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# Pycotools Documentation

## *Release 1*

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PyCoTools is a python package that was developed as an alternative interface into [COPASI](#), simulation software for modelling biochemical systems. The PyCoTools paper can be found [here](#) and describes in detail the intentions and functionality of PyCoTools. There are some important differences between the PyCoTools version that is described in the publication and the current version. The first is that PyCoTools is now a python 3 only package. If using Python 2.7 you should create a virtual Python 3.6 environment using [conda](#) or [virtualenv](#). My preference is conda. The other major difference is the interface to COPASI's parameter estimation task which is described in the tutorials and examples.



# CHAPTER 1

---

## Installation

---

First make sure you use a Python 3.6 environment.

**Warning:** Using Python 3.7 or 3.8 will not work at this time due to dependency issues (which are unfortunately out of my control).

Then use:

```
$ pip install pycotools3
```

Remember to `source activate` your python 3.6 environment if you need to.

To install from [source](#):

```
$ git clone https://github.com/CiaranWelsh/pycotools3.git
$ cd pycotools3
$ python setup.py install
```

The procedure is the same in linux, mac and windows.



# CHAPTER 2

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## Troubleshooting

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### PyCoTools3 will not install Pycotools3 is only supported in Python 3 to Python 3.6. If you are using Python 2.7 or Python 3.7 please create a new conda Python3.7 environment.

```
$ conda create -n py36 python=3.6
$ conda activate py36
$ pip install pycotools3
```

The same commands should work cross platform.

### You get errors when trying to build a model using pycotools3.model.loada () Make sure you have installed *Copasi* <<http://copasi.org/Download/>> and added the *Copasi/bin* directory to the path variable.

- On Linux <<https://unix.stackexchange.com/questions/26047/how-to-correctly-add-a-path-to-path>>
- On Windows <<https://stackoverflow.com/questions/9546324/adding-directory-to-path-environment-variable-in-windows>>
- On Mac <<https://stackoverflow.com/questions/7703041/editing-path-variable-on-mac>>



# CHAPTER 3

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## Documentation

---

This is a guide to PyCoTools version >2.0.1.

### 3.1 Getting started

As PyCoTools only provides an alternative interface into some of COPASI's tasks, if you are unfamiliar with [COPASI](#), it is a good idea to become acquainted prior to proceeding. As much as possible, arguments to PyCoTools functions follow the corresponding option in the COPASI user interface.

In addition to COPASI, PyCoTools depends on [tellurium](#) which is a Python package for modelling biological systems. While tellurium and COPASI have some of the same features, generally they are complementary and productivity is enhanced by using both together.

More specifically, tellurium uses [antimony strings](#) to define a model which is then converted into SBML. PyCoTools provides the `model.BuildAntimony` class which is a wrapper around this tellurium feature, which creates a Copasi model and parses it into a PyCoTools `model.Model`.

Since antimony is described [elsewhere](#) we will focus here on using antimony to build a copasi model.

#### 3.1.1 Build a model with antimony

```
[2]: import site, os
from pycotools3 import model

working_directory = os.path.abspath('') #create a base directory to ↴work from
```

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```

copasi_filename = os.path.join(working_directory,
    'NegativeFeedbackModel.cps') #create a string to a copasi file on_
    ↪system
antimony_string = '''
    model negative_feedback()
        // define compartments
        compartment cell = 1.0
        //define species
        var A in cell
        var B in cell
        //define some global parameter for use in reactions
        vAProd = 0.1
        kADeg = 0.2
        kBProd = 0.3
        kBDeg = 0.4
        //define initial conditions
        A      = 0
        B      = 0
        //define reactions
        APProd: => A; cell*vAProd
        ADeg: A => ; cell*kADeg*A*B
        BProd: => B; cell*kBProd*A
        BDeg: B => ; cell*kBDeg*B
    end
    '''

negative_feedback = model.loada(antimony_string, copasi_filename) #_
    ↪load the model
print(negative_feedback)
assert os.path.isfile(copasi_filename) #check that the model exists
Model(name=negative_feedback, time_unit=s, volume_unit=l, quantity_
    ↪unit=mol)

```

## 3.1.2 Create an antmiony string from an existing model

The Copasi user interface is an excellant way of constructing a model and it is easy to convert this model into an antimony string that can be pasted into a document.

```
[4]: print(negative_feedback.to_antimony())
// Created by libAntimony v2.9.4
function Constant_flux_irreversible(v)
    v;
end

function Function_for_ADeg(A, B, kADeg)
    kADeg*A*B;
end
```

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```

function Function_for_BProd(A, kBProd)
    kBProd*A;
end

model *negative_feedback()

    // Compartments and Species:
    compartment cell;
    species A in cell, B in cell;

    // Reactions:
    APProd:  ==> A; cell*Constant_flux_irreversible(vAPProd);
    ADeg: A ==> ; cell*Function_for_ADeg(A, B, kADeg);
    BProd:  ==> B; cell*Function_for_BProd(A, kBProd);
    BDeg: B ==> ; cell*kBDeg*B;

    // Species initializations:
    A = 0;
    B = 0;

    // Compartment initializations:
    cell = 1;

    // Variable initializations:
    vAPProd = 0.1;
    kADeg = 0.2;
    kBProd = 0.3;
    kBDeg = 0.4;

    // Other declarations:
    const cell, vAPProd, kADeg, kBProd, kBDeg;
end

```

One paradigm of model development is to use antimony to ‘hard code’ permanent changes to the model and the Copasi user interface for experimental changes. The `Model.open()` method is useful for this paradigm as it opens the model with whatever configurations have been defined.

[5] : `negative_feedback.open()`

### 3.1.3 Simulate a time course

Since we have used an antimony string, we can simulate this model with either tellurium or Copasi. Simulating with tellurium uses a library called roadrunner which is described in detail [elsewhere](#). To run a simulation with Copasi we need to configure the time course task, make the task executable (i.e. tick the check box in the top right of the time course task) and run the

simulation with CopasiSE. This is all taken care of by the `tasks.TimeCourse` class.

```
[6]: from pycotools3 import tasks
time_course = tasks.TimeCourse(negative_feedback, end=100, step_
    ↪size=1, intervals=100)
time_course
[6]: <pycotools3.tasks.TimeCourse at 0x1ff411e5588>
```

However a more convenient interface is provided by the `model.simulate` method, which is a wrapper around `tasks.TimeCourse` which additionally parses the resulting data from file and returns a `pandas.DataFrame`

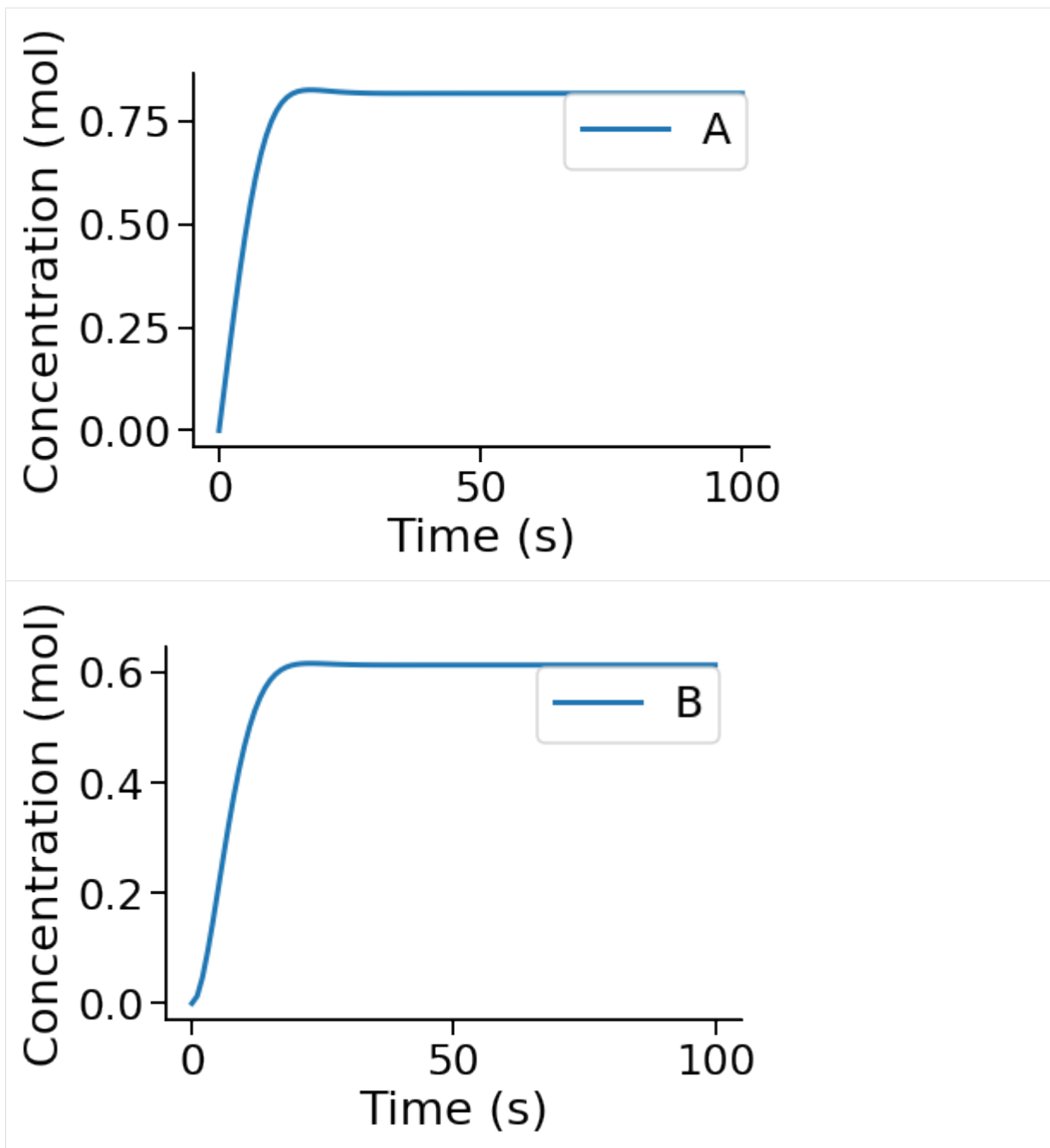
```
[7]: from pycotools3 import tasks
fname = os.path.join(os.path.abspath(''), 'timcourse.txt')
sim_data = negative_feedback.simulate(0, 100, 1, report_name=fname)
    ↪##start, end, by
sim_data.head()
[7]:          A          B
Time
0      0.000000  0.000000
1      0.099932  0.013181
2      0.199023  0.046643
3      0.295526  0.093275
4      0.387233  0.147810
```

The results are saved in a file defined by the `report_name` option, which defaults to `timcourse.txt` in the same directory as the copasi model.

## 3.1.4 Visualise a time course

PyCoTools also provides facilities for visualising simulation output. To plot a timecourse, pass the `task.TimeCourse` object to the `viz.PlotTimeCourse` object.

```
[8]: from pycotools3 import viz
viz.PlotTimeCourse(time_course, savefig=True)
[8]: <pycotools3.viz.PlotTimeCourse at 0x1ff411e53c8>
```



More information about running time courses with PyCoTools and Copasi can be found in the [time course tutorial](#)

### 3.1.5 Run Parameter Estimation

The following configures a regular copasi parameter estimation (`context='s'`) on all global and initial concentration parameters (`parameters='gm'`) using the genetic algorithm

```
[15]: from pycotools3 import tasks, viz

with tasks.ParameterEstimation.Context(negative_feedback, fname,
    context='s', parameters='gm') as context:
```

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```

context.set('randomize_start_values', True)
context.set('method', 'genetic_algorithm')
context.set('population_size', 50)
context.set('number_of_generations', 300)
context.set('run_mode', True) ##defaults to False
context.set('pe_number', 2) ## number of repeat items in scan
→task
    context.set('copy_number', 2) ## number of times to copy model
config = context.get_config()

pe = tasks.ParameterEstimation(config)
data = viz.Parse(pe)
print(data)

{'NegativeFeedbackModel': A B RSS kADeg
 → kBDeg kBProd vAProd
0 0.000004 0.000321 0.000348 0.198104 0.404187 0.298985 0.
→099467
1 0.000397 0.000009 0.000358 0.201986 0.396722 0.296337 0.
→100254
2 0.002442 0.000002 0.000399 0.197253 0.403480 0.299229 0.
→099694
3 0.110722 0.004461 0.002555 0.198449 0.402032 0.297208 0.
→097885}

```

pycotools3 supports the configuration of:

- *multiple models at once*
- *multiple parameter estimation repeats at once*
- *profile likelihoods*
- *cross validations*

Also you can checkout the [parameter estimation tutorial](#).

## 3.2 Tutorials

In this section of the documentation I provide detailed explanations of how PyCoTools works, with examples. The tutorials are split into sections which are linked to below.

### 3.2.1 Running Time-Courses

Copasi enables users to simulate their model with a range of different solvers.

## Create a model

Here we do our imports and create the model we use for the tutorial

```
[1]: import os
import site
from pycotools3 import model, tasks, viz

working_directory = r'/home/ncw135/Documents/pycotools3/docs/source/
˓→Tutorials/timecourse_tutorial'
if not os.path.isdir(working_directory):
    os.makedirs(working_directory)

copasi_file = os.path.join(working_directory, 'michaelis_menten.cps
˓→')

if os.path.isfile(copasi_file):
    os.remove(copasi_file)

antimony_string = """
model michaelis_menten()
    compartment cell = 1.0
    var E in cell
    var S in cell
    var ES in cell
    var P in cell

    kf = 0.1
    kb = 1
    kcat = 0.3
    E = 75
    S = 1000

    SBindE: S + E => ES; kf*S*E
    ESUnbind: ES => S + E; kb*ES
    ProdForm: ES => P + E; kcat*ES
end
"""

with model.BuildAntimony(copasi_file) as loader:
    mm = loader.load(antimony_string)

mm
The BuildAntimony context manager is deprecated and will be removed
˓→in future versions. Please use model.loada instead.

[1]: Model(name=michaelis_menten, time_unit=s, volume_unit=l, quantity_
˓→unit=mol)
```

## Deterministic Time Course

```
[2]: TC = tasks.TimeCourse(
    mm, report_name='mm_simulation.txt',
    end=1000, intervals=50, step_size=20
)

## check its worked
os.path.isfile(TC.report_name)

import pandas
df = pandas.read_csv(TC.report_name, sep='\t')
df.head()

[2]:   Time      [E]      [S]      [ES]      [P]  Values[kf]
  ↗Values[kb] \
0     0  75.00000  1000.00000  1.000000  1.000      0.1
  ↗
1    20  2.00306  479.797000  73.996900  448.206      0.1
  ↗
2    40  12.80010   62.104800  63.199900  876.695      0.1
  ↗
3    60  75.13160      0.119830  0.868371  1001.010      0.1
  ↗
4    80  75.99560      0.000604  0.004434  1001.990      0.1
  ↗

      Values[kcat]
0            0.3
1            0.3
2            0.3
3            0.3
4            0.3
```

When running a time course, you should ensure that the number of intervals times the step size equals the end time, i.e.:

```
- $$intervals \cdot step\_size = end$$
```

The default behaviour is to output all model variables as they can easily be filtered later in the Python environment. However, the `metabolites`, `global_quantities` and `local_parameters` arguments exist to filter the variables that are simulated prior to running the time course.

```
[3]: TC=tasks.TimeCourse(
    mm,
    report_name='mm_timecourse.txt',
    end=100,
    intervals=50,
    step_size=2,
```

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```
global_quantities = ['kf'], ##recall that antimony puts all
#parameters as global quantities
)

##check that we only kf as a global variables
pandas.read_csv(TC.report_name, sep='\t').head()
```

	Time	[E]	[S]	[ES]	[P]	Values[kf]
0	0	75.00000	1000.000	1.0000	1.0000	0.1
1	2	1.10437	881.378	74.8956	45.7263	0.1
2	4	1.16265	836.516	74.8373	90.6465	0.1
3	6	1.22737	791.698	74.7726	135.5300	0.1
4	8	1.29962	746.928	74.7004	180.3720	0.1

An alternative and more convenient interface into the `tasks.TimeCourse` class is using the `model.Model.simulate` method. This is simply a wrapper and is used like so.

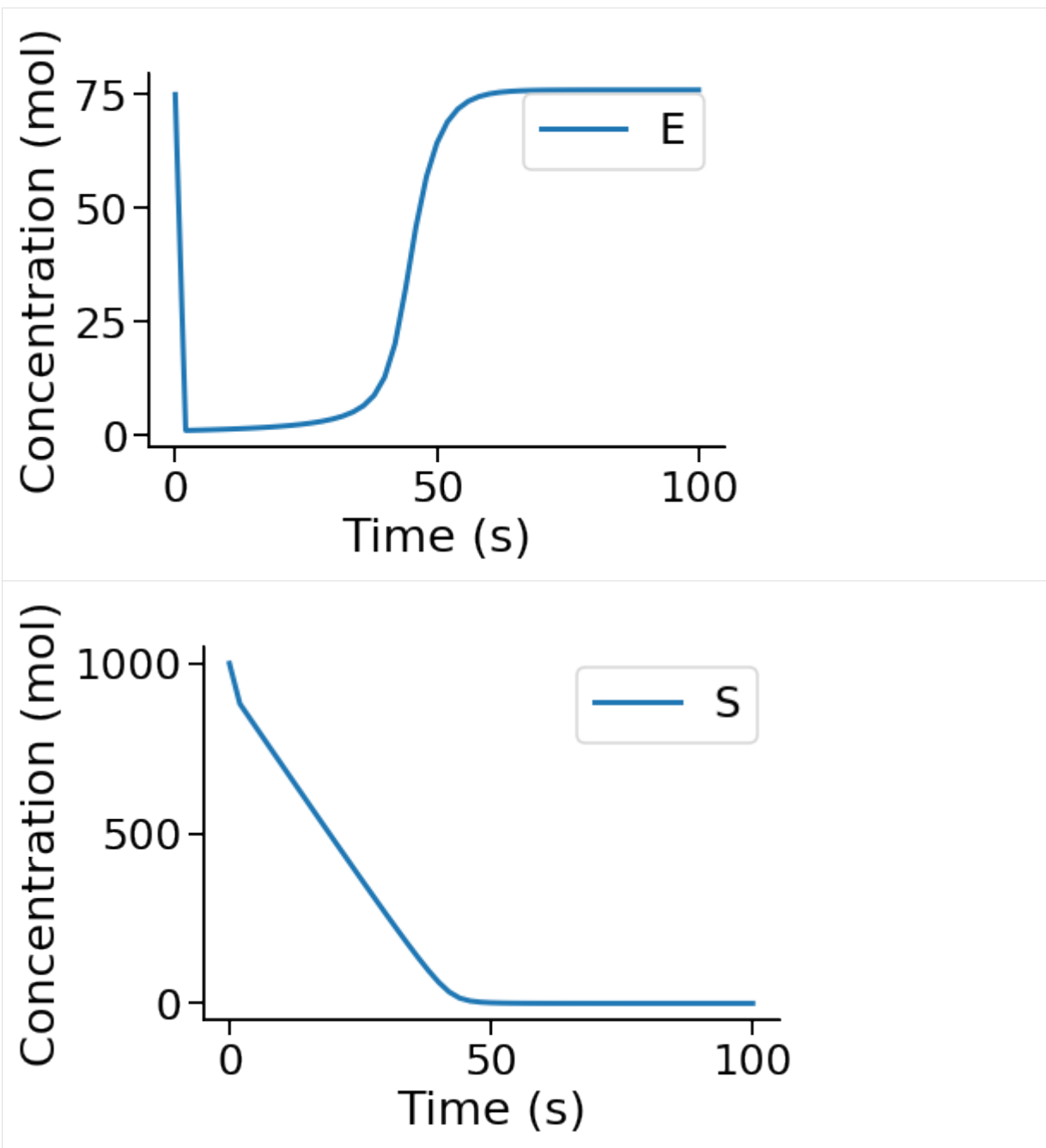
```
[4]: data = mm.simulate(start=0, stop=100, by=0.1)
data.head()
```

