

# Escherichia coli Core Metabolism Model in LIM

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## Abstract

R package **LIM** (Soetaert and van Oevelen 2009a) is designed for reading and solving linear inverse models (LIM).

A package vignette deals with the structure of the LIM input files and how to solve the problems (Soetaert and van Oevelen 2009b).

To open it, type:

```
vignette("LIM")
```

Here it is exemplified on a (relatively small) problem from systems biology, the core metabolism of E. coli (Edwards, Covert, and Palsson 2002) as from the following website:

[http://gcrg.ucsd.edu/Downloads/Flux\\_Balance\\_Analysis](http://gcrg.ucsd.edu/Downloads/Flux_Balance_Analysis)

The original input file can be found in the package subdirectory `/examples/Reactions/E_coli.lim`

If you use this package, please cite as: (van Oevelen, van den Meersche, Meysman, Soetaert, Middelburg, and Vezina 2009).

*Keywords:* Linear inverse models, flux balance analysis, linear programming, E coli, R.

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## 1. the E. coli input file

The input file consists of several sections (see package vignette).

- The header of the file (ends at first line with ###) is ignored
- The metabolic reactions
- A function to maximise
- The bounds on the reactions (inequalities)
- A measurement equation
- The name of the components
- The names of the externals

Everything following a "!" is ignored.

E.coli input file

#####

## ## REACTIONS

!gene:	Reaction	enzyme
GLK1:	GLC + ATP -> G6P + ADP	Glucokinase
PGI1:	G6P <-> F6P	Phosphoglucose isomerase
PFKA:	F6P + ATP -> FDP + ADP	Phosphofructokinase
FBP:	FDP -> F6P + PI	Fructose-1,6-bisphosphatase
FBA:	FDP <-> T3P1 + T3P2	Fructose-1,6-bisphosphatase a
TPIA:	T3P2 <-> T3P1	Triosphosphate Isomerase
GAPA:	T3P1 + PI + NAD <-> NADH + 13PDG	Glyceraldehyde-3-phosphate de
PGK:	13PDG + ADP <-> 3PG + ATP	Phosphoglycerate kinase
GPMA:	3PG <-> 2PG	Phosphoglycerate mutase 1
ENO:	2PG <-> PEP	Enolase
PPSA:	PYR + ATP -> PEP + AMP + PI	Phosphoenolpyruvate synthase
PYKA:	PEP + ADP -> PYR + ATP	Pyruvate Kinase II
!	PYKF: PEP + ADP -> PYR + ATP	Pyruvate Kinase I
ACEE:	PYR + COA + NAD -> NADH + CO2 + ACCOA	Pyruvate dehydrogenase
!Pentose Phosphate Pathway		
ZWF:	G6P + NADP <-> D6PGL + NADPH	Glucose 6-phosphate-1-dehydro
PGL:	D6PGL -> D6PGC	6-Phosphogluconolactonase
GND:	D6PGC + NADP -> NADPH + CO2 + RL5P	6-Phosphogluconate dehydrogen
RPIA:	RL5P <-> R5P	Ribose-5-phosphate isomerase
RPE:	RL5P <-> X5P	Ribulose phosphate 3-epimerase
TKTA1:	R5P + X5P <-> T3P1 + S7P	Transketolase I
!	TKTB1: R5P + X5P <-> T3P1 + S7P	Transketolase II
TKTA2:	X5P + E4P <-> F6P + T3P1	Transketolase I
!	TKTB2: X5P + E4P <-> F6P + T3P1	Transketolase II
TALA:	T3P1 + S7P <-> E4P + F6P	Transaldolase A
!The Tricarboxylic Acid Cycle		
GLTA:	ACCOA + OA -> COA + CIT	Citrate synthase
ACNA:	CIT <-> ICIT	Aconitase A
ICDA:	ICIT + NADP <-> CO2 + NADPH + AKG	Isocitrate dehydrogenase
SUCA:	AKG + NAD + COA -> CO2 + NADH + SUCCOA	2-Ketoglutarate dehydrogenase
SUCC1:	SUCCOA + ADP + PI <-> ATP + COA + SUCC	Succinyl-CoA synthetase
SDHA1:	SUCC + FAD -> FADH + FUM	Succinate dehydrogenase
FRDA:	FUM + FADH -> SUCC + FAD	Fumurate reductase

