## Stoichiometric Model of Escherichia coli Metabolism: Incorporation of **Growth-Rate Dependent Biomass Composition and Mechanistic Energy Requirements**

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Abstract: A stoichiometric model of metabolism was developed to describe the balance of metabolic reactions during steady-state growth of Escherichia coli on glucose (or metabolic intermediates) and mineral salts. The model incorporates 153 reversible and 147 irreversible reactions and 289 metabolites from several metabolic data bases for the biosynthesis of the macromolecular precursors, coenzymes, and prosthetic groups necessary for synthesis of all cellular macromolecules. Correlations describing how the cellular composition changes with growth rate were developed from experimental data and were used to calculate the drain of precursors to macromolecules, coenzymes, and prosthetic groups from the metabolic network for the synthesis of those macromolecules at a specific growth rate. Energy requirements for macromolecular polymerization and proofreading, transport of metabolites, and maintenance of transmembrane gradients were included in the model rather than a lumped maintenance energy term. The underdetermined set of equations was solved using the Simplex algorithm, employing realistic objective functions and constraints; the drain of precursors, coenzymes, and prosthetic groups and the energy requirements for the synthesis of macromolecules served as the primary set of constraints. The model accurately predicted experimentally determined metabolic fluxes for aerobic growth on acetate or acetate plus glucose. In addition, the model predicted the genetic and metabolic regulation that must occur for growth under different conditions, such as the opening of the glyoxylate shunt during growth on acetate and the branching of the tricarboxylic acid cycle under anaerobic growth. Sensitivity analyses were performed to determine the flexibility of pathways and the effects of different rates and growth conditions on the distribution of fluxes. © 1997 John Wiley & Sons, Inc. Biotechnol Bioeng 56: 398-421, 1997.

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#### INTRODUCTION

A central problem in metabolic engineering is understanding how the cell balances its energy and biosynthetic needs

Correspondence to: Jay Keasling Contract grant sponsor: NSF Contract grant number: BES-9502495 growth kinetics of the cell are incorporated into the model through the energy and biomass requirements, which are functions of the growth rate. The stoichiometry of metabolism is well defined and variations among different cells are limited to a few reactions. There are two different methods of using stoichiometry to study bioreaction networks. The first method reduces the stoichiometric matrix to an over-

ous conditions. Simple carbon sources and mineral salts are taken into the cell and transformed into the complex biopolymers and cofactors that compose the cell, while generating the metabolic energy necessary to make these complex biomolecules. There have been many attempts to study regulation of metabolism using mathematical models. The simplest description of cellular metabolism uses order of magnitude calculations to determine the metabolic yields from cell composition, measured substrate uptake and product syn-

(catabolism and anabolism) for optimal growth under vari-

thesis, and growth rates (Blanch and Clark, 1996; Savinell et al., 1989). However, this analysis provides limited formation about the fluxes of intermediary metabolism. A more detailed analysis uses the stoichiometry of biochemical pathways and cell composition data to estimate the steady-state mass and energy distributions (Nissen et al., 1997; Savinell and Palsson, 1992a; Tsai et al., 1988; Vallino and Stephanopoulos, 1993; Varma et al., 1993); experimentally determined intermediary fluxes can be used as constraints to improve the accuracy of calculations. The most detailed level of metabolic pathway analysis examines the dynamic behavior of cell metabolism and requires kinetic and thermodynamic data, most of which is currently un-

available: kinetic parameter measurements are difficult to

obtain and kinetic models developed from in vitro measure-

ments may not apply in vivo.

Metabolic modeling, based on the stoichiometry of the reactions, does not require kinetic parameters and information about the kinetic mechanism of each enzyme. The determined form and then uses linear regression to find the flux distribution (Tsai et al., 1988; Vallino and Stephanopoulos, 1993). Independent measurements must be added

or a number of reactions must be removed or constrained to render the matrix nonsingular. Unfortunately, the pathways neglected for mathematical reasons may, in fact, be active. Removing entire pathways may cause large changes in the calculated fluxes. Several previous models solved an overdetermined system by constraining the stoichiometric matrix through inspection (Papoutsakis, 1984, Tsai et al., 1988).

The second method optimizes an underdetermined matrix using different objective functions and allows retention of the entire network. Applications of this approach to a subset of hybridoma, yeast, and Escherichia coli metabolism demonstrated the utility of this technique (Majewski and Domach, 1990; Savinell et al., 1989; Savinell and Palsson, 1992a,b; van Gulik and Heijnen, 1995; Varma and Palsson, 1993, 1994a,b; 1995; Varma et al., 1993). Majewski and Domach were able to predict the secretion of acetate during growth of E. coli on glucose (Majewski and Domach, 1990). Varma and Palsson showed the effect of oxygen availability on acetate secretion and the metabolic capabilities of E. coli to overproduce amino acids and other products (Varma and Palsson, 1993; 1994a,b; 1995; Varma et al., 1993). However, in all of these cases, the metabolic pathways were not complete, numerous reactions were lumped, and there was no accounting for the effects of growth rate on cellular composition and energy requirements.

We developed a detailed stoichiometric model of E. coli metabolism that includes a more complete data base of known reactions involved in the catabolism of glucose, acetate, or tricarboxylic acid (TCA) cycle intermediates. None of the pathways are lumped to reduce the matrix so that any future simulations studying deletions or mutations in pathways would not require generation of an entirely new stoichiometric matrix. A detailed stoichiometric matrix allows us to study deletions or mutations of individual enzymes by setting constraints of the flux values for those enzymes. The model uses the precursor requirements (calculated from the known composition of the bacterial cell) (Neidhardt et al., 1990) and solves for the fluxes through the internal metabolic pathways using linear optimization. Rather than include a "maintenance energy" term, energy demands for growth are calculated from mechanistic energy requirements for macromolecular polymerization and proofreading, transport of metabolites, and maintenance of transmembrane gradients. Experimental data on nutrient uptake and secretion can be incorporated into the model. The model is solved using linear optimization and predicts the metabolic regulation observed during growth under different conditions and on different carbon sources.

#### **MODEL DEFINITION**

The basis for this flux-based model of metabolism is a mass balance on the metabolites in *E. coli*,

$$\frac{d\mathbf{x}}{dt} = \mathbf{S} \cdot \mathbf{v} - \mathbf{b} \tag{1}$$

where  $\mathbf{x}$  is the vector of metabolite concentrations  $(n \times 1)$  dimension),  $\mathbf{S}$  is the stoichiometric matrix  $(n \times m)$  dimension),  $\mathbf{v}$  is a vector of reaction rates or fluxes through the metabolic reactions  $(m \times 1)$  dimension), and  $\mathbf{b}$  is the vector for consumption and secretion rates of metabolites and for biosynthetic requirements for cellular macromolecules  $(n \times 1)$  dimension). Under balanced growth conditions, the concentrations of intracellular metabolites are constant with time:

$$\mathbf{S} \cdot \mathbf{v} = \mathbf{b}. \tag{2}$$

The goal of this model is to determine how mass and energy is allocated within the network of metabolic reactions ( $\mathbf{v}$ ). We begin with descriptions of the composition of the cell (which determines  $\mathbf{b}$ ) and of the reactions involved in synthesizing the precursors and energy required for growth (which determine  $\mathbf{S}$ ).

#### **Biomass and Energy Requirements**

An average E. coli B/r cell growing exponentially at 37°C under aerobic conditions in glucose minimal medium with a doubling time of approximately 40 min has a dry weight of  $2.8 \times 10^{-13}$  g. The dry weight is 55% protein, 20.5% RNA, 3.1% DNA, 9.1% lipids, 3.4% lipopolysaccharides, 2.5% peptidoglycan, 2.5% glycogen, 0.4% polyamines, and 3.5% other metabolites, cofactors, and ions (Neidhardt, 1987). The types and amounts of precursors required to synthesize these macromolecules at a given growth rate were determined from the composition of each of the macromolecules: the amino acid composition of proteins and the nucleotide composition of RNA and DNA are listed in Table I, the phospholipid composition in Table II, and the fatty acid composition in Table III. The amounts of cofactors and energy carriers present per gram dry weight (DW) of biomass are listed in Table IV. The b vector contains these precursor requirements to account for synthesis of the cellular macromolecules.

In addition to the precursors required to synthesize macromolecules, energy and reducing equivalents are also required for growth. Table V presents the energy requirements for 1 g of E. coli B/r cells growing aerobically with a 40-min doubling time at 37°C in glucose minimal medium. The energy requirements for DNA production include that needed by helicase to unwind the helix, the synthesis of the primer RNA to Okazaki fragments and ligation of the fragments, proofreading by DNA polymerase III, adjustment of the torsional tension of each chromosomal domain, and methylation of newly synthesized DNA (Neidhardt et al., 1990). The energy requirements for stable RNA production include that for discarding segments of primary transcripts and that for modifications (Neidhardt et al., 1990). The energy requirements for protein synthesis include that for mRNA synthesis, charging tRNAs with amino acids and incorporation of amino acids into protein, and proofreading, assembly, and modification of the protein (Neidhardt et al.,

**Table I.** Precursor requirements for synthesis of 40-min cell.

|             | •       |           |
|-------------|---------|-----------|
| Component   | Percent | μmol/g DW |
| Amino acids |         |           |
| Ala         | 9.60    | 488       |
| Arg         | 5.53    | 281       |
| Asn         | 4.51    | 229       |
| Asp         | 4.51    | 229       |
| Cys         | 1.71    | 87        |
| Glu         | 4.92    | 250       |
| Gln         | 4.92    | 250       |
| Gly         | 4.92    | 582       |
| His         | 1.77    | 90        |
| Ile         | 5.43    | 276       |
| Leu         | 8.42    | 428       |
| Lys         | 6.42    | 326       |
| Met         | 2.87    | 146       |
| Phe         | 3.46    | 176       |
| Pro         | 4.13    | 210       |
| Ser         | 4.03    | 205       |
| Thr         | 4.74    | 241       |
| Trp         | 1.06    | 54        |
| Tyr         | 2.58    | 131       |
| Val         | 7.91    | 402       |
| rNTPs       |         |           |
| ATP         | 26.2    | 165       |
| GTP         | 32.2    | 203       |
| CTP         | 20.0    | 126       |
| UTP         | 21.6    | 136       |
| dNTPS       |         |           |
| dATP        | 24.7    | 24.7      |
| dGTP        | 25.4    | 25.4      |
| dCTP        | 25.4    | 25.4      |
| TTP         | 24.7    | 24.7      |

Amino acids, ribonucleotide triphosphates (rNTPs), and deoxyribonucleotide triphosphates (dNTPs) are given in percentage of the biopolymer and as micromoles per gram dry weight ( $\mu$ mol/g DW) of cells (Neidhardt, 1987).

1990). These energy requirements were also included in the **b** vector.

#### Growth-Rate Dependence of Cell Composition

The macromolecular composition and energy requirements listed above are not the same for cells growing at different

rates. For example, RNA content increases with growth rate whereas DNA and protein contents decrease with the growth rate (Bremer and Dennis, 1996; Brunschede et al., 1977). To solve for the fluxes through the metabolic reactions for doubling times other than 40 min, correlations were developed from experimental data for RNA, DNA, protein, surface area (for membrane components), and glycogen content as a function of specific growth rate (µ) for exponentially growing cells or dilution rate for continuous cultures (Fig. 1, Table VI). As the macromolecular composition of the cell changes with growth rate, so must the energy requirements to synthesize these macromolecules, which were correlated with the macromolecular needs (Fig. 2). Because protein is one of the most energetically expensive macromolecules and because the relative amount of protein decreases with increasing growth rate, the total energy expended by the cell (per g DW) actually decreases with growth rate.

### Metabolic Pathways

The transformation of a simple carbon source and mineral salts to the biomass and energy requirements for growth is facilitated through the metabolic reactions. The stoichiometric matrix S contains the stoichiometry of all reactions incorporated into the model. Included in this model were 153 reversible and 147 irreversible reactions (Appendix A) and 289 metabolites (Appendix B) compiled primarily from three sources: the Boehringer-Mannheim wall chart (Michal, 1993), chapters 14–19, 24–41, 44, 48, 49, 67, 69, 72, 75, and 87 of Neidhardt et al. (1996), and the Ecocyc data base (Karp et al., 1996). There are discrepancies between the number of reactions included in this model and the Ecocyc data base because a number of reactions in the Ecocyc data base have the same product but alternative substrates (e.g., NH<sub>4</sub><sup>+</sup> versus glutamine as a source for nitrogen) when only one of these is known to be used under physiological conditions in E. coli. Finally, a number of transport reactions were included in the model to account for uptake or secretion of inorganics or metabolites. In many cases, these transport steps deplete or enhance the transmembrane proton gradient.

Table II. Phospholipid composition of E. coli strains.

| Specific growth rate | 1.3  |      | 1.04 |      |      | 0.92 |      |      |     |
|----------------------|------|------|------|------|------|------|------|------|-----|
| $(h^{-1})$           | PE%  | PG%  | CL%  | PE%  | PG%  | CL%  | PE%  | PG%  | CL% |
| Strains              |      |      |      |      |      |      |      |      |     |
| E. coli B/r          |      |      |      | 75.0 | 18.0 | 5.0  |      |      |     |
| E. coli B/r          | 77.0 | 21.3 | 1.1  |      |      |      | 78.0 | 20.4 | 1.5 |
| E. coli B/r          | 74.8 | 20.6 | 2.3  |      |      |      | 75.1 | 20.3 | 1.9 |
| E. coli AX14         | 71.3 | 19.2 | 3.4  |      |      |      | 69.5 | 19.4 | 4.6 |
| E. coli K-12         | 67.1 | 18.7 | 5.1  |      |      |      | 68.8 | 19.3 | 5.7 |
| Overall average      | 73.0 | 19.7 | 3.4  |      |      |      | 72.8 | 19.8 | 3.4 |
| Overall SD           | 3.9  | 1.0  | 1.8  |      |      |      | 4.4  | 0.6  | 2.0 |

Data for a specific growth rate of  $1.04\ h^{-1}$  are from Neidhardt (1987). All other data are from Ballesta and Schaechter (1971). PE, phosphatidylethanolamine; PG, phosphatidylglycerol; CL, cardiolipin.

**Table III.** Fatty acid composition of *E. coli* lipids.

| Fatty acid                     | Total fatty acid (%) |
|--------------------------------|----------------------|
| Myristic acid, 14:00           | 2.68                 |
| Myristoleic acid, 14:01        | 7.70                 |
| Palmitic acid, 16:00           | 38.23                |
| Palmitoleic acid, 16:01        | 10.74                |
| Heptadecenoic acid, $17\Delta$ | 16.11                |
| cis-Vaccenic acid, 18:00       | 0.90                 |
| Oleic acid, 18:01              | 17.91                |
| Nonadecenoic acid, $19\Delta$  | 5.73                 |

Data from Bright-Gaertner and Proulx (1972), Kanemasa et al. (1967), Mavis and Vagelos (1972), and Neidhardt (1987).

Although the stoichiometry of most biosynthetic reactions is well known, the required cofactor(s) for a particular reaction may not be so well known. When known, correct NADH or NADPH was used in the stoichiometry of a reaction. When it was known that NADP/NADPH and NAD/NADH could be used interchangeably for the same reaction, pathways for both reactions were included; however, if no information was available to determine which electron carrier is used, it was assumed that NAD/NADH was used for anabolism and NADP/NADPH was used for catabolism.