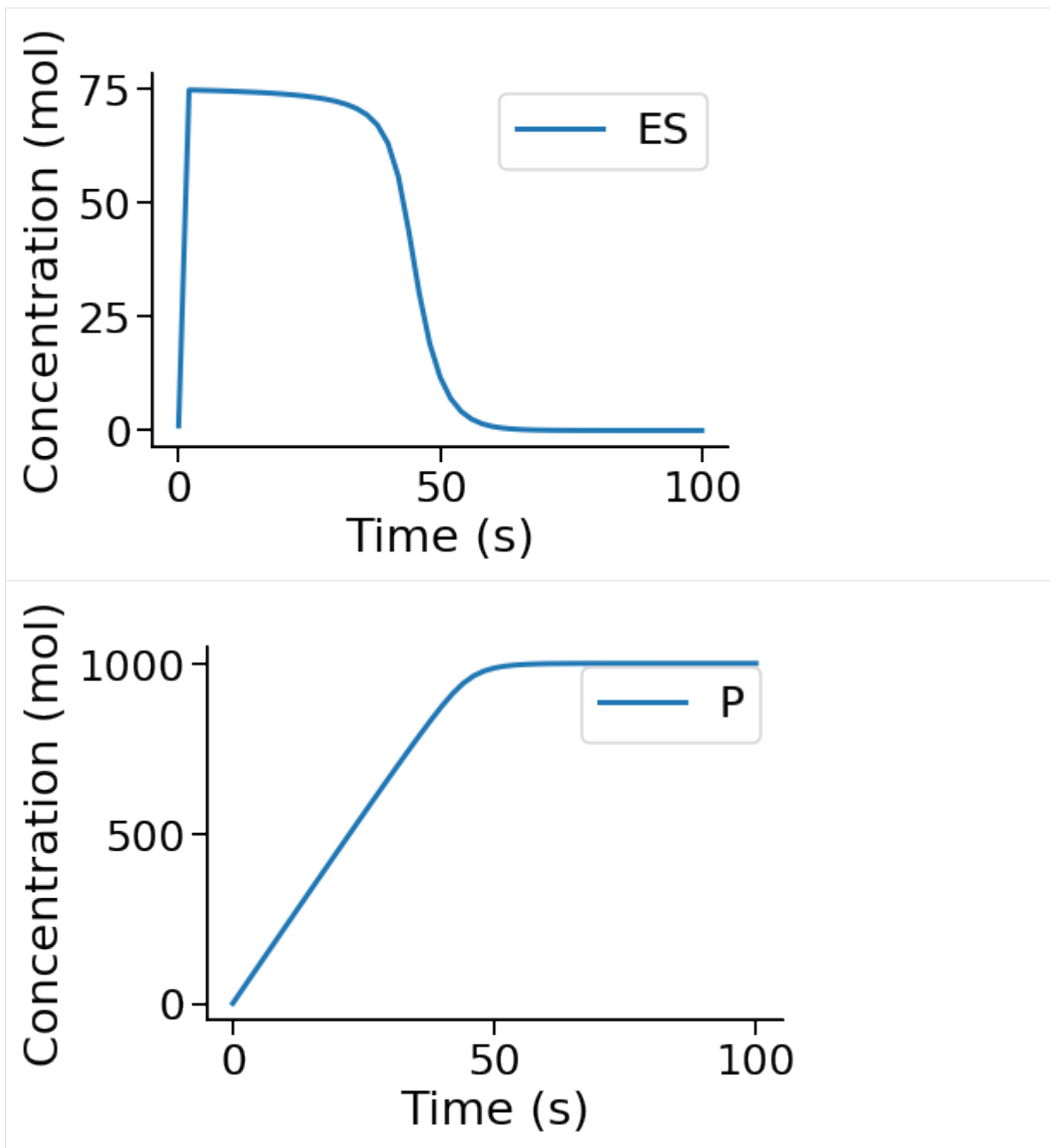
	E	ES	P	S
Time				
0.0	75.00000	1.0000	1.00000	1000.000
0.1	1.05984	74.9402	3.02124	924.039
0.2	1.05665	74.9433	5.26956	921.787
0.3	1.05920	74.9408	7.51783	919.541
0.4	1.06175	74.9382	9.76601	917.296

This mechanism of running a time course has the advantage that 1) pycotools parses the data back into python in the form of a `pandas.DataFrame` and 2) the column names are automatically pruned to remove the copasi reference information.

## Visualization

```
[5]: viz.PlotTimeCourse(TC)
[5]: <pycotools3.viz.PlotTimeCourse at 0x16c80294358>
```

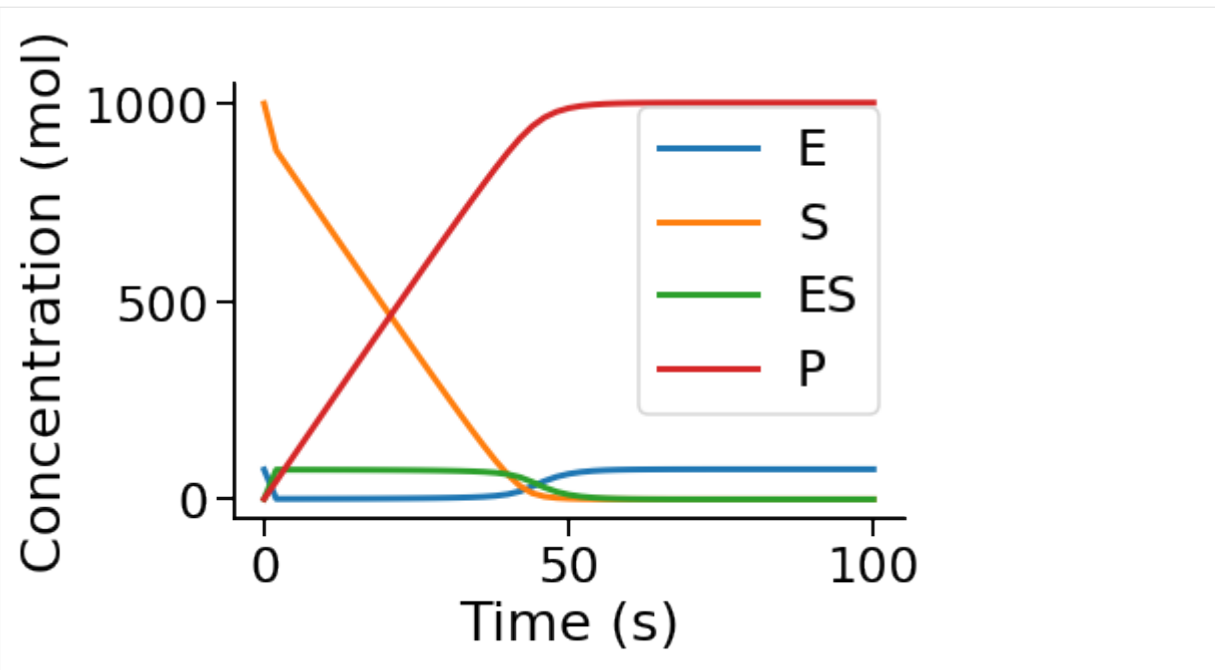




It is also possible to plot these on the same axis by specifying `separate=False`

```
[6]: viz.PlotTimeCourse(TC, separate=False)
```

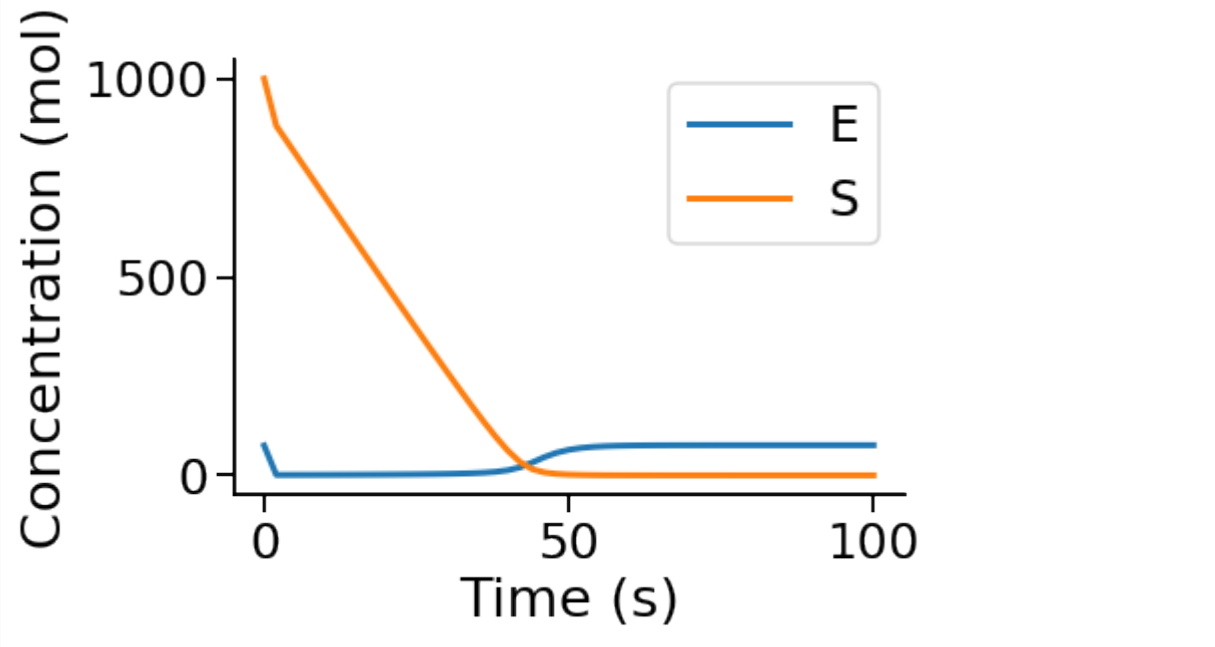
```
[6]: <pycotools3.viz.PlotTimeCourse at 0x16ca06f91d0>
```

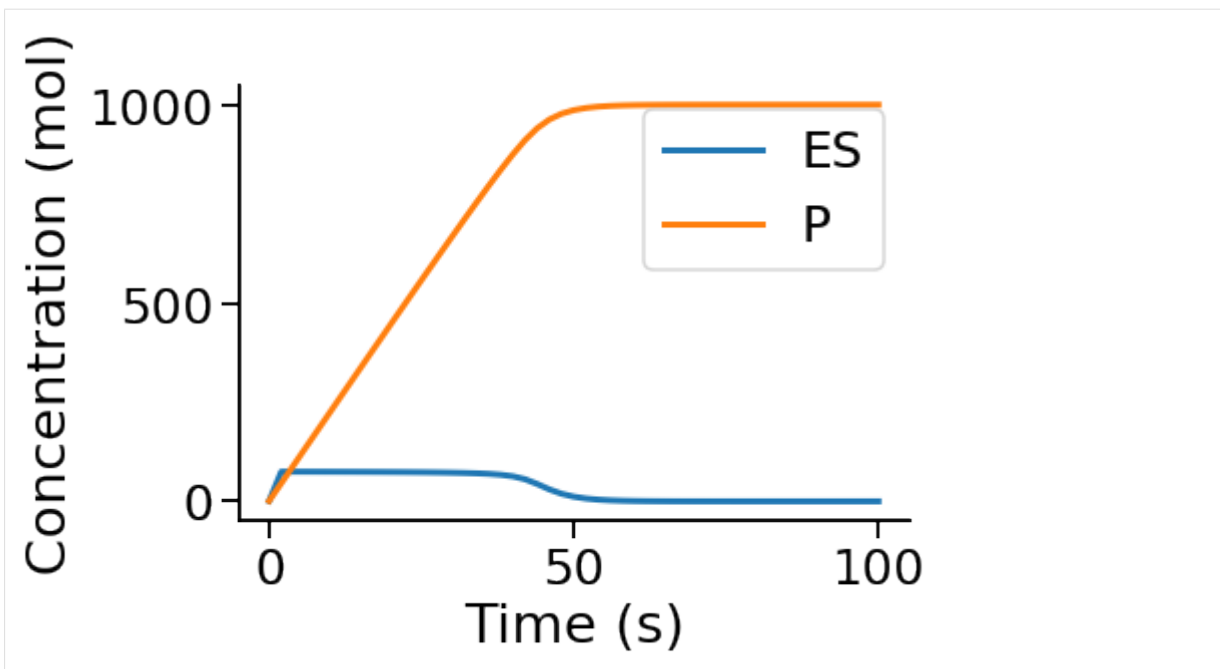


or to choose the y variables,

```
[7]: viz.PlotTimeCourse(TC, y=['E', 'S'], separate=False)
viz.PlotTimeCourse(TC, y=['ES', 'P'], separate=False)
```

[7]: <pycotools3.viz.PlotTimeCourse at 0x16ca0760748>

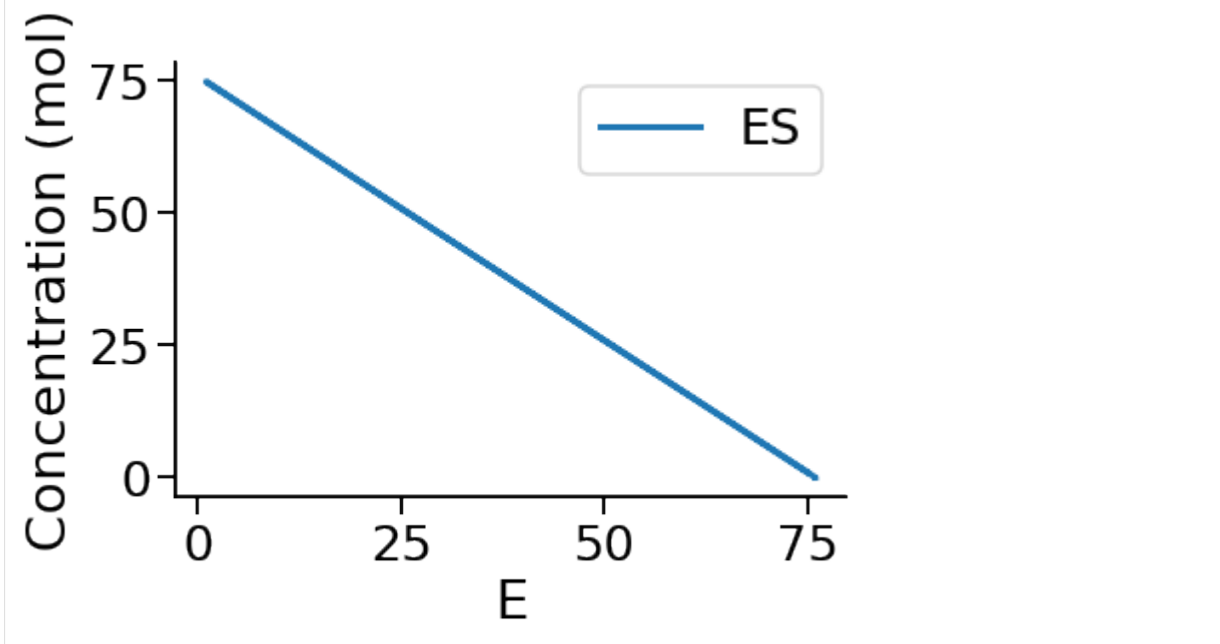




### Plot in Phase Space

Choose the x variable to plot phase space. Same arguments apply as above.

```
[8]: viz.PlotTimeCourse(TC, x='E', y='ES', separate=True)  
[8]: <pycotools3.viz.PlotTimeCourse at 0x16ca07c1b38>
```

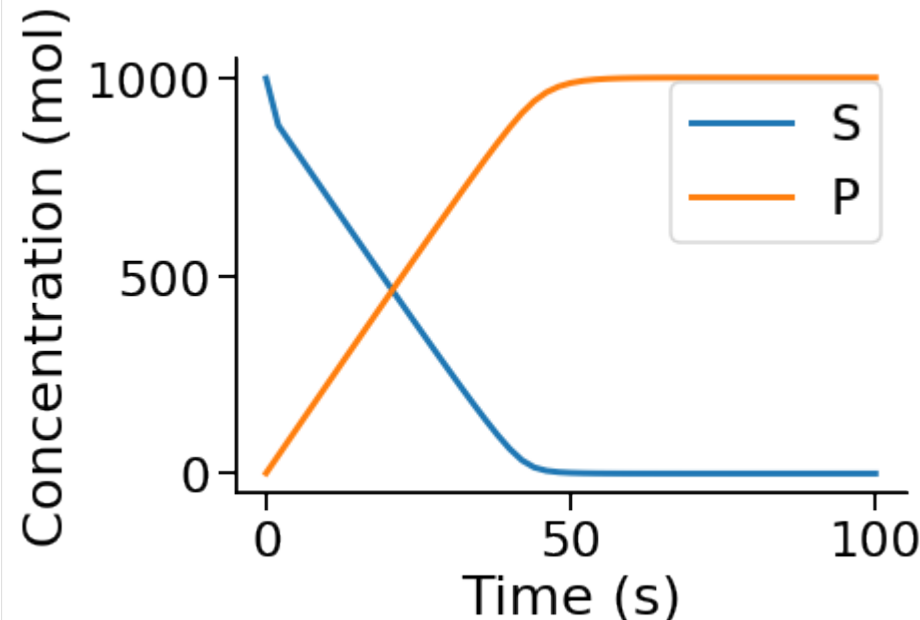


## Save to file

Use the `savefig=True` option to save the figure to file and give an argument to the `filename` option to choose the filename.

```
[9]: viz.PlotTimeCourse(TC, y=['S', 'P'], separate=False, savefig=True,   
 →filename='MyTimeCourse.png')
```

```
[9]: <pycotools3.viz.PlotTimeCourse at 0x16ca0847cf8>
```



## Alternative Solvers

Valid arguments for the `method` argument of `TimeCourse` are:

- deterministic
- direct
- gibson\_bruck
- tau\_leap
- adaptive\_tau\_leap
- hybrid\_runge\_kutta
- hybrid\_lsoda

Copasi also includes a `hybrid_rk45` solver but this is not yet supported by Pycotools. To use an alternative solver, pass the name of the solver to the `method` argument.

## Stochastic MM

For demonstrating simulation of stochastic time courses we build another michaelis-menten type reaction schema. We need to do this so we can set `unit substance = item`, or in other words, change the model to particle numbers - otherwise there are too many molecules in the system to simulate a stochastic model

```
[10]: copasi_file = os.path.join(working_directory, 'michaelis_menten_
        ↪stochastic.cps')

antimony_string = """
model michaelis_menten()
    compartment cell = 1.0;
    var E in cell;
    var S in cell;
    var ES in cell;
    var P in cell;

    kf = 0.1;
    kb = 1;
    kcat = 0.3;
    E = 75;
    S = 1000;

    SBindE: S + E => ES; kf*S*E;
    ESUnbind: ES => S + E; kb*ES;
    ProdForm: ES => P + E; kcat*ES;

    unit substance = item;

end
"""

with model.BuildAntimony(copasi_file) as loader:
    mm = loader.load(antimony_string)
```

The `BuildAntimony` context manager is deprecated and will be removed  
in future versions. Please use `model.loada` instead.

## Run a Time Course Using Direct Method

```
[11]: data = mm.simulate(0, 100, 1, method='direct')
data.head(n=10)
```

	E	ES	P	S
Time				
0	75	1	1	1000
1	0	76	18	908
2	2	74	36	892

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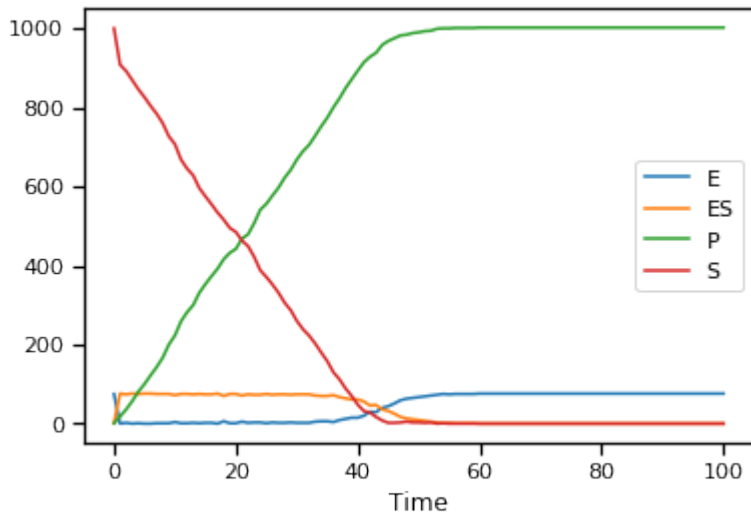
(continued from previous page)

3	0	76	57	869
4	1	75	81	846
5	0	76	100	826
6	0	76	122	804
7	1	75	143	784
8	1	75	167	760
9	1	75	200	727

## Plot stochastic time course

Note that we can also use the `pandas`, `matplotlib` and `seaborn` libraries for plotting

```
[12]: import matplotlib
import seaborn
seaborn.set_context('notebook')
data.plot()
[12]: <matplotlib.axes._subplots.AxesSubplot at 0x16ca0810c88>
```



Notice how similar the stochastic simulation is to the deterministic. As the number of molecules being simulated increases, the stochastic simulation converges to the deterministic solution. Another way to put it, stochastic effects are greatest when simulating small molecule numbers

### 3.2.2 Insert Parameters

Parameters can be inserted automatically into a Copasi model.

