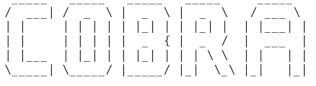
# Metabotools tutorial I

In this tutorial we generate contextualized models of two lymphoblastic leukemia cell lines, CCRF-CEM and Molt- 4 cells, will be generated by integrating semi-quantitative metabolomic data, transcriptomic data, and growth rates. We will afterwards analyze the solution space of these models by sampling analysis.

Before running a section in the tutorial, read the corresponding sections in the MetaboTools protocol and supplemental tutorial (Data sheet 2, http://journal.frontiersin.org/article/10.3389/fphys.2016.00327/ full).

Clear workspace and initialize the COBRA Toolbox





The COBRA Toolbox - 2017

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 ...
- > TranslateSBML is installed and working properly.
- > Configuring solver environment variables ...
  - [----] ILOG CPLEX PATH: /opt/ibm/ILOG/CPLEX Studio1271/cplex/matlab/x86-64 linux
  - [----] GUROBI PATH: /home/syarra/Dropbox/software/qurobi/gurobi652/linux64/matlab
  - [----] TOMLAB PATH : --> set this path manually after installing the solver ( see instructions )
  - [----] MOSEK PATH: /home/syarra/Dropbox/software/mosek/linux/8/
- > Checking available solvers and solver interfaces ... Done.
- > Setting default solvers ... Done.
- > Saving the MATLAB path ... Done.
  - The MATLAB path was saved as ~/pathdef.m.
- > Summary of available solvers and solver interfaces

Support	LP MIL	.P QP	MIQP	NLP			
cplex direct	full		0	0	0	0	-
dqqMinos	full		1	-	-	-	-
glpk	full		1	1	-	-	-
gurobi	full		1	1	1	1	-
ibm_cplex	full		1	1	1	-	-
matlab	full		1	-	-	-	1
mosek	full		1	1	1	-	-
pdco	full		1	-	1	-	-
quadMinos	full		1	-	-	-	1
tomlab_cplex	full		0	0	0	0	-
qpng	experime	ental	-	-	1	-	-
tomlab_snopt	experime	ental	-	-	-	-	0

```
0
                                                0
gurobi_mex
              legacy
                                                       0
 lindo old
              legacy
                                 0
lindo legacy legacy
                                 0
lp solve
              legacy
                                 1
                                 0
opti
              legacy
                                                0
                                                       0
                                                              0
Total
                                  9
                                         4
                                                5
                                                       1
                                                              2
+ Legend: - = not applicable, 0 = solver not compatible or not installed, 1 = solver installed.
> You can solve LP problems using: 'dqqMinos' - 'glpk' - 'gurobi' - 'ibm cplex' - 'matlab' - 'mosek' -
> You can solve MILP problems using: 'glpk' - 'gurobi' - 'ibm_cplex' - 'mosek'
> You can solve QP problems using: 'gurobi' - 'ibm cplex' - 'mosek' - 'pdco' - 'qpng'
> You can solve MIQP problems using: 'gurobi'
> You can solve NLP problems using: 'matlab' - 'quadMinos'
> Checking for available updates ...
ssh: /usr/local/MATLAB/R2016a/bin/glnxa64/libcrypto.so.1.0.0: no version information available (required
ssh: /usr/local/MATLAB/R2016a/bin/glnxa64/libcrypto.so.1.0.0: no version information available (required
OpenSSL version mismatch. Built against 1000207f, you have 100010bf
fatal: Could not read from remote repository.
Please make sure you have the correct access rights
and the repository exists.
> The changes of The COBRA Toolbox could not be fetched. > There are 169 new commit(s) on <master> and
> You can update The COBRA Toolbox by running updateCobraToolbox() (from within MATLAB).
```

### Step 0 Alter to define our own output location and LP solver!!

#### Define variables

```
global CBTDIR % set path to cobratoolbox (pathToCOBRA)
% ouputPath = 'ADD YOUR PATH TO YOUR OUTPUT FOLDER'; %
% solver = 'ADD YOUR SOLVER'; %, e.g., 'cplex_direct' for ILOG
outputPath = pwd;
solver = 'glpk';
```

#### set and check solver

```
solverOK = changeCobraSolver(solver, 'LP');

if solverOK == 1
    fprintf('Solver %s is set.\n', solver);
else
    error('Solver %s could not be used. Check if %s is in the matlab path (set path) or check end
```

Solver glpk is set.

