Sparse Flux Balance Analysis

Author: Ronan Fleming, Hoai Minh Le, Systems Biochemistry Group, University of Luxembourg.

Reviewer: Stefania Magnusdottir, Molecular Systems Physiology Group, University of Luxembourg.

INTRODUCTION

We consider a biochemical network of m molecular species and n biochemical reactions. The $S \in \mathcal{Z}^{m \times n}$

biochemical network is mathematically represented by a stoichiometric matrix $S \in \mathbb{Z}^{m \times n}$. In standard notation, flux balance analysis (FBA) is the linear optimisation problem

$$\min_{v} \rho(v) \equiv c^{T} v$$
s.t. $Sv = b$, $l \le v \le u$.

where $c \in \Re^n$ is a parameter vector that linearly combines one or more reaction fluxes to form what is termed the objective function, and where a $b_i < 0$, or $b_i > 0$, represents some fixed output, or input, of the ith molecular species. A typical application of flux balance analysis is to predict an optimal non-equilibrium steady-state flux vector that optimises a linear objective function, such biomass production rate, subject to bounds on certain reaction rates. Herein we use sparse flux balance analysis to predict a minimal number of active reactions 1 , consistent with an optimal objective derived from the result of a standard flux balance analysis problem. In this context *sparse flux balance analysis* requires a solution to the following problem

$$\min_{v} \quad ||v||_{0}$$
s.t.
$$Sv = b$$

$$l \le v \le u$$

$$c^{T}v = \rho^{*}$$

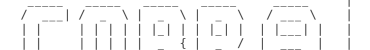
where the last constraint is represents the requirement to satisfy an optimal objective value ρ^* derived from any solution to a flux balance analysis (FBA) problem.

EQUIPMENT SETUP

Initialize the COBRA Toolbox.

If necessary, initialize The Cobra Toolbox using the initCobraToolbox function.

initCobraToolbox



```
Documentation:
                                                  http://opencobra.github.io/cobratoolbox
> Checking if git is installed ... Done.
> Checking if the repository is tracked using git ... Done.
> Checking if curl is installed ... Done.
> Checking if remote can be reached ... Done.
> Initializing and updating submodules ... Done.
> Adding all the files of The COBRA Toolbox ... Done.
> Define CB map output... set to svg.
> Retrieving models ...
                        Done.
> TranslateSBML is installed and working properly.
> Configuring solver environment variables ...
  - [---*] ILOG_CPLEX_PATH: C:\Program Files\IBM\ILOG\CPLEX Studio1271\cplex\matlab\x64 win64
  - [----] GUROBI PATH : --> set this path manually after installing the solver ( see instructions )
  - [---*] TOMLAB PATH: C:\Program Files\tomlab\
  - [----] MOSEK PATH : --> set this path manually after installing the solver ( see instructions )
  Done.
> Checking available solvers and solver interfaces ... Done.
> Setting default solvers ... Done.
> Saving the MATLAB path ... Done.
  - The MATLAB path was saved in the default location.
```

> Summary of available solvers and solver interfaces

Support	LP	MILP		QP	MIQP	N	LP		
cplex_direct	full		0		0	0	0	-	
dqqMinos	full		0		-	-	-	-	
glpk	full		1		1	-	-	-	
gurobi	full		1		1	1	1	-	
ibm_cplex	full		1		1	1	-	-	
matlab	full		1		-	-	-	1	
mosek	full		0		0	0	-	-	
pdco	full		1		-	1	-	-	
quadMinos	full		0		-	-	-	0	
tomlab_cplex	full		1		1	1	1	-	
qpng	experimental		-		-	1	-	-	
tomlab_snopt	experimental		-		-	-	-	1	
gurobi_mex	legacy		0		0	0	0	-	
lindo_old	legacy		0		-	-	-	-	
lindo_legacy	legacy		0		-	-	-	-	
lp_solve	legacy		1		-	-	-	-	
opti	legacy		0		0	0	0	0	
Total	-		7		4	5	2	2	

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

COBRA model.

In this tutorial, the model used is the generic reconstruction of human metabolism, the Recon 2.04², which is provided in the COBRA Toolbox. The Recon 2.04 model can also be downloaded from the

Virtual Metabolic Human webpage. You can also select your own model to work with. Before proceeding with the simulations, the path for the model needs to be set up:

```
if 0
% Using own model, change "if 0" to "if 1" and change the filename and directory
    filename = 'Recon3.0model';
    directory = '~/work/sbgCloud/programReconstruction/projects/recon2models/data/reconXCompan
    model = loadIdentifiedModel(filename, directory);
    % model = convertOldStyleModel(model);%convert to new COBRA format style if needed.
else
    % Default use of Recon 2.04
    global CBTDIR
    load([CBTDIR filesep 'test' filesep 'models' filesep 'Recon2.v04.mat']);
    model = modelR204;
    clear modelR204;
```

Recon 2.04 is written in the "old style" COBRA format, and we thus use the function convertOldStyleModel to convert it to the new COBRA Toolbox format.

```
model = convertOldStyleModel(model);
end
```

NOTE: The following text, code, and results are shown for the Recon 2.04 model

PROCEDURE

Set the tolerance to distinguish between zero and non-zero flux, based on the numerical tolerance of the currently installed optimisation solver.

```
feasTol = getCobraSolverParams('LP', 'feasTol');
```