#### Build a demonstration model

While antimony or the COPASI user interface are the preferred ways to build a model, PyCoTools does have a mechanism for constructing COPASI models. For variation and demonstra-

tion, this method is used here.

```
[1]: import os
import site
from pycotools3 import model, tasks, viz
## Choose a working directory for model
working_directory = os.path.abspath('')
copasi_file = os.path.join(working_directory, 'MichaelisMenten.cps')

if os.path.isfile(copasi_file):
    os.remove(copasi_file)

kf = 0.01
kb = 0.1
kcat = 0.05
with model.Build(copasi_file) as m:
    m.name = 'Michaelis-Menten'
    m.add('compartment', name='Cell')

    m.add('metabolite', name='P', concentration=0)
    m.add('metabolite', name='S', concentration=30)
    m.add('metabolite', name='E', concentration=10)
    m.add('metabolite', name='ES', concentration=0)

    m.add('reaction', name='S bind E', expression='S + E -> ES',
          rate_law='kf*S*E',
          parameter_values={'kf': kf})

    m.add('reaction', name='S unbind E', expression='ES -> S + E',
          rate_law='kb*ES',
          parameter_values={'kb': kb})

    m.add('reaction', name='ES produce P', expression='ES -> P + E',
          rate_law='kcat*ES',
          parameter_values={'kcat': kcat})

mm = model.Model(copasi_file)
mm

[1]: Model(name=Michaelis-Menten, time_unit=s, volume_unit=ml, quantity_
       unit=mmol)
```

## Insert Parameters from Python Dictionary

```
[2]: params = {'E': 100,
             'P': 150}

## Insert into model
```

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```
I = model.InsertParameters(mm, parameter_dict=params)
##format the parameters for displaying nicely
I.parameters.index = ['Parameter Value']
I.parameters.transpose()

[2]:    Parameter Value
E          100
P          150
```

Alternatively use `inplace=True` argument (analogous to the pandas library) to modify the object `inplace`, rather than needing to assign

```
[3]: model.InsertParameters(mm, parameter_dict=params, inplace=True)
[3]: <pycotools3.model.InsertParameters at 0x15c51a255c0>
```

## Insert Parameters from Pandas DataFrame

```
[4]: import pandas
params = {'(S bind E).kf': 50,
          '(S unbind E).kb': 96}
df = pandas.DataFrame(params, index=[0])
df

[4]:   (S bind E).kf  (S unbind E).kb
0           50            96
```

```
[ ]:
```

```
[5]: model.InsertParameters(mm, df=df, inplace=True)
[5]: <pycotools3.model.InsertParameters at 0x15c519f8dd8>
```

## Insert Parameters from Parameter Estimation Output

First we'll get some parameter estimation data by *fitting* a model to simulated data.

```
[6]: fname = os.path.join(os.path.abspath(''), 'timecourse.txt')
print(fname)
data = mm.simulate(0, 50, 1, report_name=fname)
assert os.path.isfile(fname)

D:\pycotools3\docs\source\Tutorials\timecourse.txt
```

```
[7]: with tasks.ParameterEstimation.Context(copasi_file, fname, context=
    ↪'s', parameters='l') as context:
    context.set('randomize_start_values', True)
    context.set('lower_bound', 0.01)
```

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```

context.set('upper_bound', 100)
context.set('run_mode', True)
config = context.get_config()
print(config)
PE = tasks.ParameterEstimation(config)

datasets:
    experiments:
        timecourse:
            affected_models:
                - MichaelisMenten
            filename: D:
            ↳\pycotools3\docs\source\Tutorials\timecourse.txt
            mappings:
                E:
                    model_object: E
                    object_type: Metabolite
                    role: dependent
                ES:
                    model_object: ES
                    object_type: Metabolite
                    role: dependent
                P:
                    model_object: P
                    object_type: Metabolite
                    role: dependent
                S:
                    model_object: S
                    object_type: Metabolite
                    role: dependent
            Time:
                model_object: Time
                role: time
            normalize_weights_per_experiment: true
            separator: "\t"
    validations: {}
items:
    fit_items:
        (ES produce P).kcat:
            affected_experiments:
                - timecourse
            affected_models:
                - MichaelisMenten
            affected_validation_experiments: []
            lower_bound: 1.0e-06
            start_value: model_value
            upper_bound: 1000000.0
        (S bind E).kf:
            affected_experiments:
                - timecourse

```

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```
affected_models:
- MichaelisMenten
affected_validation_experiments: []
lower_bound: 1.0e-06
start_value: model_value
upper_bound: 1000000.0
(S unbind E).kb:
    affected_experiments:
    - timecourse
    affected_models:
    - MichaelisMenten
    affected_validation_experiments: []
    lower_bound: 1.0e-06
    start_value: model_value
    upper_bound: 1000000.0
models:
MichaelisMenten:
    copasi_file: D:
    ↳ \pycotools3\docs\source\Tutorials\MichaelisMenten.cps
        model: Model(name=Michaelis-Menten, time_unit=s, volume_
    ↳ unit=ml, quantity_unit=mmol)
settings:
calculate_statistics: false
config_filename: config.yml
context: s
cooling_factor: 0.85
copy_number: 1
create_parameter_sets: false
cross_validation_depth: 1
fit: 1
iteration_limit: 50
lower_bound: 0.01
max_active: 3
method: genetic_algorithm
number_of_generations: 200
number_of_iterations: 100000
overwrite_config_file: false
pe_number: 1
pf: 0.475
pl_lower_bound: 1000
pl_upper_bound: 1000
population_size: 50
prefix: null
problem: Problem1
quantity_type: concentration
random_number_generator: 1
randomize_start_values: true
report_name: PEData.txt
results_directory: ParameterEstimationData
```

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```

rho: 0.2
run_mode: true
save: false
scale: 10
seed: 0
start_temperature: 1
start_value: 0.1
std_deviation: 1.0e-06
swarm_size: 50
tolerance: 1.0e-05
update_model: false
upper_bound: 100
use_config_start_values: false
validation_threshold: 5
validation_weight: 1
weight_method: mean_squared
working_directory: D:\pycotools3\docs\source\Tutorials

```

Now we can insert the estimated parameters using:

```
[8]: ##index=0 for best parameter set (i.e. lowest RSS)
model.InsertParameters(mm, parameter_path=PE.results_directory[
    ↪'MichaelisMenten'], index=0, inplace=True)

[8]: <pycotools3.model.InsertParameters at 0x15c6fb8c2e8>
```

## Insert Parameters using the `model.Model().insert_parameters` method

The same means of inserting parameters can be used from the model object itself

```
[9]: mm.insert_parameters(parameter_dict=params, inplace=True)
```

## Change parameters using `model.Model().set`

Individual parameters can also be changed using the `set` method. For example, we could set the metabolite with name S concentration or particle numbers to 55

```
[10]: mm.set('metabolite', 'S', 55, 'name', 'concentration')

## or

mm.set('metabolite', 'S', 55, 'name', 'particle_numbers')

[10]: Model(name=Michaelis-Menten, time_unit=s, volume_unit=ml, quantity_
    ↪unit=mmol)
```

### 3.2.3 Parameter Scan

Copasi supports three types of scan, a regular parameter scan, a repeat scan and sampling from a parametric distributions.

We first build a model to work with throughout the tutorial.

```
[1]: import os
import site
from pycotools3 import model, tasks, viz
import pandas

working_directory = r'/home/ncw135/Documents/pycotools3/docs/source/
˓→Tutorials/timecourse_tutorial'
if not os.path.isdir(working_directory):
    os.makedirs(working_directory)

copasi_file = os.path.join(working_directory, 'michaelis_menten.cps
˓→')

if os.path.isfile(copasi_file):
    os.remove(copasi_file)

antimony_string = """
model michaelis_menten()
    compartment cell = 1.0
    var E in cell
    var S in cell
    var ES in cell
    var P in cell

    kf = 0.1
    kb = 1
    kcat = 0.3
    E = 75
    S = 1000

    SBindE: S + E => ES; kf*S*E
    ESUnbind: ES => S + E; kb*ES
    ProdForm: ES => P + E; kcat*ES
end
"""

with model.BuildAntimony(copasi_file) as loader:
    mm = loader.load(antimony_string)

mm
```

The BuildAntimony context manager is deprecated and will be removed in future versions. Please use `model.loada` instead.

```
[1]: Model(name=michaelis_menten, time_unit=s, volume_unit=l, quantity_
       ↪unit=mol)

[2]: S = tasks.Scan(
        mm, scan_type='scan', subtask='time_course', report_type='time_
       ↪course',
        report_name = 'ParameterScanOfTimeCourse.txt', variable='S',
        minimum=1, maximum=20, number_of_steps=8, run=True,
    )

    ## Now check parameter scan data exists
os.path.isfile(S.report_name)

[2]: True
```

## Two Way Parameter Scan

By default, scan tasks are removed before setting up a new scan. To set up dual scans, set clear\_scans to False in a second call to Scan so that the first is not removed prior to adding the second.

```
[3]: ## Clear scans for setting up first scan
tasks.Scan(
    mm, scan_type='scan', subtask='time_course', report_type='time_
       ↪course',
    variable='E', minimum=1, maximum=20, number_of_steps=8, run=False, clear_scan=True,
)

## do not clear tasks when setting up the second
S = tasks.Scan(
    mm, scan_type='scan', subtask='time_course', report_type='time_
       ↪course',
    report_name = 'TwoWayParameterScanOfTimeCourse.csv', variable='S
       ↪',
    minimum=1, maximum=20, number_of_steps=8, run=True, clear_
       ↪scan=False,
)

## check the output exists
os.path.isfile(S.report_name)

[3]: True
```

An arbitrary number of scans can be setup this way. Further, its possible to chain together scans with repeat or random distribution scans.

## Repeat Scan Items

Repeat scans are very useful for running multiple parameter estimations and for running stochastic time courses.

```
[4]: ## Assume Parameter Estimation task already configured
tasks.Scan(
    mm, scan_type='repeat', subtask='parameter_estimation', report_
    ↪type='parameter_estimation',
    number_of_steps=6, run=False, ##set run to True to run via_
    ↪CopasiSE
)

## Assume model runs stochastically and time course settings are_
    ↪already configured
tasks.Scan(
    mm, scan_type='repeat', subtask='time_course', report_type=
    ↪'time_course',
    number_of_steps=100, run=False, ##set run to True to run via_
    ↪CopasiSE
)
[4]: <pycotools3.tasks.Scan at 0x1cb859370b8>
```

## 3.2.4 Parameter Estimation

```
[4]: import os, glob
import site
from pycotools3 import viz, model, misc, tasks
from io import StringIO
import pandas
%matplotlib inline
```

## Build a Model

```
[8]: working_directory = os.path.abspath('')

copasi_file = os.path.join(working_directory, 'negative_feedback.cps'
    ↪')
ant = """
    model negative_feedback
        compartment cell = 1.0
        var A in cell
        var B in cell

        vAProd = 0.1
```

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```

kADeg = 0.2
kBProd = 0.3
kBDeg = 0.4
A = 0
B = 0

AProd: => A; cell*vAProd
ADeg: A =>; cell*kADeg*A*B
BProd: => B; cell*kBProd*A
BDeg: B =>; cell*kBDeg*B
end
"""
mod = model.loada(ant, copasi_file)

## open model in copasi
#mod.open()
mod
[8]: Model(name=negative_feedback, time_unit=s, volume_unit=l, quantity_
      ↪unit=mol)

```

## Collect some experimental data

Organise your experimental data into delimited text files

```

[9]: experimental_data = StringIO(
    """
Time,A,B
0, 0.000000, 0.000000
1, 0.099932, 0.013181
2, 0.199023, 0.046643
3, 0.295526, 0.093275
4, 0.387233, 0.147810
5, 0.471935, 0.206160
6, 0.547789, 0.265083
7, 0.613554, 0.322023
8, 0.668702, 0.375056
9, 0.713393, 0.422852
10, 0.748359, 0.464639
    """.strip()
)

df = pandas.read_csv(experimental_data, index_col=0)

fname = os.path.join(os.path.abspath(''), 'experimental_data.csv')
df.to_csv(fname)

assert os.path.isfile(fname)

```

## The Config Object

The interface to COPASI's parameter estimation using pycotools3 revolves around the `ParameterEstimation.Config` object. `ParameterEstimation.Config` is a dictionary-like object which allows the user to define their parameter estimation problem. All features of COPASI's parameter estimations task are supported, including configuration of validation experiments, affected experiments, affected validation experiments and constraints as well additional features such as the configuration of multiple models simultaneously via the `affected_models` keyword.

The `ParameterEstimation.Config` object expects at the bare minimum some information about the models being configured, some experimental data, some fit items and a working directory. The remaining options are automatically filled in with defaults.

```
[10]: config = tasks.ParameterEstimation.Config(
    models=dict(
        negative_feedback=dict(
            copasi_file=copasi_file
        )
    ),
    datasets=dict(
        experiments=dict(
            first_dataset=dict(
                filename=fname,
                separator=','
            )
        )
    ),
    items=dict(
        fit_items=dict(
            A={},
            B={}
        )
    ),
    settings=dict(
        working_directory=working_directory
    )
)
config

[10]: datasets:
       experiments:
           first_dataset:
               affected_models:
                   - negative_feedback
               filename: D:
               ↵\pycotoold3\docs\source\Tutorials\experimental_data.csv
               mappings:
                   A:
                       model_object: A
                       object_type: Metabolite
```

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```

        role: dependent
B:
    model_object: B
    object_type: Metabolite
    role: dependent
Time:
    model_object: Time
    role: time
normalize_weights_per_experiment: true
separator: ','
validations: {}
items:
    fit_items:
        A:
            affected_experiments:
            - first_dataset
            affected_models:
            - negative_feedback
            affected_validation_experiments: []
            lower_bound: 1.0e-06
            start_value: model_value
            upper_bound: 1000000
        B:
            affected_experiments:
            - first_dataset
            affected_models:
            - negative_feedback
            affected_validation_experiments: []
            lower_bound: 1.0e-06
            start_value: model_value
            upper_bound: 1000000
models:
    negative_feedback:
        copasi_file: D:\pycotools3\docs\source\Tutorials\negative_
        ↪feedback.cps
        model: Model(name=negative_feedback, time_unit=s, volume_
        ↪unit=l, quantity_unit=mol)
settings:
    calculate_statistics: false
    config_filename: config.yml
    context: s
    cooling_factor: 0.85
    copy_number: 1
    create_parameter_sets: false
    cross_validation_depth: 1
    fit: 1
    iteration_limit: 50
    lower_bound: 1.0e-06
    max_active: 3

```

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```
method: genetic_algorithm
number_of_generations: 200
number_of_iterations: 100000
overwrite_config_file: false
pe_number: 1
pf: 0.475
pl_lower_bound: 1000
pl_upper_bound: 1000
population_size: 50
prefix: null
problem: Problem1
quantity_type: concentration
random_number_generator: 1
randomize_start_values: false
report_name: PEData.txt
results_directory: ParameterEstimationData
rho: 0.2
run_mode: false
save: false
scale: 10
seed: 0
start_temperature: 1
start_value: 0.1
std_deviation: 1.0e-06
swarm_size: 50
tolerance: 1.0e-05
update_model: false
upper_bound: 1000000
use_config_start_values: false
validation_threshold: 5
validation_weight: 1
weight_method: mean_squared
working_directory: D:\pycotools3\docs\source\Tutorials
```

The COPASI user will be familiar with most of these settings, though there are also a few additional options.

Once built, a `ParameterEstimation.Config` object can be passed to `ParameterEstimation` object.

```
[11]: PE = tasks.ParameterEstimation(config)
```

By default, the `run_mode` setting is set to False. To run the parameter estimation in background processes using CopasiSE, set `run_mode` to True or parallel.

```
[12]: config.settings.run_mode = True
PE = tasks.ParameterEstimation(config)
viz.Parse(PE) ['negative_feedback']
# config
```

```
[12]:      A          B          RSS
0  0.000001  0.000001  7.955450e-12
```

## Running multiple parameter estimations

With pycotools, parameter estimations are run via the scan task interface so that we have the option of running the same problem `pe_number` times. Additionally, pycotools provides a way of copying a model `copy_number` times so that the final number of parameter estimations that get executed is `pe_number * copy_number`.

```
[13]: config.settings.copy_number = 4
config.settings.pe_number = 2
config.settings.run_mode = True
PE = tasks.ParameterEstimation(config)
```

And sure enough we have ran the problem 8 times.

```
[14]: viz.Parse(PE) ['negative_feedback']
```

```
[14]:      A          B          RSS
0  0.000001  0.000001  7.955430e-12
1  0.000001  0.000001  7.955450e-12
2  0.000001  0.000001  7.955450e-12
3  0.000001  0.000001  7.955450e-12
4  0.000001  0.000001  7.955450e-12
5  0.000001  0.000001  7.955450e-12
6  0.000001  0.000001  7.955450e-12
7  0.000001  0.000001  7.955450e-12
```

### 3.2.5 A shortcut for configuring the ParameterEstimation.Config object

Manually configuring the `ParameterEstimation.Config` object can take some time as it is bulky, but necessarily so in order to enable users to configure any type of parameter estimation. The `ParameterEstimation.Config` class should be used directly when a lower level interface into COPASI configurations are required. For instance, if you want to configure different boundaries for each parameter, choose which parameters are affected by which experiment, mix timecourse and steady state experiments, define independent variables, add constraints or choose which models are affected by which experiments, you can use the `ParameterEstimation.Config` class directly.

However, if you want a more standard configuration such as all parameters estimated between the same boundaries, all experiments affecting all parameters and models etc.. then you can use the `ParameterEstimation.Context` class to build the `ParameterEstimation.Config` class for you. The `ParameterEstimation.Context` class has a `context` argument that defaults to '`s`' for simple. While not yet implemented, eventually, alternative options for context will be provided to support other common patterns, such as

cross\_validation or chaser\_estimations (global followed by local algorithm). Note that an option is no longer required for model\_selection since it is innately incorporated via the affected\_models argument.

To use the ParameterEstimation.Context object

```
[17]: with tasks.ParameterEstimation.Context(mod, fname, context='s',  
    ↪parameters='g') as context:  
    context.set('method', 'genetic_algorithm')  
    context.set('population_size', 10)  
    context.set('copy_number', 4)  
    context.set('pe_number', 2)  
    context.set('run_mode', True)  
    context.set('separator', ',')  
    config = context.get_config()  
  
pe = tasks.ParameterEstimation(config)
```

```
[18]: viz.Parse(pe) ['negative_feedback']
```

```
[18]:
```

	RSS	kADeg	kBDeg	kBProd	vAProd
0	8.851340e-13	0.2	0.4	0.3	0.1
1	8.851340e-13	0.2	0.4	0.3	0.1
2	8.851340e-13	0.2	0.4	0.3	0.1
3	8.851340e-13	0.2	0.4	0.3	0.1
4	8.851340e-13	0.2	0.4	0.3	0.1
5	8.851340e-13	0.2	0.4	0.3	0.1
6	8.851340e-13	0.2	0.4	0.3	0.1
7	8.851340e-13	0.2	0.4	0.3	0.1

The parameters keyword provides an easy interface for parameter selection. Here are the available options: - g specifies that all global variables are to be estimated - l specifies that all local parameters are to be estimated - m specifies that all metabolites are to be estimated - c specifies that all compartment volumes are to be estimated - a specifies that all of the above will be estimated

These options can also be combined. For example, parameters='cgm' means that compartment volumes, global quantities and metabolite concentrations (or particle numbers) will be estimated.

```
[ ]:
```

## 3.3 Examples

### 3.3.1 Working with an existing copasi model

You can create a new model using the antimony language. Sometimes however, you have already built a model using the CoapsiUI and want to use pycotools3 with this model. This example shows you how to use the `pycotools3.model.Model` class directly.

## Create a PyCoTools model from an existing model

```
from pycotools3 import model

# remember to input the string to your own model here
copasi_file = <'string/to/model.cps'>

mod = model.Model(copasi_file)

assert isinstance(mod, model.Model)
```

## Extracting the antimony string associated with a copasi model

It is often useful to have an antimony string generated directly from a copasi model.

```
ant_string = mod.to_antimony()
print(ant_string)
```

### 3.3.2 Integration with Tellurium

**Tellurium** is a Python package built on top of [libRoadRunner](#), a C++ package for simulation and analysis of models in systems biology. Antimony is a model definition language for SBML models that was written by the authors of *Tellurium* and *libRoadRunner*.

Since PyCoTools has adopted *Antimony*, the same model can be simulated and analysed with both COPASI and tellurium.

Here's a short example of how using the tellurium back end.

