# Thermodynamically constrain a metabolic model

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**Reviewers:** 

#### INTRODUCTION

In flux balance analysis of genome scale stoichiometric models of metabolism, the principal constraints are uptake or secretion rates, the steady state mass conservation assumption and reaction directionality. Von Bertylanffy [1,4] is a set of methods for (i) quantitative estimation of thermochemical parameters for metabolites and reactions using the component contribution method [3], (ii) quantitative assignment of reaction directionality in a multi-compartmental genome scale model based on an application of the second law of thermodynamics to each reaction [2], (iii) analysis of thermochemical parameters in a network context, and (iv) thermodynamically constrained flux balance analysis. The theoretical basis for each of these methods is detailed within the cited papers.

#### **PROCEDURE**

# Configure the environment

All the installation instructions are in a separate .md file named vonBertalanffy.md in docs/source/installation

With all dependencies installed correctly, we configure our environment, verfy all dependencies, and add required fields and directories to the matlab path.

```
initVonBertalanffy
```

# Select the model

This tutorial is tested for the E. coli model iAF1260 and the human metabolic model Recon3Dmodel. However, only the data for the former is provided within the COBRA Toolbox as it is used for testing von Bertylanffy. However, the figures generated below are most suited to plotting results for Recon3D, so they may not be so useful for iAF1260. The Recon3D example uses values from literature for input variables where they are available.

```
modelName = 'iAF1260';
%modelName='Ec_iAF1260_flux1'; %uncomment this line and comment the line below if you w
%modelName='Recon3DModel_Dec2017';
```

#### Load a model

Load a model, and save it as the original model in the workspace, unless it is already loaded into the workspace.

```
global CBTDIR
modelFileName = [modelName '.mat']
modelFileName =
'iAF1260.mat'
modelDirectory = getDistributedModelFolder(modelFileName); %Look up the folder for the
modelFileName= [modelDirectory filesep modelFileName]; % Get the full path. Necessary t
switch modelName
    case 'Ec_iAF1260_flux1'
        modelFileName = [modelName '.xml']
        model = readCbModel(modelFileName);
        if model.S(952, 350) == 0
            model.S(952, 350)=1; % One reaction needing mass balancing in iAF1260
        end
        model.metCharges(strcmp('asntrna[Cytosol]', model.mets))=0; % One reaction need
    case 'iAF1260'
        model = readCbModel(modelFileName);
        model.mets = cellfun(@(mets) strrep(mets,'_c','[c]'),model.mets,'UniformOutput
        model.mets = cellfun(@(mets) strrep(mets,'_e','[e]'),model.mets,'UniformOutput
        model.mets = cellfun(@(mets) strrep(mets,'_p','[p]'),model.mets,'UniformOutput
        bool = strcmp(model.mets, 'lipa[c]old[c]');
        model.mets{bool}='lipa_old_[c]';
        bool = strcmp(model.mets,'lipa[c]old[e]');
        model.mets{bool}='lipa_old_[e]';
        bool = strcmp(model.mets, 'lipa[c]old[p]');
        model.mets{bool}='lipa_old_[p]';
        if model.S(952, 350) == 0
            model.S(952, 350)=1; % One reaction needing mass balancing in iAF1260
        end
        model.metCharges(strcmp('asntrna[c]', model.mets))=0; % One reaction needing ch
    case 'Recon3DModel_Dec2017'
      model = readCbModel(modelFileName);
      model.csense(1:size(model.S,1),1)='E';
      %Hack for thermodynamics
      model.metFormulas{strcmp(model.mets, 'h[i]')}='H';
      model.metFormulas(cellfun('isempty',model.metFormulas)) = {'R'};
      if isfield(model, 'metCharge')
          model.metCharges = double(model.metCharge);
          model=rmfield(model,'metCharge');
      modelOrig = model;
    otherwise
            error('setup specific parameters for your model')
end
```

Each model.subSystems $\{x\}$  is a character array, and this format is retained.

clear model

## Set the directory containing the results

```
switch modelName
    case 'Ec_iAF1260_flux1'
        resultsPath=which('tutorial_vonBertalanffy.mlx');
        resultsPath=strrep(resultsPath,'/tutorial_vonBertalanffy.mlx','');
        resultsPath=[resultsPath filesep modelName '_results'];
        resultsBaseFileName=[resultsPath filesep modelName '_results'];
    case 'iAF1260'
        resultsPath=which('tutorial_vonBertalanffy.mlx');
        resultsPath=strrep(resultsPath,'/tutorial_vonBertalanffy.mlx','');
        resultsPath=[resultsPath filesep modelName '_results'];
        resultsBaseFileName=[resultsPath filesep modelName '_results'];
    case 'Recon3DModel Dec2017'
        basePath='~/work/sbgCloud';
        resultsPath=[basePath '/programReconstruction/projects/recon2models/results/the
        resultsBaseFileName=[resultsPath filesep modelName '_' datestr(now,30) '_'];
        error('setup specific parameters for your model')
end
```