In contrast to most of the biosynthetic reactions that have well-known stoichiometry, the pathways involved in electron transport and oxidative phosphorylation have variable stoichiometry due to the use of different dehydrogenases and cytochromes: the NADH dehydrogenases NDH-I and NDH-II transport 2 H<sup>+</sup>/e<sup>-</sup> and 0 H<sup>+</sup>/e<sup>-</sup>, respectively (Gennis and Stewart, 1996); the cytochromes cyt bd and cyt bo3 transport 1 H<sup>+</sup>/e<sup>-</sup> and 2 H<sup>+</sup>/e<sup>-</sup>, respectively (Gennis and Stewart, 1996); the number of H<sup>+</sup> transported into the cell by the membrane bound H<sup>+</sup>-ATPase to phosphorylate ADP has been estimated as 2-4, with 3 being the most likely (Harold and Maloney, 1996). Pathways for all possible stoichiometries were incorporated into the model. Thus, the P/O ratio can be a noninteger value (Neidhardt et al., 1990), because it is a function of multiple enzymes being used in parallel for respiration. The model simulates the noninteger P/O ratios by incorporating all known respiration pathways

**Table IV.** Amounts of cofactors, energy carriers, and other molecules in *E. coli*.

| Metabolite                 | μmol/g DW |
|----------------------------|-----------|
| 5-Methyl-THF               | 50.0      |
| Putrescine                 | 35.0      |
| Spermidine                 | 7.0       |
| NAD                        | 2.15      |
| NADH                       | 0.05      |
| NADP                       | 0.13      |
| NADPH                      | 0.4       |
| UDP-Glucose (soluble pool) | 3.0       |
| ATP (soluble pool)         | 4.0       |
| ADP (soluble pool)         | 2.0       |
| AMP (soluble pool)         | 1.0       |

Data from Lowry et al. (1971) and Penfound and Foster (1996).

Table V. Energy requirements for polymerization and processing of macromolecules.

| Process                                       | Energy required | Reference              |
|---|-----------------|------------------------|
| Protein synthesis and processing <sup>a</sup> |                 |                        |
| Activation and incorporation                  | 4.0             | Neidhardt et al., 1990 |
| mRNA synthesis                                | 0.2             | Neidhardt et al., 1990 |
| Proofreading                                  | 0.1             | Neidhardt et al., 1990 |
| Assembly and modification                     | 0.006           | Neidhardt et al., 1990 |
| RNA synthesis and processing <sup>b</sup>     |                 |                        |
| Discarding segments                           | 0.38            | Neidhardt et al., 1990 |
| Modification                                  | 0.02            | Neidhardt et al., 1990 |
| DNA synthesis and processing <sup>b</sup>     |                 |                        |
| Unwinding helix                               | 1.0             | Neidhardt et al., 1990 |
| Proofreading                                  | 0.36            | Neidhardt et al., 1990 |
| Discontinuous synthesis                       | 0.006           | Neidhardt et al., 1990 |
| Negative supercoiling                         | 0.005           | Neidhardt et al., 1990 |
| Methylation                                   | 0.001           | Neidhardt et al., 1990 |
| Membrane processes <sup>c</sup>               |                 |                        |
| Proton leakage                                | 62.9            | Maloney, 1987          |

Data from Neidhardt et al. (1990).

into the stoichiometric matrix and allowing fluxes through multiple pathways simultaneously.

#### Solution Method

For the system described above, the number of reactions (m) is greater than the number of metabolites (n). Because multiple solutions exist, linear optimization was used to determine the fluxes (v). Linear optimization requires objective functions for solution:

minimize/maximize: 
$$Z = \sum_{i} c_{i} v_{i}$$
 (3)

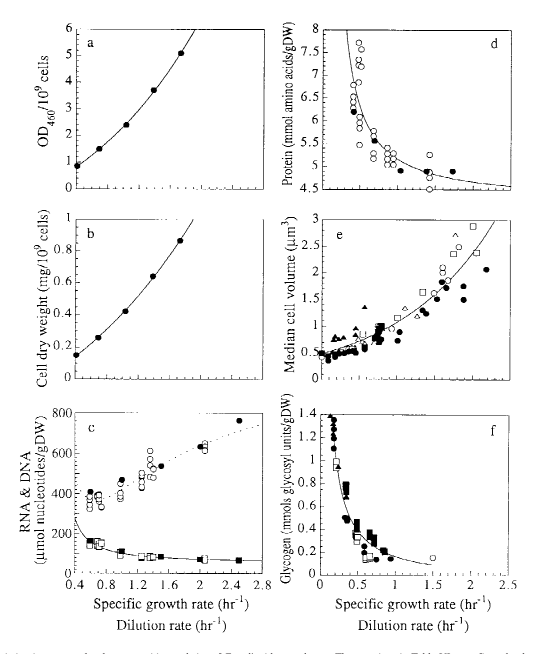
where  $c_i$  are the weights and  $v_i$  are the elements of the flux vector. Some of the objective functions that were used included minimization or maximization of ATP usage, substrate uptake, growth rate, and product synthesis. Typically, maximization of growth rate and minimization or maximization of metabolite secretion were used as objective functions.

For the objective of maximizing growth rate, the flux of precursors for a biomass composition corresponding to a particular growth rate was calculated using the relationships in Table VI. The amount of each precursor per unit time necessary for synthesis of biomass at a particular growth rate was used as the upper bound for the flux of each precursor, and the lower bound was set to zero. Then the model was forced to maximize the synthesis rate of each precursor for biomass composition, such as individual amino acids, nucleotides, and glycogen monomer units. If the model was unable to match the necessary flux of one or more of the precursors for a biomass composition consistent with a particular growth rate, the biomass composition was recalculated at another growth rate based on the precursor metabo-

<sup>&</sup>lt;sup>a</sup>μmol ATP/μmol amino acid.

<sup>&</sup>lt;sup>b</sup>μmol ATP/μmol nucleotide.

cμmol H+/g DW h.



**Figure 1.** Variation in macromolecular composition and size of *E. coli* with growth rate. The equations in Table VI were fit to the data (lines in each plot). (a) Optical density per 10<sup>9</sup> cells as a function of specific growth rate (Bremer and Dennis, 1987). (b) Dry cell weight per 10<sup>9</sup> cells as a function of specific growth rate (Bremer and Dennis, 1987). (c) RNA and DNA (μmol per g DW) as a function of specific growth rate. Circles: RNA. Squares: DNA. Open symbols are from Brunschede et al. (1977). Closed symbols are from Bremer and Dennis (1987). (d) Protein (mmol per g DW) as a function of specific growth rate. Open symbols are from Brunschede et al. (1977). Closed symbols are from Bremer and Dennis (1987). (e) Median cell volume (μm³) as a function of specific growth rate (Bremer and Dennis, 1987; Eckert and Schaechter, 1965; Shehata and Marr, 1971). (f) Glycosyl units (mmol per g DW) as a function of dilution rate. Circles are from Neidhardt (1987). All other data are from Holme (1957).

lite that was not synthesized at the rate calculated from the correlations. The model was resolved with the new biomass composition. If the synthesis fluxes were consistent with biomass composition at that growth rate, then the case was considered biologically feasible. Otherwise, the iterative procedure was repeated until the appropriate biomass requirements could be balanced for a particular carbon and energy source and growth rate.

In addition to the objective function, optimization requires constraints. Equation (2) served as one set of constraints. This constraint states that the mass balance must be

satisfied and no accumulation of metabolites is allowed. Another set of constraints included the minimum  $(\alpha_i)$  and/or maximum  $(\beta_i)$  of allowable fluxes:

$$\alpha_i < v_i < \beta_i \quad i = 1, 2, 3, \dots, m.$$
 (4)

Because fluxes were defined as positive values only, the individual fluxes had a lower limit of zero and an upper limit of infinity. Reversible reactions were divided into two positive reactions in opposite directions. Experimental data for individual fluxes can be used as either lower and/or

Table VI. Equations for growth-rate dependent biomass composition.

| Component  | Correlation  | Reference                           |
|--|--|-------------------------------------|
| Optical density (OD <sub>460</sub> /10 <sup>9</sup> cells) | $-2.688 + 2.748 \cdot 2^{0.869 \; \mu}$  | Calculated from data in Figure 1(a) |
| Cell mass $(10^{-12} \text{ g DW})$                        | $-0.636 + 0.635 \cdot 2^{0.718 \mu}$   | Calculated from data in Figure 1(b) |
| RNA <sup>a,b</sup><br>(μmol NTPs/g DW)                     | $1139.5 - \frac{966.5}{\mu} \cdot 2^{-0.665/\mu}$                                    | Calculated from data in Figure 1(c) |
| DNA <sup>b</sup> (μmol dNTPs/g DW)                         | $\frac{100 \cdot \mu}{0.023} \left( 2^{(0.017 + 0.663)/\mu} - 2^{0.663/\mu} \right)$ | Calculated from data in Figure 1(c) |
| Protein <sup>b</sup> (mmol AAs/g DW)                       | $4.228 \cdot 2^{0.288/\mu}$  | Calculated from data in Figure 1(d) |
| Cell volume (µm³)  | $0.486 \cdot 2^{1.144\mu}$   | Calculated from data in Figure 1(e) |
| Cell radius (R) (µm)                                       | $0.293 \cdot 2^{0.41\mu}$  |                                     |
| Cell length ( $L$ ) ( $\mu$ m)                             | $2\cdot 2^{0.333\mu}$  | Donachie and Robinson,<br>1987      |
| Surface area <sup>c</sup> (µm <sup>2</sup> )               | $2\pi R (L-2R) + 4\pi R^2$   |                                     |
| Glycogen (glycosyl units)                                  | $10^3(1-2^{-3.24\cdot 10^4/\mu})$  | Calculated from data in Figure 1f   |

<sup>&</sup>lt;sup>a</sup>It has been speculated that the increase in the RNA content with growth rate is due to the larger fraction of stable RNA (rRNA, tRNA) necessary for the increase in protein production rate.

cThe radius and length are then used to calculate the surface area (*A*) of the cell, assuming that the cell is a cylinder with hemispherical caps. The lipid composition was calculated from the surface area. For a cell with a 40-min doubling time, phosphatidylethanolamine makes up 75% of these lipids, phosphatidylglycerol 18%, and cardiolipin 5% with only trace amounts of phosphatidylserine (Table II) (Ballesta and Schaechter, 1971; Neidhardt, 1987). The lipopolysaccharide content of the 40-min cell is 8.4 μmol/g DW and the peptidoglycan content is 8.4 μmol/g DW (Neidhardt, 1987). The fatty acid composition found in total lipids is presented in Bright–Gaertner and Proulx (1972), Kanemasa et al. (1967), Mavis and Vagelos (1972), and Neidhardt (1987). Because the lipid and fatty acid composition was available for only a cell with 40-min doubling time, it was assumed that the composition did not change with growth rate (data from Table III averaged).

upper constraints, or the flux value can be set to the value of the experimental data. Because a range of values is allowed for the individual flux constraints, variability in experimental data can be incorporated into the model. In this model the internal fluxes were not constrained because the goal was to predict the fluxes and compare them with experimental data.

The model was solved using the Simplex subroutines in the OSL package (IBM, 1992). This package has a number of important features: it has several alternative solution methods, it has been optimized for very large problems, and it enables the performance of a extensive sensitivity analysis

## **Sensitivity Analysis**

The basis matrix for each case (B) is a nonsingular subset of the stoichiometric matrix corresponding to the elements of v  $(v_B)$ , which uniquely solve the equation.

$$\mathbf{B} \cdot \mathbf{v_R} = \mathbf{b.} \tag{5}$$

Two types of sensitivity analysis can be performed on the solution. The first type determines what changes are required in the row and column bounds to cause the optimum

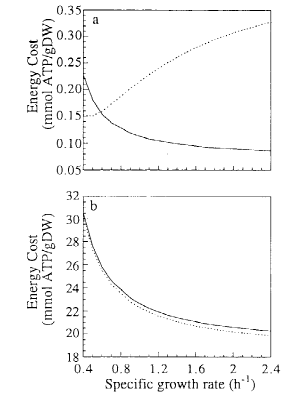
solution to occur with a different basis. That is, it examines how constrained a given reaction is in order to achieve a given objective using the same basis matrix. If a reaction is bounded tightly with respect to the basis matrix, it means that the basis matrix is very dependent on the flux value through that reaction and so is the simulation solution. The second type of analysis determines how large a change is required in the objective function coefficients to cause the optimum solution to occur with a different basis. This type of analysis gives rise to "reduced costs" (Luenberger, 1984).

#### **RESULTS**

#### Model Predictions Compared to Experimental Data

To determine the validity of the model, model predictions were compared to experimental data for two different growth conditions: aerobic growth on acetate plus glucose with a doubling time of 70 min and aerobic growth on acetate with a doubling time of 145 min (Walsh and Koshland, 1985b). The experimental flux measurements from Walsh and Koshland (1985b) were originally in units of

<sup>&</sup>lt;sup>b</sup>Although the amino acid and nucleotide compositions of protein and DNA and RNA, respectively, may vary under different growth conditions, the amino acid and nucleotide fractions were available for only a cell with a 40-min doubling time (Fig. 1); it was assumed that this composition was maintained at all growth rates.



**Figure 2.** Growth rate dependence of energy requirements. (a) RNA and DNA polymerization energy requirements (Neidhardt et al., 1990). (---) RNA, (—) DNA. (b) Total energy and protein polymerization requirements (Neidhardt et al., 1990). (---) Protein polymerization requirements. (—) Total energy requirements.

millimoles of substrate consumed per minute per liter of cytoplasmic volume. The experimental data was converted to units of millimoles of substrate consumed per hour per gram DW, assuming that cytoplasmic volume of a cell is 75% of the total volume, and the correlations for cell mass and cell volume (Table VI) were used to convert from a volume basis flux to a gram DW basis flux. The lower bound for the carbon dioxide secretion rate was set at the experimentally determined value, and the values predicted for oxygen uptake rates were in the range of experimental values (Andersen and von Meyenburg, 1980; Harrison and Loveless, 1971; Hempfling and Mainzer, 1975; Marr, 1991; Schulze and Lipe, 1964). The constraints and objective function for each case are presented in Table VII.

For aerobic growth on acetate plus glucose, the model predictions were very similar to experimental data for fluxes through the reactions of glycolysis and the TCA cycle (Fig. 3). The experimental glucose uptake rate for this case (7 mmol glucose/g DW h) for a doubling time of 70 min was taken from Herbert and Kornberg (1976), Schulze and Lipe (1964), and Tempest and Neijssel (1987). The average difference between the experimental values and the simulation results was 16%. In addition, the model was able to predict several levels of genetic regulation; for example, the glyoxylate shunt was not functional for growth on glucose and the flux through PEP carboxykinase was toward oxaloacetate (Walsh and Koshland, 1985b).

