saved as a matlab save file, and is the default save method. It has the advantage of lossless data storage even for model specific fields not supported by the Toolbox.

SBML format (.xml)

The systems biology markup language (SBML) is a very common format to store biological models. The COBRA Toolbox allows generation of models using Level 3 Version 1 and uses the fbc-package extension to encode constraint based properties. This is the format that is recommended for publication, as it can be used by many different tools and allows the best use of the model.

Excel format

Historically, models were often exchanged using excel files, and this is still in use today. Some users prefer to get an overview over a model using Excel, since it provides a general overview at a first glance. The toolbox offers an Excel export using the format described in the Documentation (see above).

Text Format

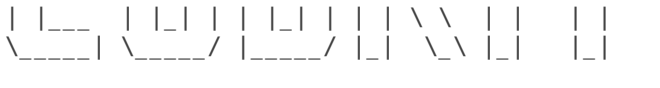
Finally the toolbox offers a simple textual export, which is essentially a tab separated file containing the reactions with their reaction formulas along with the associated GPRs, but no further information. This format only uses the required fields and will ignore any optional fields.

MATERIALS

We will use two model files for this tutorial, one matlab mat file and an xml file stored in SBML format. The following code loads these files into the tutorial directory (cleaning any old copies).

```
cd(fileparts(which('tutorial_I0.mlx')));
```





Documentation:
<http://opencobra.github.io/cobratoolbox>

```
> Checking if git is installed ... Done.
> Checking if the repository is tracked using git ... Done.
> Checking if curl is installed ... Done.
> Checking if remote can be reached ... Done.
> Initializing and updating submodules ... Done.
> Adding all the files of The COBRA Toolbox ... Done.
> Define CB map output... set to svg.
> Retrieving models ... Done.
> TranslateSBML is installed and working properly.
> Configuring solver environment variables ...
- [---*] ILOG_CPLEX_PATH: /opt/ibm/ILOG/CPLEX_Studio1271/cplex/matlab/x86-64_linux
- [--*-] GUROBI_PATH: /home/thomas/Downloads/gurobi701/linux64/matlab
- [----] TOMLAB_PATH : --> set this path manually after installing the solver ( see instructions )
- [----] MOSEK_PATH : --> set this path manually after installing the solver ( see instructions )
Done.
> Checking available solvers and solver interfaces ... Done.
> Setting default solvers ... Done.
> Saving the MATLAB path ... Done.
- The MATLAB path was saved as ~/pathdef.m.
```

> Summary of available solvers and solver interfaces

Support	LP	MILP	QP	MIQP	NLP		
cplex_direct	full			0	0	0	-
dqqMinos	full			1	-	-	-
glpk	full			1	1	-	-
gurobi	full			1	1	1	-
ibm_cplex	full			1	1	1	-
matlab	full			1	-	-	1
mosek	full			0	0	0	-
pdco	full			1	-	1	-
quadMinos	full			1	-	-	1
tomlab_cplex	full			0	0	0	-
qpng	experimental			-	-	1	-
tomlab_snopt	experimental			-	-	-	0
gurobi_mex	legacy			0	0	0	-
lindo_old	legacy			0	-	-	-
lindo_legacy	legacy			0	-	-	-
lp_solve	legacy			1	-	-	-
opti	legacy			0	0	0	0
Total	-			8	3	4	2

+ Legend: - = not applicable, 0 = solver not compatible or not installed, 1 = solver installed.

```
> You can solve LP problems using: 'dqqMinos' - 'glpk' - 'gurobi' - 'ibm_cplex' - 'matlab' - 'pdco' - 'quadMinos'
> You can solve MILP problems using: 'glpk' - 'gurobi' - 'ibm_cplex'
> You can solve QP problems using: 'gurobi' - 'ibm_cplex' - 'pdco' - 'qpng'
> You can solve MIQP problems using: 'gurobi' - 'ibm_cplex'
> You can solve NLP problems using: 'matlab' - 'pdco' - 'quadMinos'

> Checking for available updates ...
--> You cannot update your fork using updateCobraToolbox(). [80828f @ IOTutorialAndUpdate].
Please use the MATLAB.devTools (https://github.com/opencobra/MATLAB.devTools).
```

%Copy two files that can be loaded (if they are nto yet present).

```
try
delete 'ecoli_core_model.mat';
delete 'Abiotrophia_defectiva_ATCC_49176.xml'
copyfile(which('ecoli_core_model.mat'), '.');
```

```
    copyfile(which('Abiotrophia_defectiva_ATCC_49176.xml'), '.');  
end
```

PROCEDURE

The time it takes to load a model depends on the file format and the complexity of a model. While mat file loading even for large models is a question of seconds, very large SBML files can take a few minutes to load depending on the machine.