## Set the directory containing molfiles

## Set the thermochemical parameters for the model

```
switch modelName
    case 'Ec_iAF1260_flux1'
        T = 310.15; % Temperature in Kelvin
        compartments = {'Cytosol'; 'Extra_organism'; 'Periplasm'}; % Cell compartment :
        ph = [7.7; 7.7; 7.7]; % Compartment specific pH
        is = [0.25; 0.25; 0.25]; % Compartment specific ionic strength in mol/L
        chi = [0; 90; 90]; % Compartment specific electrical potential relative to cyto
    case 'iAF1260'
        T = 310.15; % Temperature in Kelvin
        compartments = ['c'; 'e'; 'p']; % Cell compartment identifiers
        ph = [7.7; 7.7; 7.7]; % Compartment specific pH
        is = [0.25; 0.25; 0.25]; % Compartment specific ionic strength in mol/L
        chi = [0; 90; 90]; % Compartment specific electrical potential relative to cyto
    case 'Recon3DModel Dec2017'
```

```
% Temperature in Kelvin
T = 310.15;
% Cell compartment identifiers
compartments = ['c'; 'e'; 'g'; 'l'; 'm'; 'n'; 'r'; 'x';'i'];
% Compartment specific pH
ph = [7.2; 7.4; 6.35; 5.5; 8; 7.2; 7.2; 7; 7.2];
% Compartment specific ionic strength in mol/L
is = 0.15*ones(length(compartments),1);
% Compartment specific electrical potential relative to cytosol in mV
chi = [0; 30; 0; 19; -155; 0; 0; -2.303*8.3144621e-3*T*(ph(compartments == 'x'
otherwise
    error('setup specific parameters for your model')
```

### Set the default range of metabolite concentrations

```
switch modelName
    case 'Ec_iAF1260_flux1'
        concMinDefault = 1e-5; % Lower bounds on metabolite concentrations in mol/L
        concMaxDefault = 0.02; % Upper bounds on metabolite concentrations in mol/L
        metBoundsFile=[];
    case 'iAF1260'
        concMinDefault = 1e-5; % Lower bounds on metabolite concentrations in mol/L
        concMaxDefault = 0.02; % Upper bounds on metabolite concentrations in mol/L
        metBoundsFile=[];
    case 'Recon3DModel_Dec2017'
        concMinDefault=1e-5; % Lower bounds on metabolite concentrations in mol/L
        concMaxDefault=1e-2; % Upper bounds on metabolite concentrations in mol/L
        metBoundsFile=which('HumanCofactorConcentrations.txt');%already in the COBRA to otherwise
        error('setup specific parameters for your model')
end
```

# Set the desired confidence level for estimation of thermochemical parameters

The confidence level for estimated standard transformed reaction Gibbs energies is used to quantitatively assign reaction directionality.

```
switch modelName
    case 'Ec_iAF1260_flux1'
        confidenceLevel = 0.95;
        DrGt0_Uncertainty_Cutoff = 20; %KJ/KMol
    case 'iAF1260'
        confidenceLevel = 0.95;
        DrGt0_Uncertainty_Cutoff = 20; %KJ/KMol
    case 'Recon3DModel_Dec2017'
        confidenceLevel = 0.95;
        DrGt0_Uncertainty_Cutoff = 20; %KJ/KMol
    otherwise
        error('setup specific parameters for your model')
end
```

## Prepare folder for results

```
if ~exist(resultsPath,'dir')
    mkdir(resultsPath)
end
cd(resultsPath)
```

# Set the print level and decide to record a diary or not (helpful for debugging)

```
printLevel=2;
diary([resultsPath filesep 'diary.txt'])
```

# Setup a thermodynamically constrained model

#### Read in the metabolite bounds

```
setDefaultConc=1;
setDefaultFlux=0;
rxnBoundsFile=[];
model=readMetRxnBoundsFiles(model,setDefaultConc,setDefaultFlux,concMinDefault,concMaxI
```

## **Check inputs**

```
model = configureSetupThermoModelInputs(model,T,compartments,ph,is,chi,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDefault,concMinDef
```

