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Doctoral Dissertation

Development of computational method for heterologous pathways design

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Chapter 1

General introduction

Biosynthesis of biofuels, diverse chemicals and sustainable synthesis of several chemicals has attracted much attention due to the potential depletion of petroleum. Natural organisms often produce target metabolites at low yields, and it is difficult to improve bio-products since the information about metabolic system and genetic manipulation tools of the organisms are generally limited.

Metabolic engineering is the key solution broadly used to redirect existing metabolic pathways and/or to incorporate heterologous pathways into well-characterized hosts, including *Escherichia coli*, *Saccharomyces cerevisiae*, and so forth, for improvement of the productions of native and/or nonnative metabolites. However, it is still difficult to know and answer the following questions; How does the global metabolism of a microbial cell respond to changes in its environment? How do we get the feasible heterologous pathways and genes to improve the productivity from huge number of possible pathways/genes? Which host is suitable for a target production? To answer these questions, an appropriate computational method/*in silico* platform focusing on metabolic design is desirable.

This chapter, the importance of bio-based process and the reason for the development of *in silico* platform will be discussed. Then, several pathway design methods previously reported will be explained and key features comparing with this

thesis are summarized. In this study, flux balance analysis (FBA) was used to estimate fluxes of the metabolic system. In the third part of this chapter, a process for constructing a model of the metabolic system and how to use FBA technique for analyzing the metabolic fluxes will be presented. Finally, the objective and outline of this thesis will be summarized.

1.1 Importance of bio-based process for valuable chemicals

The demand of crude oil is gradually increasing as shown in Fig. 1.1 as well as increasing in its prices. These data suggest that petroleum sources are unsustainable. It takes about hundred to million years to make oil by decomposition of dead organisms mostly zooplankton and algae. Fuels and chemicals are mainly produced by petroleum-based process from crude oil. Moreover, the petroleum-based process has a negative impact on earth by releasing pollutants and generating hazardous wastes. Thus alternative energy resources, such as solar, wind, hydroelectric and biomass, which are renewable and sustainable, have been increasingly used in order to preserve fossil resources and to reduce CO₂-emissions. Among renewable energies, biomass shares the most consumption in 2010 (Fig. 1.2). Biomass can be converted into three main types of products, i.e., electrical/heat energy, transport fuels, and chemical feedstocks (McKendry, 2002) by using microorganism via bio-based processes.

In addition, biomass is available as renewable resources either as natural processes (wood, pulp) or as by-products and/or wastes of human activities

(molasses, rice straw, corn stover). Therefore, bio-based process by using microorganisms as cell factories to convert biomass into valuable products is becoming an attractive way. In many years ago, bio-based process mainly focused on improving the production of native metabolites (found in a host cell) widely used in food and beverage industry, for example, amino acids produced by E. coli and ethanol produced by yeast. Nowadays, trend of bio-based process for useful products has moved to produce nonnative metabolites (not found in the host cell) basically important as chemicals used for many industrial purposes such as polymers, pharmaceuticals, fuels, solvents, and so forth. Currently, those of useful native and nonnative metabolites are able to be produced by microorganisms (Dugar and Stephanopoulos, 2011; J. W. Lee et al., 2011; S. K. Lee et al., 2008; Papini et al., 2010; Schneider and Wendisch, 2011). The examples of such compounds and global demand are shown in Table 1.1. Some of bio-fermentation products, for instance, ethanol and higher alcohols are usually being used as fuels and solvents (Wang et al., 2012). 1,3-propanediol forms the basis of polymers such as polytrimethylene terephathalate (PTT) (H. Liu et al., 2010), while isoprene is an intermediate metabolite in the production of cis-1,4-polyisoprene, a synthetic version of natural rubber (Ohya and Koyama, 2005).

As shown in Fig. 1.3, the sales of chemicals made by bio-based process in 2010 were \$118 billion. It is forecasted to increase to \$296 and \$668 billion by 2015 and 2020, respectively. These data suggest the importance of industrial chemicals made by using bio-based process. The bio-based process for chemicals would be an efficient route for reducing wastes and CO₂ emissions, and preserving fossil fuels comparing with petroleum-based process.



FIG. 1.1 Crude oil consumption and price from 1990-2011

(Source: U.S. Energy Information Administration available at http://www.eia.gov/)



FIG. 1.2 World total and energy consumption by source, 2010

(Source: U.S. Energy Information Administration available at http://www.eia.gov/)



FIG. 1.3 Trends in production of bio-based chemicals. 2015 and 2020 data are

projections based on 2010 data (Adapted from (Festel et al., 2012))

Table 1.1 Bio-based products and global demand

Compound	Host cell factory	Reference	Global market size (million ton/year)	Industrial application	
	Escherichia coli	(H. Liu et al., 2010)		Co-polymers to	
1,3-propanediol	Saccharomyces cerevisiae	(Rao et al., 2008)	0.1-0.5	produce PTT for plastics	
1,4-butanediol	Escherichia coli	(Yim et al., 2011)	1.7	Polymers, solvents	
4 hutanal	Escherichia coli	(Shen et al., 2011)		Polymers, plastics,	
1-butanoi	Synechococcus elongatus PCC7942	(Lan and Liao, 2012)	3.0	solvents	
2.3 butanodial	Escherichia coli	(S. Lee et al., 2012)	0.06	Chemical, food, fuel	
2,3-butaneuloi	Bacillus subtilis	(Biswas et al., 2012)	0.00	fibers, plastics	
3-hydroxypropanoic acid	Escherichia coli	(Rathnasingh et al., 2012)	0.5	Contact lenses, polymers for diapers, carpet fibers	
O deverine	Corynebacterium glutamicum	(Kind and Wittmann, 2011)	0.1		
Cadaverine	Escherichia coli	(ZG. Qian et al., 2011)	0.1	Polyamides for plastics	
Ethonal	Saccharomyces cerevisiae	(Guadalupe Medina et al., 2010)	<u></u>	Biofuel, food beverages, solvents	
Ethanoi	Escherichia coli	(Woodruff et al., 2013)	60		
Glucaric acid	Escherichia coli	(Moon et al., 2009)	0.06	Solvents, nylons	
Glutamic acid	Corynebacterium glutamicum	(Becker and Wittmann, 2012)	2.5	Monomers for polyester and polyamides	
Isoprene	Escherichia coli	(Lv et al., 2012)	0.1-0.5	Natural rubber, thermoplastics	
Itaconic acid	Aspergillus terreus	(Kuenz et al., 2012)	0.08	Polymers, fibers	
	Escherichia coli	(Mazumdar et al., 2013)		Polymers, plastics, fibers	
Lactic acid	Saccharomyces cerevisiae	(Pacheco et al., 2012)	0.3-0.5		
	Synnechocystis sp. PCC6803	(Angermayr et al., 2012)			
Malic acid	Escherichia coli	(Zhang et al., 2011)	0.06	Acidulent in food industry	
Muconic acid	Saccharomyces cerevisiae	(Curran et al., 2013)	2.20	Polymore plastice	
	Escherichia coli	(Niu et al., 2002)	2.30	Polymers, plastics	
	Corynebacterium glutamicum	(Litsanov et al., 2012)		Feed additives fuel	
Succinic acid	Escherichia coli	(Hoefel et al., 2012)	0.1	additives, fibers,	
	Saccharomyces cerevisiae	(Otero et al., 2013)]	polymers	

*Data retrieved from (Vennestrøm et al., 2011) and IEA Bioenergy (source:

www.ieabioenergy.com/) .

1.2 Importance of computational methods for metabolic pathway design

Several approaches leading to the production of valuable products generally use engineered microbes in which native metabolic networks of microorganisms are artificially modified to produce target products. One such standard strategy employed for producing target metabolites is the incorporation of heterologous pathways into well-characterized hosts such as B. subtilis, C. glutamicum, E. coli, and S. cerevisiae as shown in Table 1.1. Nevertheless, the selection of suitable heterologous metabolic pathways for host organisms is often difficult due to the metabolic network complexity. Besides, huge amount of information on metabolic reactions have been found in literature, and are available on public databases such as KEGG (Kanehisa et al., 2008), BRENDA (Chang et al., 2009), ENZYME (Bairoch, 2000), MetaCyc (Caspi et al., 2008), and BKM-react (Lang et al., 2011) as shown in Fig. 1.4. There are 8,507, 8,244, and 9,096 metabolic reactions available on BRENDA, KEGG, and MetaCyc, respectively, and the integration database, BKM-react, contains 18,172 unique metabolic reactions combining from the 3 databases, BRENDA, KEGG and MetaCyc. Thus, to find appropriate heterologous pathways for target production generally requires massive calculations. For example, to search for heterologous pathway for producing target nonnative metabolite, it requires users to search the heterologous reactions which are able to connect nonnative metabolite to the target host metabolism. As you can see in Fig. 1.5, there are 11 possible routes/heterologous pathways containing 4 heterologous reactions to connect nonnative metabolite to the host's metabolic network. Since the complexity of metabolic network available on database, it is difficult to manually check and search for all possible heterologous pathways. In addition, the substrate-product conversion should be primitively estimated as a reference value to be compared with experimental outcome, and could be used to evaluate the feasibility of heterologous pathways.

In Fig. 1.6, a toy metabolic network demonstrates 7 biochemical reactions (2 intracellular and 5 transport reactions) and contains 5 intracellular metabolites (A-E) and 5 extracellular metabolites (A_{ext}, C_{ext}, D_{ext}, E_{ext}, and Biomass). If biomass is a target from this network, theoretical yield can be calculated from reaction stoichiometry between reactants and products. Therefore, 1 mole of A, D, and C will be converted to 2 mole of Biomass and 1 mole of E as a by-product as shown in Fig. 1.7. This toy metabolic network is simple and contains only 7 reactions, so it is possible to calculate by hand. However, the real host cell such as *E. coli* contains 931 internal and 143 transport reactions (Reed et al., 2003), it may be impossible to manually calculate theoretical yield of target product and find feasible pathways of targets.



FIG. 1.4 Distribution of the unique metabolic reactions between BRENDA, KEGG, and

MetaCyc databases (Lang et al., 2011)



FIG. 1.5 Heterologous pathway design by human to search for possible heterologous

reactions to connect nonnative metabolite to the host metabolism



FIG. 1.6 Toy metabolic network map consists of 5 intracellular metabolites (A-E) and 5 extracellular metabolites (A_{ext} , C_{ext} , D_{ext} , E_{ext} , and Biomass), 2 intracellular reactions (v_1 - v_2) and 5 transport reactions (b_1 - b_5) and biochemical reactions present on right-hand side.



FIG. 1.7 Theoretical yield calculation based on reaction stoichiometry

Without using the computational method, it is difficult and tight to handle numerous reactions in host's metabolic network and to search heterologous reactions for the production of nonnative metabolites from databases. Additionally, to find the feasible heterologous pathway and to check whether one target nonnative metabolites produced by the host cell, are time-consuming tasks, when these tasks depend only on the researcher's knowledge and manual screen of information. From these reasons, a computational method is required to help researchers for finding the feasible heterologous pathway of target nonnative metabolite.

Recently, numerous *in silico* heterologous pathway search methods have been proposed and used in target metabolites productions (Cho et al., 2010; Dogrusoz et al., 2009; Finley et al., 2009; Flórez et al., 2011; Handorf et al., 2005; Li et al., 2004; McShan et al., 2003; Moriya et al., 2010; Pey et al., 2011; Pharkya et al., 2004; Rodrigo et al., 2008; Yousofshahi et al., 2011). Comparison of key points among those methods is shown in Table 1.2. The key points are composed of:

- 1. There is no requirement of background information for searching pathway, for example, rules of enzyme transformation between substrate to product.
- 2. The calculation time of the method is fast when search for the optimized pathway with maximum theoretical yield of target.
- 3. New/alternative pathways of target can be generated.
- 4. New reaction(s) not existing in the available databases can be generated.
- 5. The information of new reaction such as gene, sequence, protein is available.
- 6. The heterologous pathway is generated for specific host cells.
- 7. The information of heterologous genes corresponding to a heterologous pathway is available and specific for the target host cell.
- 8. All possible nonnative metabolites and heterologous pathways information are available as a catalog for each target host cell.

In Table 1.2, all approaches are able to screen the new pathway for target production. However, only PathMiner (McShan et al., 2003), Pathway generation (C. Li et al., 2004), PathPred (Moriya et al., 2010), BNICE (Finley et al., 2009), and Prioritization (Cho et al., 2010). These methods can generate new metabolic reactions since it used the concept of generalize enzyme activity based on third-level of enzyme classification, which is recommended by International Union of Pure and Applied Chemistry and International Union of Biochemistry and Molecular Biology (IUPAC-IUBMB) (NC-IUBMB, 1999). Each enzyme is assigned a four-digit (EC i.j.k.I) Enzyme Commission (EC) number. Enzyme-catalyzed reactions based on the thirdlevel enzyme classification, EC i.j.k, are not substrate specific, and thus, these described the transformation of functional groups. With this concept, those methods create rules of enzyme transformation to search for possible reactions catalyzed by the third-level enzyme. Even those methods are capable to create new metabolic reactions, but the information of gene and protein for further study is unavailable. Thus, the rule-based methods are high risk as predicted chemicals may not be possible in real experiments.

PathMiner, Pathway generation, PathPred, BNICE, and Prioritization methods require background information are categorized as the rules-based methods, while OptStrain (Pharkya et al., 2004), Network expansion (Handorf et al., 2005), DESHARKY (Rodrigo et al., 2008), Graph-based pathway (Dogrusoz et al., 2009), Path finding (Pey et al., 2011), Probabilistic (Yousofshahi et al., 2011), SPABBATS (Flórez et al., 2011), and ArtPathDesign (this thesis) (Chatsurachai et al., 2013) are categorized as the graph-based methods. The graph-based methods are capable to generate pathways of target without using any background information of enzyme activity. The rule-based methods require much time to search for the optimized pathway due to a huge number of possible pathways from rules and atomic balances, while most of graph-based methods are fast to calculate based on reaction stoichiometry. However, the OptStrain, Network expansion, and DESHARKY applied alternative ways/scores to rank and select the optimized pathways. For example, OptStrain used OptKnock (Burgard et al., 2003) algorithm for further improving yield of target by knockout genes in the host network. DESHARKY used earlier experimental data of RNA polymerase activity as a score to find the optimized pathway, which required more calculation time as well as prior information.

Among key features in Table 1.2, most of earlier methods have not yet been developed to provide specific heterologous pathways and heterologous genes for the

host cell in addition to the catalog of nonnative metabolite especially for the target host cell. These features are significant for researchers in order to improve and /or produce the desire of industrial nonnative metabolites, because expression of heterologous pathways may be low/not active. The main reasons are sometimes occurred by heterologous genes show low/no expression in the target host cell since diversity of species and cofactor balances such as NAD+/NADH and NADP+/NADPH.

Hence, in this thesis I have developed the *in silico* method according to the purposes (Fig. 1.8), to search for feasible heterologous pathways, suggest rational heterologous genes particular for each host, and provide the catalog of nonnative metabolites, which are able to be produced by the host cell. This system was named as ArtPathDesign (<u>Art</u>ificial heterologous <u>Path</u>way <u>Design</u>) and could overcome the problems about host-specific heterologous pathways and genes, as well as the catalog of nonnative metabolites (Table 1.2). In addition, this computational method greatly reduces times and costs for data analysis and also experiments.



FIG. 1.8 The schematic of the computational method developed in this thesis, named as "ArtPathDesign".

Host metabolic network and metabolic reactions collected in an in-house database were used as input data for screening heterologous pathways of nonnative metabolites. In parallel, Codon Adaptation Index (CAI) scores of all genes retrieving from the available databases was calculated and used to select candidate genes. FBA simulation was performed to select heterologous pathways. Finally, list of nonnative metabolites and heterologous genes particular for a target host will be listed as the output result in html-format easily opened by web browser such as Google Chrome, Firefox, etc.

Table 1.2 Comparison of computational metabolic pathway search methods

Method		Key points of method							
name	Reference	No requirement of Background	Optimized pathway search (fast)	New pathway	New reaction	Information of new reaction (enzyme/gene)	Host- specific pathway	Host-specific heterologous gene(s)	Catalog of nonnative metabolites (host-specific)
PathMiner	McShan et al. (2003)	×	×	~	~	×	×	×	×
Pathways generation	Li et al. (2004)	×	×	~	~	×	×	×	×
OptStrain	Pharkya et al. (2004)	~	×	~	×	×	~	×	×
Network expansion	Handorf et al. (2005)	~	×	~	×	*	×	×	×
DESHARKY	Rodrigo et al. (2008)	~	×	~	×	×	~	~	×
Graph- based pathway	Dogrusoz et al. (2009)	~	~	~	*	*	*	×	×
PathPred	Moriya et al. (2010)	×	*	~	~	×	×	×	×

Table 1.2 (Continued)

		Key points of method							
Method name	Reference	No requirement of Background	Optimized pathway search (fast)	New pathway	New reaction	Information of new reaction (enzyme/gene)	Host- specific pathway	Host-specific heterologous gene(s)	Catalog of nonnative metabolites (host-specific)
BNICE	Finley et al. (2009)	×	×	~	~	×	×	×	×
Prioritization	Cho et al. (2010)	×	×	~	~	×	×	×	×
Path finding	Pey et al. (2011)	~	~	~	×	×	×	×	×
Probabilistic	Yousofshahi et al. (2011)	~	~	~	×	×	×	×	×
SPABBATS	Flórez et al. (2011)	~	~	~	×	×	×	×	×
ArtPathDesig n (this thesis)	Chatsurachai et al. (2013)	V	V	~	×	×	~	~	v

Modeling and analysis of genome-scale metabolic networks

Generally, a model is created to simulate a process or a set of processes observed in the experiment in order to better understand mechanisms of process and to predict outcomes for a given set of specific input parameters. Therefore, to gain insight into cellular metabolism, a genome-scale metabolic network model has been an important tool. The genome-scale metabolic network is reconstructed from genome sequence annotation and biochemical reactions mining from databases (Table 1.3) and literatures. The construction of metabolic network is an iterative decision-making and time-consuming process and it could take up to one month to several months to complete a comprehensive model of a genome-scale metabolic network. In addition, an accurate model could be done by using experimental data retrieved from literatures. *E. coli* (Reed et al., 2003), *S. cerevisiae* (Mo et al., 2009), *B. subtilis* (Oh et al., 2007) and *C. glutamicum* (Shinfuku et al., 2009) models are the examples of accurate models predicting cellular phenotypes under various conditions.

Table 1.3 Database	e useful for	pathway	mining and	l curation

Database	URL
KEGG	http://www.genome.jp/kegg/pathway.html
MetaCyc	http://metacyc.org/
BRENDA	http://www.brenda-enzymes.org/
BKM-react	http://bkm-react.tu-bs.de/
BiGG	http://bigg.ucsd.edu/
CyanoBase	http://genome.microbedb.jp/cyanobase
Biochemical Pathway Maps	http://web.expasy.org/pathways/

Several modeling techniques (Raman et al., 2006) are available to analyze and to simulate cellular mechanism/response such metabolic flow when metabolic pathway or environment altered. Among these techniques, flux balance analysis (FBA) is the most commonly used. FBA can provide estimations of the metabolic fluxes on the genome-scale metabolic network, thereby making it possible to predict the growth rate of an organism or the rate of target production (Orth et al., 2010).

Theory of flux balance analysis (FBA)

FBA assumes that metabolic fluxes will reach a steady state constrained by the stoichiometry (Kauffman et al., 2003). The stoichiometric constraints lead to an underdetermined system; however, a bounded solution space of all feasible fluxes can be identified. This solution space can be further restricted by specifying maximum and minimum fluxes through any particular reaction and by specifying other physiochemical constraints. Here, the metabolic fluxes are estimated by these constraints, and the constraints can be refined by adding experimental data.

When the solution space that describes the capability of the organism is defined, the metabolic network's behavior can be studied by optimizing the steady-state behavior with respect to some objective function. The simulation results can then be experimentally verified and used to further strengthen the model. Finally, the iterative model refinement procedure can result in predictive models of cellular metabolism (Mo et al., 2009; Oh et al., 2007; Reed et al., 2003; Shinfuku et al., 2009). To better understand how to formulate a FBA problem, the steps are explained in detail and demonstrated through the toy metabolic network (Fig. 1.6). **FBA model formulation** contains 4 steps. (Adapted from (Kauffman et al., 2003; Raman and Chandra, 2009))

Step I. System definition

Development of a flux balance model requires the definition of all the metabolic reactions and metabolites. Fig. 1.6 shows the toy metabolic network, which contains 2 intracellular and 5 transport reactions. There are 5 intracellular metabolites (A-E) and 5 extracellular metabolites (A_{ext}, C_{ext}, D_{ext}, E_{ext}, and Biomass).

The biochemical reactions of the network are listed here.

 $A_{extracellular} \rightarrow A$ $A + D \rightarrow B + E$ $C \rightarrow B$ $C_{extracellular} \rightarrow C$ $D_{extracellular} \rightarrow D$ $E \rightarrow E_{extracellular}$

Step II. Mass balance

Once all reactions and transport mechanism of the system are identified, a dynamic mass balance is derived for all intracellular metabolites in the metabolic network shown here (equation 1.1a-e).

$$\frac{d[A]}{dt} = b_1 - v_1 \tag{1.1a}$$

$$\frac{d[B]}{dt} = v_1 + v_2 - b_2 \tag{1.1b}$$

$$\frac{d[C]}{dt} = b_3 - v_2 \tag{1.1c}$$

$$\frac{d[D]}{dt} = b_4 - v_1 \tag{1.1d}$$

$$\frac{d[E]}{dt} = v_1 - b_5 \tag{1.1e}$$

The mass balance is defined in terms of the flux through each reaction and the stoichiometry of that reaction, thus, a set of ordinary differential equations is obtained (equation 1.2a-1.2e). In this analysis, the steady state of the system is assumed, and corresponds to the case that input fluxes are equal to output fluxes, as follows.

$$\frac{d[A]}{dt} = b_1 - v_1 = 0 \tag{1.2a}$$

$$\frac{d[B]}{dt} = v_1 + v_2 - b_2 = 0 \tag{1.2b}$$

$$\frac{d[C]}{dt} = b_3 - v_2 = 0 \tag{1.2c}$$

$$\frac{d[D]}{dt} = b_4 - v_1 = 0 \tag{1.2d}$$

$$\frac{d[E]}{dt} = v_1 - b_5 = 0 \tag{1.2e}$$

The differential equations can be represented using a matrix notation, where "S" is the stoichiometric matrix and "v" is the vector of the fluxes. The goal of FBA is to identify the metabolic fluxes under steady-state condition of the metabolic network.

At steady-state condition, the different equations of all metabolites can display in matrix form shown in Fig. 1.9 below.

Stoichiometric maxtrix

 $\forall i \in M \\ M = a \text{ set of metabolites} \\ R = a \text{ set of fluxes (reactions)}$

Steady state Mass balance



FIG. 1.9 Matrix notations of differential equations under steady-state condition

Step III. Defining measurable fluxes and/or range of fluxes

In general, there are more reactions (or fluxes) than the number of metabolites,

and thus the steady-state solution for the metabolic fluxes is underdetermined.

Therefore, additional constraints are required to uniquely determine the steadystate flux distribution. One way to get the additional constraints for the metabolic network is measuring metabolic fluxes experimentally. In linear algebra, to solve the problem, the number of variables should be equal to the number of equations, that is, the measurement of fluxes such as substrates uptake, by-products formation rate should be used as additional constraints. For example, the toy metabolic system contains 5 equations and 7 variables; it requires at least 2 measured fluxes of substrate uptake or production rate to find the unique solution of this problem. The exact flux values are commonly not defined, but rather a range of allowable flux values. The ranges of flux values are either retrieved from experiments or literature, which are used as additional constraints.