For aerobic growth on acetate only, the model predictions were also very similar to experimentally determined fluxes (Fig. 4). The average difference between the experimental values and the simulation results for this case was 17%. The model was able to predict that the glyoxylate shunt must be active during growth on acetate to generate sufficient precursors for macromolecule synthesis. The model was also able to predict the correct directions for reactions catalyzed by PEP carboxykinase and malic enzyme (Fraenkel, 1996).

The elemental compositions for bacteria were determined experimentally to be  $CN_{0.2}O_{0.27}$ ,  $CN_{0.25}O_{0.5}$ , and CN<sub>0.24</sub>O<sub>0.49</sub> (Blanch and Clark, 1996; Characklis and Marshall, 1989) and our biomass composition data predicted elemental balances of  $CN_{0.31}O_{0.2}$  for aerobic growth on acetate plus glucose and CN<sub>0.3</sub>O<sub>0.58</sub> for aerobic growth on acetate. Because the model only accounts for a proton balance across the cell membrane and there is no accounting of protons in the cell moving between metabolites, it is not possible to calculate an intracellular proton composition necessary for the protons in the elemental composition. The model predictions of  $Y_{x/s}$  (g g<sup>-1</sup>) = 0.53 (neglecting the acetate consumed) or  $Y_{x/s}$  (g g<sup>-1</sup>) = 0.52 (including acetate and assuming that the acetate has half of the carbon value of glucose) compares favorably with the experimentally determined yield coefficient of  $Y_{x/s}$  (g g<sup>-1</sup>) = 0.53 (Characklis and Marshall, 1989). If the acetate and glucose consumed are normalized per carbon, the model predicts  $Y_{x/s}$  [g(g carbon)<sup>-1</sup>] = 0.086 compared to experimental data of  $Y_{x/s}$  [g(g  $(carbon)^{-1}$ ] = 0.088 (Characklis and Marshall, 1989). The oxygen uptake rate for a 70-min doubling time was reported as 11-33 mmol of  $O_2/g$  DW h and the model predicted a flux of 12.6 mmol of O<sub>2</sub>/g DW h. For a 145-min doubling time the oxygen uptake was reported as 8-29 mmol of O<sub>2</sub>/g DW h (Andersen and von Meyenburg, 1980; Harrison and Loveless, 1971; Hempfling and Mainzer, 1975; Marr, 1991; Schulze and Lipe, 1964) and the model predicted a flux of 16 mmol of O<sub>2</sub>/g DW h. It was observed experimentally that the oxygen consumption rate decreases by 22% for growth on acetate plus glucose relative to growth on only acetate, even though the growth rate increases (Walsh and Koshland, 1985a), and the model predicted a 21% decrease in oxygen consumption.

Anaerobic growth on glucose with a 70-min doubling time was simulated to determine if the TCA cycle would branch as observed experimentally (Fig. 5) (Cronan and LaPorte, 1996; Nimmo, 1987). The glucose uptake rate was set to 3 times the aerobic rate (Smith and Neidhardt, 1983), and secretion rates for organic acids were set to experimentally determined values (Tempest and Neijssel, 1987). The model predicted the experimental observation that the TCA cycle is not a cycle but rather branches into a reductive pathway that produces succinyl-CoA and an oxidative pathway that produces  $\alpha$ -ketoglutarate (Cronan and LaPorte, 1996). The model was also able to predict that  $\alpha$ -ketoglutarate dehydrogenase is not expressed and that the glyoxylate shunt is closed during anaerobic growth. Unfortunately, there were no available experimental data to validate the flux values.

**Table VII.** Simulation parameters: Constraints on fluxes and criteria for maximization and minimization.

|                            | Lower bound | Upper<br>bound | Objective          |                         |
|----------------------------|-------------|----------------|--------------------|-------------------------|
| Growth condition           | (mmol/g     | g DW h)        | function criterion | Note                    |
| Aerobic, glucose + acetate |             |                |                    |                         |
| Glucose uptake             | 0           | 7              | Maximize           | Measureda               |
| Acetate uptake             | 0           | 11.94          | Maximize           | Measured <sup>b</sup>   |
| Carbon dioxide secretion   | 14          | ∞              | Minimize           |                         |
| Acetate secretion          | 0           | 11.14          | Maximize           | Measured <sup>b</sup>   |
| Precursor production rate  | μ dep       | endent         | Maximize           | Calculated <sup>c</sup> |
| Aerobic, acetate           |             |                |                    |                         |
| Acetate uptake             | 0           | 33.42          | Maximize           | Uptake <sup>d</sup>     |
| Carbon dioxide secretion   | 45          | $\infty$       | Minimize           | Measured <sup>b</sup>   |
| Precursor production rate  | μ dep       | endent         | Maximize           | Calculated <sup>c</sup> |
| Anaerobic, glucose         |             |                |                    |                         |
| Glucose uptake             | 0           | 21             | Maximize           | Measurede               |
| Oxygen uptake              | 0           | 0              |                    |                         |
| Carbon dioxide secretion   | 18.48       | $\infty$       | Minimize           | Measuredf               |
| Acetate secretion          | 0           | 7.67           | Maximize           | Measurede               |
| Lactate secretion          | 0           | 16.69          | Maximize           | Measurede               |
| Formate secretion          | 0           | 0.50           | Maximize           | Measurede               |
| Succinate secretion        | 0           | 2.25           | Maximize           | Measurede               |
| Ethanol secretion          | 0           | 10.46          | Maximize           | Measurede               |
| Precursor production rate  | μ dep       | endent         | Maximize           | Calculated <sup>c</sup> |

<sup>&</sup>lt;sup>a</sup>The values present in the literature are in the same range as the value used as a lower bound for the simulation (Herbert and Kornberg, 1976; Schulze and Lipe, 1964; Tempest and Neijssel, 1987).

#### Sensitivity to Biomass Composition

To determine the sensitivity of the solutions to the biomass composition, the three cases described above were simulated for incorrect biomass compositions: the cell composition corresponding to a 145-min doubling time (rather than the experimentally determined 70-min doubling time) was used to solve for the fluxes for aerobic growth on glucose and acetate and for anaerobic growth on glucose, and the cell composition corresponding to a 70-min doubling time (rather than the experimentally determined 145min doubling time) was used to solve for the fluxes for aerobic growth on acetate. The constraints and objective functions remained unchanged. For growth on glucose plus acetate, the correct biomass composition gave rise to an average error of 16% between the experimental data and model predictions for fluxes through the TCA cycle and glycolytic pathway, whereas the incorrect biomass composition increased the average error to 80%. Similarly, for growth on only acetate the correct biomass composition gave rise to an average error of 17%, whereas the incorrect biomass composition increased the average error to 32%.

The results for the two aerobic cases were significantly affected by changing the biomass compositions (Table

VIII). The flux distribution predicted by the model for aerobic growth on acetate with the incorrect biomass composition resulted in no flux through PEP carboxykinase, contrary to experimental observations (Walsh and Koshland, 1985b). For anaerobic growth on glucose, the incorrect biomass composition gave rise to incorrect branching of the TCA cycle with fumarase and malate dehydrogenase catalyzing reactions in directions that were not observed experimentally (Cronan and LaPorte, 1996). Further, the simulation suggested that there would be flux through the glyoxylate shunt, which was not observed experimentally (Nimmo, 1987). The sensitivity analysis of the anaerobic case was not performed because there were no available experimental data for comparisons.

#### Sensitivity Analysis on Flux Constraints

Sensitivity analysis was performed on the three cases to determine which reactions were the most constrained; that is, which reactions had the least amount of flexibility in flux values for which the solution will not change. The flexibility in the flux for a reaction is the range of flux values (represented as a percentage) that can occur without changing the basis matrix (Figs. 3–5).

<sup>&</sup>lt;sup>b</sup>Data from Walsh and Koshland (1985b).

<sup>&</sup>lt;sup>c</sup>Calculated from correlations in Table VI and information presented in Tables I–V.

<sup>&</sup>lt;sup>d</sup>The acetate uptake rate from Walsh and Koshland (1985b) was 41.4 mmol/g DW h but the model predicted an acetate uptake rate of 33.42 mmol/g DW h. Any amount of acetate provided in surplus resulted in acetate secretion.

<sup>&</sup>lt;sup>e</sup>Data from Smith and Neidhardt (1983).

<sup>&</sup>lt;sup>f</sup>Data from Bock and Sawers, (1996).

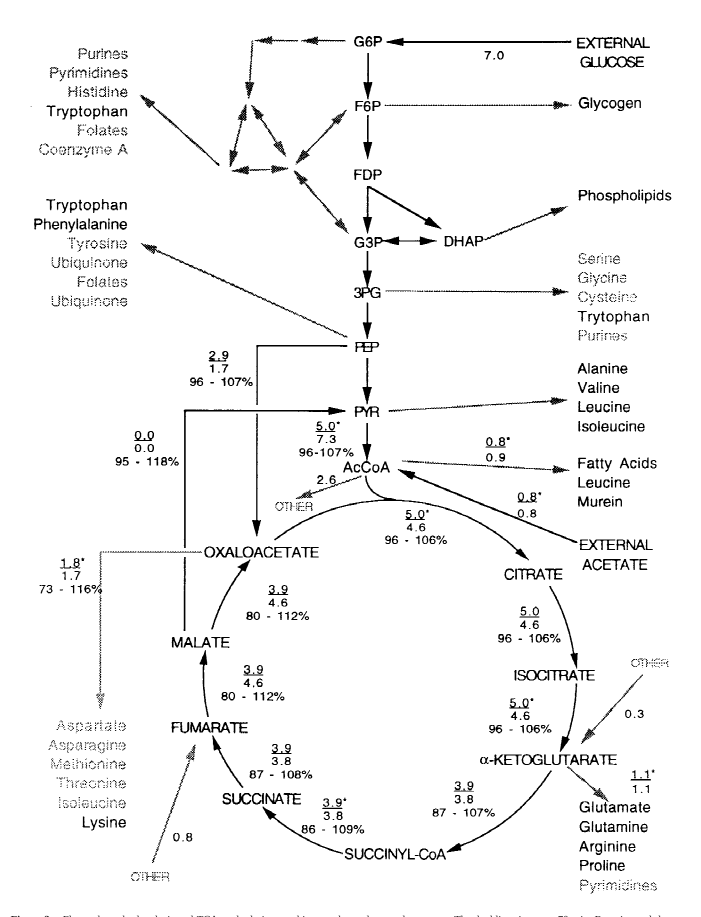
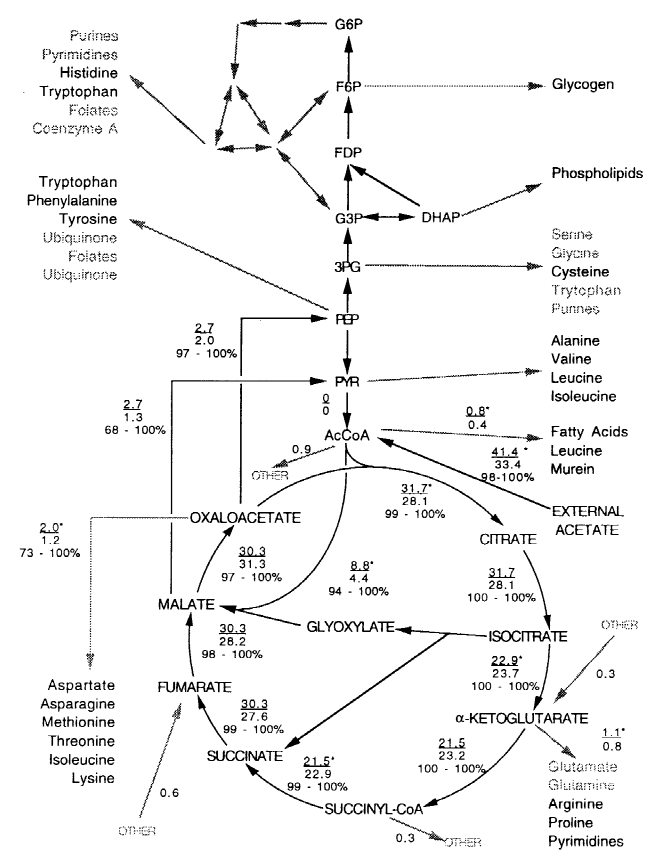


Figure 3. Fluxes through glycolysis and TCA cycle during aerobic growth on glucose plus acetate. The doubling time was 70 min. Experimental data are underlined (Walsh and Koshland, 1985b), the simulation results are the center values, and the flexibility of the reactions (as a percent of the simulation results) are on the bottom. Solid lines and words indicate highly constrained reactions and precursors. Stippled lines and words indicate less constrained reactions and precursors. The flux values measured experimentally are marked with an asterisk, and the flux values calculated from the experimental data using a simpler model are unmarked (Walsh and Koshland, 1985b). The "other" on the figure means synthesis of the metabolite due to the presence of other enzymes not depicted in the figure.



**Figure 4.** Fluxes through glycolysis and TCA cycle during aerobic growth on acetate. The doubling time was 145 min. Experimental data are underlined (Walsh and Koshland, 1985b), the simulation results are the center values, and the flexibility of the reactions (as a percent of the simulation results) are on the bottom. Solid lines and words indicate highly constrained reactions and precursors. Stippled lines and words indicate less constrained reactions and precursors. The flux values experimentally measured are marked with an asterisk, and the flux values calculated from the experimental data using a simpler model are unmarked (Walsh and Koshland, 1985b). The "other" on the figure means synthesis of the metabolite due to the presence of other enzymes not depicted in the figure.

**Table VIII.** Effect of biomass composition on the simulation results. The simulation results for fluxes through key reactions of TCA cycle and glycolysis for two different biomass compositions are compared to experimental data. The error is relative to experimental data (Walsh and Koshland, 1985b). Only the flux values presented in Table 8 were used to determine average error percentages.

|                               | Aero          | bic, glucose - | + acetate               | Aerobic acetate |                         |                |  |
|-------------------------------|---------------|----------------|-------------------------|-----------------|-------------------------|----------------|--|
|                               | Predictions   |                | Experiment <sup>a</sup> | Pred            | Experiment <sup>a</sup> |                |  |
|                               | $\tau_D = 70$ | $\tau_D = 145$ | $\tau_D = 70$           | $\tau_D = 70$   | $\tau_D = 145$          | $\tau_D = 145$ |  |
| PEP carboxykinase             | 0             | 0              | 0                       | 0<br>(100%)     | 2.0<br>(28%)            | 2.71           |  |
| PEP carboxylase               | 1.7<br>(41%)  | 0.8<br>(71%)   | 2.9                     | 0               | 0                       | 0              |  |
| Pyruvate dehydrogenase        | 7.3 (45%)     | 7.5 (50%)      | 5.0                     | 0               | 0                       | 0              |  |
| Citrate synthase              | 4.6 (8%)      | 1.0 (79%)      | 5.0                     | 25.1<br>(21%)   | 28.1<br>(11%)           | 31.7           |  |
| Aconitase                     | 4.6 (8%)      | 1.0<br>(79%)   | 5.0                     | 25.1<br>(21%)   | 28.1 (11%)              | 31.7           |  |
| Isocitrate dehydrogenase      | 4.6 (8%)      | 0.6<br>(87%)   | 5.0                     | 18.5            | 23.7                    | 22.9           |  |
| 2-Ketoglutarate dehydrogenase | 3.8 (3%)      | 0<br>(100%)    | 3.87                    | 17.7 (18%)      | 23.2 (8%)               | 21.5           |  |
| Succinate thiokinase          | 3.8 (3%)      | 0 (100%)       | 3.9                     | 17.2 (20%)      | 22.9 (7%)               | 21.5           |  |
| Succinate dehydrogenase       | 3.8 (3%)      | 0.4 (90%)      | 3.9                     | 24.3<br>(20%)   | 27.6<br>(9%)            | 30.3           |  |
| Fumarase                      | 4.6 (20%)     | 1.1 (72%)      | 39                      | 25.2<br>(17%)   | 28.2                    | 30.3           |  |
| Malate<br>dehydrogenase       | 4.6<br>(20%)  | 1.1 (72%)      | 3.87                    | 26.9<br>(11%)   | 31.3 (3%)               | 30.3           |  |
| Malic enzyme                  | 0             | 0              | 0                       | 4.9 (80%)       | 1.3 (52%)               | 2.7            |  |
| Isocitrate lyase              | 0             | 0.4            | 0                       | 6.6<br>(25%)    | 4.4<br>(50%)            | 8.8            |  |
| Overall error <sup>b</sup>    | 16%           | 80%            |                         | 32%             | 17%                     |                |  |

<sup>&</sup>lt;sup>a</sup>Data from Walsh and Koshland (1985b).