#### load and check input is loaded correctly

```
tutorialPath = [CBTDIR filesep 'tutorials' filesep 'metabotools' filesep 'tutorial_I'];
if isequal(exist([tutorialPath filesep 'starting_model.mat'], 'file'), 2) % 2 means it's a filesed ([tutorialPath filesep 'starting_model.mat']);
    fprintf('The model is loaded.\n');
else
```

```
error('The model ''starting_model'' could not be loaded.');
end
```

The model is loaded.

#### Check output path and writing permission

```
if ~exist(outputPath, 'dir') == 7
    error('Output directory in ''outputPath'' does not exist. Verify that you type it correctle
end
% make and save dummy file to test writing to output directory
A = rand(1);
try
    save([outputPath filesep 'A']);
catch ME
    error('Files cannot be saved to the provided location: %s\nObtain rights to write into %s end
```

### Step 1: Shaping the model's environment using setMediumConstraints

RPMI medium composition.

```
medium composition = {'EX ala L(e)'
                        'EX arg L(e)'
                        'EX asn L(e)'
                        'EX asp L(e)'
                       'EX cys L(e)'
                       'EX gln L(e)'
                        'EX_glu_L(e)'
                       'EX gly(e)'
                       'EX his L(e)'
                        'EX ile L(e)'
                        'EX leu L(e)'
                       'EX lys L(e)'
                       'EX met L(e)'
                        'EX_phe_L(e)'
                        'EX 4HPR0(e)'
                       'EX pro L(e)'
                        'EX ser L(e)'
                        'EX thr L(e)'
                       'EX trp L(e)'
                       'EX tyr L(e)'
                        'EX val L(e)'
                        'EX ascb L(e)'
                        'EX btn(e)'
                        'EX chol(e)'
                        'EX pnto R(e)'
                        'EX fol(e)'
                       'EX ncam(e)'
                        'EX pydxn(e)'
                        'EX ribflv(e)'
                        'EX thm(e)'
                       'EX inost(e)'
                        'EX ca2(e)'
                        'EX fe3(e)'
                        'EX k(e)'
```

```
'EX hco3(e)'
                        'EX_na1(e)'
                        'EX_pi(e)'
                        'EX_glc(e)'
                        'EX hxan(e)'
                        'EX lnlc(e)'
                        'EX_lipoate(e)'
                        'EX pyr(e)'
                        'EX thymd(e)'
                        'EX gthrd(e)'
                        'EX_anth(e)'
                       };
% Medium concentrations
met_Conc_mM = [0.1]
                1.15
                0.15
                0.379
                0.208
                2
                0.136
                0.133
                0.0968
                0.382
                0.382
                0.274
                0.101
                0.0909
                0.153
                0.174
                0.286
                0.168
                0.0245
                0.129
                0.171
                0.00863
                0.00082
                0.0214
                0.000524
                0.00227
                0.082
                0.00485
                0.000532
                0.00297
                0.194
                0.424
                0
                5.33
                23.81
                127.26
                5.63
                11.11
                0
                0
                0
                1
                0
                0.00326
                0.0073
                ];
```

```
current_inf = 1000;
set_inf = 500;

cellConc = 2.17 * 1e6;
t = 48;
cellWeight = 3.645e-12;
```

Definition of basic medium (defines uptake from the medium, not captured by the medium composition, all with same constraints)

```
mediumCompounds = {'EX_co2(e)'; 'EX_h(e)'; 'EX_h2o(e)'; 'EX_hco3(e)'; 'EX_nh4(e)'; 'EX_o2(e)';
mediumCompounds_lb = -100;
```

Additional constraints to limit the model behaviour, e.g., secretion of oxygen, essAA that need to be taken up (and that have not been measured).

```
customizedConstraints = {'EX_o2(e)'; 'EX_strch1(e)'; 'EX_acetone(e)'; 'EX_glc(e)'; 'EX_his_L(e)')
customizedConstraints_lb = [-2.3460
                             0
                             -500
                             - 100
                             - 100
                             -100
                             ];
customizedConstraints ub = [500
                             0
                             0
                             500
                             500
                             500
                             500
                             ];
[modelMedium, ~] = setMediumConstraints(starting model, set inf, current inf, medium compositi
```

