# The nutrition toolbox to generate in silico diets

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## INTRODUCTION

This is a tutorial to show how to use the nutrition toolbox to a) find database equivalents in the USDA [1] or FRIDA [2] databases, b) use the database equivalents to create *in silico* diets, c) how to set *in silico* diet constraints on whole-body metabolic models[3], and d) how to connect to the nutrition algorithm[4] to predict dietary changes. A database equivalent here means a food item in either the USDA or FRIDA database that matches the original dietary food item both on keyword and macronutrient composition.

## **MATERIALS**

- MATLAB.
- COBRA toolbox [5]. For instruction on installing the COBRA toolbox see https://opencobra.github.io/ cobratoolbox/stable/installation.html
- Food items with corresponding macro nutrient information (see template file in the nutrition toolbox paper).
- Diet file with consumed dietary items per diet (see template file in the nutrition toolbox paper).
- Total macrocomposition of diets (optional) (see template file in the nutrition toolbox paper).
- Industrial solver. IBM cplex, MOSEK, gurobi see https://opencobra.github.io/cobratoolbox/stable/installation.html# for solver installation and compatibilities.

## **PROCEDURE**

## Step 1. Prepare your input files

Before we can start using the codebase we need to properly format and fill in the input files.

The nutrition toolbox uses three input files, one of which is optional. The first input file contains the macronutrient composition and associated consumed weight of each food item, referred to as the foodDescription file (you may also use the default values found on the packaging, in which case the weight consumed is 100g or equivalent depending on the information available). An example can be found in the file demoFoodDescription.xlsx. We also provide a partially completed file with the macronutrient information already filled in (demoFoodDescriptionEmpty.xlsx), so you can test filling in keywords. The columns in the files are:

- OriginalFoodName the original name of the food item. This would be the name you, the user, would refer to the food item. It has no influence on the results.
- KeyWords semicolon (;) seperated list of words that describe the food item. Ensure that the first keyword is the most descriptive or important as the code uses that in weighting potential database alternatives. E.g., for the food item "red apple", the most descriptive word is apple, so we shall put this as

the initial keyword. A method to quickly generate keywords is to separate each word in the original food name with a semicolon.

- toExclude semicolon (;) seperated list of key words that exclude results from the database equivalent search.
- databaseID semicolon (;) seperated list of database IDs already found through previous iterations of running the algorithm.
- databaseName semicolon (;) seperated list of the name of the database corresponding to the IDs filled in the databaseID column. This is either usda or frida and nothing else.
- WeightEaten(g) Amount of food consumed (included water weight). It is important to check that the reported macronutrients correspond to this weight eaten. Always given in grams
- Alcohol(g) Amount of alcohol reported on the food label.
- TotalEnergy(kcal) Amount of energy reported on the food label. IMPORTANT given in KCAL.
- TotalFibre(g) Amount of fibre reported on the food label.
- TotalLipid(g) Amount of lipids reported on the food label, including both saturated and unsaturated fatty acids.
- TotalProtein(g) Amount of protein reported on the food label.
- TotalStarch(g) Usually not given on normal food labels, yet included here in case this information is available to the user, e.g., through consulting other sources.
- TotalSugar(g) Amount of sugar reported on the food label. Usually it will say Carbohydrates -> of which sugars.
- TotalWater(g) Amount of water in the food, usually not given on normal food labels.

Notably, total carbohydrates are excluded from the comparison due to inconsistency in definitions across databases and on food labels of dietary food items, specifically whether fibre is included in the total carbohydrate count. Instead, individual carbohydrate components (i.e., sugars, fibre, and starch) are used where available.

If information for a macronutrient column is not available, leave the cell blank.

If the label of your food is given as a volume instead of weight (e.g., /100mL) you will need to convert the volume into weight, e.g., by measuring how many grams the 100mL correspond to, or converting the volume into to grams by using the food item density (if available).

If a food item is consumed in different quantaties across different diets, you only have to enter it once foodDescriptions file. IMPORTANT is that reported macronutrients correspond to the consumed weight (includes water weight)

The weights consumed are used to to adjust the macronutrients from database equivalents to the same weight consumption as the macronutrients for database equivalents are all given per 100g of food consumed.

Users should double-check that the consumed macros correspond the consumed weight of the food item. I.e., make sure that if 80g of food is consumed, not the standard 100g of food on food labels is input in the various macronutrient columns.

**The second input file** contains the consumed weight of food items per diet which will be used to create *in silico* diets, referred to here as the foodsConsumed file. In the foodsConsumed file we will fill in the

database equivalents found with the foodDescription file. If a food item is not consumed in a diet, set the weight consumed to 0. A worked example can be found in demoFoodConsumed.xlsx, an example without the database alternatives can be found in demoFoodConsumedEmpty.xlsx. The following columns can be filled in:

- originalName The original food name. This is usefull to double check that the consumed weights per diet are correct and the correct database alternatives are put in place.
- foodName The name of the database alternative, serves as a control to make sure it matches with the original food.
- databaseID The ID of the database alternative this will be given in the output of step 2 when we find and compare database alternatives.
- databaseUsed The name of the database where the database alternative is found this will be given in the output of step 2 when we find and compare database alternatives.

**The third input file** contains the macronutrient composition of the full diets and can found in demoOriginalMacroDiet. Each column is a diet with the rows being the relevant macronutrients.

