COBRAme Documentation

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Colton Lloyd, Ali Ebrahim, Laurence Yang

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This resource is intended to:

- 1. Provide an overview of what ME-models are and how they work
- 2. Describe the architecture of the COBRAme code base.
- 3. Highlight the major object classes and how they interrelate
- 4. Demonstrate basic model building and editing procedures
- 5. Provide examples of how to query, edit and update information in a constructed ME-Model

CHAPTER 1

ME-Model Fundamentals

Models of metabolism and expression (ME-models) are unique in that they are capable of predicting the optimal macromolecular expression required to sustain a metabolic phenotype. In other words, they are capable of making novel predictions of the amount of individual protein, nucleotides, cofactors, etc. that the cell must synthesize in order to grow optimally. To enable these types of predictions, ME-models differ from metabolic models (M-models) in a few key ways:

- 1. ME-models are multi-scale in nature so they require the addition of *coupling constraints* to couple cellular processes to each other.
- 2. ME-models predict the biomass composition of a growing cell thus forgoing much of the M-model biomass composition function. For this reason, the function representing growth needs to be updated.

These two ME-model features are briefly described below. Their practical implementation is further outlined in **ME-model Reaction Building**.

Coupling Constraints

Coupling constraints are required in an ME-model in order to couple a reaction flux to the synthesis of the macromolecule catalyzing the flux. The easiest example of this is for the coupling of metabolic enzymes to metabolic reactions. This has the form:

$$\frac{\mu}{k_{eff}}(1.1)$$

where μ is the growth rate and k_{eff} is an approximation of the effective turnover rate for the metabolic process. The coupling of enzyme synthesis cost to metabolic flux scales with μ to represent the dilution of macromolecules as they are passed on the daughter cells. More macromolecules are diluted at faster growth rates. Enzyme turnover rates determine the efficiency of of an enzyme *in vivo* and are largly unknown for a majority of metabolic and expression enzymes. Optimizing the vector of k_{effs} for the cellular processes modeled in the ME-model is an ongoing area of

research. Currently, the $k_{eff}s$ are set with an average of 65 s^{-1} and scaled by their solvent accessible surface area (approximated as protein_moleculare_weight³/₄). A set ~125 metabolic k_{effs} were found by Ebrahim et. al. 2016 to be particularly important in E. coli for computing an accurate metabolic/proteomic state using proteomics data. We suspect similar observations would be seen in other organisms.

For non metabolic macromolecules such as ribosome, mRNA, tRNA and RNA polymerase, the coupling constrains coefficients ($k_{eff}s'$) are derived by essentially back-calculating the individual rates using a measured RNA-to-Protein ratio from Scott et. al. 2010 and measured mRNA, tRNA and rRNA fractions. The coupling constraints coefficients for these macromolecules are derived in detail in O'Brien et. al. 2013. Applying these constraints results in a final nonlinear optimization problem (NLP) shown below. COBRAme reformulates these coupling constraints to embed them directly into the reaction which they are used as follows:

$$\begin{split} \max_{v,\mu} \mu \\ \text{s.t. } Sv &= 0 \\ v_{\text{formation,Ribosome}} - \sum_{i \in Peptide} \left(\frac{l_{p,i}}{c_{\text{ribo}}\kappa_{\tau}} (\mu + r_{0}\kappa_{\tau}) \cdot v_{\text{translation},i} \right) = 0 \\ v_{\text{formation,RNAP}} - \sum_{i \in TU} \left(\frac{l_{\text{TU},i}}{3c_{\text{ribo}}\kappa_{\tau}} (\mu + r_{0}\kappa_{\tau}) \cdot v_{\text{transcription},i} \right) = 0 \\ v_{\text{formation,j}} \\ - \sum_{i \in \text{generic_tRNA_{AA}}} \left(\left(1 + \frac{\mu}{k_{eff,tRNA}} \right) \frac{\mu}{k_{eff,charging}} v_{\text{charging},i} \right) = 0 \\ \forall j \in Synthetase \\ v_{\text{formation,j}} - \sum_{i \in \text{enzymatic reaction}} \left(\frac{\mu}{k_{ij}^{\text{eff}}} v_{\text{usage},i} \right) = 0, \quad \forall j \in Enzyme \\ v_{\text{formation,j}} - \sum_{i \in \text{tRNA anticodons}} \frac{(\mu + \kappa_{\tau}r_{0})}{\kappa_{\tau}c_{tRNA,j}} v_{charging,i} = 0, \quad \forall j \in tRNA \\ v_{\text{degredation,j}} - \frac{3k_{deg,j}}{\kappa_{\tau}c_{mRNA}} \cdot \frac{\mu + \kappa_{\tau}r_{0}}{\mu} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation,j}} - \frac{(\mu + \kappa_{\tau}r_{0})}{3\kappa_{\tau}c_{mRNA}} v_{translation,j} = 0, \quad \forall j \in mRNA \\ v_{\text{formation}} - \frac{(\mu +$$

The biomass_dilution constraint is discussed below.

Previous ME-Model Coupling Constraint Implementation

v,

The previous iterations of ME models applied coupling constraints using the following:

a + enzyme
$$\xrightarrow{v1} \alpha \cdot \text{coupling} + \text{enzyme} + b \Rightarrow a \xrightarrow{v1} \alpha \cdot \text{coupling} + b$$

coupling + enzyme $\xrightarrow{\alpha \cdot v1} \emptyset$

 $\alpha \cdot \operatorname{coupling} + \alpha * \operatorname{enzyme} \xrightarrow{v^2} \emptyset$

Where α is the coupling coefficient applied to "coupling", a metabolite (constraint) that can be given a "constraint sense" in COBRApy of 'L' which effectively allows the second reaction to proceed at a rate equal to or greater than $v2 \ge \alpha \cdot v1$.

COBRAme Coupling Constraint Implementation

With COBRAme it is assumed that the optimal ME solution will never dilute more enzyme than is required by the coupling constraint thus constraining $v2 = \alpha \cdot v1$ and allowing us to combine the implementation of the constraint with the reaction that uses the enzyme to give.

$$\mathbf{a} + \alpha \cdot \mathbf{enzyme} \xrightarrow{v1} \mathbf{b}$$

The coupling constraints and coefficients were derived as in O'Brien et. al. 2013. As stated above, however, these were implemented in the current study as equality constraints. Effectively, this means that each ME-model solution will give the computed optimal proteome allocation for the *in silico* conditions. Previous ME-model formulations have applied the constraints as inequalities thus allowing the simulation to overproduce macromolecule components. While overproduction is seen *in vivo* in cells, this phenomenon would not be selected as the optimal solution. Furthermore, using inequality constraints greatly expands the size of the possible solution space significantly increasing the time required to solve the optimization. Reformulating the model using equality constraints thus resulted in a reduced ME-matrix with the coupling constraints embedded directly into the reaction in which they are used.

Biomass Dilution Constraints

For metabolic models (M-models), the biomass objective function has been used to represent the amount of biomass production that is required for the cell to double. The metabolites represented in the biomass function are typically building blocks for major macromolecules (e.g. amino acids and nucleotides), cell wall components and cofactors. The coefficients of the biomass objective function are determined from empirical measurements from a cell growing at a measured rate. Since ME-models explicitly compute the predicted amount of RNA, protein, cofactors, etc. necessary for growth, this concept has to be modified for ME models.

This is accomplished via the *biomass_dilution* variable (reaction), which contains the *biomass* constraint (pseudo metabolite) that represents the mass produced by the synthesis of each functional RNA or protein. This reaction essentially ensured that the ME-model can only produce biomass at the rate it is being diluted (via growth and division).

Implementation

For each ME-model transcription or translation reaction in an ME-model an amount of a *biomass* constraint (pseudo metabolite) is created with a stoichiometry equal to the molecular weight of the mRNA or protein being made (in kDA). The below figure shows an example of this where a translation reaction produces both the catalytic protein as well as the protein biomass constraint. The formed *protein_biomass* constraint is a participant in the overall ME *Biomas_Dilution* reaction which restricts the total production of the major biomass components to equal the rate at which biomass is diluted (eg. the cell's growth rate, μ).

Some biomsas constituents do not have a mechanistic function in the ME-model (e.g. cell wall components, DNA and glycogen). These metabolites are included in the *biomass_dilution* reaction indentical to the M-model biomass reaction.

The units for this contraint work out as follows:

The units of a given reaction in the ME-mdoel are in molecules per hour.

$$v_i: \frac{mmol}{gDW \cdot hr}$$

The individual components of the biomass dilution constrain are in units of kDa.

$$molecular_weight: kDA[rac{g}{mmol}]$$

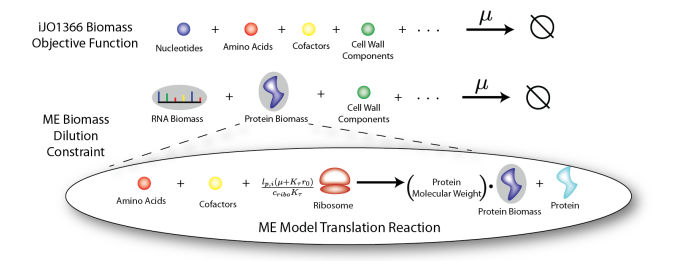


Fig. 1.1: Biomass Dilution for iJO1366 and for a ME-Model

Therefore, when the *biomass dilution* variable (reaction) carries flux it gives units of hr^{-1} representing the growth rate of the cell, μ

 $molecular_weight \cdot v_i : hr^{-1}[\mu]$

CHAPTER 2

ME-Model Architecture

The COBRAme codebase is constructed with the intention of both including all of the metabolic processes associated with gene expression, while also giving a final ME-model that was not prohibitively difficult to use and interpret. To accomplish this we decided to separate the information associated with each cellular process from the actual ME-model reaction in which the process is modeled. We called these two major object classes ProcessData and MEReaction, respectively. The logic behind each of these classes is briefly presented below, and a description of the class attributes and properties is presented in xxxx.

ProcessData

The previous ME-models for *E. coli* and *T. Maritima* established a database containing, for instance, all of the enzyme complex stoichiometries or the *E. coli* transcription unit architecture. Then, when building the ME-model, the database was queried to obtain any relevant information and incorporate this into the appropriate reactions. For the COBRAme formulation, this database was replaced by the ProcessData "information storage" class. The ProcessData class generally consists of attributes which use simple python types (string, dictionary, etc.) to describe features of a biological process.

The use of the ProcessData class has the advantage of:

- 1. Simplifying the process of querying information associated with a cellular function
- 2. Allowing edits to this information to easily be applied throughout the model without rebuilding from scratch
- 3. Enabling additional computations to be performed and seamlessly accessed as ProcessData class properties

The ProcessData classes are broken into the following subclasses:

ProcessData		
Subclass	Information Contained	Example
	Metabolite stoichiometry of a metabolic reaction	
StoichiometricData	(often equivalent to M-model reaction)	HISTD
	Protein subunit stoichiometry of an enzyme complex	
ComplexData	as well as the modifications required for its activity	ATPSYN_complex
	Metabolite stoichiometry of an enzyme complex	
	modification and information about any chaperone	
ModificationData	catalyzing the modification process	Mod_2fe2s_c
	Some processes occur in multiple steps (e.g.	
	translation reactions). This class details the metabolic	
	stoichiometry and catalytic enzyme associated with	
SubreactionData	the process.	ala_addition_at_GCA
	Nucleotide sequence, RNA products, sigma factor	
Transcription Data	usage, etc. for a given transcription unit	TU_xxxxxx
	Subreactions (tRNA mediated amino acid additions),	
	sequence of mRNA/protein, etc. for a given mRNA to	
Translation Data	be translated	b2020
	Codon, Amino acid, tRNA, and modifications required	
tRNAData	to make a functioning tRNA	tRNA_b0202_AUU
	Keff, enzymes, metabolite stoichiometry of a	
Translocation Data	particular protein translocation pathway	srp_translocation
	Details the translocation pathways, protein	
	modifications (for lipoproteins), etc. required to	
PostTranslationData	produce a function protein.	translocation_protein_b0733
	Redundant complexes or metabolites are represented	generic_Tuf (elongation
GenericData	as generics	factor)

MEReaction

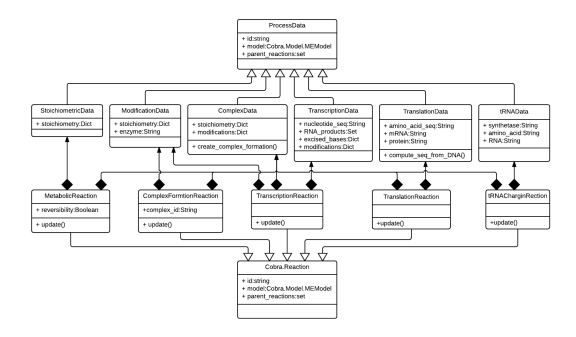
COBRAme compartmentalizes the major reaction types into their own MEReaction classes. Each of these classes contains a single *update* function which effectively reads in the appropriate ProcessData types, applies the coupling constraints on the macromolecules and assembles these into a complete model reaction. This allows changes made to the ProcessData of a cellular process to easily be incorporated into the reactions which it was used.

Overview

Using the major classes described above, building a ME-model can then be broken down into two steps:

- 1. Define and construct all necessary ProcessData objects
- 2. Link the ProcessData to the appropriate MEReaction

The overall codebase architecture is displayed below in the following UML diagram. This includes the ways which MEReactions and ProcessData are often linked in the model.