In general, the sensitivity analysis indicated that glycolysis was the most constrained pathway during aerobic and anaerobic growth on glucose, whereas the TCA cycle was the most constrained pathway during growth on acetate (Table IX). The rigid constraints on glycolysis during growth on glucose are due, in part, to the sensitivity of the leucine and valine biosynthetic pathways, which diverge from pyruvate. For growth on acetate, the rigid constraints on the TCA cycle are partly due to the rigidity in isocitrate dehydrogenase and in the cysteine and methionine synthesis rates. In the anaerobic case, the TCA cycle reactions had a lower bound of 59% of the flux value and no upper bound except for reactions catalyzed by fumarase and malate dehydrogenase, which had bounds of 92–142%; these are the two enzymes responsible for the TCA cycle branching under anaerobic growth conditions by changing direction of the reactions.

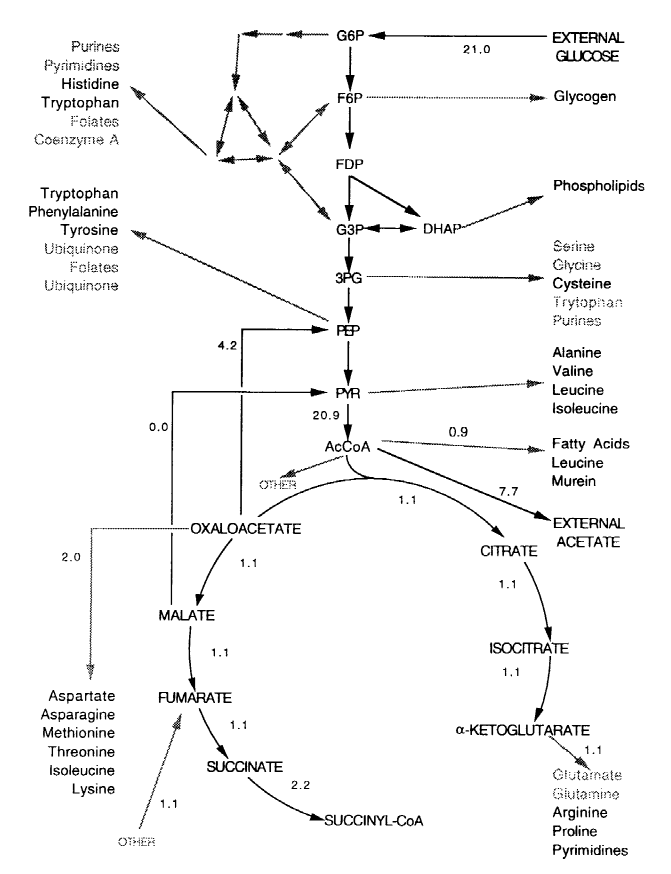
#### **DISCUSSION**

A steady-state, flux-based model was developed to study the distribution of mass and energy fluxes through the *E. coli* 

metabolic reaction network. The contributions of this model are threefold: a relatively complete data base of reactions from glucose to precursors, coenzymes, and prosthetic groups was used; correlations for experimentally observed changes in biomass composition with growth rate were included; and energy requirements for growth were based on mechanistic requirements rather than on a lumped maintenance energy requirement.

The model incorporated 153 reversible and 147 irreversible reactions using 289 metabolites. The relatively complete nature of the reactions will allow one to examine the effect of mutations in specific genes on fluxes through pathways by constraining fluxes through relevant enzymes. It will also allow more flexibility in choosing which metabolites or pathways might be most amenable to alterations to engineer metabolism. In genetically manipulating cells for the production of a desired compound or protein, the growth rate of the cell may change significantly. Therefore, incorporation of growth-rate dependent biomass composition as well as energy requirements in such a model would allow one to accurately simulate the effect of the growth rate changes on the distribution of resources throughout the cell.

<sup>&</sup>lt;sup>b</sup>The error is calculated as (|experiment - prediction|)/experiment.



**Figure 5.** Fluxes through glycolysis and TCA cycle during anaerobic growth on glucose. The doubling time was 70 min. Solid lines and words indicate highly constrained reactions and precursors. Stippled lines and words indicate less constrained reactions and precursors. The ''other'' on the figure means synthesis of the metabolite due to the presence of other enzymes not depicted in the figure.

Experimental data were used to develop correlations for biomass composition at different growth rates. The biomass sensitivity analysis indicates the importance of incorporating changes in biomass composition with growth rate. For growth on acetate plus glucose, the biomass composition greatly affected the fluxes. Using the correct composition, the predicted fluxes differed from experimental measurements by 16%. Using an incorrect composition, the predicted fluxes differed from experimental measurements by 80%. For growth on acetate the predicted fluxes using a correct composition differed from experimental measurements by 17%, and for the incorrect composition the predicted fluxes differed from experimental measurements by 32%.

Maintenance energy terms have often been incorporated into models to balance energy requirements that could not be accounted for through biomass synthesis requirements (Pirt, 1965, 1982). However, the use of the maintenance energy term does not account mechanistically for this energy drain. To improve the predictive power of stoichiometric models, we accounted for the energy drain using a mechanistic approach and correlated this with growth rate. Besides incorporating the obvious energy requirements, such as those for polymerization of biopolymers, we also incorporated energy requirements, such as those for proofreading of DNA and protein, RNA processing, and proton leakage across membranes. Because the composition of the cell changes with growth rate, the energy requirements were scaled with cell composition. In a similar manner, proton leakage was scaled with the cell's surface area, which increases with growth rate.

There was close agreement between the predicted and experimentally determined flux values, and the solutions agreed with observed regulation under the different growth conditions. The model was able to predict the opening and closing of the glyoxylate shunt in the presence and absence of acetate as the sole carbon source. The O<sub>2</sub> uptake rate under aerobic growth conditions predicted by the model agreed with experimental data (Andersen and von Meyenburg, 1980; Harrison and Loveless, 1971; Hempfling and

Mainzer, 1975; Marr, 1991; Schulze and Lipe, 1964). The model also predicted the branching of the TCA cycle with no flux through  $\alpha$ -ketoglutarate dehydrogenase under anaerobic growth conditions. The largest errors occurred at the branch points of glycolysis and the TCA cycle, where there were drains for the synthesis of cellular constituents. Because the data used to develop the correlations for the biomass components came from many different experiments conducted under various growth conditions using various strains of *E. coli*, differences in biomass composition could be responsible for these discrepancies. A more complete set of experimental data is required to improve model predictions.

Sensitivity analysis was also performed to determine how changes in the fluxes would affect the basis matrix that was used to arrive at a given solution. This type of analysis indicates how much the flux through a given reaction can change and still allow the model to arrive at the optimal solution using the same basis matrix. This analysis indicates that during growth on glucose the reactions of glycolysis are the most constrained, with the TCA cycle and the pentose phosphate shunt reactions close behind. In contrast, the reactions of the TCA cycle are the most highly constrained during growth on acetate. Because the glycolysis and the TCA cycle reactions are involved in both energy and precursor production, one would expect them to be the most highly constrained reactions.

This metabolic model should be a useful tool for studying the effects of reengineering pathways. It can provide information about how the overall flux distribution will be affected if an organism is forced to synthesize a product or to degrade a pollutant. The model can also elucidate cases under which it will not be possible to satisfy growth demands and secrete a product of interest at a desired level. The sensitivity analysis can be used to study the stiffness of the solutions and the regions where bottlenecks may form under certain growth conditions. The model shows a high degree of sensitivity to the biomass information, and therefore the dependence of biomass composition on growth rates is an important aspect of a flux-based metabolic model.

Table IX. Sensitivity analysis of basis matrix with respect to flux constraints.

|                                | Allowable fluxes <sup>a</sup> (% solution value) |                          |                          |  |  |  |
|--------------------------------|--|--------------------------|--------------------------|--|--|--|
| Pathway/enzyme                 | Glucose + acetate, aerobic                       | Acetate, aerobic         | Glucose, anaerobic       |  |  |  |
| Glycolysis                     | 94–103   | 84–100                   | 100–101                  |  |  |  |
| Pentose phosphate              | 95-109   | 81–100                   | 94–101                   |  |  |  |
| TCA cycle                      | 90–106   | 100-100                  | 92-142                   |  |  |  |
| Carbon dioxide secretion       | 68–108   | 50-100                   | 72–106                   |  |  |  |
| Ammonia uptake                 | 85–109   | 85-100                   | 87–103                   |  |  |  |
| Oxygen uptake                  | 87–108   | 96-100                   |                          |  |  |  |
| Acetate secretion              | 48–116   |                          | 46–117                   |  |  |  |
| Precursors tightly constrained | Phosphatidylethanolamine                         | Phosphatidylethanolamine | Phosphatidylethanolamine |  |  |  |
|                                | Arginine   | Cardiolipin              | •                        |  |  |  |
|                                | Lysine   | Cysteine                 |                          |  |  |  |
|                                | Leucine  | Methionine               |                          |  |  |  |
|                                | Valine   |                          |                          |  |  |  |

<sup>&</sup>lt;sup>a</sup>Values refer to the amount the flux value can change without changing the basis matrix. The most tightly bounded reaction in a particular pathway determines the bounds on that pathway.

# APPENDIX A: METABOLIC PATHWAYS IN STOICHIOMETRIC MATRIX GENE NAMES ARE SUPPLIED WHERE POSSIBLE.

| Enzyme                                   | Gene       | Pathway   | Reference  |
|--|------------|---|--|
| Glycolysis                               |            |   |  |
| Phosphotransferase system                | ptsI, ptsH | $GLC + PEP \rightarrow PYR + G6P$                     | Fraenkel, 1996; Postma et al., 1996                |
| Phosphoglucose isomerase                 | pgi        | $G6P \leftrightarrow F6P$                             | Fraenkel, 1996                                     |
| Phosphofructokinase                      | pfk        | $F6P + ATP \leftrightarrow ADP + F16P$                | Fraenkel, 1996                                     |
| Fructose-1,6-bisphosphate aldolase       | fbp        | $F16P + PI \rightarrow F6P$                           | Fraenkel, 1996                                     |
| Fructose-1,6-bisphosphatase              | fba        | $F16P \leftrightarrow T3P1 + T3P2$                    | Fraenkel, 1996                                     |
| Triosphosphate isomerase                 | tpi        | $T3P1 \leftrightarrow T3P2$                           | Fraenkel, 1996                                     |
| Glyceraldehyde-3-phosphate dehydrogenase | gap        | $T3P1 + PI + NAD \leftrightarrow NADH + 13P2DG$       | Fraenkel, 1996                                     |
| Phosphoglycerate kinase                  | pgk        | $13P2DG + ADP \leftrightarrow ATP + 3PDGL$            | Fraenkel, 1996                                     |
| Phosphoglycerate mutase                  | gpm        | $3PDGL \leftrightarrow 2PDGL$                         | Fraenkel, 1996                                     |
| Enolase                                  | eno        | $2PDGL \leftrightarrow PEP$                           | Fraenkel, 1996                                     |
| Pyruvate kinase                          | pyk        | $PEP + ADP \rightarrow ATP + PYR$                     | Fraenkel, 1996                                     |
| PEP carboxykinase                        | pck        | $OA + ATP \leftrightarrow PEP + CO2 + ADP$            | Fraenkel, 1996                                     |
| PEP carboxylase                          | ppc        | $PEP + CO2 \rightarrow OA + PI$                       | Fraenkel, 1996                                     |
| Pyruvate dehydrogenase                   | pdh        | $PYR + COA + NAD \rightarrow NADH + CO2 + ACCOA$      | Fraenkel, 1996                                     |
| PEP synthase                             | pps        | $PYR + ATP \rightarrow PEP + AMP + PI$                | Fraenkel, 1996                                     |
| Pentose phosphate shunt                  |            |   |  |
| Glucose-6-phosphate dehydrogenase        | zwf        | $G6P + NADP \leftrightarrow D6PGL + NADPH$            | Fraenkel, 1996                                     |
| 6-Phosphogluconolactonase                | pgl        | $D6PGL \rightarrow D6PGC$                             | Fraenkel, 1996                                     |
| 6-Phosphogluconate dehydrogenase         | gnd        | $D6PGC + NADP \leftrightarrow NADPH + CO2 + RL5P$     | Fraenkel, 1996                                     |
| Ribose-5-phosphate isomerase             | rpi        | $RL5P \leftrightarrow R5P$                            | Fraenkel, 1996                                     |
| Ribose-5-phosphate epimerase             | rpe        | $RL5P \leftrightarrow X5P$                            | Fraenkel, 1996                                     |
| Transketolase 1                          | tktA       | $R5P + X5P \leftrightarrow T3P1 + S7P$                | Fraenkel, 1996                                     |
| Transaldolase                            | tal        | $T3P1 + S7P \leftrightarrow E4P + F6P$                | Fraenkel, 1996                                     |
| Transketolase 2                          | tktB       | $X5P + E4P \leftrightarrow F6P + T3P1$                | Fraenkel, 1996                                     |
| 6-Phosphogluconate dehydrase             | edd        | $D6PGC \rightarrow 2K3D6PG$                           | Fraenkel, 1996                                     |
| 2-Keto-3-deoxy-6-phospho-                | eda        | $2K3D6PG \rightarrow T3P1 + PYR$                      | Fraenkel, 1996                                     |
| gluconate aldolase                       |            |   |  |
| Glycogen metabolism                      |            |   |  |
| Phosphogluconomutase                     | pgm        | $G6P \leftrightarrow G1P$                             | Fraenkel, 1996                                     |
| Glycogen synthase                        | glgA       | $G1P + ATP \rightarrow ADP + PPI + GLYCOGEN$          | Preiss, 1996                                       |
| Glycogen phosphorylase                   | glgP       | $GLYCOGEN + PI \rightarrow G1P$                       | Preiss, 1996                                       |
| Dissimilation of pyruvate                |            |   |  |
| Lactate dehydrogenase                    | ldh        | $PYR + NADH \leftrightarrow NAD + LAC$                | Bock and Sawers, 1996; Kessler and Knappe, 1996    |
| Alcohol dehydrogenase                    | adh        | $ACAL + NADH \leftrightarrow ETHANOL + NAD$           | Bock and Sawers, 1996; Kessler and Knappe, 1996    |
| Acetaldehyde dehydrogenase               | adh        | $AC + NADH \leftrightarrow NAD + ACAL$                | Bock and Sawers, 1996; Kessler and Knappe, 1996    |
| Pyruvate formate lyase                   | pfl        | $PYR + COA \rightarrow ACCOA + FORMATE$               | Bock and Sawers, 1996; Kessler<br>and Knappe, 1996 |
| Phosphotransacetylase                    | pta        | $ACCOA + PI \leftrightarrow ACTP + COA$               | Bock and Sawers, 1996; Kessler<br>and Knappe, 1996 |
| Acetate kinase                           | ackA       | $ACTP + ADP \leftrightarrow ATP + AC$                 | Bock and Sawers, 1996; Kessler<br>and Knappe, 1996 |
| Formate hydrogen lyase                   | fhl        | $FORMATE \rightarrow CO2$                             | Bock and Sawers, 1996; Kessler<br>and Knappe, 1996 |
| TCA cycle and glyoxylate bypass          |            |   | ана кнарре, 1990                                   |
| Citrate synthase                         | gltA       | $ACCOA + OA \leftrightarrow COA + CIT$                | Cronan and LaPorte, 1996                           |
| Aconitase                                | acn        | $CIT \leftrightarrow ICIT$                            | Cronan and LaPorte, 1996                           |
| Isocitrate dehydrogenase                 | idh        | $ICIT + NAD \leftrightarrow CO2 + NADH + AKG$         | Cronan and LaPorte, 1996                           |
| 2-Ketoglutarate dehydrogenase            | sucAB      | $AKG + NAD + COA \leftrightarrow CO2 + NADH + SUCCOA$ | Cronan and LaPorte, 1996                           |
| Succinate thiokinase                     | sucCD      | $SUCCOA + GDP + PI \leftrightarrow GTP + COA + SUCC$  | Cronan and LaPorte, 1996                           |
| Succinate dehydrogenase                  | sdhABCD    | $SUCC + FAD \rightarrow FADH2 + FUM$                  | Cronan and LaPorte, 1996                           |
| Fumurate reductase                       | frdABCD    | $FUM + FADH2 \rightarrow SUCC + FAD$                  | Cronan and LaPorte, 1996                           |
| Fumarase                                 | fumAB      | $FUM \leftrightarrow MAL$                             | Cronan and LaPorte, 1996                           |
| Malate dehydrogenase                     | mdh        | $MAL + NAD \leftrightarrow NADH + OA$                 | Cronan and LaPorte, 1996                           |
| Malic enzyme                             | mez        | $MAL + NADP \rightarrow CO2 + NADPH + PYR$            | Cronan and LaPorte, 1996                           |
| Malic enzyme                             | mez,       | $MAL + NAD \rightarrow CO2 + NADH + PYR$              | Cronan and LaPorte, 1996                           |