```
import tellurium as te

antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
    B in Cell;
    C in Cell;

    // reactions
    R1: A => B ; Cell * k1 * A;
    R2: B => A ; Cell * k2 * B * C;
    R3: B => C ; Cell * k3 * B;
    R4: C => B ; Cell * k4 * C;

    // initial concentrations
    A = 100;
    B = 1;
```

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```
C = 1;

// reaction parameters
k1 = 0.1;
k2 = 0.1;
k3 = 0.1;
k4 = 0.1;
end
"""

mod = te.loada(antimony_string)

# simulate a time series with 11 time points between 0 and 10
timeseries_data = mod.simulate(0, 10, 11)

# compute the steady state
mod.conservMoietyAnalysis = True
mod.setSteadyStateSolver('nleq')
mod.steadyState()
mod.steadyStateSelections = ['A', 'B', 'C']
print(dict(zip(*mod.steadyStateSelections,
    mod.getSteadyStateValues())))

```

### 3.3.3 Simple Parameter Estimation

This is an example of how to configure a simple parameter estimation using pycotools. We first create a toy model for demonstration, then simulate some experimental data from it and fit it back to the model, using pycotools for configuration.

```
import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz
seaborn.set_context(context='talk')           # set seaborn context_
→for formatting output of plots

## Choose a directory for our model and analysis. Note this can be_
→anywhere.
working_directory = os.path.abspath('')

## In this model, A gets reversibly converted to B but the_
→backwards reaction is additionally regulated by C.
## B is reversibly converted into C.
antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;
```

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```

A in Cell;
B in Cell;
C in Cell;

// reactions
R1: A => B ; Cell * k1 * A;
R2: B => A ; Cell * k2 * B * C;
R3: B => C ; Cell * k3 * B;
R4: C => B ; Cell * k4 * C;

// initial concentrations
A = 100;
B = 1;
C = 1;

// reaction parameters
k1 = 0.1;
k2 = 0.1;
k3 = 0.1;
k4 = 0.1;
end
"""

# Create a path to a copasi file
copasi_file = os.path.join(working_directory, 'example_model.cps')

## build model
mod = model.loada(antimony_string, copasi_file)
assert isinstance(mod, model.Model)

## simulate some data, returns a pandas.DataFrame
data = mod.simulate(0, 20, 1)

## write data to file
experiment_filename = os.path.join(working_directory, 'experiment_
→data.txt')
data.to_csv(experiment_filename)

## We now have a model and some experimental data and can
## configure a parameter estimation

```

Parameter estimation configuration in pycotools3 revolves around the `tasks.ParameterEstimation.Config` object which is the input to the parameter estimation task. The object necessarily takes a lot of manual configuration to ensure it is flexible enough for any parameter estimation configuration. However, the `ParameterEstimation.Context` class is a tool for simplifying the construction of a `Config` object.

```
with tasks.ParameterEstimation.Context(mod, experiment_filename,_
    ↪context='s', parameters='g') as context:
    context.set('separator', ',')
    context.set('run_mode', True)
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 100)
    context.set('lower_bound', 1e-1)
    context.set('upper_bound', 1e1)

    config = context.get_config()

pe = tasks.ParameterEstimation(config)

data = viz.Parse(pe).data
print(data)
```

### 3.3.4 Simple Parameter with Steady State Data

The short story here is that PyCoTools distinguishes time series and steady state data automatically, using the presence or absence of the *time* column.

Here's an example.

```
import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz
seaborn.set_context(context='talk')           # set seaborn context_
    ↪for formatting output of plots

## Choose a directory for our model and analysis. Note this can be_
    ↪anywhere.
working_directory = os.path.abspath('')

## In this model, A gets reversibly converted to B but the_
    ↪backwards reaction is additionally regulated by C.
## B is reversibly converted into C.
antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
    B in Cell;
    C in Cell;

    // reactions
    R1: A => B ; Cell * k1 * A;
```

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```

R2: B => A ; Cell * k2 * B * C;
R3: B => C ; Cell * k3 * B;
R4: C => B ; Cell * k4 * C;

// initial concentrations
A = 100;
B = 1;
C = 1;

// reaction parameters
k1 = 0.1;
k2 = 0.1;
k3 = 0.1;
k4 = 0.1;
end
"""

# Create a path to a copasi file
copasi_file = os.path.join(working_directory, 'example_model.cps')

## build model
mod = model.loada(antimony_string, copasi_file)
assert isinstance(mod, model.Model)

## create some made up data
data = pandas.DataFrame({'A': 30, 'B': 10, 'C': 10})

## write data to file
experiment_filename = os.path.join(working_directory, 'experiment_
→data.txt')
data.to_csv(experiment_filename)

```

We now have a model and some experimental data and can configure a parameter estimation. Configuring steady state data is semantically identical to configuring time series data. The difference is that our *data* no longer has a *time* column and so PyCoTools assumes that it is steady state data.

Now, as usual, we configure the parameter estimation with the *Context* manager.

```

with tasks.ParameterEstimation.Context(mod, experiment_filename, ↴
→context='s', parameters='g') as context:
    context.set('separator', ',')
    context.set('run_mode', True)
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 100)
    context.set('lower_bound', 1e-1)
    context.set('upper_bound', 1e1)

```

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```
config = context.get_config()

pe = tasks.ParameterEstimation(config)

data = viz.Parse(pe).data
print(data)
```

### 3.3.5 Parameter Estimation using Prefix Argument

Sometimes we would like to select a set of parameters to estimate and leave the rest fixed. One way to do this is to compile a list of parameters you would like to estimate and enter them into the `ParameterEstimation.Config` class. A quicker alternative is to use the *prefix* setting.

```
import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz
seaborn.set_context(context='talk')

## Choose a directory for our model and analysis
working_directory = os.path.abspath('')
```

The *prefix* argument will setup the configuration of a parameter estimation containing only parameters that start with *prefix*. While this can be anything, its quite useful to use the `_` character and then add an `_` to any parameters that you want estimated. In this way you can keep your estimated parameters marked.

```
antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
    B in Cell;
    _C in Cell;

    // reactions
    R1: A => B ; Cell * _k1 * A;
    R2: B => A ; Cell * k2 * B * _C;
    R3: B => C ; Cell * _k3 * B;
    R4: C => B ; Cell * k4 * _C;

    // initial concentrations
    A = 100;
    B = 1;
    _C = 1;
```

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```
// reaction parameters
_k1 = 0.1;
k2 = 0.1;
_k3 = 0.1;
k4 = 0.1;
end
"""

copasi_file = os.path.join(working_directory, 'example_model.cps')

## build model
mod = model.loada(antimony_string, copasi_file)

assert isinstance(mod, model.Model)

fname = os.path.join(working_directory, 'experiment_data.txt')
data = mod.simulate(0, 20, 1, report_name=fname)

## write data to file
```

And now we configure a parameter estimation like normal but set *prefix* to `_`.

```
with tasks.ParameterEstimation.Context(mod, experiment_filename, 
    ↪context='s', parameters='a') as context:
    context.set('separator', ',')
    context.set('run_mode', True)
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 100)
    context.set('lower_bound', 1e-1)
    context.set('upper_bound', 1e1)
    context.set('prefix', '_')
    config = context.get_config()

pe = tasks.ParameterEstimation(config)

data = viz.Parse(pe).data
print(data)
```

### 3.3.6 Parameter Estimation with Independent Variables

The concept of independent variables is important for parameter fitting because it enables us to simultaneously fit multiple datasets to a single model. This is achieved by iterating over all the datasets in your objective function and changing variables (such as initial concentration parameters) to whatever they should be for that dataset. These variables are independent variables and they basically define the initial conditions for fitting the dataset.

Independent variables are handled in PyCoTools by appending the string ‘*\_indep*’ after a variable in the data file itself. PyCoTools will then recognize the variable and set it as independent rather than dependent.

Here’s an example:

```
import os, glob
import pandas, numpy
from pycotools3 import model, tasks, viz
seaborn.set_context(context='talk')           # set seaborn context_
→for formatting output of plots

## Choose a directory for our model and analysis. Note this can be_
→anywhere.
working_directory = os.path.abspath('')

## In this model, A gets reversibly converted to B but the_
→backwards reaction is additionally regulated by C.
## B is reversibly converted into C.
antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
    B in Cell;
    C in Cell;

    // reactions
    R1: A => B ; Cell * k1 * A;
    R2: B => A ; Cell * k2 * B * C;
    R3: B => C ; Cell * k3 * B;
    R4: C => B ; Cell * k4 * C;

    // initial concentrations
    A = 100;
    B = 1;
    C = 1;

    // reaction parameters
    k1 = 0.1;
    k2 = 0.1;
    k3 = 0.1;
    k4 = 0.1;
end
"""

# Create a path to a copasi file
copasi_file = os.path.join(working_directory, 'example_model.cps')

# build model
mod = model.loada(antimony_string, copasi_file)
```

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```
assert isinstance(mod, model.Model)

# simulate some data, returns a pandas.DataFrame
data = mod.simulate(0, 20, 1)

# creates a new column in the dataset called A_indep
data['A_indep'] = 50
# the initial abundance of A will now be set to 50 prior to_
→estimation

# write data to file
experiment_filename = os.path.join(working_directory, 'experiment_'
→data.txt')
data.to_csv(experiment_filename)
```

We now configure a parameter estimation like normal.

```
with tasks.ParameterEstimation.Context(mod, experiment_filename,_
→context='s', parameters='g') as context:
    context.set('separator', ',')
    context.set('run_mode', True)
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 100)
    context.set('lower_bound', 1e-1)
    context.set('upper_bound', 1e1)

    config = context.get_config()

pe = tasks.ParameterEstimation(config)

data = viz.Parse(pe).data
print(data)
```

### 3.3.7 Advanced Parameter Estimation

Parameter estimation configuration in PyCoTools revolves around the `ParameterEstimation.Config` object, which is merely a nested data storage class. When you use the `ParameterEstimation.Context` object, you are using a high level interface to create a `ParameterEstimation.Config` object. However, it is also possible to build it manually. It should be stressed however, that the `Context` method should be preferred when possible. The low level interface can be tedious but gives you more flexibility for more fine tuned parameter estimation configurations. While there is no getting around the fact that building the `ParameterEstimation.Config` object can take time, I have built in a ‘default system’ whereby if a (non-compulsory) section is left blank, then PyCoTools tries to use the default values.

The `ParameterEstimation.Config` Object is a bit like a Struct from other languages or

a tree (see `bunch.Bunch`). You can access attributes using either dot notation or dictionary like access.

There are four main sections to the Config object:

- *Models*
- *Datasets*
- *Items*
- *Settings*

Each of these are described in detail below. Note that all branches in the config object described here are themselves nodes in the tree with children unless they are labelled with *leaf*. These nodes expect the usual key: value pairs.

## Models

This is a nested structure containing an arbitrary number of models for simultaneous configuration. It is the users responsibility that it makes sense to configure multiple models for parameter estimation in the same way.

- `models`
  - `model1_name` (*leaf*): full string to model 1 copasi file
  - `model2_name` (*leaf*): full string to model 2 copasi file
  - ...

```
models = dict(
    model1=dict(copasi_file='</full/path/to/copasi_file1.cps'),
    model2=dict(copasi_file='</full/path/to/copasi_file2.cps')
)
```

---

**Note:** The nested structure of the `models` section remains a deprecated version of PyCoTools. Since it does not need to be nested, this will be changing to non-nested in the near future.

---

## Datasets

This is where we tell PyCoTools about the data were using for parameter estimation. The datasets attribute is itself nested with the following structure:

- `datasets`
  - `experiments`
  - `validations`

Both *experiments* and *validations* are nested structures and they are pretty much identical. The difference is that those configured via the *experiments* section are

used for calibrating/training the model while those in the validation section are only used for validation (or testing). For simplicity, the structure of both is described in one section.

- experiments (or validation)
  - experiment\_name. An arbitrary string representing the name of the experiment.
  - \* filename (leaf). The full path to the dataset
  - \* affected\_models. Analogous to affected\_experiments or affected\_validation\_experiments, you can have an experiment target only one (or more) model. This feature is a superset of COPASI. Defaults to the string ‘all’ which is translated to all models.
  - \* mappings. Another nested structure for mapping arguments. If left blank, PyCoTools will assume 1:1 mappings between experimental data file headers and model variables. Independent variables are assumed to contain a trailing *\_indep*, i.e. *PI3K\_indep*. This should have as many elements as there are columns in the data file.
    - Experimental variable name (or time). These should be the same as used for data column headers.
    - model\_object (leaf node). The object that corresponds to the experimental variable name.
    - role (leaf node). Either *time*, *ignored* (default), *dependent* or *independent*
    - \* separator (leaf). Overrides the separator in the settings section, for when they are different. However, good practice is to always use the same separator and set the separator in the settings section.
    - \* normalize\_weights\_per\_experiment (leaf): boolean, default=True.

Here’s an example of the datasets section.

```
datasets=dict(
    experiments= dict(
        report1 = dict(
            filename='full/path/to_datafile1.csv',
            affected_models='all',
            mappings=dict(
                Time=dict(
                    model_object='Time',
                    role='time'
                ),
                A=dict(
                    model_object='A'),
                    role='dependent'),
                )
            )
        )
)
```

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```
) ,  
    # note the absence of the mappings field. This tells  
    # PyCoTools that you have used the suggested convention  
    # of matching model variables with data file headers and  
→using  
    # '_indep' suffix for independent variables.  
    report2=dict(  
        filename='full/path/to_datafile2.csv',  
        separator='\t'  #overrides separator from main settings  
→menu  
    )  
) ,  
    # This data will not be used for parameter estimation. Only  
→validation.  
    validations=dict(  
        report3=dict(  
            filename='full/path/to_datafile3.csv',  
            affected_models='modell',  # this validation experiment  
→only affects modell  
            # were excepting default mapping convention  
        ),  
    )  
)
```

## Items

This is where we configure the parameters to be estimated, their boundaries, start values and affected experiments. The *items* structure is composed of *fit\_items* and *constraint\_items*.

- items
    - fit\_items
    - constraint\_items

Similarly to the experiment section, fit\_items and constraint\_items are nearly identical. The difference is that whilst fit items are used to define the parameter space constraints are used to restrict the parameter space to a subset of the full parameter space. An estimation with constraints can explore beyond the restrictions imposed by the constraints but solutions that violate the constraints will not be excepts. In contrast, the solution cannot go beyond the boundaries of the boundaries set by the fit\_items.

```
items = dict(
    fit_items=dict(
        A=dict(
            affected_experiments=['report1'],
            affected_models=['model1'],
            affected_validation_experiments=['report3'],
            lower_bound=15,
```

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```
        start_value=0.1,
        upper_bound=35
    ),
    B=dict (
        affected_experiments=['report1', 'report2'],
        affected_models=['model1'],
        affected_validation_experiments=['report3'],
        lower_bound=0.05,
        start_value=1.05,
        upper_bound=36
    ),
    C=dict (
        affected_experiments=['report1', 'report2'],
        affected_models=['model1'],
        affected_validation_experiments=['report3'],
        lower_bound=0.05,
        start_value=1.0,
        upper_bound=36
    )
),
constraint_items=dict (
    C=dict (
        affected_experiments=['report1', 'report2'],
        affected_models=['model1'],
        affected_validation_experiments=['report3'],
        lower_bound=16,
        start_value=1.05,
        upper_bound=26
    )
)
)
```

**Note:** *affected\_experiments*, *affected\_models*, and *affected\_validation\_experiments* all accept the special string *all* which resolves to all of your data files. This is default behaviour for both *affected\_experiments* and *affected\_models* whereas the default behaviour for *affected\_validation\_experiments* is None.

## Settings

These are global settings for the parameter estimation.

```
settings = dict(
    calculate_statistics=False,      # Corresponds to the `calculate_
→statistics` flag in copasi
    config_filename=config.yml      # Filename for saving config to_
→file
```

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```

    context=s,                                # Alters the behaviour of the
→configuration. See :py:class:`ParameterEstimation.Context`.
    cooling_factor=0.85                      # Parameter estimation

→algorithm setting
    copy_number=1,                            # How many times to copy the
→copasi file for simultaneous runs
    create_parameter_sets=False,             # Corresponds to the create_
→parameter_sets flag in copasi
    cross_validation_depth=1,                # depth of cross validation.
→Corresponds to COPASI, (though this feature was buggy during
development)
    fit=1,                                    # This is an index of parameter
→estimation. Increment by 1 to repeat a similar parameter
→estimation to test alternative configurations
    iteration_limit=50,                      # Parameter estimation

→algorithm setting
    lower_bound=0.05                         # Default lower boundary for
→all parameters in the estimation. Can be overwritten under the
→fit_items section to have different boundaries for every fit item.
    max_active=3,                            # When running
    method=genetic_algorithm_sr,            # which algorithm to use
    number_of_generations=100,              # Parameter estimation

→algorithm setting
    number_of_iterations=100000,            # Parameter estimation

→algorithm setting
    overwrite_config_file=False,           # Set to True to explicitly
→overwrite existing configuration file.
    pe_number=1,                            # How many parameter estimations
    pf=0.475                               # Parameter estimation

→algorithm settings
    pl_lower_bound=1000,                   # When context is set to 'pl'
→for profile likelihood configurations, this defines the upper
→boundary of the analysis. The upper boundary is the best
→estimated parameter multiplied by this value.
    pl_upper_bound=1000,                   # When context is set to 'pl'
→for profile likelihood configurations, this defines the lower
→boundary of the analysis. The lower boundary is the best
→estimated parameter divided by this value.
    population_size=38,                  # Parameter estimation

→algorithm setting
    prefix=None,                           # Prefix used to automatically
→locate parameters to be estimated. For instance, you can 'tag'
→each parameter you want to include in the estimation with an
→underscore at the begining (i.e. _kAktPhosphorylation) to filter
→the parameters for estimation.
    problem=Problem1,                     # This is the name of the
→folder that will be created to contain the results.
    quantity_type=concentration,        # either 'concentration' or
→'particle_numbers' to switch between the two.

```

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```

    random_number_generator=1,           # Parameter estimation
    ↪algorithm setting.
    randomize_start_values=False,      # Corresponds to the 'randomize_
    ↪start_values' flag in copasi
    report_name=PEData.txt            # The base report name for the
    ↪parameter estimation output. This is automatically modified when
    ↪copy_number is > 1. The results have as many rows as `pe_number`.
    results_directory=ParameterEstimationData, # This folder
    ↪stores the actual parameter estimation results, within the fit_
    ↪directory (which is within the Problem directory)
    rho=0.2                           # Parameter estimation
    ↪algorithm setting
    run_mode=False,                   # Switch between False
    save=False,                      # Save the model to file after
    ↪configuration or not.
    scale=10,                         # Parameter estimation
    ↪algorithm setting
    seed=0,                           # Parameter estimation
    ↪algorithm setting
    start_temperature=1,             # Parameter estimation
    ↪algorithm setting
    start_value=0.1                  # Parameter estimation
    ↪algorithm setting
    starting_parameter_sets=None,     # Experimental feature.
    std_deviation=1.0e-06            # Parameter estimation
    ↪algorithm setting
    swarm_size=50,                   # Parameter estimation
    ↪algorithm setting
    tolerance=1.0e-05                # Parameter estimation
    ↪algorithm setting
    update_model=False,              # Corresponds to the update_
    ↪model flag in copasi
    upper_bound=36,                  # Default upper boundary for_
    ↪all parameters in the estimation. Can be overwritten under the
    ↪fit_items section to have different boundaries for every fit item.
    use_config_start_values=False,   # If True, parameter estimation
    ↪will start from the start values specified under the `fit_items`_
    ↪section.
    validation_threshold=8.5         # Corresponds to the validation_
    ↪threshold in COPASI. This is the default value that can be
    ↪overwritten by giving this argument to the validation dataset_
    ↪section.
    validation_weight=4,             # Corresponds to the validation_
    ↪weight in COPASI. This is the default value that can be
    ↪overwritten by giving this argument to the validation dataset_
    ↪section.
    weight_method=value_scaling,    # Which weight method to use.
    ↪Default='mean_squared'. Other options: mean, standard_deviati
    ↪or value_scaling

```

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```
working_directory=/home/ncw135/Documents/pycotoold3/Tests #  
→The overall directory for the whole analysis. Defaults to the  
→same directory containing the first copasi file found for  
→configuration.  
)
```

## Building a ParameterEstimation.Config object

When you have configured the relevant sections, you can simply call the ParameterEstimation.Config constructor to create your object.