The examples of additional constraints for the toy metabolic network are demonstrated below. Here, b_1 , b_3 and b_4 represent substrate uptake rates (equation 1.3 h-j) that can be observed from experiments.

 $0 \le v_{1} \le 10 \quad (1.3a)$ $0 \le v_{2} \le 10 \quad (1.3b)$ $0 \le b_{1} \le 10 \quad (1.3c)$ $0 \le b_{2} \le 10 \quad (1.3d)$ $0 \le b_{3} \le 10 \quad (1.3e)$ $0 \le b_{4} \le 10 \quad (1.3f)$ $0 \le b_{5} \le 10 \quad (1.3g)$

$b_1 = 4.2$	(1.3h)
$b_3 = 5.5$	(1.3 <i>i</i>)
$b_4 = 3.2$	(1.3i)

Step IV. Optimization

A genome-scale metabolic network always has more reactions (fluxes) in the system than the number of metabolites corresponding to the underdetermined system, thus there are allowable solution spaces/flux distributions (Fig. 1.10). To find the unique/optimal solution, an optimization technique is commonly applied for measuring the internal fluxes in the metabolic network. Fig. 1.10 shows the flux distribution of the toy metabolic network predicted by using the optimization technique with the objective function to maximize Biomass production (the flux of b_2).

Using the optimization technique, the metabolic network is assumed to be optimized with respect to a target objective function. This allows the underdetermined system to be formulated as the optimization problem. The objective functions such as maximization of biomass, minimization of substrate uptake, maximization of ATP production, etc. are widely used to estimate cellular metabolisms and to provide predictions which can be verified by experimental data (Mo et al., 2009; Oh et al., 2007; Reed et al., 2003; Schuetz, Kuepfer, & Sauer, 2007; Shinfuku et al., 2009). The optimization problem of the genome-scale metabolic network is formulated below. Problem:

Maximize/Minimize $\sum_{j=1}^{R} c_j \cdot v_j$ (Problem 1)

Subject to.

1. $\sum_{i}^{R} S_{ii} \cdot v_{i} = 0, \forall i \in M \text{ and } \forall j \in R$ (mass balance constraints)

2. Linear inequality or equality constraints (additional constraints)

where,

R is a number of reactions (fluxes) in the system or cell.

M is a number of metabolites in the system or cell.

 c_j represents weight of the individual flux of the j^{th} reaction that contributed to the objective function.

 v_i represents metabolic fluxes of the j^{th} reaction.

 S_{ij} represents the stoichiometric coefficient indicating the amount of the i^{th} metabolite produced per unit of flux of the j^{th} reaction.

The aim of FBA is to maximize or minimize the objective function (Problem 1) that is subject to mass balance and additional constraints. The output of this problem is a particular flux distribution of vector, v, which maximizes/minimizes the objective function (Orth et al., 2010).

(1.4)

In general, for the genome-scale metabolic network of the host, the biomass equation is generated based on the ratios of cellular components such as amino acids, RNA, DNA, etc., which are either estimated from experiments and/or genome information. Here is the example of the *E. coli*'s biomass reaction (equation 1.4) retrieved from (Reed et al., 2003):

(0.05) 5mthf + (5.0E-5) accoa + (0.488) ala-L + $(0.0010) \operatorname{amp} + (0.281) \operatorname{arg-L} + (0.229) \operatorname{asn-L} +$ (0.229) asp-L + (45.7318) atp + (1.29E-4) clpn_EC + $(6.0E-6) \cos + (0.126) \operatorname{ctp} + (0.087) \operatorname{cys-L} + (0.0247)$ datp + (0.0254) dctp + (0.0254) dgtp + (0.0247) dttp+ (1.0E-5) fad + (0.25) gln-L + (0.25) glu-L + (0.582)gly + (0.154) glycogen + (0.203) gtp + (45.5608) h20+ (0.09) his-L + (0.276) ile-L + (0.428) leu-L +(0.0084) lps_EC + (0.326) lys-L + (0.146) met-L + (0.00215) nad + (5.0E-5) nadh + (1.3E-4) nadp + (4.0E-4) nadph + (0.001935) pe_EC + (0.0276) $peptido_EC + (4.64E-4) pg_EC + (0.176) phe-L +$ (0.21) pro-L + (5.2E-5) ps_EC + (0.035) ptrc + (0.205)ser-L + (0.0070) spmd + (3.0E-6) succoa + (0.241)thr-L + (0.054) trp-L + (0.131) tyr-L + (0.0030) udpg+ (0.136) utp + (0.402) val-L= (45.5608) adp + (45.56035) h + (45.5628) pi +(0.7302) ppi + (1) Biomass



FIG. 1.10 Optimization of the system with the objective function to maximize the flux of b_2 . The b_2 is the biomass production of the toy network (see in Fig. 1.5), which is applied to obtain one optimal solution represented as red dot. The flux distribution of the optimal point is labeled in red on the toy metabolic network map.

1.4 Microorganisms used as industrial cell factories

Since the sequencing of the first complete microbial genome of *Haemophilus influenza* (Fleischmann et al., 1995), a hundred of microbial genomes have been sequenced and archived for public research in GenBank database (Benson et al., 2009). This availability of data provides the scientists to make a genome-scale metabolic model for discovering new information for better understanding cellular properties and processes. In last decade, a genome-scale metabolic model, which is a mathematical model to represent cellular metabolism in linear algebra form, has been used for metabolic engineering by integration of laboratory data such as genome, transcriptome, proteome, metabolome and so forth. Generally, a genomescale metabolic model was constructed by using genome information such as gene, protein, and metabolic network. As shown in Fig. 1.11, the rapid proliferation of genome sequencing projects over the last decade has resulted in an exponential growth in the amount of genomic DNA sequences and information available for reconstructing genome-scale metabolic models.

The complete genome sequence for a number of microorganisms has been established. Thus, the genome information is available to construct the model that helps for the novel metabolic engineering strategies. Well-characterized hosts would be used as cell factories to yield value-added products because of the availability of genome information and genetic manipulation tools. The examples of these famous host models widely used as cell factories for industrial production are briefly summarized as follows.



FIG. 1.11 Statistical information of genome projects from GOLD database until October, 2011, Total projects are 10,031 projects (Pagani et al., 2012).

Escherichia coli

E. coli is an aerobic, gram-negative, rod shaped bacteria that can be commonly found in animal feces, lower intestines of mammals, and even on the edge of hot springs. The complete genome sequence of *E. coli* strain K-12 was finished in 1997 (Blattner et al., 1997). Its genome sequence contains about 4.6 Mbps and 4,288 protein-coding genes. The main reasons why *E. coli* becomes famous host for numerous of products are easy for cultivation and fast growth. Currently, *E. coli* has been engineered to produce valuable compounds such as 1,3-propanediol, 1-butanol, lactic and so forth (in Table 1.1). In order to analyze, interpret, and predict cellular behavior, a genomescale model of *E. coli* was constructed (Reed et al., 2003). This model, named as iJR904, constructed based on *E. coli* K-12 genome annotation data, and showed well predictive simulation comparing with experimental data.

Saccharomyces cerevisiae

A budding yeast *S. cerevisiae* is a eukaryote model for producing alcohols and organic acids such as lactic acid and succinic acid (Table 1.1). The genome sequence of the yeast *S. cerevisiae* was completed in 1996 (Goffeau et al., 1996) and contains about 12 Mbps and defines 5,885 protein-coding genes. Besides, the genome metabolic model of yeast *S. cerevisiae* is also available (Mo et al., 2009). This model named as iMM904 predicted intracellular flux changes consistent with published measurements.

Corynebacterium glutamicum

C. glutamicum, a gram-positive microorganism, is one of the most important bacteria in industrial biotechnology with an annual production of more than 2 million tons of amino acids mainly, L- glutamate, L-serine and L-lysine (Becker and Wittmann, 2012). The *C. glutamicum* genome consists of a single circular chromosome with 3.3 Mbps in size and comprises 3,002 protein-coding genes (Kalinowski et al., 2003). Additionally, its genome-scale metabolic model was constructed and demonstrated metabolic profiles that correponding with experimental data (Shinfuku et al., 2009).

Bacillus subtilis

B. subtilis, a rod-shaped gram-positive bacterium naturally found in soil and plants, is well-recognized as a producer of enzymes such as proteases and amylases (Zweers et al., 2008). In addition, industrial compound like 2,3-butanediol and isobutanol have also been produced by *B. subtilis* (Biswas et al., 2012; Jia et al., 2012). Its genome is about 4.2 Mbps and comprises 4,100 protein-coding genes (Kunst et al., 1997).

Genome-scale metabolic model of *B. subtilis* was published in 2007 and this *in silico* model could predict growth phenotypes of knock-out strains that found to be quite consistent with experimental observations (Oh et al., 2007).

The microorganisms mentioned above are ideal hosts for bioengineered products such as industrial chemicals, since they exhibit high growth activity under various conditions as well as easy to genetically manipulated (Christina, 2010). Moreover, the genome-scale metabolic models of these microorganisms are available for scientists to use as the tools to identify metabolic engineering strategies such as gene amplification and deletion for strain improvements. For example, target genes to improve lycopene production in *E. coli* were successfully identified using *in silico* simulation and corresponding to enhance the lycopene production in ni vivo experiment (Choi et al., 2010). Another example is gene knockout simulation to guide target genes for improving L-valine in *E. coli* (Park et al., 2007).

1.5 Objectives of the work

The current demands of fuels and chemical feedstocks are critically increased, while petroleum resources are limited and unsustainable. Moreover, fuels and industrial chemicals by petroleum-based process show negative impacts on environment. The alternative route to produce energy and valuable chemicals using microorganisms becomes an attractive way. However, some microorganisms are not easy to cultivate and produce high level of target products. In addition, the very large amount of possible heterologous pathways is generated without any background in formation and impossible to handle by human. Therefore, a computational or *in silico* platform to design and select such suitable heterologous pathways is required. Several pathway design methods had been reported (Table 1.2), however, it still lacked of the method that can provide host-specific heterologous pathways, host-specific heterologous genes as well as a catalog of nonnative metabolites particular for each host.

Since the *in silico* platform to design and select suitable heterologous pathways into particular hosts is still not developed, the goal of this thesis is to develop the system aiming to provide necessary information to scientists for producing target metabolites in cell factories hosts such as *B. subtilis, C. glutamicum, E. coli* and *S. cerevisiae*. The algorithm to screen heterologous pathways for the specific host was developed. This algorithm can provide all possible pathways for the production of nonnative metabolites that are non-existent in the host. Then, parameters used for selection of heterologous genes were applied to select feasible pathways for the production of nonnative metabolites. Thus, a rational heterologous pathway design system named as "ArtPathDesign" (<u>Art</u>ificial heterologous <u>Path</u>ways <u>Design</u>) was proposed for an efficient production of nonnative metabolites.
1.6 Outline of the thesis

The thesis consists of 4 chapters, and a schematic outline of the thesis is shown in Fig. 1.12.

Chapter 1 deals with the background and motivation of this thesis. Literature review of metabolic pathway design methods is summarized, and comparison of those methods is demonstrated. The objectives and schematic of this thesis are also described.

Chapter 2 deals with data collection, an in-house database construction and the development of the algorithm for screening heterologous pathways of nonnative metabolites in 3 host cell factories, *E. coli, C. glutamicum*, and *S. cerevisiae* as templates. In this chapter, the K_m value is applied in order to select candidate heterologous genes based on enzyme-substrate affinity corresponding to the heterologous pathway. However, the information of K_m value is depending on experimental data and several enzymes show no information of K_m values. Besides, to obtain or improve the production of nonnative metabolites, host-specific heterologous pathways and host-specific heterologous genes are important features. Thus, a new selection parameter could be applied for selection of suitable heterologous pathways and genes.

Chapter 3 deals with the new score that used for selection of candidate heterologous pathways and genes which are specific for each host. Codon Adaptation Index (CAI) is accepted as a measurement of synonymous codon usage bias, which is one of the most important factors effecting on heterologous enzymes expression. Thus, CAI was applied in this study. With the CAI as the selection score, the host-specific pathways and host-specific genes features are included in the improved *in silico* system, ArtPathDesign. Furthermore, a catalog of nonnative metabolites especially for the target host is improved and available as html file that is well-formed representations.

Chapter 4 deals with the general conclusion and future perspective of this research.



FIG. 1.12 Outline of the thesis

4

Chapter 2

Development of an algorithm to design heterologous pathway

2.1 Introduction

Recognizing the potential depletion of petroleum resources, researchers have become increasingly interested in production of fuels and industrial chemicals by microorganisms (Dugar and Stephanopoulos, 2011; S. K. Lee et al., 2008; Schneider and Wendisch, 2011). Such biosynthesized materials include fuels, plastics, polymers, solvents and drugs (Becker and Wittmann, 2012; J. W. Lee et al., 2011; Papini et al., 2010; B.-W. Wang et al., 2012). To produce such industrially useful materials, modifications of host's metabolic networks are generally required. Target metabolites are frequently produced by incorporating heterologous metabolic pathways into well-characterized host microorganisms, such as E. coli, S. cerevisiae, and so on, that I mentioned in previous chapter. However, the selection of suitable heterologous metabolic pathways for host organisms is often difficult due to the complexity of metabolic network. Although copious data on metabolic reactions and enzymes have been available in the literature and available databases such as KEGG, ENZYME, and BRENDA, constructing a target production pathway using a host's metabolic network with satisfying the metabolic balances requires a scientist' s experience and intuition. Thus, the development of an appropriate in silico platform can facilitate industry-focused metabolic network design by providing possible heterologous pathways for target metabolite production.

Currently, several *in silico* pathway search methods have been developed and used to produce target metabolites (Cho et al., 2010; Dogrusoz et al., 2009; Finley et al., 2009; Flórez et al., 2011; Handorf et al., 2005; C. Li et al., 2004; McShan et al., 2003; Moriya et al., 2010; Pey et al., 2011; Pharkya et al., 2004; Rodrigo et al., 2008; Yousofshahi et al., 2011). In previous chapter, the comparison of these methods is summarized. Even numerous methods are available for screening heterologous pathways of target metabolites, there still remains a lack of agreement on how to choose heterologous pathways and host microorganisms for target production.

In this chapter, I first developed a novel pathway search algorithm that identifies the shortest pathway between a host's metabolic network and target metabolites when heterologous reactions are added to the host's metabolic network. Using this algorithm, all producible target metabolites listed in databases were screened. In addition, to select candidate heterologous enzymes, K_m value was utilized as a selection score. That is, among the heterologous genes that coding enzyme having minimum K_m value was selected for the construction of heterologous pathways. Then, for all producible target metabolites, the production yields were estimated by using flux balance analysis (FBA), assuming the steady-state conditions and the maximization of target or biomass production rate. By analyzing the entire list of producible target metabolites in several different hosts, a set of rational heterologous pathways and host microorganisms that will likely produce desired targets were selected.

2.2 Methods

2.2.1 Construction of an in-house database of metabolic reactions

All known metabolic reactions were considered as candidate heterologous reactions that could be added to the host metabolic network. First, an in-house database of metabolic reactions was constructed based on data stored in KEGG ligand section (Kanehisa et al., 2008) and BRENDA (Chang et al., 2009) databases. All metabolic reaction information regarding genes, enzymes, pathways, and organisms in the KEGG database was collected into the database, which was developed using PostgreSQL 9.0 (The PostgreSQL Global Development Group). The Michaelis-Menten constants (K_m) of the enzymatic reaction data were retrieved from BRENDA.

2.2.2 Genome-scale metabolic model of host microorganisms

In this chapter, 3 well-characterized and industry-used microorganisms, namely, *E. coli, C. glutamicum*, and *S. cerevisiae* were adopted as host microorganisms to be engineered for the target metabolite productions. *E. coli* has been exploited for such industrially valuable compounds as L-phenylalanine, L-tyrosine, 1-butanol, and 1,2-propanediol (Clomburg and Gonzalez, 2011; Juminaga et al., 2012; Shen et al., 2011). *C. glutamicum* is widely used in amino acid production (Becker and Wittmann, 2012). *S. cerevisiae* is an important producer of alcohols and organic acids such as lactate (Hong and Nielsen, 2012). These 3 organisms are widely used for bioengineering since they exhibit high growth activity under several conditions and are easily genetically manipulated

(Christina, 2010). Genome-scale metabolic models of *E. coli* (iJR904)(Reed et al., 2003), *S. cerevisiae* (iMM904) (Mo et al., 2009), and *C. glutamicum* (Shinfuku et al., 2009), based on earlier metabolic constructions with slight modification were used in this study. Because the pathway search algorithm developed in this chapter uses the heterologous reactions listed in the KEGG database, all metabolite IDs in the earlier genome-scale metabolic models were converted to the KEGG compound ID format using metabolite name matching and manually checking.

2.2.3 Heterologous pathway identification for target production

An algorithm to identify heterologous reaction(s) producing a target metabolite within a host microorganism was developed. The algorithm expands the host's metabolic network by sequentially adding heterologous metabolic reactions from the constructed in-house database. The concept of the heterologous pathway identification is shown in Fig. 2.1, and the procedure is as follows:

- 1. A set of metabolites M_0 and a set of metabolic reactions Rxn_0 are defined as those present in the genome-scale metabolic network of the host microorganism.
- From the in-house database, heterologous reactions that satisfy the following conditions are collected:
 - i. The reaction does not exist in Rxn_0 , and
 - ii. It can produce metabolites that do not exist in M_0 from a metabolite in M_0 . A set of these heterologous reactions is defined

as Rxn_1 , and a set of metabolites produced by reactions in Rxn_1 is defined as M_1 .

3. In the same way, Rxn_i is the set of reactions not present in $\{Rxn_0, Rxn_1, ..., Rxn_i\}$ which can produce metabolites not existing in $\{M_0, M_1, ..., M_i\}$ from metabolites included in those sets. This expansion procedure is iterated until no further reaction is connectable to the expanded metabolic network.





 $i = iteration, \forall i \in R$

If a target metabolite is included in a nonnative metabolite set M_i , a set of heterologous reactions that are necessary to produce the target metabolite can be identified. For simplicity, all metabolic reactions in the database were assumed to be reversible. Of course some reactions are known to be irreversible, such as the carboxylation and decarboxylation reactions classified by Nomenclature Committee of the international Union of Biochemistry and Molecular Biology (NC-IUBMB, 1999). However, for the majority of reactions in the database, directional information is limited and thus the reversibility of the reactions is difficult to judge. To avoid the risk of missing important heterologous pathways due to misjudgment of their reaction reversibility, all reactions are assumed to be reversible. This strategy here is to initially screen all possible heterologous pathways regardless of reaction irreversibility, then decide whether the predicted pathway is plausible based on physiological knowledge of the reaction irreversibility.

2.2.4 Flux balance analysis (FBA)

FBA is based on a genome-scale metabolic model and optimization of a specific objective flux by linear programming (Kauffman et al., 2003; Orth et al., 2010). FBA was used to estimate the metabolic flux profile of metabolic networks expanded with heterologous reactions. A pseudo steady-state is assumed, that is, the net sum of all production and consumption fluxes for each internal metabolite is zero. In matrix notation, this condition is represented as $S \cdot v = 0$, where S is the stoichiometric matrix representing the stoichiometry of metabolic reactions in the network and v is the vector of metabolic fluxes. In FBA, the flux profile (constrained by steady-state) is determined by optimizing a specific objective function. The biomass production flux is one of several widely used objective functions that can be maximized. The flux profiles obtained by maximizing biomass production fluxes are known to be well correlated with those

obtained experimentally (Mo et al., 2009; Schuetz et al., 2007; Shinfuku et al., 2009).

For simulation, the coefficients of metabolites representing biomass production flux were extracted from earlier studies (Mo et al., 2009; Reed et al., 2003; Shinfuku et al., 2009). Another objective function, the production flux of the target metabolite, was applied to judge whether the target metabolite was producible by the metabolic network. In all of the FBA simulations, glucose was chosen as the sole carbon source and the following external metabolites were allowed to freely transport through the cell membrane: CO₂, H₂O, SO₄ or SO₃, and NH₃. All calculations were performed using MATLAB 2009b (MathWorks Inc., Natick, MA). The linear programming problem was solved using GLPK 4.34 (GNU Linear Programming Kit) via MATLAB interface.

2.3 Results and discussion

2.3.1 Identification of heterologous pathway(s)

Table 2.1 shows information of metabolic reactions and metabolites obtained from 1,139 species were collected from KEGG database and deposited in the constructed in-house database (published as the Additional file 1 of (Chatsurachai et al., 2012)). To screen the target metabolites that are producible by the host microorganisms *S. cerevisiae, E. coli,* and *C. glutamicum,* the host's metabolic network was iteratively expanded by adding heterologous metabolic reactions as described in the method section.

	In-house database
	(version 1.0)
Source database	KEGG
No. of reactions	7,769
No. of compounds	6,635
No. of reversible reactions	7,769
No. of irreversible reactions	0

Table 2.1 Statistics of the constructed in-house database used in this study

Fig. 2.2 displays the number of nonnative metabolites connected to the host's metabolic network as a function of the number of heterologous reactions. Fewer than 33 heterologous reactions are required to connect 3,154, 3,244, and 3,112 nonnative metabolites to the host's metabolic networks of S. cerevisiae, E. coli, and C. glutamicum respectively. The list of metabolites connected to the host's metabolic networks is presented in the Additional files 2, 3, 4 provided at supplementary section of the publication (Chatsurachai et al., 2012). To this list, the K_m values of heterologous enzymes were added. Knowing the K_m assists in deciding which heterologous enzymes originating from various organisms in the BRENDA database displaying minimum K_m of the corresponding heterologous enzymes are also listed, since the enzyme from this organism is expected to have highest affinity among the orthologous enzymes to the corresponding substrate. Importantly, these identified heterologous reactions of nonnative metabolite production agreed well with those widely used in metabolic engineering and which are important to the industry (Table 2.2), such as isoprene, α -farnesene, poly- β -hydroxybutyrate (PHB), and cadaverine.



FIG. 2.2 Number of connected nonnative metabolites produced by heterologous reactions in 3 host microorganisms. The first vertical axis (solid line) shows the number of connected metabolites in each iteration, while the second vertical axis (dotted line) shows the cumulative number of the connected metabolites.

As an example, the production pathway of 1,3-propanediol (C02457) by *E. coli* and *S. cerevisiae*, which were adopted in earlier studies (Cameron et al., 1998; Nakamura and Whited, 2003), are shown in Fig. 2.3. In previous studies, C02457 production proceeded via conversion of glycerol to 3-hydroxypropanal using glycerol dehydratase (encoded by *dhaB1-B3*). 1,3-Propanediol was then produced, aided by 1,3-propanediol oxidoreductase (encoded by *dhaT*). In this study, the screened heterologous pathways for C02457 production exactly matched those of earlier studies. In *E. coli*, the screened production pathways of isoprene, α -farnesene, and PHB derived by this algorithm were also identical to those of the

earlier studies, while similar heterologous genes introduced to the alternative hosts (C. glutamicum and S. cerevisiae) additionally produced these targets (see Table 2.2). Moreover, both reported and alternative production pathways were screened by the proposed algorithm. For instance, It was found that E. coli cells (R)-propane-1,2-diol when methylglyoxal can produce reductase and lactaldehyde reductase are added to the metabolic network, which has not been reported so far. Similar alternative pathways were found for the production of itaconate, cis, cis-muconate, 2,3-dihydroxybenzoate, and so forth. These results suggest that the developed algorithm successfully identified the metabolic reactions necessary for the target productions and could assist in screening for heterologous pathways of target productions.