Then we initialise the COBRA toolbox and set our industrial solver

```
% Initialise the toolbox
initCobraToolbox();
                                                      COnstraint-Based Reconstruction and Analysis
                                                      The COBRA Toolbox - 2025
                                                      Documentation:
                                                      http://opencobra.github.io/cobratoolbox
> Checking if git is installed ... Done (version: 2.37.3).
\gt 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 (this may take a while)... Done.
> Adding all the files of The COBRA Toolbox ... Done.
> Define CB map output... set to svg.
> TranslateSBML is installed and working properly.
> Configuring solver environment variables ...
   - [---*] ILOG CPLEX PATH: C:\Program Files\ibm\ILOG\CPLEX Studio1210\cplex\matlab\x64 win64
   - [*---] GUROBI PATH: C:\gurobi1201\win64\matlab
   - [*---] TOMLAB PATH: --> set this path manually after installing the solver ( see instructions )
   - [*---] MOSEK_PATH: --> set this path manually after installing the solver ( see instructions )
  Done.
> Checking available solvers and solver interfaces ... Could not find installation of dqqMinos, so it cannot be tes
Could not find installation of matlab, so it cannot be tested
Could not find installation of mosek, so it cannot be tested
Could not find installation of quadMinos, so it cannot be tested
Could not find installation of tomlab_snopt, so it cannot be tested
Done.
> Setting default solvers ...Could not find installation of mosek, so it cannot be tested
Could not find installation of mosek, so it cannot be tested
Could not find installation of matlab, so it cannot be tested
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	Q	P	MIQP	NLP	EP	CLP
dqqMinos	active		0 0	-	0	-	-	_	-
glpk	active		1	1	-	-	-	-	-
gurobi	active		1	1	1	1	-	-	-
lp_solve	legacy		1	-	-	-	-	-	-
matlab	active		0	-	-	-	0	-	-
mosek	active		0	-	0	-	-	0	0
pdco	active		1	-	1	-	-	1	-
qpng	passive		-	-	1	-	-	-	-
quadMinos	active		0	-	-	-	-	-	-
tomlab_snopt	passive		-	-	-	-	0	-	-
Total	-		4	2	3	1	0	1	0

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

```
> You can solve LP problems using: 'gurobi' - 'glpk' - 'pdco'
> You can solve MILP problems using: 'gurobi' - 'glpk'
> You can solve QP problems using: 'gurobi' - 'pdco'
> You can solve MIQP problems using: 'gurobi'
> You can solve NLP problems using:
> You can solve EP problems using: 'pdco'
> You can solve CLP problems using:
> Checking for available updates ...
> There are 132 new commit(s) on <master> and 28 new commit(s) on <develop> [b75356 @ develop]
> You can update The COBRA Toolbox by running updateCobraToolbox() (from within MATLAB).
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\componentContribution\new
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\groupContribution\new
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\inchi\new
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\molFiles\new
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\protons\new
removing: C:\Users\MSPG\Documents\cobratoolbox\src\analysis\thermo\trainingModel\new
```

```
% Change the solver, can be gurobi, ibm_cplex, tomlab_cplex, or mosek
changeCobraSolver('gurobi');
```

> changeCobraSolver: Gurobi interface added to MATLAB path.

## Step 2: Finding database alternatives

The first step in making an *in silico* diet is to find alternatives in databases that have metabolite compositions measured. Here we use both the the U.S. Department of Agriculture, FoodData Central (USDA) database and the National Food Institute, Technical University of Denmark FRIDA (FRIDA) database. The function we use to find database alternatives is *vmhFoodFinder.m* and is available in the COBRA toolbox. The function takes the following inputs:

- templateFilePath **Required** The full file path to where the input file is stored. IMPORTANT! remember to include a file extension. The demo file for this step is the demoFoodDescription.xlsx file.
- searchType Optional Method of searching keywords in the food database. Either sequential or cumulative. Explanation about the difference is given below.

- addStarch **Optional** Boolean, indicates if additional starch should be added based on the reported macros. It will add on additional starch based on the equation starch = total carbohydrates fibre total sugars, as starch is often not measured in the USDA database. Defaults to false.
- databaseType Optional Character, which database should be used either 'usda' for USDA FoodData database or 'frida' for the Danish food institute database. Can be 'mixed' if both databases should be used. Defaults to mixed.
- maxItems **Optional** Numeric, value indicating the maximum amount of VMH food alternatives to be analysed for macros. Defaults to 50.
- outputDir **Optional** Path to where the output file will be stored. Defaults to [pwd filesep 'NT Result']
- foodSources2Use **Optional** Cell of strings, dictates which food sources from the USDA database will be used to find food items. Defaults to {'sr\_legacy\_food'; 'foundation\_food'; 'survey\_fndds\_food'};

There are two search methods employed to find potential database equivalents: sequential and cumulative

In the sequential search method, the first keyword is used to identify database equivalents containing that keyword, resulting in sub-selection 1. The second keyword is then used to filter sub-selection 1, retaining only those entries that contain the second keyword, thereby producing sub-selection 2. This process continues until all keywords have been used. If the final sub-selection is empty, the results from the second to last sub-selection are used, if applicable.

In the cumulative search method, each keyword independently generates a sub-selection of database equivalents that contain the respective keyword. All resulting database equivalents are then tallied. Database equivalents with a total count of total number of keywords -1 or more are returned. If no database equivalents reach that threshold, the results for total number of keywords -2 matches are returned.

First we will define some of the input arguments

```
% Path to the input file, change to where the file is on your computer
templateFilePath = 'C:\nutritionToolbox\demoFoodDescription.xlsx';

% Path to where we want our results stored so we can easily find them,
% change to an existing path on your computer
outputDir = 'C:\nutritionToolbox\results';
```

If we want to adjust more input arguments we can define them by pressing enter to add additional lines after line 5, and copy and pasting the input argument we want to alter. Add an equal sign (=) and the new value and end with a semicolon. E.g., if you want to decrease the number of items to be compared, you would put maxItems = 30; underneath line 5.

To call the function (Note for the demo it will take likely between 10 to 20 minutes for the initial run):