# Check elemental balancing of metabolic reactions

```
ignoreBalancingOfSpecifiedInternalReactions=1;
if ~exist('massImbalance','var')
    if isfield(model,'Srecon')
        model.S=model.Srecon;
    end
    % Check for imbalanced reactions
    fprintf('\nChecking mass and charge balance.\n');
    Heuristically identify exchange reactions and metabolites exclusively involved in
    if ~isfield(model,'SIntMetBool') | ~isfield(model,'SIntRxnBool') | ignoreBalance
        %finds the reactions in the model which export/import from the model
        %boundary i.e. mass unbalanced reactions
        %e.g. Exchange reactions
              Demand reactions
              Sink reactions
       model = findSExRxnInd(model,[],printLevel);
    end
    if ignoreBalancingOfSpecifiedInternalReactions
```

```
[nMet,nRxn]=size(model.S);
        ignoreBalancingMetBool=false(nMet,1);
              if strcmp(model.mets{m}, 'Rtotal3coa[m]')
응
응
                  pause(0.1);
응
              end
            if ~isempty(model.metFormulas{m})
                ignoreBalancingMetBool(m,1)=numAtomsOfElementInFormula(model.metFormula
            end
        end
        ignoreBalancingRxnBool=getCorrespondingCols(model.S,ignoreBalancingMetBool,mode
        SIntRxnBool=model.SIntRxnBool;
        model.SIntRxnBool=model.SIntRxnBool & ~ignoreBalancingRxnBool;
    end
   printLevelcheckMassChargeBalance=-1; % -1; % print problem reactions to a file
    %mass and charge balance can be checked by looking at formulas
    [massImbalance,imBalancedMass,imBalancedCharge,imBalancedRxnBool,Elements,missingFo
        = checkMassChargeBalance(model,printLevelcheckMassChargeBalance,resultsBaseFile
    model.balancedRxnBool=~imBalancedRxnBool;
    model.balancedMetBool=balancedMetBool;
    model.Elements=Elements;
    model.missingFormulaeBool=missingFormulaeBool;
    %reset original boolean vector
    if ignoreBalancingOfSpecifiedInternalReactions
        model.SIntRxnBool=SIntRxnBool;
    end
end
```

```
Checking mass and charge balance.

Assuming biomass reaction is: BIOMASS_Ec_iAF1260_core_59p81M

ATP maintenance reaction is not considered an exchange reaction by default. It should be mass balanced:

ATPM atp[c] + h2o[c] -> adp[c] + h[c] + pi[c]

There are mass imbalanced reactions, see /home/rfleming/work/sbgCloud/code/fork-COBRA.tutorials/analysis/v
```

# Check that the input data necessary for the component contribution method is in place

```
model = setupComponentContribution(model,molfileDir);

Creating MetStructures.sdf from molfiles.
Percentage of metabolites without mol files: 100.0%
Converting SDF to InChI strings.
Estimating metabolite pKa values.
Assuming that metabolite species in model.metFormulas are representative for metabolites where pKa could representative.
```

# Prepare the training data for the component contribution method

```
training_data = prepareTrainingData(model,printLevel);
```

Successfully added 3914 values from TECRDB Successfully added 223 formation energies

```
Successfully added 13 redox potentials
Loading the InChIs for the training data from: /home/rfleming/work/sbgCloud/code/fork-cobratoolbox/src/ana
Successfully created balanced training-data structure: 672 compounds and 3061 reactions
Loading the pKa values for the training data from: cache/kegg_pkas.mat
Mapping model metabolites to nist compounds
Creating group incidence matrix
Performing reverse transform
```

### Call the component contribution method

```
if ~isfield(model,'DfG0')
    [model,~] = componentContribution(model,training_data);
end
```

Running Component Contribution method

## Setup a thermodynamically constrained model

```
if ~isfield(model,'DfGt0')
   model = setupThermoModel(model,confidenceLevel);
end
```

Estimating standard transformed Gibbs energies of formation.

Estimating bounds on transformed Gibbs energies.

Additional effect due to possible change in chemical potential of Hydrogen ions for transport reactions. Additional effect due to possible change in electrical potential for transport reactions.

# Generate a model with reactants instead of major microspecies

```
if ~isfield(model,'Srecon')
    printLevel_pHbalanceProtons=-1;

model=pHbalanceProtons(model,massImbalance,printLevel_pHbalanceProtons,resultsBaseRend
```

Warning: vonBertalanffy:pHbalanceProtons 'Hydrogen unbalanced reconstruction reactions exist!