FUMA:	FUM <-> MAL	!	Fumarase A
MDH:	MAL + NAD <-> NADH + OA	!	Malate dehydrogenase

## !Pyruvate Metabolism

DLD1:	PYR + NADH <-> NAD + LAC	!	D-Lactate dehydrogenase 1
ADHE2:	ACCOA + 2*NADH <-> ETH + 2*NAD + COA	!	Acetaldehyde dehydrogenase
PFLA:	PYR + COA -> ACCOA + FOR	!	Pyruvate formate lyase 1
PTA:	ACCOA + PI <-> ACTP + COA	!	Phosphotransacetylase
ACKA:	ACTP + ADP <-> ATP + AC	!	Acetate kinase A
ACS:	ATP + AC + COA -> AMP + PPI + ACCOA	!	Acetyl-CoA synthetase

## !Anaplerotic Reactions

PCKA:	OA + ATP -> PEP + CO2 + ADP	!	Phosphoenolpyruvate carboxykinase
PPC:	PEP + CO2 -> OA + PI	!	Phosphoenolpyruvate carboxylase
MAEB:	MAL + NADP -> CO2 + NADPH + PYR	!	Malic enzyme (NADP)
SFCA:	MAL + NAD -> CO2 + NADH + PYR	!	Malic enzyme (NAD)
ACEA:	ICIT -> GLX + SUCC	!	Isocitrate lyase
ACEB:	ACCOA + GLX -> COA + MAL	!	Malate synthase A
PPA:	PPI -> 2*PI	!	Inorganic pyrophosphatase
GLPK:	GL + ATP -> GL3P + ADP	!	Glycerol kinase
GPSA1:	GL3P + NADP <-> T3P2 + NADPH	!	Glycerol-3-phosphate-dehydrogenase-
RBSK:	RIB + ATP -> R5P + ADP	!	Ribokinase

## !Respiration

Note: the P/O ratio is set to 1.5 currently

NUOA:	NADH + Q -> NAD + QH2 + 3.5*HEXT	!	NADH dehydrogenase I
FDOH:	FOR + Q -> QH2 + CO2 + 2*HEXT	!	Formate dehydrogenase-0
GLPD:	GL3P + Q -> T3P2 + QH2	!	Glycerol-3-phosphate dehydrogenase
CYOA:	QH2 + 0.5*O2 -> Q + 2.5*HEXT	!	Cytochrome oxidase bo3
SDHA2:	FADH + Q <-> FAD + QH2	!	Succinate dehydrogenase complex
PNT1A:	NADPH + NAD -> NADP + NADH	!	Pyridine nucleotide transhydrogenase
PNT2A:	NADP + NADH + 2*HEXT -> NADPH + NAD	!	Pyridine nucleotide transhydrogenase
ATPA:	ATP <-> ADP + PI + 4*HEXT	!	FOF1-ATPase