```
vmhFoodFinder(templateFilePath, 'outputDir', outputDir);
Warning: The following USDA food ID does not seem to have a KCAL assocatiated with it. Please take note:321505
Warning: The following USDA food ID does not seem to have a KCAL assocatiated with it. Please take note:2685568
Warning: The following USDA food ID does not seem to have a KCAL assocatiated with it. Please take note:321505
```

If you defined multiple input arguments, add them after outputdir as 'name as defined in the list above' and the variable name where the new value is stored or the value directly. E.g., if you want to adjust maxItems, you can either set ,'maxItems', maxItems if you defined maxItems before OR set ,'maxItems', 30 behind outputdir. The resulting input would look like vmhFoodFinder(templateFilePath, 'outputDir', outputDir', outputDir, 'maxItems, maxItems); OR vmhFoodFinder(templateFilePath, 'outputDir', outputDir, maxItems, 30);

The outputs will be stored in a folder called NT\_results in the defined directory set in the outputDir variable. If left empty, it will be made in your MATLAB current working directory.

Four output files will be stored in the NT results folder

- fullComparisonFoodItems.xlsx The file contains the full comparison for each food item. Food items that had no, or too many, database alternatives have also been indicated. The file can be used to select database alternatives to create *in silico* diets with.
- tooManyDatabaseHits.xlsx This file contains all the food items that had more than the maximum allowed database alternatives to be compared (set in the maxItems input variable). The columns contain the food items and the rows contain every database alternative found for that food item. The file can be used to refine the key words used to describe food items, or to populate the toExclude column in the input file.
- noDatabaseHits.txt A list of food items for which no database alternatives were found. For these items, check the spelling of key words, alternative names, or reduce the amount of key words. If that does not help, try to find a database alternative on VMH.life.
- topResultsComparisonFoodItems.xlsx Similar to fullComparisonFoodItems.xlsx, containing only the top 10 results for a more concise overview of the results.

Either the fullComparisonFoodItemsn or topResutIsComparisonFoodItems can be used to select database alternatives. The code compares the macros provided by the user for each food item and compares them to the given/measured macros from the database alternatives (so-called label macros) and then uses Euclidean distance to evaluate how similar the two are. The Euclidean distance is given in the SimilarityScore column of these two files. Besides the Euclidean distance, the percentage of the macronutrients measured as metabolites is also reported. The measured metabolites are grouped into macronutrient groups and give an indication of how well the macronutrients are represented in measured metabolite content. Database equivalents less similar to dietary food items may be chosen if they have a higher measured metabolite representation in their macronutrients. By choosing the database equivalents that may be less similar to the dietary food item but with a higher measured metabolite representation, it increases the metabolic content of the *in silico* diets and give more overall information as opposed to choosing database alternatives with lower metabolite representation that might be more similar to the dietary food item.

For example, if we were to compare two database equivalents: "Dates, medjool" and "Date", both from the USDA database, the reported macronutrient values are near identical. However, "Dates, medjool" has higher percentages of macronutrients covered by measured metabolites than "Date", making it a better choice due to its greater metabolite coverage.

It is very likely that you will need to perform 2 or 3 runs with updated keywords to obtain suitable database equivalents for each dietary food item. To reduce time, it is recommended to fill in the column databaseID and databaseName for the dietary food items for which appropriate database equivalents have been identified. To improve keyword performance in subsequent runs, users may consider changing the number of keywords, exploring alternative spellings, or naming conventions. Additionally, looking at the tooManyDatabaseHits may help finding keywords for the toExclude column. Alternatively, you can increase the max items compared, set by putting ,'maxItems', x in the function call (where x is the new number of max items compared).

## Step 3: Generating in silico diets

With step 1 and step 2 performed, we can now prepare the second input file, referred to as "demoCreateDiet" here. In step 1, we already filled in the original food items and the consumed weight per diet in the "demoCreateDiet" file. Now we can fill in the database id and database name for each dietary food iem. The two fields can be directly copied from fullComparisonFoodItemsn or topResutlsComparisonFoodItems where the relevant database alternatives were chosen. Once completed, we can use the function *generateInSilicoDiet.m* to create our *in silico* diets and do some quick analyses. The function takes the following inputs:

- toCreateDiet **Required**: Path to the file with the diets that need to be created. consist of column "originalName" where the original food names are set. "databaseID" the ID of the fooditem in their database. "databaseUsed" which database was used to find that database alternative. Each column after is a diet where the values in gram show the food items consumed for each diet. The demo file for this is called 'demoCreateDiet.xlsx'.
- outputDir **Optional**: Path to the directory where the results should be stored. Defaults to current working directory.
- originalDietMacros **Optional**: Path to the file with the total macros for the original diets. Should contain the rows lipids, carbohydrate, protein and energy. Defaults to an empty string, i.e., not provided. Template how the file should look like can be found with the name 'demoOriginalMacroDiet.xlsx'.
- analyseMacros **Optional**: Boolean, indicates if analysis on macros should be performed. Defaults to true.
- addStarch **Optional**: Boolean, indicates if additional starch should be added based on the reported macros. It will add on additional starch based on the equation starch = total carbohydrates fibre total sugars, as starch is often not measured in the USDA database. Defaults to false.

#### First let us define our variables

```
% The path to your diet file. The demo is called demoCreateDiet.xlsx.
% Change to the path where the file is on your computer
toCreateDiet = 'C:nutritionToolbox\demoCreateDiet.xlsx';

% We initialised outputDir in step 2 already. If you want to store it in a
% different directory, remove the % from the code below and put in the
% desired location.
% outputDir = '';