Assuming you have nested dictionaries containing the appropriate information detailed above:

```
config = tasks.ParameterEstimation.Config(  
    models=models,  
    datasets=datasets,  
    items=items,  
    settings=settings  
)
```

The config is formatted using yaml for ease of inspection.

---

**Note:** It is possible to load from yaml file on disk. Documentation to come.

---

## Using a ParameterEstimation.Context as a template

The most effective way to use the low level interface is to let the ParameterEstimation.Context do most of the work and then retrieve the mostly configured config string and then make your desired amendments.

### 3.3.8 Multiple Parameter Estimations

Configuring multiple parameter estimations is easy with COPASI because you can configure a parameter estimation task, configure a scan repeat item with the *subtask* set to *parameter estimation* and hit run. This is what pycotools does under the hood to configure a parameter estimation, even if the desired number of parameter estimations is 1.

Moreover, pycotools additionally supports the running of multiple copies of your copasi file in separate processes, or on a cluster.

```
from pycotools3 import model, tasks  
  
working_directory = os.path.abspath('')
```

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```

antimony_string = '''
model negative_feedback()
    // define compartments
    compartment cell = 1.0
    //define species
    var A in cell
    var B in cell
    //define some global parameter for use in reactions
    vAProd = 0.1
    kADeg = 0.2
    kBProd = 0.3
    kBDeg = 0.4
    //define initial conditions
    A      = 0
    B      = 0
    //define reactions
    AProd:  => A; cell*vAProd
    ADeg: A => ; cell*kADeg*A*B
    BProd: => B; cell*kBProd*A
    BDeg: B => ; cell*kBDeg*B
end
'''

copasi_file = os.path.join(working_directory, 'negative_fb.cps')
mod = model.loada(antimony_string, copasi_file )

data_fname = os.path.join(working_directory, 'timecourse.txt')
mod.simulate(0, 10, 1, report_name=data_fname)

assert os.path.isfile(data_fname)

```

## Increasing Parameter Estimation Throughput

The *pe\_number* argument specifies the number that gets entered into the copasi scan repeat item while the *copy\_number* argument specifies the number of identical model copies to make.

```

with tasks.ParameterEstimation.Context(mod, data_fname, context='s',
    ↳ parameters='g') as context:
    context.set('copy_number', 2)
    context.set('pe_number', 2)
    context.set('randomize_start_values', True)
    context.set('run_mode', True)
    config = context.get_config()

pe = tasks.ParameterEstimation(config)
data = viz.Parse(pe)

```

---

**Note:** The `copy_number` argument here doesn't really do anything useful because `run_mode=True`. This tells pycotools to run the parameter estimations in series (i.e. back to back) and therefore the `copy_number` argument here does nothing.

---

However, it is also possible to give `run_mode='parallel'` and in this case, each of the model copies will be run simultaneously.

```
with tasks.ParameterEstimation.Context(mod, data_fname, context='s',
    ↪ parameters='g') as context:
    context.set('copy_number', 2)
    context.set('pe_number', 2)
    context.set('randomize_start_values', True)
    context.set('run_mode', 'parallel')
    config = context.get_config()

pe = tasks.ParameterEstimation(config)
data = viz.Parse(pe)
```

**Warning:** Users should not use `run_mode='parallel'` in combination with a high `copy_number` as it will slow your system.

Your system has a limited amount of resources and can only handle a number of parameter estimations being run at once. For this reason, be careful when choosing the `copy_number`. For reference, my computer can run approximately 8 parameter estimations in different processes before slowing.

If you have access to a cluster running either SunGrid Engine or Slurm then each of the `copy_number` models will be submitted as separate jobs. To do this set `run_mode='slurm'` or `run_mode='sge'` (see `tasks.Run`).

**Warning:** The cluster functions are fully operational on the Newcastle University clusters but untested on other clusters. If you run into trouble, contact me for help.

It is easy to support other cluster systems by adding a method to `tasks.Run` using `tasks.Run.run_sge()` and `tasks.Run.run_slurm()` as examples.

### 3.3.9 Parameter estimation with multiple models

This is an example of how to configure a parameter estimation for multiple COPASI models using pycotools. We first create two similar but different toy models for demonstration, then simulate some experimental data from one of them and fit it back to both models.

```

import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz

# set seaborn context
seaborn.set_context(context='talk')

## Choose a directory for our model and analysis
working_directory = os.path.abspath('')

model1_string = """
model model1()

    R1:    => A ; k1*S;
    R2: A =>      ; k2*A;
    R3:    => B ; k3*A;
    R4: B =>      ; k4*B*C; //feedback term
    R5:    => C ; k5*B;
    R6: C =>      ; k6*C;

    S = 1;
    k1 = 0.1;
    k2 = 0.1;
    k3 = 0.1;
    k4 = 0.1;
    k5 = 0.1;
    k6 = 0.1;
end
"""

model2_string = """
model model2()
    R1:    => A ; k1*S;
    R2: A =>      ; k2*A*C; //feedback term
    R3:    => B ; k3*A;
    R4: B =>      ; k4*B;
    R5:    => C ; k5*B;
    R6: C =>      ; k6*C;

    S = 1;
    k1 = 0.1;
    k2 = 0.1;
    k3 = 0.1;
    k4 = 0.1;
    k5 = 0.1;
    k6 = 0.1;
end
"""

```

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```

# create paths to where we want the two models
copasi_file1 = os.path.join(working_directory, 'model1.cps')
copasi_file2 = os.path.join(working_directory, 'model2.cps')

# Assemble into lists
antimony_strings = [model1_string, model2_string]
copasi_files = [copasi_file1, copasi_file2]

# create models
model_list = []
for i in range(len(copasi_files)):
    model_list.append(model.loada(antimony_strings[i], copasi_
        ↴files[i]))

## simulate some data, returns a pandas.DataFrame
data = model_list[0].simulate(0, 20, 1)

## write data to file
experiment_filename = os.path.join(working_directory, 'data_from_'
    ↴model1.txt')
data.to_csv(experiment_filename)

# Create the context, passing the model list rather than the Model_
↪object
with tasks.ParameterEstimation.Context(model_list, experiment_
    ↴filename, context='s', parameters='g') as context:
    context.set('separator', ',')
    context.set('run_mode', True)
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 25)
    context.set('lower_bound', 1e-1)
    context.set('upper_bound', 1e1)

    config = context.get_config()

# Do the parameter estimation
pe = tasks.ParameterEstimation(config)

# Parse the resulting data
data = viz.Parse(pe).data
print(data)

```

### 3.3.10 Working from the model.Model class directly

The pycotools3 tasks module contains classes for a bunch of copasi tasks that can be configured from python using pycotools. To simplify some of these tasks, wrappers have been build around these task classes in the `model.Model` class so that they can be used like a regular method.

Here I demonstrate some of these.

We first configure a model for the demonstration

```

import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz
seaborn.set_context(context='talk')

## Choose a directory for our model and analysis
working_directory = os.path.abspath('')

## In this model, A gets reversibly converted to B but the ↶
## backwards reaction is additionally regulated by C.
## B is reversibly converted into C.
antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
    B in Cell;
    C in Cell;

    // reactions
    R1: A => B ; Cell * k1 * A;
    R2: B => A ; Cell * k2 * B * C;
    R3: B => C ; Cell * k3 * B;
    R4: C => B ; Cell * k4 * C;

    // initial concentrations
    A = 100;
    B = 1;
    C = 1;

    // reaction parameters
    k1 = 0.1;
    k2 = 0.1;
    k3 = 0.1;
    k4 = 0.1;
end
"""

copasi_file = os.path.join(working_directory, 'example_model.cps')

## build model
mod = model.loada(antimony_string, copasi_file)

assert isinstance(mod, model.Model)

```

## Inserting parameters

```
dct = {  
    'k1': 55,  
    'k2': 36  
}  
mod.insert_parameters(parameter_dict=dct, inplace=True)
```

or

```
mod = mod.insert_parameters(parameter_dict=dct)
```

or

```
import pandas  
df = pandas.DataFrame(dct, index=[0])  
mod.insert_parameters(df=df, inplace=True)
```

or if the dataframe *df* has more than one parameter set we can specify the rank using the *index* argument.

```
import pandas  
##insert second best parameter set  
mod.insert_parameters(df=df, inplace=True, index=1)
```

---

**Note:** This is most useful when using `viz.Parse` output dataframes, which are `pandas.DataFrame` objects containing parameters in the columns and parameter sets in the rows, sorted by best RSS

---

or, assuming the variable *results\_directory* is a directory to a folder containing parameter estimation results.

```
mod.insert_parameters(parameter_path=results_directory, ↴  
    →inplace=True)
```

## Simulating a time course

```
data = mod.simulate(0, 10, 11)
```

Simulates a deterministic time course, 11 time points between 0 and 10. *data* contains a `pandas.DataFrame` object with variables along the columns and time points down the rows.

```
fname = os.path.join(os.path.dirname(__file__), 'simulation_data.csv'  
    ↴')  
## write data to file named fname  
data = mod.simulate(0, 10, 11, report_name=fname)
```

Like with the other shortcuts, arguments for the `tasks.TimeCourse` class are pass on.

```
data = mod.simulate(0, 10, 11, method='direct')
```

```
fname = ps.path.join(os.path.dirname(__file__), 'scan_results.csv')
mod.scan(variable='A', minimum=5, maximum=10, report_name=fname)
```

By default the scan type is set to ‘scan’. We can change this

```
fname = ps.path.join(os.path.dirname(__file__), 'scan_results.csv')
mod.simulate(0, 10, 11, method='direct', run_mode=False)
mod.scan(variable='A', scan_type='repeat',
         number_of_steps=10, report_name=fname,
         subtask='timecourse')
```

---

**Note:** In the `mod.simulate` we configure copasi to run a stochastic time course but do not execute. We then configure the repeat scan task to run the stochastic time course 10 times.

---

## Sensitivities

```
sens = mod.sensitivities(
    subtask='steady_state', cause='all_parameters',
    effect='all_variables'
)
```

### 3.3.11 Profile Likelihoods

Since a profile likelihood is just a parameter scan of parameter estimations, all we need to do to configure a profile likelihood analysis is to setup an appropriate `ParameterEstimation`. `Config` object and feed it into the `ParameterEstimation` class. This would be tedious to do manually but is easy with `ParameterEstimation.Context`

```
import os, glob
import pandas, numpy
import matplotlib.pyplot as plt
import seaborn
from pycotools3 import model, tasks, viz

working_directory = os.path.abspath('')

antimony_string = """
model simple_parameter_estimation()
    compartment Cell = 1;

    A in Cell;
```

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```

B in Cell;
C in Cell;

// reactions
R1: A => B ; Cell * k1 * A;
R2: B => A ; Cell * k2 * B * C;
R3: B => C ; Cell * k3 * B;
R4: C => B ; Cell * k4 * C;

// initial concentrations
A = 100;
B = 1;
C = 1;

// reaction parameters
k1 = 0.1;
k2 = 0.1;
k3 = 0.1;
k4 = 0.1;
end
"""

copasi_file = os.path.join(working_directory, 'example_model.cps')

## build model
mod = model.loada(antimony_string, copasi_file)

assert isinstance(mod, model.Model)

## simulate some data, returns a pandas.DataFrame
data = mod.simulate(0, 20, 1)

## write data to file
experiment_filename = os.path.join(working_directory, 'experiment_
→data.txt')
data.to_csv(experiment_filename)

```

The profile likelihood is calculated around the current parameter set in the model. If you want to change the current parameter set, maybe to the best fitting parameter set from a parameter estimation you can use the `InsertParameters` class. For now, we'll assume the best parameter set is already in the model.

```

with ParameterEstimation.Context(
    mod, experiment_filename,
    context='pl', parameters='gm'
) as context:
    context.set('method', 'hooke_jeeves')
    context.set('pl_lower_bound', 1000)
    context.set('pl_upper_bound', 1000)

```

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```
context.set('pe_number', 25) # number of steps in each profile
→likelihood
context.set('run_mode', True)
config = context.get_config()
```

We set the method to hooke and jeeves, a local optimiser which does well with profile likelihoods. We also set the *pl\_lower\_bound* and *pl\_upper\_bound* arguments to 1000 (which are defaults anyway). These are multipliers, not boundaries, of the profile likelihood. For instance, if the best estimated parameter for *A* was 1, then the profile likelihood would stretch from 1-e3 to 1e3.

Now, like with other parameter estimations we can simply do

```
ParameterEstimation(config)
```

Because the *context=pl* was used, pycotools knows to copy the model for each parameter, remove the parameter of interest from the parameter estimation task and create a scan of the parameter of interest.

### 3.3.12 Cross validation

Validation experiments are not used in model calibration. Instead the objective function is evaluated on validation data to see if the model can predict data it has not already seen. This can then be used as stopping criteria for the algorithm as we give a threshold for the closeness of the validation fits to simulations. This idea is common practice in machine learning and is used to prevent overfitting.

Cross validation is a new feature of pycotools3 but has been supported by COPASI for some years. The idea is to rotate calibration and validation datasets until you have tried all the combinations.

Cross validation can help identify datasets which do and don't fit well together. Here we create a model, simulate 3 datasets, make a data set up and use cross validation to infer the dataset that is made up.

```
# imports and create our antimony model string
from pycotools3 import model, tasks

working_directory = os.path.abspath('')
antimony_string = '''
    model negative_feedback()
        // define compartments
        compartment cell = 1.0
        //define species
        var A in cell
        var B in cell
        //define some global parameter for use in reactions
        vAProd = 0.1
```

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```

kADeg = 0.2
kBProd = 0.3
kBDeg = 0.4
//define initial conditions
A = 0
B = 0
//define reactions
AProd: => A; cell*vAProd
ADeg: A => ; cell*kADeg*A*B
BProd: => B; cell*kBProd*A
BDeg: B => ; cell*kBDeg*B
end
'''

# Create string to where we want to copasi model to go
copasi_file = os.path.join(os.path.dirname(__file__), 'negative_fb.
˓→cps')
mod = model.loada(antimony_string, copasi_file) # create the
˓→pycotools model

# create some filenames for experimental data
tc_fname1 = os.path.join(working_directory, 'timecourse1.txt')
tc_fname2 = os.path.join(working_directory, 'timecourse2.txt')
ss_fname1 = os.path.join(working_directory, 'steady_state1.txt')
ss_fname2 = os.path.join(working_directory, 'steady_state2.txt')

# simulate/create some experimental data
model.simulate(0, 5, 0.1, report_name=self.tc_fname1)
model.simulate(0, 10, 0.5, report_name=self.tc_fname2)
dct1 = {
    'A': 0.07,
    'B': 0.06,
    'C': 2.8
}
dct2 = {
    'A': 846,
    'B': 697,
    'C': 739
}
ss1 = pandas.DataFrame(dct1, index=[0])
ss1.to_csv(self.ss_fname1, sep='\t', index=False)
ss2 = pandas.DataFrame(dct2, index=[0])
ss2.to_csv(self.ss_fname2, sep='\t', index=False)

```

Configuring a cross validation experiment is similar to running parameter estimation or profile likelihoods: the difference is that you use `context='cv'` as argument to `ParameterEstimation.Context`.

`with tasks.ParameterEstimation.Context(`

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```

model, experiments, context='cv', parameters='gm'
) as context:
    context.set('randomize_start_values', True)
    context.set('method', 'genetic_algorithm')
    context.set('population_size', 20)
    context.set('number_of_generations', 50)
    context.set('validation_threshold', 500)
    context.set('cross_validation_depth', 1) ## 3/4 datasets
    →calibration; 1/4 for validation.
    context.set('copy_number', 3) #3 per model (5 models here)
    context.set('run_mode', True)
    context.set('lower_bound', 1e-3)
    context.set('upper_bound', 1e2)
    config = context.get_config()

pe = ParameterEstimation(config)
data = pycotools3.viz.Parse(pe).concat()

```

---

**Note:** The *cross\_validation\_depth* argument specifies how far to go combinatorially. For instance, when *cross\_validation\_depth*=2 and there are 4 datasets, all combinations of 2 datasets for experiments and 2 for validation will be applied.

---

**Warning:** While validation experiments are correctly configured with pycotools, there seems to be some instability in the current release of Copasi regarding multiple experiments in the *validation datasets* feature. Validation experiments work well when only one validation experiment is specified, but can crash when more than one is given.

---

**Note:** The *copy\_number* applies per model here. So 4 datasets, *cross\_validation\_depth*=1 means four models are configured for validation. Also configured is the model without any validation experiments for convenience.

---

The *validation\_weight* and *validation\_threshold* arguments are specific for validations. The copasi docs are vague on precisely what these mean but the higher the threshold, the more rigorous the validation.

## 3.4 API documentation

Here you will find detailed information about every module class and method in the pycotools3 package.

### 3.4.1 The model module

The pycotools3 model is of central importance in pycotools.

<i>Model</i>	Construct a pycotools3 model from a copasi file
<i>ImportSBML</i>	Import from sbml file
<i>InsertParameters</i>	Parse parameters into a copasi model
<i>BuildAntimony</i>	Build a copasi model using antimony
<i>Build</i>	Build a copasi model.

#### pycotools3.model.Model

```
class pycotools3.model.Model(copasi_file, quantity_type='concentration',
                             new=False, **kwargs)
```

Construct a pycotools3 model from a copasi file

The Model object is of central importance in pycotools as it extracts relevant information from a copasi file file into python.

These are *Model* attributes and properties:

#### Examples

```
>>> from pycotools3.model import Model
>>> model_path = r'/full/path/to/model.cps'
>>> model = Model(model_path) ##work in concentration units
>>> model = Model(model_path, quantity_type='particle_numbers')
→## work in particle numbers
```

Property	Description
copasi_file	Full path to model
root	Full path directory containing model
reference	Copasi model reference
time_unit	Time unit
name	Model name
volume_unit	Volume unit
quantity_unit	Quantity unit
area_unit	Area Unit
length_unit	Length unit
avagadro	Avagadro's number
key	Model key
states	List of states in correct order defined by copasi StateTemplate element.
fit_item_order	Order in which fit items appear
all_variable_names	list of reactions, metabolites, global_quantities local_parameters, compartment names as string
number_of_reactions	Number of reactions in model.Model

---

`__init__(copasi_file, quantity_type='concentration', new=False, **kwargs)`

#### Parameters

- `copasi_file (str)` – full path to a copasi file
- `quantity_type (str)` – either ‘concentration’ (default) or ‘particle\_numbers’
- `new (bool)` – True when constructing a new model

#### Methods

<code>__init__(copasi_file[, quantity_type, new])</code>	<code>param copasi_file</code> full path to a copasi file
<code>add(component_name, **kwargs)</code>	add a model component to the model
<code>add_compartment(compartment)</code>	Add compartment to model
<code>add_component(component_name, component[, ...])</code>	add a model component to the model
<code>add_function(function)</code>	Add function to model
<code>add_global_quantity(global_quantity)</code>	Add global quantity to model
<code>add_local_parameter(local_parameter)</code>	Add a local parameter to the model, specifically into the String=’kinetic Parameters’ section of parameter sets

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Table 2 – continued from previous page

<code>add_metabolite</code> (metab)	Add a metabolite to the model xml
<code>add_reaction</code> (reaction[, expression, rate_law])	<b>param reaction</b> py:class: <i>Reaction</i> or str. If str then
<code>add_state</code> (state, value)	Append state on to end of state template.
<code>convert_molar_to_particles</code> (mole, mol_unit, ...)	Convert molarity to particle numbers
<code>convert_particles_to_molar</code> (particles)	Converts particle numbers to Molarity.
<code>get</code> (component, value[, by])	Factory method for getting a model component by a value of a certain type
<code>get_variable_names</code> ([which, ...])	Get the names of variables in the model.
<code>insert_parameters</code> (**kwargs)	Wrapper around the InsertParameters class
<code>open</code> ([copasi_file, as_temp])	Open model with the gui.
<code>refresh()</code>	Save the file then reload the Model.
<code>remove</code> (component, name)	General factor method for removing model components
<code>remove_compartment</code> (value[, by])	Remove a compartment with the attribute given as the ‘by’ and value arguments
<code>remove_function</code> (value[, by])	remove a function from model
<code>remove_global_quantity</code> (value[, by])	Remove a global quantity from your model
<code>remove_metabolite</code> (value[, by])	Remove metabolite from model.
<code>remove_reaction</code> (value[, by])	Remove reaction
<code>remove_state</code> (state)	Remove state from StateTemplate and InitialState fields.
<code>reset_cache</code> (prop)	Delete property from cache then reset it
<code>save</code> ([copasi_file])	Save copasiML to copasi_filename.
<code>scan</code> ([inplace])	Perform a parameter scan on model
<code>set</code> (component, new_value[, ...])	Set a model components attribute to a new value
<code>simulate</code> (start, stop, by[, variables])	
<code>to_antimony()</code>	Returns antimony string of model.
<code>to_df()</code>	Convert kwargs to 1D df :return: pandas.DataFrame
<code>to_dict()</code>	get kwargs as dictionary :return: dict
<code>to_sbml</code> ([sbml_file])	convert model to sbml
<code>to_string()</code>	Produce kwargs as string format for using in __str__ methods in subclasses.