Display the constraints

```
minInf = -1000;
maxInf = 1000;
printConstraints(model, minInf, maxInf);
```

```
MinConstraints:
DM_T_antigen_g_
                - 1
EX 10fthf(e) -1
EX 10fthf5glu(e) -1
EX_10fthf6glu(e) -1
EX_10fthf7glu(e) -1
EX_11_cis_retfa(e) -1
EX_13_cis_retnglc(e) -1
EX 1glyc hs(e) -1
EX 1mncam(e) -1
EX 2425dhvitd2(e) -1
EX 2425dhvitd3(e) -1
EX 24nph(e) -1
EX 25hvitd2(e) -1
EX 25hvitd3(e) -1
EX 2hb(e) -1
EX 2mcit(e) -1
EX_34dhoxpeg(e) -1
EX 34dhphe(e) -1
EX_35cgmp(e) -1
EX 3aib(e) -1
```

```
EX_3aib_D(e) -1
EX 3mlda(e) -1
EX 4abut(e) -1
EX_4hdebrisoquine(e) -1
EX 4hphac(e) -1
EX 4mptnl(e) -1
EX 4mtolbutamide(e) -1
EX 4nph(e) -1
EX 4nphsf(e) -1
EX_4pyrdx(e) -1
EX 5adtststerone(e) -1
EX 5adtststeroneglc(e) -1
EX 5adtststerones(e) -1
EX 5dhf(e) -1
EX 5fthf(e) -1
EX 5homeprazole(e) -1
EX 5htrp(e) -1
EX 5thf(e) -1
EX_6dhf(e) -1
EX_6htststerone(e) -1
EX_6thf(e) -1
EX_7dhf(e) -1
EX_7thf(e) -1
EX_9_cis_retfa(e) -1
EX abt(e) -1
EX_ac(e) -1
EX acac(e) -1
EX acald(e) -1
EX acetone(e) -1
EX acgalfucgalacgalfuc12gal14acglcgalgluside hs(e) -1
EX_acgalfucgalacgalfucgalacglcgal14acglcgalgluside_hs(e) -1
EX acgam(e) -1
EX ach(e) -1
EX_acn13acngalgbside_hs(e) -1
EX acn23acngalgbside hs(e) -1
EX_acnacngal14acglcgalgluside_hs(e) -1
EX_acnacngalgbside_hs(e) -1
EX acngalacglcgal14acglcgalgluside hs(e) -1
EX ade(e) -1
EX adp -1
EX adprbp(e) -1
EX_adprib(e) -1
EX_adrn(e) -1
EX_adrnl(e) -1
EX aflatoxin(e) -1
EX ahandrostanglc(e) -1
EX_ak2lgchol_hs(e) -1
EX_akg(e) -1
EX_ala_B(e) -1
EX ala D(e) -1
EX ala L(e) -1
EX aldstrn(e) -1
EX amp(e) -1
EX andrstrn(e) -1
EX andrstrnglc(e) -1
EX_antipyrene(e) -1
EX apnnox(e) -1
EX appnn(e) -1
EX_aprgstrn(e) -1
EX aqcobal(e) -1
EX_arab_L(e) -1
EX arachd(e) -1
EX_arg_L(e) -1
EX_ascb_L(e) -100
EX_asn_L(e) -1
EX_asp_D(e) -1
EX asp L(e) -1
EX atp(e) -1
```

```
EX_avite1(e) -1
EX avite2(e) -1
EX bhb(e) -1
EX_bildglcur(e) -1
EX bilglcur(e) -1
EX bilirub(e) -1
EX biocyt(e) -1
EX btn(e) -100
EX but(e) -1
EX bvite(e) -1
EX_bz(e) -1
EX ca2(e) -1
EX_camp(e) -1
EX caro(e) -1
EX carveol(e) -1
EX cca d3(e) -1
EX cgly(e) -1
EX chsterol(e) -1
EX_chtn(e) -1
EX_cit(e) -1
EX_CLPND(e) -1
EX_{cmp}(e) -1
EX_co(e) -1
EX_co2(e) -100
EX_coumarin(e) -1
EX_creat(e) -1
EX crmp hs(e) -1
EX crtsl(e) -1
EX_crtstrn(e) -1
EX crvnc(e) -1
EX csn(e) -1
EX cspg a(e) -1
EX_cspg_b(e) -1
EX_cspg_c(e) -1
EX cspg d(e) -1
EX_cspg_e(e) -1
EX cyan(e) -1
EX_dcsptn1(e) -1
EX debrisoquine(e) -1
EX dqchol(e) -1
EX dheas(e) -1
EX_dhf(e) -1
EX_digalsgalside_hs(e) -1
EX_dlnlcg(e) -1
EX dmantipyrine(e) -1
EX dmhptcrn(e) -1
EX dopa(e) -1
EX dopasf(e) -1
EX_drib(e) -1
EX duri(e) -1
EX_eaflatoxin(e) -1
EX ebastine(e) -1
EX ebastineoh(e) -1
EX_eicostet(e) -1
EX_elaid(e) -1
EX_estradiol(e) -1
EX estradiolglc(e) -1
EX_estriolglc(e) -1
EX_estroneglc(e) -1
EX estrones(e) -1
EX etoh(e) -1
EX_fe2(e) -1
EX_fe3(e) -1
EX_for(e) -1
EX fru(e) -1
EX_fuc13galacglcgal14acglcgalgluside_hs(e) -1
EX fuc14galacglcgalgluside hs(e) -1
```