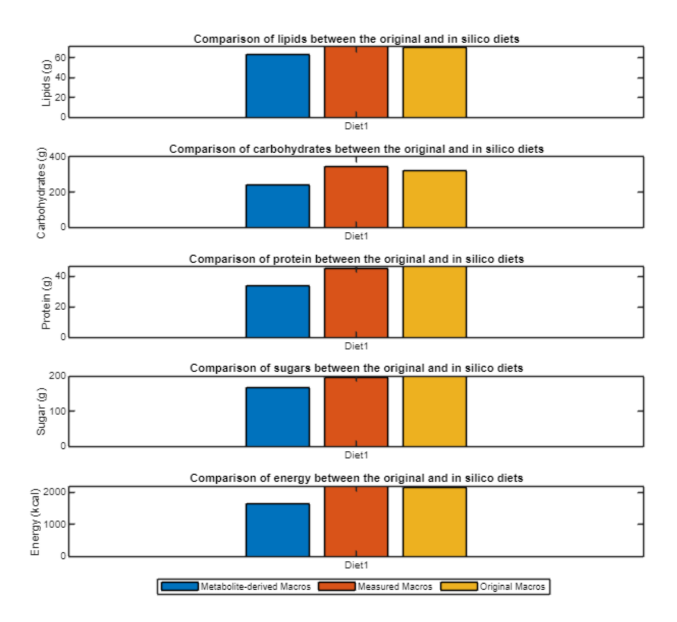
% If you have the macros for the original diets fill the
% demoOriginalMacroDiet.xlsx file and give the path here. If you dont have
```

```
% it leave it as is. Change to the path where the file is on your computer.
originalDietMacros = 'C:\nutritionToolbox\demoOriginalMacroDiet.xlsx';

% Put to false if you do not want analysis on the macros
analyseMacros = true;

% Set to true if you want to add on additional starch for USDA based
% database alternatives
addStarch = false;

% Now we can the function
[dietFlux] = generateInSilicoDiet(toCreateDiet, 'outputDir', outputDir, ...
    'originalDietMacros', originalDietMacros, ...
    'analyseMacros', analyseMacros, 'addStarch', addStarch);
```



The figure generated by the code will show you how well various macro-nutrients match the original diet. In blue are the macronutrients calculated from measured metabolites, in orange the macronutrients as measured from the database equivalents and in yellow the reported macronutrients from the original diet. From the demo we can see that the database equivalents and the original diet match well in macronutrient composition, with only a slight overestimation of lipid content. We see that the macronutrients calculated from measured metabolites (blue bar) is lower. This is because the metabolite content for every database equivalent is not perfectly represented and some metabolites might be missed or not measured. There are also measurement errors that could play a small role in the deviations we see.

We can look at the first 10 rows of dietFlux and see that indeed the dietary metabolites are returend for the diet that we created. It is normal that not all dietary metabolites have an asocciated flux value as not every metabolite is a) measured or b) found in every food item / database. The flux values are given in mmol/human/day

## dietFlux(1:10,:)

ans =  $10 \times 2$  table

	VMHID	Diet1
1	'Diet_EX_25hvitd3[d]'	"0"
2	'Diet_EX_4hpro_LT[d]'	"0"
3	'Diet_EX_M01582[d]'	"0"
4	'Diet_EX_M02053[d]'	"0"
5	'Diet_EX_M03045[d]'	"0"
6	'Diet_EX_M03051[d]'	"0"
7	'Diet_EX_ala_L[d]'	"18.62491657"
8	'Diet_EX_arach[d]'	"0.5792149427"
9	'Diet_EX_arachd[d]'	"0"
10	'Diet_EX_arg_L[d]'	"11.66127926"

All results are stored in the **inSilicoDiet** folder created in the output directory defined at the start with the following files:

- fluxDiets.csv This file contains the flux values for all the dietary components for each diet.
- macrosCalculatedFromMetabolites.csv The macronutrients as calculated from the measured metabolites for each diet.
- macrosAsReported.csv The macronutrients as reported for all the food items that make up each diet.
- macroComparison.png The figure made output by MATLAB, showing the comparison for each diet between the various macronutrients and how well the metabolite macronutrients align with the reported/ label macronutrients and, if provided, the original diet macronutrients.
- energyFractionPerMacro.csv For the lipids, carbohydrates and the proteins the fraction of their contribution to the total energy of the diet. Calculated for the metabolite macronutrients, reported/label macronutrients and, if provided, the original macronutrients.

These files can be used to gain a deeper understanding of diet. The total metabolite macronutrients (the macronutrients calculated from measured metabolites) compared to the actual measured macronutrients of the *in silico* diets give an understanding on how well the overall macronutrients are represented by measured metabolites. Comparing the measured macronutrients of the *in silico* diets and the total macronutrients from the original diet will give an insight as to how well the *in silico* diet matches the original diet.

The file fluxDiets.csv or the output fluxDiets can be used to constrain WBMs with. However we can also use the function *setInSilicoDiet.m* to easily set the diet with all the correct checks and save them. The function can also check if the WBMs are feasible on the given diet and add on required micronutrients where required. Besides setting the diet altering the dietary exchange reactions, we can also have the WBMs "consume" the food items themselves. In order to do that the toolbox adds in novel reactions to allow for food intake and food breakdown. The food intake is straightforward, as it will be an exchange reaction of the food item (now treated as a metabolite) as in the novel food ([f]) compartment

## foodItem[f] <-

This allows for food items to enter the WBMs. Then we to breakdown the food item into its metabolite components.

Where x is the number of metabolites measured for the food item. We also incorporate macronutrients such as energy and total lipids so we can track them in the diet. The macronutrients here are treated as metabolites. In order to comply with mass balance rules, we need to remove the macronutrients we are treating as metabolites from the system. We do this by setting another exchange reaction.

## macronutrient[d] ->

With these new reactions in place the WBMs can consume and breakdown food items into their metabolite components and we can track the total amount of various macronutrients in the diet. Important to note is that the food exchange, breakdown, and macronutrient exchange reactions are in g/person/day (kcal/person/day for energy consumption) instead of the usual (mmol/person/day). You can choose either or both options to constrain your WBMs with in setInSilicoDiet.m but we advise to only use the food item-constrained WBMs in conjuction with the nutrition algorithm or to do simulation on diet composition. The additional food item reaction can make the vastly increase the reaction content of the WBMs (by over 30.000 reactions) and make the WBM overly complex for questions that do not involve the optimisation of the actual food items. To study the effect of diet on the various reactions in the human body - we suggest to just use metabolite-constrained WBMs. If no metadata is provided, the function will create both a female and a male WBM for each diet. If metadata is provided the diets will be matched to the corresponding sex-specific WBMs. You also have the possibility to give your own WBMs if you do not want to use the standard Harvey/Harvetta (e.g., community microbiome WBMs). If you have sample specific WBMs, ensure that the sample ID is in the file name of your WBM flanked by underscores. E.g., WBM sample1 female.mat. If you have altered WBMs that are not sample specific, i.e., one altered male WBM and one altered female WBM, ensure the filename include " male" and " female", respectrively. I.e., WBM altered male.mat or WBM altered female.mat.

The *setInSilicoDiet.m* function takes the following inputs:

- dietToSet Required Path to the file with the diets that need to be created. Consists of column
  "originalName" where the original food names are set. "databaseID" the ID of the fooditem in a database.
  "databaseUsed" which database was used to find that database alternative. Each column after is a diet
  where the values in gram show the food items consumed for each diet. The demo file for this is called
  'demoCreateDiet.xlsx'.
- metadataPath **Optional** Path to the metadata file, should contain the columns ID, sex and diet. Defaults to an empty string, i.e., not provided.
- outputDir **Optional** Path to the directory where the results should be stored. Defaults to an empty string, i.e., not provided.
- constrainFoodWBM **Optional** Boolean, indicates if the code should alter WBMs to 'consume' food items and set the diet as food items instead of dietary fluxes. Defaults to false.
- constrainFluxWBM **Optional** Boolean, indicates if WBM should be constrained with the dietary exchange fluxes. Defaults to true.
- checkFeasibility **Optional** Boolean, indicates if the newly constrained WBMs should be tested for feasibility. If they are not feasible, dietary exchange reactions are added to try and make them feasible. Defaults to true.
- wbmVersion **Optional** If a specific version of the WBMs should be used. Do not start with \_. Defaults to an empty string, i.e., not provided.
- pathToWBMs Optional If non-default WBMs are to be used. Default WBMs are unchanged Harvetta/
  Harvey. Path to where the WBMs are stored that need to be constrained. If you have sample
  specific WBMs, ensure that the sample ID is in the file name of your WBM flanked by underscores.
  E.g., WBM\_sample1\_female.mat. If you have altered WBMs that are not sample specific, i.e., one
  altered male WBM and one altered female WBM, ensure the filename include "\_male" and "\_female",
  respectrively. I.e., WBM altered male.mat or WBM altered female.mat.

Let us now define the variables. Setting more diets and setting diets on more models will increase the time of this step. For the demo this step should take ~5 minutes