CHAPTER 3

Building a ME-Model

```
In [1]: from __future__ import print_function
```

```
import cobrame
from cobrame.util import dogma, building
import cobrame.util.building
import cobra
import cobra.test
from collections import defaultdict
```

```
#import warnings
#warnings.filterwarnings('ignore')
```

Overview

COBRAme is constructed entirely over COBRApy. This means that ME-model reactions will have all of the same properties, methods, and functions as a COBRApy reaction. However, one key difference between M and ME models is that many reactions involved in gene expression are effectively templates that are constructed identically but vary based on characteristics of the gene being expressed. For example, a gene with a given nucleotide sequence is always translated following the same rules provided by the codon table for that organism.

In order to facilliate the template nature of many gene expression reactions, COBRAme reactions are constructed and their components are manipulated through the use of ProcessData classes. These act as information vessels for holding the information assocatied with a cellular process in simple, standard datatypes such as dictionaries and strings.

This tutorial will go step-by-step through the process of creating a generic enzyme catalyzed reaction (i.e. metabolic reaction):

 $a \to b$

which requires the formation and coupling of complex_ab in order to proceed.

In order for this reaction to carry flux in the model we will additionally need to first add the corresponding:

- 1. Transcription reactions
- 2. Translation reactions
- 3. tRNA charging reactions
- 4. Complex formation reactions

Once these are added we will add in the synthesis of key macromolecular components (ribosome, RNA polymerase, etc.) and show how they are coupled to their respective reactions. The derived coupling coefficients will also be described. For more on the derivation of the coupling coefficients, reference the supplemental text of O'brien et. al. 2013

Initializing new ME-Models

When applying some constraints in the ME-model, metabolite properties are required. For instance, to calculate the total biomass (by molecular weight) produced by a particular macromolecule, the amino acid, nucleotide, etc. molecular weights are required. To enable these calculations, all metabolites from *i*JO1366, along with their metabolite attributes are added to the newly initialized ME-model.

Further the reactions from *i*JO1366 will be added to the ME-model to demonstrate ME-model solving procedures.

```
In [2]: # create empty ME-model
  me = cobrame.MEModel('test')
    ijo = cobra.test.create_test_model('ecoli')
In [3]: # Add all metabolites and reactions from iJ01366 to the new ME-model
  for met in ijo.metabolites:
        me.add_metabolites(met)
    for rxn in ijo.reactions:
        me.add_reaction(rxn)
```

The ME-model contains a "global_info" attribute which stores information used to calculate coupling constraints, along with other functions. The specifics of each of these constraints will be discussed when they are implemented.

Note: k_deg will initially be set to 0. We will apply RNA degradation later in the tutorial.

```
In [4]: # "Translational capacity" of organism
  me.global_info['kt'] = 4.5 # (in h-1)scott 2010, RNA-to-protein curve fit
  me.global_info['ro'] = 0.087 # scott 2010, RNA-to-protein curve fit
  me.global_info['k_deg'] = 1.0/5. * 60.0 # 1/5 1/min 60 min/h # h-1
  # Molecular mass of RNA component of ribosome
  me.global_info['m_rr'] = 1453. # in kDa
  # Average molecular mass of an amino acid
  me.global_info['m_aa'] = 109. / 1000. # in kDa
  # Proportion of RNA that is rRNA
  me.global_info['f_rRNA'] = .86
  me.global_info['m_nt'] = 324. / 1000. # in kDa
  me.global_info['f_mRNA'] = .02
  # tRNA associated global information
```

Define sequence of gene that will be expressed in tutorial

Adding Reactions without utility functions

We'll first demonstrate how transcription, translation, tRNA charging, complex formation, and metabolic reactions can be added to a model without using any of the utility functions provided in cobrame.util.building.py. The second half of the tutorial will show how these utility functions can be used to add these reactions.

The basic workflow for adding any reaction to a ME-model using COBRAme occurs in three steps:

- 1. Create the ProcessData(s) associated with the reaction and populate them with the necessary information
- 2. Create the MEReaction and link the appropriate ProcessData
- 3. Execute the MEReaction's update method

Add Transcription Reaction

Add TranscribedGene metabolite to model

Transcription reactions is unique in that they occur at a transcription unit level and can code for multiple transcript products. Therefore the nucleotide sequence of both the transcription unit and the RNA transcripts must be defined in order to correctly construct a transcription reaction.

class cobrame.core.component.**TranscribedGene**(*id*, *rna_type*, *nucleotide_sequence*)

Metabolite class for gene created from cobrame.core.reaction.TranscriptionReaction

Parameters

- id (str) Identifier of the transcribed gene. As a best practice, this ID should be prefixed with 'RNA + _'
- **rna_type** (*str*) Type of RNA encoded by gene sequence (mRNA, rRNA, tRNA, or ncRNA)
- **nucleotide_sequence** (*str*) String of base pair abbreviations for nucleotides contained in the gene

left_pos

int – Left position of gene on the sequence of the (+) strain

```
right_pos
```

int - Right position of gene on the sequence of the (+) strain

strand

str –

•(+) if the RNA product is on the leading strand

•(-) if the RNA product is on the comple(mentary strand

```
In [6]: gene = cobrame.TranscribedGene('RNA_a', 'mRNA', sequence)
    me.add_metabolites([gene])
```

When adding the TranscribedGene above, the RNA_type and nucleotide_sequence was assigned to the gene. This sequence cannot be determined from the transcription unit (TU) sequence because a single TU often contains several different RNAs.

Add TranscriptionData to model

```
class cobrame.core.processdata.TranscriptionData(id, model, rna_products=set([]))
Class for storing information needed to define a transcription reaction
```

Parameters

- id (str) Identifier of the transcription unit, typically beginning with 'TU'
- model (*cobrame.core.model.MEModel*) ME-model that the TranscriptionData is associated with

nucleotide_sequence

str - String of base pair abbreviations for nucleotides contained in the transcription unit

RNA_products

set – IDs of *cobrame.core.component.TranscribedGene* that the transcription unit encodes. Each member should be prefixed with "RNA + _"

RNA_polymerase

str – ID of the *cobrame.core.component.RNAP* that transcribes the transcription unit. Different IDs are used for different sigma factors

subreactions

collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata. SubreactionData ID: num_usages} required for the transcription unit to be transcribed

Add TranscriptionReaction to model

And point TranscriptionReaction to TranscriptionData

```
class cobrame.core.reaction.TranscriptionReaction (id)
Transcription of a TU to produced TranscribedGene.
```

RNA is transcribed on a transcription unit (TU) level. This type of reaction produces all of the RNAs contained within a TU, as well as accounts for the splicing/excision of RNA between tRNAs and rRNAs. The appropriate RNA_biomass constrain is produced based on the molecular weight of the RNAs being transcribed

Parameters id (*str*) – Identifier of the transcription reaction. As a best practice, this ID should be prefixed with 'transcription + _'

```
In [8]: transcription_rxn = cobrame.TranscriptionReaction('transcription_TU_a')
    transcription_rxn.transcription_data = transcription_data
    me.add_reactions([transcription_rxn])
```

Update TranscriptionReaction

TranscriptionReaction.update(verbose=True)

Creates reaction using the associated transcription data and adds chemical formula to RNA products

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for *cobrame.core.processdata.TranscriptionData*):

1.RNA_polymerase from data.RNA_polymerase w/ coupling coefficient (if present)

2.RNA products defined in data.RNA_products

3.Nucleotide reactants defined in data.nucleotide_counts

4.If tRNA or rRNA contained in data.RNA_types, excised base products

5.Metabolites + enzymes w/ coupling coefficients defined in data.subreactions (if present)

6.Biomass cobrame.core.component.Constraint corresponding to data.RNA_products and their associated masses

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

In [9]: transcription_rxn.update()
 print(transcription_rxn.reaction)

```
86 atp_c + 38 ctp_c + 12 gtp_c + 50 utp_c --> RNA_a + 59.172286 mRNA_biomass + 186 ppi_c
/home/sbrg-cjlloyd/cobrame/cobrame/core/reaction.py:813 UserWarning: RNA Polymerase () not found
```

Note: the RNA_polymerase complex is not included in the reaction. This will be added later

This reaction now produces a small amount of the a mRNA_biomass metabolite (constraint). This term has a coefficient corresponding to the molecular weight (in kDA) of the RNA being transcribed. This constraint will be implemented into a $v_{biomasss_dilution}$ reaction with the form:

 $\mu \leq v_{biomass_dilution} \leq \mu$

A mathematical description of the biomass constraint can be found in *Biomass Dilution Constraints* in **ME-Model** Fundamentals.

Note: This is not a complete picture of transcription because the RNA polymerase is missing.

Incorporate RNA Polymerase

For the purposes of this tutorial, we'll skip the steps required to add the reactions to form the RNA_polymerase. The steps however are identical to those outlined in **add enzyme complexes** below

class cobrame.core.component.RNAP (id)

Metabolite class for RNA polymerase complexes. Inherits from cobrame.core.component.Complex

Parameters id (*str*) – Identifier of the RNA Polymerase.

In [10]: RNAP = cobrame.RNAP('RNA_polymerase')
 me.add_metabolites(RNAP)

Associate RNA_polymerase with all TranscriptionData and update

print (me.reactions.transcription_TU_a.reaction)

```
0.00088887053605567*mu + 0.000347992814865795 RNA_polymerase + 86 atp_c + 38 ctp_c + 12 gtp_c + 50 ut
```

The coefficient for RNA_polymerase is the first instance in this tutorial where a coupling constraint is imposed. In this case the constraint couples the formation of a RNA_polymerase metabolite to its transcription flux. This constraint is formulated as in O'brien et. al. 2013, with assumption that $k_{rnap} = 3 \cdot k_{ribosome}$ based on data from Proshkin et al. 2010:

$$v_{dilution,RNAP,j} = \frac{l_{TU,j}}{3c_{ribo}\kappa_{\tau}} v_{transcription,j}(\mu + r_0\kappa_{\tau}), \forall j \in TU$$
(3.1)

where:

- κ_{τ} and r_0 are phenomenological parameters from Scott et. al. 2010 that describe the linear relationship between the observed RNA/protein ratio of *E. coli* and its growth rate (μ)
- $c_{ribo} = \frac{m_{rr}}{f_{rRNA} \cdot m_{aa}}$ where: m_{nt} is the mass of rRNA per ribosome. f_{rRNA} is the fraction of total RNA that is rRNA m_{aa} is the molecular weight of an average amino acid
- $v_{transcription,j}$ is the rate of transcription for TU_j
- $l_{TU,j}$ is number of nucleotides in TU_j

Add Translation Reaction

Add TranslationData to model

In order to add a TranslationData object to a ME-model the user must additionally specifify the mRNA id and protein id of the translation reaction that will be added. This information as well as a nucleotide sequence is the only information required to add a translation reaction.

class cobrame.core.processdata.TranslationData (id, model, mrna, protein)

Class for storing information about a translation reaction.

Parameters

- id (str) Identifier of the gene being translated, typically the locus tag
- model (*cobrame.core.model.MEModel*) ME-model that the TranslationData is associated with
- mrna (str) ID of the mRNA that is being translated
- **protein** (*str*) ID of the protein product.

mRNA

str - ID of the mRNA that is being translated

protein

str – ID of the protein product.

subreactions

```
collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata.
SubreactionData.id: num_usages} required for the mRNA to be translated
```

nucleotide_sequence

str - String of base pair abbreviations for nucleotides contained in the gene being translated

Add TranslationReaction to model

By associating the TranslationReaction with its corresponding TranslationData object and running the update function, COBRAme will create a reaction reaction for the nucleotide sequence given based on the organisms codon table and prespecified translation machinery.

```
class cobrame.core.reaction.TranslationReaction (id)
Reaction class for the translation of a TranscribedGene to a TranslatedGene
```

```
Parameters id (str) – Identifier of the translation reaction. As a best practice, this ID should be prefixed with 'translation + _'
```

Update TranslationReaction

TranslationReaction.update(verbose=True)

Creates reaction using the associated translation data and adds chemical formula to protein product

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.TranslationData):

- 1.Amino acids defined in data.amino_acid_sequence. Subtracting water to account for condensation reactions during polymerization
- 2. Ribosome w/ translation coupling coefficient (if present)

3.mRNA defined in data.mRNA w/ translation coupling coefficient

- 4.mRNA + nucleotides + hydrolysis ATP cost w/ degradation coupling coefficient (if kdeg (defined in model.global_info) > 0)
- 5.RNA_degradosome w/ degradation coupling coefficient (if present and kdeg > 0)
- 6.Protein product defined in data.protein
- 7. Subreactions defined in data.subreactions
- 8.protein_biomass cobrame.core.component.Constraint corresponding to the protein product's mass
- 9.Subtract mRNA_biomass cobrame.core.component.Constraint defined by mRNA degradation coupling coefficinet (if kdeg > 0)

Parameters verbose (bool) – Prints when new metabolites are added to the model when execut-

ing update()

```
In [14]: rxn.update()
    print(rxn.reaction)
```

```
/home/sbrg-cjlloyd/cobrame/cobrame/core/reaction.py:1051 UserWarning: ribosome not found
/home/sbrg-cjlloyd/cobrame/cobrame/core/reaction.py:1094 UserWarning: RNA_degradosome not found
```

```
0.000498399634202103*mu + 0.000195123456790123 + 0.00598079561042524*(mu + 0.3915)/mu RNA_a + 12 asp
```