EX fucacgalfucgalacglcgalgluside hs(e) -1

```
EX_fucacngal14acglcgalgluside_hs(e) -1
EX fucacngalacglcgalgluside hs(e) -1
EX fucfuc12gal14acglcgalgluside hs(e) -1
EX_fucfuc132galacglcgal14acglcgalgluside_hs(e) -1
EX fucfucfucgalacglc13galacglcgal14acglcgalgluside hs(e) -1
EX fucfucfucgalacglcgal14acglcgalgluside hs(e) -1
EX fucfucgalacglcgalgluside hs(e) -1
EX fucgal14acglcgalgluside hs(e) -1
EX fucgalfucgalacglcgalgluside_hs(e) -1
EX fucgalgbside hs(e) -1
EX fuc L(e) -1
EX galacglcgalgbside hs(e) -1
EX galfuc12gal14acglcgalgluside hs(e) -1
EX_galfucgalacglcgal14acglcgalgluside_hs(e) -1
EX_galgalfucfucgalacglcgalacglcgal14acglcgalgluside_hs(e) -1
EX galgalgalthcrm_hs(e) -1
EX qbside hs(e) -1
EX gchola(e) -1
EX_gd1b2_hs(e) -1
EX gd1c hs(e) -1
EX_gdp(e) -1
EX_glc(e) -1
EX_gln_L(e) -1
EX_gluala(e) -1
EX_glu_L(e) -1
EX_glyb(e) -1
EX glyc S(e) -1
EX glygn2(e) -1
EX glygn4(e) -1
EX glygn5(e) -1
EX gmp(e) -1
EX gplc hs(e) -1
EX gp1calpha hs(e) -1
EX_gq1b_hs(e) -1
EX gqlbalpha hs(e) -1
EX_gt1a_hs(e) -1
EX gthox(e) -1
EX gthrd(e) -1
EX gtp(e) -1
EX gua(e) -1
EX h(e) -100
EX h2o(e) -100
EX h2o2(e) -1
EX_ha(e) -1
EX ha prel(e) -1
EX hco3(e) -100
EX hcoumarin(e) -1
EX hdca(e) -1
EX hestratriol(e) -1
EX hexc(e) -1
EX hista(e) -1
EX hom L(e) -1
EX hpdca(e) -1
EX hspg(e) -1
EX htaxol(e) -1
EX_hxan(e) -1
EX i(e) -1
EX idp(e) -1
EX_ile_L(e) -1
EX imp(e) -1
EX inost(e) -1
EX k(e) -1
EX ksi(e) -1
EX_ksi_deg1(e) -1
EX_ksii_core2(e) -1
EX_ksii_core4(e) -1
EX lac D(e) -1
EX lac L(e) -1
```

```
EX_lcts(e) -1
EX Lcystin(e) -1
EX leuktrA4(e) -1
EX_leuktrB4(e) -1
EX leuktrC4(e) -1
EX leuktrD4(e) -1
EX leuktrE4(e) -1
EX leuktrF4(e) -1
EX leu L(e) -1
EX lgnc(e) -1
EX_limnen(e) -1
EX lipoate(e) -1
EX_lneldc(e) -1
EX lnlc(e) -1
EX_lnlnca(e) -1
EX lnlncg(e) -1
EX lys L(e) -1
EX malttr(e) -1
EX_{meoh(e)} -1
EX mepi(e) -1
EX_mercplaccys(e) -1
EX met L(e) -1
EX_mthgxl(e) -1
EX_n2m2nmasn(e) -1
EX nac(e) -1
EX_nad(e) -1
EX nadp(e) -1
EX ncam(e) -1
EX nh4(e) -100
EX nifedipine(e) -1
EX no(e) -1
EX npthl(e) -1
EX nrpphr(e) -1
EX_nrpphrsf(e) -1
EX nrvnc(e) -1
EX o2s(e) -1
EX oagd3 hs(e) -1
EX oagt3 hs(e) -1
EX ocdcea(e) -1
EX omeprazole(e) -1
EX onpthl(e) -1
EX_orn(e) -1
EX_oxa(e) -1
EX_paf_hs(e) -1
EX pchol hs(e) -1
EX peplys(e) -1
EX perillyl(e) -1
EX pglyc hs(e) -1
EX_pheacgln(e) -1
EX_phyQ(e) -1
EX_phyt(e) -1
EX pi(e) -100
EX_pnto_R(e) -100
EX_ppa(e) -1
EX_prgstrn(e) -1
EX_pro_D(e) -1
EX pro L(e) -1
EX_prostgd2(e) -1
EX_prostge1(e) -1
EX prostge2(e) -1
EX_prostgf2(e) -1
EX_ps_hs(e) -1
EX_ptdca(e) -1
EX_pyr(e) -1
EX_rbt(e) -1
EX_retfa(e) -1
EX retinol(e) -100
EX_retinol_9_cis(e) -1
```

```
EX_retinol_cis_11(e) -1
EX_retn(e) -100
EX_retnglc(e) -1
EX_Rtotal(e) -1
EX Rtotal2(e) -1
EX Rtotal3(e) -1
EX s2l2fn2m2masn(e) -1
EX s2l2n2m2masn(e) -1
EX_sarcs(e) -1
EX sel(e) -1
EX_ser_D(e) -1
EX ser L(e) -1
EX_{sl} L(e) -1
EX_so4(e) -100
EX spc hs(e) -1
EX sph1p(e) -1
EX sphs1p(e) -1
EX srtn(e) -1
EX_strch1(e) -1
EX_strch2(e) -1
EX_strdnc(e) -1
EX succ(e) -1
EX_sucr(e) -1
EX_tag_hs(e) -1
EX_tagat_D(e) -1
EX_taur(e) -1
EX taxol(e) -1
EX tchola(e) -1
EX tcynt(e) -1
EX tdchola(e) -1
EX tethex3(e) -1
EX tetpent3(e) -1
EX_tetpent6(e) -1
EX_tettet6(e) -1
EX thf(e) -1
EX_{thmmp(e)} -1
EX thmtp(e) -1
EX thr L(e) -1
EX_{thym}(e) -1
EX thymd(e) -1
EX_thyox_L(e) -1
EX_tmndnc(e) -1
EX_tolbutamide(e) -1
EX_tre(e) -1
EX triodthy(e) -1
EX_triodthysuf(e) -1
EX trp L(e) -1
EX tststerone(e) -1
EX_tststeroneglc(e) -1
EX tststerones(e) -1
EX tsul(e) -1
EX txa2(e) -1
EX tymsf(e) -1
EX_Tyr_ggn(e) -1
EX_tyr_L(e) -1
EX_udp(e) -1
EX ump(e) -1
EX_ura(e) -1
EX_urate(e) -1
EX urea(e) -1
EX_uri(e) -1
EX_utp(e) -1
EX_vacc(e) -1
EX_val_L(e) -1
EX_vitd2(e) -100
EX_whddca(e) -1
EX whhdca(e) -1
EX whtststerone(e) -1
```

```
EX xoltri25(e) -1
EX xoltri27(e) -1
EX xyl D(e) -1
EX yvite(e) -1
sink_pre_prot(r) -1
EX 4abutn(e) -1
EX_acmana(e) -1
EX ahdt(e) -1
EX ctp(e) -1
EX_dgmp(e) -1
EX dgtp(e) -1
EX_dha(e) -1
EX dhap(e) -1
EX dtmp(e) -1
EX_dttp(e) -1
EX_fad(e) -1
EX_fald(e) -1
EX_g1p(e) -1
EX_HC00229(e) -1
EX_HC00250(e) -1
EX_HC01104(e) -1
EX_HC01361(e) -1
EX HC01440(e) -1
EX HC01441(e) -1
EX HC01444(e) -1
EX HC01446(e) -1
EX HC01577(e) -1
EX HC01609(e) -1
EX HC01610(e) -1
EX_HC01700(e) -1
EX HC02160(e) -1
EX_HC02161(e) -1
EX itp(e) -1
EX orot(e) -1
EX prpp(e) -1
EX ptrc(e) -1
EX_pydx5p(e) -1
EX spmd(e) -1
EX_udpg(e) -1
EX_no2(e) -1
EX so3(e) -1
EX sprm(e) -1
EX prostgh2(e) -1
EX prostgi2(e) -1
EX_ppi(e) -1
EX cdp(e) -1
EX dtdp(e) -1
EX HC00955(e) -1
EX HC00001(e) -1
EX_HC00002(e) -1
EX_HC00003(e) -1
EX_HC00004(e) -1
EX citr L(e) -1
EX_HC01787(e) -1
EX_C02470(e) -1
EX HC01852(e) -1
EX_HC01939(e) -1
EX_HC01942(e) -1
EX_HC01943(e) -1
EX_HC01944(e) -1
EX_HC00822(e) -1
EX_C02528(e) -1
EX HC02192(e) -1
EX HC02193(e) -1
```