```
% This is the same as toCreateDiet set previously. Change to where the file
% is on your computer.
dietToSet = 'C:\nutritionToolbox\demoCreateDiet.xlsx';

% The path to the metadata file if available. Otherwise standard female and
% male WBMs will be constrained for each diet.
metadataPath = '';

% We initialised outputDir in step 2 already. If you want to store it in a
% different directory, remove the % from the code below and put in the
% desired location.
% outputDir = '';

% Change to false if you do not want the WBMs to be constrained with food items
constrainFoodWBM = true;

% Change to false if you want to WBMs to be constrained with dietary flux
% values
```

```
constrainFluxWBM = true;
% Change to false if you do not want to check if WBMs are feasible on the
% constrained diets.
checkFeasibility = true;
% Change if you want to use a specific version of the WBMs, otherwise it
% will take the latest version available. If you want to use a specific
% version do not start with
% The results in this tutorial were made with wbmVersion 1 03d in the COBRA
st toolbox and for testing to see if the same results are achieved, we
% recommend using this version.
wbmVersion = '1 03d';
% Change if non-default WBMs are to be used. Path to where the WBMs are stored
% that need to be constrained. If general models, use female or male
% to indicate the sex of the generic model. Otherwise ensure that the
% sample ID is flanked by underscores (e.g., _sample1_).
pathToWBMs = '';
% Set all the inputs and run the code
setInSilicoDiet(dietToSet, 'metadataPath', metadataPath, 'outputDir', outputDir, ...
    'constrainFoodWBM', constrainFoodWBM, 'constrainFluxWBM', constrainFluxWBM, ...
    'checkFeasibility', checkFeasibility, 'wbmVersion', wbmVersion, ...
    'pathToWBMs', pathToWBMs);
Setting dietary flux vector on WBMs for Diet1
Setting food items on WBM for Diet1
> changeCobraSolver: Gurobi interface added to MATLAB path.
EnsureGrowthOnDiet -- STEP 1: Test which models are feasible on the given diet
Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model
Warning: Reaction Excretion EX microbiota LI biomass[fe] not in model
Model Diet1 fWBM female is feasible on the given diet
Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model
```

```
EnsureGrowthOnDiet -- STEP 1: Test which models are feasible on the given diet Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Model Diet1_fWBM_female is feasible on the given diet Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Model Diet1_fWBM_male is feasible on the given diet All models are feasible on the diet. No dietary alterations are needed.