## Attributes

<code>active_parameter_set</code>	get active parameter set
<code>all_variable_names</code>	The names of all compartments, metabolites, global quantities, reactions and local parameters in the model.
<code>area_unit</code>	<i>str</i> . The currently defined area unit.
<code>avagadro</code>	Not really needed but good to check the consistancy of avagadros number.
<code>compartments</code>	Get list of model compartments
<code>constants</code>	Get list of constants from xml attribute ‘cn=”String=Kinetic Parameters”’
<code>copasi_file</code>	<code>Model</code>
<code>fit_item_order</code>	Get names of parameters being fitted in the order they appear
<code>functions</code>	get model functions
<code>global_quantities</code>	<i>list</i> each element is <code>GlobalQuantity</code>
<code>key</code>	Get the model reference - the ‘key’ from self.get_model_units
<code>length_unit</code>	<i>str</i>
<code>local_parameters</code>	Get local parameters in model.
<code>metabolites</code>	<i>list</i> . Each element is <code>Metabolite</code>
<code>name</code>	<i>str</i> . The model name
<code>number_of_reactions</code>	<i>int</i> number of reactions
<code>parameter_descriptions</code>	<i>list</i> . Each element a <code>ParameterDescription</code>
<code>parameter_sets</code>	Here for potential future implementation of easy switching between parameter sets :return:
<code>parameters</code>	get all locals, globals and metabs as pandas dataframe
<code>quantity_unit</code>	<i>str</i> . The currently defined quantity unit
<code>reactions</code>	assemble a list of reactions
<code>reference</code>	Get model reference from xml
<code>root</code>	Root directory for model.
<code>states</code>	The states (metabolites, globals, compartments) in the order they are read by Copasi from the StateTemplate element.
<code>time_unit</code>	<i>str</i> current time unit defined by copasi
<code>volume_unit</code>	<i>str</i> . The currently defined volume unit

**active\_parameter\_set**

get active parameter set

**Not really in use****Returns** `etree.Element`

Args:

Returns:

**add**(*component\_name*, *\*\*kwargs*)  
add a model component to the model

#### Parameters

- **component\_name** – str. i.e. ‘reaction’, ‘function’, ‘metabolite’
- **component** – py:class:*model.<component>*. The component class to add i.e. Metabolite
- **reaction\_expression** – When adding reaction using string as first arg, this argument takes the reaction expression (i.e. A -> B)
- **reaction\_rate\_law** – When adding reaction using string as first argument this argument takes the reaction rate law (i.e. k\*A)
- **\*\*kwargs** –

**Returns** py:class:‘Model’

**add\_compartment**(*compartment*)  
Add compartment to model

**Parameters** **compartment** – py:class:*Compartment*

**Returns** py:class:*Model*

**add\_component**(*component\_name*, *component*, *reaction\_expression=None*,  
*reaction\_rate\_law=None*)  
add a model component to the model

#### Parameters

- **component\_name** – str. i.e. ‘reaction’, ‘function’, ‘metabolite’
- **component** – py:class:*model.<component>*. The component class to add i.e. Metabolite
- **reaction\_expression** – When adding reaction using string as first arg, this argument takes the reaction expression (i.e. A -> B) (Default value = None)
- **reaction\_rate\_law** – When adding reaction using string as first argument this argument takes the reaction rate law (i.e. k\*A) (Default value = None)

**Returns** class:‘Model’

**Return type** py

**add\_function**(*function*)  
Add function to model

**Parameters** `function` – py:class:*Function*.

**Returns** py:class:*Model*

**add\_global\_quantity** (*global\_quantity*)

Add global quantity to model

**Parameters** `global_quantity` – str‘ or GlobalQuantity. If str

is the name of global\_quantity to add and default GlobalQuantity properties are adopted. If GlobalQuantity, a GlobalQuantity instance must be pre-built and passes as arg.

**Returns** py:class:*Model*

**add\_local\_parameter** (*local\_parameter*)

Add a local parameter to the model, specifically into the String='kinetic Parameters' section of parameter sets

**Parameters** `local_parameter` – py:class:*LocalParameter*

**Returns** py:class:*Model*

**add\_metabolite** (*metab*)

Add a metabolite to the model xml

**Parameters** `metab` – str‘ or Metabolite. If str

is the name of metabolite to add and default Metabolite properties are adopted. If Metabolite, a Metabolite instance must be prebuilt and passes as arg.

**Returns** py:class:*Model*

**add\_reaction** (*reaction*, *expression=None*, *rate\_law=None*)

**Parameters** `reaction` – py:class:*Reaction* or str. If str then

**must be the name of the reaction.** expression: (Default value = None) rate\_law: (Default value = None)

**Returns** py:class:*Model*

**add\_state** (*state*, *value*)

Append state on to end of state template. Used within add\_metabolite and add\_global\_quantity. Shouldn't need to use manually

**Parameters**

- `state` – str‘. A valid key
- `value` – int‘, float. Value for state

Returns:

## **all\_variable\_names**

The names of all compartments, metabolites, global quantities, reactions and local parameters in the model.

**Returns** *list*. Each element is *str*

Args:

Returns:

## **area\_unit**

*str*. The currently defined area unit.

Args:

Returns:

**Type** return

## **avagadro**

Not really needed but good to check the consistency of avogadros number.

This number was changed between version 16 and 19 and caused a bug

Args:

Returns:

## **compartments**

Get list of model compartments

**Returns** *list*. Each element is Compartmet

Args:

Returns:

## **constants**

Get list of constants from xml attribute ‘cn=”String=Kinetic Parameters”’

**Returns** *list* each element LocalParameter

## **static convert\_molar\_to\_particles(moles, mol\_unit, compartment\_volume)**

Convert molarity to particle numbers

### **Parameters**

- **moles** – int‘ or float. Number of moles in mol\_unit to convert
- **mol\_unit** – str‘. Mole unit to convert from.

**suppoorted: fmol, pmol, nmol, umol, mmol or mol** compartment\_volume: int‘ or float. Volume of compartment containing specie to convert

**Returns** int‘. number of particles

```
static convert_particles_to_molar(particles, mol_unit, compartment_volume)
```

Converts particle numbers to Molarity.

##TODO build support for copasi's newest units

#### Parameters

- **particles** – int‘ Number of particles to convert
- **mol\_unit** – str‘. The quantity unit, i.e:

**fmol, pmol, nmol, umol, mmol or mol** compartment\_volume: int‘, float. Volume of compartment containing specie to convert

**Returns** float‘. Molarity

### **copasi\_file**

*Model*

**Type** Returns

#### **fit\_item\_order**

Get names of parameters being fitted in the order they appear

**Returns** list

Args:

Returns:

#### **functions**

get model functions

**Returns** list each element a py:class:‘Function

#### **get (component, value, by='name')**

Factory method for getting a model component by a value of a certain type

#### Parameters

- **component** – str‘. The component i.e. metabolite or local\_parameter
- **value** – str‘. Value of the attribute to match by i.e. metabolite called A
- **by** – str‘. Which attribute to search by. i.e. name or key or value (Default value = ‘name’)

#### Returns

py:class:Model.<component>‘

Get reaction called A2B:

Get metabolite called A:

Get all reactions which have a fixed simulation\_type:

Get all compartments with an initial value of 15 (concentration or particles depending on quantity\_type):

Get metabolites in the nucleus compartment:

**Return type** Instance of ‘

```
>>> model.get('reaction', 'A2B', by='name')
```

```
>>> model.get('metabolite', 'A', by='name')
```

```
>>> model.get('global_quantity', 'fixed', by='simulation_<br>type')
```

```
>>> model.get('compartment', 15, by='initial_value')
```

```
>>> model.get('metabolite', 'nuc', by='compartment')
```

## **get\_model\_object** (*string*)

Retrieve a model object, such as a parameter or metabolite :param string: name of parameter :type string: str

**Returns** correct model object

## **get\_parameters\_as\_antimony** ()

get parametes as antimony string Returns (string):

## **get\_variable\_names** (*which*=’a’, *include\_assignments*=True, *prefix*=None)

Get the names of variables in the model. If include\_assignments is off these are ommited from the results (this is useful for ParameterEstimation) as they are not generally estimated. Prefix provides a way of filtering the returned list

### Parameters

- **which** – string. Default=’a’. A string containing any or all of characters ‘a’, ‘m’, ‘g’, ‘l’, ‘c’ for all, metabolites, global\_quantities, local\_parameters and compartments respectively
- **include\_assignments** – Boolean. Default=True. If True, return global variables with assignments
- **prefix** – str. Default=None. If given, returned parameter names are filtered to only include parameter with *prefix* at the begining.

**Returns** ‘list’ of variable names

## **global\_quantities**

*list* each element is GlobalQuantity

Args:

Returns:

Type return

**insert\_parameters** (\*\*kwargs)

Wrapper around the InsetParameters class

**Parameters**

- **kwargs** – Arguments for InsertParameters
- **\*\*kwargs** –

**Returns** py:class:Model

**key**

Get the model reference - the ‘key’ from self.get\_model\_units

**Returns** str

Args:

Returns:

**length\_unit**

str

Args:

Returns:

Type return

**local\_parameters**

Get local parameters in model. local\_parameters are those which are actively used in reactions and do not have a global variable assigned to them. The constant property returns all local parameters regardless of simulation type (fixed or assignment)

**Returns** list. Each element is LocalParameter

Args:

Returns:

**metabolites**

list. Each element is Metabolite

Args:

Returns:

Type return

**name**

str. The model name

Args:

Returns:

**Type** return

## **number\_of\_reactions**

*int* number of reactions

Args:

Returns:

**Type** return

## **open (copasi\_file=None, as\_temp=False)**

Open model with the gui. In order to work the environment variables must be properly set so that the command *CopasiUI* in the terminal or command prompt opens the model.

First [\*Model.save\(\)\*](#) the model to copasi\_file then open with CopasiUI. Optionally open with a temporary filename.

### **Parameters**

- **copasi\_file** – str‘ or *None*. Same as `model.Save()` (Default value = *None*)
- **as\_temp** – bool‘. Use temp file to open the model and remove afterwards (Default value = *False*)

**Returns** None‘

## **parameter\_descriptions**

*list*. Each element a ParameterDescription

Args:

Returns:

**Type** return

## **parameter\_sets**

Here for potential future implementation of easy switching between parameter sets  
:return:

Args:

Returns:

## **parameters**

get all locals, globals and metabs as pandas dataframe

**Returns** pandas.DataFrame

Args:

Returns:

## **quantity\_unit**

*str*. The currently defined quantity unit

Args:

Returns:

**Type** return

### **reactions**

assemble a list of reactions

**Returns** *list* each element a Reaction

### **reference**

Get model reference from xml

**Returns** str

Args:

Returns:

### **refresh()**

Save the file then reload the Model. Can't use the save method though because the save method uses the refresh method. :return:

Args:

Returns:

### **remove(component, name)**

General factor method for removing model components

#### **Parameters**

- **component** – str‘ which component to remove (i.e. metabolite)
- **name** – str‘ name of component to remove

**Returns** py:class:Model

### **remove\_compartment(value, by='name')**

Remove a compartment with the attribute given as the ‘by’ and value arguments

#### **Parameters**

- **value** – str‘. Value of attribute to match i.e. ‘Nucleus’
- **by** – str‘ which attribute to match i.e. ‘name’ or ‘key’ (Default value = ‘name’)

**Returns** py:class:Model

### **remove\_function(value, by='name')**

remove a function from model

#### **Parameters**

- **value** – str‘ value of attribute to match (i.e the functions name)

- **by** – str‘ which attribute to match by. default=’name’

**Returns** py:class:*model.Model*

**remove\_global\_quantity** (*value*, *by*=’name’)

Remove a global quantity from your model

#### Parameters

- **value** – value to match by (i.e. ProteinA or ProteinB)
- **by** – attribute to match (i.e. name or key) (Default value = ‘name’)

**Returns** py:class:*model.Model*

**remove\_metabolite** (*value*, *by*=’name’)

Remove metabolite from model.

#### Parameters

- **value** – str‘. Attribute value to remove
- **by** – str‘ Any metabolite attribute type to match (Default value = ‘name’)

#### Returns

py:class:*Model*

Usage: ## Remove attribute called ‘A’

## Remove metabolites with initial concentration of 0

```
>>> model.remove_metabolite('A', by='name')
```

```
>>> model.remove_metabolite(0, by='concentration')
```

**remove\_reaction** (*value*, *by*=’name’)

Remove reaction

#### Parameters

- **value** – str‘. Value of attribute
- **by** – attribute of reaction to match default=’name

*str* which :py:class‘Reaction‘ atrribute to match

**Returns** py:class:*Model*

**remove\_state** (*state*)

Remove state from StateTemplate and InitialState fields. USed for deleting metabolites and global quantities.

**Parameters** **state** – str‘. key of state to remove (i.e. Metabolite\_1)

**Returns** py:class:*Model*

**reset\_cache** (*prop*)

Delete property from cache then reset it

**Parameters** **prop** – str‘. property to reset

**Returns** py:class:*Model*

**root**

Root directory for model. The directory where copasi\_file is saved.

Does not need a setter since root is derived from copasi\_file property

**Returns** *str*

Args:

Returns:

**save** (*copasi\_file=None*)

Save copasiML to copasi\_filename.

**Parameters** **copasi\_filename** – str‘ or *None*. Default is *None*.

When *None* defaults to same filepath the model came from. If another path, saves to that path. *copasi\_file*: (Default value = None)

**Returns** py:class:*Model*

**scan** (*inplace=False, \*\*kwargs*)

Perform a parameter scan on model

This is a wrapper around tasks.Scan and accepts all of the same arguments, except the model which is already provided.

**Parameters** **\*\*kwargs** –

Returns:

**sensitivities** (*inplace=False, \*\*kwargs*)

Perform a sensitivity analysis on model

This is a wrapper around tasks.Sensitivities and accepts all of the same arguments, except the model which is already provided.

**Parameters** **\*\*kwargs** –

Returns:

**set** (*component, match\_value, new\_value, match\_field='name', change\_field='name'*)

Set a model components attribute to a new value

**Parameters**

- **component** – str‘ type of component to change (i.e. metabo-lite)

- **match\_value** – str‘, *int*, *float* depending on value of *match\_field*.

**The value to match.** new\_value: str‘, int or float depending on value of *match\_field*

**new value for component attribute** match\_field: str‘. The attribute of component to match by. (Default value = ‘name’) change\_field: str‘ The attribute of the component matched that you want to change? (Default value = ‘name’)

## Returns

py:class:*Model*

Set initial concentration of metabolite called ‘X’ to 50:

Set name of global quantity called ‘G’ to ‘H’:

```
>>> model.set('metabolite', 'X', 50, match_field='name',
   ↪change_field='concentration')
```

```
>>> model.set('global_quantity', 'G', 'H', match_field='name
   ↪', change_field='name')
```

## states

The states (metabolites, globals, compartments) in the order they are read by Co-pasi from the StateTemplate element.

## Returns *OrderedDict*

Args:

Returns:

## time\_unit

str current time unit defined by copasi

Args:

Returns:

Type return

## to\_antimony()

Returns antimony string of model. Wrapper around tellurium functions :return:

Args:

Returns:

## to\_sbml(*sbml\_file=None*)

convert model to sbml

**Parameters** **sbml\_file** – str‘. Path for SBML. Defaults to same as copasi filename

**Returns** str‘. Path to smbl file

**to\_tellurium()**

return a roadrunner model via the tellurium package Returns:

**volume\_unit**

*str*. The currently defined volume unit

Args:

Returns:

**Type** return

**pycotools3.model.ImportSBML****class pycotools3.model.ImportSBML (sbml\_file, copasi\_file=None)**

Import from sbml file

Accepts an SBML file, converts it to copasi format and reads it into a Model object

**\_\_init\_\_(sbml\_file, copasi\_file=None)****Parameters**

- **sbml\_file** (*str*) – path to sbml
- **copasi\_file** (*None, str*) – Default is None and pyco-tools automatically creates a copasi model with the same name as the sbml file. Otherwise, a path to copasi\_file.

**Methods****\_\_init\_\_(sbml\_file[, copasi\_file])**

**param sbml\_file** path to  
sbml

**convert()**

Perform conversion using CopasiSE :return:

**copasi\_filename()****load\_model()****convert()**

Perform conversion using CopasiSE :return:

Args:

Returns:

**copasi\_filename()****load\_model()**

## pycotools3.model.InsertParameters

```
class pycotools3.model.InsertParameters(model, parameter_dict=None, df=None, parameter_path=None, index=0, quantity_type='concentration', inplace=False)
```

Parse parameters into a copasi model

Insert parameters from a file, dictionary or a pandas dataframe into a copasi file.

```
__init__(model, parameter_dict=None, df=None, parameter_path=None, index=0, quantity_type='concentration', inplace=False)
```

### Parameters

- **model** ([Model](#)) – The model to parse parameters into
- **parameter\_dict** ([dict](#)) – Default None. If not None, dict[parameter\_name] = parameter\_value
- **df** ([pandas.DataFrame](#)) – Default None. If not None, a dataframe containing parameters to insert
- **parameter\_path** ([str](#)) – Default None. If not None a path to parameter estimation output file
- **index** ([int](#)) – Default 0 (best RSS). When multiple parameter sets available, rank of best fit you want to insert
- **quantity\_type** ([str](#)) – concentration (default) or particle\_numbers
- **inplace** ([bool](#)) – Whether to operate inplace or return a new model

### Methods

---

```
__init__(model[, parameter_dict, df, ...])
```

**param model** The model to parse parameters into

---

```
insert()
```

User other methods defined in this class to insert parameters into the model :return:

---

```
insert_compartments()
```

insert new parameters into compartment :return:

---

```
insert_global_quantities()
```

insert new parameters into compartment :return:

Continued on next page

Table 5 – continued from previous page

---

<code>insert_locals()</code>	<b>return</b>
<code>insert_metabolites()</code>	insert new parameters into compartment :return:
<code>read_model(m)</code>	<b>param m</b>
<code>to_dict()</code>	Args:

---

## Attributes

---

<code>parameters</code>	Get parameters depending on the type of input.
-------------------------	--

---

### `insert()`

User other methods defined in this class to insert parameters into the model :return:

Args:

Returns:

### `insert_compartments()`

insert new parameters into compartment :return:

Args:

Returns:

### `insert_global_quantities()`

insert new parameters into compartment :return:

Args:

Returns:

### `insert_locals()`

#### Returns

### `insert_metabolites()`

insert new parameters into compartment :return:

Args:

Returns:

### `parameters`

Get parameters depending on the type of input. Converge on a pandas dataframe.  
Columns = parameters, rows = parameter sets

Use check parameter consistency to see whether headers have been pruned or not.  
If not try pruning them

Args:

Returns:

**to\_dict()**

Args:

**Returns** return:

## pycotools3.model.BuildAntimony

**class** pycotools3.model.**BuildAntimony**(*copasi\_file*: str)

Build a copasi model using antimony

A context manager to create a copasi model *copasi\_file* using the antimony language

### Examples

```
working_directory = os.path.dirname(__file__)
copasi_filename = os.path.join(working_directory,
    ↪'NegativeFeedbackModel.cps')
with model.BuildAntimony(copasi_filename) as loader:
    negative_feedback = loader.load(
        '''
        model negative_feedback()
            // define compartments
            compartment cell = 1.0
            //define species
            var A in cell
            var B in cell
            //define some global parameter for use in reactions
            vAProd = 0.1
            kADeg = 0.2
            kBProd = 0.3
            kBDeg = 0.4
            //define initial conditions
            A = 0
            B = 0
            //define reactions
            AProd: => A; cell*vAProd
            ADeg: A =>; cell*kADeg*A*B
            BProd: => B; cell*kBProd*A
            BDeg: B =>; cell*kBDeg*B
        end
        '''
    )
print(negative_feedback)
```

**\_\_init\_\_**(*copasi\_file*: str)

**Parameters** `copasi_file` (`str`) – Path to a valid location on disk to store the copasi file

## Methods

---

`__init__(copasi_file)`

**param** `copasi_file` Path to a valid location on disk to store the copasi file

---

`load(antimony_str)`

Load the antimony string `antimony_str` into a `copasi_file` and `Model`.

**load** (`antimony_str`)

Load the antimony string `antimony_str` into a `copasi_file` and `Model`.