## !Membrane Transport

GLCUP:	GLCxt + HEXT -> GLC	!	Glucose/galactose transporter
GLCPTS:	GLCxt + PEP -> G6P + PYR	!	Glucose
GLUP:	GLxt <-> GL	!	Glycerol
RIBUP:	RIBxt + ATP <-> RIB + ADP + PI	!	Ribose
ACUP:	ACxt + HEXT <-> AC	!	Acetate transport
LACUP:	LACxt + HEXT <-> LAC	!	L-Lactate
FORUP:	FORxt <-> FOR	!	Formate transport
ETHUP:	ETHxt + HEXT <-> ETH	!	Ethanol transport
SUCCUP:	SUCCxt + HEXT <-> SUCC	!	Succinate transport

```

PYRUP: PYRxt + HEXt <-> PYR      !      Pyruvate transport
PIUP:  PIxt <-> PI                !      Phosphate transport
O2TX:  O2xt <-> O2                !      Oxygen transport
CO2TX: CO2xt <-> CO2              !      Carbon dioxide transport

ATPM:  ATP -> ADP + PI            !      ATP drain flux for constant m
ADK:   ATP + AMP-> 2*ADP          !      ADK

Growth:                                &
41.25*ATP +3.54*NAD +18.22*NADPH +0.2*G6P      &
+0.07*F6P +0.89*R5P +0.36*E4P +0.12*T3P1      &
+1.49*3PG +0.51*PEP +2.83*PYR +3.74*ACCOA +1.78*OA +1.07*AKG  &
-> 3.74*COA +41.25* ADP +41.25* PI            &
+3.54* NADH +18.22* NADP +1.00* Biomass
### END REACTION

## MAXIMISE
maxgrowth: Growth
## END MAXIMISE

### INEQUALITIES
! Carbon sources...
O2TX = [0,20]      ! Oxygen input
GLCUP = [0,10]     ! glucose input
GLUP  = [-1000,0]  ! glycerol
RIBUP = [-1000,0]  ! Ribose uptake   - strange!
SUCCUP= [-1000,0]  ! succinate
ACUP  = [-1000,0]  ! acetate
LACUP = [-1000,0]  ! lactate
PYRUP = [-1000,0]  ! pyruvate
! Carbon byproducts
FORup = [-1000,0]  ! formate
ETHup = [-1000,0]  ! ethanol
CO2TX = [-1000,0]  ! CO2
! phosphate
PIUP = [-1000,1000]

SDHA1 <100
FRDA  <100
FORup+ LACUP=[-10,-10]
### END INEQUALITIES

### EQUATIONS

```

ATPM = 5.87 ! Non-growth associated ATP drain flux for constant maintenance requirements  
### END EQUATIONS

## ### COMPONENTS

GLC ! a-D-Glucose  
G6P ! Glucose 6-phosphate  
F6P ! Fructose 6-phosphate  
FDP ! Fructose 1,6-diphosphate  
T3P2 ! /DHAP Dihydroxyacetone phosphate  
T3P1 ! Glyceraldehyde 3-phosphate  
13PDG ! 1,3-bis-Phosphoglycerate  
3PG ! 3-Phosphoglycerate  
2PG ! 2-Phosphoglycerate  
PEP ! Phosphoenolpyruvate  
PYR ! Pyruvate  
ACCOA ! Acetyl-CoA  
CIT ! Citrate  
! ACO ! cis-Aconitate  
ICIT ! Isocitrate  
AKG ! a-Ketoglutarate  
SUCCOA ! Succinate CoA  
SUCC ! Succinate  
FUM ! Fumarate  
MAL ! Malate  
OA ! Oxaloacetate  
! ACAL ! Acetaldehyde  
ACTP ! Acetyl-phosphate  
ETH ! Ethanol  
AC ! Acetate  
LAC ! D-Lactate  
FOR ! Formate  
D6PGL ! D-6-Phosphate-glucono-delta-lactone  
D6PGC ! D-6-Phosphate-gluconate  
RL5P ! Ribulose 5-phosphate  
X5P ! Xylulose-5-phosphate  
R5P ! Ribose 5-phosphate  
S7P ! sedo-Heptulose  
E4P ! Erythrose 4-phosphate  
RIB ! Ribose  
GLX ! Glyoxylate  
NAD ! Nicotinamide adenine dinucleotide  
NADH ! Nicotinamide adenine dinucleotide (reduced)  
NADP ! Dihyronicotinamide adenine dinucleotide phosphate

NADPH ! Dihydronicotinamide adenine dinucleotide phosphate (reduced)  
HEXT ! External Hydrogen Ion (Proton)  
Q ! Ubiquinone