> changeCobraSolver: Gurobi interface added to MATLAB path.
EnsureGrowthOnDiet -- STEP 1: Test which models are feasible on the given diet Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Model Diet1_WBM_female is feasible on the given diet Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Warning: Reaction Excretion_EX_microbiota_LI_biomass[fe] not in model Model Diet1_WBM_male is feasible on the given diet
All models are feasible on the diet. No dietary alterations are needed.
```

New directories will be created in the chosen output directory. One is called fluxDietWBMs, if you chose to constrain WBMs with dietary flux reactions, and one called foodDietWBMs, if you chose to constrain with food items. In each folder, the various diet-constrained WBMs are stored. If the feasibilities were checked, an Excel file named dietGrowthStats.xlsx is created ONLY if any of the WBMs are infeasible on the given diet. The dietGrowthStats file contains WBM feasibility on a rich diet (all dietary metabolites available) and on

the constrained diet, as well as the feasibility on the constrained diet with adjustments made try and make infeasibile WBMs feasible. The feasible WBMs can now be used for modelling purposes. For an indepth tutorial on how to solve WBMs, we refer you to tutorial 2 of Persephone [6]. Here we will further demonstrate how to use WBMs in combination with the nutrition algorithm to predict changes in the diet to achieve desired adjustments in human metabolism.

## Step 4: Using the nutrition algorithm

We can use either the food item-constrained WBMs or flux-constrained WBMs with the nutrition algorithm to ask for the minimum changes in diet to achieve to biggest change in chosen reactions in human metabolism. Here we will show three scenarios on how to work with the nutrtion algorithm with food-constrained WBM. The same principles apply for flux-constrained WBMs.

The nutrition algorithm has 6 inputs:

- model A WBM model
- obj the objective function. Usually this is 'Whole\_body\_objective\_rxn' for WBMs or 'Excretion EX microbiota LI biomass[fe]' for community microbiome WBMs.
- objMinMax if the objective function has to be minimised or maximised. Use either 'min' or 'max', respectively.
- rois Reactions of interest, a cell array. A list of reactions or metabolites that you want to minimise or
  maximise and which will be the main drivers to modulate changes in the diet. Metabolites, with their
  corresponding compartments, will be created as demand reactions. It is important to note that we can at
  this time only maximimse demand reactions. If existing reactions are picked, double-check the reaction
  directionality with printRxnFormula.m as it could influence the decision to pick whether to minimise or
  maximise the reaction.
- roisMinMax A cell array of the same size as rois. For every reaction of interest chosen, give 'max' or 'min' to maximise or minimise, respectively.
- options A structure, where every field has an option that modulates how the nutrition algorithm works. We will go over the most relevant options in the various scenarios. For a full list of option please refer to the nutritionAlgorithmWBM.m function description.

### And outputs the following:

- newDietModel A WBM model, with the updated dietary constraints calculated by the algorithm.
- pointsModel The adjusted model used to arrive to the dietary solution.
- roiFlux A structure with the solution structures for the original and adjusted diet reaction optimisations of interest .
- pointsModelSIn The solution structure for the optimisation that found the dietary changes.
- menuChanges A table with the changes in diet food items or dietary metabolites as printed on the terminal.
- macroChanges A table with the original and new macronutrient distrubutions and the relative changes between the two.

• roiChanges - A table with the range of the fluxes carried by the reactions of interest before and after dietary changes, as printed in the terminal during the run.

We shall use the food-constrained model created in step 3. Let us assume that we want to find dietary alterations that will result in an a) increase of glucose the in blood, b) increase dopamine in the cerebrospinal fluid, and c) minimise the total lipid intake.

If we only set these constraints on the system, we will get unrealistic results as we have not set any bounds on how much food is allowed to be removed or added ,or what the allowed ranges in macronutrient composition are. Also, it may happen that the nutrition algorithm will output unrealistic dietary considerations without setting further details (e.g., to optimise dopamine levels, a recommendation may be to supplement a diet by 9kg of egg whites). Therefore, to ensure obtaining realistic dietary changes, we shall use the (optional) "options" variable to constrain our system. There are many options we can set, but the most relevant for us are:

- options.foodOrMets indicates if the dietary changes will be calculated through food items or dietary metabolites. Accepted inputs are 'Food', 'allMets', and 'dietMets'. Food will enable food item calculations. allMets will enable calculation of all dietary metabolites. dietMets will enable calculation of metabolites only found in the the given diets. The default is set to 'Food'.
- options.roiWeights numeric array. Indicates the 'importance' of a reaction of interest. The higher the value, the more the dietary solution will be focused on the minimisation or maxisimation of that particular reactions. By default all reactions of interest are weighted with 10.
- options.caloricRange a 1x2 numeric array. Specifies what the minimum and maximum range of the consumed calories are allowed to be. It is important to know what the original caloric content of the diet is. If calories of the current diet do not fall within the range you set here, the dietary changes will be altered to fit inside that range. Defaults to [0 1e6].
- options.carbohydrateRange a 1x2 numeric array. Specifies what the minimum and maximum range of the consumed carbohydrates are allowed to be. It is important to know what the original carbohydrate content of the diet is. If the carbohydrates of the current diet do not fall within the range you set here, the dietary changes will be altered to fit inside that range. Defaults to [0 1e6].
- options.lipidRange a 1x2 numeric array. Specifies what the minimum and maximum range of the consumed lipids are allowed to be. It is important to know what the original lipid content of the diet is. If the lipids of the current diet do not fall within the range you set here, the dietary changes will be altered to fit inside that range. Defaults to [0 1e6].
- options.proteinRange a 1x2 numeric array. Specifies what the minimum and maximum range of the consumed proteins are allowed to be. It is important to know what the original protein content of the diet is. If the proteins of the current diet do not fall within the range you set here, the dietary changes will be altered to fit inside that range. Defaults to [0 1e6].
- options.sugarsRange a 1x2 numeric array. Specifies what the minimum and maximum range of the consumed sugars are allowed to be. It is important to know what the original sugar content of the diet is. If the sugars of the current diet do not fall within the range you set here, the dietary changes will be altered to fit inside that range. Important to note is that the lower bound set for sugars CANNOT be higher than the upper bound of the carboyhydrate range. Defaults to [0 1e6].

- options.foodAddedLimit A numeric value, specifies how many units of an item (metabolite or food) are allowed to be added to the diet. For food items the unit is in grams, for metabolites it is in mmol. Defaults to 1e6.
- options.foodRemovedLimit A numeric value, specifies how many units of an item (metabolite or food) are allowed to be removed from the diet. For food items the unit is in grams, for metabolites it is in mmol. Defaults to 1e6.
- options.removeFoodItem: A nx2 cell array with the first column being the food item ID and the second column the name of the database of the food items that shall be removed from the analysis.
- options.addPrice: a nx3 cell array with the first column being the food item ID, the second column the name of the database, and the third column the price (any currency) associated with that food item.

  Allows for optimising the price of a diet, but only if enough information is available.

These constraints, especially the macronutrient ranges, are only interesting when looking at food item-constrained WBMs as metabolite dietary changes will not take those into account. In theory, it is be possible to set the macronutrient range contraints on flux-constrained WBMs where you calculate additional food items. However, the ranges have to be lower as you can only calculate the added macronutrients from the new food items.

Now that we know the various inputs and constraints we need to take into account let us put it into practice.