Next, I investigated whether these connectable metabolites are producible from glucose as a sole carbon source, by using FBA simulations. In this simulation, the production flux of each nonnative metabolite was used as an objective function to be maximized under the steady-state assumption. When the maximum production flux of a nonnative metabolite is zero, this metabolite is non-producible under the given condition. The maximum production fluxes of all connectable nonnative metabolites were calculated. 28% of the connectable nonnative metabolites of *E. coli* could not be produced using glucose as a sole carbon source. Similarly, 33% of the connectable nonnative metabolites of *S. cerevisiae* and 16% of the connectable nonnative metabolites of *C. glutamicum* were non-producible under this condition. These metabolites could not be produced since they are disconnected from glycolysis. In *E. coli*, these metabolites included trans-aconitase (C02341), butyrate (C00246), acetoacetate (C00164), and L-lactaldehyde (C00424).



FIG. 2.3 Heterologous pathways for 1,3-propanediol production: (a) the production pathway described in earlier studies, in *E. coli* (Cameron et al., 1998; Nakamura and Whited, 2003); (b) the pathway identified by my algorithm in either *E. coli* or *S. cerevisiae* as the host.

2.3.2 Evaluation of production feasibility

To evaluate the feasibility of nonnative target metabolite production, FBA simulations were performed under conditions of maximizing biomass production following heterologous reaction expansion of the genome-scale metabolic model. Metabolic flux profiles calculated by maximization of biomass production rates have been shown to closely represent those in real microorganisms (Edwards and Palsson, 2000a; Edwards et al., 2001; Feist and Palsson, 2010; Schuetz et al.,

2007; Varma and Palsson, 1994). Such agreement can be explained by the growth optimization of microorganisms through evolutionary dynamics (Fong et al., 2003). Furthermore, for the mutant strains constructed in the laboratory, the cells could achieve the near-optimal metabolic state calculated by the FBA simulation after long-term cultivation (Cornelius et al., 2011; Edwards and Palsson, 2000b; Gerdes et al., 2003; Soyer and Pfeiffer, 2010), via the selection of faster growing cells. Thus, it is expected that if a nonnative target metabolite is produced in the FBA simulation under maximized biomass production, that target may be feasibly manufactured.

In Fig. 2.4, the number of target metabolites produced under maximized biomass production was plot, versus the number of heterologous reactions necessary for metabolite production. A threshold yield (1%) was set to identify the produced metabolites because the production yields of some metabolites were positive but extremely small. Sometimes the FBA simulation was underdetermined under biomass maximization conditions; that is, the solution was not unique. In such cases, following the maximization of biomass production, the production flux of the target metabolites was further maximized with fixing the maximized biomass production, to obtain a unique flux profile that would produce the target. In the simulations, a micro-aerobic condition was used to screen the target metabolites produced under the biomass maximization condition, in which significantly a larger number of target metabolites were produced than under anaerobic conditions, and at the same time all anaerobically produced metabolites were included.

Table 2.3 shows the representative target metabolites produced under biomass maximization, together with their corresponding heterologous reactions. The mechanisms involved in these reactions can be classified into two categories. One is based on the production of oxygen as a by-product of the targets. Since the simulations were performed under the micro-aerobic condition, oxygen supply increased the biomass production by activating the electron transfer system and facilitating adenosine triphosphate production. Therefore, if the heterologous reactions used to produce the target are accompanied by oxygen production, the target can be produced under minimum biomass production flux. For example, pentane-2,4-dione was produced by introducing a single heterologous reaction into *E. coli* and *S. cerevisiae*, whereas two heterologous reactions were necessary to produce this metabolite in *C. glutamicum*. Vanillin can be produced under the same mechanism by introducing 4 heterologous reactions into the *E. coli* and *C. glutamicum* metabolic networks.

Another mechanism is associated with NADH oxidation. Under micro-aerobic condition, the cellular growth of microorganisms can be limited by NAD regeneration, which is necessary for glycolysis activity, and which occurs through NADH oxidization. Thus, when the heterologous reactions producing the targets are associated with NADH oxidization, these heterologous reactions are activated when the biomass production is maximized. This phenomenon occurs, for example, in the production of (R)-propane-1,2-diol and 2-propyn-1-al.

It is found that some metabolites are produced only by *E. coli* under conditions of maximum biomass production, such as (R)-propane-1,2-diol and

adipate semialdehyde. Unlike *S. cerevisiae* and *C. glutamicum*, *E. coli* possesses NAD transhydrogenase, which can convert NADP and NADH to NADPH and NAD respectively (and vice versa). In *E. coli* cells, the excess NADH is converted to NADPH which can then enter the target production pathway.



FIG. 2.4 The number of metabolites producible under biomass maximization conditions with the addition of < 10 heterologous reactions.

Commonword		listencie recetion (c) from the literature	Deferrence	Evolution of in cilico decim
compound (synonym separated by a semicolon)	KEGG ID	neterologous reaction(s) from the literature	Reference	Evaluation of in sinco design
Isoprene; 2-methyl-1,3-butadiene	C16521	Introduced ispS gene from <i>Populus nigra</i> to <i>Escherichia coli</i>	(Zhao et al., 2011)	Identical reaction found in <i>E. coli</i> and in <i>Saccharomyces cerevisiae</i> and <i>Cerevisiae</i> <i>glutamicum</i> as the host
α-Farnesene	C09665	Introduced farnesene synthase from plant to <i>E.</i> coli	(C. Wang et al., 2011)	Identical reaction found in <i>E. coli</i> and in <i>S. cerevisiae</i> and <i>C. glutamicum</i> as the host
Poly-β-hydroxybutyrate; PHB	C06143	Introduced phbC and phbB from <i>Streptomyces</i> aureofaciens to <i>E. coli</i>	(Mahishi et al., 2003)	Identical reaction found in <i>E. coli</i> and in <i>S. cerevisiae</i> and <i>C. glutamicum</i> as the host
Cadaverine; 1,5-pentanediamine; 1,5-diaminopentane	C01672	Introduced IdcC from E. coli to C. glutamicum	(Becker and Wittmann, 2012; Kind et al., 2010)	Identical reaction found in <i>C. glutamicum</i> and in <i>S. cerevisiae</i> as the host
Amorpha-4,11-diene	C16028	Introduced AMS1 from the plant <i>Artemisia annua L</i> . to <i>E. coli</i>	(Lindahl et al., 2006; Wallaart et al., 2001)	Identical reaction found in <i>E. coli</i> and <i>S. cerevisiae</i> and in <i>C. glutamicum</i> as the host
Propane-1,3-diol; 1,3propanediol; trimethylene glycol	C02457	Introduced glycerol dehydratase and 1,3- propanediol oxidoreductase from <i>Klebsiella</i> pneumoniae to <i>E. coli</i> .	(Cameron et al., 1998; Nakamura and Whited, 2003)	Identical reaction found in <i>E. coli</i> and in <i>S. cerevisiae</i> as the host
Ethanol; ethyl alcohol; methylcarbinol	C00469	Introduced pyruvate decarboxylase and alcohol dehydrogenase genes from <i>Zymomonas mobilis</i> to <i>C. glutamicum</i>	(Inui et al., 2004)	Identical reaction found in <i>C. glutamicum</i> as the host
(R,R)-Butane-2,3-diol; (R,R)-2,3-Butanediol; (R,R)-2,3-Butylene glycol	C03044	Introduced acetolactate decarboxylase and butanediol dehydrogenase genes to <i>E. coli</i>	(Nielsen et al., 2010)	Identical reaction found in <i>E. coli</i> as the host

Table 2.2 Examples of nonnative metabolites for which my algorithm detected heterologous reactions matching those of previous studies

(R)-Propane-1,2-diol; (R)-1,2-propanediol; (R)-propylene glycol	C02912	Introduced glycerol dehydrogenase gene from <i>Klebsiella pneumoniae</i> and used aldehyde dehydrogenase to produce product in <i>E. coli</i>	(Altaras and Cameron, 1999)	Alternative pathway found to produce target by adding methylglyoxal reductase and lactaldehyde reductase to <i>E. coli</i>
		Introduced glycerol dehydrogenase and methylglyoxal synthase genes from <i>E. coli</i> to <i>S. cerevisiae</i>	(W. Lee and Dasilva, 2006)	Alternative pathway found to produce target by adding methylglyoxal reductase and lactaldehyde reductase to <i>S.</i> <i>cerevisiae</i>
Itaconate; itaconic acid; methylenesuccinic acid	C00490	No information	No information	EC 4.2.1.4-citrate dehydratase and EC 4.1.1.6-aconitate decarboxylase were found to be added to <i>E. coli</i> as the host.
<i>cis,cis</i> -Muconate; <i>cis,cis</i> -hexadienedioate; <i>cis,cis</i> -2,4- hexadienedioic acid	C02480	Introduced aroZ, aroY, and catA to <i>E. coli</i>	(Niu et al., 2002)	Alternative pathways from antharnilate or 2,3-dihydroxybenzoate to produce catechol, which is a substrate for <i>cis,cis</i> -muconate production
Adipate; adipic acid; hexanedioate; hexan-1,6-dicarboxylate	C06104	Introduced aroZ, aroY, and catA to <i>E. coli</i> for producing <i>cis,cis</i> -muconate and then convert to adipic acid by chemical synthesis	(Niu et al., 2002)	Alternative pathway found to produce the target by adding 5 heterologous reactions to <i>E. coli</i> or <i>C. glutamicum</i> as the hosts (see Additional files 4 and 5 in (Chatsurachai et al., 2012) for enzyme information)

Nonnative metabolites	Host network	By-product	No. of reaction(s)	Heterologous reaction(s)	EC number
Pentane-2,4-dione	E. coli, S. cerevisiae	Oxygen	1	Pentane-2,4-dione + oxygen ↔ acetate + methylglyoxal	1.13.11.50
	C. glutamicum	Oxygen	2	Glycerone phosphate ↔ methylglyoxal + orthophosphate	4.2.3.3
				Pentane-2,4-dione + oxygen ↔ acetate + methylglyoxal	1.13.11.50
Vanillin	E. coli, C. glutamicum	Oxygen, NADH	4	Formaldehyde + NAD+ + H ₂ O \leftrightarrow formate + NADH + H ⁺	1.2.1.46
(4-hydroxy-3-	Ū			3-Dehydroshikimate ↔ 3,4-dihydroxybenzoate + H2O	4.2.1.118
metnoxy- benzaldehyde)				Vanillate + oxygen + NADH + $H^+ \leftrightarrow 3,4$ -dihydroxybenzoate + NAD+ + H_2O + formaldehyde	1.14.13.82
				Vanillate + NAD+ + H2O \leftrightarrow 4-hydroxy-3-methoxy-benzaldehyde + oxygen + NADH + H ⁺	1.2.3.9
(R)-Propane-1,2-diol	E. coli	NAD^{+}	2	(R)-Lactaldehyde + NAD ⁺ + H ₂ O \leftrightarrow (R)-lactate + NADH + H ⁺	1.2.1.23
				(R)-Propane-1,2-diol + NAD ⁺ \leftrightarrow (R)-lactaldehyde + NADH + H ⁺	1.1.1.77
2-Propyn-1-al	S. cerevisiae	NAD^{+}	3	3-Oxopropanoate \leftrightarrow acetaldehyde + CO ₂	4.1.1
				3-Oxopropanoate ↔ propynoate + H ₂ O	4.2.1.27
				2-Propyn-1-al + NAD ⁺ + H ₂ O ↔ propynoate + NADH + H ⁺	1.2.1.3

Table 2.3 Examples of producible nonnative metabolites under conditions of maximized biomass production

ser	Adipate nialdehyde	E. coli	NADP+	6	Succinyl-CoA + acetyl-CoA ↔ CoA + 3-oxoadipyl-CoA	2.3.1.174
					(3S)-3-Hydroxyadipyl-CoA + NAD ⁺ \leftrightarrow 3-Oxoadipyl-CoA + NADH + H ⁺	1.1.1.35
					5-Carboxy-2-pentenoyl-CoA + H2O \leftrightarrow (3S)-3-hydroxyadipyl-CoA	4.2.1.17
					$\label{eq:Adipyl-CoA + FAD} \leftrightarrow \text{5-carboxy-2-pentenoyl-CoA + FADH}_2$	1.3.99
					Adipate + CoA + ATP ↔ Adipyl-CoA + AMP + diphosphate	6.2.1
					Adipate semialdehyde + NADP+ + $H_2O \leftrightarrow adipate + NADPH + H^+$	1.2.1.4

2.3.3 Differences in target production capacity among host microorganisms

While screening for heterologous pathways to produce the target metabolites discussed earlier, differences in production capacity between the 3 host microorganisms emerged; for example, a group of metabolites was producible by the addition of heterologous reactions to one of the hosts, but was not produced by the other hosts. To characterize the differences in target production capacity, the producible metabolites (the Additional files 5, 6, 7 of supplementary of the publication (Chatsurachai et al., 2012)) were categorized using the KEGG Orthology database (Kanehisa et al., 2008). A chi-square statistical analysis was then performed to identify the categories in which the frequency of producible metabolites is significantly higher than expected. Fig. 2.5 shows the 10 categories that demonstrated significant differences (P<0.001). As shown in the figure, metabolites belonging to 5 categories, namely, "tyrosine metabolism", "dioxin degradation", "benzoate degradation", "chlorocyclohexane and chlorobenzene degradation", and "xylene degradation", tended to be producible by S. cerevisiae and C. glutamicum, but were non-producible in E. coli cells.

Similarly, the metabolites in "flavonoid biosynthesis", "phenylpropanoid biosynthesis", and "nicotinate and nicotinamide metaoblism" were preferentially generated by *E. coli* and *C. glutamicum*. Metabolites assigned to "porphyrin and chlorophyll metabolism" also tended to be produced in *C. glutamicum* cells. Likewise, the metabolites assigned to "biosynthesis of 12-, 14-, and 16-membered macrolides" were produced preferentially in *E. coli* cells. Such differences in production capabilities result from the different metabolic

pathways by which the hosts produce necessary substrates, and from cellular compartmentalization in the yeast strain (which is absent in the bacterial strains).

In yeast cells, the compartments present barriers to metabolite transport. For instance, mitochondrial/cytoplasmic interfaces prohibit the production of certain target metabolites when sugar is used as a carbon source. Similarly, the production of metabolites in the "flavonoid biosynthesis" category was inhibited in yeast cells because the transportation of 4-coumarate between the mitochondria and the cytosol is not permitted; therefore, the yeast strain could not produce p-coumaroyl-CoA (required for making chalconoid, an important ingredient in flavonoid biosynthesis). The genome-scale metabolic model used in this study does not account for transportation capabilities between compartments, which are currently unclear for many metabolites, and which might influence the production capacities of target metabolites in real cell systems.



FIG. 2.5 The number of producible and non-producible metabolites in functional categories that exhibit significant differences between host microorganisms. The blue and red bars represent the non-produced and produced metabolites respectively, under conditions of maximized target production.

 \sim

2.4 Summary

In this chapter, the *in silico* pathway search algorithm for target production was developed, in which iterative additions of heterologous metabolic reactions to host's metabolic networks enable target productions. Biosynthetic capabilities are also evaluated by pathway design and metabolic flux calculations. The 3 industrial host cell factories, *E. coli, S. cerevisiae*, and *C. glutamicum* were used as templates. The screened heterologous pathways by using the developed algorithm were validated. The results were consistent with earlier reports. In addition, alternative heterologous pathways that are no reports so far were also suggested, including the production pathways of itaconate, cis,cis-muconate, adipic acid, and so on. Since these compounds are important in industrial chemicals (Table 1.1), these alternative pathways could be options of metabolic engineering strategies in order to produce/improve industrial metabolites by bio-based process.

The computational platform developed in this chapter has included the features which are host-specific heterologous pathways and selection of heterologous genes by using K_m value and provides a catalog of nonnative metabolites including industrial chemicals for specific host cell. However, K_m values are generally obtained from experiments and many enzymes have no information on K_m in the database. Furthermore, in some cases, the heterologous genes are not expressed or low expressed, which can result low production rate of target metabolites. In addition, the catalog of nonnative metabolites in this chapter is available as a table in Microsoft® Excel, which is difficult to search for the information of heterologous genes.

In chapter 3, I will deal with these problems for improving the selection scores of heterologous genes particular for each host cell and improving the catalog of nonnative metabolites into well-format to provide heterologous genes and compounds information.

Chapter 3

Selection of heterologous genes using CAI score

3.1 Introduction

The production of industrial compounds using microorganisms as cell factories has presently become attractive due to the potential depletion of petroleum resources. Metabolic engineering to incorporate heterologous pathways to host cell factories is one major way to produce and improve target chemicals and fuels via bio-fermentation process. In chapter 2, the development of the computational platform are presented, which enables to screen heterologous pathways of nonnative metabolites and used K_m as a parameter to select heterologous genes corresponding to heterologous pathways. Still, a large number of enzymes has no information on K_m values, thus a new and efficient parameter is desirable to select candidate heterologous genes from among numerous of orthologous enzymes with similar enzymatic activities. In addition, the inefficient target production by the introduced heterologous genes could be caused by a low or lack of expression of heterologous enzymes in the host cell. To overcome this problem, heterologous enzymes which have the potential to be highly expressed should be chosen from the screened candidates. It has been established that several factors namely, genome GC content, codon usage, and mRNA secondary structures influence the expression of heterologous enzymes. Amongst these factors, codon usage is known to cause a

relatively high impact on the enzymatic expression levels. In fact, it is demonstrated there is the bias of codon used in high and low expressed genes in several organisms (Gupta et al., 2004; G. Liu et al., 2010; Sharp et al., 1986). In addition, Botzman and Margalit discovered that the global extent of codon usage bias of an organism plays a crucial role in the adaptation of prokaryotes to their environments (Botzman and Margalit, 2011). A few species such as *E. coli* (M.-S. Lee et al., 2011) and *S. cerevisiae* (Norkiene and Gedvilaite, 2012) exhibited enhanced heterologous protein expressions when the codons usage of heterologous genes were replaced with a set of more suitable host codons. Several methods for analyzing codon usage bias have been developed in order to study molecular evolution and heterologous gene/protein expression (Ingvarsson, 2008; Olivares-Hernández et al., 2011; W. Qian et al., 2012; Sharp and Li, 1987; Tao et al., 2009). Codon Adaptation Index (CAI) (Sharp and Li, 1987) is one of the most widely used to estimate the extent of codon usage bias in genes (Martín-Galiano et al., 2004; Nayak, 2012) and proteins (Futcher et al., 1999; Washburn et al., 2001) based on their expression levels. Previous reports employed CAI for optimizing DNA vaccines (Mani et al., 2011), enhancing and altering exogenous and endogenous protein expressions (W. Li et al., 2011; Redemann et al., 2011). These studies suggested that CAI can be used for the selection of heterologous enzymes which is expected to be highly expressed in the host microorganisms.

In this chapter, the computational platform for heterologous pathway search is expanded by incorporating the CAI measure of heterologous genes to select appropriate heterologous enzymes, whose introduction into host microorganisms would enable efficient target metabolite production. Furthermore, I design the optimized gene sequences for such enzymes by substituting the most frequent codons found in host's highly expressed genes. By integrating the simple in silico screening platform (for identifying feasible heterologous pathways developed in chapter 2) with the selected heterologous genes as well as the optimized gene sequences, based on a target host's preferable codons, a rational heterologous pathway design system named "ArtPathDesign" (Artificial heterologous Pathway Design) is proposed for an efficient production of nonnative metabolites. The ArtPathDesign system applied to 3 hosts E. coli, S. cerevisiae and B. subtilis which are recognized as industrial host producers. Using this system, all producible nonnative metabolites of E. coli, S. cerevisiae, and B. subtilis were obtained along with specific information regarding the feasible heterologous enzymes. Furthermore, the catalog of nonnative metabolites which can be produced by each host is improved from Microsoft[®] Excel-format to Hyper Text Markup Language (HTML)-format that is a well-demonstration of heterologous pathway, gene, and compound information.

3.2 Materials and methods

3.2.1 Constructing an in-house database of metabolic reactions

All known metabolic reactions were considered as candidate heterologous reactions that could be added to the host's metabolic network. An in-house database of metabolic reactions was firstly constructed from data stored in BKM-react (Lang et al., 2011), which is an integrated database resulting from a

combination of database namely, BRENDA (Chang et al., 2009), KEGG (Kanehisa et al., 2008), and MetaCyc (Caspi et al., 2008). This in-house database is the update version of previously constructed that used metabolic reactions from retrieved only from KEGG database. All metabolic reaction information regarding genes, enzymes, pathways, and organisms in the KEGG and the EMBL (Kulikova et al., 2004) databases were collected into the new version of the in-house database developed using PostgreSQL 9.0 (The PostgreSQL Global Development Group).

3.2.2 Screening for artificial heterologous pathways involved in the production of targets

Genome-scale metabolic models of *E. coli* (iJR904)(Reed et al., 2003), *S. cerevisiae* (iMM904) (Mo et al., 2009), and *B. subtilis* (Oh et al., 2007), were used based on earlier metabolic reconstructions with slight modifications. In order to incorporate known metabolic pathways listed in BKM-react database, all metabolite IDs in the earlier genome-scale metabolic models were converted to the KEGG compound ID and MetaCyc compound ID format using metabolic name matching using python scripts and manually search. In this study, the algorithm presented in chapter 2 was applied in order to search heterologous reaction(s) that produce target metabolites within a host microorganism.

3.2.3 Codon Adaptation Index (CAI)

CAI was proposed by Sharp and Li as a measure of synonymous codon bias calculated based on the codon preference of highly expressed proteins, such as ribosomal proteins, and elongation factors (Sharp and Li, 1987). CAI score of a

gene is a measure of favorable codons or optimal codons, which are frequently used by the highly expressed proteins in a given genome, which takes a value from 0 to 1. CAI for a gene with L-amino acids is formulated as follows:

$$CAI = \left(\prod_{i=1}^{L} w_i\right)^{1/L}$$
$$w_i = \frac{f_i}{max(f_j)}$$

where, f_i denotes the frequency of i^{th} codon among other synonymous codons coding the same amino acids in the set of highly expressing genes. The relative adaptiveness w_i is defined by the ratio between the frequency of i^{th} codon and that of the most frequently used codon for the amino acid. CAI of a protein is defined as the geometric mean of the relative adaptiveness of all the codons in the coding sequence. For the calculation of CAI, a set of highly expressed genes in the hosts, *E. coli*, *S. cerevisiae*, and *B. subtilis* was collected from literature (Das et al., 2009) and database (Puigbò et al., 2008). The w_i table was created for each host by using sequence of highly expressed genes in the corresponding host strain (see in Table B.1 in Appendix B). The CAI score of orthologous genes of all organisms in KEGG and EMBL database was then calculated. The overall strategy of the computational platform integrating a new selection score, named as ArtPathDesign, developed in this thesis is summarized in Fig. 3.1.



FIG. 3.1 The schematic representation of ArtPathDesign platform.

Host metabolic network and metabolic reactions collected in the in-house database were used as input data for screening heterologous pathways of nonnative metabolites. In parallel, CAI scores of all genes retrieving from the available databases was calculated and used to select candidate genes. Finally, the catalog of nonnative metabolites will be generated and available in html-format.

3.3 Results and discussion

3.3.1 Identification of heterologous pathways

Statistics of the updated version of the in-house database is shown in Table 3.1. The updated in-house database contains 17,488 known metabolic reactions, 17,617 metabolites of 2,056 organisms retrieved from BKM-react (shown in Table S1 of supplementary data). All supplementary data (Table S1-S4) in this chapter are available on the web site below.

http://www-shimizu.ist.osaka-u.ac.jp/supplementaryData_artpathdesign.zip

Necessary information regarding genes, enzymes, nucleotide sequences, organisms as well as the reversibility data from KEGG, BRENDA, and EMBL databases were parsed and stored in a version 2.0 of the in-house database.