FAD ! Flavin adenine dinucleotide  
FADH ! Flavin adenine dinucleotide (reduced)  
AMP ! Adenosine monophosphate  
ADP ! Adenosine diphosphate  
ATP ! Adenosine triphosphate  
GL3P ! Glycerol 3-phosphate  
CO2 ! Carbon dioxide  
PI ! Inorganic Phosphate  
PPI ! Pyrophosphate

O2 ! Oxygen  
COA  
GL  
QH2 !  
### END COMPONENTS

### EXTERNALS  
Biomass  
GLCxt  
GLxt  
RIBxt  
ACxt  
LACxt  
FORxt  
ETHxt  
SUCCxt  
PYRxt  
PIxt  
O2xt  
CO2xt  
### END EXTERNALS

## 2. Reading the E.coli input file

Assuming that the input file is called "E\_coli.lim" and the working directory has been set, it can be read as follows:

```
require(LIM)
LIMEcoli <- Setup("E_coli.lim")
```

This creates a list of type `lim`, that contains all information necessary to solve the problem

## 3. The parsimonious and optimized solution, ranges

Once the input file has been read, we can generate the "simplest" solution, i.e. the one where  $\sum(x^2)$  is minimal, where  $x$  are the unknown reaction rates. This is called the "parsimonious solution". It is common to calculate this in foodweb ecology (where it is unclear which characteristics of a foodweb is optimized); it may be less relevant from a system's biology perspective.

Function `Ldei` estimates the parsimonious solution

```
> pars <- Ldei(LIMEcoli)
```

It makes more sense to optimize the growth. That growth has to be maximised was inputted in the file (by the `## maximize` statement).

The optimal value is found by linear programming, using function `Linp`:

```
> LP <- Linp(LIMEcoli)
```

It is also simple to estimate the ranges of all unknown reaction rates:

```
> xr <- Xranges(LIMEcoli)
```

Now for every reaction rate, the "simplest", "optimal", "minimal" and "maximal" value has been estimated:

```
> data.frame(simplest = pars$X, optimal = LP$X, xr)
```

	simplest	optimal	min	max
GLK1	-2.026294e-03	0.000000	0.0000000	10.000000
PGI1	4.184964e+00	807.532745	-15.8333333	807.532745
PFKA	4.188066e+00	781.590686	0.8333333	2229.130000
FBP	2.420527e-03	0.000000	0.0000000	1604.130000
FBA	4.185645e+00	781.590686	0.8333333	781.590686
TPIA	4.185708e+00	781.590686	0.8333333	781.590686
GAPA	8.371775e+00	1541.434199	5.0000000	1541.434199