| Enzyme                                     | Gene  | Pathway   | Reference   |
|--|-------|---|---|
| Isocitrate lyase                           | aceA  | $ICIT \rightarrow GLX + SUCC$   | Cronan and LaPorte, 1996                            |
| Malate synthase                            | aceB  | $ACCOA + GLX \rightarrow COA + MAL$   | Cronan and LaPorte, 1996                            |
| Respiration                                |       |   |   |
| NADH dehydrogenase II                      | ndh   | $NADH + Q \rightarrow NAD + QH2$  | Gennis and Stewart, 1996                            |
| NADH dehydrogenase I                       | ndh   | $NADH + Q \rightarrow NAD + QH2 + 4 HEXT$   | Gennis and Stewart, 1996                            |
| Formate dehydrogenase                      |       | FORMATE + Q $\rightarrow$ 2 HEXT + QH2 + CO2  | Gennis and Stewart, 1996                            |
| Cytochrome oxidase bo3                     |       | $QH2 + 1/2 O2 \rightarrow Q + 4 HEXT$   | Gennis and Stewart, 1996                            |
| Cytochrome oxidase bd                      |       | $QH2 + 1/2 O2 \rightarrow Q + 2 HEXT$   | Gennis and Stewart, 1996                            |
| Succinate dehydrogenase complex            |       | $FADH2 + Q \leftrightarrow FAD + QH2$   | Gennis and Stewart, 1996                            |
| ATP synthesis                              |       | 100 100 DY 0.110110   |   |
| F0F1-ATPase                                | unc   | $ATP \leftrightarrow ADP + PI + 3 HEXT$   | Harold and Maloney, 1996                            |
| Biosynthesis of aspartate                  |       | 0.4 GLU 40D 41/G  | D : 1006  |
| Aspartate transaminase                     | aspC  | $OA + GLU \leftrightarrow ASP + AKG$  | Reitzer, 1996                                       |
| Biosynthesis of asparagine                 | -     | 1.65 1.75 GY 1. GY 1. 1.65 DD   | B 1 4005  |
| Glutamine-dependent asparagine synthetase  | asnB  | $ASP + ATP + GLN \rightarrow GLU + ASN + AMP + PPI$   | Reitzer, 1996                                       |
| Ammonia-dependent asparagine synthetase    | asnA  | $ASP + ATP + NH3 \rightarrow ASN + AMP + PPI$   | Reitzer, 1996                                       |
| Biosynthesis of glutamate                  |       |   |   |
| Glutamate dehydrogenase                    | gdhA  | $AKG + NH3 + NADPH \rightarrow GLU + NADP$  | Reitzer, 1996                                       |
| Glutamine synthatase                       | glnA  | $GLU + NH3 + ATP \rightarrow GLN + ADP + PI$  | Reitzer, 1996                                       |
| Glutamate synthase                         | gltBD | $AKG + GLN + NADPH \rightarrow NADP + 2 GLU$  | Reitzer, 1996                                       |
| Biosynthesis of alanine                    |       |   |   |
| Glutamic-pyruvic transaminase              |       | $PYR + GLU \leftrightarrow AKG + ALA$   | Reitzer, 1996                                       |
| Biosynthesis of arginine, putrescine,      |       |   |   |
| and spermidine                             |       |   |   |
| N-Acetylglutamate synthase                 | argA  | $GLU + ACCOA \rightarrow COA + NAGLU$   | Glansdorf, 1996                                     |
| N-Acetylglutamate kinase                   | argB  | $NAGLU + ATP \rightarrow ADP + NAGLUYP$   | Glansdorf, 1996                                     |
| N-Acetylglutamate phosphate reductase      | argC  | $\begin{aligned} \text{NAGLUYP} + \text{NADPH} &\leftrightarrow \text{NADP} + \text{PI} \\ + \text{NAGLUSAL} \end{aligned}$   | Glansdorf, 1996                                     |
| Acetylornithine aminotransferase           | argD  | $NAGLUSAL + GLU \leftrightarrow AKG + NAARON$   | Glansdorf, 1996                                     |
| N-Acetylornithinase                        | argE  | $NAARON \rightarrow AC + ORN$   | Glansdorf, 1996                                     |
| Carbamoyl phosphate synthetase             | carAB | $GLN + 2 ATP + CO2 \rightarrow GLU + CAP + 2 ADP + PI$  | Glansdorf, 1996                                     |
| Ornithine carbamoyl transferase            | argFI | $ORN + CAP \leftrightarrow CITR + PI$   | Glansdorf, 1996                                     |
| Argininosuccinate synthase                 | argG  | $CITR + ASP + ATP \leftrightarrow AMP + PPI + ARGSUCC$  | Glansdorf, 1996                                     |
| Argininosuccinase                          | argH  | $ARGSUCC \leftrightarrow FUM + ARG$   | Glansdorf, 1996                                     |
| Ornithine decarboxylase                    | speC  | $ORN \rightarrow PTRSC + CO2$   | Glansdorf, 1996                                     |
| Spermidine synthase                        | spE   | $PTRSC + DSAM \rightarrow SPRMD + 5MTA$   | Glansdorf, 1996                                     |
| Adenosylmethionine decarboxylase           | speD  | $SAM \leftrightarrow DSAM + CO2$  | Glansdorf, 1996; Greene, 1996;<br>Karp et al., 1996 |
| Unknown pathway                            |       | $5MTA \rightarrow ADN + MET$  | Greene, 1996  |
| Agmatine decarboxylase                     | speA  | $ARG \rightarrow CO2 + AGM$   | Glansdorf, 1996                                     |
| Agmatine ureohydrolase                     | speB  | $AGM \rightarrow UREA + PTRSC$  | Glansdorf, 1996                                     |
| Biosynthesis of proline                    |       |   |   |
| Glutamyl kinase                            | proB  | $GLU + ATP \rightarrow ADP + GLUP$  | Leisinger, 1996                                     |
| Glutamate-5-semialdehyde dehydrogenase     | proA  | $\operatorname{GLUP} + \operatorname{NADPH} \leftrightarrow \operatorname{NADP} + \operatorname{PI} + \operatorname{GLUGSAL}$ | Leisinger, 1996                                     |
| Pyrroline-5-carboxylate reductase          | proC  | $GLUGSAL + NADPH \leftrightarrow PRO + NADP$  | Leisinger, 1996                                     |
| Ornithine oxoacid transaminase             |       | $ORN + AKG \leftrightarrow GLU + GLUGSAL$   | Leisinger, 1996                                     |
| Biosynthesis of branched-chain amino acids |       |   |   |
| Isopropylmalate synthase                   | leuA  | $ACCOA + OIVAL \rightarrow COA + CBHCAP$  | Umbarger, 1996                                      |
| Isopropylmalate isomerase                  | leuCD | $CBHCAP \leftrightarrow IPPMAL$   | Umbarger, 1996                                      |
| 3-Isopropylmalate dehydrogenase            | leuB  | $IPPMAL + NAD \rightarrow NADH + OICAP + CO2$   | Umbarger, 1996                                      |
| Transaminase C                             | ilvE  | $OICAP + GLU \rightarrow AKG + LEU$   | Umbarger, 1996                                      |
| Acetyohydroxy acid synthase                | ilvB  | $2 \text{ PYR} \rightarrow \text{CO2} + \text{ACLAC}$   | Umbarger, 1996                                      |
| Acetohydroxy acid isomeroreductase         | ilvC  | $ACLAC + NADPH \rightarrow NADP + DHVAL$  | Umbarger, 1996                                      |
| Dihydroxy acid dehydratase                 | ilvD  | $DHVAL \rightarrow OIVAL$   | Umbarger, 1996                                      |
| Transaminase C                             | ilvE  | $OIVAL + GLU \leftrightarrow AKG + VAL$   | Umbarger, 1996                                      |
| Threonine deaminase                        | ilvA  | $THR \rightarrow NH3 + OBUT$  | Umbarger, 1996                                      |
| Acetohydroxy acid synthase                 | ilvB  | $OBUT + PYR \rightarrow ABUT + CO2$   | Umbarger, 1996                                      |

| Enzyme   | Gene           | Pathway  | Reference  |
|--|----------------|--|--|
| Acetohydroxy acid isomeroreductase                         | ilvC           | $ABUT + NADPH \rightarrow NADP + DHMVA$  | Umbarger, 1996   |
| Dihydroxy acid dehydratase                                 | ilvD           | $DHMVA \rightarrow OMVAL$  | Umbarger, 1996   |
| Transaminase B   | ilvE           | $OMVAL + GLU \leftrightarrow AKG + ILE$  | Umbarger, 1996   |
| Amino acid oxidase   |                | $O2 + ILE \rightarrow OMVAL + NH3$   | Umbarger, 1996   |
| Biosynthesis of aromatic amino acids                       |                |  |  |
| 3-Deoxy-D-arabinoheptulosonate-7-<br>phosphate synthase    | aroFGH         | $E4P + PEP \rightarrow PI + 3DDAH7P$   | Pittard, 1996  |
| 3-Dehydroquinate synthase                                  | aroB           | $3DDAH7P \rightarrow DQT + PI$   | Pittard, 1996  |
| 3-Dehydroquinate dehydratase                               | aroD           | DQT ↔ DHSK   | Pittard, 1996  |
| Shikimate dehydrogenase                                    | aroE           | $DHSK + NADPH \leftrightarrow SME + NADP$  | Pittard, 1996  |
| Shikimate kinase   | aroKL          | $SME + ATP \rightarrow ADP + SME5P$  | Pittard, 1996  |
| 5-Enolpyruvoylshikimate-3-phosphate synthase               | aroA           | $SME5P + PEP \leftrightarrow 3PSME + PI$   | Pittard, 1996  |
| Chorismate synthase  | aroC           | $3PSME \rightarrow PI + CHOR$  | Pittard, 1996  |
| Chorismate mutase  | pheA           | $CHOR \rightarrow PHEN$  | Pittard, 1996  |
| Prephenate dehydratase                                     | pheA           | $PHEN \rightarrow CO2 + PHPYR$   | Pittard, 1996  |
| Phenylalanine aminotransferase                             | tyrB           | $PHPYR + GLU \leftrightarrow AKG + PHE$  | Pittard, 1996  |
| Prephanate dehydrogenase                                   | tyrA           | PHEN + NADP $\rightarrow$ HPHPYR + CO2 + NADPH   | Pittard, 1996  |
| Tyrosine aminotransferase                                  | tyrB           | $\begin{array}{c} \text{HPHPYR} + \text{GLU} \leftrightarrow \text{AKG} + \text{TYR} \\ \end{array}$   | Pittard, 1996  |
| Anthranilate synthase                                      | trpDE          | CHOR + GLN $\rightarrow$ GLU + PYR + AN  | Pittard, 1996  |
| Anthranilate phosphoribosyl transferase                    | trpD           | $AN + PRPP \rightarrow PPI + NPRAN$  | Pittard, 1996  |
| Phosphoribosyl anthranilate isomerase                      | trpC           | $NPRAN \rightarrow CPAD5P$   | Pittard, 1996  |
| Indoleglycerol phosphate synthetase                        | trpC           | $CPAD5P \rightarrow CO2 + IGP$   | Pittard, 1996  |
| Tryptophan synthetase                                      | trpAB          | $IGP + SER \rightarrow T3P1 + TRP$   | Pittard, 1996  |
| Biosynthesis of histidine synthesis                        | v.p.12         | 101 - 1021 - 110   | 111111111111111111111111111111111111111                          |
| Phosphoribosyl pyrophosphate synthetase                    | prs            | $R5P + ATP \leftrightarrow PRPP + AMP$   | Zalkin and Nygaard, 1996   |
| Phosphoribosyl pyrophosphate                               | hisG           | $PRPP + ATP \rightarrow PPI + PRBATP$  | Winkler, 1996  |
| PR-ATP pyrophosphohydrolase                                | hisI           | $PRBATP \rightarrow PPI + PRBAMP$  | Winkler, 1996  |
| PR-AMP cyclohydrolase                                      | hisI           | PRBAMP → PRFP  | Winkler, 1996  |
| 5-ProFAR isomerase   | hisA           | PRFP → PRLP  | Winkler, 1996  |
| Imidazoleglycerol phosphate synthase                       | hisFH          | $PRLP + GLN \rightarrow GLU + AICAR + DIMGP$   | Winkler, 1996  |
| IGP dehydratase  | hisB           | DIMGP → IMACP  | Winkler, 1996  |
| L-Histidinol phosphate                                     | hisC           | $IMACP + GLU \rightarrow AKG + HISOLP$   | Winkler, 1996  |
| aminotransferase   | 7775           | marer / oze / me / moozi   | ,, mmer, 1330  |
| Hol- <i>P</i> -phosphatase                                 | hisB           | $HISOLP \rightarrow PI + HISOL$  | Winkler, 1996  |
| Hol dehydrogenase  | hisD           | HISOL + 2 NAD $\rightarrow$ 2 NADH + HIS   | Winkler, 1996  |
| Biosynthesis of serine, glycine, and 1-carbon units        | nisb           | INDOE   E IVID   / E IVIDII   IND  | Winard, 1990   |
| 3-Phosphoglycerate dehydrogenase                           | serA           | $3PDGL + NAD \rightarrow NADH + PHP$   | Stauffer, 1996   |
| Phosphoserine transaminase                                 | serC           | $PHP + GLU \rightarrow AKG + 3PSER$  | Stauffer, 1996   |
| Phosphoserine phosphatase                                  | serB           | $3PSER \rightarrow PI + SER$   | Stauffer, 1996   |
| Serine hydroxymethyltransferase<br>Glycine cleavage system | glyA<br>gcvHTP | $\begin{aligned} & \text{GLY} + \text{METTHF} &\leftrightarrow \text{THF} + \text{SER} \\ & \text{GLY} + \text{THF} + \text{NAD} &\to \text{METTHF} + \text{NADH} \end{aligned}$ | Matthews, 1996; Stauffer, 1996<br>Matthews, 1996; Stauffer, 1996 |
|  |                | + CO2 + NH3  | 10.5   |
| Threonine dehydrogenase                                    | thd            | $THR + NAD \leftrightarrow AABK + NADH$  | Matthews, 1996   |
| Amino-b-ketobutyrase                                       | kbl            | $AABK + COA \leftrightarrow GLY + ACCOA$   | Matthews, 1996   |
| Formate dehydrogenase                                      |                | $FORMATE + NAD \rightarrow NADH + CO2$   | Kadner, 1996   |
| Formate THF ligase   |                | THF + FORMATE + ATP $\rightarrow$ ADP + PI + FTHF  | Michal, 1993   |
| Formyl THF deformylase                                     |                | $FTHF \rightarrow FORMATE + THF$   | Michal, 1993   |
| Dihydrofolate reductase (STEP 2)                           |                | $DHF + NADPH \leftrightarrow NADP + THF$   | Michal, 1993   |
| Biosynthesis of cysteine                                   |                |  |  |
| ATP sulfhydrolase  | cysDN          | $H2SO4 + ATP \rightarrow PPI + APS$  | Kredich, 1996  |
| ATS kinase   | cysC           | $APS + ATP \rightarrow ADP + PAPS$   | Kredich, 1996  |
| PAPS sulfotransferase                                      | cysH           | $PAPS + NADPH \rightarrow NADP + H2SO3 + PAP$  | Kredich, 1996  |
| NAHPH-sulfite reductase                                    | cysGIJ         | $H2SO3 + 3 NADPH \leftrightarrow 3 NADP + H2S$   | Kredich, 1996  |
| Serine transacetylase                                      | cysE           | $SER + ACCOA \leftrightarrow COA + ASER$   | Kredich, 1996  |
| O-Acetylserine (thiol)-lyase                               | cysKM          | $ASER + H2S \rightarrow AC + CYS$  | Kredich, 1996  |
| Sulfotransferase   |                | $PAP + H2SO3 \leftrightarrow PAPS$   |  |
| Adenylyl sulfate kinase                                    |                | $PAPS + ADP \rightarrow ATP + APS$   |  |