EX_whttdca(e) -1 EX_xolest_hs(e) -1 EX_xolest2_hs(e) -1 EX_xoltri24(e) -1

```
EX_HC02195(e) -1
EX HC02196(e) -1
EX HC02220(e) -1
EX_HC02154(e) -1
EX HC02175(e) -1
EX HC02176(e) -1
EX HC02199(e) -1
EX HC02200(e) -1
EX HC02201(e) -1
EX HC02172(e) -1
EX_HC02191(e) -1
EX_HC02194(e) -1
EX HC02197(e) -1
EX_HC02198(e) -1
EX HC02187(e) -1
EX HC02180(e) -1
EX HC02179(e) -1
EX HC02202(e) -1
EX_HC02203(e) -1
EX HC02204(e) -1
EX_HC02205(e) -1
EX HC02206(e) -1
EX_HC02207(e) -1
EX_HC02208(e) -1
EX_HC02210(e) -1
EX_HC02213(e) -1
EX HC02214(e) -1
EX HC02216(e) -1
EX HC02217(e) -1
EX malcoa(e) -1
EX_arachcoa(e) -1
EX coa(e) -1
EX CE2250(e) -1
EX_CE1935(e) -1
EX CE1940(e) -1
EX_CE1943(e) -1
EX CE2011(e) -1
EX_CE1936(e) -1
EX_CE1939(e) -1
EX maltttr(e) -1
EX_maltpt(e) -1
EX_malthx(e) -1
EX_CE2915(e) -1
EX_CE4722(e) -1
EX CE2916(e) -1
EX CE4723(e) -1
EX CE2917(e) -1
EX CE4724(e) -1
EX_malthp(e) -1
EX CE2839(e) -1
EX CE2838(e) -1
EX CE1950(e) -1
EX_cynt(e) -1
EX_23cump(e) -1
EX_3ump(e) -1
EX_CE5786(e) -1
EX CE5788(e) -1
EX_CE5789(e) -1
EX_CE5797(e) -1
EX CE5798(e) -1
EX CE5787(e) -1
EX CE5791(e) -1
EX CE5867(e) -1
EX_CE5868(e) -1
EX_CE5869(e) -1
EX_CE4633(e) -1
EX CE4881(e) -1
EX CE5854(e) -1
```

```
EX_glcur(e) -1
EX_CE1926(e) -1
EX_udpgal(e) -1
EX_crm_hs(e) -1
EX_galside_hs(e) -1
EX_CE0074(e) -1
EX_cdpea(e) -1
EX_12dgr120(e) -1
EX_CE5853(e) -1
EX_CE5853(e) -1
EX_CC5965(e) -1
EX_C05965(e) -1
EX_C05965(e) -1
maxConstraints:
```