PGK	8.371775e+00	1541.434199	5.0000000	1541.434199
GPMA	8.372227e+00	1492.089090	5.0000000	1492.089090
ENO	8.372227e+00	1492.089090	5.0000000	1492.089090
PPSA	-1.506859e-03	0.000000	0.0000000	1604.130000
PYKA	2.518254e+00	466.657964	0.0000000	2136.630000
ACEE	-1.395991e-04	1149.295284	0.0000000	1158.949190
ZWF	-1.381780e-03	0.000000	0.0000000	75.000000
PGL	-1.381780e-03	0.000000	0.0000000	75.000000
GND	-1.381780e-03	0.000000	0.0000000	75.000000
RPIA	-2.041909e-03	23.623833	0.0000000	28.202015
RPE	6.601289e-04	-23.623833	-23.6238328	50.000000
TKTA1	2.754418e-04	-5.850762	-5.8507623	25.000000
TKTA2	3.846871e-04	-17.773070	-17.7730705	25.000000
TALA	2.754418e-04	-5.850762	-5.8507623	25.000000
GLTA	1.593211e+00	35.435749	0.0000000	40.847149
ACNA	1.593211e+00	35.435749	0.0000000	40.847149
ICDA	-4.829526e-04	35.435749	0.0000000	40.847149
SUCA	-1.582513e-04	0.000000	0.0000000	30.000000
SUCC1	-1.582513e-04	0.000000	0.0000000	30.000000
SDHA1	1.592902e+00	0.000000	0.0000000	100.000000
FRDA	-1.265199e-03	100.000000	0.0000000	100.000000
FUMA	1.594167e+00	-100.000000	-100.0000000	8.333333
MDH	-7.440225e-02	-100.000000	-1168.3150000	16.666667
DLD1	3.448268e+00	0.000000	0.0000000	10.000000
ADHE2	3.331147e+00	1000.000000	0.0000000	1000.000000
PFLA	6.518954e+00	10.000000	0.0000000	150.000000
PTA	1.772074e+00	0.000000	0.0000000	1660.380000
ACKA	1.772074e+00	0.000000	0.0000000	1660.380000
ACS	1.770176e+00	0.000000	0.0000000	1604.130000
PCKA	6.329015e-01	0.000000	0.0000000	1604.130000
PPC	2.299974e+00	194.384939	0.0000000	1704.130000
MAEB	6.548934e-01	0.000000	0.0000000	1068.315000
SFCA	2.607370e+00	0.000000	0.0000000	1068.315000
ACEA	1.593694e+00	0.000000	0.0000000	30.000000
ACEB	1.593694e+00	0.000000	0.0000000	30.000000
PPA	1.770176e+00	0.000000	0.0000000	1604.130000
GLPK	6.306863e-05	0.000000	0.0000000	0.000000
GPSA1	-1.450648e+00	0.000000	-140.0000000	0.000000
RBSK	2.047272e-03	0.000000	0.0000000	0.000000
NUOA	-6.649777e-04	140.000000	0.0000000	140.000000
FDOH	2.647923e-03	0.000000	0.0000000	140.000000
GLPD	1.450711e+00	0.000000	0.0000000	140.000000
CYOA	3.046861e+00	40.000000	0.0000000	40.000000



SDHA2	1.594167e+00	-100.000000	-100.000000	8.333333
PNT1A	3.296669e+00	0.000000	0.000000	3208.260000
PNT2A	4.090142e+00	567.965512	0.000000	3208.260000
ATPA	-1.555974e+00	-145.466329	-460.000000	1144.130000
GLCUP	-2.026294e-03	0.000000	0.000000	10.000000
GLCPTS	4.185548e+00	814.156250	0.000000	814.156250
GLUP	6.306863e-05	0.000000	0.000000	0.000000
RIBUP	2.047272e-03	0.000000	0.000000	0.000000
ACUP	-1.898141e-03	0.000000	-75.000000	0.000000
LACUP	-3.448268e+00	0.000000	-10.000000	0.000000
FORUP	-6.516306e+00	-10.000000	-10.000000	0.000000
ETHUP	-3.331147e+00	-1000.000000	-1000.000000	0.000000
SUCCUP	6.318284e-04	-100.000000	-130.000000	0.000000
PYRUP	-1.347754e-03	-27.796342	-150.000000	0.000000
PIUP	-1.104592e-03	120.547782	0.000000	120.547782
O2TX	1.523431e+00	20.000000	0.000000	20.000000
CO2TX	-1.595676e+00	-990.346093	-1000.000000	0.000000
ATPM	5.870000e+00	5.870000	5.870000	5.870000
ADK	1.768669e+00	0.000000	0.000000	1604.130000
Growth	-3.034592e-04	33.117523	0.000000	33.117523

The range solutions can be plotted; as there are many reactions, we plot them in two figures.  
The "optimal" solution is added as a black dot.

```
> par(mfrow = c(1, 2))
> nr <- LIMEcoli$NUnknowns
> ii <- 1:(nr/2)
> dotchart(LP$X[ii], xlim = range(xr), pch = 16, cex = 0.8)
> segments(xr[ii,1], 1:nr, xr[ii,2], 1:nr)
> ii <- (nr/2+1):nr
> dotchart(LP$X[ii], xlim = range(xr), pch = 16, cex = 0.8)
> segments(xr[ii,1], 1:nr, xr[ii,2], 1:nr)
> mtext(side = 3, cex = 1.5, outer = TRUE, line = -1.5,
+       "E coli Core Metabolism, optimal solution and ranges")
```