### Step 2 calculateLODs

```
'EX trp L(e)'
            'EX orn(e)'
            'EX_arg_L(e)'
            'EX_thr_L(e)'
            'EX fol(e)'
            'EX_gln_L(e)'
            'EX_4pyrdx(e)'
            'EX ser L(e)'
            'EX glc(e)'
            'EX ribflv(e)'
            'EX_glu_L(e)'
            'EX tyr L(e)'
            'EX phe L(e)'
            'EX inost(e)'
            'EX_Lcystin(e)'
            'EX leu L(e)'
            'EX met L(e)'
            'EX cys L(e)'
            'EX asn L(e)'
            'EX mal L(e)'
            'EX ile L(e)'
            'EX_pyr(e)'
            'EX lys L(e)'
            'EX ala L(e)'
            'EX cit(e)'
            'EX_pro_L(e)'
            'EX gly(e)'
            'EX asp L(e)'
            'EX_34hpp'
            'EX octa(e)'
            'EX 4mop(e)'
            'EX glyb(e)'
            'EX_val_L(e)'
            'EX ade(e)'
            'EX hxan(e)'
            'EX gua(e)'
            'EX_ins(e)'
            'EX orot(e)'
            'EX ura(e)'
            'EX ahcys(e)'
            'EX_cbasp(e)'
            'EX Lcystin(e)'
            'EX ser L(e)'
            'EX_cys_L(e)'
            'EX thm(e)'
            'EX arg L(e)'
            'EX ncam(e)'
           };
theo mass = [298.0974]
              243.0617
              104.1075
              123.0558
              129.0552
              117.0188
              220.1185
              128.0348
              265.1123
              138.0555
              132.0661
              89.0239
```

```
115.0395
              156.0773
              205.0977
              133.0977
              175.1195
              120.0661
              440.1319
              147.077
              182.0453
              106.0504
              179.0556
              377.1461
              148.061
              182.0817
              166.0868
              179.0556
              241.0317
              132.1025
              150.0589
              122.0276
              133.0613
              133.0137
              132.1025
              87.0082
              147.1134
              90.0555
              191.0192
              116.0712
              74.0242
              134.0453
              180.157
              172.265
              130.142
              118.0868
              118.0868
              136.0623
              137.0463
              152.0572
              267.0729
              155.0093
              111.0195
              385.1294
              175.0355
              241.0317
              106.0504
              122.0276
              265.1123
              175.1195
              123.0558
              ];
lod ngmL = [0.3]
             1.7
             2.8
             3
             3.5
             3.9
             4
             4.8
             6.1
             7.7
```

```
8.1
             10.9
             11.2
             13.6
             15.7
             16.9
             24.8
             25.6
             25.7
             28.4
             32.7
             37.5
             44
             45
             45
             47.4
             48.4
             59
             59.7
             68.9
             74.1
             77
             82.1
            99.2
             112.9
             121.3
             131.7
             133.5
             150.8
             169.2
             214.3
             229.5
             537.3
             10.9
             3.5
             2.8
             28.2
             1.6
             0.8
             48.9
             8.8
             37.1
             52.4
             50
             229.5
             59.7
             37.5
             77
             6.1
            24.8
             3
             ];
[lod mM] = calculateLODs(theo mass, lod ngmL);
```