Select the biomass reaction to optimise

```
model.biomassBool = strcmp(model.rxns, 'biomass_reaction');
model.c(model.biomassBool) = 1;
```

Display the biomass reaction

```
rxnAbbrList={'biomass_reaction'};
printFlag = 1;
formulas = printRxnFormula(model, rxnAbbrList, printFlag);
```

```
biomass reaction 20.6508 h2o[c] + 20.7045 atp[c] + 0.385872 glu L[c] + 0.352607 asp L[c] + 0.036117 gtp
```

Sparse flux balance analysis

We provide two options to run sparse flux balance analysis. A: directly in one step, no quality control, and B: two steps, all approximations, with a heuristic sparsity test.

TIMING

The time to compute a sparse flux balance analysis solution depends on the size of the genome-scale model and the option chosen to run sparse flux balance analysis. Option A: directly in one step, no quality control, can take anything from <0.1 seconds for a 1,000 reaction model, to 1,000 seconds for a model with 20,000 reactions. Option B: two steps, all approximations, with a sparsity test could take hours for a model with >10,000 reactions because the length of time for the heuristic sparsity test is proportional to the number of active reactions in an approximate sparse solution.

A. Sparse flux balance analysis (directly in one step, no quality control)

This approach computes a sparse flux balance analysis solution, satisfing the FBA objection, with the default approach to approximate the solution to the cardinality minimisation problem³ underling sparse FBA. This approach does not check the quality of the solution, i.e., whether indeed it is the sparsest flux vector satisfing the optimality criterion $c^Tv = \rho^*$.

First choose whether to maximize ('max') or minimize ('min') the FBA objective. Here we choose maximise

```
osenseStr='max';
```

Choose to minimize the zero norm of the optimal flux vector

```
minNorm='zero';
```

Run sparse flux balance analysis

```
sparseFBAsolution = optimizeCbModel(model, osenseStr, minNorm);
```

Obtain the vector of reaction rates from the solution structure

```
v = sparseFBAsolution.v;
```

Display the sparse flux solution, but only the non-zero fluxes

```
nonZeroFlag = 1;
printFluxVector(model, v, nonZeroFlag);
```

```
3MOBt2im 0.127657
3MOPt2im 49.4471
4ABUTtm 0.0439253
ABTArm 0.0439253
ABUTt2r 0.0439253
ADEt -3.17842
ADK1 -3.03312
ADK1m -0.0837315
ADK3 -0.27054
ADNtm -0.0837315
ALAt2r 1
ALATA_L -0.137857
AMPDA 0.147159
ARGtiDF 1
R_group_phosphotase_1 0.0559212
ASNt4 0.893617
ASPGLUm 0.127657
ASPTAm -0.127657
BTNDe 0.893613
CATm 0.024882
CDIPTr -0.0372829
CHOLt4 0.493981
CLS hs 0.0372829
CYOR u10m 9.95278
CYSTA 0.150649
CYTK4 0.155035
DAGK hs 0.0559212
DATPtn 0.04216
DCMPDA -0.155035
DCTPtn 0.030196
DESAT18 4 0.11773
DESAT18 7 0.11773
DGTPtn 0.0316544
DNDPt9m 0.0418657
DOPACHRMISO 2.29026
DSAT 0.0559212
DTTPtn 0.0418657
DURIK1 0.0932265
ENO 5.22241
EX 4abut(e) -0.0439253
EX ade(e) 2.17842
EX ala L(e) -1
EX_amp(e) 1
EX arg L(e) -1
EX_asn_L(e) -0.893617
EX_asp_L(e) -1
EX_atp(e) -1
```

```
EX_biocyt(e) -0.893613
```