| Enzyme Gene Pathway  |                        | Reference   |  |
|--|------------------------|---|--|
| Biosynthesis of threonine and lysine                           |                        |   |  |
| Aspartate kinase   | thrA                   | $ASP + ATP \leftrightarrow ADP + BASP$  | Patte, 1996  |
| Aspartate semialdehyde   | asd                    | $BASP + NADPH \leftrightarrow NADP + PI + ASPSA$                                | Patte, 1996  |
| dehydrogenase  |                        |   |  |
| Homoserine dehydrogenase                                       | thrA                   | $ASPSA + NADPH \leftrightarrow NADP + HSER$                                     | Patte, 1996  |
| Homoserine kinase  | thrB                   | $HSER + ATP \rightarrow ADP + PHSER$  | Patte, 1996  |
| Threonine synthase   | thrC                   | $PHSER \rightarrow PI + THR$  | Patte, 1996  |
| Dihydrodipicolinate synthase                                   | dapA                   | $ASPSA + PYR \rightarrow D23PIC$  | Patte, 1996  |
| Dihydrodipicolinate reductase                                  | dapB                   | D23PIC + NADPH $\rightarrow$ NADP + PIP26DX                                     | Patte, 1996  |
| Tetrahydrodipicolinate succinylase<br>Succinyl diaminopimelate | dapD<br>dapC           | $PIP26DX + SUCCOA \rightarrow COA + NS2A6O$ $NS2A6O + GLU(A)AVG + NS26DB$       | Patte, 1996<br>Patte, 1996                           |
| aminotransferase   | аарс                   | $NS2A6O + GLU \leftrightarrow AKG + NS26DP$                                     | ratie, 1990  |
| Succinyl diaminopimelate                                       | dapE                   | NS26DP → SUCC + D26PIM  | Patte, 1996  |
| desuccinylase  | ширЕ                   | NS20DI -> SUCC + D20I IWI   | 1 atte, 1990   |
| Diaminopimelate epimerase                                      | dapF                   | $D26PIM \leftrightarrow MDAP$   | Patte, 1996  |
| Diaminopimelate decarboxylase                                  | lysA                   | $MDAP \rightarrow CO2 + LYS$  | Patte, 1996  |
| Biosynthesis of methionine                                     | 1 9 3 2 1              | MDM 7 CO2 + E15   | 1 atte, 1990   |
| Homoserine transsuccinylase                                    | metA                   | $HSER + SUCCOA \rightarrow COA + OSLHSER$                                       | Greene, 1996   |
| Cystathionine synthase   | metB                   | OSLHSER + CYS $\rightarrow$ SUCC + HCYS + PYR + NH3                             | Greene, 1996   |
| Cystathionase  | metC                   | HCYS + ADN ↔ SAH  | Greene, 1996   |
| Methionine synthase  | metEH                  | $HCYS + MTHF \rightarrow MET + THF$   | Greene, 1996; Matthews, 1996                         |
| Methionyl adenosyl transferase                                 | metK                   | $MET + ATP \rightarrow PPI + PI + SAM$  | Greene, 1996; Matthews, 1996                         |
| Biosynthesis of purine nucleotides                             |                        |   | , ., .,  |
| Glutamine PRPP amidotransferase                                | purF                   | $PRPP + GLN \rightarrow PPI + GLU + PRAM$                                       | Zalkin and Nygaard, 1996                             |
| GAR synthetase   | purD                   | $PRAM + ATP + GLY \leftrightarrow ADP + PI + GAR$                               | Zalkin and Nygaard, 1996                             |
| GAR transformylase   | purNT                  | $GAR + FTHF \rightarrow THF + FGAR$   | Zalkin and Nygaard, 1996                             |
| FGAM synthetase  | purL                   | $FGAR + ATP + GLN \rightarrow GLU + ADP + PI + FGAM$                            | Zalkin and Nygaard, 1996                             |
| AIR synthetase   | purM                   | $FGAM + ATP \rightarrow ADP + PI + AIR$   | Zalkin and Nygaard, 1996                             |
| RCAIM synthetase   | purK                   | $AIR + CO2 \leftrightarrow RCAIM$   | Zalkin and Nygaard, 1996                             |
| PRSCAIM synthetase   | purE                   | $RCAIM + ATP + ASP \leftrightarrow ADP + PI + PRSCAIM$                          | Zalkin and Nygaard, 1996                             |
| Adenylosuccinate lyase   | purB                   | $PRSCAIM \leftrightarrow FUM + AICAR$   | Zalkin and Nygaard, 1996                             |
| AICAR transformylase   | purH                   | $AICAR + FTHF \leftrightarrow THF + PRFICA$                                     | Zalkin and Nygaard, 1996                             |
| IMP cyclohydrolase   | purH                   | $PRFICA \leftrightarrow IMP$  | Zalkin and Nygaard, 1996                             |
| Adenylosuccinate synthetase                                    | purA                   | $IMP + GTP + ASP \rightarrow GDP + PI + ASUC$                                   | Zalkin and Nygaard, 1996                             |
| Adenylosuccinate lyase   | purB                   | $ASUC \leftrightarrow FUM + AMP$  | Zalkin and Nygaard, 1996                             |
| AMP phosphatase  |                        | $AMP \rightarrow PI + ADN$  | Zalkin and Nygaard, 1996                             |
| Adenylate kinase   | adk                    | $ATP + ADN \rightarrow ADP + AMP$   | Zalkin and Nygaard, 1996                             |
| Adenylate kinase   | adk                    | $ATP + AMP \rightarrow 2 ADP$   | Zalkin and Nygaard, 1996                             |
| IMP dehydrogenase  | guaB                   | $IMP + NAD \rightarrow NADH + XMP$  | Zalkin and Nygaard, 1996                             |
| GMP synthetase   | guaA                   | $XMP + ATP + GLN \rightarrow GLU + AMP + PPI + GMP$                             | Zalkin and Nygaard, 1996                             |
| GMP kinase   | gmk                    | $GMP + ATP \leftrightarrow GDP + ADP$   | Zalkin and Nygaard, 1996                             |
| GDP kinase   | gmk                    | $GDP + ATP \leftrightarrow GTP + ADP$   | Zalkin and Nygaard, 1996                             |
| Ribonucleotide reductase (ADP)                                 | deoD                   | $ADP + NADPH \rightarrow DADP + NADP$   | Zalkin and Nygaard, 1996                             |
| Ribonucleotide reductase (GDP)                                 | deoD                   | $GDP + NADPH \rightarrow DGDP + NADP$   | Zalkin and Nygaard, 1996                             |
| Ribonucleotide reductase (ATP)                                 | deoD                   | $ATP + NADPH \rightarrow NADP + DATP$ $CTP + NADPH \rightarrow NADP + DCTP$     | Zalkin and Nygaard, 1996                             |
| Ribonucleotide reductase (GTP)                                 | deoD                   | $GTP + NADPH \rightarrow NADP + DGTP$   | Zalkin and Nygaard, 1996                             |
| dADP kinase  | nck                    | $DADP + ATP \leftrightarrow DATP + ADP$   | Zalkin and Nygaard, 1996                             |
| dGDP kinase<br>dAMP kinase                                     | ndk<br>ndk             | $DGDP + ATP \leftrightarrow DGTP + ADP$ $DAMP + ATP \leftrightarrow ADP + DADP$ | Zalkin and Nygaard, 1996<br>Zalkin and Nygaard, 1996 |
| dGMP kinase  |                        | $DAMP + ATP \leftrightarrow ADP + DADP$ $DCMP + ATP \leftrightarrow DCDP + ADP$ | Zalkin and Nygaard, 1996<br>Zalkin and Nygaard, 1996 |
| Biosynthesis of pyrimidines                                    | ndk                    | $DGMP + ATP \leftrightarrow DGDP + ADP$   | Zaikiii alid Nygaard, 1990                           |
| Aspartate carbamoyl transferase                                | pyrBI                  | $CAP + ASP \rightarrow CAASP + PI$  | Neuhard and Klein, 1996                              |
| Dihydroorotase   | pyr <b>B</b> 1<br>pyrC | $CAASP \leftrightarrow DOROA$   | Neuhard and Klein, 1996                              |
| Dihydroorotate dehydrogenase                                   | pyrC<br>pyrD           | $OROA + NAD \leftrightarrow NADH + OROA$  | Neuhard and Klein, 1996                              |
| Orotate phosphoribosyl transferase                             | pyrE<br>pyrE           | $OROA + PRPP \leftrightarrow PPI + OMP$   | Neuhard and Klein, 1996                              |
| OMP decarboxylase  | pyrE<br>pyrF           | $OMP \rightarrow CO2 + UMP$   | Neuhard and Klein, 1996                              |
| UMP kinase   | pyrH<br>pyrH           | $UMP + ATP \leftrightarrow UDP + ADP$   | Neuhard and Klein, 1996                              |
| UDP kinase   | ndk                    | $UDP + ATP \leftrightarrow UTP + ADP$   | Neuhard and Klein, 1996                              |
| CTP synthetase   | pyrG                   | $UTP + GLN + ATP \rightarrow GLU + CTP + ADP + PI$                              | Neuhard and Klein, 1996                              |
| CMP kinase   | ndk                    | $CMP + ATP \leftrightarrow CDP + ADP$   | Neuhard and Klein, 1996                              |
| CDP kinase   | ndk                    | $CDP + ATP \leftrightarrow CTP + ADP$   | Neuhard and Klein, 1996                              |
| Deoxycytidilate deaminase                                      | cdd                    | $DCMP \rightarrow NH3 + DUMP$   | Neuhard and Klein, 1996                              |