```
% Load in a food item-constrained WBM. Change to where the file is on your
% computer.
model = load('C:\nutritionToolbox\foodDietWBMs\Diet1_fWBM_male.mat');
```

**IMPORTANT**: To get accurate results, it is imperative that you ensure that the lower and upper bounds of the dietary reactions or food exchange reactions, whichever you use, are the same. If the upper bound is set to 0 or to a higher amount than the lower bound (they are all negative values), the correct macronutrient composition or metabolite composition will not be captured by the model as it can choose to consume less. That will lead to results that only add new foods or metabolites without telling you that it actually consumed less of a particular item. To easily set the upper and lower bounds to the same value use:

for food item-constrained WBMs

```
model.ub(contains(model.rxns, 'Food_EX_')) = model.lb(contains(model.rxns,
'Food_EX_'));
```

for flux-constrained WBMs

```
% Find all the dietary exchange reactions
dietaryMetExchanges = model.rxns(contains(model.rxns, 'Diet_EX_'));

% Remove the dietary exchange of macronutrients as we want those to be
% opened fully
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_energy[d]')) = [];
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_carbohydrate[d]')) = [];
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_protein[d]')) = [];
```

```
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_lipid[d]')) = [];
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_sugars[d]')) = [];
dietaryMetExchanges(strcmp(dietaryMetExchanges, 'Diet_EX_money[d]')) = [];

% Find the index in the model for the remaining dietary exchange reactions
rxnIdx = findRxnIDs(model, dietaryMetExchanges);

% Set the lower and upper bound to the same value
model.ub(rxnIdx) = model.lb(rxnIdx);
```

Then we cann set the remaining variables

```
% Set the objective function to the whole body objective reaction. It does
% not matter if we maxisimise or minimise as the reaction is constrained to
% [1,1].
obj = 'Whole_body_objective_rxn';
objMinMax = 'max';

% Set the reactions of interest and if they have to minimised or maximised
rois = {'glc_D[bc]', 'dopa[csf]', 'Diet_EX_lipid[d]'};
roisMinMax = {'max', 'max', 'min'};

% We want to calculate food item dietary changes
options.foodOrMets = 'Food';

% We will not weight our reaction of interest more then our dietary
% reactions
options.roiWeights=1*ones(1,length(rois));
```

First we will show what happens when we do not use any additional dietary constraints on the system.

[newDietModel,pointsModel,roiFlux,pointsModelSln,menuChanges, macroChanges,
roiChanges] = nutritionAlgorithmWBM(model,obj,objMinMax,rois,roisMinMax,options);

```
Reaction of Interest 1 = DM_glc_D[bc]
Reaction of Interest 2 = DM_dopa[csf]
Reaction of Interest 3 = Diet_EX_lipid[d]
Solution points =-4049.1245
850.5981 come from diet
-4899.7226 come from roi
Food items of interest are:
```

foodNames	status	Flux	food
{["Alcoholic beverage, distilled, all (gin, rum, vodka, whiskey) 100 proof"]} {["Cherries, sweet, dark red, raw"	{'Added'} {'Added'} {'Added'} {'Added'} {'Added'} {'Added'} {'Added'}	-0.39132 -36.153 -187.36 -451.72 -21.706 -11.728 -141.53	{'1736 {'2346 {'225 {'1353 {'1396 {'1418

Diet Energy = 4790.3713

```
f1 =1 & f2=1
DM_glc_D[bc]
Original Diet RoI range = 0:1328.4361
New Diet RoI range = 0:4987.4572
DM_dopa[csf]
Original Diet RoI range = 0:10.7923
New Diet RoI range = 0:104.0193
Diet_EX_lipid[d]
Original Diet RoI range = 71.282:71.282
New Diet RoI range = 104.5768:105.2495
```

As we can see from the printed output, the food items that were added were: Alcoholic beverages, cherries, egg whites, sorbitol, cod liver oil and linseed oil and dried lactose powder. The total amount of food added to the diet is ~850g, which on top of an already normal diet is quite some extra food. When we look further in the dietary composition, 180g of egg white and 141g of dried lactose powder does not seem appetising to eat and difficult to incorporate in an already existing diet. This further asks for additional bounds on food intake and macro-nutrient composition. Interestingly, we see an increase of almost 20g of lipids in the diet, while we asked for the lipid conusmption to be as small as possible. This is likely because we asked for the maximal storage of glucose in the blood. That means that the points generated by adding more glucose in the blood outweigh the points generated to by limiting lipid consumption. This shows to highlight that various reactions of interest can counteract each other as long if one reaction can generate vastly more points than other reactions. A way to counteract this is by adjusting the reaction weights which we show in a section further down.

We do see that the maximum storage of glucose is blood is increases with ~3000 mmol/person/day, the dopamine in the cerebral spine fluuid with ~95 mmol/person/day and consumed lipids with 33 g/person/day. The changes in diet, macronutrient composition, and reaction of interest fluxes can be found in the output variables dietChanges, macroChanges, and roiChanges, respectively.

To obtain a more palatable dietary change we will now add additional constraints to food consumed and macronutrient composition.

From the diet analysis performed in step 3, we see that we have ~345g of carbohydrates, ~45g of proteins, ~70g of lipids, ~195g of sugar and ~2200 kcal (found in the macrosAsReported.csv table). Depending on your modelling objectives, you can use these values as a starting point to set constraints. If you want the diet to remain in a similar range of macronutrient composition, we suggest to have the constraints 10-20% of the original value to allow for some leeway. Otherwise, if you want to drastically alter the composition, you can adjust the constraints however you like it. You may also choose to adjust only one or two macronutrient constraints. Just be aware that diet will only be as realistic as the constraints you put on the system.

Here we will set the constraints to keep more or less the same dietary composition, and we shall only add 150g of new food items at maximum. We also demonstrate here the capability to remove food items from the analysis. We saw in the previous calculation that we should add 130g of sorbitol to the diet, an item that is not typically consumed in its raw form. Here we remove the item with the option "removeFoodItems". To discard further items from analysis, add new rows with;