EX btn(e) 0.893613

EX_chol(e) -0.493981

EX_chsterol(e) -0.0652435

EX_duri(e) -0.0932265

EX gln L(e) -1

 $EX_gly(e) -0.974083$

EX h2o2(e) -1

EX hco3(e) -0.0837315

EX his L(e) -0.404253

EX_ile_L(e) -0.914893

EX inost(e) -0.0745627

EX leu L(e) -1

EX lneldc(e) 0.11773

EX_lpchol_hs(e) -0.0559212

EX lys L(e) -1

EX_mercplaccys(e) 0.150649

EX met L(e) - 0.48936

 $EX_{no}(e) - 0.148933$

EX o2(e) -6.68552

EX_ocdca(e) -0.11773

EX orn(e) -0.84642

 $EX_pe_hs(e) -0.177089$

EX_pglyc_hs(e) -0.0466021

 $EX_phe_L(e) -0.829787$

EX_pi(e) 7.60762

EX pro L(e) -1

EX ps hs(e) -0.624468

EX_pyr(e) 4.20007

EX ser L(e) -1

EX sph1p(e) -0.0559212

EX thr L(e) -1

 $EX_{thymd}(e) - 0.0418657$

 $EX_{trp_L(e)} - 0.0425533$

 $EX_tyr_L(e) -0.510637$

EX ura(e) 0.767268

EX_utp(e) -1

EX val L(e) -1

FACOAL1822 -0.11773

FATP3t -0.11773

FBA 0.435143

FUMm 0.0439253

G3PD2m 0.0559212

GAPD 5.22241

GK1 0.147159

GLNS 0.0425533

GLNt4 1

GLUDxm 0.647824

GLUt2m 48.4859

GLYt2r 0.974083

GPDDA1 0.0559212

GTHRDt -49.6274

H202t 1

H20t 1.56714

H20tm -7.19359

HISt4 0.404253

HPYRRy 0.35051 ILEt4 0.914893

ILEt5m -49.4471

ILETA 49.4471

ILETAm -49.4471

INSTt2r 0.0745627

LEUt4 1

LNELDCt -0.11773

LPASE 0.0559212

LPCH0Lt 0.0559212

LYSt4 1.89361

MCLACCYSR 0.150649

MCLOR -0.150649

MDHm 0.0439253

MERCPLACCYSt 0.150649

METtec 0.48936

NADH2 u10m 6.18955

NDPK6 -0.705459

NTD7 3.09469

02t 6.68552

02tm 4.90175

ORNt3m -0.84642

ORNTArm 0.84642

ORNtiDF 0.84642

P5CDm 0.430173

P5CRm 0.0837315

PCm 0.0837315

PEt 0.177089

PGI -0.880086

PGK -5.22241

PGLYCt 0.0466021

PGM -5.22241

PHEtec 0.829787

PPM 3.94569

PR01xm 4.17729

PROt2r 1

PSSA1 hs -0.549902

PSt3 0.624468

PUNP1 3.17842

PYK 5.06486

PYNP2r 0.767268

PYRt2m 0.0837315

PYRt2r -4.20007

RNDR1 0.125891

RNDR2 0.0316544

RNDR4 0.0618088

RPI 2.63046

SMS 0.0559212

SPH1Pte -0.0559212

SPODMm 0.0497639

THMDt4 0.0418657

THRt4 1

TKT1 1.31523

TKT2 1.31523

TMDK1 0.0418657

TPI 1.80629

TRDR 0.0945153

TRIOK 0.35051

TRPt 0.0425533

TYRt 0.510637

UMPK -0.612233

UMPK6 -0.155035

URAt -0.767268

VALt4 1

VALt5m -0.127657

VALTAm -0.127657

EX ahdt(e) 1

EX_dgmp(e) 0.539243

EX dgtp(e) -0.539243

EX_HC00250(e) -0.450235

 $EX_HC01361(e) -1$

EX prpp(e) -1

r0010 0.5

r0047 0.0837315

r0051 -1

r0074 0.84642

r0145 -0.148933

r0160 0.35051

r0178 0.0439253

r0191 0.435143

r0193 -0.450235 r0276 0.147159 r0280 0.0840257 r0392 -0.35051 r0407 1.31523 r0408 0.921296 r0409 0.393933 r0410 0.539243 r0413 0.0316544 r0474 0.124839 r0509 0.0439253 r0707 -1 r0787 0.0559212 r0817 -0.148933 r0838 -0.647824 r0885 0.167463 r0892 -1 r0911 0.430173 r0940 -0.450235 r0941 0.0837315 r1050 0.0652435 r1116 -3.53924 r1117 0.0418657 r1143 1 r1156 -0.767268 r1423 5.66708 r1431 0.0837315 r1433 0.0837315 r1453 -3.66338 r2447 0.0932265 r2471 1 r2520 49.4599 RE0344C -0.11773 RE0452M 0.0418657 RE2675C 0.0559212 RE2954C 0.0418657 RE3198C 4.58051 RE3273C -0.111846 RE3301C 0.0559244 EX ppi(e) -1EX citr L(e) -0.148933 PIt9 -4.94055 CY00m3 4.97639 biomass reaction 3.19806 ALAALACNc 0.0643263 ALAALAPEPT1tc 0.0643263 LEULEULAPc 0.37234 LEULEUPEPT1tc 0.37234 PROGLYPEPT1tc 0.74932 PROGLYPRO1c 0.74932 3HC03 NAt 0.0279105 DATPtm 0.0418657 DUTPDP 0.0618088 EX alaala(e) -0.0643263 EX_leuleu(e) -0.37234 EX_progly(e) -0.74932 EX dpcoa(e) -2.53924 EX pan4p(e) 2.53924 FADH2ETC 0.0559212 NODe -0.148933 PTPATe -2.53924 RPEc 1.31523 3MOBte 0.127657 $EX_3mob(e) - 0.127657$

```
fprintf('%u%s\n',nnz(v),') active reactions in the sparse flux balance analysis solution.');
```

805 active reactions in the sparse flux balance analysis solution.