## **Step 3 define Uptake and Secretion Profiles**

```
exclude_upt = {'EX_gln_L(e)'; 'EX_cys_L(e)'; 'EX_ala_L(e)'; 'EX_mal_L(e)'; 'EX_fol(e)'};
exclude_secr = {'EX_gln_L(e)'; 'EX_cys_L(e)'; 'EX_ala_L(e)'};
```

```
add secr = {'EX mal L(e)'};
add upt = \{\};
essAA_excl = {'EX_his_L(e)'; 'EX_ile_L(e)'; 'EX_leu_L(e)'; 'EX_lys_L(e)'; 'EX_met_L(e)'; 'EX_ret_L(e)'; 'EX_r
tol = 0.05;
data RXNS = {'EX_orn(e)'
                                                            'EX mal L(e)'
                                                            'EX lac L(e)'
                                                            'EX_gly(e)'
                                                            'EX glu L(e)'
                                                           'EX cit(e)'
                                                            'EX 5oxpro(e)'
                                                            'EX 4mop(e)'
                                                           'EX 3mop(e)'
                                                           'EX 3mob(e)'
                                                            'EX_tyr_L(e)'
                                                            'EX trp L(e)'
                                                           'EX thr L(e)'
                                                            'EX pyr(e)'
                                                            'EX_phe_L(e)'
                                                            'EX lys L(e)'
                                                           'EX leu L(e)'
                                                            'EX ile L(e)'
                                                            'EX_glc(e)'
                                                           'EX chol(e)'
                                                            'EX anth(e)'
                                                            'EX val L(e)'
                                                            'EX met L(e)'
                                                            'EX his L(e)'
                                                            'EX gln L(e)'
                                                            'EX_cys_L(e)'
                                                           'EX ala L(e)'
                                                           'EX pi(e)'
                                                            'EX asp L(e)'
                                                            'EX 4HPR0(e)'
                                                           'EX pnto R(e)'
                                                            'EX pro L(e)'
                                                            'EX fol(e)'
                                                           };
```

#### Molt-4

```
input_A = [
% control TP 1 control TP 2 Cond TP 1 Cond TP 2
65245.09667 68680.93 54272.41667 65159.50333
3000 30970.784 20292.406 27226.6555
2038946.433 1917042.967 5654513.467 101768253
163882.9467 186682.92 121762.3567 310547.7
473539.8667 455197.4667 462903.8333 1024508.5
8681.527333 8704.7345 9459.837 34177.945
29168.15 21808.73 120655.9867 2060525.467
3000 3000 34436.50433 113668.5123
3000 3000 25108.829 121927.3673
3000 3000 3000 14717.55667
4142302 4063607.667 3934639.333 3075783.333
2153692 2132723.667 2037735.333 1387754.333
406102.2667 417512.6333 381085.2333 259555.2667
```

```
465074.6 387569.1333 439148.0667 210407.8333
    8087955 8345511.333 8215168.333 5360276
    198435.8 195675.8 188473.1 112386.1667
    20823770.33 20801258.67 19725086.67 15148808
    21229254.67 21225778.33 20799761 17160163
    76555640.67 71459886.33 61697085.33 34981419.33
    876300.4333 905132.5 892182.2 541860.4667
    159124.46 178538.2167 162567.13 3000
    2857012.667 2900419.667 2853523.667 1793173.667
    2995910.333 3018536.333 3024630.333 2266832.333
    69077.16333 67843.12 69406.69 95624.28
    3000 3000 824549.3667 2283200.867
    45304.84667 52977.77333 56566.27667 60759.23
    1613345.1 1258710.1 3430342.067 25970024.1
    216828142.3 221118425 223518663 216863897.3
    632160.0333 612562.3 590881.7333 940705.6
   814465.8333 786011.5667 630513.4 622493.9
   84638.70667 86751.96 89717.10667 68882.68333
    5107317.333 5168599.333 5163708.333 5263614.333
   95419.73667 105904.7067 97550.78667 102678.49
];
```