# Determine quantitative directionality assignments

```
if ~exist('directions','var')
    fprintf('Quantitatively assigning reaction directionality.\n');
    [modelThermo, directions] = thermoConstrainFluxBounds(model,confidenceLevel,DrGt0_tend
```

```
Quantitatively assigning reaction directionality. 3/2382 reactions with DrGtMin=DrGtMax=0 4 inactive reactions (lb = ub = 0) The following reactions have DrGtMax=DrGtMin=0: H2Otex h2o[e] <=> h2o[p] H2Otpp h2o[p] <=> h2o[c] Htex h[e] <=> h[p]
```

## Analyse thermodynamically constrained model

Choose the cutoff for probablity that reaction is reversible

```
cumNormProbCutoff=0.2;
```

#### Build Boolean vectors with reaction directionality statistics

[modelThermo,directions]=directionalityStats(modelThermo,directions,cumNormProbCutoff,p

```
3/2382 reactions with DrGtMin=DrGtMax=0
Qualitative internal reaction directionality:
     2077
              internal reconstruction reaction directions.
     1520
             forward reconstruction assignment.
        0
             reverse reconstruction assignment.
      553
             reversible reconstruction assignment.
Quantitative internal reaction directionality:
     2077
              internal reconstruction reaction directions.
      549
              of which have a thermodynamic assignment.
     1525
              of which have no thermodynamic assignment.
       17
              forward thermodynamic only assignment.
              reverse thermodynamic only assignment.
        0
      532
              reversible thermodynamic only assignment.
Qualitiative vs Quantitative:
      335
              Reversible -> Reversible
        0
              Reversible -> Forward
        0
              Reversible -> Reverse
      215
              Reversible -> Uncertain
       16
              Forward -> Forward
              Forward -> Reverse
      196
              Forward -> Reversible
     1308
              Forward -> Uncertain
        0
              Reverse -> Reverse
        0
              Reverse -> Forward
              Reverse -> Reversible
        0
              Reversible -> Uncertain
        0
Breakdown of relaxation of reaction directionality, Qualitiative vs Quantitative:
      196
              qualitatively forward reactions that are quantitatively reversible (total).
      136
              of which are quantitatively reversible by range of dGt0. P(\Delta_{r}^{r}G^{r}) > 0.7
              of which are quantitatively reversible by range of dGt0. 0.3< P(\Delta_{r})^{0} < 0
        1
              of which are quantitatively reversible by range of dGt0. P(\Delta_{r}G^{\circ}) < 0.3
              of which are quantitatively forward by fixed dGr0t, but reversible by concentration alone (
              of which are quantitatively reverse by dGr0t, but reversible by concentration (negative fix
              of which are quantitatively forward by dGr0t, but reversible by concentration (positve fixed)
        0
        3
              of which are quantitatively reverse by dGr0t, but reversible by concentration (uncertain ne
              of which are quantitatively forward by dGr0t, but reversible by concentration (uncertain po
% directions
                   a structue of boolean vectors with different directionality
응
                   assignments where some vectors contain subsets of others
응
응
  qualtiative -> quantiative changed reaction directions
응
    .forward2Forward
    .forward2Reverse
응
응
    .forward2Reversible
응
    .forward2Uncertain
응
    .reversible2Forward
응
    .reversible2Reverse
응
    .reversible2Reversible
```

```
.reversible2Uncertain
응
    .reverse2Forward
응
    .reverse2Reverse
    .reverse2Reversible
응
응
    .reverse2Uncertain
왕
    .tightened
%
% subsets of qualtiatively forward -> quantiatively reversible
응
    .forward2Reversible_bydGt0
응
    .forward2Reversible_bydGt0LHS
    .forward2Reversible_bydGt0Mid
응
응
    .forward2Reversible_bydGt0RHS
응
응
    .forward2Reversible_byConc_zero_fixed_DrG0
    .forward2Reversible_byConc_negative_fixed_DrG0
응
응
    .forward2Reversible_byConc_positive_fixed_DrG0
    .forward2Reversible_byConc_negative_uncertain_DrG0
응
응
    .forward2Reversible_byConc_positive_uncertain_DrG0
```

Write out reports on directionality changes for individual reactions to the results folder.

```
fprintf('%s\n','directionalityChangeReport...');
```

directionalityChangeReport...