In this case the constraint couples the formation of a mRNA metabolite to its translation flux. This constraint is formulated as in O'brien et. al. 2013:

$$v_{dilution,j} = \frac{3}{\kappa_{\tau} c_{mRNA}} \cdot (\mu + \kappa_{\tau} r_0) v_{translation,j}, \qquad \forall j \in mRNA$$
(3.2)
(3.3)

where:

- κ_{τ} and r_0 are phenomenological parameters from Scott et. al. 2010 that describe the linear relationship between the observed RNA/protein ratio of *E. coli* and its growth rate (μ)
- $c_{mRNA} = \frac{m_{nt}}{f_{mRNA} \cdot m_{aa}}$ where: m_{nt} is the molecular weight of an average mRNA nucleotide. f_{mRNA} is the fraction of total RNA that is mRNA m_{aa} is the molecular weight of an average amino acid
- $v_{translation,j}$ is the rate of translation for $mRNA_j$

Incorporate Ribosome

```
class cobrame.core.component.Ribosome (id)
```

Metabolite class for Ribosome complexes. Inherits from cobrame.core.component.Complex

Parameters id (*str*) – Identifier of the Ribosome.

```
In [15]: ribosome = cobrame.Ribosome('ribosome')
    me.add_metabolites([ribosome])
    me.reactions.translation_a.update()
    print(me.reactions.translation_a.reaction)
```

0.000498399634202103*mu + 0.000195123456790123 + 0.00598079561042524*(mu + 0.3915)/mu RNA_a + 12 asp

This imposes a new coupling constraint for the ribosome. In this case the constraint couples the formation of a ribosome to its translation flux. This constraint is formulated as in O'brien et. al. 2013:

$$v_{dilution,ribo,j} = \frac{l_{p,j}}{c_{ribo}\kappa_{\tau}} v_{translation,j}(\mu + r_0\kappa_{\tau}), \forall j \in mRNA$$
(3.4)

where:

- κ_{τ} and r_0 are phenomenological parameters from Scott et. al. 2010 that describe the linear relationship between the observed RNA/protein ratio of *E. coli* and its growth rate (μ)
- $c_{ribo} = \frac{m_{rr}}{f_{rRNA} \cdot m_{aa}}$ where: m_{nt} is the mass of rRNA per ribosome. f_{rRNA} is the fraction of total RNA that is rRNA m_{aa} is the molecular weight of an average amino acid
- $v_{translation,j}$ is the rate of translation for $mRNA_j$
- $l_{p,j}$ is number of amino acids in peptide translated from $mRNA_j$

Note: The above reactions do not provide a complete picture of translation in that it is missing charged tRNAs to facilitate tRNA addition.

Below, we'll correct this by adding in an tRNA charging reaction.

Add tRNA Charging Reaction

Add tRNAData to model

```
In [16]: # Must add tRNA metabolite first
gene = cobrame.TranscribedGene('RNA_d', 'tRNA', sequence)
me.add_metabolites([gene])
```

class cobrame.core.processdata.**tRNAData** (*id*, *model*, *amino_acid*, *rna*, *codon*) Class for storing information about a tRNA charging reaction.

Parameters

- id (*str*) Identifier for tRNA charging process. As best practice, this should be follow "tRNA + _ + <tRNA_locus> + _ + <codon>" template. If tRNA initiates translation, <codon> should be replaced with START.
- model (*cobrame.core.model.MEModel*) ME-model that the tRNAData is associated with
- amino_acid (str) Amino acid that the tRNA transfers to an peptide
- rna (str) ID of the uncharged tRNA metabolite. As a best practice, this ID should be prefixed with 'RNA + _'

subreactions

```
collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata.
SubreactionData.id: num_usages} required for the tRNA to be charged
```

synthetase

str - ID of the tRNA synthetase required to charge the tRNA with an amino acid

synthetase_keff

float – Effective turnover rate of the tRNA synthetase

```
In [17]: data = cobrame.tRNAData('tRNA_d_GUA', me, 'val__L_c', 'RNA_d', 'GUA')
```

Add tRNAChargingReaction to model

And point tRNAChargingReaction to tRNAData

class cobrame.core.reaction.**tRNAChargingReaction** (*id*) Reaction class for the charging of a tRNA with an amino acid

Parameters id (*str*) – Identifier for the charging reaction. As a best practice, ID should follow the template "charging_tRNA + _ + <tRNA_locus> + _ + <codon>". If tRNA initiates translation, <codon> should be replaced with START.

```
In [18]: rxn = cobrame.tRNAChargingReaction('charging_tRNA_d_GUA')
    me.add_reaction(rxn)
    rxn.tRNA_data = data
```

Update tRNAChargingReaction

tRNAChargingReaction.update (verbose=True) Creates reaction using the associated tRNA data

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for *cobrame.core.processdata.tRNAData*):

- 1.Charged tRNA product following template: "generic_tRNA + _ + <data.codon> + _ + <data.amino_acid>"
- 2.tRNA metabolite (defined in data.RNA) w/ charging coupling coefficient
- 3. Charged amino acid (defined in data.amino_acid) w/ charging coupling coefficient
- 5.Synthetase (defined in data.synthetase) w/ synthetase coupling coefficient found, in part, using data.synthetase_keff
- 6.Post transcriptional modifications defined in data.subreactions

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

In [19]: #Setting verbose=False suppresses print statements indicating that new metabolites were creation.
print(rxn.reaction)

0.00011626666666666667*mu + 4.55184e-5 RNA_d + 0.000116266666666666667*mu + 4.55184e-5 val__L_c --> gener.

This reaction creates one generic_charged_tRNA equivalement that can then be used in a translation reaction

The coefficient for RNA_d and lys__L_c are defined by:

$$v_{dilution,j} \ge \frac{1}{\kappa_{\tau} c_{tRNA,j}} (\mu + \kappa_{\tau} r_0) v_{charging,j}, \forall j \in tRNA$$
(3.5)

where:

- κ_{τ} and r_0 are phenomenological parameters from Scott et. al. 2010 that describe the linear relationship between the observed RNA/protein ratio of *E. coli* and its growth rate (μ)
- $c_{tRNA,j} = \frac{m_{tRNA}}{f_{tRNA} \cdot m_{aa}}$ where: m_{tRNA} is molecular weight of an average tRNA. f_{tRNA} is the fraction of total RNA that is tRNA m_{aa} is the molecular weight of an average amino acid
- $v_{charging,j}$ is the rate of charging for $tRNA_j$

Note: This tRNA charging reaction is still missing a tRNA synthetase which catalyzes the amino acid addition to the uncharged tRNA.

Incorporate tRNA Synthetases

.. autoclass:: cobrame.core.component.Complex :noindex:

```
In [20]: synthetase = cobrame.Complex('synthetase')
            me.add_metabolites(synthetase)
```

Associate synthetase with tRNAData and update

The synthetase coupling was reformulated from O'brien et. al. 2013 enable more modularity in the ME-model. A more complete mathematical description of the tRNA synthetase coupling constraints can be found in the tRNA.ipynb

Add tRNAs to Translation

Here we take advantage of an additional subclass of ProcessData, called a SubreactionData object. This class is used to lump together processes that occur as a result of many individual reactions, including translation elongation, ribosome formation, tRNA modification, etc. Since each of these steps often involve an enzyme that requires its own coupling constraint, this process allows these processes to be lumped into one reaction while still enabling each subprocess to be modified.

TranslationData objects have an subreaction_from_sequence method that returns any subreactions that have been added to the model and are part of translation elongation (i.e. tRNA). Since no tRNA-mediated amino acid addition subreactions have been added to the model, the below call returns nothing.

In [22]: print(me.process_data.a.subreactions_from_sequence)

```
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction the
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction phy
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction asp
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction ty:
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction ty:
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:826 UserWarning: tRNA addition subreaction ty:
```

UserWarnings are returned to indicate that tRNA subreactions have not been added for each codon.

Below, we add the SubreactionData (excluding enzymes) for the addition of an amino acid using information from the *E. coli* codon table. The charge tRNA does not act as an enzyme in this case because it's coupling is handled in the tRNAChargingReaction

Add Subreactions for tRNA addition to model

class cobrame.core.processdata.SubreactionData (id, model)

Parameters

- id (*str*) Identifier of the subreaction data. As a best practice, if the subreaction data details a modification, the ID should be prefixed with "mod + _"
- model (*cobrame.core.model.MEModel*) ME-model that the SubreactionData is associated with

enzyme

list or str or None – List of cobrame.core.component.Complex.id s for enzymes that catalyze this process

or

String of single cobrame.core.component.Complex.id for enzyme that catalyzes this process

keff

float - Effective turnover rate of enzyme(s) in subreaction process

_element_contribution

dict – If subreaction adds a chemical moiety to a macromolecules via a modification or other means, net element contribution of the modification process should be accounted for. This can be used to mass balance check each of the individual processes.

Dictionary of {element: net_number_of_contributions}

Now calling subreactions_from_sequence returns the number of tRNA subreactions that should be added to the TranslationData

```
In [24]: translation_subreactions = me.process_data.a.subreactions_from_sequence
    print(translation_subreactions)
```

```
'asp_addition_at_GAU': 12
```

Updating TranslationData.subreactions with the tRNA subreactions incorporates this information into the TranslationReaction

```
_____
```

0.000498399634202103*mu + 0.000195123456790123 + 0.00598079561042524*(mu + 0.3915)/mu RNA_a + 12 asp /home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:229 **UserWarning:** No element contribution input

Add Complex Formation Reaction

Add ComplexData to model

For COBRAme models, the reaction gene-protein-reaction rule (GPR) is replaced with a metabolite representing the synthesis of the enzyme(s) catalyzing a reaction. This metabolite is formed explicitly in a ME model by separate reaction to transcribe the gene(s) and translate the protein(s) the compose the complex.

class cobrame.core.processdata.**ComplexData**(*id*, *model*)

Contains all information associated with the formation of an functional enzyme complex.

This can include any enzyme complex modifications required for the enzyme to become active.

Parameters

- id (str) Identifier of the complex data. As a best practice, this should typically use the same ID as the complex being formed. In cases with multiple ways to form complex '_ + alt' or similar suffixes can be used.
- model (cobrame.core.model.MEModel) ME-model that the ComplexData is associated with

stoichiometry

collections.DefaultDict(int) - Dictionary containing {protein_id: count} for all protein subunits comprising enzyme complex

subreactions

dict – Dictionary of {subreaction_data_id: count} for all complex formation subreactions/modifications. This can include cofactor/prosthetic group binding or enzyme side group addition.

Add ComplexFormation reaction to model

And point ComplexFormation to ComplexData .. autoclass:: cobrame.core.reaction.ComplexFormation :noindex:

```
In [27]: rxn = cobrame.ComplexFormation('formation_complex_ab')
        me.add_reaction(rxn)
        rxn.complex_data_id = data.id
        rxn._complex_id = data.id
```

Update ComplexFormation reaction

ComplexFormation.update(verbose=True)

Creates reaction using the associated complex data and adds chemical formula to complex metabolite product.

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for *cobrame.core.processdata.ComplexData*):

1.Complex product defined in self._complex_id

2. Protein subunits with stoichiometery defined in data.stoichiometry

- 3.Metabolites and enzymes w/ coupling coefficients defined in data.subreactions. This often includes enzyme complex modifications by coenzymes or prosthetic groups.
- 4.Biomass cobrame.core.component.Constraint corresponding to modifications detailed in data.subreactions, if any

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

```
In [28]: rxn.update(verbose=False)
    print(me.reactions.formation_complex_ab.reaction)
```

protein_a + protein_b --> complex_ab

Apply modification to complex formation reaction

Many enzyme complexes in an ME-model require cofactors or prosthetic groups in order to properly function. Information about such processes are stored as ModificationData.

For instance, we can add the modification of an iron-sulfur cluster, a common prosthetic group, by doing the following:

```
In [29]: # Define the stoichiometry of the modification
    mod_data = cobrame.SubreactionData('mod_2fe2s_c', me)
    mod_data.stoichiometry = {'2fe2s_c': -1}
    # this process can also be catalyzed by a chaperone
    mod_data.enzyme = 'complex_ba'
    mod_data.keff = 65. # default value
```

Associate modification to complex and update() its formation

Update ComplexFormation reaction

```
In [31]: print('Before adding modification')
    print('-------')
    print(me.reactions.formation_complex_ab.reaction)
    me.reactions.formation_complex_ab.update()
    print('\n')
    print('After adding modification')
    print('-------')
    print(me.reactions.formation_complex_ab.reaction)
```

Before adding modification

```
protein_a + protein_b --> complex_ab
Created <Complex complex_ba at 0x7f4f3b7aac90> in <ComplexFormation formation_complex_ab at 0x7f4ef0a
```

After adding modification

```
2fe2s_c + 4.27350427350427e-6*mu complex_ba + protein_a + protein_b --> complex_ab + 0.17582 prosther
```

Add Metabolic Reaction

Add StoichiometricData to model

MetabolicReactions require, at a minimum, one corresponding StoichiometricData. StoichiometricData essentially holds the information contained in an M-model reaction. This includes the metabolite stoichiometry and the upper and lower bound of the reaction. As a best practice, StoichiometricData typically uses an ID equivalent to the M-model reaction ID.

So first, we will create a StoichiometricData object to define the stoichiometry of the conversion of *a* to *b*. Only one StoichiometricData object should be created for both reversible and irreversible reactions

class cobrame.core.processdata.**StoichiometricData** (*id*, *model*) Encodes the stoichiometry for a metabolic reaction.