| Enzyme  | Gene   | Pathway   | Reference                               |
|---|--------|---|---|
| Ribonucleotide reductase (CDP)                      | nrdAB  | $CDP + NADPH \rightarrow DCDP + NADP$   | Neuhard and Klein, 1996                 |
| Ribonucleotide reductase (UDP)                      | nrdAB  | $UDP + NADPH \rightarrow DUDP + NADP$   | Neuhard and Klein, 1996                 |
| Ribonucleotide reductase (CTP)                      | nrdAB  | $CTP + NADPH \rightarrow DCTP + NADP$   | Neuhard and Klein, 1996                 |
| Ribonucleotide reductase (UTP)                      | nrdAB  | $UTP + NADPH \rightarrow NADP + DUTP$   | Neuhard and Klein, 1996                 |
| dCMP kinase   | ndk    | $DCMP + ATP \leftrightarrow ADP + DCDP$   | Neuhard and Klein, 1996                 |
| dCDP kinase   | ndk    | $DCDP + ATP \leftrightarrow DCTP + ADP$   | Neuhard and Klein, 1996                 |
| dUDP kinase   | ndk    | $DUDP + ATP \leftrightarrow DUTP + ADP$   | Neuhard and Klein, 1996                 |
| dUTP pyrophosphatase                                | dut    | $DUTP \rightarrow PPI + DUMP$   | Neuhard and Klein, 1996                 |
| dUMP kinase   | ndk    | $DUMP + ATP \leftrightarrow ADP + DUDP$   | Neuhard and Klein, 1996                 |
| Thymidilate synthetase                              | thyA   | $DUMP + METTHF \rightarrow DHF + TMP$   | Matthews, 1996; Neuhard and Klein, 1996 |
| TMP kinase  | tmk    | $TMP + ATP \leftrightarrow ADP + TDP$   | Neuhard and Klein, 1996                 |
| TDP kinase  | ndk    | $TDP + ATP \leftrightarrow ADP + TTP$   | Neuhard and Klein, 1996                 |
| Biosynthesis of THF                                 |        |   |   |
| Methylene THF reductase                             | metF   | $METTHF + NADH \rightarrow NAD + MTHF$  | Matthews, 1996                          |
| Methylene THF dehydrogenase                         | folD   | $METTHF + NADP \leftrightarrow METHF + NADPH$   | Matthews, 1996                          |
| Methenyl tetrahydrofolate cyclehydrolase            | folD   | $METHF \leftrightarrow FTHF$  | Matthews, 1996                          |
| Biosynthesis of membrane lipids                     |        |   |   |
| Acetyl-CoA carboxylase                              | acc    | $ACCOA + ATP + CO2 \leftrightarrow MALCOA + ADP + PI$   | Cronan and Rock, 1996                   |
| Malonyl-CoA:ACP transacylase                        | mta    | $MALCOA + ACP \leftrightarrow MALACP + COA$   | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I                           | kas1   | $MALACP \rightarrow ACACP + CO2$  | Cronan and Rock, 1996                   |
| Acetyl-CoA:ACP transacylase                         | ata    | $ACACP + COA \leftrightarrow ACCOA + ACP$   | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I (C14:0)                   | fab    | ACACP + 6 MALACP + 12 NADPH $\rightarrow$ C14:0ACP<br>+ 6 CO2 + 12 NADP + 6 ACP                               | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I (C14:1) fab               |        | $ACACP + 6 MALACP + 11 NADPH \rightarrow C14:1ACP + 6 CO2 + 11 NADP + 6 ACP$                                  | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I (C16:0) fab               |        | ACACP + 7 MALACP + 14 NADPH $\rightarrow$ C16:0ACP<br>+ 7 CO2 + 14 NADP + 7 ACP                               | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I (C16:1)                   |        |   | Cronan and Rock, 1996                   |
| b-Ketoacyl-ACP synthase I (C18:1)                   | fab    | ACACP + 8 MALACP + 15 NADPH $\rightarrow$ C18:1ACP + 8 CO2 + 15 NADP + 8 ACP                                  | Cronan and Rock, 1996                   |
| Glycerol-3-phosphate dehydrogenase                  | gpsA   | $NADH + T3P2 \leftrightarrow GL3P + NAD$  | Cronan and Rock, 1996                   |
| 1-Acyl-G3P acyltransferase                          | pls    | GL3P + 0.03 C14:0ACP + 0.086 C14:1ACP + 0.607<br>C16:0ACP + 0.12 C16:1ACP + 0.85 C18:1ACP<br>→ PA + 1.693 ACP | Cronan and Rock, 1996                   |
| CDP-Diacylglycerol synthetase                       | cdsA   | $PA + CTP \leftrightarrow CDPDG + PPI$  | Cronan and Rock, 1996                   |
| Phosphatidylserine synthase                         | pssA   | $CDPDG + SER \leftrightarrow CMP + PS$  | Cronan and Rock, 1996                   |
| PS decarboxylase                                    | psd    | $PS \rightarrow PE + CO2$   | Cronan and Rock, 1996                   |
| Phosphatidylglycerol phosphate synthase             | pgsA   | $CDPDG + GL3P \leftrightarrow CMP + PGP$  | Cronan and Rock, 1996                   |
| Phosphatidylglycerol phosphate phosphate            | pgpA   | $PGP \rightarrow PI + PG$   | Cronan and Rock, 1996                   |
| Cardiolipin synthase<br>Biosynthesis of isoprenoids | cls    | $PG + CDPDG \leftrightarrow CL + CMP$   | Cronan and Rock, 1996                   |
| Aldose reductase                                    |        | $GL + NADP \leftrightarrow NADPH + GLAL$  | White, 1996                             |
| Glyceraldehyde kinase                               |        | $GLAL + ATP \rightarrow ADP + T3P1$   | White, 1996                             |
| Hydroxymethyl-glutaryl-CoA synthase                 |        | $3 \text{ ACCOA} \rightarrow 2 \text{ COA} + \text{HMGCOA}$   | White, 1996                             |
| 3-Methyl-glutaconyl-CoA hydratase                   |        | $HMGCOA \leftrightarrow TMGCOA$   | White, 1996                             |
| IPP synthase  |        | HMGCOA + 2 NADPH + 3 ATP $\rightarrow$ COA<br>+ 2 NADP + 3 ADP +1 PI + CO2 + IPPP                             | White, 1996                             |
| GGPP synthase                                       |        | $4 \text{ IPPP} \rightarrow 3 \text{ PPI} + \text{GGPP}$  | White, 1996                             |
| Methylcrotonyl-CoA carboxylase                      |        | MCCOA + ATP + CO2 ↔ ADP + TMGCOA + PI   | Michal, 1996                            |
| Acyl-CoA dehydrogenase                              |        | $ISOVCOA + Q \leftrightarrow QH2 + MCCOA$   | Michal, 1996                            |
| 2-Keto-isocaproate decarboxylase                    |        | OICAP + COA + NADP → NADPH + CO2 + ISOVCOA  | Michal, 1996                            |
| Biosynthesis of quinone                             | .1:0   |   | Managed 1005                            |
| Chorismate pyruvate-lyase                           | ubiC   | $CHOR \rightarrow 4HBZ + PYR$   | Meganathan, 1996                        |
| Hydroxybenzoate octaprenyl-<br>transferase          | ubiADX | $4HBZ + GGPP \rightarrow 2PPP + CO2 + PPI$  | Meganathan, 1996                        |

| Enzyme   | Gene       | Pathway   | Reference                           |
|--|------------|---|-------------------------------------|
| 2O6H synthetase  | ubiB       | 2PPP + O2 → 2O6H  | Meganathan, 1996                    |
| QH2 synthetase   | ubiEFGH    | $2O6H + 2 O2 + 3 SAM \rightarrow 3 SAH + QH2$   | Meganathan, 1996                    |
| Biosynthesis of riboflavin                               |            |   |                                     |
| GTP cyclohydrolase                                       | ribA       | $GTP \rightarrow D6RP5P + CO2 + PPI$  | Bacher et al., 1996                 |
| Pyimidine deaminase                                      | ribD       | $D6RP5P \rightarrow A6RP5P + NH3$   | Bacher et al., 1996                 |
| Pyrimidine reductase                                     | ribD       | $A6RP5P + NADPH \rightarrow A6RP5P2 + NADP$   | Bacher et al., 1996                 |
| Phosphatase  |            | $A6RP5P2 \rightarrow A6RP + PI$   | Bacher et al., 1996                 |
| 3,4-Dihydroxy-2-butanone-4-phosphate synthase            | ribB       | A6RP → DB4P + FORMATE   | Bacher et al., 1996                 |
| 6,7-Dimethyl-8-ribityllumazine synthase                  | ribE       | $DB4P + A6RP \rightarrow D8RL + PI$   | Bacher et al., 1996                 |
| Riboflavin synthase                                      | ribC       | $2 D8RL \rightarrow RIBOFLAVIN + A6RP$  | Bacher et al., 1996                 |
| Riboflavin kinase  | ribF       | $RIBOFLAVIN + ATP \rightarrow FMN + ADP$  | Bacher et al., 1996                 |
| FAD synthetase   | ribF       | $FMN + ATP \rightarrow FAD + PPI$   | Bacher et al., 1996                 |
| Biosynthesis of folate                                   |            |   |                                     |
| GTP cyclohydrolase                                       | folE       | $GTP \rightarrow FORMATE + AHTD$  | Green et al., 1996                  |
| H2Neopterin triphosphate<br>pyrophosphatase              |            | $AHTD \rightarrow 3 PI + DHP$   | Green et al., 1996                  |
| H2Neopterin aldolase                                     |            | $DHP \rightarrow AHHMP + GLAL$  | Green et al., 1996                  |
| 6-Hydroxymethyl H2pterin<br>pyrophosphokinase            | fol K      | $AHHMP + ATP \rightarrow AMP + AHHMD$   | Green et al., 1996                  |
| H2pteroate synthase                                      | fol P      | $AN + AHHMD \rightarrow PPI + DHD$  | Green et al., 1996                  |
| Dihydrofolate reductase                                  | folA       | $DHD + ATP + GLU \rightarrow ADP + PI + DHF$  | Green et al., 1996                  |
| Biosynthesis of coenzyme A                               |            |   |                                     |
| CoA Synthase   | panBCDE    | OIVAL + METTHF + NADPH + ALA + CTP<br>+ 4 ATP + CYS $\rightarrow$ THF + NADP + AMP + 2 PPI<br>+ 2 ADP + CO2 + COA   | Jackowski, 1996<br>(Lumped pathway) |
| ACP Synthase   | acpS       | $COA \rightarrow 35ADP + ACP$   | Jackowski, 1996                     |
| 3,5-ADP phosphatase                                      | •          | $35ADP \rightarrow AMP + PI$  | •                                   |
| Biosynthesis of NAD                                      |            |   |                                     |
| Quinolate synthase                                       | nadAB      | $ASP + FAD + T3P2 \rightarrow FADH2 + PI + QNL$   | Penfound and Foster, 1996           |
| Quinolate phosphoribosyl transferase                     | nadC       | $QNL + PRPP \rightarrow PPI + NICNT + CO2$  | Penfound and Foster, 1996           |
| NAMN adenylyl tranferase                                 | nadD       | $NICNT + ATP \rightarrow PPI + DANAD$   | Penfound and Foster, 1996           |
| Deamido-NAD ammonia ligase                               | nadE       | $DANAD + ATP + NH3 \rightarrow AMP + PPI + NAD$   | Penfound and Foster, 1996           |
| NAD kinase   |            | $NAD + ATP \rightarrow NADP + ADP$  | Penfound and Foster, 1996           |
| NADP phosphatase   |            | $NADP \rightarrow NAD + PI$   | Penfound and Foster, 1996           |
| Biosynthesis of porphyrins and hemes                     |            |   |                                     |
| GSA synthetase   | gltX, hemA | $\begin{aligned} \text{GLU} + \text{ATP} + \text{NADPH} &\rightarrow \text{GSA} + \text{AMP} + \text{PPI} \\ + \text{NADP} \end{aligned}$                             | Beale, 1996                         |
| GSA aminotransferase                                     | hemL       | $GSA \rightarrow ALAV$  | Beale, 1996                         |
| Porphyrinogen synthetase                                 | hemBCD     | $8 \text{ ALAV} \rightarrow \text{PORPH} + \text{NH3}$  | Beale, 1996                         |
| Biosynthesis of lippopolysaccharide and murein           |            |   |                                     |
| Glutamine fructose-6-P transaminase                      | glmS       | $F6P + GLN \rightarrow GLU + GA6P$  | Raetz, 1996                         |
| Glucosamine-P acetyl transferase                         | glmU       | $GA6P + ACCOA \rightarrow AGA6P + COA$  | Raetz, 1996                         |
| Acetyl glucosamine mutase                                | glmU       | $AGA6P \leftrightarrow AGA1P$   | Raetz, 1996                         |
| UDP N-acetylglucosamine                                  |            | $AGA1P + UTP \leftrightarrow PPI + UDPGA$   |                                     |
| pyrophorylase UDP N-acetylglucosamine                    |            | $UDPGA \leftrightarrow UDPGLN$  |                                     |
| 4-epimerase  | 1.1.4      | HDDGA - DED - DI HDDGG  | D 4 1006                            |
| N-Acylglucosamine-6-P 2-epimerase UDP-N-acetylmuramate   | kdsA       | $\begin{array}{l} \text{UDPGA} + \text{PEP} \leftrightarrow \text{PI} + \text{UDPGC} \\ \text{UDPGC} + \text{NADH} \rightarrow \text{UDPAM} + \text{NAD} \end{array}$ | Raetz, 1996                         |
| dehydrogenase<br>CMP-2-keto-3-deoxyoctanoate<br>synthase | kdsB       | $R5P + PEP + CTP \rightarrow 2 PI + PPI + CMPKDO$   | Raetz, 1996                         |
| Isomerase+mutase+pyropho-<br>phorylase+epimerase         |            | $S7P + ATP \leftrightarrow ADPHEP + PPI$  |                                     |
| Ethanolamine phosphotransferase                          |            | $PE + CMP \leftrightarrow CDPETN + DGR$   | Michal, 1993                        |
| Phosphatidate phosphatase                                |            | $PA \leftrightarrow PI + DGR$   | Michal, 1993                        |
| Lyposaccharide synthetase                                |            | 3 UDPGLN + 5 C14:0 + 2 ATP + 3 CMPKDO<br>+ CDPETN + 3 ADPHEP + 2 UDPG + PE+ 2   | ,                                   |
|  |            | UDPGAL $\rightarrow$ 2 AC + UMP + 6 UDP + 5 ADP<br>+ 3 CMP + DGR + LPS  |                                     |

| Enzyme                                     | Gene | Pathway  | Reference         |
|--|------|--|-------------------|
| UDP glucose synthase                       |      | $G1P + UTP \rightarrow PPI + UDPG$                                 |                   |
| UDP galactose synthase                     |      | $G1P + UTP \leftrightarrow PPI + UDPGAL$                           |                   |
| Murein synthetase                          |      | UDPGA + UDPAM + 2 ALA + D26PIM + GLU                               |                   |
| •  |      | $+$ 5 ATP $\rightarrow$ PEPTIDO $+$ 5 ADP $+$ 5 PI $+$ 2 UDP       |                   |
| Polyphosphate and pyrophosphate metabolism |      |  |                   |
| Pyrophosphatase                            | ppa  | $PPI \rightarrow 2 PI$   | Wanner, 1996      |
| Polyphosphate kinase                       | ppk  | $1000 \text{ ATP} \leftrightarrow 1000 \text{ ADP} + \text{POLYP}$ | Wanner, 1996      |
| Polyphosphatase                            | ppx  | $POLYP \rightarrow 1000PI$   | Wanner, 1996      |
| Transport reactions                        |      |  |                   |
| Ammonia transport                          |      | $NH3ext + Hext \leftrightarrow NH3$                                | Silver, 1996      |
| Sulfate transport                          |      | $H2SO4ext \leftrightarrow H2SO4$                                   |                   |
| Phosphate transport                        | pit  | $PIext + Hext \leftrightarrow PI$                                  | Wanner, 1996      |
| Acetate transport                          |      | $ACext + Hext \leftrightarrow AC$                                  |                   |
| Lactate transport                          |      | $LACext + Hext \leftrightarrow LAC$                                |                   |
| Formate transport                          |      | $FORMATEext + Hext \leftrightarrow FORMATE$                        |                   |
| Ethanol transport                          |      | $ETHANOLext \leftrightarrow ETHANOL$                               |                   |
| Succinate transport                        |      | $SUCCext + Hext \leftrightarrow SUCC$                              |                   |
| D-Glyceraldehyde transport                 |      | $GLALext + Hext \leftrightarrow GLAL$                              |                   |
| Glucose transport                          |      | $GLCext \leftrightarrow GLC$                                       |                   |
| Carbon dioxide transport                   |      | $CO_2$ ext $\leftrightarrow CO_2$                                  |                   |
| Oxygen transport                           |      | $O_2$ ext $\leftrightarrow$ $O_2$                                  |                   |
| Glycerol metabolism                        |      |  |                   |
| Glycerol kinase                            |      | $GL + ATP \leftrightarrow GL3P + ADP$                              | Karp et al., 1996 |
| Glycerol-3-phosphate dehydrogenase         |      | $GL3P + FAD \rightarrow T3P2 + FADH2$                              | Karp et al., 1996 |