```
\% Let us clear the options from the previous run clear options
```

```
% Set the new options for the new run
% We want to calculate food item dietary changes
options.foodOrMets = 'Food';
% We will not weight our reaction of interest more than our dietary
% reactions
options.roiWeights=[1 1 1];
% Set the dietary limits
options.caloricRange=[1900 2300];
options.carbohydrateRange = [300 370];
options.lipidRange = [60 80];
options.proteinRange = [40 60];
options.sugarsRange = [170 210];
% Set the maximum amount of food addded
options.foodAddedLimit=150;
% Remove food item sorbitol
options.removeFoodItem = {'1351', 'frida'};
% Run the nutrition algorithm again
[newDietModel,pointsModel,roiFlux,pointsModelSln,menuChanges, macroChanges,
roiChanges] = nutritionAlgorithmWBM(model,obj,objMinMax,rois,roisMinMax,options);
```

```
Reaction of Interest 1 = DM_glc_D[bc]
Reaction of Interest 2 = DM_dopa[csf]
Reaction of Interest 3 = Diet_EX_lipid[d]
Solution points =-2818.5508
180 come from diet
-2998.5508 come from roi
Food items of interest are:
```

†oodNames	status	Flux	+00
{["Alcoholic beverage, distilled, all (gin, rum, vodka, whiskey) 100 proof"]} {["Oil, soybean"	{'Added' }	-2.0068 -11.974 -15.126 -72.136 -16.928 -27.356 -4.4748	{'17; {'74; {'17; {'23; {'22; {'13; {'14;

```
Diet Energy = 2300
f1 =1 & f2=1
DM_glc_D[bc]
Original Diet RoI range = 0:1328.4361
New Diet RoI range = 0:3083.4015
DM_dopa[csf]
Original Diet RoI range = 0:10.7923
New Diet RoI range = 0:19.215
Diet_EX_lipid[d]
Original Diet RoI range = 71.282:71.282
```

We can see that the dietary changes are different, although some items remained the same. We now add ~150g of food to the diet while removing ~30g of food from the diet. The removed food was likely removed to make space for the carbohydrate macronutrient constraints. We can also see that, indeed, sorbitol does no longer show up in the solution, as we removed it in the options. To double-check if it would show up in the solution with these new constraints you can comment out (put % in front of) the line where we set which food items we want to have removed and rerun.

The main difference we see in our reactions of interest is for dopamine in the cerebrospinal fluid, where we now only see an increase of ~9 mmol/person/day. A possbile explanation is that the solution in the nutrition algorithm is biased by how many points are generated by the reactions of interest. It sees that more points can be achieved with glucose (up to ~3000 points), which is far higher than for dopamine. This leads to the algorithm preferring dietary changes to favour increased glucose in the blood, as this will garner more points. A way to combat this bias is by adjusting the weight of the dopamine reaction in roiWeights option to 10 or 100, so that every point made in that reaction will be counted 10 or 100 times as much. This way the reactions of interest can have the same influence on the solution.

If we look at the macro nutrient distribution:

### disp(macroChanges);

macroNames	originalMacro	newMacro	relChange	
{'energy' }	2190.8	2302.4	111.55	
{'carbohydrate'}	344.36	361.4	17.043	
{'protein' }	45.317	60.132	14.815	
{'lipid' }	71.282	80.384	9.1024	
{'sugars' }	194.9	209.87	14.972	

We see indeed that all the new macros are within the ranges we set. It might slighty pass over or under the set bounds as we set the diet with a 1% margin of the calculated optimal dietary changes. We also see that despite asking to minimise the lipids in the diet, it still calculated that the optimal solution would be to add the highstest allowed amount of lipids. This means that its more important for the model to get the glucose maximised than it is to minimise the lipid consumption. Again we change the roiWeights to give more importance to minimising the lipid content in the diet to limit the effect glucose has on the solution.

To give the diets an additional layer of realism, we can add on the money food items costs. This can be done through the options variable:

options.addPrice: a nx3 cell array with the first column the food item ID, the second column the name of the database, and in the third column the price (any currency) associated with that food item. This allows for optimising the cost of a diet, but only if sufficient information is available.

Important is that you remove all other food items that you do not have price information for. Otherwise the algorithm will include those items as they "do not cost anything". Note that this will be a time-intensive option as you will have to link price to the various database items of interest.

This concludes the tutorial on how to work with the nutrition toolbox. If more information is required on the nutrition algorithm, we recommend reading the original paper [2] and looking at the tutorial on just the nutrition algorithm, present in the COBRA toolbox. For any questions, please use the COBRA toolbox google community <a href="https://groups.google.com/g/cobra-toolbox">https://groups.google.com/g/cobra-toolbox</a>, the issues on the COBRA toolbox github, or contact us directly through the corresponding author from the publication associated with this tutorial. **%Include actual reference here.**%

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