Table 3.1 Statistics of the in-house database version 1.0 (chapter 2) and version2.0 constructed in this chapter

	In-house database	In-house database
	(version 1.0)	(version 2.0)
Source database	KEGG	BKM-react
No. of reactions	7,769	17,488
No. of reactions (no EC number)	1,274	0
No. of compounds	6,635	17,617
No. of reversible reactions	7,769	17,488

To archive nonnative production in *E. coli*, *S. cerevisiae*, and *B. subtilis* as the hosts, the algorithm developed in chapter 2, was used to screen heterologous

pathways from the metabolic reactions collected in the in-house database. In brief, by employing this algorithm the host's metabolic network were iteratively expanded by adding heterologous reactions step-by-step in order to link nonnative metabolites. The host's metabolic network expanded until no further nonnative metabolites could connect to it. Fewer than 32 heterologous reactions were required to connect 3,226, 3,298, and 3,211 nonnative metabolites to the host's metabolic networks of E. coli, S. cerevisiae, and B. subtilis respectively. Comparison of results between version 1.0 and 2.0 of the in-house databases are shown in Table 3.2. From this table, it was found in all hosts excluding *E. coli*, the update version of the in-house database was improved as for the number of connectable metabolites. The total number of connectable metabolites when E. coli as the host decreased from 3,244 to 3,226 because, removing metabolic reactions, which are no information of EC number, previously used to connect to E. coli's metabolic network in chapter 2. Those metabolites included D-Ribulose (C00309), Hyaluronate (C00518), etc. If no EC number information, it is not possible to select heterologous genes. Therefore, the in-house database version 2.0 contains only metabolic reactions that have information of EC number. The list of metabolites connected to the host's metabolic networks by using the new version in-house database is shown in Table S2 of supplementary data.

Host cell	In-house database	In-house database
	(version 1.0)	(version 2.0)
E. coli	3,244	3,226
S. cerevisiae	3,154	3,298
C. glutamicum	3,112	3,321
B. subtilis	3,063	3,211

 Table 3.2 Comparison of total connectable nonnative metabolites between 2

 versions of the in-house databases

3.3.2 Relationship between CAI score and protein abundance

To analyze the relationship between CAI scores and protein expression levels, protein abundance data were obtained from PaxDb (M. Wang et al., 2012), which is a repository containing information from different proteome experiments of several organisms. 1,560 coding gene sequences (CDSs) of *E. coli* were obtained from KEGG database. Likewise, 5,004 CDSs of *S. cerevisiae* and protein abundance were collected for correlation analysis. CAI scores of genes were then computed and compared them with the protein abundance data as shown in Fig. 3.2. A statistically significant correlation was observed between CAI scores and log (protein abundance values) (correlation coefficient r=0.60 in *E. coli* and r=0.65 in *S. cerevisiae*). The observed positive correlation of CAI and protein abundances suggested that the CAI could be used as the index representing potential protein expression levels in host organisms.



FIG. 3.2 The relationship between the normalized log protein abundance data (ppm: part per million) and CAI scores in (A) *E. coli* (B) *S. cerevisiae*. CAI scores of 1,560 genes in *E. coli* and 5,004 genes in *S. cerevisiae* from PaxDb (Protein Abundance Across Organisms) database are shown. Protein abundance in "ppm" essentially refers to each protein with reference to the complete proteome expression. In other words, it implies the amount of each protein relative to all other protein molecules present in the sample.

3.3.3 Screening of heterologous genes with higher CAI scores

The expression levels of heterologous enzymes are often low (Kleber-Janke and Becker, 2000; Lakey et al., 2000), probably due to the differences in the codon usage between the source and the host organisms. However, based on the positive correlation obtained between CAI scores and protein abundance values (Fig. 3.2), it can be inferred that, amongst the multiple candidate heterologous enzymes involved in the production of a target metabolite, selecting the one
whose corresponding gene sequence has a higher CAI score would result in an enhanced expression of the heterologous enzyme.

In this chapter, potentially highly expressed genes that could enhance the expression level of the corresponding encoded enzymes were identified, and thereby result in a higher productivity of the target metabolites. The process involved determining the CAI scores for all the screened candidate heterologous genes which could lead to the production of all possible target metabolites, and subsequently selecting the candidate heterologous genes with higher CAI scores. Expectedly, for all host microorganisms used in this study, the distributions of CAI scores of the native genes were found to be higher when compared to those of the heterologous genes from other organisms (Fig. 3.3).

It was reported that human genes selected to express in *E. coli* as the host showed low expression (Dai et al., 2013; Gvritishvili et al., 2010; Q. Wang et al., 2012). Thus, I collected the wide type coding sequences of human kallistatin, human Pigment epithelium-derived factor (PEDF), and human cystatin from the previous reports, and calculated CAI scores of these genes. As results, CAI scores of the 3 human genes selected to be expressed in *E. coli* are 0.37, 0.32, and 0.45, respectively. These data suggests that the human genes show low expression due to low CAI scores. In addition, these previous reports also successfully improved the production of proteins by replacing the human genes with the codons that suitable for host *E. coli* without changing amino acids.

In Fig. 3.3, the dashed line represents the distribution of CAI scores of different native genes that originally exist in the host, while the solid line

corresponds to the CAI scores of all possible heterologous genes which could be utilized for the production of the target metabolite. The dotted line represents the CAI scores of the selected heterologous genes. It was observed that the highest CAI scores correspond to the set of heterologous genes which encode enzymes having same functions. As evident, a significant increase in CAI scores was achieved by repeated selection process, thus demonstrating that this approach can overcome the problem of low expression of heterologous enzymes which occur due to differences in the codon usage. The selected heterologous genes required for the production of all possible target metabolites are presented in Table S3 of supplementary data.

Fig. 3.4 illustrates 2 different heterologous pathways where the higher CAI score selection technique was applied for choosing suitable heterologous genes. The production of 1,3-propanediol by *S. cerevisiae* (Rao et al., 2008) is generally achieved by introducing *dhaB* (glycerol dehydratase) from *Klebsiella pneumoniae* and *yqhD* (an alcohol dehydrogenase possessing the activity of 1,3-propanediol dehydrogenase) from *E. coli*. As shown in Fig. 3.4A, the CAI scores of *dhaB* and *yqhD* in *S. cerevisiae* is relatively low in comparison with the native genes, which might result in low expression levels of these heterologous genes. However, this screening platform demonstrated that the productivity of 1,3-propanediol by the host *S. cerevisiae* could be enhanced by introduction of *dhaB* and *dhaT* from *Clostridium perfringens* and *Lactobacillus reuteri*, respectively instead of *dhaB* and *yqhD* from *K. pneumoniae* and *E. coli*, respectively since they have relatively high CAI scores. Another classical example is the heterologous pathway for (R,R)-2,3-butanediol in a host *E. coli*, which has been shown to utilize heterologous

genes from *B. subtilis* (Yan et al., 2009). In this case, the *in silico* system suggested that *alsS* from *Pseudomonas putida*, *alsD* from *Enterobacter cloacae* and *bdh* from *Klebsiella oxytoca* were comparatively better in terms of CAI, shown in Fig. 3.4B. Although other enzymatic characteristics such as K_m value could influence the production of target metabolites, the expression level of a particular heterologous enzyme is crucial for the activation of a specific heterologous pathway in order to achieve a higher productivity of the desired target. Therefore, these findings highlight that the selection of heterologous genes with higher CAI scores is an effective approach to enhance the productivity of the target metabolites.

In addition, the distributions in Fig. 3.3 reveal that in spite of selecting heterologous genes with the highest CAI scores (shown in dotted line) from many organisms, some heterologous genes still exhibit relatively low CAI scores which could result in a low expression level due to the mismatch of codon usage. One possible strategy to overcome this is to calculate an optimized gene sequence encoding a specific heterologous enzyme which has preferable codon usage in the host microorganism (i.e., CAI = 1), and subsequently synthesize this artificial gene experimentally for its introduction into the host. With the growing availability of low-cost de novo gene synthesis (e.g., \$300/1kb in 2012), such optimized gene sequences could be easily employed for bio-production. Fig. 3.5 shows the example of optimized gene sequence comparing with the original gene sequences encoding for catechol 1,2-dioxygenase (*catA*).The optimized sequences encoding all the heterologous enzymes that are required for the production of target metabolites in the 3 host microorganism are given in Table S4 of

supplementary data. Collectively, these data suggest that artificial gene sequences when considered in combination with optimal CAI are valuable, only when the artificial gene synthesis technique is applied to the expansion of the host's metabolic network for the production of targets.



FIG. 3.3 Distribution of CAI scores for each organism (A) *E. coli*, (B) *S. cerevisiae*, and (C) *B. subtilis* as the hosts. Solid line denotes the CAI score distribution of all possible heterologous genes that can be used for nonnative metabolite productions. Dotted line represents the CAI score distribution of all selected heterologous genes with the highest CAI score encoding enzymes having the

same enzymatic function. Dashed line denotes the CAI score distribution of native genes originally from the host microorganism.



FIG. 3.4 (A) Heterologous pathway for the production of 1,3-propanediol in *S. cerevisiae* (Rao et al., 2008) as well as CAI scores of orthologous genes. The heterologous genes from different organism with the higher CAI scores (shown in box) were selected by employing the improved computational system. (B) Heterologous pathway for the production of (R,R)-2,3-butanediol in *E. coli* (Yan et al., 2009) as well as CAI scores of orthologous gens. The heterologous genes from different organism with the higher CAI scores denote the production of (R,R)-2,3-butanediol in *E. coli* (Yan et al., 2009) as well as CAI scores of orthologous gens. The heterologous genes from different organism with the higher CAI scores (shown in box) were selected by employing the improved system. *alsD* encodes acetolactate decarboxylase, *alsS* encodes acetolactate synthase, and *bdh* encodes butanediol dehydrogenase.

Psest_2692_org	а	t	g	alo	t	g	t	а	a	ala	a	t	с	t	с	с	с	a a	a a	c	с	а	g	c	gla	a	g	t	t	с	a	ga	ala	I C	t	t	с	t	t	с	[4	5]
Psest_2692_opt	а	t	g	a c	c	g	t	t	a	a a	a	t	с	t	с	t	с	a g	<mark>j</mark> a	ı c	с	t	c	t	g a	ac	g	t	t	с	a (g a	aa	I C	t	t	с	t	t	с	[4	5]
Psest_2692_org	а	a	g	g a	a	g	с	с	a	g o	c g	g	с	t	t	с	g	g c	a	a	с	g	a	c	g o	c c	g	g	с	a	g (са	a g	j c	с	g	с	а	t	g	[9)0]
Psest_2692_opt	а	a	a	g a	a	g	с	g	t (c t	g	g	t	t	t	с	g	g t	а	a	с	g	a	c	g o	e g	g	g	t	t (c t	t t	c	: t	с	g	t	а	t	g	[9	0]
Psest_2692_org	а	a	g	ac	с	g	t	g	a t	t	a	a	c	с	g	с	a t	t	: t	t	g	g	t	c	g a	ac	а	с	t	g	cļ	g a	a	ı a	a	t	с	g	t	с	[1:	35]
Psest_2692_opt	а	a	a	ac	с	g	t	t	a t	t	a	a	c	с	g	t	a t	t	c	t	g	g	t	t	g a	a	а	с	с	g	c (ga	a a	ı a	a	t	с	g	t	t	[1:	35]
Psest_2692_org	g	a	a	g a	c	с	t	g	g	a a	a	t	с	а	с	с	с	a g	9 9	, a	с	g	a	a t	t	c	t	g	g	a	a	g g	g c	: c	g	t	g	g	а	с	[1	80]
Psest_2692_opt	g	a	a	g a	c	с	t	g	g	a a	a	t	с	а	с	с	с	a g	9 9) a	с	g	a	a t	t t	c	t	g	g	a	a a	a g	g o	: g	g	t	t	g	а	с	[1	80]
Psest_2692_org	t	at	t	c t	с	а	а	с	c	g o	c c	t	g	g	g	с	g	g t	c	g	с	с	a	c	g a	a	g	с	с	g	g t	t	: t	g	, c	t	g	g	t	t	[22	25]
Psest_2692_opt	t	a	c	c t	g	а	а	с	c	g t	с	t	g	g	g	t	g g	g t	c	; g	t	с	а	c	g a	a	g	с	g	g	g t	t	: t	g	, c	t	g	g	t	t	[2;	25]
Psest_2692_org	g	с	c	g g	с	с	t	g	g	g o	c c	t	g	g	а	g	с	ac	: t	t	с	с	t	c	g a	a	с	t	g	c t	t g	g (g a	ı ə	g	a	с	g	с	с	[2]	70]
Psest_2692_opt	g	с	g	g g	t	с	t	g	g	g t	с	t	g	g	а	a	с	ac	: t	t	с	с	t	g	g a	ac	с	t	g	c t	t g	g	g a	1 8	g	a	с	g	с	g	[2]	70]

FIG. 3.5 The catA sequence (ORF ID: Psest_2692) from *Pseudomonas stutzeri* RCH2 was replaced with the host's favorable codons, for example codon "act" was replace with "acc" coding for threonine. The GC content of original sequence is 64.65%, while GC content of optimized sequence is 57.64%.

Finally, FBA was performed to investigate whether all connected nonnative metabolites are producible by using glucose as a sole carbon source. The maximum fluxes of all connectable nonnative metabolites corresponding to each host were calculated. It was observed that 11% of the connectable nonnative metabolites of *E. coli* could not be produced using glucose as the sole carbon source. Likewise, 27% and 11% of connectable nonnative metabolites of *S. cerevisiae* and *B. subtilis* respectively could not be produced under this condition.

3.3.4 Examples of results in hyper-text based user interface

HTML (Hyper Text Markup Language) is a language for describing web pages, thus outputs in html-format easily open by using web browser such as Google Chrome and Firefox. Python script function was created in order to provide information of heterologous pathways of nonnative metabolites in html-based format. The hyper-text is automatically generated by these programs and output results are summarized in the file named "index.html" found in archive file available at: http://www-shimizu.ist.osaka-u.ac.jp/APD.zip

In order to search for target metabolite, the index.html can be opened via web browser for example Google Chrome as shown in Fig. 3.6. In this figure, how to find heterologous pathway(s) for the production of 1,3-propanediol using search function (Ctrl+F) of Google Chrome is shown. By typing compound name or ID, the possible pathways will be found as shown in Step 3 of Fig. 3.6. By clicking target pathway (for example Pathway-1), the detail of the heterologous pathway will be displayed as shown in Step 4 of Fig. 3.6. Additional example of (R,R)-2,3-butanediol is shown in Fig. 3.7. By using this computational system, the list of all possible metabolites able to connect to host's metabolic network will be provided as a catalogue for user to search for target production as well as necessary information such as candidate genes, open reading frame ID, and organisms.

Step 1. Open "index.html" The index.html can open via Google Chrome

ArtPathDesign

Index of nonnative metabolites for E. coli

Page created by: Sunisa Chatsurachai

Date generated: 2013-06-07

Nonnative metabolite	Heterologous pathway(s)				
(E)-1,2,4-But-1-enetricarboxylic acid trans-Homoaconitate C16579	Pathway-1]			
Ribulose C05052	Pathway-1]			
GDP-D-glucose GDP-alpha-D-glucose GDP-glucose C00394	Pathway-1	Pathway- 2	Pathway- <u>3</u>	Pathway- 4	Pathway <u>5</u>
N6-Hydroxy-L-lysine C01028	Pathway-1	Pathway- 2	<u>Pathway-</u> <u>3</u>		
ditrans,heptacis-Decaprenyl diphosphate di-trans,poly-cis-Decaprenyl diphosphate C04509	Pathway-1]			
N,N-Dimethylglycine Dimethylglycine C01026	Pathway-1	Pathway- 2	Pathway- <u>3</u>	<u>Pathway-</u> 4	<u>Pathway</u> <u>5</u>
Thiocyanic acid Thiocyanate C01755	Pathway-1	Pathway- 2	Pathway- <u>3</u>		

FIG. 3.6 The output result of heterologous pathways for the production of 1,3propanediol in *E. coli* is provided in html-format. To search for target metabolite, 4 steps are required and shown here. Step 1 shows "index.html", which is the summarized results of ArtPathDesign (the html file can be easily opened using Google Chrome and Firefox). In this example, the index.html was open by using Google Chrome. Step 2 shows how to search target metabolite using Ctrl+F (search function of Google Chrome) and type compound name or compound ID of target. Step 3 shows the result of target search and two heterologous pathways are available for 1,3-propanediol. Step 4 shows result page of the first candidate pathway in the new page including a structure of compound and detail information of heterologous pathway that are reactions, genes, open reading frame ID, and organisms.

Step 2. Press Ctrl+F (search function of Google Chrome) and Type 1,3-propanediol

← → C [] file:///E:/APD_eco/index.html					© ☆
8047		· ·	4 1,1	3-propanediol	1 of 2 🥎 🖌
Manganese(2+) Mn(II) Mn2+ C19610		Pathway-1	<u>Pathway-</u> 2	Zoom sea	arch
Mn(III) Manganese(3+) Mn3+ C19611		Pathway-1	Pathway 2		
dTDP-3-dehydro-6-deoxy-alpha-D-galactopyranose dTDP-6-deoxy ulose dTDP-6-deoxy-D-xylo-hex-3-ulose C19960	1,3-propa	nediol		1 of 2	
allopurinol CPD-9024		<u>с ангway-1</u>			
9-Fluorenol Fluoren-9-ol 9-Hydroxyfluorene Diphenylene carbin	ol C06711	Pathway-1	Pathway- 2	Pathway- Pat 3 4	thway- Pathw 5
Indole-3-acetamide (Indol-3-yl)acetamide C02693		Pathway-1			
4-Hydroxyphenylacetyl-CoA C05338		Pathway-1]		
Trimethylene glycol <mark>1,3-Propanediol</mark> Propane-1,3-diol C02457		Pathway-1	Pathway- 2		
(S)-Scoulerine (13aS)-5,8,13,13a-Tetrahydro-3,10-dimethoxy-6H- dibenzo[a,g]quinolizine-2,9-dio1 C02106		Pathway-1	<u>Pathway-</u> 2	Pathway-Pat 3 4	thway- Pathw 5
Butein C08578		Pathway-1	<u>Pathway-</u> 2		
Phenylethylamine Phenethylamine 2-Phenylethylamine beta- Phenylethylamine C05332		Pathway-1			
2-O-[2-O-(alpha-D-Mannopyranosyl)-alpha-D-glucopyranosyl]-3-p glycerate C19793	hospho-D-	Pathway-1]		

Step 3. Red box shows 2 heterologous pathways for 1,3 propanediol in *E. coli*.

← → C D file:///E:/APD_eco/index.html					Q 😭	=
8047		4	1,3-propanediol	10	f2 ^ ~ ×	1
Manganese(2+) Mn(II) Mn2+ C19610	<u>Pathway-1</u>	<u>Pathwa</u> 2	<u>y-</u>			
Mn(III) Manganese(3+) Mn3+ C19611	<u>Pathway-1</u>	Pathwa 2	<u>y-</u>			-
dTDP-3-dehydro-6-deoxy-alpha-D-galactopyranose dTDP-6-deoxy-D-xylo-hexos-3- ulose dTDP-6-deoxy-D-xylo-hex-3-ulose C19960	Pathway-1					
allopurinol CPD-9024	Pathway-1					
9-Fluorenol Fluoren-9-ol 9-Hydroxyfluorene Diphenylene carbinol C06711	<u>Pathway-1</u>	Pathwa 2	y- Pathway- <u>3</u>	Pathway- 4	Pathwa 5	2
Indole-3-acetamide (Indol-3-yl)acetamide C02693	Pathway-1					
4-Hydroxyphenylacetyl-CoA C05338	Pathway-1]				
Trimethylene glycol <mark>1,3-Propanediol</mark> Propane-1,3-diol C02457	<u>Pathway-1</u>	<u>Pathwa</u> 2	<u>v-</u>			
(S)-Scoulerine (13aS)-5,8,13,13a-Tetrahydro-3,10-dimethoxy-6H- dibenzo[a,g]quinolizine-2,9-diol C02106	Pathway-1	<u>Pathwa</u> 2	<u>y-</u> Pathway- <u>3</u>	Pathway- 4	<u>Pathwa</u> <u>5</u>	ż
Butein C08578	Pathway-1	<u>Pathwa</u> 2	<u>y-</u>			
Phenylethylamine Phenethylamine 2-Phenylethylamine beta- Phenylethylamine C05332	Pathway-1					
2-O-[2-O-(alpha-D-Mannopyranosyl)-alpha-D-glucopyranosyl]-3-phospho-D- glycerate C19793	Pathway-1]				

FIG. 3.6 (Continued)



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Step 4. Click Pathway-1, the result page will display.

FIG. 3.6 (Continued)

Heterologous Pathway for C03044 in E. coli (Glucose as a carbon source)

Page created by: Sunisa Chatsurachai

Date: 2013-06-07

Target compound ID	000044
Target compound Name	(R,R):Butane-2,3-diol/(R,R)-2,3-Butanediol/(R,R)-2,3-Butylene glycol
List of Heterologous reaction(s)	R07477]R01563[R10025
Detail of heterologous reaction	R07477 (R.R)-butane-2,3-diol + NAD+ <=> (R)-acetoin + NADH + H+ 1 C03044 + 1 C00003 <=> 1 C00810 + 1 C00080
	Erzyme information: 1.1.1.4 Gene symbol: butB;butA;BDH from database: KB0G Ist-candidate gene(s): CAI= 0.647;ES15_1123;csk;Cronobacter sakazakii ES15 (R;R)-butanediol dehydrogenase;diacetyl reductase 2nd-candidate gene(s): CAI= 0.647;KOX_01940;KOX_22375;kox;Klebsiella oxytoca KCTC 1686 (R;R)-butanediol dehydrogenase;diacetyl reductase;(R;R)-butanediol dehydrogenase;diacetyl reductase 2nd-candidate gene(s): CAI= 0.645;KOX_01940;KOX_22375;kox;Klebsiella oxytoca KCTC 1686 (R;R)-butanediol dehydrogenase;diacetyl reductase;(R;R)-butanediol dehydrogenase;diacetyl reductase 3rd-candidate gene(s): CAI= 0.645;SSP291_04195;csx;Cronobacter sakazakii Sp291 (R;R)-butanediol dehydrogenase;diacetyl reductase;(R;R)-butanediol dehydrogenase;diacetyl reductase
Detail of heterologous reaction	R01565 2-Acetolactate <=> (R)-Acetoin + CO2 1 C00900 <=> 1 C00810 + 1 C00011 4.1.1.5 Gene symbol: alsD,E4.1.1.5 from database: KEOO 1st-candidate gene(s): CAI= 0.614ECL_03127,enc,Enterobacter cloacae subsp. cloacae ATCC 13047 acetolactate decarboxylase 2nd-candidate gene(s): CAI= 0.596,APA32_03800,apx,Acetobacter pasteurianus IFO 3283-01-42C acetolactate decarboxylase 3rd-candidate gene(s): CAI= 0.596,APA32_03800,apx,Acetobacter pasteurianus IFO 3283-32 acetolactate decarboxylase
Detail of heterologous reaction	R10025 2 pyruvate <=> 2-acetolactate + CO2 2 c00202 <=> 1 C00900 + 1 C00011

FIG. 3.7 Heterologous pathway for the production of (R,R)-2,3-butanediol shown as html-format

3.3.5 Examples of new heterologous pathways for the production of nonnative metabolites in the specific host

In previous part, the evaluation of the ArtPathDesign system has been done, thus, this part will demonstrate the advantage of this system that could identify a new heterologous pathway of nonnative metabolites in the target host cell. For example, muconic acid (C02480) previously reported to produce in E. coli by adding 3 heterologous reactions by enzymes 3-dehydroshikimate dehydratase, protocatechuic acid decarboxylase, and catechol 1,2-dioxygenase encoded by heterologous genes aroZ, aroY and catA, respectively (Niu et al., 2002). The new heterologous pathway was suggested by ArtPathDesign. This pathway contained 2 heterologous reactions by enzymes anthranilate 1,2-dioxygenase (EC number: 1.14.12.1) and catechol 1,2-dioxygenase (EC number: 1.13.11.1) encoded by heterologous genes antABC and catA, respectively. In addition, the new heterologous pathway of muconic acid was just reported by Sun et al., 2013 and it is identical with the new pathway suggested by ArtPathDesign. Sun and colleagues successfully produced muconic acid in *E. coli* by introducing the new pathway containing antABC and catA from Pseudomonas aeruginosa and Pseudomonas putida KT2440 (Sun et al., 2013). As the result, it is confirmed that ArtPathDesign successfully identified the new heterologous pathway for muconic acid and validated by the recently reported experiment (Sun et al., 2013).