#### **CCRF-CEM**

```
input B = [
    % control 2 TP 1 control 2 TP 2 Cond 2 TP 1 Cond 2 TP 2
   65245.09667 68680.93 73850.77 98489.89
    3000 30970.784 3000 94181.77233
    2038946.433 1917042.967 5222377.933 134980059.9
    163882.9467 186682.92 219683.7 460476.5267
    473539.8667 455197.4667 437398.3667 630407.2667
    8681.527333 8704.7345 8317.144 86546.77933
    29168.15 21808.73 62146.47333 1012932.38
    3000 3000 9918.992 129433.4973
    3000 3000 7222.259333 145547.7347
    3000 3000 3000 17641.55667
    4142302 4063607.667 4023284.333 3489981.333
    2153692 2132723.667 2068977 1570648
    406102.2667 417512.6333 386495.2 303808.2
    465074.6 387569.1333 376779.1 249036.3333
    8087955 8345511.333 8237784.667 6540301.667
    198435.8 195675.8 196447.1 149861.6667
    20823770.33 20801258.67 21119935.67 16346765.67
    21229254.67 21225778.33 20790535.33 17219085
    76555640.67 71459886.33 65009057.67 24330565.33
    876300.4333 905132.5 884112.5667 259273.9333
    159124.46 178538.2167 158271.14 60631.19333
    2857012.667 2900419.667 2668140 2790196.333
    2995910.333 3018536.333 2890029.333 2538211
    69077.16333 67843.12 74035.24 86165.55
    3000 3000 323185.6667 2063962.067
    45304.84667 52977.77333 62076.23333 64524.22333
    1613345.1 1258710.1 2788313.567 30868376.53
    216828142.3 221118425 212276379 208623151.3
   632160.0333 612562.3 680373.4333 770903.9333
    814465.8333 786011.5667 679862.7 582257.4667
    84638.70667 86751.96 88002.12 99449.36667
    5107317.333 5168599.333 5134219 4445918.333
   95419.73667 105904.7067 100629.24 84807.62333
```

```
];
[cond1_uptake, cond2_uptake, cond1_secretion, cond2_secretion, slope_Ratio] = defineUptakeSecretion
```

MANIPULATE OUTPUT: Add secretion without data points to secretion condition 2.

### **Step 4 Calculate Quantitative Diffs**

```
[cond1_upt_higher, cond2_upt_higher, cond2_secr_higher, cond1_secr_higher, cond1_uptake_LODs,
```

MANIPULATE OUTPUT: Remove the metabolites from the uptake and secretion profiles that you adjusted in the previous steps, e.g. those for which you assume a different directionality as in the data, for metabolites that have inconclusive data (e.g., in case of the anth the metabolite was not detected in the 48 hr samples. It could be assumed that all of it (down to the LOD) was consumed, however in the case of the two cell lines, the relative difference between the cell lines based on the slope ratio (of consumption) would have been 1975% higher in Molt-4 compared the CCRF-CEM cells. In order to prevent that this extreme point distorts the results, these metabolites need t be removed from the input for semi-quantitative adjustment unless such large differences are justified, and make biological sense).

```
remove = {'EX_anth(e)'; 'EX_ile_L(e)'};

A = [];
for i = 1:length(cond2_upt_higher)
    if find(ismember(remove, cond2_upt_higher{i, 1})) > 0
        A = [A; i];
    end
end
cond2_upt_higher(A, :) = [];
```

### Step 5 setQualitativeConstraints

```
cellConc = 2.17 * 1e6;
t = 48;
cellWeight = 3.645e-12;
```

#### Molt-4 model

```
ambiguous_metabolites = {'EX_ala_L(e)'; 'EX_gln_L(e)'; 'EX_cys_L(e)'}; % for ala and gln, we basisMedium = {'EX_o2(e)'; 'EX_strch1(e)'; 'EX_acetone(e)'; 'EX_glc(e)'; 'EX_his_L(e)'; 'EX_caetone(e)'; 'EX_glc(e)'; 'EX_caetone(e)'; 'EX_caeton
```

#### CCRF-CEM model ModelB

```
ambiguous_metabolites = {'EX_ala_L(e)'; 'EX_gln_L(e)'; 'EX_pydxn(e)'; 'EX_cys_L(e)'};
basisMedium = {'EX_ca2(e)'; 'EX_cl(e)'; 'EX_co(e)'; 'EX_fe2(e)'; 'EX_fe3(e)'; 'EX_k(e)'; 'EX_ready.
[model_B] = setQualitativeConstraints(modelMedium, cond2_uptake, cond2_uptake_LODs, cond2_secretary.)
```

### Step 5 setSemiQuantConstraints

This function applies the constraints to the models. It takes two condition specific models into consideration.

```
[modelA_QUANT, modelB_QUANT] = setSemiQuantConstraints(model_A, model_B, cond1_upt_higher, cor
```

## **Step 6 Apply growth constraints**

```
of = 'biomass_reaction2';
tolerance = 20;
```

set constraints on MOLT4 model

```
fprintf('Set contraints on MOLT4 model\n');

Set contraints on MOLT4 model

dT = 19.6;

[model_A_BM] = setConstraintsOnBiomassReaction(modelA_QUANT, of, dT, tolerance);
```

set constraints on CCRF-CEM model

```
dT = 22;
[model_B_BM] = setConstraintsOnBiomassReaction(modelB_QUANT, of, dT, tolerance);
```