**Args** `antimony_str` (`str`): A valid antimony string encoding a model

**return**

`model` (`Model`) A PyCoTools model containing the model defined in the :parameter:`'antimony\_str'`.

**Parameters** `antimony_str` –

Returns:

## pycotools3.model.Build

**class** `pycotools3.model.Build(copasi_file)`

Build a copasi model.

Context manager for building a copasi model with only PyCoTools Functions.

Users should also see `BuildAntimony`

---

`__init__(copasi_file)`

Initialize self. See help(type(self)) for accurate signature.

## Methods

---

`__init__(copasi_file)`

Initialize self.

### 3.4.2 The tasks module

<code>TimeCourse</code>	Simulate a time course
<code>ParameterEstimation.Config</code>	A parameter estimation configuration class
<code>ParameterEstimation</code>	Interface to COPASI's parameter estimation task
<code>ParameterEstimation.Context</code>	A high level interface to create a <code>ParameterEstimation.Config</code> object.
<code>Sensitivities</code>	Interface to COPASI sensitivity task
<code>Scan</code>	Interface to COPASI scan task
<code>Reports</code>	Creates reports in copasi output specification section.

## pycotools3.tasks.TimeCourse

```
class pycotools3.tasks.TimeCourse (model, **kwargs)
```

Simulate a time course

A class for running a time course from python using a copasi model. All but one of copasi's solvers are supported and available via the *method* kwarg.

TimeCourse Kwargs	Description
intervals	Default: 100
step_size	Default: 0.01
end	Default: 1
start	Default: 0
update_model	Default: False
method	Default: deterministic
output_event	Default: False
scheduled	Default: True
automatic_step_size	Default: False
start_in_steady_state	Default: False
inte- grate_reduced_model	Default: False
relative_tolerance	Default: 1e-6
absolute_tolerance	Default: 1e-12
max_internal_steps	Default: 10000
max_internal_step_size	Default: 0
subtype	Default: 2
use_random_seed	Default: True
random_seed	Default: 1
epsilon	Default: 0.001
lower_limit	Default: 800
upper_limit	Default: 1000
partitioning_interval	Default: 1
runge_kutta_step_size	Default: 0.001
run	Default: True
correct_headers	Default: True
save	Default: False
<report_kwargs>	Arguments for :ref:`report_kwargs` <report_kwargs> are also accepted here

Args:

Returns:

`__init__(model, **kwargs)`

Initialize self. See help(type(self)) for accurate signature.

## Methods

<code>__init__(model, **kwargs)</code>	Initialize self.
<code>adaptive_tau_leap()</code>	<code>return</code>

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<code>check_integrity(allowed, given)</code>	Method to raise an error when a wrong kwarg is passed to a subclass
<code>convert_bool_to_numeric(dct)</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>convert_bool_to_numeric2()</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>create_task()</code>	Begin creating the segment of xml needed for a time course.
<code>deterministic()</code>	:return:lxml.etree._Element
<code>direct()</code>	<b>return</b>
<code>get_report_key()</code>	cross reference the timecourse task with the newly created time course report to get the key
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pycotools3 variable
<code>gibson_bruck()</code>	<b>return</b>
<code>hybrid_lsoda()</code>	<b>return</b>
<code>hybrid_rk45()</code>	<b>return</b>
<code>hybrid_runge_kutta()</code>	<b>return</b>
<code>read_model(m)</code>	<b>param m</b>
<code>set_report()</code>	Set a time course report containing time and all species or global quantities defined by the user.
<code>set_timecourse()</code>	Set method specific sections of xml.
<code>simulate()</code>	
<code>tau_leap()</code>	<b>return</b>
<code>update_properties(kwags)</code>	method for updating properties from kwags

## Attributes

---

schema

---

**adaptive\_tau\_leap()**

**Returns**

**create\_task()**

Begin creating the segment of xml needed for a time course. Define task and problem definition. This section of xml is common to all methods :return: lxml.etree.\_Element

Args:

Returns:

**deterministic()**

:return:lxml.etree.\_Element

**direct()**

**Returns**

**get\_report\_key()**

cross reference the timecourse task with the newly created time course report to get the key

Args:

Returns:

**gibson\_bruck()**

**Returns**

**hybrid\_lsoda()**

**Returns**

**hybrid\_rk45()**

**Returns**

**hybrid\_runge\_kutta()**

**Returns**

**set\_report()**

Set a time course report containing time and all species or global quantities defined by the user.

**Returns** pycotools3.model.Model

Args:

Returns:

**set\_timecourse()**

Set method specific sections of xml. This is a method element after the problem element that looks like this:

**Returns** lxml.etree.\_Element

Args:

Returns:

**simulate()**

**tau\_leap()**

**Returns**

## pycotools3.tasks.ParameterEstimation.Config

```
class pycotools3.tasks.ParameterEstimation.Config(models,
                                                    datasets,
                                                    items,    set-
                                                    tings={}, de-
                                                    faults=None)
```

A parameter estimation configuration class

Stores all the settings needed for configuration of a parameter estimation using COPASI. Base class is a Bunch

**Structure of a ParameterEstimation.Config Object has four main sections:**

- models
- datasets
- items
- settings

## Examples

```
>>> ## create a model
>>> antimony_string = """
...     model TestModel1()
...         R1: A => B; k1*A;
...         R2: B => A; k2*B
...         A = 1
...         B = 0
...         k1 = 4;
...         k2 = 9;
...
...     end
...
...
>>> copasi_filename = os.path.join(os.path.dirname(__file__),
...                                 'example_model.cps')
>>> mod = moddel.loada(antimony_string, copasi_filename)
>>> ## Simulate some data from the model and write to file
```

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```

>>> fname = os.path.join(os.path.dirname(__file__), 'timeseries.
   ↵txt')
>>> data = self.model.simulate(0, 10, 11)
>>> data.to_csv(fname)
>>>
>>> ## create nested dict containing all the relevant arguments
   ↵for your configuration
>>> config_dict = dict(
...     models=dict(
...         ## model name is the users choice here
...         example1=dict(
...             copasi_file=copasi_filename
...         )
...     ),
...     datasets=dict(
...         experiments=dict(
...             ## experiment names are the users choice
...             report1=dict(
...                 filename=self.TC1.report_name,
...             ),
...         ),
...         ## our validations entry is empty here
...         ## but if you have validation data this should
...         ## be the same as the experiments section
...         validations=dict(),
...     ),
...     items=dict(
...         fit_items=dict(
...             A=dict(
...                 affected_experiments='report1'
...             ),
...             B=dict(
...                 affected_validation_experiments=['report2
   ↵']
...             ),
...             k1={},
...             k2={},
...         ),
...         constraint_items=dict(
...             k1=dict(
...                 lower_bound=1e-2,
...                 upper_bound=10
...             )
...         )
...     ),
...     settings=dict(
...         method='genetic_algorithm_sr',
...         population_size=2,
...         number_of_generations=2,

```

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```

...
working_directory=os.path.dirname(__file__),
...
copy_number=4,
pe_number=2,
weight_method='value_scaling',
validation_weight=2.5,
validation_threshold=9,
randomize_start_values=True,
calculate_statistics=False,
create_parameter_sets=False
)
...
)
>>> config = ParameterEstimation.Config(**config_dict)

```

## `__init__(models, datasets, items, settings={}, defaults=None)`

Initialisation method for Config class

### Parameters

- **models** (*dict*) – Dict containing model names and paths to copasi files
- **datasets** (*dict*) – Dict containing experiments and validation experiments
- **items** (*dict*) – Dict containing fit items and constraint items
- **settings** (*dict*) – Dict containing all other settings for parameter estimation
- **defaults** (*ParameterEstimation.\_Defaults*) – Custom set of Defaults to use for unspecified arguments

Returns:

## Methods

<code>__init__(models, datasets, items[, ...])</code>	Initialisation method for Config class
<code>check_integrity(allowed, given)</code>	Method to raise an error when a wrong kwarg is passed to a subclass
<code>clear()</code>	
<code>configure()</code>	Configure the class for production of parameter estimation config
<code>convert_bool_to_numeric(dct)</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>convert_bool_to_numeric2()</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>copy()</code>	

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<code>fromDict(d)</code>	Recursively transforms a dictionary into a Munch via copy.
<code>fromYAML(*args, **kwargs)</code>	
<code>from_json(string)</code>	Create config object from json format :param string: a valid json string :type string: Str
<code>from_yaml(yml)</code>	Read config object from yaml file :param yml: full path to text file containing configuration arguments in yaml format :type yml: str
<code>fromkeys</code>	Returns a new dict with keys from iterable and values equal to value.
<code>get(k,d)</code>	
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pyco-tools3 variable
<code>items()</code>	
<code>keys()</code>	
<code>pop(k,d)</code>	If key is not found, d is returned if given, otherwise KeyError is raised
<code>popitem()</code>	2-tuple; but raise KeyError if D is empty.
<code>set_default_fit_items_dct()</code>	Configure missing entries for items.fit_items when they are in nested dict format
<code>set_default_fit_items_str()</code>	Configure missing entries for items.fit_items when they are strings pointing towards model variables
<code>setdefault(k,d)</code>	
<code>toDict()</code>	Recursively converts a munch back into a dictionary.
<code>toJSON(**options)</code>	Serializes this Munch to JSON.
<code>toYAML(**options)</code>	Serializes this Munch to YAML, using <code>yaml.safe_dump()</code> if no <code>Dumper</code> is provided.
<code>to_json()</code>	Output arguments as json
<code>to_yaml([filename])</code>	Output arguments as yaml
<code>update([E, ]**F)</code>	If E is present and has a .keys() method, then does: for k in E: D[k] = E[k] If E is present and lacks a .keys() method, then does: for k, v in E: D[k] = v In either case, this is followed by: for k in F: D[k] = F[k]
<code>update_properties(kwargs)</code>	method for updating properties from kwargs
<code>values()</code>	

## Attributes

<code>constraint_items</code>	The constraint items as nested dict
<code>experiment_filenames</code>	A list of experiment filenames
<code>experiment_names</code>	A list of experiment names
<code>experiments</code>	The experiments property :returns: datasets.experiments as dict
<code>fit_items</code>	The fit items as nested dict
<code>model_objects</code>	A list of model objects for mapping
<code>schema</code>	
<code>validation_filenames</code>	a list of validation filenames
<code>validation_names</code>	A list of validation names
<code>validations</code>	The validations property :returns: datasets.validations as dict

### `configure()`

Configure the class for production of parameter estimation config

Like a main method for this class. Uses the other methods in the class to configure a ParameterEstimation.Config object

**Returns** Operates inplace and returns None

### `constraint_items`

The constraint items as nested dict

**Type** Returns

### `experiment_filenames`

A list of experiment filenames

**Type** Returns

### `experiment_names`

A list of experiment names

**Type** Returns

### `experiments`

The experiments property :returns: datasets.experiments as dict

### `fit_items`

The fit items as nested dict

**Type** Returns

### `from_json(string)`

Create config object from json format :param string: a valid json string :type string: Str

**Returns** ParameterEstimation.Config

**from\_yaml** (*yml*)

Read config object from yaml file :param *yml*: full path to text file containing configuration arguments in yaml format :type *yml*: str

**Returns** ParameterEstimation.Config

**model\_objects**

A list of model objects for mapping

**Type** Returns

**models\_affected\_experiments**

Get which experiment datasets affect which models

**Returns** dict. Keys are model names, value are list of affected models

**models\_affected\_validation\_experiments**

Get which experiment datasets affect which models

**Returns** dict. Keys are model names, value are list of affected models

**multi\_experiments**

List of experiment names that have more than one experiment separated by blank line Returns (tuple): (list, int)

**set\_default\_fit\_items\_dct()**

Configure missing entries for items.fit\_items when they are in nested dict format

**Returns** None. Method operates inplace on class attributes

**set\_default\_fit\_items\_str()**

Configure missing entries for items.fit\_items when they are strings pointing towards model variables

**Returns** None. Method operates inplace on class attributes

**to\_json()**

Output arguments as json

**Returns:** str All arguments in json format

**to\_yaml** (*filename=None*)

Output arguments as yaml

**Parameters** **filename** (*str*, *None*) – If not None (default), path to write yaml configuration to

**Returns** Config object as string in yaml format

**validation\_filenames**

a list of validation filenames

**Type** Returns

**validation\_names**

A list of validation names

**Type** Returns

## validations

The validations property :returns: datasets.validations as dict

## pycotools3.tasks.ParameterEstimation

**class** pycotools3.tasks.**ParameterEstimation** (*config*)

Interface to COPASI's parameter estimation task

### Examples

Assuming a *ParameterEstimation.Config* object has been configured and is called *config* >>> pe = ParameterEstimation(config)

**\_\_init\_\_** (*config*)

Configure a the parameter estimation task in copasi

Pycotools supports all the features of parameter estimation configuration as copasi, plus a few additional ones (such as the affected models setting).

**Parameters config** (*ParameterEstimation.Config*) – An appropriately configured *ParameterEstimation.Config* class

### Examples

See *ParameterEstimation.Config* or *ParameterEstimation.Context* for detailed information on how to produce a *ParameterEstimation.Config* object. Note that the *ParameterEstimation.Context* class is higher level and should be the preferred way of constructing a *ParameterEstimation.Config* object while the *ParameterEstimation.Config* class gives you the same level of control as copasi but is bulkier to write.

Assuming the *ParameterEstimation.Config* class has already been created >>> pe = ParameterEstimation(config)

### Methods

<b>__init__(config)</b>	Configure a the parameter estimation task in copasi
<b>check_integrity(allowed, given)</b>	Method to raise an error when a wrong kwarg is passed to a subclass
<b>convert_bool_to_numeric(dct)</b>	CopasiML uses 1's and 0's for True or False in some but not all places.
<b>convert_bool_to_numeric2()</b>	CopasiML uses 1's and 0's for True or False in some but not all places.

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<code>do_checks()</code>	validate integrity of user input
<code>get_component(model, component, string)</code>	Get component called string from model
<code>get_model_objects_from_string(string)</code>	Get model objects from the strings provided by the user in the Config class :return: list of <i>model.Model</i> objects
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pycotools3 variable
<code>read_model(m)</code>	<b>param m</b>
<code>run(models)</code>	Run a parameter estimation using command line copasi.
<code>update_properties(kwargs)</code>	method for updating properties from kwargs

## Attributes

<code>fit_dir</code>	Property holding the directory where the parameter estimation fitting occurs.
<code>global_quantities</code>	list of strings of global quantities present in the models
<code>local_parameters</code>	list of strings of local parameters in the model
<code>metabolites</code>	list of strings of metabolites in the model
<code>models</code>	Get models
<code>models_dir</code>	A directory containing models
<code>problem_dir</code>	Property holding the directory where the parameter estimation problem is stored :returns: str.
<code>results_directory</code>	A directory containing results, parameter estimation report files from copasi
<code>schema</code>	
<code>valid_methods</code>	

**class Config** (*models, datasets, items, settings={}, defaults=None*)

A parameter estimation configuration class

Stores as attributes all the settings needed for configuration of a parameter estimation using COPASI. Base class is a Bunch

**Structure of a *ParameterEstimation.Config* Object has four main sections:**

- models
- datasets

- items
- settings

## Examples

```
>>> ## create a model
>>> antimony_string = """
...             model TestModel1()
...                 R1: A => B; k1*A;
...                 R2: B => A; k2*B
...                 A = 1
...                 B = 0
...                 k1 = 4;
...                 k2 = 9;
...
...             end
...
...
>>> copasi_filename = os.path.join(os.path.dirname(__file__),
...                                  'example_model.cps')
>>> mod = moddel.loada(antimony_string, copasi_filename)
>>> ## Simulate some data from the model and write to file
>>> fname = os.path.join(os.path.dirname(__file__), 
...                      'timeseries.txt')
>>> data = self.model.simulate(0, 10, 11)
>>> data.to_csv(fname)
>>>
>>> ## create nested dict containing all the relevant arguments for your configuration
>>> config_dict = dict(
...     models=dict(
...         ## model name is the users choice here
...         example1=dict(
...             copasi_file=copasi_filename
...         )
...     ),
...     datasets=dict(
...         experiments=dict(
...             ## experiment names are the users choice
...             report1=dict(
...                 filename=self.TC1.report_name,
...             ),
...         ),
...         ## our validations entry is empty here
...         ## but if you have validation data this should
...         ## be the same as the experiments section
...         validations=dict(),
...     ),
...     items=dict(
...         fit_items=dict(
...             
```

(continues on next page)

(continued from previous page)

```

...
        A=dict(
            ...
                affected_experiments='report1'
            ),
            B=dict(
                ...
                    affected_validation_experiments=[
            ↵'report2']
                ...
            ),
            k1={ },
            k2={ },
        ),
        constraint_items=dict(
            ...
            k1=dict(
                ...
                    lower_bound=1e-2,
                    upper_bound=10
                )
            )
        ),
        settings=dict(
            ...
            method='genetic_algorithm_sr',
            population_size=2,
            number_of_generations=2,
            working_directory=os.path.dirname(__file__),
            copy_number=4,
            pe_number=2,
            weight_method='value_scaling',
            validation_weight=2.5,
            validation_threshold=9,
            randomize_start_values=True,
            calculate_statistics=False,
            create_parameter_sets=False
        )
    )
)
>>> config = ParameterEstimation.Config(**config_dict)

```

**configure()**

Configure the class for production of parameter estimation config

Like a main method for this class. Uses the other methods in the class to configure a `ParameterEstimation.Config` object

**Returns** Operates inplace and returns None

**constraint\_items**

The constraint items as nested dict

**Type** Returns

**experiment\_filenames**

A list of experiment filenames

**Type** Returns

**experiment\_names**

A list of experiment names

**Type** Returns

**experiments**

The experiments property :returns: datasets.experiments as dict

**fit\_items**

The fit items as nested dict

**Type** Returns

**from\_json (string)**

Create config object from json format :param string: a valid json string :type string: Str

**Returns** ParameterEstimation.Config

**from\_yaml (yml)**

Read config object from yaml file :param yml: full path to text file containing configuration arguments in yaml format :type yml: str

**Returns** ParameterEstimation.Config

**model\_objects**

A list of model objects for mapping

**Type** Returns

**models\_affected\_experiments**

Get which experiment datasets affect which models

**Returns** dict. Keys are model names, value are list of affected models

**models\_affected\_validation\_experiments**

Get which experiment datasets affect which models

**Returns** dict. Keys are model names, value are list of affected models

**multi\_experiments**

List of experiment names that have more than one experiment separated by blank line Returns (tuple): (list, int)

**set\_default\_fit\_items\_dct ()**

Configure missing entries for items.fit\_items when they are in nested dict format

**Returns** None. Method operates inplace on class attributes

**set\_default\_fit\_items\_str ()**

Configure missing entries for items.fit\_items when they are strings pointing towards model variables

**Returns** None. Method operates inplace on class attributes

**to\_json()**

Output arguments as json

**Returns:** str All arguments in json format

**to\_yaml (filename=None)**

Output arguments as yaml

**Parameters** `filename (str, None)` – If not None (default),  
path to write yaml configuration to

**Returns** Config object as string in yaml format

**validation\_filenames**

a list of validation filenames

**Type** Returns

**validation\_names**

A list of validation names

**Type** Returns

**validations**

The validations property :returns: datasets.validations as dict

```
class Context(models, experiments, working_directory=None, con-  
text='s', parameters='mg', filename=None, valida-  
tion_experiments={}, settings={})
```

A high level interface to create a `ParameterEstimation.Config` object.

Enables the construction of a `ParameterEstimation.Config` object assuming one of several common patterns of usage.

## Examples

Assuming that we have two copasi models (`mod1` and `mod2`) and two experimental data files (`fname1`, `fname2`), correctly formatted according to the copasi specification. We can generate a config object that specifies the fitting of both experiments to both models and to fit all global and local parameters `parameters='gl'` in each.

```
with ParameterEstimation.Context(  
    [mod1, mod2], [fname1, fname2],  
    context='s', parameters='gl') as context:  
    context.set('method', 'genetic_algorithm_sr')  
    context.set('number_of_generations', 25)  
    context.set('population_size', 10)  
    config = context.get_config()  
  
pe = ParameterEstimation(config)
```

Or for profile likelihoods on the first model `mod1`

**get\_config\_cv()**  
configure for cross validation Returns:

**get\_config\_pl()**  
configure for profile likelihoods Returns:

**set** (*parameter, value*)  
Set the value of *parameter* to *value*.  
Looks for the first instance of *parameter* and sets its value to *value*.