Another example is Isomultiflorenol production by using *E. coli* as the host (Fig. 3.8). Isomultiflorenol (Hayashi et al., 2001) is a triterpene showed similar structure with α -Amyrin and β -Amyrin (Fig. 3.9). In addition, α -Amyrin and β -

Amyrin (Liliana et al., 2012) have various pharmacological activities *in vitro* and *in vivo* conditions against various health-related conditions, including conditions such as anti-inflammation, antimicrobial, antifungal and antiviral infections. By using the ArtPathDesign system, it was demonstrated that 3 heterologous reactions are required to produce isomultiflorenol from trans,trans-Farnesyl diphosphate and 3 heterologous genes encoding 3 enzymes have activities of EC 2.5.1.21, EC 1.14.13.132, and EC 5.4.99.36. The candidate heterologous genes and open reading frame (ORF) ID are also suggested.

One more typical example is heterologous pathway for vanillin production in E. coli (Fig. 3.10). Vanillin is recognized as one of the most widely used flavoring agents in the world (Kaur and Chakraborty, 2013) which is extracted from the orchid Vanilla planifolia, Vanilla tahitiensis, and Vanilla pompona. In addition, pure vanillin used in food and beverage industry and also as a bio-preservative agent since its antimicrobial and antioxidant properties. According to vanillin properties, previous studies had design the heterologous pathway to incorporate into well-characterized hosts, E. coli, S. cerevisiae, etc. (Kaur and Chakraborty, 2013). By using the ArtPathDesign system, the new heterologous pathway and heterologous genes for E. coli as the host, which has no report today, are suggested. This heterologous pathway contains 4 heterologous reactions and 4 heterologous enzymes that have enzyme activities according to the following EC numbers: EC 1.14.13.33, EC 1.2.1.46, EC 1.14.13.82, and EC 1.2.1.67. The information of heterologous genes is shown below. Top substrates of this heterologous pathway are formate (C00058) and 4-hydroxybenzoate (C00156) which are able to be produced from the central metabolic pathway of *E. coli*.

Heterologous Pathway for C19801 in *E. coli* (Glucose as a carbon source)

н₃с сн₃ CH-HO H₃C CH₃

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Date: 2013-06-07

C19801

Target compound ID	C19801
Target compound Name	Isomultiflorenol
List of Heterologous reaction(s)	R06507 R04061 R14915
	R06507 (S)-squalene-2,3-epoxide <=> isomultiflorenol 1 C01054 <=> 1 C19801
Detail of heterologous reaction	Enzyme information: 5.4.99.36 Gene symbol: NONE from database: EMBL
	IIst-candidate gene(s): CAI= 0.277,BAB08529,NONE,Lutra aegyptiaca (smooth loofan) isomulufiorenoi synthase 2nd-candidate gene(s): CAI= NONE 3rd-candidate gene(s): CAI= NONE
	R04061 squalene + NADPH + H+ + O2 <=> (3S)-2,3-epoxy-2,3-dihydrosqualene + NADP+ + H2O 1 C00751 + 1 C00005 + 1 C00080 + 1 C0007 <=> 1 C01054 + 1 C00006 + 1 C00001
Detail of heterologous reaction	Enzyme information: 1.14.13.132 Gene symbol: SQLE;ERG1 from database: KEGG
	1st-candidate gene(s): CAI= 0.47,LBRM_13_1480,Ibz,Leishmania braziliensis squalene monooxygenase 2nd-candidate gene(s): CAI= 0.458,LMJF_13_1620,Ima,Leishmania major squalene monooxygenase 3rd-candidate gene(s): CAI= 0.454,LDBPK_131360,Ido,Leishmania donovani squalene monooxygenase
	R14915 2 trans.trans-Farnesyl diphosphate + NADPH + H+ <=> Squalene + 2 Diphosphate + NADP+ 2 C00448 + 1 C00005 + 1 C00080 <=> 1 C00751 + 2 C00013 + 1 C00006
Detail of heterologous reaction	Enzyme information: 2.5.1.21 Gene symbol: FDFT1 from database: KEGG Est-candidate represe): CAL= 0.514 LMIE_31_2040 ima Leistmania maior famesul.dinhosphate famesultransferase
	2nd-candidate gene(s): CAI= 0.502,LMXM_30_2940,lmi,Leishmania megic rainesyl-diphosphate farnesyltransferase 3rd-candidate gene(s): CAI= 0.501,Bcen_4104,bcn,Burkholderia cenocepacia AU1054 farnesyl-diphosphate farnesyltransferase

FIG. 3.8 Heterologous pathway for Isomultiflorenol (C19801) production in E. coli

was produced via trans, trans-Farnesyl diphosphate (C00448).

FIG. 3.9 Compound structures between α -Amyrin, β -Amyrin and Isomultiflorenol

(compound structures retrieved from KEGG database)

Heterologous Pathway for C00755 in E. coli (Glucose as a carbon source)

Page created by: Sunisa Chatsurachai

Date: 2013-06-07

Target compound ID	C00755
Target compound Name	Vanillin∥Vanillaldehyde∥4Hydroxy-3-methoxy-benzaldehyde∥4-Hydroxy-3-methoxybenzaldehyde
List of Heterologous reaction(s)	R13873 R09195 R00285 R01792
Detail of heterologous reaction	R13873 4 Hydroxy-3-methoxy-benzaldehyde + NAD+ + H2O <=> Vanillate + NADH + H+ 1 C00755 + 1 C00003 + 1 C00001 <=> 1 C06672 + 1 C00004 + 1 C00080
Detail of heterologous reaction	R09195 vanilate + O2 + NADH <=> 3,4 dihydroxybenzoate + NAD+ + H2O + formaldehyde 1 C06672 + 1 C00007 + 1 C00004 <=> 1 C00230 + 1 C00003 + 1 C00001 + 1 C00067
Detail of heterologous reaction	R00285 Formaldehyde + NAD+ + H2O <=> Formate + NADH + H+ 1 C00067 + 1 C00003 + 1 C00001 <=> 1 C00058 + 1 C00080 +
Detail of heterologous reaction	R01792 p-hydroxybenzoate + O2 <=> protocatechuic acid + H2O 1 C00156 + 1 C00007 <=> 1 C00230 + 1 C00001 Enzyme information: 1.1.1.24 Gene symbol: qa-3;QUIB from database: KEGG 1st-candidate gene(s): CAI= 0.492,MGG_07781,mgr,Magnaporthe oryzae quinate dehydrogenase 2nd-candidate gene(s): CAI= 0.431,ANI_1_1306034, ang.A spergillus niger quinate dehydrogenase 3rd-candidate gene(s): CAI= 0.43,ACLA_024050, act,A spergillus clavatus quinate dehydrogenase

FIG. 3.10 Heterologous pathway for vanillin production in E. coli

3.4 Summary

An improved computational or *in silico* platform, named as ArtPathDesign, was developed to screen heterologous pathways suitable for the production of nonnative metabolites particular for host cell factories such as E. coli, S. cerevisiae, and B. subtilis. Owing to the presence of a vast number of candidate genes encoding similar enzymes and problem of low/no heterologous enzyme expression, the selection of suitable heterologous genes particular for the target host become difficult. Therefore, the applicability of the previously designed platform in chapter 2 was extended to select suitable candidate heterologous genes encoding enzymes. CAI score was used to screen the target genes whose introduction to host microorganisms would enable efficient target metabolite production. This ArtPathDesign system was improved to overcome the problems about host-specific heterologous pathways, host-specific heterologous genes, and enables to provide a catalog of nonnative metabolites in the specific host. Besides, the ArtPathDesign system could suggest the new heterologous pathways, for example Isomultiflorenol, vanillin, and so on. Furthermore, optimized nucleotide sequences of heterologous enzymes consisting of only the most preferred codons of hosts were calculated and these optimized heterologous genes may improve the production of metabolites if all optimized gene sequences applied for the corresponding heterologous pathway. It is expected that the *in silico* platform, ArtPathDesign, is proved as a valuable tool by providing essential information for improving cell factories. This aids researchers in developing strategies for strain improvement and facilitates the rational design of metabolic pathways for the production of value-added chemicals by host microorganisms.

Chapter 4

4.1 General conclusion and discussion

Nowadays, the demands of fuels and chemical feedstocks are largely increased, while petroleum resources are limited and unsustainable. In addition, petroleumbased process for energy and industrial chemicals shows negative impacts on environment. The alternative route to produce fuels and valuable chemicals using microorganisms becomes a striking way. However, some microorganisms are not easy to cultivate and produce high level of target products, and lack of metabolic information and genetic manipulation tools.

Metabolic engineering is one of the most widely used techniques to modify and/or integrate heterologous pathways into the well-developed hosts, such as *E. coli, S. cerevisiae*, etc. for producing and/or improving target compounds. It is difficult for scientists to search for the feasible heterologous to produce target metabolites since the following reasons. There are a huge number of metabolic reactions available from databases and literature, and the complexity of host's metabolic network. To search for heterologous pathways connecting to the host's metabolism is tough and time-consuming tasks, and it is difficult to handle by human.

Accordingly, the computational or *in silico* system is required to solve those problems. In previous studies, several pathway design approaches (Cho et al., 2010; Dogrusoz et al., 2009; Finley et al., 2009; Flórez et al., 2011; Handorf et al., 2005; Li et

al., 2004; McShan et al., 2003; Moriya et al., 2010; Pey et al., 2011; Pharkya et al., 2004; Rodrigo et al., 2008; Yousofshahi et al., 2011) for the production of target metabolites have been developed (comparison of among methods are summarized in Table 1.2 in chapter 1). Nevertheless, there is no consensus or general approach to select heterologous pathways and genes especially for target host cells.

Therefore, the aim of this thesis was to develop the *in silico* method, named as the ArtPathDesign (<u>Art</u>ificial heterologous <u>Path</u>way <u>Design</u>), for screening hostspecific heterologous pathways, host-specific heterologous genes and providing a catalog of nonnative metabolites of each host. These key features are important for producing/improving target compound in a specific host cell and still not yet be developed by the earlier reports.

In chapter 2, in order to produce target nonnative metabolites in the wellcharacterized hosts, *E. coli*, *S. cerevisiae*, and *C. glutamicum*, the algorithm to search for heterologous pathways was developed. With this algorithm, all possible nonnative metabolites that are able to connect to the 3 hosts were screened as well as information of heterologous reactions. Still, it was found that numerous orthologous genes encoding enzymes with similar function are available. Thus, a minimum of K_m value was applied with the expectation to have the highest affinity among orthologous genes to select candidate heterologous genes corresponding to heterologous pathway for the production of nonnative metabolites. Combining with the K_m selection score, the identical heterologous reactions of nonnative metabolites agreed well with widely used in metabolic engineering of industrial products were successfully screened (Table 2.2 and 2.3). The examples of those nonnative metabolites are isoprene, α -farnesene, poly- β -hydroxybutyrate (PHB), and cadaverine. Furthermore, by comparing the number of producible nonnative metabolites among host microorganisms, it was found that yeast (*S. cerevisiae*) cell has several compartments presenting barriers to metabolite transport. For instance, mitochondrial/cytoplasmic interfaces prohibit the production of certain target metabolites when sugar is used as a carbon source. In addition, the genome-scale metabolic model used in this thesis does not account for transportation capabilities between compartments, which are currently unclear for many metabolites, and which might influence the production capacities of target metabolites in the real cell systems. By using the screening algorithm and K_m as the selection score, times and costs for searching suitable heterologous pathways agreed well with previous reports were reduced when comparing with the possible heterologous pathways generated without any background information. In addition, this computational method is applicable for any genome-scale metabolic models.

Although K_m value was used as the score for selection of candidate heterologous genes among numerous orthologous genes, the K_m information is limited since it required *in vitro* experiments to observe kinetic of enzymes and activities itself depending on substrate concentration. However, the expression level of a particular heterologous enzyme is crucial for the activation of a specific heterologous pathway to yield a higher productivity of the desired target metabolite.

In chapter 3, the improvement of the *in silico* platform was developed to facilitate the screening of host-specific pathways and genes by applying a new selection score. Firstly, the in-house database was updated from version 1.0 to 2.0 by

using the newest metabolic reaction database called BKM-react (Lang et al., 2011). The alternative score namely Codon Adaptation Index (CAI) was applied to be used as a selection score for finding candidate heterologous genes. The CAI score of a gene is calculated based on the preference codons found in the highly expressed genes of the host. It is expected that a higher CAI score resulting in a high expression level of the enzyme which could increase the production of target metabolite. With the CAI score as the selection score, the alternative heterologous genes were screened that could improve the productivity of nonnative metabolites such as 1, 3-propanediol in S. cerevisiae as the host. The heterologous pathway for (R, R)-2, 3 -butanediol in E. coli was identified and alternative heterologous genes were suggested. This in silico system, named as ArtPathDesign, is successfully identified the new heterologous pathways and genes of useful compounds such as cis, cis muconic acid, vanillin, Isomultiflorenol, etc. and could open the new biotransformation route to produce nonnative metabolites in the target host such as E. coli, S. cerevisiae, B. subtilis, etc. In case of cis, cis muconic acid, Sun and colleagues (Sun et al., 2013) was just reported the novel heterologous pathway which is the identical heterologous pathway suggested by the in silico system developed in this thesis. This example shows the proof of the algorithm successfully identified the new heterologous pathway that can be used to produce useful chemical in *E. coli* cell.

In conclusion, the computational platform, ArtPathDesign, was developed as a metabolic engineering tool in order to provide strategies for producing target metabolites in specific host cell factories, *E. coli*, *S. cerevisiae*, *C. glutamicum*, and *B. subtilis*. The ArtPathDesign platform is applicable for any genome-scale metabolic models. The strategy of current ArtPathDesign is shown in Fig. 4.1. Host's metabolic

networks and metabolic reactions from available databases were parsed into an inhouse database, which will be used as the input data for generating all possible heterologous pathways. To select and rank heterologous genes for those possible heterologous pathways, the Codon Adaptation Index (CAI) was calculated for all orthologous genes retrieved from KEGG and EMBL databases and the optimized gene sequences were created by replacing original codons with favorable codons from the host's highly expressed genes without changing amino acid. Then, the rank of candidate heterologous genes and optimized gene sequences were included into the in-house database. Once all possible heterologous pathways were generated for the host cell, flux balance analysis (FBA) simulation was performed by including constraints that are glucose as a carbon source and mass balance. FBA was applied to observe capacity of the nonnative metabolites after introduction of heterologous pathway to the host metabolism under specific constraints. The heterologous pathway of the nonnative metabolite, which can produce more than 1% molproduct/mol-glucose, was selected and the heterologous genes corresponding to those heterologous pathways were ranked by using CAI score. Finally, list of nonnative metabolites, heterologous pathways, and genes was obtained to provide necessary information to users for conducting trial experiments. However, this platform was not included such factors thermodynamics of heterologous reactions, toxicity of products on the host cell, those features will be developed in the future as expansion features for suggesting more rational heterologous pathways.

The ArtPathDesign platform successfully identified host-specific heterologous pathways, host-specific heterologous genes, and provided the catalog of all nonnative metabolites are able to be produced by each host. Also, the new heterologous pathways and genes were also suggested from this platform which would be a new route to engineer the host strain for target productions. Finally, the optimized gene sequences, which substituted original coding sequences with the preferable codons from highly expressed genes of each host, were also included. As mentioned in chapter 3, the cost for the artificial gene synthesis is gradually decreased. It is possible to improve the productivity of target metabolites by including optimized heterologous genes into the target host cell.

FIG. 4.1 Overall strategy of heterologous pathway identification and selection

(current version of ArtPathDesign system)

4.2 Future perspective

The current version of ArtPathDesign generates all possible heterologous pathways by assuming all known metabolic reactions to be reversible. However, there are a few class of reactions such as carboxylation/decarboxylation reaction, ATP consuming reactions (controlled by kinases), which are thermodynamically irreversible. In order to identify a more rational heterologous pathway, thermodynamics of heterologous reactions should be included to judge and select heterologous pathway for the production of nonnative metabolites. However, to manually incorporate all information on the reaction irreversibility is a hard task and it will be an expansion feature (shown in Fig. 4.2). Thermodynamics of metabolic reactions is calculated by using kinetic parameters retrieving from experiments, which are limitation. Several computational methods have been developed to estimate Gibbs free energy for biochemical reactions(Jankowski et al., 2008; Noor et al., 2012; Rother et al., 2010; Sabatini et al., 2012). One of these methods called group contribution is widely used. This group contribution method is demonstrated to be capable of estimating Gibbs free energy for the majority of the biochemical compounds and reactions found in E. coli's metabolic network (Jankowski et al., 2008). Therefore, the reversibility of heterologous reactions will be calculated using this group contribution method and can be used to select more reasonable heterologous pathways. The thermodynamics feature should further develop for integrating with the current version of ArtPathDesign, and will help to improve the performance for selection of heterologous pathways.

The current version of ArtPathDesign platform was developed based on metabolic reactions data from available databases such as KEGG, BRENDA, BKMreact and so on. However, the input data for ArtPathDesign is depending on metabolic reactions from databases. Generating new metabolic reactions (nonexistent in any databases) is a desire feature of heterologous pathway design. Therefore, to complete the ArtPathDesign system it requires a strategy to find new metabolic reactions, including information of enzymes/proteins and genes. Thus, another expansion feature is to create new heterologous reactions (Fig. 4.2 shown in red dashed box).

It was reported that several enzymes in *E. coli* (about 37% of enzymatic enzymes) that found to be generalist enzymes that promiscuously catalyze reactions on a variety of substrates (Nam et al., 2012). The idea of generalist enzymes leads to the possibility to create the novel metabolic reaction since these enzymes can bind to multi-substrates. Therefore, the design of novel metabolic reactions based on substrate similarity of generalist enzymes will be added to the ArtPathDesign as shown in Fig. 4.2 (red dashed box). Tanimoto or Jaccard coefficient is one of the most popular scores used to measure similarity between chemical structures represented by means of fingerprints (Willett et al., 1998). By using compound similarity score such as Tanimoto coefficient, the substrate-like compound will be screened. The possible new metabolic reaction will suggest from the substrate-like compound used as the alternative substrate in that metabolic reaction. The schematic of novel metabolic reaction deign is shown in Fig. 4.3.

All metabolic enzymes and reactions from several databases such as KEGG, BRENDA, EMBL, ENZYMES, PDB, etc. will be compiled. Next, an enzyme that is able to catalyzed more than one reaction will be classified as generalists. Then, all substrate compounds of those generalist enzymes will be collected. Structures of those compounds can be retrieved from MetaCyc and KEGG databases. Consequently, Tanimoto coefficient will be calculated between the substrates of generalist enzymes and other compounds from available databases. If the similarity scores of such compounds can pass the threshold of the substrate-like compound, the compound will be used as the target for next step. After that, molecular docking technique will be used for observing enzyme-substrate affinity and a binding score will be calculated. Several docking tools (Yuriev and Ramsland, 2013) are available and successfully used for screening drug targets based on free energy binding and inhibition constant. In order to do enzyme-compound docking, target enzyme structures can be downloaded from PDB database. Finally, enzyme-binding scores between the substrate-like compound and target enzyme will be calculate by using tools such as AutoDock (Morris et al., 2009), GOLD (Jones et al., 1997), Glide (Friesner et al., 2004; Halgren et al., 2004), etc. However, the most important parameters are thresholds of substrate-like compounds and enzyme-compound binding scores; it required a computational system to repeat the process until finding a well-represented score which fits to some experimental data. Finally, new heterologous reactions will be identified and will be integrated to the ArtPathDesign system. Thus, to further complete ArtPathDesign system, the thermodynamics of heterologous reactions and the computational methods to design the new heterologous reaction(s) features will integrate to the current version of ArtPathDesign. The complete version of ArtPathDesign would be the useful tool that enables the scientists for improvement of target metabolites in their target hosts by providing alternative metabolic engineering strategies.

FIG. 4.2 The schematic of the complete ArtPathDesign integrated with the expansion features.

Thermodynamics of heterologous reactions will be expanded to improve the current version for selection of heterologous pathways. Red dashed box represents another expansion feature will be developed for creation of new heterologous reactions based on compound similarity and enzyme-substrate binding scores. (An image of protein structure was retrieved from http://www.rcsb.org/pdb/images/2yow bio r 500.jpg)

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FIG. 4.3 The schematic of strategy to design new heterologous reactions

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Appendix A

Source code

A.1 Heterologous pathways detection by Python™

A.1.1 Main script "runningScript.py"

Runing script by using this file with IDLE

Before run the script, you need to change the path of input file that you want to obtain all possible connected metabolites

kegg_metabolic_reaction2010mar.csv -> is the reference reactions from
KEGG ligand database

host_metabolites.txt -> contains all metabolites in terms of KEGG ID if available (you can get this list fromn the supplementary of Genome-scale metabolic model

nonNativeMetabolites.txt -> contains all metabolites (retriving from reference reactions that are not in the host metabolic network)

import sys,pickle,re,os,csv,glob
from heterologousReactionDetection import *
from addInfoOutput import *
from createIndexFile import *

current directory
cur_dir = os.getcwd()

os.path.join(cur_dir,"input/bkm_metabolic_reactions_2012.csv")## KEGG
reaction in tab delimited format

gemMet_file = os.path.join(cur_dir,"input/eco_nativeMetbolites.csv")## host
metabolites if no KEGG ID used the original ID

nonNative_file =

list of metabolites from host metabolic model
gemMetList = [line.upper().rstrip() for line in open(gemMet_file)]

list of nonnative metabolites (found in KEGG reference reactions, but not found in host network) nonNativeMetList = [line.upper().rstrip() for line in open(nonNative_file)]

result = Reaction(bkm_rn_file)## class KEGG metabolic reaction with
reaction ID, left metabolites, right metabolite
bkm_rns ={}
for item in result.getReaction():
 rec =
 [item.rx_id,item.leftMet,item.rightMet,item.equation,item.orgEquation,item.
direction]
 bkm_rns[item.rx_id] = rec

Input the number of iteration that you want to observe numOfiteration = raw_input("Please input total iterations (such as 10,20, or 30) and then press Enter = ")

To write pre-results that contain the possible reaction to be used maxIteration = writePreResult(numOfiteration,nonNativeMetList,bkm rns,gemMetList)

To write all heterolgous reactions (equation in term of KEGG compound ID [Cxxxxx]) for each nonnative metabolites in each iteration writeAllHeterologousPathwayToFile(maxIteration,bkm_rns) print "Finished searching !!!!!! for the current host organism"

To remove preResult file (*.csv) removePreResultFile()

To generate output files in html-format

input_dir = os.path.join(cur_dir,"input")

bkmReactDict =
getBKMreaction(os.path.join(input_dir,"bkm_metabolic_reactions_2012.csv"))
keggDict = getKEGGcompound(os.path.join(input_dir,"kegg_compounds.csv"))
metaCycDict =
getMetaCycCompound(os.path.join(input_dir,"metacyc_compounds.csv"))
ecGenesDict =
getOptGenes(os.path.join(input_dir,"2012_optGenes_ecoHost_bothDB.txt"))

print "Still writing all results in output folder (html-file) !!!!!"

writeResultFiles(maxIteration,bkmReactDict,keggDict,metaCycDict,ecGenesDict)

print "Finished all nonnative metabolites and Heterologous reactions in output folder !!!!!!!"