# Step 7 integrateGeneExpressionData

```
dataGenes = [535
```

```
1548
              2591
              3037
              4248
              4709
              6522
              7167
              7367
              8399
              23545
              129807
              221823
              ];
[model A GE] = integrateGeneExpressionData(model A BM, dataGenes);
dataGenes = [239]
              443
              535
              1548
              2683
              3037
              4248
              4709
              5232
              6522
              7364
              7367
              8399
              23545
              54363
              66002
              129807
              221823
              ];
[model B GE] = integrateGeneExpressionData(model B BM, dataGenes);
```

### Step 8 extractConditionSpecificModel

```
theshold = 1e-6;
model = model_A_GE;
[model_Molt] = extractConditionSpecificModel(model, theshold);

Starting parallel pool (parpool) using the 'local' profile ...
connected to 12 workers.
```

[model CEM] = extractConditionSpecificModel(model B GE, theshold);

```
[MetConn, RxnLength] = networkTopology(modelMedium);
[MetConnA, RxnLengthA] = networkTopology(model_Molt);
```

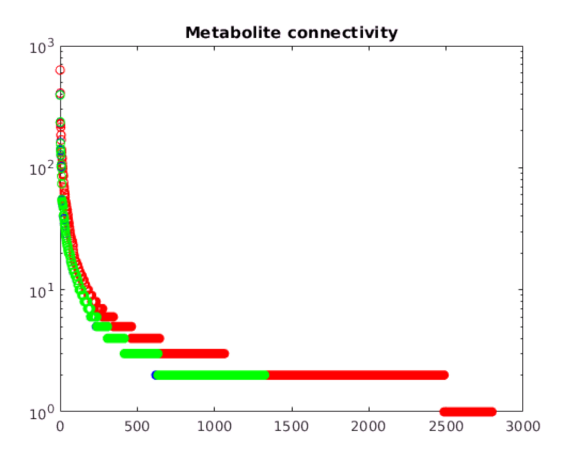
```
[MetConnB, RxnLengthB] = networkTopology(model_CEM);
MetConnCompare = sort(MetConn, 'descend');
MetConnCompareA = sort(MetConnA, 'descend');
MetConnCompareB = sort(MetConnB, 'descend');
```

## Plot metabolite connectivity

```
figure
semilogy(sort(MetConnCompare, 'descend'), 'ro')
hold
```

Current plot held

```
semilogy(sort(MetConnCompareA, 'descend'), 'bo')
semilogy(sort(MetConnCompareB, 'descend'), 'go')
title('Metabolite connectivity')
```



# Step 9 perform sampling analysis

```
fprintf('Perform sampling analysis\n');
```

Perform sampling analysis

```
warmupn = 2000;
nFiles = 10;
pointsPerFile = 1000;
```

```
stepsPerPoint = 500;
fileBaseNo = 0;
maxTime = 3600000;
fileName = 'modelA';
performSampling(model_Molt, warmupn, fileName, nFiles, pointsPerFile, stepsPerPoint,
fileBaseNo

Warning: Need a minimum of 3828 warmup points
Creating warmup points ...
:

fileName = 'modelB';
performSampling(model_CEM, warmupn, fileName, nFiles, pointsPerFile, stepsPerPoint, fileBaseNo

Warning: Need a minimum of 3876 warmup points
Creating warmup points ...
:
:
:

Creating warmup points ...
:
```

### summarize sampling results

```
fonts = 8;
nFiles = 10;
pointsPerFile = 1000;
starting Model = modelMedium;
hist per page = 4;
bin = 30;
modelA = model Molt;
modelB = model CEM;
dataGenes = [32]
              205
              411
              412
              1537
              1608
              1632
              1645
              1737
              1757
              2108
              2184
              2224
              2539
              ];
show rxns = {'PYK'
              'SUCD1m'
              'ATPS4m'
              'ETF'
              };
```

[stats, statsR] = summarizeSamplingResults(modelA, modelB, outputPath, nFiles, pointsPerFile,

```
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