## Parameters

- **parameter** – A key somewhere in the nested structure of the config object
- **value** – A value to replace the current value with

## Returns

None

**setd** (*dct*)  
Set the value of multiple settings using a *dict[setting] = value*.  
Iterates over *ParameterEstimation.Context.set()* with key value pairs

**Parameters** **dct** (*dict*) – a settings dict where keys are settings and values are setting values

## Returns

None

**do\_checks()**  
validate integrity of user input

**duplicate\_for\_every\_experiment** (*model, fit\_items, lower\_bounds, start\_values, upper\_bounds*)  
Replicate Copasi's duplicate for every experiment button.

## Parameters

- **model** –
- **fit\_items** –
- **lower\_bounds** –
- **start\_values** –
- **upper\_bounds** –

Returns:

**fit\_dir**  
Property holding the directory where the parameter estimation fitting occurs. This can be enumerated under a single problem directory to group similar parameter estimations :returns: str. A directory.

**get\_model\_objects\_from\_strings()**

Get model objects from the strings provided by the user in the Config class :return: list of *model.Model* objects

**Returns** list of model objects

**global\_quantities**

list of strings of global quantities present in the models

**Type** Returns

**local\_parameters**

list of strings of local parameters in the model

**Type** Returns

**metabolites**

list of strings of metabolites in the model

**Type** Returns

**models**

Get models

**Returns** the models entry of the *ParameterEstimation.Config* object

**models\_dir**

A directory containing models

Each model will be configured in a different directory when multiple models are being configured simultaneously :returns: dct. Location of models directories

**problem\_dir**

Property holding the directory where the parameter estimation problem is stored :returns: str. A directory.

**results\_directory**

A directory containing results, parameter estimation report files from copasi

Each model configured will have their own results directory :returns: dict[model] = results\_directory

**run(models)**

Run a parameter estimation using command line copasi.

**Parameters** **models** – dict of models. Output from \_setup()

**Returns** dict of models. Output from \_setup()

**Return type** param models

## pycotools3.tasks.ParameterEstimation.Context

```
class pycotools3.tasks.ParameterEstimation.Context(models, experiments, working_directory=None, context='s', parameters='mg', filename=None, validation_experiments={}, settings={})
```

A high level interface to create a ParameterEstimation.Config object.

Enables the construction of a ParameterEstimation.Config object assuming one of several common patterns of usage.

### Examples

Assuming that we have two copasi models (*mod1* and *mod2*) and two experimental data files (*fname1*, *fname2*), correctly formatted according to the copasi specification. We can generate a config object that specifies the fitting of both experiments to both models and to fit all global and local parameters *parameters='gl'* in each.

```
with ParameterEstimation.Context(
    [mod1, mod2], [fname1, fname2],
    context='s', parameters='gl') as context:
    context.set('method', 'genetic_algorithm_sr')
    context.set('number_of_generations', 25)
    context.set('population_size', 10)
    config = context.get_config()

pe = ParameterEstimation(config)
```

Or for profile likelihoods on the first model *mod1*

```
__init__(models, experiments, working_directory=None, context='s', parameters='mg', filename=None, validation_experiments={}, settings={})
```

Initialize self. See help(type(self)) for accurate signature.

### Methods

---

<code>__init__(models, experiments[, ...])</code>	Initialize self.
<code>get_config()</code>	
<code>set(parameter, value)</code>	Set the value of parameter to value.

---

## Attributes

---

<code>acceptable_context_args</code>
<code>acceptable_parameters_args</code>
<code>experiment_filetypes</code>

---

### `get_config_cv()`

configure for cross validation Returns:

### `get_config_pl()`

configure for profile likelihoods Returns:

### `set(parameter, value)`

Set the value of parameter to value.

Looks for the first instance of parameter and sets its value to value.

#### Parameters

- **parameter** – A key somewhere in the nested structure of the config object
- **value** – A value to replace the current value with

#### Returns

None

### `setd(dct)`

Set the value of multiple settings using a `dict[setting] = value`.

Iterates over `ParameterEstimation.Context.set()` with key value pairs

**Parameters** `dct` (`dict`) – a settings dict where keys are settings and values are setting values

#### Returns

None

## pycotools3.tasks.Sensitivities

### `class pycotools3.tasks.Sensitivities(model, **kwargs)`

Interface to COPASI sensitivity task

### `__init__(model, **kwargs)`

Initialize self. See `help(type(self))` for accurate signature.

## Methods

<code>__init__(model, **kwargs)</code>	Initialize self.
<code>add_list_of_variables_element()</code>	
<code>check_integrity(allowed, given)</code>	Method to raise an error when a wrong kwarg is passed to a subclass
<code>convert_bool_to_numeric(dct)</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>convert_bool_to_numeric2()</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>create_new_report()</code>	
<code>create_problem()</code>	
<code>create_sensitivity_task()</code>	
<code>get_component(model, component, string)</code>	Get component called string from model
<code>get_report_key()</code>	
<code>get_single_object_references()</code>	
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pyco-tools3 variable
<code>process_data()</code>	
<code>read_model(m)</code>	<b>param m</b>
<code>replace_sensitivities_task()</code>	
<code>run_task()</code>	
<code>sensitivity_task_key()</code>	Get the sensitivity task as it currently is in the model as etree.Element :return:
<code>set_cause()</code>	
<code>set_effect()</code>	
<code>set_method()</code>	
<code>set_report()</code>	
<code>set_secondary_cause()</code>	
<code>set_subtask()</code>	
<code>update_properties(kwargs)</code>	method for updating properties from kwargs

## Attributes

<code>cross_section_cause</code>
<code>cross_section_effect</code>
<code>evaluation_cause</code>
<code>evaluation_effect</code>
<code>optimization_cause</code>
<code>optimization_effect</code>
<code>parameter_estimation_cause</code>

Continued on next page

Table 19 – continued from previous page

---

parameter_estimation_effect
schema
sensitivity_number_map
steady_state_cause
steady_state_effect
subtasks
time_series_cause
time_series_effect
update_model

---

```

add_list_of_variables_element()
create_new_report()
create_problem()
create_sensitivity_task()
get_report_key()
get_single_object_references()
process_data()
replace_sensitivities_task()
run_task()
sensitivity_task_key()

```

Get the sensitivity task as it currently is in the model as etree.Element :return:

Args:

Returns:

```

set_cause()
set_effect()
set_method()
set_report()
set_secondary_cause()
set_subtask()

```

## pycotools3.tasks.Scan

```

class pycotools3.tasks.Scan(model, **kwargs)
    Interface to COPASI scan task

```

Scan Kwargs	Description
update_model	Default: False
subtask	Default: parameter_estimation
report_type	Default: profile_likelihood. Name of report from <i>Reports</i> class
output_in_subtask	Default: False
ad-just_initial_conditions	Default: False
number_of_steps	Default: 10
maximum	Default: 100
minimum	Default: 0.01
log10	Default: False
distribution_type	Default: normal
scan_type	Default: scan
scheduled	Default: True
save	Default: False
clear_scans	Default: True. If true, will remove all scans present then add new scan
run	Default: False
<report_kwargs>	Arguments for report_kwargs are also accepted here

Args:

Returns:

`__init__(model, **kwargs)`

#### Parameters

- **model** – Model
- **kwargs** –

#### Methods

---

`__init__(model, **kwargs)`

##### param model

check_integrity(allowed, given)	Method to raise an error when a wrong kwarg is passed to a subclass
convert_bool_to_numeric(dct)	CopasiML uses 1's and 0's for True or False in some but not all places.
convert_bool_to_numeric2()	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>create_scan()</code>	metabolite cn:
<code>define_report()</code>	Use Report class to create report :return:
<code>execute()</code>	

Continued on next page

Table 20 – continued from previous page

<code>get_report_key()</code>	
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pycotools3 variable
<code>read_model(m)</code>	
	<b>param m</b>
<code>remove_scans()</code>	Remove all scans that have been defined.
<code>set_scan_options()</code>	
<code>update_properties(kwags)</code>	method for updating properties from kwags

## Attributes

---

`schema`

---

**create\_scan ()****metabolite cn:** CN=Root,Model=New Model,Vector=Compartments[nuc],Vector=Metabolites[A]**Returns**

Args:

Returns:

**define\_report ()**

Use Report class to create report :return:

Args:

Returns:

**execute ()****get\_report\_key ()****remove\_scans ()**

Remove all scans that have been defined.

**Returns**

Args:

Returns:

**set\_scan\_options ()**

## pycotools3.tasks.Reports

**class** pycotools3.tasks.**Reports** (*model*, \*\**kwargs*)

Creates reports in copasi output specification section. Which report is controlled by the

report\_type key word. The following are valid types of report:

Report Types	Description
time_course	Report definition for collection of time course data.
parameter_estimation	Collect parameter estimates from parameter estimations run from the parameter estimation task
multi_parameter_estimation	Collect parameter estimation data from parameter estimations run from the scan task with copasi's repeat feature
profile_likelihood	Collect both the parameter being scanned value and the parameter estimates

Here are the keyword arguments accepted by the Reports class.

Args:

Returns:

**`__init__(model, **kwargs)`**

A class for configuring reports :param model: Model to add report configuration to :type model: model.Model :param \*\*kwargs: Arguments for report

## Methods

<code>__init__(model, **kwargs)</code>	A class for configuring reports :param model: Model to add report configuration to :type model: model.Model :param **kwargs: Arguments for report
<code>check_integrity(allowed, given)</code>	Method to raise an error when a wrong kwarg is passed to a subclass
<code>clear_all_reports()</code>	Having multile reports defined at once can be really annoying and give you unexpected results.
<code>convert_bool_to_numeric(dct)</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>convert_bool_to_numeric2()</code>	CopasiML uses 1's and 0's for True or False in some but not all places.
<code>get_component(model, component, string)</code>	Get component called string from model
<code>get_variable_from_string(m, v[, glob])</code>	Use model entity name to get the pycotools3 variable
<code>multi_parameter_estimation()</code>	Define a parameter estimation report and include the progression of the parameter estimation (function evaluations).
<code>parameter_estimation()</code>	Define a parameter estimation report and include the progression of the parameter estimation (function evaluations).

Continued on next page

Table 22 – continued from previous page

<code>profile_likelihood()</code>	Create report of a parameter and best value for a parameter estimation for profile likelihoods
<code>read_model(m)</code>	<b>param m</b>
<code>remove_report(report_name)</code>	remove report called report_name
<code>run()</code>	Execute code that builds the report defined by the kwargs
<code>scan()</code>	creates a report to collect scan time course results.
<code>sensitivity()</code>	
<code>timecourse()</code>	creates a report to collect time course results.
<code>update_properties(kwargs)</code>	method for updating properties from kwargs

## Attributes

---

`schema`

---

### `clear_all_reports()`

Having multile reports defined at once can be really annoying and give you unexpected results. Use this function to remove all reports before defining a new one to ensure you only have one active report any once. :return:

Args:

Returns:

### `multi_parameter_estimation()`

Define a parameter estimation report and include the progression of the parameter estimation (function evaluations). Defaults to including all metabolites, global variables and local variables with the RSS best value These can be over-ridden with the global\_quantities, LocalParameters and metabolites keywords.

Args:

Returns:

### `parameter_estimation()`

Define a parameter estimation report and include the progression of the parameter estimation (function evaluations). Defaults to including all metabolites, global variables and local variables with the RSS best value These can be over-ridden with the global\_quantities, LocalParameters and metabolites keywords.

Args:

Returns:

### **profile\_likelihood()**

Create report of a parameter and best value for a parameter estimation for profile likelihoods

Args:

Returns:

### **remove\_report(*report\_name*)**

remove report called *report\_name*

**Parameters** **report\_name** – return: pycotools3.model.Model

**Returns** pycotools3.model.Model

### **run()**

Execute code that builds the report defined by the kwargs

### **scan()**

creates a report to collect scan time course results.

By default all species and all global quantities are used with Time on the left most column. This behavior can be overwritten by passing lists of metabolites to the metabolites keyword or global quantities to the global quantities keyword

Args:

Returns:

### **sensitivity()**

### **timecourse()**

creates a report to collect time course results.

By default all species and all global quantities are used with Time on the left most column. This behavior can be overwritten by passing lists of metabolites to the metabolites keyword or global quantities to the global quantities keyword

Args:

Returns:

### 3.4.3 The viz module

The viz module exists to make visualising simulation output quick and easy for common patterns, such as plotting time courses or comparing parameter estimation output to experimental data. However it should be emphasised that the *matplotlib* and *seaborn* libraries are always close to hand in Python.

The viz module is currently in a state of rebuilding and so I only describe here the features which currently work.

---

<i>Parse</i>	General class for parsing copasi output into Python.
<i>PlotTimeCourse</i>	Plot time course data
<i>Boxplots</i>	Plot a boxplot for multi parameter estimation data.

---

## pycotools3.viz.Parse

```
class pycotools3.viz.Parse (cls_instance, log10=False, copasi_file=None, alpha=0.95, rss_value=None, num_data_points=None)
```

General class for parsing copasi output into Python.

First argument is an instance of a pycotools3 class.

instance	Description
<code>tasks.TimeCourse</code>	Parse time course data from TC.report_name into pandas.df
<code>tasks.ParameterEstimation</code>	Parse parameter estimation data from PE.report_name into pandas.df
<code>tasks.Scan</code>	Parse scan data from scan.report_name
<code>Parse</code>	enable parsing from a parse instance. Just returns itself
<code>str</code>	Parse data from folder of parameter estimation data into pandas.df. Requires the copasi file argument.

Args:

Returns:

```
__init__ (cls_instance, log10=False, copasi_file=None, alpha=0.95, rss_value=None, num_data_points=None)
```

### Parameters

- **cls\_instance** – A instance of pycotools3 class
- **log10** – bool. Whether to work on log10 scale
- **copasi\_file** – str. Optional but necessary when `cls_instance` is string. Must be the copasi\_file which produced the parameter estimation data as Parse extracts data headers from the copasi file
- **rss\_value** – float When `cls` is a profile likelihood with the `current_parameters` setting,  
    `rss_value` may not be empty. It is not automatically inferable from the COPASI model and must be specified separately.
- **num\_data\_points** – int When `cls` is a profile likelihood with `current_parameters` setting, the number of data points can-

not be automatically inferred for the calculation of likelihood ratio based confidence intervals. Therefore, this must be specified by the user.

## Methods

---

<code>__init__(cls_instance[, log10, co-pasi_file, ...])</code>	<b>param</b> <code>cls_instance</code>
<code>from_chaser_estimations(cls_instance[, folder])</code>	<b>return</b>
<code>from_folder()</code>	<b>param</b> <code>folder</code> <b>return:</b>
<code>from_multi_parameter_estimation()</code>	<code>Results instance</code> without headers - parse the results give them the proper headers then overwrite the file again
<code>from_profile_likelihood()</code>	Parse data from tasks. ProfileLikelihood :return: <code>pandas.DataFrame</code>
<code>from_timecourse()</code>	read time course data into pandas dataframe.
<code>parse()</code>	determine class type of self.cls_instance and call the appropriate method for parsing the data type :return:
<code>parse_scan()</code>	read scan data into pandas Dataframe.

---

## Attributes

---

<code>from_parameter_estimation</code>	Parse parameter estimation data.
--	----------------------------------

---

**from\_chaser\_estimations (cls\_instance, folder=None)**

**Returns**

**Parameters**

- `cls_instance` –
- `folder` – (Default value = None)

Returns:

**from\_folder ()**

**Parameters** `folder` – return:

Returns:

**static from\_multi\_parameter\_estimation(*cls\_instance*)**

Results come without headers - parse the results give them the proper headers then overwrite the file again

**Parameters**

- **cls\_instance** – instance of MultiParameterEstiamtion
- **folder** – alternative folder to parse from. Useful for tests  
(Default value = None)

Returns:

**from\_parameter\_estimation**

Parse parameter estimation data. Store the data in a cache. :return:

Args:

Returns:

**from\_profile\_likelihood()**

Parse data from tasks.ProfileLikelihood :return:

pandas.DataFrame

Args:

Returns:

**from\_timecourse()**

read time course data into pandas dataframe. Remove copasi generated square brackets around the variables :return: pandas.DataFrame

Args:

Returns:

**parse()**

determine class type of self.cls\_instance and call the appropriate method for parsing the data type :return:

Args:

Returns:

**parse\_scan()**

read scan data into pandas Dataframe. :return: pandas.DataFrame

Args:

Returns:

## pycotools3.viz.PlotTimeCourse

**class pycotools3.viz.PlotTimeCourse(*cls*, \*\*kwargs)**

Plot time course data

Time course kwargs:

kwarg	Description
x	<i>str</i> . Parameter to go on x axis. defaults to ‘Time’. If not ‘Time’ then plot is a phase space plot
y	<i>str or list of str</i> . Parameters for the y axis.
log10	<i>bool</i> plot on log10 scale
sepa- rate	<i>bool</i> separate time courses onto different axes. Default: True
<b>**kwargs</b>	See kwargs for more options

Args:

Returns:

`__init__(cls, **kwargs)`

#### Parameters

- **cls** – Instance of tasks.TimeCourse class
- **kwargs** –

#### Methods

---

`__init__(cls, **kwargs)`

**param cls**

---

`context([font_scale, rc])`

**param context** (Default  
value = ‘talk’)

---

`create_directory(results_directory)` create directory for results and switch to it

---

`parse(cls, log10[, copasi_file])` Mixin method interface to parse class :return:

---

`plot()`

**return**

---

`plot_kwargs()`

---

`save_figure(directory, filename[, dpi])`

**param directory**

---

`truncate(data, mode, theta)`

---

`update_properties(kwargs)` mixin method interface to truncate data  
method for updating properties from kwargs

---

`plot()`

#### Returns

## pycotools3.viz.Boxplots

---

**class** pycotools3.viz.Boxplots(*cls*, *\*\*kwargs*)

Plot a boxplot for multi parameter estimation data.

kwarg	Description
num_per_plot	Number of parameter per plot. Remainder fills up another plot.
<b>**kwargs</b>	see kwargs options

Args:

Returns:

---

**\_\_init\_\_**(*cls*, *\*\*kwargs*)

### Parameters

- **cls** – instance of tasks.MultiParameterEstimation or string .  
Same as PlotTimeCourseEnsemble
- **kwargs** –

## Methods

---

**\_\_init\_\_**(*cls*, *\*\*kwargs*)

**param** **cls**

---

**context**([*font\_scale*, *rc*])

**param** **context** (Default  
value = ‘talk’)

---

**create\_directory()**

**return**

---

**divide\_data()**

split data into multi plot :return:

**parse**(*cls*, *log10*[, *copasi\_file*])

Mixin method interface to parse class :return:

---

**plot()**

Plot multiple parameter estimation data as boxplot :return:

---

**plot\_kwargs()**

**save\_figure**(*directory*,     *filename*[,  
*dpi*])

**param** **directory**

---

**truncate**(*data*, *mode*, *theta*)

mixin method interface to truncate data

---

**update\_properties**(*kwargs*)

method for updating properties from kwargs

**create\_directory()**

**Returns**

**divide\_data()**

    split data into multi plot :return:

    Args:

    Returns:

**plot()**

    Plot multiple parameter estimation data as boxplot :return:

    Args:

    Returns:

# CHAPTER 4

---

## Support

---

Users can post a question on stack-overflow using the `pycotools` tag. I get email notifications for these questions and will respond.



# CHAPTER 5

---

## People

---

PyCoTools has been developed by Ciaran Welsh in Daryl Shanley's lab at Newcastle University.



# CHAPTER 6

---

## Caveats

---

- Non-ascii characters are minimally supported and can break PyCoTools
- Do not use unusual characters or naming systems (i.e. A reaction name called “A -> B” will break pycotools)
- In COPASI we can have (say) a global quantity and a metaboltie with the same name because they are different entities. This is not supported in Pycotools and you must use unique names for every model component

## 6.1 Citing PyCoTools

If you made use of PyCoTools, please cite [this article](#) using:

- Welsh, C.M., Fullard, N., Proctor, C.J., Martinez-Guimera, A., Isfort, R.J., Bascom, C.C., Tasseff, R., Przyborski, S.A. and Shanley, D.P., 2018. PyCoTools: a Python toolbox for COPASI. *Bioinformatics*, 34(21), pp.3702-3710.

And also please remember to cite [COPASI](#):

- Hoops, S., Sahle, S., Gauges, R., Lee, C., Pahle, J., Simus, N., Singhal, M., Xu, L., Mendes, P. and Kummer, U., 2006. COPASI—a complex pathway simulator. *Bioinformatics*, 22(24), pp.3067-3074.

and [tellurium](#):

- Medley, J.K., Choi, K., König, M., Smith, L., Gu, S., Hellerstein, J., Sealfon, S.C. and Sauro, H.M., 2018. Tellurium notebooks—An environment for reproducible dynamical modeling in systems biology. *PLoS computational biology*, 14(6), p.e1006220.



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