To create index.html file that summarized all connectable nonnative metabolites

A.1.2 "heterologousReactionDetection.py"

```
#!/usr/bin/env python
# Author: Sunisa Chatsurachai
# Purpose: For finding all additional reactions (Heterologous pathway) for
each non-native metabolite
# Created: 11/26/2010
```

import sys,pickle,re,os,csv,glob from candidateReaction import * from createHeterologousPathwayFile import writeAllHeterologousPathwayToFile


```
class Record:
    """Blank class"""
    pass
```



```
class Reaction():
    .....
```

```
def __init__(self,reaction_file):
      reaction file = "finalBKM ir.csv"
      EX:
      R00004 2.3.1.48BS407080acetyl-CoA + c-Myc <=> CoA + acetylated c-
      1 C00024 + 1 c-Myc <=> 1 C00010 + 1 acetylated c-Myc brenda r
Myc
       .....
      self.reaction_file = open(reaction_file)
      self.data =[]
      self.process()
   def getReaction(self):
       ....
      get each reaction from file
       ......
      return self.data
   def checkGlycanReaction(self,list equation item,default = "No glycan"):
      for item in list equation item:
          if item.startswith("G"):
               print "found Glvcan"
##
```

```
default = None
       break
return default
```

```
def checkMetabolite(self,item):
```

Check if it is metabolite or not if not return None

```
if item in ["<=>","<-->","<=","<--","=>","-->","+"]:
    return None
```

```
elif re.findall(r'[a-z]|[A-Z]',item) == []:## found only numeric
   return None
elif item != "":
   return None
```

else: return item

def getMetabolitesFromEquation(self,equation):

```
input equation
return [leftmets,rightmets]
.....
```

```
result = []
        pattern = re.search(r'(\<\=\>)|(\=\>)|(\<\-\->)|(\-\-
\>)',equation)
```

```
if pattern is not None:
    #print pattern.group()
    data = equation.split(pattern.group())
    leftMet = data[0]
    rightMet = data[1]
    #print leftMet,rightMet
```

result.append(leftMet) result.append(rightMet)

else:

print "not found direction" return result

def getLeftMetabolite(self,leftMets):

```
metItem = leftMets.lstrip().rstrip().split(" + ")
       for m in metItem:
            m item = m.split()
            if self.checkMetabolite(m item[0]) is None and
re.findall(r'^[1-9]$|^[1-9][0-9]$|^[1-9][0-9][0-9]$',m_item[0]) != []:
               #print m_item[0], "##", " ".join(m_item[1:])
                stoi = m item[0]
               met = " ".join(m item[1:])
            else:
               if re.findall(r'^n$|^m$|^\(n\+[1-9]\)$|^\(n\+m\)|^\(m\+[1-
9]\)$',m item[0]) !=[]:
                    #print m_item[0], "##"," ".join(m_item[1:])
                    stoi = '1'
                    met = " ".join(m item[1:])
                elif re.findall(r'^[2-9]n$|^[2-9]m$|^[1-9][0-9]n$|^[1-9][0-
9]m$',m item[0]) != []:
                    stoiData = m item[0]
                    if len(stoiData) >= 2:
                        stoi = stoiData[:-1]
                        met = " ".join(m item[1:])
                else:
                    stoi = '1'
                   met = " ".join(m_item[:])
            #print stoi,met
            result.append(met)
       return result
   def getRigthMetabolite(self,rightMets):
       result = []
       metItem = rightMets.lstrip().rstrip().split(" + ")
       for m in metItem:
            m item = m.split()
            if self.checkMetabolite(m item[0]) is None and
re.findall(r'^[1-9]$|^[1-9][0-9]$|^[1-9][0-9][0-9]$',m_item[0]) != []:
               #print m_item[0],"##"," ".join(m_item[1:])
               stoi = m item[0]
                met = " ".join(m item[1:])
            else:
```

result = []



if re.findall(r'^n\$|^m\$|^\(n\+[1-9]\)\$|^\(n\+m\)|^\(m\+[1-

elif re.findall(r'^[2-9]n\$|^[2-9]m\$|^[1-9][0-9]n\$|^[1-9][0-

#print m_item[0], "##"," ".join(m_item[1:])

9]\)\$',m item[0]) !=[]:

9]m\$',m item[0]) != []:

else:

return result

result.append(met)

stoi = '1'

stoi = '1'

met = " ".join(m_item[1:])

stoiData = m_item[0]

if len(stoiData) >= 2:

stoi = stoiData[:-1]

met = " ".join(m_item[:])

met = " ".join(m item[1:])

rRecord.direction = "<=>"

self.data.append(rRecord)

def writePreResult(maxIt,nonNativeMetList,kegg rns,gemMetList): cur dir = os.getcwd() for it in range(1, int(maxIt)+1): if it ==1: ## 1st searching result = getSingleAddedRn(nonNativeMetList,kegg_rns,gemMetList) set_r1 =[]## keep reaction used to connect to host GEM set c1 =[]## keep non native compound found connecting f = open(os.path.join(cur dir, "preResult iteration "+str(it)+'.csv'), 'w') for cpd, list rn, min rn in result: set r1.extend(list rn) set c1.append(cpd) write result = createListOfAddedRns(cpd,list rn,it) for rec in write result: f.write("\t".join(map(str,rec))) f.write("\n") f.close() pre Rn = set r1 pre Cpd = set(set c1) ## remove compound duplicate (for next

iteration)

```
nextNonNativeMetList =
list(set(nonNativeMetList).difference(pre_Cpd))
            print "Iteration = ",it,"\t","No. of connected metabolites =
",len(pre Cpd), "\t", "No. of nonnative metabolties for next step =
", "\t", len(nextNonNativeMetList)
```

elif it==2: second result = getAddedRns(pre_Rn,list(pre_Cpd),gemMetList,nextNonNativeMetList,kegg_rns,i t)

set r2 = [] set c2 = [] f = open(os.path.join(cur_dir, "preResult_iteration_"+str(it)+'.csv'), 'w') for cpd,list rn,min rn in second result: set_r2.extend(list_rn) set c2.append(cpd)

write_result = createListOfAddedRns(cpd,list_rn,it) for rec in write result: f.write("\t".join(map(str,rec))) f.write("\n") f.close() set_c2 = set(set c2)

```
pre Cpd = set(set c2)
            pre Rn = set r2
            nextNonNativeMetList =
list(set(nextNonNativeMetList).difference(pre Cpd))
```

print "Iteration = ",it,"\t","No. of connected metabolites = ",len(pre Cpd),"\t","No. of nonnative metabolties for next step = ","\t",len(nextNonNativeMetList)

```
else:
```

next result = getAddedRns(pre Rn,list(pre Cpd),gemMetList,nextNonNativeMetList,kegg rns,i t) pre Rn =[] <pre_Cpd =[]</pre> f =

open(os.path.join(cur_dir,"preResult_iteration_"+str(it)+'.csv'),'w') for cpd,list rn,min rn in next result: pre_Rn.extend(list_rn) pre Cpd.append(cpd) write result = createListOfAddedRns(cpd,list rn,it) for rec in write result: f.write("\t".join(map(str,rec))) f.write("\n") f.close()

pre_Cpd = set(pre_Cpd) nextNonNativeMetList = list(set(nextNonNativeMetList).difference(pre_Cpd)) print "Iteration = ",it,"\t","No. of connected metabolites = ",len(pre_Cpd),"\t","No. of nonnative metabolties for next step = ","\t",len(nextNonNativeMetList) if len(pre Cpd) ==0: break return it-1 ## return maximum iteration that found connecting nonnative

metabolite

def removePreResultFile():

import glob cur_dir = os.getcwd() preResultFiles = glob.glob(os.path.join(cur_dir,"*.csv")) for f in preResultFiles: os.remove(f)

def main():

pass

if __name__=='__main__':main()

A.1.3 "candidateReaction.py"

.....

#!/usr/bin/env python # Author: Sunisa Chatsurachai # Purpose: functions for finding candidate reaction(s) # Created: 11/26/2010 import os,re,sys from glob import glob def allPrestepsMapping(preStepPath,can rxn metList): ## preStepPath = "D:\\SUNISA data\\2012 additional reaction\\2012JUL bkmReact\\2012JUL addRe action\\assume_r\\eco\\" metNotFound = [] connectMets = [] metFound = [] for met in can rxn metList: list csv = glob("*.csv")## list of preResult File for f in list csv: for line in open(f, 'r'): if line.startswith(met) == True: metFound.append(met) break metNotFound = set(can rxn metList).difference(set(metFound)) return [list(metNotFound),metFound] def hostMetaboliteMaping(gemMetList, can rn metList): Comparing metabolite from candidate reaction with genome-scale model(GEM) metabolite list and return list of metabolite does not found in GEM list met not found =[] for item in can rn metList: if item not in gemMetList: met_not_found.append(item) return list(set(met not found)) def findAllPossibleCandidateRn(kegg rns,nonNativeMet,gemMetList):

 \geq

kegg rns (all metabolic reaction from KEGG (no glycan reaction here) Type is Dictionary each record as key -> item.rx id value list-> [item.rx id.item.leftMet.item.rightMet.item.equation.item.equation name.ite m.direction] pre_candidate_rn = [] for key,value in kegg rns.iteritems(): rx id = value[0] leftMet = value[1] rightMet = value[2] direction = value[-1] if direction == "<=>": ## found reversible reaction if nonNativeMet in leftMet: rightMet = hostMetaboliteMaping(gemMetList,rightMet)## compare and return met not found in GEM pre candidate rn.append([rx id,rightMet]) elif nonNativeMet in rightMet: leftMet = hostMetaboliteMaping(gemMetList,leftMet) pre candidate rn.append([rx id,leftMet]) else:## found irreversible reaction if nonNativeMet in rightMet: leftMet = hostMetaboliteMaping(gemMetList,leftMet) pre_candidate_rn.append([rx_id,leftMet]) result = $\{\}$ for k,m in pre candidate rn: result[k] = m return result def getSingleAddedRn(nonNativeMetList,kegg_rns,gemMetList):

At 1st iteration -> finding non native metabolite which can connect to
host by adding single reaction
"""
result ={}
final_result = []
for nonNativeMetList:

eachNonNativeMet = findAllPossibleCandidateRn(kegg_rns,nonNativeMet,gemMetList) if [] in eachNonNativeMet.itervalues():## [] found in possible pathway of connected metabolite candidate_rn = [] for key,val in eachNonNativeMet.iteritems(): if val == []: ##all substrate found in GEM list candidate_rn.append(key) ## append reaction id result[nonNativeMet] = candidate_rn for k,v in result.iteritems(): final_result.append([k,v,len(v)]) ## create list of all possible reaction return final_result

def

getAddedRns(preStepRn,preStepMet,gemMetList,nonNativeMetList,kegg_rns,itera tion):

```
....
```

From 2nd iteration -> finding at least two reactions to be added to host for connecting non-native metabolite

3rd iteration -> finding at least three reactions to be added to host for connecting non-native metabolite

```
....
```

next_step_result = []

for nonNativeMet in nonNativeMetList:
 pre_rn =
findAllPossibleCandidateRn(kegg_rns,nonNativeMet,gemMetList)
 pre_result = {}
 possible rns = []

```
for rx_id,met_not_found in pre_rn.iteritems():
    min_rn_found = 0+iteration ## at least start from iteration
number ( to connect to host metabolite)
```

min_rn_found = min_rn_found

```
preMets =[]
                                                                                                            for cMet in connectedMets:
possible rns.append([rx id,met not found[0],min rn found])
                else:
                                                                                                                cur cMet = cMet.split(" ")[0]
                                                                                                                listReaction = re.findall("R[0-9][0-9][0-9][0-
metNotFoundInAllPresteps,connectedMets=allPrestepsMapping(os.getcwd(),met n
                                                                                    9][0-9]", cMet)
ot found)
                                                                                                                preMets.append(cur cMet)
                    if metNotFoundInAllPresteps == []:## all metabolite
found in presteps
                                                                                                            for m in met found in preStep:
                        min_rn_found = min_rn_found+len(met_not_found)
                                                                                                                preMets.append(m)
                        preMets =[]
                        for cMet in connectedMets:
                                                                                                            preMets = list(set(preMets))
                            cur_cMet = cMet.split("_")[0]
                                                                                                            possible rns.append([rx id,preMets,min rn found])
                            listReaction = re.findall("R[0-9][0-9][0-9][0-
9][0-9]", cMet)
                            preMets.append(cur cMet)
                                                                                            pre result[nonNativeMet] = possible rns
                        preMets = list(set(preMets))
                                                                                            if pre result[nonNativeMet] != []:
possible rns.append([rx id,preMets[0],min rn found])
                                                                                    next step result.append([nonNativeMet,possible rns,len(possible rns)])
            else:## more than one metabolite does not found in GEM list
               met found in preStep = []
                                                                                        return next step result
                for i,m in enumerate(met not found):
                   if m in preStepMet:
                                                                                    def createListOfAddedRns(cpd id,ListRns,iteration):
                        met found in preStep.append(m)
                                                                                        rec = []
                                                                                        if iteration ==1: ## 1st iteration result
                cur metNotFound =
                                                                                            if len(ListRns) == 1:
set(met not found).difference(set(met found in preStep))
                                                                                                rn = ListRns[0]
                cur_metNotFound = list(cur_metNotFound)
                                                                                                rec.append([cpd_id,rn,iteration])
                                                                                            else:
               if cur metNotFound == []:## all metabolites already found
                                                                                                for rn in ListRns:
in prestep metabolite
                                                                                                    rec.append([cpd id,rn,iteration])
                    min_rn_found = min_rn_found +len(met_not_found)## at
                                                                                            return rec
least reaction should be added
                                                                                        else:
                                                                                            ## ListRns=[['R01424', 'C00180']]
possible rns.append([rx id,met found in preStep,min rn found])
                                                                                            if len(ListRns) == 1:
                                                                                                rn = ListRns[0][0] ## rx_id
                else:
                                                                                                if rn.find("##") != -1:
                                                                                                    rn = rns.split('##')
                                                                                                else:
metNotFoundInAllPresteps,connectedMets=allPrestepsMapping(os.getcwd(),cur m
                                                                                                    rn
etNotFound)
                                                                                                cpd pre = ListRns[0][1] ## compound id
                                                                                                min_rn_found = ListRns[0][2]
                    if metNotFoundInAllPresteps == []:## all metabolite
found in presteps
                                                                                                if type(cpd_pre) != list:## only one previous step metabolite
                        min rn found = min rn found+len(met not found)
                                                                                    (in iteration -1)
```

 \geq

rec.append([cpd_id,rn,iteration,cpd_pre,min_rn_found])
else:
 rec.append([cpd_id,rn,iteration,cpd_pre,min_rn_found])

```
else:
        ## [['R07922', 'C16353'], ['R07921', 'C07481'], ['R07954',
'C07481'], ['R07939', 'C07481'], ['R07943', 'C16358']]
        ## [['R03186', ['C00072', 'C05422']], ['R00095', 'C00072']]
           for r in ListRns:
               rn = r[0]
               if rn.find('##') != -1:
                   rn = rn.split('##')
               else:
                   rn
               cpd pre = r[1]
               min_rn_found = r[-1]
               if type(cpd pre) != list:
                   rec.append([cpd id,rn,iteration,cpd pre,min rn found])
               else:
                   rec.append([cpd id,rn,iteration,cpd pre,min rn found])
       return rec
```

A.1.4 "createHeterologousPathwayFile.py"

#!/usr/bin/env python # Author: Sunisa Chatsurachai # Purpose: Create additional pathway detection for simulation (txt-file) # Created: 11/29/2010 import sys,os,csv,re from glob import glob def addCompartmentToMet(reaction, compartment="[c]"): result = [] reaction = reaction.upper().split() for item in reaction: if item not in ["<=>","<=","=>","+"] and re.findall(r'[0-9]',item[0]) ==[]: ## only metabolite item = item+compartment result.append(item) else: result.append(item) return result

```
def addTargetMetOutputRn(nonNativeMet):
    met_ext = nonNativeMet.upper()+"[e]"
    sink_met_rn = " ".join([met_ext,'=>'])
    return sink_met_rn
```

```
def createTransportRn(nonNativeMet):
    met_cytosol = nonNativeMet.upper()+"[c]"
    met_extracellular = nonNativeMet.upper()+"[e]"
    transport_rn = " ".join([met_cytosol,"=>",met_extracellular])
    return transport_rn
def getPreviousStepRn(preMet,pre_it):
    """
```

get all reactions of previous metabolite
"""
cur_dir = os.getcwd()

for iteration in range(pre_it):

```
targetDir = os.path.join(cur_dir,"iteration_"+str(iteration+1))
```

```
fileList = glob(targetDir+"/*.txt")
results = []
for fName in fileList:
    file_name = fName.split("\\")[-1]
    met_name = file_name.split("_")[0]
    if met_name == preMet:
        rns = [line.rstrip() for line in open(fName)]
        previous_rns_name = rns[0]
        list_rns = rns[1:-2]
        results.append([previous_rns_name,list_rns])
if results != []:
        break
return results
```

```
def getAllPossiblePreviousStepRn(preMetList,cur it):
   get all reaction of all previous metabolites(as list)
   preMetList rns ={}
   for preMet in preMetList:
       rns = []
       pre_rn_set = getPreviousStepRn(preMet,cur_it)
       if pre rn set != []:
            if len(pre_rn_set) == 1:## one possible set of reaction
               pre cid rid = pre rn set[0][0]
               pre list rn = pre rn set[0][1][:]
               rns.append([pre cid rid,pre list rn])
            else:
               for pre cid rid, pre list rn in pre rn set:
                   rns.append([pre cid rid,pre list rn])
       preMetList rns[preMet] =rns
   return preMetList rns
def changeFormatPreviousListMet(preMet):
    ## nonNativeMet addedRn itearation previousMets minimum rn
                                ['C02336', 'C00668']
   ##C16688 R03921 2
                                                        4##
```

```
##C16688 R03921 2 ['C02336', 'C00668']
preMetList = ""
for char in preMet:
    if char not in ["[","]","'"," "]:
        preMetList = preMetList +char
preMetList = sorted(preMetList.split(","))
```

return preMetList

```
def writeResultToTextFile(targetDirectory,iteration,List result):
    ## List result =
[cur rn name, pre rn name, cur rn, pre rn, cur tran rn, sink met rn, path no]
   if iteration > 2:
        f =
open(targetDirectory+str(iteration)+"/"+List result[0]+" "+str(List result[
-1])+".txt","w")
       f.write(List_result[0]+"_"+List_result[1])
        f.write("\n")
        if len(List result[3]) == 1: # only one heterologous pathway
            f.write(List result[3][0])
            f.write("\n")
        else:
            rns = List_result[3]
            for r in rns:
                f.write(r)
                f.write("\n")
        f.write("\n".join([List_result[2],List_result[4],List_result[5]]))
        f.write("\n")
   else:
       f =
open(targetDirectory+str(iteration)+"/"+List result[0]+" "+List result[1]+"
.txt","w")
        f.write(List result[0]+" "+List result[1])
        f.write("\n")
        if len(List result[3]) == 1: # only one heterologous pathway
            f.write(List result[3][0])
            f.write("\n")
```

```
else:
    rns = List_result[3]
    for r in rns:
        f.write(r)
        f.write("\n")
f.write("\n".join([List_result[2],List_result[4],List_result[5]]))
f.write("\n")
```

def writeAllHeterologousPathwayToFile(maxIt,kegg_rns):

cur_dir = os.getcwd()
for iteration in range(1,maxIt+2):

if not

if iteration ==1:

f =
open(os.path.join(cur_dir,"preResult_iteration_"+str(iteration)+".csv"),"r"
)

```
for nonNativeMet,r_id,it in csv.reader(f,delimiter= "\t"):
    if kegg_rns.has_key(r_id):
        eq_with_compartment = "
".join(addCompartmentToMet(kegg_rns[r_id][3],"[c]"))
        add_rn = [nonNativeMet,r_id,eq_with_compartment]
        try:
        f1 =
    open(cur_dir+"/iteration_1/"+nonNativeMet+"_"+r_id+".txt",'w')
        fName = nonNativeMet+"_"+r_id
```

except IOError:

f1 =
open(cur_dir+"/iteration_1/"+nonNativeMet+"_"+r_id+".txt",'w')
fName = nonNativeMet+" "+r id

if iteration > 1 and iteration <= maxIt+2:</pre>

f =
open(os.path.join(cur_dir, "preResult_iteration_"+str(iteration)+".csv"), "r"
)

for nonNativeMet,r_id,it,preMet,min_rn in
csv.reader(f,delimiter="\t"):
 path_no = 0
 if int(min_rn) == iteration:## means -> connecting by one
to one metabolite since other found in GEM List
 cur_rn_with_com = "
".join(addCompartmentToMet(kegg_rns[r_id][3],"[c]"))
 cur_rn_name = nonNativeMet+"_"+r_id
 cur_trans_rn = createTransportRn(nonNativeMet)
 sink_met_rn = addTargetMetOutputRn(nonNativeMet)
 #pre_rn_set =
getPreviousStepRn(preMet,cur_dir+"/iteration_"+str(iteration-1))
 pre_rn_set = getPreviousStepRn(preMet,iteration-1)

if len(pre_rn_set)==1: ## only one possible added

pathway

path_no = path_no+1
pre_cid_rid = pre_rn_set[0][0]
pre_list_rn = pre_rn_set[0][1][:]

List_result =[cur_rn_name,pre_cid_rid,cur_rn_with_com,pre_list_rn,cur_trans_rn,sink_met _rn,path_no]

writeResultToTextFile(cur_dir+"/iteration_",iteration,List_result)

else:

_rn,path_no]

writeResultToTextFile(cur_dir+"/iteration_",iteration,List_result)

else:

```
pre_metList = changeFormatPreviousListMet(preMet)
                                                                                   if __name__=='__main__':main()
                   pre metListRn =
getAllPossiblePreviousStepRn(pre_metList,iteration-1)
                   first_substrate_rn = pre_metListRn[pre_metList[0]]
                   del pre_metListRn[pre_metList[0]] ## update list of
previous step metabolite
                   for p_rn in first_substrate_rn:
                        pre_cid_rid = p_rn[0]
                        pre_list_rn = p_rn[1][:]
                        for key,value in pre_metListRn.iteritems():
                           if len(value)== 1: ##only one possible pathway
                                ##key = C02557
                                ##value = [['C02557 R05373', ['C02557[c]]
<=> C00100[c] + C00011[c]']]]
                               path_no = path_no+1
                                new_pre_list_rn = []
                                v cid rid = value[0][0]
                                v_list_rn = value[0][1]
                                new pre list rn.extend(pre list rn)
                                new_pre_list_rn.extend(v_list_rn)
                                List_result
=[cur_rn_name,pre_cid_rid+"_"+v_cid_rid,cur_rn_with_com,new_pre_list_rn,cur
_trans_rn,sink_met_rn,path_no]
writeResultToTextFile(cur_dir+"/iteration_",iteration,List_result)
                           else:
                                for v_cid_rid,v_list_rn in value:
                                   new pre list rn = []
                                   path no = path no+1
                                   new_pre_list_rn.extend(pre_list_rn)
                                   new_pre_list_rn.extend(v_list_rn)
                                   List result
=[cur_rn_name,pre_cid_rid+" "+v_cid_rid,cur_rn_with_com,new_pre_list_rn,cur
trans rn,sink met rn,path no]
writeResultToTextFile(cur dir+"/iteration ",iteration,List result)
def main():
   pass
```

Appendix A

A.1.5 "addInfoOutput.py"