Reading a Model (TIMING 1s to a few minutes)

The most straightforward way to import a model into the Toolbox is to use the readCbModel function. To e.g. load a model from a mat file, you can simply use the filename (with or without file extension).

```
fileName = 'ecoli_core_model'
```

```
fileName = Acidaminococcus_intestini_RyC_MR95
```

```
model = readCbModel(fileName);
```

readCbModel assumes, that .mat files are a matlab save file, .xml files are models in SBML format, .sto are SimPheny models, and .xls or .xlsx are models in Excel format.

```
%This code is to avoid execution in non gui-environments  
if usejava('desktop')
```

You can also call the function without a fileName to get a file selection dialog

```
model = readCbModel();
```

The model loaded in this way can directly be used with COBRA ToolBox functions. You can view the data stored in the model by e.g. using

```
open model
```

```
%This code is to avoid execution in non gui-environments  
end
```

Writing a model (TIMING: 1s to a few minutes)

```
%This code is to avoid execution in non gui-environments  
if usejava('desktop')
```

To write files, use the writeCbModel function. The function can be called directly with a model.

```
writeCbModel(model)
```

This will call a file selection dialog, which allows the selection of a filename and, depending on the selected format from the dropdown, the output will be generated.

```
%This code is to avoid execution in non gui-environments  
end
```

```
ans =  
  
      S: [952×1069 double]  
      b: [952×1 double]  
  csense: [952×1 char]  
      lb: [1069×1 double]  
      ub: [1069×1 double]  
      c: [1069×1 double]  
  osense: -1  
    rxns: {1069×1 cell}  
    mets: {952×1 cell}  
   genes: {598×1 cell}  
   rules: {1069×1 cell}  
description: '/home/thomas/cobra_devel/fork-cobratoolbox/tutorials/I0/Abiotrophia_defectiva_  
   grRules: {1069×1 cell}  
 metCharges: [952×1 double]  
metFormulas: {952×1 cell}  
   metNames: {952×1 cell}  
modelVersion: [1×1 struct]  
  rxnGeneMat: [1069×598 double]  
   rxnNames: {1069×1 cell}  
   rxnNotes: {1069×1 cell}  
 subSystems: {1069×1 cell}  
  metChEBIID: {952×1 cell}  
   methMDBID: {952×1 cell}  
metInChIString: {952×1 cell}  
   metKEGGID: {952×1 cell}  
   metPubChemID: {952×1 cell}  
rxnConfidenceScores: {1069×1 cell}  
   rxnECNumbers: {1069×1 cell}  
   rxnReferences: {1069×1 cell}
```

```
%This code is to avoid execution in non gui-environments  
if usejava('desktop')
```

If no fileName is provided, a popup will ask you to provide a fileName with the specified format.

```
writeCbModel(model, 'text')
```

The available format options are:

- 'mat' - for a matlab save file
- 'sbml' - for a SBML model
- 'xls' - for a model in Excel format
- 'text' - for a textual export.

```
%This code is to avoid execution in non gui-environments  
end
```

```
ans =  
  
      S: [952×1069 double]
```

```

        b: [952×1 double]
        csense: [952×1 char]
        lb: [1069×1 double]
        ub: [1069×1 double]
        c: [1069×1 double]
        osense: -1
        rxns: {1069×1 cell}
        mets: {952×1 cell}
        genes: {598×1 cell}
        rules: {1069×1 cell}
        description: '/home/thomas/cobra_devel/fork-cobratoolbox/tutorials/I0/Abiotrophia_defectiva_
        grRules: {1069×1 cell}
        metCharges: [952×1 double]
        metFormulas: {952×1 cell}
        metNames: {952×1 cell}
        modelVersion: [1×1 struct]
        rxnGeneMat: [1069×598 double]
        rxnNames: {1069×1 cell}
        rxnNotes: {1069×1 cell}
        subSystems: {1069×1 cell}
        metChEBIID: {952×1 cell}
        metHMDBID: {952×1 cell}
        metInChIString: {952×1 cell}
        metKEGGID: {952×1 cell}
        metPubChemID: {952×1 cell}
        rxnConfidenceScores: {1069×1 cell}
        rxnECNumbers: {1069×1 cell}
        rxnReferences: {1069×1 cell}

```

It is also possible to specify the format and filename explicitly:

```
writeCbModel(model, 'SBML', 'Acidaminococcus.xml')
```

```
ans =
    1
```

Document written

```

ans =
    constraint: [1×0 struct]
    functionDefinition: [1×0 struct]
    event: [1×0 struct]
    rule: [1×0 struct]
    unitDefinition: [1×1 struct]
    initialAssignment: [1×0 struct]
    SBML_level: 3
    SBML_version: 1
    annotation: ''
    areaUnits: ''
    avogadro_symbol: ''
    conversionFactor: ''
    delay_symbol: ''
    extentUnits: ''
    fbc_activeObjective: 'obj'
    fbc_version: 2
    id: 'COBRAModel'
    lengthUnits: ''
    metaid: 'COBRAModel'
    name: ''
    notes: ''
    sboTerm: -1
    substanceUnits: ''
    timeUnits: ''
    time_symbol: ''

```

```
        typecode: 'SBML_MODEL'
    volumeUnits: ''
    species: [1×952 struct]
    compartment: [1×2 struct]
    parameter: [1×15 struct]
    reaction: [1×1069 struct]
    fbc_fluxBound: [1×2 struct]
    fbc_geneProduct: [1×598 struct]
    fbc_objective: [1×1 struct]
    namespaces: [1×2 struct]
    fbc_strict: 1
```

which will write the model to the file Acidaminococcus.xml.

```
%Some Cleanup
currentDir = pwd;
cd(fileparts(which('tutorial_I0.mlx')));

%Copy two files that can be loaded (if they are nto yet present).
try
    delete 'ecoli_core_model.mat';
    delete 'Abiotrophia_defectiva_ATCC_49176.xml'
    delete 'Acidaminococcus.xml'
end
cd(currentDir)
```