## APPENDIX B: METABOLITE DEFINITIONS

| 2.123, 22222      |  | 7.1 LINDIX B. GOITHINGED |   |  |
|-------------------|--|--------------------------|---|--|
| Abbreviation      | Compound   | Abbreviation             | Compound  |  |
| 13P2DG<br>2K3D6PG | 1,3- <i>P</i> -D glycerate<br>2-Dehydro-3-deoxy-6- <i>P</i> -gluconate | AHTD                     | 2-Amino-4-hydroxy-6-(erythro-1-2-3-trihydroxypropyl) dihydropteridine-phosphate |  |
| 2O6H              | 2-Octaprenol 6-hydroxyphenol   | AICAR                    | 5-Phosphoribosyl-5-amino-4-imidazole carboxamide                                |  |
| 2PDGL             | 2-P-D glycerate  | AIR                      | 5-Phosphoribosyl-5-aminoimidazole   |  |
| 2PPP              | 2-Polyprenylphenol   | AKG                      | α-Ketoglutarate   |  |
| 3DDAH7P           | 3-Deoxy-D-arabinoheptulosonate-7-phosphate                             | ALA                      | Alanine   |  |
| 3PDGL             | 3 P-D glycerate  | ALAV                     | δ-Amonolevulinate   |  |
| 3PSER             | 3-Phosphoserine  | AMP                      | Adenosine monophosphate   |  |
| 3PSME             | o (1-Carboxyvinyl)-3-D-shikimate                                       | AN                       | Anthranilate  |  |
| 4HBZ              | 4-Hydroxybenzoate  | APS                      | Adenylyl sulfate  |  |
| 5MTA              | 5'-Methylthioadenosine   | ARG                      | Arginine  |  |
| A6RP              | 5-Amino-6-ribitylamino-2,4(1H,3H)-pyrimidinedione                      | ARGSUCC                  | 1-Arginiosuccinate  |  |
| A6RP5P            | 5-Amino-6-ribosylamino-2,4(1H,3H)-pyrimidinedione                      | ASER                     | o-Acetylserine  |  |
|                   | 5'-phosphate   | ASN                      | Asparagine  |  |
| A6RP5P2           | 5-Amino-6-ribitylamino-2,4(1H,3H)-pyrimidinedione                      | ASP                      | Aspartate   |  |
|                   | 5'-phosphate   | ASPSA                    | Aspartate β-semialdehyde  |  |
| AABK              | 2-Amino-3-ketobutyrate   | ASUC                     | Adenylsuccinate   |  |
| ABUT              | 2-Aceto-2-hydroxybutyrate  | ATP                      | Adenosine triphosphate  |  |
| AC                | Acetate  | BASP                     | β-Aspartyl-phosphate  |  |
| ACACP             | Acetyl-ACP   | C14:0ACP                 | Myristic acid ACP   |  |
| ACAL              | Acetaldehyde   | C14:1ACP                 | β-Hydroxymyristic acid ACP  |  |
| ACCOA             | Acetyl-CoA   | C16:0ACP                 | Palmitic acid ACP   |  |
| ACTP              | Acetyl-phosphate   | C16:1ACP                 | Palmitoleic acid ACP  |  |
| ADN               | Adenosine  | C18:1ACP                 | cis-Vaccenic acid ACP   |  |
| ADP               | Adenosine diphosphate  | CAASP                    | Carbamoyl aspartate   |  |
| ADPHEP            | ADP-mannoheptose   | CAP                      | Carbamoyl phosphate   |  |
| AGA1P             | N-Acetyl-D-glucosamine-1-phosphate                                     | CBHCAP                   | 3-Carboxy-3-hydroxy-isocaproate   |  |
| AGA6P             | <i>N</i> -Acetyl-D-glucosamine-6-phosphate                             | CDP                      | Cytidine diphosphate  |  |
| AHHMD             | 2-Amino-4-hydroxy-6-hydroxymethyl                                      | CDPDG                    | CDP-2-Diacylglycerol  |  |
|                   | dihydropteridine-pyrophosphate   | CDPETN                   | CDP-Ethanolamine  |  |
| AHHMP             | 2-Amino-4-hydroxy-6-hydroxymethyl dihydropteridine                     | CHOR                     | Chorismate  |  |

| Abbreviation | Compound   | Abbreviation | Compound                                    |
|--------------|--|--------------|---|
| CITR         | Citrulline   | GLX          | Glyoxylate                                  |
| CL           | Cardiolypin  | GLY          | Glycine                                     |
| CMP          | Cytidine monophosphate                             | GLYCOGEN     | Glycogen                                    |
| CMPKDO       | CMP-2-Keto-3-deoxyoctanoate                        | GMP          | Guanosine monophosphate                     |
| CO2          | Carbon dioxide                                     | GSA          | Glutamate 1-semialdehyde                    |
| COA          | Coenzyme A-SH                                      | GTP          | Guanosine triphosphate                      |
| CPAD5P       | 1-o-Carboxyphenylamino 1-deoxyribulose-5-phosphate | H2S          | Hydrogen sulfide                            |
| CTP          | Cytidine triphosphate                              | H2SO3        | Hydrogen sulfite                            |
| CYS          | Cysteine   | H2SO4        | Hydrogen sulfate                            |
| D23PIC       | 2,3-Dihydrodipicolinate                            | HCYS         | Homocysteine                                |
| D26PIM       | L,L-2,6-Diaminopimelate                            | HEXT         | External H <sup>+</sup>                     |
| D6PGC        | D-6-Phosphoglucono-δ-lactone                       | HIS          | Histidine                                   |
| D6PGL        | D-6-Phosphogluconate                               | HISOL        | Histidinol                                  |
| D6RP5P       | 2,5-Diamino-6-ribosylamino-4(3H)-pyrimidinedione   | HISOL-P      | 1-Histidinol-phosphate                      |
|              | 5'-phosphate                                       | HMGCOA       | 3-Hydroxy-3-methyl-glutaryl CoA             |
| D8RL         | 6,7-Dimethyl-8-ribityllumazine                     | HPHPYR       | para-Hydroxy phenyl pyruvate                |
| DADP         | Deoxyadenosine diphosphate                         | HSER         | Homoserine                                  |
| DANAD        | Deamido-NAD  | ICIT         | Isocitrate                                  |
| DATP         | Deoxyadenosine triphosphate                        | IGP          | Indole glycerol phosphate                   |
| DB4P         | L-3,4-Dihydroxy-2-butanone-4-phosphate             | ILE          | Isoleucine                                  |
| DCDP         | Deoxycytidine diphosphate                          | IMACP        | Imidazole acetyl-phosphate                  |
| DCTP         | Deoxycytidine triphosphate                         | IMP          | Inosine monophosphate                       |
| DGDP         | Deoxyguanosine diphosphate                         | IPPP         | Isopentyl pyrophosphate                     |
| DGMP         | 2-Deoxy-guanosine-5-phosphate                      | IPPMAL       | 3-Isopropylmalate                           |
| DGR          | D-1,2-Diacylglycerol                               | ISOVCOA      | Isovaleryl-CoA                              |
| DGTP         | Deoxyguanosine triphosphate                        | LA           | Lipoamide                                   |
| DHD          | 7,8-Dihydropteroate                                | LAC          | Lactate                                     |
| DHF          | Dihydrofolate                                      | LEU          | Leucine                                     |
| DHMVA        | 2,3-Dihydroxy-3-methyl-valerate                    | LPS          | Lipposaccharide                             |
| DHP          | Dihydroneopterin                                   | LYS          | <i>l</i> -Lysine                            |
| DHSK         | Dehydroshikimate                                   | MAL          | Malate                                      |
| DHVAL        | α,β-Dihydroxy-isovalerate                          | MALACP       | Malonyl-ACP                                 |
| DIMGP        | d-Erythroimidazoleglycerol-phosphate               | MALCOA       | Malonyl-CoA                                 |
| DOROA        | Dihydroorotic acid                                 | MCCOA        | 3-Methyl crotonyl-CoA                       |
| DQT          | 3-Dehydroquinate                                   | MDAP         | meso-Diaminopimelate                        |
| DSAM         | Decarboxylated SAM                                 | MET          | Methionine                                  |
| DUDP         | Deoxyuridine diphosphate                           | METHF        | 5,10-Methenyl tetrahydrofolate              |
| DUMP         | Deoxyuridine monophosphate                         | METTHF       | 5,10-Methylene tetrahydrofolate             |
| DUTP         | Deoxyuridine triphosphate                          | MTHF         | 5-Methyl tetrahydrofolate                   |
| E4P          | Erythrose 4-phosphate                              | NAARON       | $N$ - $\alpha$ -Acetyl ornithine            |
| EFF          | Efficiency   | NADH         | Nicotinamide adenine dinucleotide           |
| ETHANOL      | Ethanol  | NADPH        | Nicotinamide adenine dinucleotide phosphate |
| F16P         | Fructose 1,6-diphosphate                           | NAGLU        | N-Acetyl glutamate                          |
| F6P          | Fructose 6-phosphate                               | NAGLUYP      | N-Acetyl glutamyl-phosphate                 |
| FADH         | Flavin adenine dinucleotide                        | NAGLUSAL     | N-Acetyl glutamate semialdehyde             |
| FGAM         | 5-Phosphoribosyl- <i>N</i> -formylgycineamidine    | NH3          | Ammonia                                     |
| FGAR         | 5-Phosphoribosyl- <i>N</i> -formylglycineamide     | NICNT        | Nicotinate nucleotide                       |
| FMN          | Riboflavin 5'-phosphate                            | NPRAN        | N-5-Phosphoribosyl-antranilate              |
| FORMATE      | Formate  | NS26DP       | N-Succinyl-1,1-2,6-diaminopimelate          |
| FTHF         | 10-Formyl-tetrahydrofolate                         | NS2A6O       | N-Succinyl-2-amino-6-ketopimelate           |
| FUM          | Fumarate   | 02           | Oxygen                                      |
| G1P          | Glucose 1-phosphate                                | OA           | Oxaloacetate                                |
| G6P          | Glucose 6-phosphate                                | OBUT         | Oxobutyrate                                 |
| GA6P         | Glucosamine-6-phosphate                            | OICAP        | 2-Oxoisocaproate                            |
| GAR          | 5-Phosphoribosyl glycineamide                      | OIVAL        | Oxoisovalerate                              |
| GDP          | Guanosine diphosphate                              | OMP          | Orotidylate                                 |
| GGPP         | Geranylgeranyl pyrophosphate                       | OMVAL        | Oxomethylvalerate                           |
| GL           | Glycerol   | ORN          | Ornithine                                   |
| GL3P         | Glycoden 3-phosphate                               | OROA         | Orotic acid                                 |
| GLAL         | D-Glyceraldehyde                                   | OSLHSER      | o-Succinyl-L-homoserine                     |
| GLC          | Glucose  | PA           | Phosphatidyl acid                           |
| GLN          | Glutamine  | PAP          | Adenosine-3,5-diphosphate                   |
| GLUCSAL      | Glutamate  | PAPS         | 3-Phosphoadenylyl sulfate                   |
| GLUGSAL      | L-Glutamate γ-semialdehyde                         | PE           | Phosphatidyl ethanolamine                   |

| Abbreviation   | Compound  |
|----------------|---|
| PEP            | Phosphoenolpyruvate                                       |
| PEPTIDO        | Peptidoglycan   |
| PG             | Phosphatidyl glycerol                                     |
| PGP            | 1-1-Phosphatidyl-glycerol-phosphate                       |
| PHE            | Phenylalanine   |
| PHEN           | Prephenate  |
| PHP            | 3-Phosphohydroxypyruvate                                  |
| PHPYR          | Phenyl pyruvate   |
| PHSER          | o-Phospho-l-homoserine                                    |
| PI             | Phosphate (inorganic)                                     |
| PIP26DX        | δ-Pieperidine-2,6-dicarboxylate                           |
| POLYP<br>PORPH | Polyphosphate   |
| PPI            | Porphyrinogen  Pyrophyrinogen                             |
| PRAM           | Pyrophosphate<br>5-Phospho-β-D-ribosyl amine              |
| PRBAMP         | Phosphoribosyl-AMP  |
| PRBATP         | Phosphoribosyl-ATP  |
| PRFICA         | 5-Phosphoribosyl-formamido-4-imidazole carboxamido        |
| PRFP           | Phosphoribosyl-formimino-AICAR-phosphate                  |
| PRLP           | Phosphoribulosyl-formimino-AICAR-phosphate                |
| PRO            | Proline   |
| PRPP           | Phosphoribosyl pyrophosphate                              |
| PRSCAIM        | 5-Phosphoribosy-14- <i>N</i> -succinocarboxyamide-5-amino |
| PS             | Phosphatidyl serine                                       |
| PTRSC          | Putrescine  |
| PYR            | Pyruvate  |
| Q              | Ubiquinone  |
| QH2            | Ubiquinol   |
| QNL            | Quinolate   |
| R5P            | Ribose 5-phosphate  |
| RCAIM          | 5-p-Ribosyl-4-carboxy-5-aminoimidazole                    |
| RIBOFLAVIN     | Riflavin  |
| RL5P           | D-Ribulose 5-phosphate                                    |
| S7P            | D-Sedoheptulose-7-P                                       |
| SAH<br>SAM     | s-Adenosyl homocystine<br>s-Adenosyl methionine           |
| SER            | Serine  |
| SME            | Shikimate   |
| SME5P          | Shikimate-5-phosphate                                     |
| SPRMD          | Spermidine  |
| SUCC           | Succinate   |
| SUCCOA         | Succinyl-CoA  |
| T3P1           | Glyceraldehyde-3-phosphate                                |
| T3P2           | Dihydroxyacetone phosphate                                |
| TDP            | Thymidine-5-diphosphate                                   |
| THF            | Tetrahydrofolate  |
| THR            | Threonine   |
| TMGCOA         | trans-3-Methyl-glutaconyl-CoA                             |
| TMP            | Thymidine-5-monophosphate                                 |
| TRP            | Tryptophan  |
| TTP            | Thymidine-5-triphosphate                                  |
| TYR            | Tyrosine  |
| UDP            | Uridine diphosphate                                       |
| UDPAM          | UDP-Acetylmuramate  |
| UDPGA          | UDP-N-Acetylglucosamine                                   |
| UDPGAL         | UDP N-Acetylgalactosamine                                 |
| UDPGC          | UDP N-Acetylglucosamine-enolpyruvate                      |
| UDPGLN         | UDP N-Actylglucosamine                                    |
| UMP            | Uridine monophosphate                                     |
| UREA<br>UTP    | Urea<br>Uridine triphosphate                              |
| VAL            | Valine  |
| VAL            | Vululose 5 mhosmbets                                      |

Xylulose-5-phosphate

Xantosine monophosphate

X5P

XMP

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