StoichiometricData defines the metabolite stoichiometry and upper/lower bounds of metabolic reaction

Parameters

- id (*str*) Identifier of the metabolic reaction. Should be identical to the M-model reactions in most cases.
- model (cobrame.core.model.MEModel) ME-model that the StoichiometricData is associated with

```
_stoichiometry
```

dict – Dictionary of {metabolite_id: stoichiometry} for reaction

subreactions

collections.DefaultDict(int) - Cases where multiple enzymes (often carriers ie. Acyl Carrier Protein) are involved in a metabolic reactions.

upper_bound

int – Upper reaction bound of metabolic reaction. Should be identical to the M-model reactions in most cases.

lower_bound

int – Lower reaction bound of metabolic reaction. Should be identical to the M-model reactions in most cases.

Add MetabolicReaction to model

The StoichiometricData for this reversible reaction is then assigned to two different MetabolicReactions (Due to the enzyme dilution constraint, all enzyme catalyzed reactions must be reversible; more on this later). The MetabolicReactions require: - The associated StoichiometricData - The *reverse* flag set to True for reverse reactions, False for forward reactions - Enzyme K_{eff} for reaction (discussed later, dafault=65)

These fields are then processed and the actual model reaction is created using the MetabolicReaction's update() function

class cobrame.core.reaction.MetabolicReaction (id)

Irreversible metabolic reaction including required enzymatic complex

This reaction class's update function processes the information contained in the complex data for the enzyme that catalyzes this reaction as well as the stoichiometric data which contains the stoichiometry of the metabolic conversion being performed (i.e. the stoichiometry of the M-model reaction analog)

```
Parameters id (str) – Identifier of the metabolic reaction. As a best practice, this ID should use the following template (FWD=forward, REV=reverse): "<StoichiometricData.id> + _ + <FWD or REV> + _ + <Complex.id>"
```

keff

float – The turnover rete (keff) couples enzymatic dilution to metabolic flux

reverse

boolean – If True, the reaction corresponds to the reverse direction of the reaction. This is necessary since all reversible enzymatic reactions in an ME-model are broken into two irreversible reactions

```
In [33]: # Create a forward ME Metabolic Reaction and associate the stoichiometric data to it
rxn_fwd = cobrame.MetabolicReaction('a_to_b_FWD_complex_ab')
me.add_reaction(rxn_fwd)
rxn_fwd.stoichiometric_data = data
rxn_fwd.reverse = False
rxn_fwd.keff = 65.
# Create a reverse ME Metabolic Reaction and associate the stoichiometric data to it
rxn_rev = cobrame.MetabolicReaction('a_to_b_REV_complex_ab')
me.add_reaction(rxn_rev)
rxn_rev.stoichiometric_data = data
rxn_rev.reverse = True
rxn_rev.keff = 65.
```

Update MetabolicReactions

MetabolicReaction.update(verbose=True)

Creates reaction using the associated stoichiometric data and complex data.

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.StoichiometricData):

1.Complex w/ coupling coefficients defined in self.complex_data.id and self.keff

2.Metabolite stoichiometry defined in data.stoichiometry. Sign is flipped if self.reverse == True

Also sets the lower and upper bounds based on self.reverse and data.upper_bound and data.lower_bound.

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

Note: the k_{eff} and complex_ab is not included in the reaction since no complex has been associated to it yet

Associate enzyme with MetabolicReaction

The ComplexData object created in the previous cell can be incorporated into the MetabolicReaction using code below.

Note: the update() function is required to apply the change.

```
In [35]: data = me.process_data.complex_ab
        me.reactions.a_to_b_FWD_complex_ab.complex_data = data
        print('Forward reaction (before update): %s' %
               (me.reactions.a_to_b_FWD_complex_ab.reaction))
         me.reactions.a_to_b_FWD_complex_ab.update()
         print('Forward reaction (after update): %s' %
               (me.reactions.a_to_b_FWD_complex_ab.reaction))
         print('')
        me.reactions.a_to_b_REV_complex_ab.complex_data = data
        print('Reverse reaction (before update): %s' %
               (me.reactions.a_to_b_REV_complex_ab.reaction))
         me.reactions.a_to_b_REV_complex_ab.update()
        print('Reverse reaction (after update): %s' %
               (me.reactions.a_to_b_REV_complex_ab.reaction))
Forward reaction (before update): a --> b
Forward reaction (after update): a + 4.27350427350427e-6*mu complex_ab --> b
Reverse reaction (before update): b \rightarrow a
Reverse reaction (after update): b + 4.27350427350427e-6*mu complex_ab --> a
```

The coefficient for complex_ab is determined by the expression

$$\frac{\mu}{k_{eff}}$$

which in its entirety represents the dilution of an enzyme following a cell doubling. The coupling constraint can be summarized as followed

$$v_{dilution,j} = \mu \sum_{i} \left(\frac{1}{k_{eff,i}} v_{usage,i} \right), \qquad \forall j \in Enzyme \qquad (3.6)$$

Where

- $v_{usage,i}$ is the flux through the metabolic reaction
- k_{eff} is the turnover rate for the process and conveys the productivity of the enzyme complex. Physically, it can be thought of as the number of reactions the enzyme can catalyze per cell division.

By default the k_{eff} for a MetabolicReaction is set to 65 but this can be changed using the code below.

Different Keff for forward reaction

Adding Reactions using utility functions

Add reactions using some of the utility functions provided in cobrame.util.building.py

Transcription

Using the utility functions to create the TranscribedGene metabolite has the advantage of forcing the assignment of sequence, strand and RNA_type.

Creates a TranscribedGene metabolite object and adds it to the ME-model

Parameters

• **me_model** (*cobrame.core.model.MEModel*) – The MEModel object to which the reaction will be added

- **locus_id** (*str*) Locus ID of RNA product. The TranscribedGene will be added as "RNA + _ + locus_id"
- left_pos (int or None) Left position of gene on the sequence of the (+) strain
- right_pos (int or None) Right position of gene on the sequence of the (+) strain
- **seq** (*str*) Nucleotide sequence of RNA product. Amino acid sequence, codon counts, etc. will be calculated based on this string.
- strand(str or None)-
 - (+) if the RNA product is on the leading strand
 - (-) if the RNA product is on the complementary strand
- rna_type (str) Type of RNA of the product. tRNA, rRNA, or mRNA Used for determining how RNA product will be processed.

Returns Metabolite object for the RNA product

Return type cobrame.core.component.TranscribedGene

```
In [37]: building.create_transcribed_gene(me, 'b','tRNA', 'ATCG')
    building.add_transcription_reaction(me, 'TU_b', {'b'}, sequence)
    print(me.reactions.transcription_TU_b.reaction)
    me.reactions.transcription_TU_b.update()
```

```
86 atp_c + 38 ctp_c + 12 gtp_c + 182 h2o_c + 50 utp_c --> RNA_b + 85 amp_c + 37 cmp_c + 11 gmp_c + 14
```

Translation

add_translation_reaction assumes that the RNA and protein have the same locus_id. It creates the appropriate TranslationData and TranslationReaction instance, links the two together and updates the TranslationReaction.

cobrame.util.building.add_translation_reaction(me_model, locus_id, dna_sequence, up-

date = False)

Creates and adds a TranslationReaction to the ME-model as well as the associated TranslationData

A dna_sequence is required in order to add a TranslationReaction to the ME-model

Parameters

- **me_model** (cobra.core.model.MEModel) The MEModel object to which the reaction will be added
- **locus_id** (*str*) Locus ID of RNA product. The TranslationReaction will be added as "translation + _ + locus_id" The TranslationData will be added as "locus_id"
- **dna_sequence** (*str*) DNA sequence of the RNA product. This string should be reverse transcribed if it originates on the complement strand.
- update (bool) If True, use TranslationReaction's update function to update and add reaction stoichiometry

```
In [38]: building.add_translation_reaction(me, 'b', dna_sequence=sequence, update=True)
    print(me.reactions.translation_b.reaction)
```

```
0.000498399634202103*mu + 0.000195123456790123 + 0.00598079561042524*(mu + 0.3915)/mu RNA_b + 12 asp
```

Complex Formation

Alternatively, ComplexData has a create_complex_formation() function to create the sythesis reaction following the naming conventions. It contains an update() function which incorporates changes in the ComplexData

```
ComplexData.create_complex_formation(verbose=True)
creates a complex formation reaction
```

This assumes none exists already. Will create a reaction (prefixed by "formation") which forms the complex

Parameters verbose (bool) – If True, print if a metabolite is added to model during update

protein_a + protein_b --> complex_ba

Metabolic Reaction

```
cobrame.util.building.add_metabolic_reaction_to_model (me_model, stoichiomet-
ric_data_id, directionality,
complex_id=None, sponta-
neous=False, update=False,
keff=65)
```

Creates and add a MetabolicReaction to a MEModel.

Parameters

- **me_model** (*cobrame.core.model.MEModel*) **MEModel** that the MetabolicReaction will be added to
- stoichiometric_data_id (str) ID of the StoichiometricData for the reaction being added
- directionality (str) -
 - Forward: Add reaction that occurs in the forward direction
 - Reverse: Add reaction that occurs in the reverse direction
- **complex_id** (*str or None*) ID of the ComplexData for the enzyme that catalyze the reaction being added.
- spontaneous (bool) -
 - If True and complex_id=" add reaction as spontaneous reaction
 - If False and complex_id=" add reaction as orphan (CPLX_dummy catalyzed)

Created <Metabolite c at 0x7f4f3b812f10> in <MetabolicReaction b_to_c_FWD_complex_ab at 0x7f4ef0c7df Reaction b_to_c

b + 4.27350427350427e-6*mu complex_ab --> c

CHAPTER 4

Reaction Properties

The division of the ME-model into MEReaction and ProcessData classes allows the user to essentially have access to the entire database of information used to construct the model. The following will show how this can be leveraged to easily query, edit and update aspects of the model reactions.

```
In [1]: import pickle
        from collections import defaultdict
        from os.path import abspath, dirname, join
        import pandas as pd
        import cobra.test
        import cobrame
        import ecolime
/home/sbrg-cjlloyd/cobrapy/cobra/io/sbml3.py:24: UserWarning: Install lxml for faster SBML I/O
 warn("Install lxml for faster SBML I/O")
/home/sbrg-cjlloyd/cobrapy/cobra/io/__init__.py:12: UserWarning: cobra.io.sbml requires libsbml
 warn("cobra.io.sbml requires libsbml")
In [2]: # Load E. coli ME-model
        ecoli_dir = dirname(abspath(ecolime.__file__))
        model_dir = join(ecoli_dir, 'me_models/iJL1678b.pickle')
        with open(model_dir, 'rb') as f:
           me = pickle.load(f)
        # Load E. coli M-model
```

```
iJ01366 = cobra.test.create_test_model('ecoli')
```

Metabolic Reactions

These are the ME-model representations of all reactions in the metabolic reconstruction, in this case iJO1366

Through a MetabolicReaction, the user has direct access to the StoichiometricData, ComplexData and keff used to construct the reaction.

This reactions is formed using the StoichiometricData (E4PD) and ComplexData (GAPDH-A-CPLX) with kef

As a best practice, the ComplexData and StoichiometricData themselves should not be changed. If these need changed then a new MetabolicReaction should be created.

Edit keffs

Further, the keff is an attribute of the MetabolicReaction itself and not of the ComplexData changing the ComplexData will not affect the form of the MetabolicReaction.

Edit stoichiometry

Aspects of the StoichiometricData, however, can be changed. This includes: - Reaction stoichiometry - Reaction upper & lower bounds

Currently the stoichiometry is:

'h_c': 2.0, 'nad_c': -1.0, 'nadh_c': 1.0}

This can be updated to, for instance, translocate a hydrogen by performing the following

```
In [7]: stoich_data.stoichiometry['h_c'] = -1
    stoich_data.stoichiometry['h_p'] = 1
    rxn.update()
    print('%s: \n\t%s' % (rxn.id, rxn.reaction))
```

```
E4PD_FWD_GAPDH-A-CPLX:
4.27350427350427e-6*mu GAPDH-A-CPLX + e4p_c + h2o_c + h_c + nad_c --> 4per_c + h_p + nadh_c
```

This change can be usd to update both the forward and reverse reaction

A simpler approach is to updated the parent reactions for StoichiometricData. This will update any instances of the reaction catalyzed by an isozyme.

```
In [9]: stoich_data.stoichiometry['h_c'] = -2
    stoich_data.stoichiometry['h_p'] = 2
    for r in stoich_data.parent_reactions:
        r.update()
        print('$s: \n\t$s' % (r.id, r.reaction))
E4PD_FWD_GAPDH-A-CPLX:
        4.27350427350427e-6*mu GAPDH-A-CPLX + e4p_c + h2o_c + 2.0 h_c + nad_c --> 4per_c + 2.0 h_p +
E4PD_REV_ERYTH4PDEHYDROG-CPLX:
        4per_c + 3.09490345954754e-6*mu ERYTH4PDEHYDROG-CPLX + 2.0 h_p + nadh_c --> e4p_c + h2o_c + 2.0 h_c
E4PD_REV_GAPDH-A-CPLX:
        4per_c + 3.20942472231275e-6*mu GAPDH-A-CPLX + 2.0 h_p + nadh_c --> e4p_c + h2o_c + 2.0 h_c
E4PD_FWD_ERYTH4PDEHYDROG-CPLX:
        0.0135143204065698*mu ERYTH4PDEHYDROG-CPLX + e4p_c + h2o_c + 2.0 h_c + nad_c --> 4per_c + 2.0
```