## E coli Core Metabolism, optimal solution and ranges

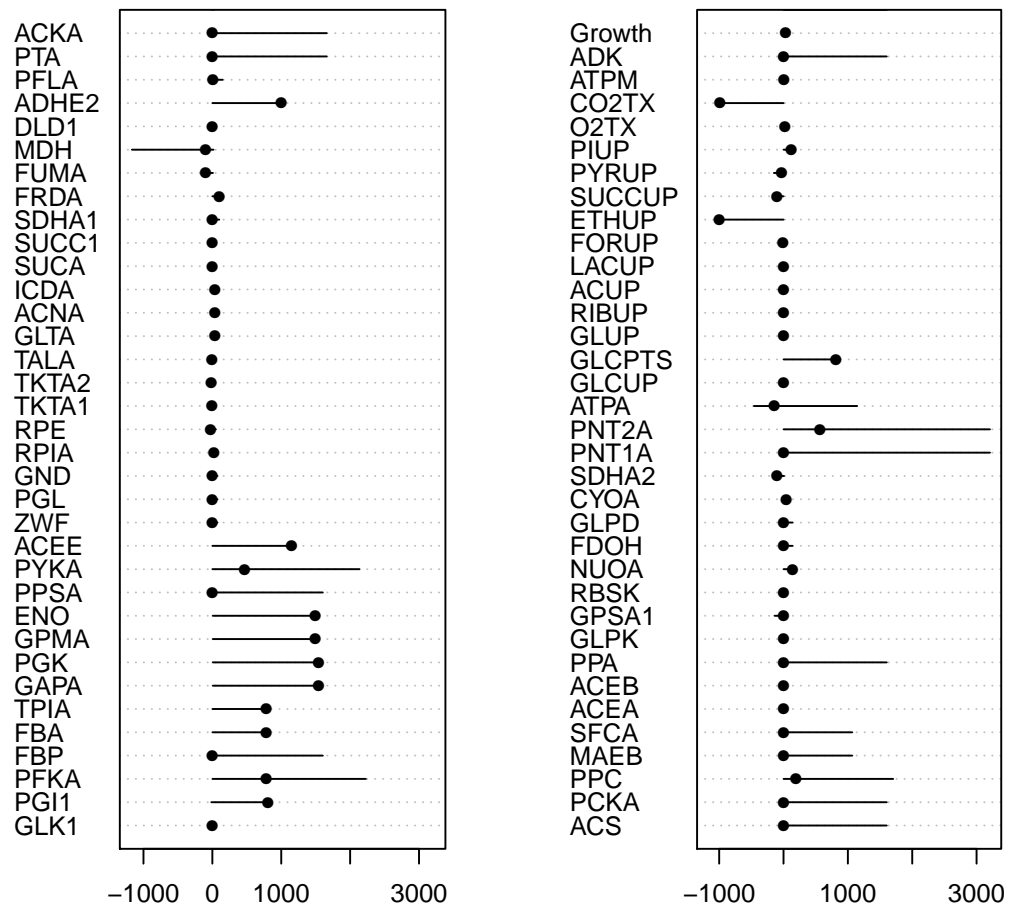


Figure 1: Ranges, and optimal solution of the E.coli central core metabolism - see text for R-code

## 4. Generating multiple plausible solutions

The E.coli model is underdetermined, such that an infinite amount of solutions are likely, given the data. By optimising growth, we selected one "optimal" solution; by estimating the ranges, we calculated the minimal and maximal values of each reaction.

It is also possible to sample the solution space in a random way. Function `xsample` does that; each point it generates is equally valid and equally likely.