#!/usr/bin/env python
Author: Sunisa Chatsurachai --<>
Purpose: To add more information of the output files
Created: 27-Feb-13

import sys,os,re
import glob
from heterologousReactionDetection import *

```
def getKEGGcompound(fName):
    result = {}
    for line in open(fName,'r'):
        line = line.lstrip().rstrip().split('\t')
        cpd_id = line[0]
        cpd_syn = line[1]
        cpd_common = line[2]
        cpd_charge = line[3]
        cpd_formula = line[4]
        result[cpd_id] = [cpd_syn,cpd_formula]
    return result
```

```
def getMetaCycCompound(fName):
    result = {}
    for line in open(fName,'r'):
        line = line.lstrip().rstrip().split('\t')
        m_id = line[0]
        k_id = line[1]
        cpd_syn = "##".join([line[2],line[3]])
        cpd_charge = line[5]
        cpd_formula = line[6]
        result[m_id] = [cpd_syn,cpd_formula,k_id]
    return result
```

```
def getBKMreaction(fName):
    result = {}
    for line in open(fName,'r'):
        line = line.lstrip().rstrip().split('\t')
        r_id = line[0]
        ec_num = line[1]
```

refID = line[2] eq_name = line[3] eq = line[4] type_rxn = line[5] result[r_id] =[eq_name,eq,ec_num,refID,type_rxn] return result

```
def getReactions(listReactions,bkmDict):
    rxns = listReactions.split('##')
    result = {}
    for r in rxns:
        eqName,eq,ec_id,refID,revData = bkmDict[r]
        result[r] = eqName
    return result
```

def getOptGenes(fName):
 result = {}
 for line in open(fName,'r'):
 line = line.lstrip().rstrip().split('\t')
 ec = line[0]
 #db = line[1]
 #genes = line[2]
 #orfList = line[3]
 #org_id = line[4]
 #org_name = line[5]
 result[ec] = line[1:6]
 return result

```
def getCompoundName(target,keggDict,MetacycDict):
    if keggDict.has_key(target):
        cpd_syn,cpd_formula = keggDict[target]
    elif MetacycDict.has_key(target):
        cpd_syn,cpd_formula,k_id = MetacycDict[target]
    else:
        cpd_syn = "None"
        cpd_formula = "None"
    return [prepareCompoundName(cpd_syn),cpd_formula]
```

def prepareCompoundName(cpdName):

cpdName = cpdName.split("##")
cpdName = set(cpdName)
cpdName = list(cpdName)

if len(cpdName)==1: met = cpdName[0] if met.upper() == "NONE" or met.upper() == "NULL": return "No information" else: return met else: data = [] for m in cpdName: if m.upper() != "NONE" and m.upper()!="NULL": data.append(m) return "||".join(data)

def

writeResultFiles(maxIteration,bkmReactDict,keggDict,metaCycDict,ecGenesDict):

cur_dir = os.getcwd()

- if os.path.exists(os.path.join(cur_dir,"output")) == False:
 os.mkdir(os.path.join(cur_dir,"output"))
- for it in range(1,maxIteration+1+1): curRentPath = os.path.join(os.getcwd(),"iteration_"+str(it)) listTextfiles = glob(curRentPath+"/*.txt")

```
outputDir = os.path.join(cur_dir,"output")
```

```
if os.path.exists(outputDir)== False:
    os.mkdir(outputDir)
```

for f in listTextfiles: f_op = open(f,'r') line = f_op.readline() f_op.close()

```
foutName = f.split("\\")[-1][:-4]
```

```
connMet= line.lstrip().rstrip().split("_")
targetName,targetFormula =
getCompoundName(connMet[0],keggDict,metaCycDict)
```

```
outputFile = os.path.join(outputDir,foutName+".html")
```

f_out = open(outputFile,'w')
f_out.write('<html>\n')
f_out.write('<body>\n')
f_out.write('\n")
f_out.write('<h1> Result </h1>\n')
f_out.write('\n')
f_out.write('Target compound ID \n')
f_out.write('Target compound ID \n')
f_out.write('Target compound ID \n')

f_out.write('\n')
f_out.write(' Target compound Name \n')
f_out.write(''+targetName+'\n')
f_out.write('

listRxns = re.findall(r'R[0-9][0-9][0-9][0-9][0-9]',line)

f_out.write('\n')
f_out.write(' List of Heterologous reaction(s) \n')
f_out.write(''+"||".join(listRxns)+'\n')
f_out.write('

for r_id in listRxns:
 f_out.write('\n')
 f_out.write(' Detail of heterologous reaction

r_data = bkmReactDict[r_id] eq_name = r_data[0] eq_id = r_data[1] ec_id = r_data[2]

f_out.write(''+r_id+'
'+eq_name+'
'+eq_id+'
')

listEC = ec_id.split("##")
if len(listEC) ==1: ## found only one EC number

try:

e = listEC[0] data_hetGenes = ecGenesDict[e] db = data_hetGenes[0] geneSym = data_hetGenes[1] orfList = data_hetGenes[2] org id = data_hetGenes[3]

except KeyError,err:

	+_out.write("
	'+' '+'Enzyme information:')
	<pre>f_out.write(' '+listEC[0]+' '+'No information</pre>
about genes')	<pre>f out.write('\n')</pre>

else:

try:

```
data_hetGenes = ecGenesDict[e]
db = data_hetGenes[0]
geneSym = data_hetGenes[1]
orfList = data_hetGenes[2]
org_id = data_hetGenes[3]
org_name = data_hetGenes[4]
f_out.write('<br>'+e+'<br>'+'Gene symbol:
'+geneSym.replace("##",";")+'<br>'+'ORF: "+orfList.replace("##","||")+'
from database '+db+'<br>'+'Organism: '+org_name+'<br>')
```

except KeyError,err:

f_out.write('
'+e+'
'+'No information

about genes'+'
')

f_out.write('_____')

f_out.write('\n')

f_out.write('\n')

f_out.write('<footer>')
f_out.write('Created by: Sunisa Chatsurachai')
f_out.write('Date: 2012-02-28')
f_out.write('</footer>\n')
f_out.write('\n')
f_out.write('</body>\n')
f_out.write('</html>\n')
f out.close()

def main():

cur_dir = os.getcwd()
input_dir = os.path.join(cur_dir,"input")

bkmReactDict =
getBKMreaction(os.path.join(input_dir, "bkm_metabolic_reactions_2012.csv"))
keggDict =
getKEGGcompound(os.path.join(input_dir, "kegg_compounds.csv"))
metaCycDict =
getMetaCycCompound(os.path.join(input_dir, "metacyc_compounds.csv"))
ecGenesDict =
getOptGenes(os.path.join(input_dir, "2012_optGenes_ecoHost_bothDB.txt"))

maxIteration = 3

writeResultFiles(maxIteration,bkmReactDict,keggDict,metaCycDict,ecGenesDict)

if __name__=='__main__':main()

A.1.6 "createIndexFile.py"

#!/usr/bin/env python
Author: Sunisa Chatsurahcai --<>
Purpose: To create index.html for link result of nonnative metabolites
Created: 28-Feb-13

import sys,os,re

```
def getKEGGcompound(fName):
    result = {}
    for line in open(fName,'r'):
        line = line.lstrip().rstrip().split('\t')
        cpd_id = line[0]
        cpd_syn = line[1]
        cpd_common = line[2]
        cpd_charge = line[3]
        cpd_formula = line[4]
        result[cpd_id] = [cpd_syn,cpd_formula]
    return result
```

```
def getMetaCycCompound(fName):
    result = {}
    for line in open(fName,'r'):
        line = line.lstrip().rstrip().split('\t')
        m_id = line[0]
        k_id = line[1]
        cpd_syn = "##".join([line[2],line[3]])
        cpd_charge = line[5]
        cpd_formula = line[6]
        result[m_id] = [cpd_syn,cpd_formula,k_id]
    return result
```

```
def getCompoundName(target,keggDict,MetacycDict):
    if keggDict.has_key(target):
        cpd_syn,cpd_formula = keggDict[target]
    elif MetacycDict.has_key(target):
        cpd_syn,cpd_formula,k_id = MetacycDict[target]
    else:
        cpd_syn = "None"
        cpd_formula = "None"
    return [prepareCompoundName(cpd_syn),cpd_formula]
```

```
def prepareCompoundName(cpdName):
   cpdName = cpdName.split("##")
   cpdName = set(cpdName)
   cpdName = list(cpdName)
   if len(cpdName)==1:
        met = cpdName[0]
        if met.upper() == "NONE" or met.upper() == "NULL":
            return "No information"
        else:
            return met
   else:
        data = []
        for m in cpdName:
            if m.upper() != "NONE" and m.upper()!="NULL":
                data.append(m)
        return "||".join(data)
```

```
def getAllConnectedMetaboites(targetDir):
    import glob
    allHtmls = glob.glob(os.path.join(targetDir+"/*.html"))
    allTargets = []
    for f in allHtmls:
        target = f.split("\\")[-1]
        target = target.split("__")[0]
```

```
allTargets.append(target)
return list(set(sorted(allTargets)))
```

def getHtmlLinks(cpdId,targetDir):### limited to 5 possible way only ###
 import glob,random
 htmls = glob.glob(os.path.join(targetDir,"%s*.html"%cpdId))

```
if len(htmls) < 5:
    targetFiles = htmls[:]
else:
    targetFiles = []
    targetFiles.append(random.choice(htmls))
    targetFiles.append(random.choice(htmls))
    targetFiles.append(random.choice(htmls))
    targetFiles.append(random.choice(htmls))
    targetFiles.append(random.choice(htmls))
    targetFiles.append(random.choice(htmls))
```

def writeIndexFile(output_dir,keggDict,metaCycDict):
 import os
 cur_dir = os.getcwd()
 allNonMets = getAllConnectedMetaboites(output_dir)
 indexFile = os.path.join(cur_dir,"index.html")
 fopen = open(indexFile,'w')
 fopen.write('<html>\n')
 fopen.write('<h1>'+'Index'+'</h1>\n')

fopen.write('<body>\n')
fopen.write('\n")
fopen.write('\n')
fopen.write('Nonnative metabolite ')
fopen.write('Heterologous pathway(s)')
fopen.write('

for m in allNonMets[:]:

targetName,targetFormula = getCompoundName(m,keggDict,metaCycDict)
htmls = getHtmlLinks(m,output_dir)

fopen.write('\n')
fopen.write(''+targetName+"||"+m+'')

for i,htm in enumerate(htmls):

```
targetFile = htm.split("\\")[-1]### name of html file
fopen.write('<d>')
fopen.write('<a
href='+"output\\"+targetFile+'>'+"Pathway_"+str(i+1)+'</a>'+'<br>')
fopen.write('')
#<a href="filename.html">text that responds to link</a>
#fopen.write('')
fopen.write('')
```

fopen.write('<footer>')
fopen.write('Created by: Sunisa Chatsurachai')
fopen.write('Date generated: 2012-02-28')
fopen.write('</footer>\n')

fopen.write("\n")
fopen.write('</body>\n')
fopen.write('</html>\n')
fopen.close()

def main():

```
cur_dir = os.getcwd()
input_dir = os.path.join(cur_dir,"input")
keggDict =
getKEGGcompound(os.path.join(input_dir,"kegg_compounds.csv"))
metaCycDict =
getMetaCycCompound(os.path.join(input_dir,"metacyc_compounds.csv"))
```

output_dir = os.path.join(cur_dir,"output")
writeIndexFile(output_dir,keggDict,metaCycDict)

if __name__=='__main__':main()

A.2 Codon Adaptation Index by Python[™] and Biopython

A.2.1 "ecoHost_cai.py"

```
# Author: SUNISA Chatsurachai --<>
# Purpose: Calculate CAI when E. coli as the host
# Created: 18-Nov-11
```

import sys,os,re

```
from Bio.Seq import Seq
from Bio.SegRecord import SegRecord
from Bio.SeqIO import *
from Bio.SeqUtils.CodonUsage import *
from Bio.SeqUtils.CodonUsageIndices import *
from glob import glob
aminoDict = { 'ala':['GCA', 'GCC', 'GCG', 'GCT'],
             'arg':['AGA','AGG','CGA','CGC','CGG','CGT'],
             'asn':['AAC','AAT'],
             'asp':['GAC','GAT'],
             'cys':['TGC','TGT'],
             'gln':['CAA','CAG'],
             'glu':['GAA','GAG'],
             'gly':['GGA','GGC','GGG','GGT'],
             'his':['CAC','CAT'],
             'ile': 'ATA', 'ATC', 'ATT'],
             'lys':['AAA','AAG'],
             'met':['ATG'],
             'phe':['TTC','TTT'],
             'pro':['CCA','CCC','CCG','CCT'],
             'ser':['AGC', 'AGT', 'TCA', 'TCC', 'TCG', 'TCT'],
             'stop':['TAA','TAG','TGA'],
             'thr':['ACA','ACC','ACG','ACT'],
             'trp':['TGG'],
             'tyr':['TAC','TAT'],
             'val':['GTA','GTC','GTG','GTT'],
             'leu':['CTA','CTC','CTG','CTT','TTA','TTG']}
```

def findKOId(orf_desc):
 orf_desc = orf_desc.split()

```
def findGenes(orf_desc):
    orf_desc = orf_desc.split(";")[0]
    if orf_desc.find(",") != -1:#found gene information
        print orf_desc
```

else:

```
pass
def printListRecordsToFile(fName, records, delimiter='\t'):
    f = open(fName,'a')
    f.write(delimiter.join(map(str,records)))
    f.write('\n')
    f.close()
```

def main():

```
cur_dir = os.getcwd()
```

```
heg_ecoli = CodonAdaptationIndex()
heg_ecoli.generate_index(os.path.join(cur_dir, 'heg_ecoli_seq.fasta'))
## w value in Sharp's Method
##heg_ecoli.print_index()
#for item,val in heg_ecoli.index.iteritems():
    #print item,val
#sys.exit()
```

```
kegg_seq_dir =
```

```
"D:\\SUNISA_data\\2012_CAI_calculation\\KEGG_cds\\2012JUNE_kegg_cds\\"
    list_seq_file = glob(kegg_seq_dir+"*.csv")
```

```
for fName in list_seq_file:
    org_id = fName.split("\\")[-1].split("_")[0]
```

```
for s in parse(fName, 'fasta'):
                                                                                        print "Finished calculation!!!!!!!!!"
            orf id = s.id[4:]
                                                                                    if __name__=='__main__':main()
            orf_desc = s.description
            seq = str(s.seq)
            outputFile =
os.path.join(cur_dir,"output/"+org_id+"_ecoHost.csv")
            if seq[:3].upper() in ['ATG','GTG','TTG'] and seq[-3:].upper()
in ['TAA','TAG','TGA']:
                try:
                    s cai = heg ecoli.cai for gene(seq)
                except TypeError,e:
                    print e
                    s_cai = "None"
            else:
                print 'NOT CDS', orf_id
                s_cai = "None"
            resultGenes = findGenes(orf_desc)
            resultK0 = findKOId(orf desc)
            if resultKO != None:
                if len(resultKO) == 1:## only one KO id for this orf
                    record = [orf_id,orf_desc,len(s.seq),s_cai,org_id]
                    ko_id = resultK0[0]
                    record.append(ko_id)
                    #print record
                    printListRecordsToFile(outputFile, record, "\t")
                else:
                    for ko id in resultKO:## one orf has more than one KO
id
                        record = [orf_id,orf_desc,len(s.seq),s_cai,org_id]
                        record.append(ko id)
                        #print record
                        printListRecordsToFile(outputFile, record, "\t")
            else:
                record = [orf_id,orf_desc,len(s.seq),s_cai,org_id]
                record.append("None")
                printListRecordsToFile(outputFile, record, "\t")
```

Appendix A

A.2.2 "maxCAIscoreECnumbers_ecoHost.py"

```
# Author: Sunisa Chatsurachai --<>
# Purpose: To collecte maximum CAI score for each EC number -> suggest
heterolgous gene(s)
# Created: 17-May-12
import sys,os,re
import glob,time
def printListToFile(fName, record, delimiter="\t"):
   f = open(fName, 'a')
   f.write(delimiter.join(map(str,record)))
   f.write("\n")
   f.close()
def getECDict(fName):
   result = {}
   for line in open(fName):
       line = line.lstrip().rstrip().split("\t")
       ec id = line[0]
       ko ids = line[1].split("##")
       gene_ids = line[2].split("##")
       data = zip(ko ids,gene ids)
       data = list(set(data))
       result[ec id] = data
   return result
def checkNoCAI(cai data):
   if cai data == "None":
       return None
   else:
       return float(cai_data)
def getAllGenesFromOrganisms(targetDir,cur_koId):
   caiList = []
   orgList = []
   orfList = []
   for f in glob.glob(targetDir+'/*.csv'):
       for line in open(f):
            line = line.lstrip().rstrip().split("\t")
            cai data = line[-3]
```

org_id = line[-2] ko id = line[-1] orf_id = line[0] if ko id == cur koId: if checkNoCAI(cai data) != None: cur_cai = cai_data else: cur_cai = 0.00 caiList.append(cur cai) orgList.append(org id) orfList.append(orf id) return sorted(set(zip(caiList,orgList,orfList))) def getCAIFromAllSubunit(caiDict): allKK = [] allCai = [] allGene = [] for kk,vv in caiDict.iteritems(): keggID = kk caiTuples = vv[0] keggGene = vv[1]allKK.append(keggID) allGene.append(keggGene) allCai.extend(caiTuples) orgDict = {} for i,v,k in allCai: cai = float(i) orgID = vorfID = kif orgDict.has key(orgID):## already have this organism caiData,listOrf = orgDict[orgID] caiData.append(cai) listOrf.append(orfID) orgDict[orgID] =[caiData,listOrf] else: orgDict[orgID] = [[cai],[orfID]] result = [] for o, caiOrgs in orgDict.iteritems():

oId = o

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```
caiOrf = "##".join(caiOrgs[1])
       avg_cai = sum(caiOrgs[0],0.0)/len(caiOrgs[0])
       #print oId,avg cai,caiOrf,"##".join(allKK),"##".join(allGene)
       rec = (avg cai,oId,caiOrf,"##".join(allKK),"##".join(allGene))
       result.append(rec)
   return list(sorted(set(result)))
def getOrgDict(fName):
   result = {}
   for line in open(fName):
       line = line.lstrip().rstrip().split("\t")
       org id = line[0]
       org name = line[1]
       result[org id] = org name
   return result
def main():
   start = time.clock()
   cur dir = os.getcwd()
   targetDir = os.path.join(cur dir,"output")
   orgNameDict = getOrgDict(os.path.join(cur dir,"input/orgName.csv"))
   ecDict = getECDict(os.path.join(cur_dir,"input/ecLinkKOIDs.csv"))
```

```
ko_id, gene = kdata
                #print k,ko id,gene
               caiScores = getAllGenesFromOrganisms(targetDir,ko id)
               caiDict[ko id] = [caiScores,gene]
            allResults =
getCAIFromAllSubunit(caiDict)##(avg cai,oId,caiOrf,"##".join(allKK),"##".jo
in(allGene))
```

if allResults != []:

result = allResults[-1] maxCai = result[0] oId = result[1] ORFids = result[2]koIDs = result[3] genes = result[4]orgName = orgNameDict[oId]

rec = [ec number, 'KEGG', genes, maxCai, ORFids, oId, orgName, koIDs] printListToFile(outputFile,rec,"\t") else:

print "Empty CAI not Found!!!!!!!","\t",ec_number

end = time.clock()

print "Time elapsed = ", (end - start)/3600, "hrs" if __name__=='__main__':main()

outputFile = os.path.join(cur dir,"20120CT maxCAI ecoHost kegg.csv") for k,v in ecDict.iteritems(): ec number = kif len(v) == 1:## found only one KO id -> 1 EC number ko id, gene = v[0]#print k,ko id,gene caiScores = getAllGenesFromOrganisms(targetDir,ko id) if caiScores != []: maxCai,oId,ORFid = caiScores[-1] orgName = orgNameDict[oId] rec = [ec_number, 'KEGG',gene,maxCai,ORFid,oId,orgName,ko_id]

```
printListToFile(outputFile,rec,"\t")
```

else:

print "Empty CAI not Found!!!!!!!","\t",ec_number