Edit upper and lower reaction bounds

The upper and lower bounds can be edited through the stoichiometric data and updated to the metabolic reaction

Important: do not change the upper and lower bounds of a MetabolicReaction directly. If this is done than the change will be overwritten when the update function is ran (shown below)

```
In [10]: rxn.lower_bound = -1000
    print('Lower bound = %d' %rxn.lower_bound)
    rxn.update()
    print('Lower bound = %d' %rxn.lower_bound)
Lower bound = -1000
Lower bound = 0
```

Editing the reaction bounds of the StoichiometricData, however, will edit the bounds of the forward and reverse reaction, as well as any instances of the reaction catalyzed by isozymes

```
In [11]: stoich_data.lower_bound = 0.
    print('Upper Bounds\n------')
    for r in stoich_data.parent_reactions:
        direction = 'Forward' if r.reverse is False else 'Reverse'
```

print('%s Before Update \n\t%s: %s' % (direction, r.id, r.upper_bound)) r.update() print('%s After Update \n\t%s: %s' % (direction, r.id, r.upper_bound)) Upper Bounds _____ Forward Before Update E4PD_FWD_GAPDH-A-CPLX: 1000.0 Forward After Update E4PD_FWD_GAPDH-A-CPLX: 1000.0 Reverse Before Update E4PD_REV_ERYTH4PDEHYDROG-CPLX: 1000.0 Reverse After Update E4PD REV ERYTH4PDEHYDROG-CPLX: 0 Reverse Before Update E4PD_REV_GAPDH-A-CPLX: 1000.0 Reverse After Update E4PD_REV_GAPDH-A-CPLX: 0 Forward Before Update E4PD FWD ERYTH4PDEHYDROG-CPLX: 1000.0 Forward After Update E4PD_FWD_ERYTH4PDEHYDROG-CPLX: 1000.0

Transcription Reactions

TranscribedGene (RNA) metabolite properties

Transciption occurs via operons contained within the organisms genome or transcription unit (TU). This means that often, a transcribed region will code for multiple RNAs. The E. coli ME-model has 4 possible RNA types that can be transcribed: - mRNA - tRNA - ncRNA (noncoding RNA)

mRNAs can then translated directly from the full transcribed TU, while rRNA, tRNA and ncRNA are spliced out of the TU by endonucleases. In these cases, in order to know which bases need excized, the RNA metabolites (TranscribedGene) themselves have to store information such as: - **DNA strand, left and right genome position** to identify which TU the RNA is a part of - **RNA type** to determine whether it needs excised from the TU - **nucleotide sequence** to determine bases that do/do not need excised if not mRNA and the RNA mass for biomass constraint

An example of a TranscribedGene's attributes is shown below

```
left_pos 3341965
nucleotide_sequence ATGGCAACATTAACTGCAAAGAACCTTGCAAAAGCCTATAAAGGCC...
right_pos 3342691
strand +
```

TranscriptionReaction/TranscriptionData properties

Each TranscriptionReaction in a COBRAme ME-model is associated with exactly one TranscriptionData which includes everything necessary to define a reaction. This includes: - **subreactions** To handle enzymatic processes not performed by RNA polymerase - **RNA Polymerase** Different RNA polymerase metabolite for different sigma factors - **RNA Products** TUs often contain more than one RNA in sequence - **Nucleotide sequence**

The TranscriptionData for TU containing the gene above is shown below

```
In [14]: rxn = me.reactions.transcription_TU_8398_from_RPOE_MONOMER
         data = rxn.transcription_data
         pd.DataFrame({i: str(v) for i, v in data.__dict__.items()}, index=['Atribute Values']).T
Out[14]: Atribute Values
         RNA_polymerase
                                                                      RNAPE-CPLX
         RNA_products
                                                      {'RNA_b3201', 'RNA_b3202'}
                                                                     iJL1678b-ME
        _model
        _parent_reactions
                                    { 'transcription_TU_8398_from_RPOE_MONOMER' }
                                                       TU_8398_from_RPOE_MONOMER
         id
         nucleotide_sequence ACAAACTCAGCCTTAATCTTGTGCTTGCCAGCTCACTTCTGGCCGC...
         subreactions.
                              defaultdict(<class 'int'>, {'Transcription_nor...
```

This reaction currently uses a subreaction called *Transcription_normal_rho_dependent* to account for the elongation factors etc. associated with transcription. This TU also requires a rho factor to terminate transcription. These complexes can be removed from the reaction by running the following

```
0.0218585689888099*mu + 0.00855762975911906 RNAPE-CPLX + 1017 atp_c + 1181 ctp_c + 1190 gtp_c + 1186
```

This poses a problem where, if RNA_b3201 and RNA_b3202 are not required in equal amounts, the model will become infeasible. To accound for this, all RNAs have a demand reaction associated with them. *mRNA_biomass* is consumed for each demand reaction with a coefficient equal to the molecular weight of each RNA (in kDa). This prevents the model from overproducing RNA to increase biomass production, and therefore growth rate, in some instances. More on the implications of the *biomass* constraint can be found in **ME-Model Fundamentals**

```
In [16]: for rna in data.RNA_products:
    r = me.reactions.get_by_id('DM_' + rna)
    print('%s: %s' % (r.id, r.reaction))
DM_RNA_b3201: RNA_b3201 + 232.671391 mRNA_biomass -->
DM_RNA_b3202: RNA_b3202 + 459.03124199999996 mRNA_biomass -->
```

As is, this reaction produces two mRNAs so no nucleotides are excised. If one or both is changed to a stable RNA (rRNA, tRNA or ncRNA) bases will be excised.

```
In [17]: me.metabolites.RNA_b3202.RNA_type = 'rRNA'
    for r in data.parent_reactions:
        r.update()
        print(r.reaction)
0.0218585689888099*mu + 0.00855762975911906 RNAPE-CPLX + 1017 atp_c + 1181 ctp_c + 1190 gtp_c + 2414
```

Changing RNA_b3202 to an rRNA and updating the transcription reaction causes both of the RNAs to now be excised from the TU, as indicated by the nucleotide monophosphates that appear in the products. This is not a complete picture because this process is catalyzes by an endonuclease, whose activity can be incorporated as ModificationData. Updating the reaction after adding these processes incorporates

Translation Reactions

TranslatedGene (Protein) metabolite properties

For COBRAme ME-models, proteins are translated directly from mRNA metabolites not from TUs. This means that all information required to construct a TranslationReaction can be found in its TranslationData therefore no extra information is contained in a TranslatedGene object.

TranslationReaction/TranslationData properties

Each TranslationReaction in a COBRAme ME-model is associated with exactly one TranslationData which includes everything necessary to define the reaction. This includes: - **subreactions** To handle enzymatic processes not performed by ribosome and incorporate tRNAs - **mRNA** ID of mRAN being translated - **term_enzyme** Enzyme that catalyzes translation termination - **Nucleotide sequence**

The TranslationData for a TranslationReaction is shown below

```
id b2020
mRNA RNA_b2020
nucleotide_sequence ATGAGCTTTAACACAATCATTGACTGGAATAGCTGTACTGCGGAGC...
protein protein_b2020
subreactions defaultdict(<class 'int'>, {'met_addition_at_A...
```

The rest of the information required to define a translation reaction can be dynamically computed from these attributes.

For example, the amino acid sequence is calculated from the nucleotide sequence with:

```
In [21]: str(data.amino_acid_count)
```

```
Out[21]: "defaultdict(<class 'int'>, {'met_L_c': 7, 'ser_L_c': 34, 'phe_L_c': 12, 'asn_L_c': 13,
```

The codon count from the sequence can be used to determine the subreaction required for charged tRNA-mediated amino acid addition.

```
In [22]: str(data.subreactions_from_sequence)
```

```
Out[22]: "{'met_addition_at_AUG': 6, 'ser_addition_at_AGC': 12, 'phe_addition_at_UUU': 6, 'asn_addit:
```

Changing the nucleotide sequence will automically update these values.

```
In [23]: data.nucleotide_sequence = 'ATGAGCTTTAAC'
    print('Elongation Subreactions')
    print(str(data.subreactions_from_sequence))
    print('\nOne of each start subreaction')
    print(str(data.add_initiation_subreactions()))
    print('\nNo termination subreaction (AAC) not valid stop codon')
    print(str(data.add_termination_subreactions()))
Elongation Subreactions
'ser_addition_at_AGC': 1, 'phe_addition_at_UUU': 1
One of each start subreaction
None
No termination subreaction (AAC) not valid stop codon
None
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:888 UserWarning: RNA_b2020 starts with 'AUG' y
/home/sbrg-cjlloyd/cobrame/cobrame/core/processdata.py:925 UserWarning: No termination enzyme for RNA
```

Stop codons are defined for the organism beforehand. Changing the sequence to a valid stop codon corrects this

```
In [24]: data.nucleotide_sequence = 'ATGAGCTTTTAA'
    print('Termination subreaction')
    print(str(data.add_termination_subreactions()))
```

Termination subreaction None

/home/sbrg-cjlloyd/cobrame/core/processdata.py:925 UserWarning: No termination enzyme for RNA

ComplexFormation Reactions

some complexes are modified in a metabolic process (e.g. Acyl Carrier Protein sidechain reactions) ### Complex Metabolite Properties Like TranslatedGenes, Complex metabolites do not need to store any additional information. ### ComplexFormation / ComplexData Properties Each ComplexFormation reaction in a COBRAme ME-model is associated with exactly one ComplexData which includes everything necessary to define the reaction. This includes: - **modification** To define the modifications by prosthetic groups or cofactors that can be required for the complex to catalyze cellular processes - **stoichiometry** Stoichiometry of the protein subunits

The ComplexData for a ComplexFormation reaction is shown below

This complex has two modification mod_lipo_c and mod_mg_c. You can view the properties of these modifications by accessing their ModificationData objects.

```
In [27]: for mod in data.subreactions:
    mod_data = me.process_data.get_by_id(mod)
    print(mod_data.id)
    for key, value in mod_data.__dict__.items():
        if not key.startswith('_') and value:
            print('\t', key, value)
mod_mg2_c
        id mod_mg2_c
        stoichiometry 'mg2_c': -1
        keff 65.0
mod_lipo_c
        id mod_lipo_c
        stoichiometry 'lipoamp_c': -1, 'amp_c': 1, 'h_c': 2
        enzyme EG11796-MONOMER
        keff 65.0
```

The information in the ModificationData and ComplexData are assembled in the ComplexFormation reaction shown below

```
In [28]: print(rxn.reaction)
4.27350427350427e-6*mu EG11796-MONOMER + lipoamp_c + mg2_c + 2.0 protein_b0116 + 12.0 protein_b0726 -
```

CHAPTER 5

cobrame package

Subpackages

cobrame.core package

Submodules

cobrame.core.reaction module

```
class cobrame.core.reaction.ComplexFormation(id)
    Bases: cobrame.core.reaction.MEReaction
```

Formation of a functioning enzyme complex that can act as a catalyst for a ME-model reaction.

This reaction class produces a reaction that combines the protein subunits and adds any coenyzmes, prosthetic groups or enzyme modifications to form complete enzyme complex.

Parameters id (*str*) – Identifier of the complex formation reaction. As a best practice, this ID should be prefixed with 'formation + _ + <complex_id>'. If there are multiple ways of producing complex, this can be suffixed with '_ + alt'

_complex_id

str - Name of the complex being produced by the complex formation reaction

complex_data_id

str – Name of ComplexData that defines the subunit stoichiometry or subreactions (modifications). This will not always be the same as the _complex_id. Sometimes complexes can be modified using different processes/enzymes

complex

Get the metabolite product of the complex formation reaction

Returns Instance of complex metabolite from self._complex_id

Return type cobrame.core.component.Complex

update (verbose=True)

Creates reaction using the associated complex data and adds chemical formula to complex metabolite product.

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.ComplexData):

- 1.Complex product defined in self._complex_id
- 2. Protein subunits with stoichiometery defined in data.stoichiometry
- 3.Metabolites and enzymes w/ coupling coefficients defined in data.subreactions. This often includes enzyme complex modifications by coenzymes or prosthetic groups.
- 4.Biomass cobrame.core.component.Constraint corresponding to modifications detailed in data.subreactions, if any

Parameters verbose (*bool*) – Prints when new metabolites are added to the model when executing update()

class cobrame.core.reaction.GenericFormationReaction(id)
 Bases: cobrame.core.reaction.MEReaction

Some components in an ME-model can perform exactly the same function. To handle this, GenericFormation-Reactions are used to create generic forms of these components.

Parameters id (*str*) – Identifier of the generic formation reaction. As a best practice, this ID should be prefixed with 'metabolite_id + _to_ + generic_metabolite_id'

class cobrame.core.reaction.MEReaction(id=None, name='')
Bases: cobra.core.Reaction.Reaction

MEReaction is a general reaction class from which all ME-Model reactions will inherit

This class contains functionality that can be used by all ME-model reactions

Parameters id (*str*) – Identifier of the MEReaction. Should follow best practices of child class

add_biomass_from_subreactions (process_data, biomass=0.0)

Account for the biomass of metabolites added to macromolecule (protein, complex, etc.) due to a modification such as prosthetic group addition.

Parameters

- process_data (cobrame.core.processdata.ProcessData) ProcessData that is used to construct MEReaction
- **biomass** (float) Initial biomass value in kDa

Returns Initial biomass value + biomass added from subreactions in kDa

Return type float

add_subreactions (process_data_id, stoichiometry, scale=1.0)

Function to add subreaction process data to reaction stoichiometry

Parameters

• **process_data_id** (*str*) – ID of the process data associated with the metabolic reaction.