We take 500 random samples; it takes a while to do this; `print(system.time())` estimates the time, in seconds.

```
> print(system.time(  
+   xs <- Xsample(LIMEcoli, iter = 500, type = "mirror", test = TRUE) #))  
+ ))
```

```
      user  system elapsed  
5.592    0.020    5.611
```

```
>
```

With 70 variables, it is not possible to plot all pairwise relationships.

Here we plot them for 12 of them.

```
> pairs(xs[, 1:12], pch = ".", cex = 2, gap = 0, upper.panel = NULL)
```

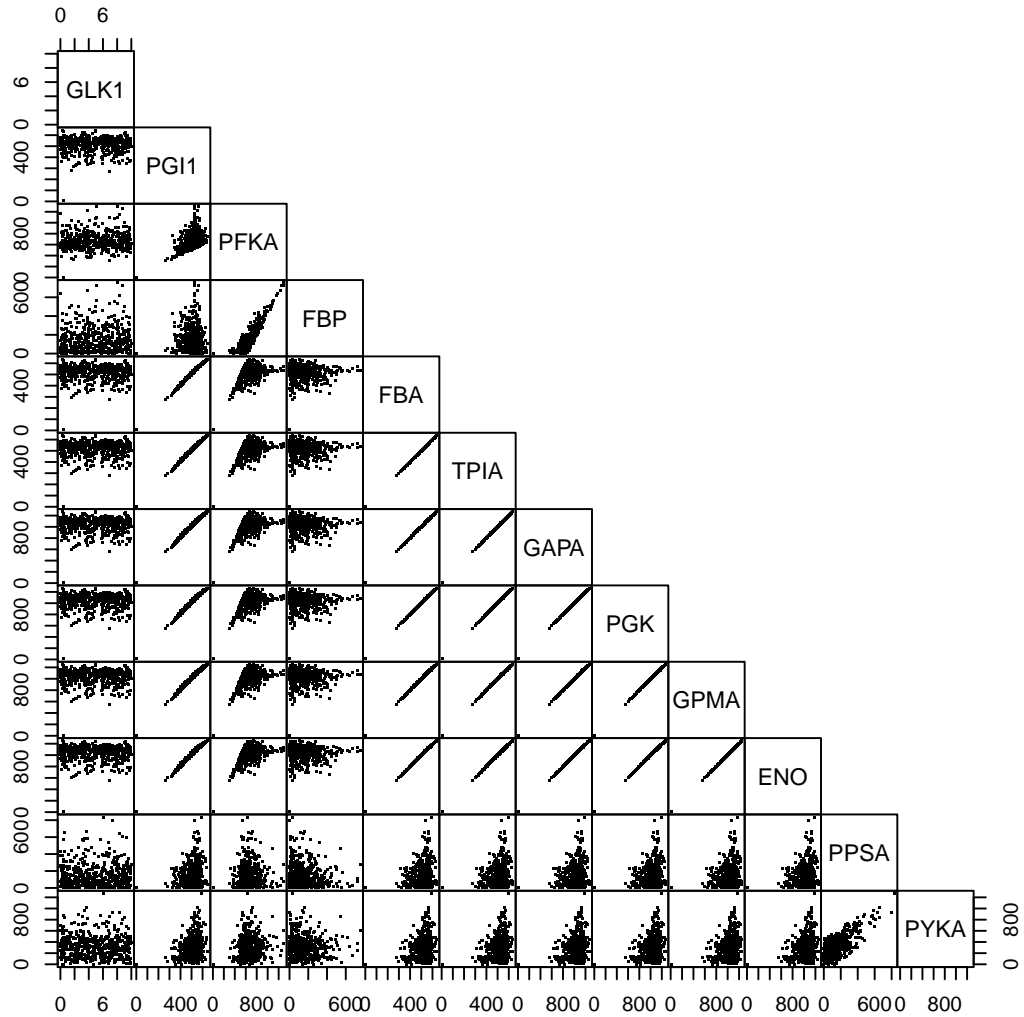


Figure 2: A random sample of plausible solutions of the E.coli central core metabolism - plotted as a pairwise plot for the first 12 reaction rates see text for R-code

## References

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