ANTICIPATED RESULTS

Typically, a sparse flux balance analysis solution will have a small fraction of the number of reactions active than in a flux balance analysis solution, e.g., Recon 2.04 model has 7,440 reactions. When maximising biomass production, a typical flux balance analysis solution might have approximately 2,000 active reactions (this is LP solver dependent) whereas for the same problem there are 247 active reactions in the sparse flux balance analysis solution from optimizeCbModel (using the default capped L1 norm approximate step function, see below).

B. Sparse flux balance analysis (two steps, all approximations, with a sparsity test)

This approach computes a sparse flux balance analysis solution, satisfing the FBA objection, with the default approach to approximate the solution to the cardinality minimisation problem³ underling sparse FBA. This approach does not check the quality of the solution, i.e., whether indeed it is the sparsest flux vector satisfing the optimality criterion $c^Tv = \rho^*$.

Solve a flux balance analysis problem

Build a linear programming problem structure (LPproblem) that is compatible with the interfacefunction (solveCobraLP) to any installed linear optimisation solver.

```
[c,S,b,lb,ub,csense] = deal(model.c,model.S,model.b,model.lb,model.ub,model.csense);
[m,n] = size(S);

LPproblem = struct('c',c,'osense',-1,'A',S,'csense',csense,'b',b,'lb',lb,'ub',ub);
```

Now solve the flux balance analysis problem

```
LPsolution = solveCobraLP(LPproblem);
if LPsolution.stat == 1
    vFBA = LPsolution.full(1:n);
else
    vFBA = [];
    error('FBA problem error!')
end
```

Display the number of active reactions

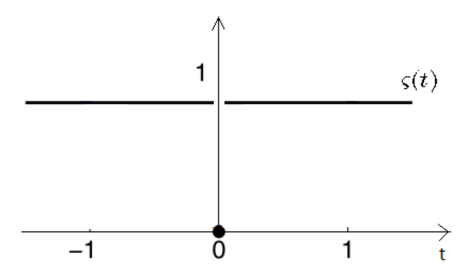
```
fprintf('%u%s\n',nnz(vFBA),' active reactions in the flux balance analysis solution.');
```

1876 active reactions in the flux balance analysis solution.

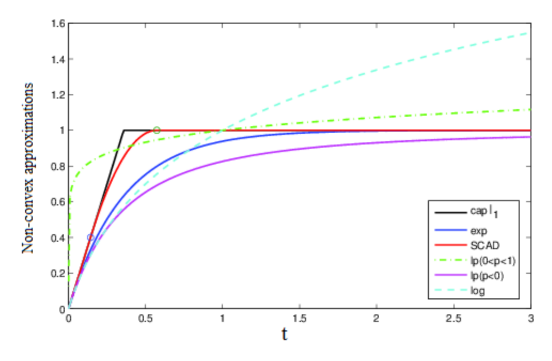
Approimations underlying sparse flux balance analysis

Due to its combinatorial nature, minimising the zero norm explicitly is an NP-hard problem. Therefore we approximately solve the problem. The approach is to replace the zero norm with a separable sum of

step functions, which are each approximated by anther function. Consider the step function $\zeta(t)$: R \to R where $\zeta(t)$ =1 if t \neq 0 and $\zeta(t)$ =0 otherwise, illustrated in the Figure below:



There are then many different approximate step functions that can be minimised. The figure below illustrates the many different approximate step functions that can be chosen to be minimised instead of an explicit step function.



Depending on the application, and the biochemical network, one or other approximation may outperform the rest, therefore a pragmatic strategy is to try each and select the most sparse flux vector. The step set of function approximations⁴ available are

* 'cappedL1' : Capped-L1 norm

* 'exp' : Exponential function

* 'log': Logarithmic function

* 'SCAD' : SCAD function

```
* 'lp-' : L_p norm with p<0
```

Here we prepare a cell array of strings which indicate the set of step function approximations we wish to compare.

```
approximations = {'cappedL1','exp','log','SCAD','lp-','lp+'};
```

Run the sparse linear optimisation solver

First we must build a problem structure to pass to the sparse solver, by adding an additional constraint requiring that the sparse flux solution also statisfy the optimal objective value from flux balance analysis

```
constraint.A = [S ; c'];
constraint.b = [b ; c'*vFBA];
constraint.csense = [csense; 'E'];
constraint.lb = lb;
constraint.ub = ub;
```