For example, if the modifications are being added to a complex formation reaction, the process data id would be the name of the complex.

- stoichiometry (dict) Dictionary of {metabolite_id: float} or {metabolite_id: float
 * (sympy.Symbol)}
- **scale** (*float*) Some processes (ie. tRNA charging) are reformulated such that other involved metabolites need scaling

Returns Stoichiometry dictionary with updated entries

Return type dict

check_me_mass_balance()

Checks the mass balance of ME reaction, ignoring charge balances

Returns {element: number_of_elemental_imbalances}

Return type dict

clear_metabolites()

Remove all metabolites from the reaction

 get_components_from_ids (id_stoichiometry, brame.core.component.Metabolite'>, verbose=True)

 Function to convert stoichiometry dictionary entries from strings to cobra objects.

 'co-

{metabolite_id: value} to {cobrame.core.component.Metabolite: value}

Parameters

- id_stoichiometry (Dict {string: float}) Input Dict of {metabolite_id: value}
- **default_type** (*String*) The type of cobra.Metabolite to default to if the metabolite is not yet present in the model
- **verbose** (*Boolean*) If True, print metabolites added to model if not yet present in model

Returns { cobrame.core.component.Metabolite: float }

Return type dict

objective_coefficient

Get and set objective coefficient of reaction

Overrides method in parent class in order to enable use of optlang interfaces.

Returns Objective coefficient of reaction

Return type float

class cobrame.core.reaction.MetabolicReaction (id)

Bases: cobrame.core.reaction.MEReaction

Irreversible metabolic reaction including required enzymatic complex

This reaction class's update function processes the information contained in the complex data for the enzyme that catalyzes this reaction as well as the stoichiometric data which contains the stoichiometry of the metabolic conversion being performed (i.e. the stoichiometry of the M-model reaction analog)

Parameters id (*str*) – Identifier of the metabolic reaction. As a best practice, this ID should use the following template (FWD=forward, REV=reverse): "<StoichiometricData.id> + _ + <FWD or REV> + _ + <Complex.id>"

keff

float - The turnover rete (keff) couples enzymatic dilution to metabolic flux

reverse

boolean – If True, the reaction corresponds to the reverse direction of the reaction. This is necessary since all reversible enzymatic reactions in an ME-model are broken into two irreversible reactions

complex_data

Get or set the ComplexData instance that details the enzyme that catalyzes the metabolic reaction. Can be set with instance of ComplexData or with its id.

Returns Complex data detailing enzyme that catalyzes this reaction

Return type cobrame.core.processdata.ComplexData

stoichiometric_data

Get or set the StoichiometricData instance that details the metabolic conversion of the metabolic reaction. Can be set with instance of StoichiometricData or with its id.

Returns Stoichiometric data detailing enzyme that catalyzes this reaction

Return type :class: cobrame.core.processdata.StoichiometricData '

update (verbose=True)

Creates reaction using the associated stoichiometric data and complex data.

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.StoichiometricData):

1.Complex w/ coupling coefficients defined in self.complex_data.id and self.keff

2.Metabolite stoichiometry defined in data.stoichiometry. Sign is flipped if self.reverse == True

Also sets the lower and upper bounds based on self.reverse and data.upper_bound and data.lower_bound.

Parameters verbose (*bool*) – Prints when new metabolites are added to the model when executing update()

class cobrame.core.reaction.PostTranslationReaction(id)

Bases: cobrame.core.reaction.MEReaction

Reaction class that includes all posttranslational modification reactions (translocation, protein folding, modification (for lipoproteins) etc)

There are often multiple different reactions/enzymes that can accomplish the same modification/function. In order to account for these and maintain one translation reaction per protein, these processes need to be modeled as separate reactions.

Parameters id (str) – Identifier of the post translation reaction

add_translocation_pathways (process_data_id, protein_id, stoichiometry=None) Add complexes and metabolites required to translocate the protein into cell membranes.

Parameters

- process_data_id (str) ID of translocation data defining post translation reaction
- protein_id (str) ID of protein being translocated via post translation reaction
- **stoichiometry** (*dict*) Dictionary of {metabolite_id: float} or {metabolite_id: float * (sympy.Symbol)}

Returns Stoichiometry dictionary with updated entries from translocation

Return type dict

posttranslation_data

Get or set PostTranslationData that defines the type of post translation modification/process (folding/translocation) that the reaction accounts for. Can be set with instance of PostTranslationData or with its id.

Returns The PostTranslationData that defines the PostTranslationReaction

Return type cobrame.core.processdata.PostTranslationData

update (verbose=True)

Creates reaction using the associated posttranslation data and adds chemical formula to processed protein product

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.PostTranslationData):

1. Processed protein product defined in data.processed_protein_id

2.Unprocessed protein reactant defined in data.unprocessed_protein_id

3. Metabolites and enzymes defined in data. subreactions

- 4. Translocation pathways defined in data.translocation
- 5.Folding mechanism defined in data.folding_mechanims w/ coupling coefficients defined in data.keq_folding, data.k_folding, model.global_info['temperature'], data.aggregation_propensity, and data.propensity_scaling
- 6.Surface area constraints defined in data.surface_are

7.Biomass if a significant chemical modification takes place (i.e. lipid modifications for lipoproteins)

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

```
class cobrame.core.reaction.SummaryVariable(id=None)
Bases: cobrame.core.reaction.MEReaction
```

Summary Variables are reactions that impose global constraints on the model.

The primary example of this is the biomass_dilution SummaryVariable which forces the rate of biomass production of macromolecules, etc. to be equal to the rate of their dilution to daughter cells during growth.

Parameters id (*str*) – Identifier of the Summary Variable

class cobrame.core.reaction.TranscriptionReaction(id)
 Bases: cobrame.core.reaction.MEReaction

Transcription of a TU to produced TranscribedGene.

RNA is transcribed on a transcription unit (TU) level. This type of reaction produces all of the RNAs contained within a TU, as well as accounts for the splicing/excision of RNA between tRNAs and rRNAs. The appropriate RNA_biomass constrain is produced based on the molecular weight of the RNAs being transcribed

Parameters id(str) – Identifier of the transcription reaction. As a best practice, this ID should be prefixed with 'transcription + _'

transcription_data

Get or set the *cobrame.core.processdata.TranscriptionData* that defines the transcription unit architecture and the features of the RNAs being transcribed.

```
update (verbose=True)
```

Creates reaction using the associated transcription data and adds chemical formula to RNA products

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for *cobrame.core.processdata.TranscriptionData*):

1.RNA_polymerase from data.RNA_polymerase w/ coupling coefficient (if present)

2.RNA products defined in data.RNA_products

- 3.Nucleotide reactants defined in data.nucleotide_counts
- 4.If tRNA or rRNA contained in data.RNA_types, excised base products
- 5.Metabolites + enzymes w/ coupling coefficients defined in data.subreactions (if present)
- 6.Biomass cobrame.core.component.Constraint corresponding to data.RNA_products and their associated masses

Parameters verbose (bool) – Prints when new metabolites are added to the model when executing update()

class cobrame.core.reaction.TranslationReaction(id)

Bases: cobrame.core.reaction.MEReaction

Reaction class for the translation of a TranscribedGene to a TranslatedGene

Parameters id (str) – Identifier of the translation reaction. As a best practice, this ID should be prefixed with 'translation + _'

translation_data

Get and set the cobra.core.processdata.TranslationData that defines the translation of the gene. Can be set with instance of TranslationData or with its id.

Returns

Return type cobra.core.processdata.TranslationData

update (verbose=True)

Creates reaction using the associated translation data and adds chemical formula to protein product

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for cobrame.core.processdata.TranslationData):

- 1.Amino acids defined in data.amino_acid_sequence. Subtracting water to account for condensation reactions during polymerization
- 2.Ribosome w/ translation coupling coefficient (if present)
- 3.mRNA defined in data.mRNA w/ translation coupling coefficient
- 4.mRNA + nucleotides + hydrolysis ATP cost w/ degradation coupling coefficient (if kdeg (defined in model.global_info) > 0)
- 5.RNA_degradosome w/ degradation coupling coefficient (if present and kdeg > 0)
- 6.Protein product defined in data.protein
- 7. Subreactions defined in data.subreactions
- 8.protein_biomass cobrame.core.component.Constraint corresponding to the protein product's mass
- 9.Subtract mRNA_biomass cobrame.core.component.Constraint defined by mRNA degradation coupling coefficinet (if kdeg > 0)
 - **Parameters verbose** (*bool*) Prints when new metabolites are added to the model when executing update()

class cobrame.core.reaction.tRNAChargingReaction (id)

Bases: cobrame.core.reaction.MEReaction

Reaction class for the charging of a tRNA with an amino acid

Parameters id (*str*) – Identifier for the charging reaction. As a best practice, ID should follow the template "charging_tRNA + _ + <tRNA_locus> + _ + <codon>". If tRNA initiates translation, <codon> should be replaced with START.

tRNA_data

Get and set the cobra.core.processdata.tRNAData that defines the translation of the gene. Can be set with instance of tRNAData or with its id.

Returns

Return type cobra.core.processdata.tRNAData

update (verbose=True)

Creates reaction using the associated tRNA data

This function adds the following components to the reaction stoichiometry (using 'data' as shorthand for *cobrame.core.processdata.tRNAData*):

- 1.Charged tRNA product following template: "generic_tRNA + _ + <data.codon> + _ + <data.amino_acid>"
- 2.tRNA metabolite (defined in data.RNA) w/ charging coupling coefficient
- 3. Charged amino acid (defined in data.amino_acid) w/ charging coupling coefficient
- 5.Synthetase (defined in data.synthetase) w/ synthetase coupling coefficient found, in part, using data.synthetase_keff
- 6.Post transcriptional modifications defined in data.subreactions
- **Parameters verbose** (bool) Prints when new metabolites are added to the model when executing update()

cobrame.core.processdata module

class cobrame.core.processdata.ComplexData(id, model)
Bases: cobrame.core.processdata.ProcessData

Contains all information associated with the formation of an functional enzyme complex.

This can include any enzyme complex modifications required for the enzyme to become active.

Parameters

- id (*str*) Identifier of the complex data. As a best practice, this should typically use the same ID as the complex being formed. In cases with multiple ways to form complex '_+ alt' or similar suffixes can be used.
- **model** (*cobrame.core.model.MEModel*) ME-model that the ComplexData is associated with

stoichiometry

collections.DefaultDict(int) - Dictionary containing {protein_id: count} for all protein subunits comprising enzyme complex

subreactions

dict – Dictionary of {subreaction_data_id: count} for all complex formation subreactions/modifications. This can include cofactor/prosthetic group binding or enzyme side group addition.

complex

Get complex metabolite object

Returns Instance of complex metabolite that ComplexData is used to synthesize

Return type cobrame.core.component.Complex

complex_id

Get and set complex ID for product of complex formation reaction

There are cases where multiple equivalent processes can result in the same final complex. This allows the equivalent final complex_id to be queried. This only needs set in the above case

Returns ID of complex that ComplexData is used to synthesize

Return type str

create_complex_formation(verbose=True)

creates a complex formation reaction

This assumes none exists already. Will create a reaction (prefixed by "formation") which forms the complex

Parameters verbose (bool) – If True, print if a metabolite is added to model during update

formation

Get the formation reaction object

Returns Complex formation reaction detailed in ComplexData

Return type cobrame.core.reaction.ComplexFormation

class cobrame.core.processdata.GenericData (id, model, component_list)
Bases: cobrame.core.processdata.ProcessData

Class for storing information about generic metabolites

Parameters

- id (str) Identifier of the generic metabolite. As a best practice, this ID should be prefixed with 'generic + _'
- model (*cobrame.core.model.MEModel*) ME-model that the GenericData is associated with
- **component_list** (*list*) List of metabolite ids for all metabolites that can provide identical functionality

create_reactions()

Adds reaction with id "<metabolite_id> + _ + to + _ + <generic_id>" for each metabolite in self.component_list.

Creates generic metabolite and generic reaction, if they do not already exist.

class cobrame.core.processdata.PostTranslationData (id, model, processed_protein, prepro-

cessed_protein)

Bases: cobrame.core.processdata.ProcessData

Parameters

• id (*str*) – Identifier for post translation process.

- **model** (*cobrame.core.model.MEModel*) ME-model that the PostTranslationData is associated with
- processed_protein (str) ID of protein following post translational process
- preprocessed_protein (str) ID of protein before post translational process

translocation

set – Translocation pathways involved in post translation reaction.

Set of {cobrame.core.processdata.TranslocationData.id}

translocation_multipliers

dict - Some proteins require different coupling of translocation enzymes.

Dictionary of {cobrame.core.processdata.TranslocationData.id: float}

surface_area

dict – If protein is translated into the inner or outer membrane, the surface area the protein occupies can be accounted for as well.

