# Representation of Metabolic Networks in ScrumPy

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## End of Part 1

We have now covered enough fundamentals to think about how to use it for modelling.

What do we want to represent and act upon?

## What is ScrumPy?

A collection of modules (a **package**) providing the ability to define and analyse models.

Everything revolves around the use of model objects:

```
>>> m = ScrumPy.Model("FileName.spy")
```

Where "FileName.spy" is the name of file describing the model.

The ".spy" extension is conventional and convenient, but not mandatory.

and "m" is the model object. In these talks, "m" will always be used to denote the model.



## **Model Definition**

In ScrumPy, a model is defined by one or more text files, defining:

Comments Ignored by ScrumPy, but are useful to the human reader.

Directives Not part of the model *per se*, but specify how the model is to be read.

Reactions Define the metabolic network.

Initialisations Define parameter values and initial metabolite concentrations (only in kinetic models)



## **Model Definition**

```
# comment, everything from #
# to the end of the line is ignored
Structural()
# a Directive. Do not do any kinetic processing.
Rubisco:
                                    # a reaction name
    x CO2 + RuBP ch -> 2 PGA ch # stoichiometry
                                    # default kinetic
PGK:
    PGA_ch + ATP_ch <> BPGA_ch + ADP_ch
G3Pdh:
    BPGA ch + x NADPH ch + x Proton ch <>
                         x NADP_ch + GAP_ch + Pi_ch
```

## Model Definition - identifiers

Identifiers = Names

#### Either:

Any sequence of alphanumeric characters and \_ (underscore), not starting with a number e.g.

#### Valid:

Fructose6\_Phosphate AlphaAnaline

#### Invalid:

2,3-bisphosphoglycerate
TRANS-23-DEHYDROADIPYL-COA

#### Or:

Any quoted (") sequence of characters.

<sup>&</sup>quot;Saturated-Fatty-Acyl-CoA"

<sup>&</sup>quot;3-oxo-cis-vaccenoyl-ACPs"

Accessed as m.sm (internal) and m.smx (external):

	Ru5Pk	Aldo2	TPT_DHAP	Light_react	TKL
RuBP_ch	1/1	0/1	0/1	0/1	0/1
ATP_ch	-1/1	0/1	0/1	1/1	0/1
ADP_ch	1/1	0/1	0/1	-1/1	0/1
GAP_ch	0/1	0/1	0/1	0/1	-1/1
Pi_ch	0/1	0/1	1/1	-1/1	0/1
DHAP_ch	0/1	-1/1	-1/1	0/1	0/1
F6P_ch	0/1	0/1	0/1	0/1	-1/1
E4P_ch	0/1	-1/1	0/1	0/1	1/1
X5P_ch	0/1	0/1	0/1	0/1	1/1
SBP_ch	0/1	1/1	0/1	0/1	0/1
Ru5P_ch	-1/1	0/1	0/1	0/1	0/1

By default values in the stoichiometry matrices are *rational* numbers (ie fractions).

They can are represented as (e.g) 1/2 or mpq(1,2).

This can be changed with the EIType() directive (earlier slide).

For large (genome scale) models it is more common to use *real* numbers

```
( EIType(float) )
```

Stoichiometry matrices behave as a list of rows:

```
>>> print m.sm[0] [mpq(1,1), mpq(0,1), mpq(0,1), mpq(0,1), ...]
```

Or as a dictionary of rows:

```
>>> print m.sm["RuBP_ch"]
[mpq(1,1), mpq(0,1), mpq(0,1), mpq(0,1), mpq(0,1), ...]
```

Individual elements can be accessed as matrix[row,col]:

```
>>> print m.sm[0,0]
1/1
>>> print m.sm["RuBP_ch","Ru5Pk"]
1/1
```

The null-space is obtained the matrix.NullSpace() method:

```
>>> k = m.sm.NullSpace()
>>> print k
```

	c_0	c_1	c_2	c_3	c_4
Ru5Pk	0/1	0/1	0/1	0/1	-3/1
Aldo2	0/1	0/1	0/1	0/1	-1/1
TPT_DHAP	0/1	2/1	1/1	1/1	-1/1
Light_react	-1/1	-1/1	0/1	1/1	-9/1
TKL	0/1	0/1	0/1	0/1	-1/1
G3Pdh	0/1	0/1	0/1	1/1	-6/1
PGK	0/1	0/1	0/1	1/1	-6/1
TPI	0/1	1/1	1/1	1/1	-3/1
TKL2	0/1	0/1	0/1	0/1	-1/1

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### And now

We have covered enough to start the practical.