Now we call the sparse linear step function approximations

```
bestResult = n;
bestAprox = '';
for i=1:length(approximations)
    solution = sparseLP(char(approximations(i)),constraint);
    if solution.stat == 1
        nnzSol=nnz(abs(solution.x)>feasTol);
        fprintf('%u%s%s',nnzSol,' active reactions in the sparseFBA solution with ', char(approximations);
        if bestResult > nnzSol
            bestResult=nnzSol;
            bestAprox = char(approximations(i));
        solutionL0 = solution;
    end
end
end
```

```
247 active reactions in the sparseFBA solution with cappedL1 247 active reactions in the sparseFBA solution with exp 247 active reactions in the sparseFBA solution with log 247 active reactions in the sparseFBA solution with SCAD 247 active reactions in the sparseFBA solution with lp-247 active reactions in the sparseFBA solution with lp+
```

Select the most sparse flux vector, unless there is a numerical problem.

```
if ~isequal(bestAprox,'')
   vBest = solutionL0.x;
else
   vBest = [];
   error('Min L0 problem error !!!!')
end
```

Report the best approximation

```
display(strcat('Best step function approximation: ',bestAprox));
```

^{* &#}x27;lp+' : L_p norm with 0<p<1

```
Best step function approximation:cappedL1
```

Report the number of active reactions in the most sparse flux vector

```
fprintf('%u%s',nnz(abs(vBest)>feasTol),' active reactions in the best sparse flux balance anal
247 active reactions in the best sparse flux balance analysis solution.
```

Warn if there might be a numerical issue with the solution

```
feasError=norm(constraint.A * solutionL0.x - constraint.b,2);
if feasError>feasTol
    fprintf('%g\t%s\n',feasError, ' feasibily error.')
    warning('Numerical issue with the sparseLP solution')
end
```

Heuristically check if the selected set of reactions is minimal

Each step function approximation minimises a different problem than minimising the zero norm explicitly. Therefore it is wise to test, at least heuristically, if the most sparse approximate solution to minimising the zero norm is at least locally optimal, in the sense that the set of preicted reactions cannot be reduced by omitting, one by one, an active reaction. If it is locally optimal in this sense, one can be more confident that the most sparse approximate solution is the most sparse solution, but still there is no global guarantee, as it is a combinatorial issue.

Identify the set of predicted active reactions

```
activeRxnBool = abs(vBest)>feasTol;
nActiveRnxs = nnz(activeRxnBool);
activeRxns = false(n,1);
activeRxns(activeRxnBool) = true;
minimalActiveRxns=activeRxns;
```

Close all predicted non-active reactions by setting their lb = ub = 0

```
lbSub = model.lb;
ubSub = model.ub;
lbSub(~activeRxns) = 0;
lbSub(~activeRxns) = 0;
```

Generate an LP problem to be reduced

```
% Check if one still can achieve the same objective LPproblem = struct('c',-c,'osense',-1,'A',S,'csense',csense,'b',b,'lb',lbSub,'ub',ubSub);
```

For each active reaction in the most sparse approximate flux vector, one by one, set the reaction bounds to zero, then test if the optimal flux balance analysis objective value is still attained. If it is, then that reaction is not part of the minimal set. If it is not, then it is probably part of the minimal set.

```
for i=1:n
   if activeRxnBool(i)
      LPproblem.lb = model.lb;
   LPproblem.ub = model.ub;
```

```
%close bounds on this reaction
        LPproblem.lb(i) = 0;% Close the reaction
        LPproblem.ub(i) = 0;% Close the reaction
        %solve the LP problem
        LPsolution = solveCobraLP(LPproblem);
        %check if the optimal FBA objective is attained
        if LPsolution.stat == 1 && abs(LPsolution.obj + c'*vFBA)<1e-8</pre>
            minimalActiveRxns(i) = 0;
            vBestTested = LPsolution.full(1:n);
        else
            %relax those bounds if reaction appears to be part of the minimal set
            LPproblem.lb(i) = model.lb(i);
            LPproblem.ub(i) = model.ub(i);
        end
    end
end
```

Report the number of active reactions in the approximately most sparse flux vector, or the reduced approximately most sparse flux vector, if it is more sparse.

```
if nnz(minimalActiveRxns)if nnz(minimalActiveRxns)fprintf('%u%s',nnz(abs(vBestTested)>feasTol),' active reactions in the best sparseFBA solution
nonZeroFlag = 1;
printFluxVector(model, vBestTested, nonZeroFlag);
else
    fprintf('%u%s',nnz(abs(vBest)>feasTol),' active reactions in the best sparseFBA solution (end)
```

247 active reactions in the best sparseFBA solution (tested).

REFERENCES

- [1] Meléndez-Hevia, E., Isidoro, A. (1085). The game of the pentose phosphate cycle. Journal of Theoretical Biology 117, 251-263.
- [2] Thiele, I., Swainston, N., Fleming, R.M., Hoppe, A., Sahoo, S., Aurich, M.K., Haraldsdottir, H., Mo, M.L., Rolfsson, O., Stobbe, M.D., et al. (2013). A community-driven global reconstruction of human metabolism. Nat Biotechnol 31, 419-425.
- [3] Fleming, R.M.T., et al. (*submitted*, 2017). Cardinality optimisation in constraint-based modelling: illustration with Recon 3D.
- [4] Le Thi, H.A., Pham Dinh, T., Le, H.M., and Vo, X.T. (2015). DC approximation approaches for sparse optimization. European Journal of Operational Research 244, 26-46.