```
else:## found more than one KO id -> 1 EC number
    caiDict = {}
    for kdata in v:
```

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A.3 Model simulations by MATLAB®

A.3.1 Main script "iJR904_maxTarget_r_glpk_par.m"

clear
clc
recycle('on');
curr_path= pwd;
cd(curr_path);
funct_folder = addpath(strcat(curr_path,'\function'));
start_time = clock;

%%% clock returns a 6-element date vector containing the current date and time in decimal form: %%% [year month day hour minute seconds]

% lb(v.glucose,1) = -20; % ub(v.glucose,1) = -20; % % lb(v.atpm,1) = 7.6; % ub(v.atpm,1) = 7.6; % % lb(v.o2,1) = -20;% ub(v.o2,1) = 0;% % lb(v.nh3,1) = -100; % ub(v.nh3,1) = 0;% % lb(v.so3,1) = -100; % ub(v.so3,1) = 0;% lb(v.pi,1) = -100;

% ub(v.pi,1) = 0;[wt ,v] = createLPMatrixBasicCon(rxn c nh3 file,rxn ext file nh3); save wt data basic condition.mat wt; % [wt ,v] = createLPMatrixMicroaerobic(rxn_c_nh3_file,rxn_ext_file_nh3); % save wt eco microaerobic condition.mat wt; %%%%%%%% Write simulation flux of wild type to text file %%%%%%%%%% % fid = fopen('iJR904 wt flux microaerobic.txt','w'); % for aa =1:1:size(wt.reaction id,1); % if iscell(wt.reaction id{aa,1}) ==0 || iscell(wt.reaction id{aa,2})== 0; % rxn data = wt.reaction id{aa,1}; % rxn id = wt.reaction_id{aa,2}; % rxn no = wt.reaction id{aa,3}; % rxn_flux = num2str(wt_flux(aa)); % else % rxn data = wt.reaction id{aa,1}{:}; % rxn id = wt.reaction id{aa,2}{:}; % rxn no = wt.reaction id{aa,3}; % rxn flux = num2str(wt_flux(aa)); % end % fprintf(fid,rxn data()); % fprintf(fid,'\t'); % fprintf(fid,rxn_id()); % fprintf(fid,'\t'); fprintf(fid,num2str(rxn_no)); % % fprintf(fid,'\t'); % fprintf(fid,rxn flux); % fprintf(fid, '\n'); % % end % fclose(fid); totalIteration = 23; matlabpool open 12 timeCals=zeros(totalIteration,3); for iteration=1:totalIteration;

if exist(strcat(curr_path,'iteration_',num2str(iteration)),'dir')==0;%% if no Directory, make directory for data mkdir(curr path, strcat('iteration ',num2str(iteration))); end nonNativeMetPath = strcat('C:\Users\sunisa\Desktop\sunisa\2012SEP BKM react\addReactions\assum e r\eco add\iteration ',num2str(iteration)); listAllFiles = dir(fullfile(nonNativeMetPath,'*.txt')); each iteration result = fullfile(strcat('iJR904 maxTarget producible basicCon r '.num2str(iteration),'.csv')); load wt data basic condition.mat; tic: parfor fName=1:length(listAllFiles); % parfor fName=1:10; targetFnamePath = fullfile(nonNativeMetPath,listAllFiles(fName).name); curFname = listAllFiles(fName).name; rxn_data = textread(targetFnamePath, '%s'); canRxnName = rxn data(1); addRxns = getAdditionalBKMrxn mod(targetFnamePath,size(wt.reaction matrix ce,1)); mutant reaction id = [wt.reaction id;addRxns]; canRxns = cellstr(rxn data((2:end),1));%% first line shows the information of connectable metabolites (BKM-react ID) [mutant rxns,mutant metabolites,met row] = convertStoiAsNumber(canRxns,size(wt.metabolites,1),wt.metabolites); [reactionMatrixExpand,lb,ub,~] = reactionMatrix(mutant rxns,size(wt.reaction matrix ce,1)+1,wt.lb,wt.ub,wt.r eaction matrix ce); %%% add lower and upper of target reaction %%% lb(size(reactionMatrixExpand,1),1) =0.0; ub(size(reactionMatrixExpand,1),1)=1000.0;

%%% Maximize biomass first %%%%

biomass_rxn = wt.biomass_rn; mutant_metabolites = unique(mutant_metabolites); mutant_lb = lb; mutant_ub = ub; targetReaction = size(reactionMatrixExpand,1);%% target reaction
of nonNative metabolite

[LP_matrixExpand] = makeLPMatrix_mod(reactionMatrixExpand,mutant_metabolites); [s,t] = size(LP_matrixExpand); %% s = no of metabolites, t = no of reactions; b = zeros(s,1); c(1,v.biomass+1) = 1; %% determine objective function objType = -1 %% for gurobi solver -> 1 = minimize, -1 = maximize obj function conType = repmat('S',1,s); varType = repmat('C',1,t); % [mt_flux,mt_fval,mt_exitflag,mt_output,mt_lambda] = gurobi_mex(c',objType,LP_matrixExpand,b,conType,mutant_lb,mutant_ub,varType);

[mt_flux,mt_fval, mt_status, mt_extra] =glpk (c, LP_matrixExpand, b, mutant_lb,mutant_ub, conType, varType, objType); mt_growth = num2str(mt_flux(v.biomass,1)); disp(['Expand model maximize E. coli growth (assume r) = ' mt_growth 'solution status if 2 , unique == ' num2str(mt_status)])

%%% Maximize Target production of mutant without fixing
growth %%%%%%%%%%%%%%%%%%
mt_lb = mutant_lb;
mt_ub = mutant_ub;
mt_lb(v.biomass+1,1) = 0;
mt_lb(v.biomass+1,1) = 0;
mt_ub(v.biomass+1,1) = mt_flux(v.biomass+1,1);
v_carbon = mt_flux(v.glucose,1);

[lpMatrixExpand] =
makeLPMatrix_mod(reactionMatrixExpand,mutant_metabolites);
[m,n] = size(lpMatrixExpand);
b_target = zeros(m,1);
c_target = zeros(1,n);
c_target(1,targetReaction) = 1; %% Target reaction
objType_target = -1; %% to maximize targer reaction flux
conType_target = repmat('S',1,m);

varType_target = repmat('C',1,n);
% [mt2_flux,mt2_fval,mt2_exitflag,mt2_output,mt2_lambda] =
gurobi_mex(c_target',objType_target,lpMatrixExpand,b_target,conType_target,
mt_lb,mt_ub,varType_target);

[mt2_flux,mt2_fval, mt2_status, mt2_extra] =glpk (c_target,lpMatrixExpand, b_target, mt_lb,mt_ub, conType_target, varType_target, objType_target);

disp(['Expand model maximize target of E. coli growth (r)= '
num2str(mt2_flux(v.biomass)) 'At iteration = ' num2str(iteration)
'Solution status if 2, unique == ' num2str(mt2_status)])
 disp(['Expand model maximize target of E. coli taget (r)= '
num2str(mt2_flux(targetReaction)) 'At iteration = ' num2str(iteration)
'Solution station if 2, unique == ' num2str(mt2 status)])

%% check target production higher than 1% of carbon source%%
if mt2_status ==5 && mt2_flux(targetReaction) > 0.00;

t_flux = num2str(mt2_flux(targetReaction)); t_growth = num2str(mt2_flux(v.biomass));

wfile= fopen(each_iteration_result,'a');
 fprintf(wfile,'%s\t%s\t%s\t%s\t^s\t%s\n',curFname(1:end4),t_flux,t_growth,num2str(iteration));
 fclose(wfile);

end

outputFile =
strcat(curr_path,'\iteration_',num2str(iteration),'\',listAllFiles(fName).n
ame(1:end-4),'_Flux.txt');
uniteOutputFluxTefile_pap(outputFile_mutant_papetion_id_mt2_flux);

writeOutputFluxToFile_par(outputFile,mutant_reaction_id,mt2_flux);

end

timeCals(iteration,1) = iteration; timeCals(iteration,2) = length(listAllFiles); % timeCals(iteration,2) = 10; timeCals(iteration,3) = toc;

fTime = fopen('eco_maxTarget_r_glpk_time.txt','a');

fprintf(fTime,'%s\t\%s\t%s\n',num2str(iteration),num2str(length(listAllFile s)),num2str(toc)); fclose(fTime);

end

matlabpool close
xlswrite('eco_maxTarget_r_glpk.xls',timeCals);
disp('Finised calculation !!!! E. coli assume r BY GLPK solver ')

%% clock ==> [year month day hour minute seconds] %%
%% start_time(1,1) = year
%% start_time (1,2) = month
%% start_time (1,3) = day
%% start_time (1,4) = hour
%% start_time (1,5) = minute
%% start_time(1,6) = seconds

end_time = clock; s_year = start_time(1,1); s_month = start_time(1,2); s_day = start_time(1,3); s_hour = start_time(1,4); s_min = start_time(1,5);

e_year = end_time(1,1); e_month = end_time(1,2); e_day = end_time(1,3); e_hour = end_time(1,4); e_min = end_time(1,5);

disp(['starting time Year Month Day Hour Minute = ' num2str(s_year)
num2str(s_month) num2str(s_day) num2str(s_hour) num2str(s_min)])
disp(['End time Year Month Day Hour Minute = ' num2str(e_year)
num2str(e_month) num2str(e_day) num2str(e_hour) num2str(e_min)])

ub(v.nh3,1) = 0;A.3.2 "createLPMatrixBasicCon.m" 1b(v.so3,1) = -20;function [wt,v] =createLPMatrixBasicCon(rxn_c_nh3_file,rxn_ext_file_nh3) ub(v.so3,1) = 0;lb(v.pi,1) = -20;[reaction_cytosol_data,reaction.cytosol_id] = getReactions(rxn_c_nh3_file); ub(v.pi,1) = 0;[reaction c.eq,reaction c.metabolite,m row]=convertStoiAsNumber(reaction cy tosol data,0,[]); [reaction_matrix_cytosol, lower_bound, upper_bound, ~] = reactionMatrix(reaction_c.eq,1,[],[],{}); wt.lpMatrix = LP matrix original; wt.biomass rn =v.biomass; wt.lb = lb; [~,rxn_extracellular] = xlsread(rxn_ext_file_nh3); wt.ub = ub; [reaction matrix withEXT, reaction ext id, lb, ub] = end addExtReaction(rxn_extracellular,reaction_matrix_cytosol,lower_bound,upper_ bound); met ext = reaction ext id(1:end,1); wt.cytosolRxns = reaction.cytosol id; wt.reaction_matrix_ce = reaction_matrix_withEXT; wt.metabolite ext = met ext; wt.reaction id =[reaction.cytosol_id;reaction_ext_id]; wt.metabolite c = reaction c.metabolite; wt.metabolites = unique([reaction c.metabolite;met ext]); [~,cc rns] = xlsread(rxn ext file nh3); [~,v]=keepTransportRn(wt.reaction_id,cc_rns); [LP matrix original] = makeLPMatrix_mod(reaction_matrix_withEXT,wt.metabolites); %%%%%% Case Basic condition %%%%%%%%%%%%%% lb(v.glucose,1) = -20;ub(v.glucose,1) = -20;lb(v.o2,1) = -20;ub(v.o2,1) = 0;

lb(v.nh3,1) = -20;

A.3.3 "getReactions.m"

```
function [rn_eq,rn_tag]= getReactions(target_file)
```

%%Second to keep metabolic information by split text data into cellstr data %%by using space as delimiter.

%% Ex:

%%

{'FADH2_TCA[c]';'+';'VITAMIN_K_{2}[c]';'+';'2.0';'C00080[e]';'=>';'FAD_TCA[c]';'+';'REDUCED-MENAQUINONE[c]';'+';'2.0';'C00080[c]';}

for k=1:size(rxns,1);%% loop for number of reaction in rxns

```
if k==1;
```

remain = rxns{1,1}; %% start from first or stoichometry coefficient
in first reaction.

position = 0;

kk= 0;

while kk < k;</pre>

%% token = strtok('str',delimiter) returns the first token in %% the text string str, that is, the first set of characters before a delimiter is encountered. %%The vector delimiter contains valid delimiter characters. Any leading delimiters are ignored. [str,remain] = strtok(remain); if strcmp(str,{''}); kk=2;%% update kk cause no more string in this reaction end position = position+1; result{position,1} = str; %% update position for collecting str found in each reaction

end

```
elseif k>1;%% not first reaction
  remain = rxns{k,1};
  kkk = 0;
  while kkk ==0;
    [str,remain] = strtok(remain);
    if strcmp(str,{''});
        kkk = 1;%% update kkk cause no more string in this reaction
    else
        result{position,1} = str;
        position = position+1;
    end
```

end

end % if k==1
end % for k=1:size(rxns,1)
rn_eq = result;
rn_tag = rxns;
end

A.3.4 "convertStoiAsNumber.m"

```
function [reaction metabolite met_rows] =
convertStoiAsNumber(rxn_cytosol,m_rows, metabolite_list)
```



```
array
```

```
if iscell(rxn cytosol{i,1});%%% convert cell to char for comparing with
string
%
          rxn_cytosol{i,1} = char(rxn_cytosol{i,1}{:});
        temp = char(rxn cytosol{i,1}{:});
       if str2num(temp) ~= 0;%% if temp is numeric:
            temp = str2num(temp);
            rxn cytosol{i,1} = temp;
        else
            pat = '\w+\[+[a-z]+\]'; %% find pattern of compartment such
as[c], [m] ..
            if regexp(temp, pat) ~= 0; %% found metabolites from the
compartment
                m_rows = m_rows+1; %% set position of metabolite in array
                metabolite list{m rows,1} = temp;
                rxn_cytosol{i,1} = temp;
            else
                temp = temp;
                rxn_cytosol{i,1} = temp;
            end
        end
    else
```

```
temp = rxn cytosol{i,1};
```

```
if str2num(temp) ~= 0;%% if temp is numeric:
            temp = str2num(temp);
            rxn cytosol{i,1} = temp;
        else
            pat = '\w+\[+[a-z]+\]'; %% find pattern of compartment such
as[c], [m] ..
            if regexp(temp, pat) ~= 0; %% found metabolites from the
compartment
               m rows = m rows+1; %% set position of metabolite in array
               metabolite_list{m_rows,1} = temp;
               rxn cytosol{i,1} = temp;
            else
               temp = temp;
               rxn cytosol{i,1} = temp;
            end
        end
```

```
end
```

```
end
```

```
metabolite = metabolite_list;
reaction = rxn_cytosol;
met rows = m rows;
```

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A.3.5 "reactionMatrix.m"

function [matrix, lb, ub, row] =
reactionMatrix(rxn_cytosol,rows,lower_bound,upper_bound,reaction_matrix)

% create reaction matrix % input reaction in cytosol % retrun reaction matrix, lowerbound, upperbound and row of reaction

% reaction_matrix = {}; % rows = 1; % lower_bound = []; % upper_bound = []; columns = 1; % Making reaction matrix %%%

for i=1:size(rxn_cytosol,1)-1;%% minus one because last metabolite
determine at line 61
 reaction_matrix{rows,columns} = rxn_cytosol{i};
 if isnumeric(rxn_cytosol{i}) ==1;
 rows = rows; % found stoichiometry so row number not increase

elseif strcmp(rxn_cytosol{i},'+') == 1 | strcmp(rxn_cytosol{i},'=>')
== 1 | strcmp(rxn_cytosol{i},'<=>') == 1;
 rows = rows;

if strcmp(rxn_cytosol{i},'=>') == 1;

lower_bound(rows,1)= 0.00; upper_bound(rows,1) = 1000.0;

elseif strcmp(rxn_cytosol{i},'<=>') == 1;

lower_bound(rows,1)=(-1)*(1000.0); upper_bound(rows,1) = 1000.0;

end

elseif isnumeric(rxn_cytosol{i}) == 0 & strcmp(rxn_cytosol{i},'+') == 0 & strcmp(rxn_cytosol{i},'=>') == 0 ... & strcmp(rxn_cytosol{i},'<=>')== 0; %%% final metabolite name (cpd_id) from previous reaction %%%

end

end end

columns = columns+1; %% if not metabolite have to extend column for equal size of all reaction

```
end
```

reaction_matrix {rows, columns} = rxn_cytosol{size(rxn_cytosol,1)}; %%
determine the last metabolite of reaction
rows = rows+1; % adding one row for starting next reaction from external
reaction
matrix = reaction_matrix;
lb = lower_bound;
ub = upper_bound;
row = rows;

A.3.6 "makeLPMatrix_mod.m"

```
function [LP_matrix] =
makeLPMatrix_mod(reaction_matrix_withEXT,Allmetabolites)
```

%%%% Making LP matrix %%%%%%

```
[m,n] = size(reaction_matrix_withEXT);
LP_matrix =sparse(length(Allmetabolites),m);
```

%%% construct stoi matrix %%%%%

```
nonMetsList = { '<=','=>','<=>' };
for i =1:m
    oneRxn = reaction_matrix_withEXT(i,1:end);
    for sign=1:3
        pos = find(strcmp(nonMetsList(sign),oneRxn));
        if (~isempty(pos))
            sign_pos = pos;
        end
end
```

end

end

```
for k=1:length(product_mets);
    item = product_mets{1,k};
    m_pos = find(strcmpi(item,Allmetabolites));
    if (~isempty(m_pos)) && k==1; %% first of product
        LP_matrix(m_pos,i) = 1;
    elseif (~isempty(m_pos)) && k >1;
        if isempty(product_mets{1,k-1})||(strcmp(product_mets{1,k-1}, +'))==1;
        LP_matrix(m_pos,i) = 1;
        elseif isnumeric(product_mets{1,k-1});
        LP_matrix(m_pos,i) = (1)*(product_mets{1,k-1});
        end
    end
end
```

```
reactant_mets = reaction_matrix_withEXT(i,1:sign_pos-1);
product_mets = reaction_matrix_withEXT(i,sign_pos+1:end);
```

```
for k=1:length(reactant_mets);
    item = reactant_mets{1,k};
    m_pos = find(strcmpi(item,Allmetabolites));
```

```
if (~isempty(m_pos)) && k==1; %% found start metabolite %%
```

```
LP_matrix(m_pos,i) = -1;
```

```
elseif (~isempty(m_pos)) && k > 1; %% found metabolite %%
```

 \geq

0

σ

D

A.3.7 "addExtReaction.m"

function [reaction_matrix_withEXT,reaction_ext_id,lower_bound,upper_bound]
= addExtReaction(rxn_ext_item,reaction_matrix,lb,ub)

```
%% add extracellular reaction %%%
%% rxn_ext_item(1,1:end) = {'[e]12ppd-S','C02917','=>';}
```

```
reaction_ext_id = cell(size(rxn_ext_item,1),2);
rows = size(reaction_matrix,1);
rows = rows+1;
columns = 1;
```

for i=1:size(rxn_ext_item,1);

reaction_ext_id{i,1} = strcat(rxn_ext_item{i,2},'[e]');%% metabolite reaction_ext_id{i,2} = rxn_ext_item{i,1};%% reaction id reaction_ext_id{i,3} = rows;%% no_reaction in reaction_matrix

reaction_matrix{rows,columns} =strcat(rxn_ext_item{i,2},'[e]');%%
external metabolite in KEGG ID

```
if strcmp(rxn_ext_item{i,3},'=>')==1;
    lb(rows,1) = 0.0;
    ub(rows,1) = 1000.0;
    columns = columns+1;
    reaction_matrix{rows,columns} = rxn_ext_item{i,3};
elseif strcmp(rxn_ext_item{i,3},'<=>')==1;
    lb(rows,1) = -1000.0;
    ub(rows,1) = 1000.0;
```

```
columns = columns+1;
reaction_matrix{rows,columns} = rxn_ext_item{i,3};
end
rows = rows+1;
columns=1;
lower_bound = lb;
upper_bound = ub;
reaction_matrix_withEXT = reaction_matrix;
```

A.3.8 "getAdditionalBKMrxn_mod.m"

```
function no_add_rxns =
getAdditionalBKMrxn_mod(target_fName_path,original_reaction_no)
f_target = fopen(target_fName_path,'r');
tLine = fgetl(f_target);
fLine = tLine;
findRxnPos = regexpi(fLine,'R[0-9][0-9][0-9][0-9][0-9]');
rxnsList = cell(size(findRxnPos,2),1);
```

for item=1:size(findRxnPos,2);
 rn_id = fLine(findRxnPos(item):findRxnPos(item)+5);
 rxnsList{item} = rn_id;
end

rxnsList = unique(rxnsList); rxnsList = [rxnsList;'addRxn1';'addRxn2'];%% add transporter reactions (by diffusion) no_add_rxns = cell(length(rxnsList),3);

```
pos = 0;
while pos < length(no_add_rxns);
    pos = pos+1;
    tLine = fgetl(f_target);
    no_add_rxns{pos,1} = tLine;
    no_add_rxns{pos,2} = rxnsList{pos};
    no_add_rxns{pos,3} = original_reaction_no+pos;
```

end

fclose(f_target);
A.3.9 "keepTransportRn.m"

<pre>function [central_reaction,v]= keepTransportRn(reaction_id,t_reaction) central_reaction = cell(size(t_reaction,1),4); for vv=1:size(t_reaction,1); target_rnID = t_reaction(vv,1);</pre>	
<pre>for k=1:size(reaction_id,1); x= reaction_id(k,2); r_id = x{:};</pre>	
<pre>if strcmpi(r_id,target_rnID) ==1; central_reaction{vv,1} =reaction_id{k,3}; central_reaction{vv,2} = t_reaction(vv,1);%% rn_id central_reaction{vv,3} = t_reaction(vv,2);%% ec_id central_reaction{vv,4} = t_reaction(vv,3);%% pathway name</pre>	
end	and
<pre>if strcmp(r_id,'BIOMASS_TRANS')==1; v.biomass =reaction_id{k,3};</pre>	end
<pre>elseif strcmp(r_id, '[e]etoh')==1; v.eth = reaction_id{k,3};</pre>	end
<pre>elseif strcmp(r_id,'[e]glc-D')==1; v.glucose = reaction_id{k,3};</pre>	

v.co2=reaction_id{k,3};

elseif strcmp(r_id,'[e]nh3')==1; v.nh3=reaction_id{k,3};

elseif strcmp(r_id,'[e]pi')==1; v.pi =reaction_id{k,3};

elseif strcmp(r_id,'[e]so4')==1; v.so3 =reaction_id{k,3};

elseif strcmp(r_id, '[e]h2o')==1; v.h2o =reaction_id{k,3}; elseif strcmp(r_id, 'ATPM') == 1; v.atpm = reaction_id{k,3};

nd

end

elseif strcmp(r_id,'[e]lac-D')==1;

elseif strcmp(r_id,'[e]succ')==1;

v.succ =reaction_id{k,3};

v.lactate =reaction_id{k,3};

elseif strcmp(r_id, '[e]o2')==1; v.o2 =reaction_id{k,3}; elseif strcmp(r_id, '[e]ac')==1; v.acetate =reaction_id{k,3};

A.3.10 "writeOutputFluxToFile.m"

function

```
writeOutputFluxToFile_par(fileName,mutant_reaction_id,mutant_maxTargetFlux)
```

```
f_flux = fopen(fileName,'w');
flux = mutant_maxTargetFlux;
parfor aa =1:1:size(mutant_reaction_id,1);
   if iscell(mutant_reaction_id{aa,1}) ==0 ||
iscell(mutant reaction id{aa,2})== 0;
       rxn_data = mutant_reaction_id{aa,1};
       rxn id = mutant reaction id{aa,2};
       rxn_no = mutant_reaction_id{aa,3};
       rxn_flux = num2str(flux(aa));
   else
       rxn_data = mutant_reaction_id{aa,1}{:};
       rxn_id = mutant_reaction_id{aa,2}{:};
       rxn_no = mutant_reaction_id{aa,3};
       rxn_flux = num2str(flux(aa));
    end
    fprintf(f_flux,rxn_data());
   fprintf(f_flux,'\t');
   fprintf(f_flux,rxn_id());
    fprintf(f_flux,'\t');
    fprintf(f_flux,num2str(rxn_no));
   fprintf(f_flux,'\t');
   fprintf(f_flux,rxn_flux);
    fprintf(f_flux,'\n');
end
fclose(f_flux);
```

Appendix B

Table B.1 Values of relative adaptiveness (w) of codon generated from highlyexpressed genes of *B. subtilis, E. coli,* and *S. cerevisiae*.

		B. subtilis	E. coli	S. cerevisiae			B. subtilis	E. coli	S. cerevisiae
	GCA	0.926	0.715	0.094		СТА	0.120	0.007	0.149
ala	GCC	0.263	0.504	0.428	leu	СТС	0.142	0.091	0.015
	GCG	0.700	1.000	0.020		CTG	0.353	1.000	0.043
	GCT	1.000	0.815	1.000		сп	1.000	0.060	0.066
						ΤΤΑ	0.508	0.034	0.310
arg	AGA	0.304	0.005	1.000		ΠG	0.263	0.051	1.000
	AGG	0.004	0.002	0.052					
	CGA	0.059	0.009	0.005	lys	AAA	1.000	1.000	0.328
	CGC	0.783	0.529	0.015		AAG	0.205	0.244	1.000
	CGG	0.023	0.004	0.004					
	CGT	1.000	1.000	0.213	met	ATG	1.000	1.000	1.000
asn	AAC	1.000	1.000	1.000	phe	πс	1.000	1.000	1.000
	AAT	0.480	0.189	0.309		Π	0.726	0.400	0.402
	GAC	0.759	1.000	1.000		CCA	0.622	0.179	1.000
ash	GAT	1.000	0.844	0.896	pro	CCC	0.022	0.016	0.034
					pro	CCG	0.829	1.000	0.014
cys	TGC	1.000	1.000	0.242		ССТ	1.000	0.122	0.212
	TGT	0.783	0.550	1.000					
						AGC	0.567	0.814	0.120
gln	CAA	1.000	0.210	1.000		AGT	0.187	0.123	0.128
	CAG	0.461	1.000	0.091		TCA	0.645	0.146	0.165
					301	тсс	0.191	0.873	0.634
	GAA	1.000	1.000	1.000		TCG	0.046	0.218	0.044
giu	GAG	0.288	0.290	0.122		тст	1.000	1.000	1.000
	GGA	0.820	0.033	0.042	stop	TAA	1.000	1.000	1.000
-1.	GGC	1.000	0.869	0.095		TAG	0.057	0.009	0.173
gıy	GGG	0.128	0.071	0.022		TGA	0.028	0.135	0.204
	GGT	0.840	1.000	1.000					
						ACA	1.000	0.067	0.167
his	CAC	0.966	1.000	1.000	thr	ACC	0.078	1.000	0.840
	CAT	1.000	0.382	0.526		ACG	0.322	0.195	0.035
						ACT	0.552	0.488	1.000
ile	ATA	0.031	0.004	0.076					
	ATC	1.000	1.000	0.959	trp	TGG	1.000	1.000	1.000
	ΑΤΤ	0.895	0.487	1.000					
						TAC	1.000	1.000	1.000
					tyr	TAT	0.843	0.530	0.274
						GTA	0.742	0 4 9 4	0.064

GTC

GTG

GTT

val

0.399

0.410

1.000

0.324

0.721

1.000

0.712

0.081

1.000

List of publications

- <u>Chatsurachai, S.</u>, Furusawa, C., and Shimizu, H. (2012). An *in silico* platform for the design of heterologous pathways in nonnative metabolite production. *BMC Bioinformatics*, 13, 93.
- <u>Chatsurachai, S.</u>, Furusawa, C., and Shimizu, H. (2013). ArtPathDesign Rational heterologous pathway design system for the production of nonnative metabolites. *J Biosci Bioeng* (in press)

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