Dictionary of {SA_+<inner_membrane or outer_membrane>: float}

subreactions

collections.DefaultDict(float) - If a protein is modified following translation, this is accounted for here

Dictionary of {subreaction_id: float}

biomass_type

str – If the subreactions add biomass to the translated gene, the biomass type (cobrame.core. compontent.Constraint.id) of the modification must be defined.

folding_mechanism

str - ID of folding mechanism for post translation reaction

aggregation_propensity

float – Aggregation propensity for the protein

keq_folding

dict - Temperature dependant keq for folding protein

Dictionary of {str(temperature): value}

k_folding

dict - Temperature dependant rate constant (k) for folding protein

Dictionary of {str(temperature): value}

propensity_scaling

float – Some small peptides are more likely to be folded by certain chaperones. This is accounted for using propensity_scaling.

class cobrame.core.processdata.ProcessData (id, model)

Bases: object

Generic class for storing information about a process

This class essentially acts as a database that contains all of the relevant information needed to construct a particular reaction. For example, to construct a transcription reaction, following information must be accessed in some way:

•nucleotide sequence of the transcription unit

•RNA_polymerase (w/ sigma factor)

•RNAs transcribed from transcription unit

•other processes involved in transcription of RNAs (splicing, etc.)

ME-model reactions are built from information in these objects.

Parameters

- id (*str*) Identifier of the ProcessData instance.
- **model** (*cobrame.core.model.MEModel*) ME-model that the ProcessData is associated with

model

Get the ME-model the process data is associated with

Returns ME-model that uses this process data

Return type class: cobrame.core.model.MEModel

parent_reactions

Get reactions that the ProcessData instance is used to construct.

Returns Parent reactions of ProcessData

Return type set

update_parent_reactions()

Executes the update() function for all reactions that the ProcessData instance is used to construct.

class cobrame.core.processdata.StoichiometricData(id, model)

Bases: cobrame.core.processdata.ProcessData

Encodes the stoichiometry for a metabolic reaction.

StoichiometricData defines the metabolite stoichiometry and upper/lower bounds of metabolic reaction

Parameters

- id (*str*) Identifier of the metabolic reaction. Should be identical to the M-model reactions in most cases.
- **model** (*cobrame.core.model.MEModel*) ME-model that the StoichiometricData is associated with

_stoichiometry

dict - Dictionary of {metabolite_id: stoichiometry} for reaction

subreactions

collections.DefaultDict(int) - Cases where multiple enzymes (often carriers ie. Acyl Carrier Protein) are involved in a metabolic reactions.

upper_bound

int – Upper reaction bound of metabolic reaction. Should be identical to the M-model reactions in most cases.

lower_bound

int – Lower reaction bound of metabolic reaction. Should be identical to the M-model reactions in most cases.

stoichiometry

Get or set metabolite stoichiometry for reaction.

Returns Dictionary of {metabolite_id: stoichiometry}

Return type dict

class cobrame.core.processdata.SubreactionData(id, model)
Bases: cobrame.core.processdata.ProcessData

Parameters

- id (str) Identifier of the subreaction data. As a best practice, if the subreaction data details a modification, the ID should be prefixed with "mod + _"
- model (*cobrame.core.model.MEModel*) ME-model that the SubreactionData is associated with

enzyme

list or str or None – List of cobrame.core.component.Complex.id s for enzymes that catalyze this process

or

String of single cobrame.core.component.Complex.id for enzyme that catalyzes this process

keff

float – Effective turnover rate of enzyme(s) in subreaction process

_element_contribution

dict – If subreaction adds a chemical moiety to a macromolecules via a modification or other means, net element contribution of the modification process should be accounted for. This can be used to mass balance check each of the individual processes.

Dictionary of {element: net_number_of_contributions}

calculate_biomass_contribution()

Calculate net biomass increase/decrease as a result of the subreaction process.

If subreaction adds a chemical moiety to a macromolecules via a modification or other means, the biomass contribution of the modification process should be accounted for and ultimately included in the reaction it is involved in.

Returns Mass of moiety transferred to macromolecule by subreaction

Return type float

calculate_element_contribution()

Calculate net contribution of chemical elements based on the stoichiometry of the subreaction data

Returns Dictionary of {element: net_number_of_contributions}

Return type dict

element_contribution

Get net contribution of elements from subreaction process to macromolecule

If subreaction adds a chemical moiety to a macromolecules via a modification or other means, net element contribution of the modification process should be accounted for. This can be used to mass balance check each of the individual processes.

Returns Dictionary of {element: net_number_of_contributions}

Return type dict

get_all_usages()

Get all process data that the subreaction is involved in

Yields cobrame.core.processdata.ProcessData – ProcessData that subreaction is involved in

get_complex_data()

Get the complex data that the subreaction is involved in

Yields cobrame.core.processdata.ComplexData - ComplexData that subreaction is involved in

class cobrame.core.processdata.TranscriptionData(id, model, rna_products=set([]))
Bases: cobrame.core.processdata.ProcessData

Class for storing information needed to define a transcription reaction

Parameters

- id (str) Identifier of the transcription unit, typically beginning with 'TU'
- model (*cobrame.core.model.MEModel*) ME-model that the TranscriptionData is associated with

nucleotide_sequence

str - String of base pair abbreviations for nucleotides contained in the transcription unit

RNA_products

set – IDs of *cobrame.core.component.TranscribedGene* that the transcription unit encodes. Each member should be prefixed with "RNA + _"

RNA_polymerase

str – ID of the *cobrame.core.component.RNAP* that transcribes the transcription unit. Different IDs are used for different sigma factors

subreactions

collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata. SubreactionData ID: num_usages} required for the transcription unit to be transcribed

RNA_types

Get generator consisting of the RNA type for each RNA product

Yields *str* – (mRNA, tRNA, rRNA, ncRNA)

codes_stable_rna

Get whether transcription unit codes for a stable RNA

Returns True if tRNA or rRNA in RNA products False if not

Return type bool

excised_bases

Get count of bases that are excised during transcription

If a stable RNA (e.g. tRNA or rRNA) is coded for in the transcription unit, the transcript must be spliced in order for these to function.

This determines whether the transcription unit requires splicing and, if so, returns the count of nucleotides within the transcription unit that are not accounted for in the RNA products, thus identifying the appropriate introns nucleotides.

Returns

{nucleotide_monophosphate_id: number_excised}

i.e. {"amp_c": 10, "gmp_c": 11, "ump_c": 9, "cmp_c": 11}

Return type dict

mass

Get mass of transcription unit in kDa from nucleotide sequence

If transcription unit is spliced, the mass of the excised bases are subtracted

Returns Mass of transcription unit in kDas

Return type float

nucleotide_count

Get count of each nucleotide contained in the nucleotide sequence

Returns {nuclotide_id: number_of_occurances}

Return type dict

class cobrame.core.processdata.TranslationData (id, model, mrna, protein)
Bases: cobrame.core.processdata.ProcessData

Class for storing information about a translation reaction.

Parameters

- id (str) Identifier of the gene being translated, typically the locus tag
- model (cobrame.core.model.MEModel) ME-model that the TranslationData is associated with
- mrna (str) ID of the mRNA that is being translated
- **protein** (*str*) ID of the protein product.

mRNA

str – ID of the mRNA that is being translated

protein

str – ID of the protein product.

subreactions

```
collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata.
SubreactionData.id: num_usages} required for the mRNA to be translated
```

nucleotide_sequence

str – String of base pair abbreviations for nucleotides contained in the gene being translated

add_elongation_subreactions (elongation_subreactions=set([]))

Add all subreactions involved in translation elongation.

This includes:

•tRNA activity subreactions returned with *subreactions_from_sequence()* which is called within this function.

•Elongation subreactions passed into this function. These will be added with a value of len(amino_acid_sequence) - 1 as these are involved in each amino acid addition

Some additional enzymatic processes are required for each amino acid addition during translation elongation

Parameters elongation_subreactions (*set*) – Subreactions that are required for each amino acid addition

add_initiation_subreactions (start_codons=set([]), start_subreactions=set([]))

Add all subreactions involved in translation initiation.

Parameters

• **start_codons** (*set*, *optional*) – Start codon sequences for the organism being modeled

• **start_subreactions** (*set*, *optional*) – Subreactions required to initiate translation, including the activity by the start tRNA

add_termination_subreactions(translation_terminator_dict=None)

Add all subreactions involved in translation termination.

Parameters translation_terminator_dict (dict or None) - {stop_codon : enzyme_id_of_terminator_enzyme}

amino_acid_count

Get number of each amino acid in the translated protein

Returns {amino_acid_id: number_of_occurrences}

Return type dict

amino_acid_sequence

Get amino acid sequence from mRNA's nucleotide sequence

Returns Amino acid sequence

Return type str

codon_count

Get the number of each codon contained within the gene sequence

Returns {codon_sequence: number_of_occurrences}

Return type dict

first_codon

Get the first codon contained in the mRNA sequence. This should correspond to the start codon for the gene.

Returns First 3 nucleotides comprising the first codon in the mRNA gene sequence

Return type str

last_codon

Get the last codon contained in the mRNA sequence. This should correspond to the stop codon for the gene.

Returns Last 3 nucleotides comprising the last codon in the mRNA gene sequence

Return type str

mass

Get mass of protein from amino acid composition

Returns Mass of protein (in kDa)

Return type float

subreactions_from_sequence

Get subreactions associated with each tRNA/AA addition.

tRNA activity is accounted for as subreactions. This returns the subreaction counts associated with each amino acid addition, based on the sequence of the mRNA.

Returns {cobrame.core.processdata.SubreactionData.id: num_usages}

Return type dict

class cobrame.core.processdata.TranslocationData(id, model)
 Bases: cobrame.core.processdata.ProcessData

Class for storing information about a protein translocation pathway

Parameters

- **id** (*str*) Identifier for translocation pathway.
- model (*cobrame.core.model.MEModel*) ME-model that the TranslocationData is associated with

keff

float - Effective turnover rate of the enzymes in the translocation pathway

enzyme_dict

dict – Dictionary containing enzyme specific information about the way it is coupled to protein translocation

{enzyme_id: {length_dependent: <True or False>, fixed_keff: <True or False>}}

length_dependent_energy

bool - True if the ATP cost of translocation is dependent on the length of the protein

stoichiometry

dict – Stoichiometry of translocation pathway, typically ATP/GTP hydrolysis

class cobrame.core.processdata.tRNAData(id, model, amino_acid, rna, codon)
Bases: cobrame.core.processdata.ProcessData

Class for storing information about a tRNA charging reaction.

Parameters

- id (*str*) Identifier for tRNA charging process. As best practice, this should be follow "tRNA + _ + <tRNA_locus> + _ + <codon>" template. If tRNA initiates translation, <codon> should be replaced with START.
- model (cobrame.core.model.MEModel) ME-model that the tRNAData is associated with
- **amino_acid** (*str*) Amino acid that the tRNA transfers to an peptide
- rna (str) ID of the uncharged tRNA metabolite. As a best practice, this ID should be prefixed with 'RNA + _'

subreactions

collections.DefaultDict(int) - Dictionary of {cobrame.core.processdata. SubreactionData.id: num_usages} required for the tRNA to be charged

synthetase

str - ID of the tRNA synthetase required to charge the tRNA with an amino acid

synthetase_keff

float - Effective turnover rate of the tRNA synthetase

cobrame.core.component module

class cobrame.core.component.**Complex**(*id*)

Bases: cobrame.core.component.MEComponent

Metabolite class for protein complexes

Parameters id (*str*) – Identifier of the protein complex.

mass

Get mass of complex based on protein subunit stoichiometries

Returns Mass of complex based on protein subunit composition in kDa

Return type int

metabolic_reactions

Get metabolic reactions catalyzed by complex

Returns List of cobrame.core.reaction.MetabolicReactions catalyzed by complex.

Return type list

class cobrame.core.component.Constraint (id)

Bases: cobrame.core.component.MEComponent

Metabolite class for global constraints such as biomass

Parameters id (str) – Identifier of the constraint

class cobrame.core.component.GenericComponent(id)

Bases: cobrame.core.component.MEComponent

Metabolite class for generic components created from cobrame.core.reaction. GenericFormationReaction

Parameters id (str) – Identifier of the generic tRNA. As a best practice should follow template: 'generic + _ + <generic metabolite id>'

class cobrame.core.component.GenerictRNA(id)
 Bases: cobrame.core.component.MEComponent

Metabolite class for generic tRNAs created from cobrame.core.reaction. tRNAChargingReaction

Parameters id (str) – Identifier of the generic tRNA. As a best practice should follow template: 'generic_tRNA + _ + <codon> + _ + <amino acid metabolite id>'

class cobrame.core.component.MeComponent (id) Bases: cobra.core.Metabolite.Metabolite

COBRAme component representation. Inherits from cobra.core.metabolite.Metabolite

Parameters id (str) – Identifier of the component. Should follow best practices of child classes

remove_from_me_model (method='subtractive')

Remove metabolite from me model along with any relevant *cobrame.core.processdata*. *ProcessData*

Parameters method (str) -

- destructive: remove metabolite from model and remove reactions it is involved in
- subtractive: remove only metabolite from model

class cobrame.core.component.Metabolite(id)
Bases: cobrame.core.component.MEComponent

COBRAme metabolite representation

Parameters id (str) – Identifier of the metabolite

class cobrame.core.component.ProcessedProtein(id, unprocessed_protein_id)

Bases: cobrame.core.component.MEComponent

Metabolite class for protein created from cobrame.core.reaction.PostTranslationReaction

Parameters

• **id** (*str*) – Identifier of the processed protein

• **unprocessed_protein_id** (*str*) – Identifier of protein before being processed by PostTranslationReaction

unprocessed_protein

Get unprocessed protein reactant in PostTranslationReaction

Returns Unprocessed protein object

Return type cobrame.core.component.TranslatedGene

unprocessed_protein_mass

Get mass of unprocessed protein

Returns Mass of unprocessed protein metabolite in kDa

Return type float

class cobrame.core.component.RNAP (*id*)