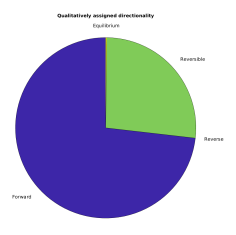
 ${\tt directionalityChangeReport(modelThermo, directions, cumNormProbCutoff, printLevel, results Boundary ChangeReport(modelThermo, directions, cumNormProbCutoff, cumNorm$ 

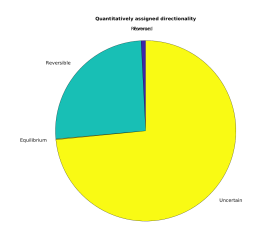
Generate pie charts with proportions of reaction directionalities and changes in directionality

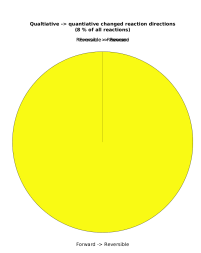
```
fprintf('%s\n','directionalityStatFigures...');
```

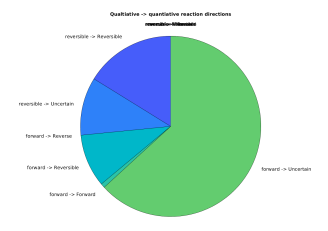
directionalityStatFigures...

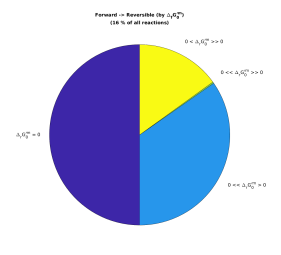
directionalityStatsFigures(directions,resultsBaseFileName)

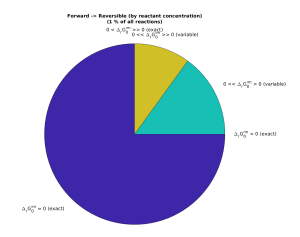








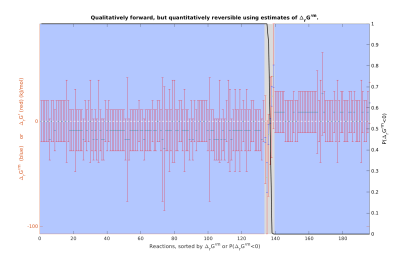


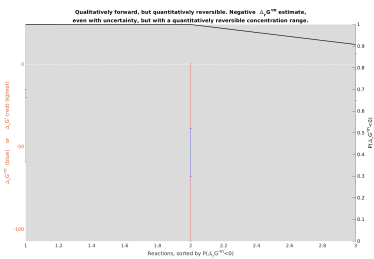


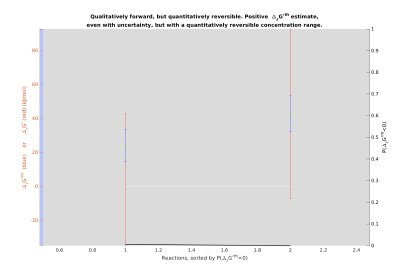
Generate figures to interpret the overall reasons for reaction directionality changes for the qualitatively forward now quantiatiavely reversible reactions

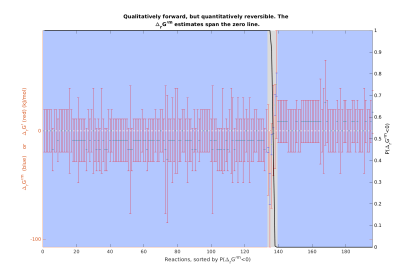
```
if any(directions.forward2Reversible)
    fprintf('%s\n','forwardReversibleFigures...');
    forwardReversibleFigures(modelThermo,directions,confidenceLevel)
end
```

forwardReversibleFigures...









Write out tables of experimental and estimated thermochemical parameters for the model

generateThermodynamicTables(modelThermo,resultsBaseFileName);

#### **REFERENCES**

- [1] Fleming, R. M. T. & Thiele, I. von Bertalanffy 1.0: a COBRA toolbox extension to thermodynamically constrain metabolic models. Bioinformatics 27, 142–143 (2011).
- [2] Haraldsdóttir, H. S., Thiele, I. & Fleming, R. M. T. Quantitative assignment of reaction directionality in a multicompartmental human metabolic reconstruction. Biophysical Journal 102, 1703–1711 (2012).
- [3] Noor, E., Haraldsdóttir, H. S., Milo, R. & Fleming, R. M. T. Consistent Estimation of Gibbs Energy Using Component Contributions. PLoS Comput Biol 9, e1003098 (2013).
- [4] Fleming, R. M. T., Predicat, G., Haraldsdóttir, H. S., Thiele, I. von Bertalanffy 2.0 (in preparation).