Bases: cobrame.core.component.Complex

Metabolite class for RNA polymerase complexes. Inherits from cobrame.core.component.Complex

Parameters id (*str*) – Identifier of the RNA Polymerase.

class cobrame.core.component.Ribosome (id)

Bases: cobrame.core.component.Complex

Metabolite class for Ribosome complexes. Inherits from cobrame.core.component.Complex

Parameters id (*str*) – Identifier of the Ribosome.

class cobrame.core.component.TranscribedGene(id, rna_type, nucleotide_sequence)
Bases: cobrame.core.component.MEComponent

Metabolite class for gene created from cobrame.core.reaction.TranscriptionReaction

Parameters

- id (str) Identifier of the transcribed gene. As a best practice, this ID should be prefixed with 'RNA + _'
- **rna_type** (*str*) Type of RNA encoded by gene sequence (mRNA, rRNA, tRNA, or ncRNA)
- **nucleotide_sequence** (*str*) String of base pair abbreviations for nucleotides contained in the gene

left_pos

int - Left position of gene on the sequence of the (+) strain

right_pos

int – Right position of gene on the sequence of the (+) strain

strand

str –

•(+) if the RNA product is on the leading strand

•(-) if the RNA product is on the comple(mentary strand

mass

Calculate mass of transcribed gene from nucleotide_sequence

Returns Mass of gene (in kDA)

Return type float

nucleotide_count

Get number of each nucleotide monophosphate

Returns {nucleotide_monophosphate_id: count}

Return type dict

class cobrame.core.component.TranslatedGene (id)

Bases: cobrame.core.component.MEComponent

Metabolite class for protein created from cobrame.core.reaction.TranslationReaction

Parameters id (*str*)-Identifier of the translated protein product. Should be prefixed with "protein + "

amino_acid_sequence

Get amino acid sequence of protein

Returns Amino acid sequence of protein

Return type str

complexes

Get the complexes that the protein forms

Returns List of cobrame.core.component.Complex s that the protein is a subunit of

Return type list

mass

Get mass of protein

Returns Mass of translated protein based on amino acid sequence in kDa

Return type float

metabolic_reactions

Get the mtabolic reactions that the protein helps catalyze

Returns List of cobrame.core.reactions.MetabolicReaction s that the protein helps catalyze

Return type list

translation_data

Get translation data that defines protein.

Assumes that TranslatedGene is "protein + _ + <translation data id>"

Returns Translation data used to form translation reaction of protein

Return type cobrame.core.processdata.TranslationData

cobrame.core.component.create_component(component_id, default_type=<class</pre>

brame.core.component.MEComponent'>,

rnap_set=set([]))

creates a component and attempts to set the correct type

cobra.core.model module

class cobrame.core.model.MEModel(*args)

 $Bases: \verb"cobra.core.Model.Model"$

```
add_biomass_constraints_to_model(biomass_types)
```

'co-

complex_data

- compute_solution_error(solution=None)
- **construct_attribute_vector** (*attr_name*, *growth_rate*) build a vector of a reaction attribute at a specific growth rate

Mainly used for upper and lower bounds

construct_s_matrix (growth_rate) build the stoichiometric matrix at a specific growth rate

gam

generic_data

get_metabolic_flux (solution=None)
 extract the flux state for metabolic reactions

- get_transcription_flux (solution=None)
 extract the transcription flux state
- get_translation_flux (solution=None)
 extract the translation flux state

ngam

posttranslation_data

```
prune (skip=None)
```

remove all unused metabolites and reactions

This should be run after the model is fully built. It will be difficult to add new content to the model once this has been run.

skip: list List of complexes/proteins/mRNAs/TUs to remain unpruned from model.

```
remove_genes_from_model (gene_list)
```

```
set_sasa_keffs(avg_keff)
```

stoichiometric_data

subreaction_data

tRNA_data

transcription_data

translation_data

translocation_data

unmodeled_protein

```
unmodeled_protein_biomass
```

unmodeled_protein_fraction

update()

updates all component reactions

Module contents

cobrame.util package

Submodules

cobrame.util.building module

cobrame.util.building.add_complex_to_model(me_model, complex id, complex_stoichiometry, complex_modifications=None)

Adds ComplexData to the model for a given complex.

Parameters

- me model (cobrame.core.model.MEModel) -
- complex_id (str) ID of the complex and thus the model ComplexData
- complex_stoichiometry (dict) {complex_id: {protein_<locus_tag>: stoichiometry}}
- **complex_modifications** (*dict*) {subreaction_id: stoichiometry}

cobrame.util.building.add_dummy_reactions (me_model, dna_seq, update=True) Add all reactions necessary to produce a dummy reaction catalyzed by "CPLX_dummy".

Parameters

- me model (cobrame.core.model.MEModel) The MEModel object to which the content will be added
- dna_seq (str) DNA sequence of dummy gene. Should be representative of the average codon composition, amino acid composition, length of a gene in the organism being modeled
- update (bool) If True, run update functions on all transcription, translation, complex formation, and metabolic reactions added when constructing dummy reactions.

cobrame.util.building.add_m_model_content (me_model, m model. complex_metabolite_ids=None)

Add metabolite and reaction attributes to me_model from m_model. Also creates StoichiometricData objects for each reaction in m_model, and adds reactions directly to me_model if they are exchanges or demands.

Parameters

- me_model (cobrame.core.model.MEModel) The MEModel object to which the content will be added
- m_model (cobra.core.model.Model) The m_model which will act as the source of metabolic content for MEModel
- complex metabolite ids (list) List of complexes which are 'metabolites' in the m-model reaction matrix, but should be treated as complexes

cobrame.util.building.add_metabolic_reaction_to_model (me_model, stoichiomet*ric_data_id*, directionality, complex_id=None, spontaneous=False, update=False, keff=65)

Creates and add a MetabolicReaction to a MEModel.

Parameters

- **me_model** (*cobrame.core.model.MEModel*) **MEModel** that the MetabolicReaction will be added to
- stoichiometric_data_id (str) ID of the StoichiometricData for the reaction being added
- directionality (str) -
 - Forward: Add reaction that occurs in the forward direction
 - Reverse: Add reaction that occurs in the reverse direction
- **complex_id** (*str or None*) ID of the ComplexData for the enzyme that catalyze the reaction being added.
- spontaneous (bool) -
 - If True and complex_id=" add reaction as spontaneous reaction
 - If False and complex_id=" add reaction as orphan (CPLX_dummy catalyzed)

cobrame.util.building.add_model_complexes (me_model, complex_stoichiometry_dict, complex_modification_dict, verbose=True)

Construct ComplexData for complexes into MEModel from its subunit stoichiometry, and a dictionary of its modification metabolites.

It is assumed that each modification adds one equivalent of the modification metabolite. Multiple

Intended to be used as a function for large-scale complex addition.

For adding individual ComplexData objects, use add_complex_to_model

Parameters

- me_model(cobrame.core.model.MEModel)-
- complex_stoichiometry_dict (dict) {unmodified_complex_id: {protein_<locus_tag>: stoichiometry}}
- complex_modification_dict (dict) -

{modified_complex_id:{core_enzyme: unmodified_complex_id, 'modifications:
 {mod_metabolite: stoichiometry}}}

cobrame.util.building.add_reactions_from_stoichiometric_data (me_model,

rxn_to_cplx_dict, rxn_info_frame, update=False, keff=65)

Creates and adds MetabolicReaction for all StoichiometricData in model.

Intended for use when adding all reactions from stoichiometric data for the first time.

For adding an individual reaction use add_metabolic_reaction_to_model()

Parameters

- **me_model** (*cobrame.core.model.MEModel*) **MEModel** that the MetabolicReaction will be added to
- **rxn_to_cplx_dict** (*dict*) {StoichiometricData.id: catalytic_enzyme_id}
- **rxn_info_frame** (pandas.Dataframe) Contains the ids, names and reversibility for each reaction in the metabolic reaction matrix as well as whether the reaction is spontaneous

cobrame.util.building.add_subreaction_data(me_model, modification_id, modification_stoichiometry, modifica-

tion_enzyme=None, *verbose=True*)

Creates a SubreactionData object for each modification defined by the function inputs.

It's assumed every complex modification occurs spontaneously, unless a modification_enzyme argument is passed.

If a modification uses an enzyme this can be updated after the SubreactionData object is already created

Parameters me_model (cobrame.core.model.MEModel) -

cobrame.util.building.add_transcription_reaction(me_model, tu_name, locus_ids, sequence, update=True)

Create TranscriptionReaction object and add it to ME-Model. This includes the necessary transcription data.

Parameters

- **me_model** (*cobrame.core.model.MEModel*) The MEModel object to which the reaction will be added
- tu_name (*str*) ID of TU being transcribed. The TranscriptionReaction will be added as "transcription_+TU_name" The TranscriptionData will be added as just "TU_name"
- **locus_ids** (*set*) Set of locus IDs that the TU transcribes
- **sequence** (*str*) Nucleotide sequence of the TU.
- **update** (*bool*) If True, use TranscriptionReaction's update function to update and add reaction stoichiometry

date = False)

Returns TranscriptionReaction for the TU

Return type cobrame.core.reaction.TranscriptionReaction

cobrame.util.building.add_translation_reaction(me_model, locus_id, dna_sequence, up-

Creates and adds a TranslationReaction to the ME-model as well as the associated TranslationData

A dna_sequence is required in order to add a TranslationReaction to the ME-model

Parameters

- **me_model** (cobra.core.model.MEModel) The MEModel object to which the reaction will be added
- **locus_id** (*str*) Locus ID of RNA product. The TranslationReaction will be added as "translation + _ + locus_id" The TranslationData will be added as "locus_id"
- **dna_sequence** (*str*) DNA sequence of the RNA product. This string should be reverse transcribed if it originates on the complement strand.
- **update** (*bool*) If True, use TranslationReaction's update function to update and add reaction stoichiometry

Creates and adds transcription and translation reactions using genomic information from the organism's genbank file. Adds in the basic requirements for these reactions. Organism specific components are added ...

Parameters

- **me_model** (*cobrame.core.model.MEModel*) The MEModel object to which the reaction will be added
- **gb_filename** (*str*) Local name of the genbank file that will be used for ME-model construction
- tu_frame (pandas.DataFrame) DataFrame with indexes of the transcription unit name and columns containing the transcription unit starting and stopping location on the genome and whether the transcription unit is found on the main (+) strand or complementary (-) strand.

If no transcription unit DataFrame is passed into the function, transcription units are added corresponding to each transcribed gene in the genbank file.

- **element_types** (*set*) Transcription reactions will be added to the ME-model for all RNA feature.types in this set. This uses the nomenclature of the genbank file (gb_filename)
- **verbose** (*bool*) If True, display metabolites that were not previously added to the model and were thus added when creating charging reactions
- **frameshift_dict** (*dict*) {locus_id: genome_position_of_TU}

If a locus_id is in the frameshift_dict, update it's nucleotide sequence to account of the frameshift

Adds tRNA charging reactions for all tRNAs in ME-model

Parameters

- **me_model** (cobra.core.model.MEModel) The MEModel object to which the reaction will be added
- trna_aa (dict) Dictionary of tRNA locus ID to 3 letter codes of the amino acid that the tRNA contributes

{tRNA identifier (locus_id): amino_acid_3_letter_code}

- trna_to_codon (dict) Dictionary of tRNA identifier to the codon which it associates {tRNA identifier (locus_id): codon_sequence}
- **verbose** (*bool*) If True, display metabolites that were not previously added to the model and were thus added when creating charging reactions

Creates a TranscribedGene metabolite object and adds it to the ME-model

Parameters

• **me_model** (*cobrame.core.model.MEModel*) – The MEModel object to which the reaction will be added

- **locus_id** (*str*) Locus ID of RNA product. The TranscribedGene will be added as "RNA + _ + locus_id"
- left_pos (int or None) Left position of gene on the sequence of the (+) strain
- right_pos (int or None) Right position of gene on the sequence of the (+) strain
- **seq** (*str*) Nucleotide sequence of RNA product. Amino acid sequence, codon counts, etc. will be calculated based on this string.
- strand(str or None)-
 - (+) if the RNA product is on the leading strand
 - (-) if the RNA product is on the complementary strand
- rna_type (str) Type of RNA of the product. tRNA, rRNA, or mRNA Used for determining how RNA product will be processed.

Returns Metabolite object for the RNA product

Return type cobrame.core.component.TranscribedGene

cobrame.util.dogma module

cobrame.util.dogma.extract_sequence(full_seq, left_pos, right_pos, strand)
cobrame.util.dogma.get_amino_acid_sequence_from_dna(dna_seq)
cobrame.util.dogma.return_frameshift_sequence(full_seq, frameshift_string)
cobrame.util.dogma.reverse_transcribe(seq)

cobrame.util.mass module

amino_acid_count: {amino_acid: number}

nucleotide_count: {nucleotide: number}

cobra.core.massbalance module

cobrame.util.massbalance.check_me_model_mass_balance(model0)
cobrame.util.massbalance.check_transcription_mass_balance(reaction)
cobrame.util.massbalance.elements_to_formula(obj, elements)
cobrame.util.massbalance.eval_reaction_at_growth_rate(reaction, growth_rate)
cobrame.util.massbalance.get_elements_from_process_data(reaction, process_data, elements)
If a modification is required to form a functioning macromolecule update the element dictionary according

If a modification is required to form a functioning macromolecule, update the element dictionary accordingly. cobrame.util.massbalance.stringify(element, number)

Module contents

Module contents

CHAPTER 6

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