Abstract

The dynamic modelling of metabolic networks aims at describing the temporal evolution of metabolite concentrations in cells. This area has attracted increasing attention in recent years due to the availability of high-throughput data and the general development of systems biology as a promising approach to study living organisms. Biochemical Systems Theory (BST) provides an accurate formalism to describe biological dynamic phenomena. However, knowledge about the molecular organization level, used in these models, is not enough to explain phenomena such as the driving forces of these metabolic networks. Dynamic Energy Budget (DEB) theory captures the quantitative aspects of the organization of metabolism at the organism level in a way that is non-species specific. This imposes constraints on the sub-organismal organization that are not present in the bottom-up approach of systems biology. We use in vivo data of lactic acid bacteria under various conditions to compare some aspects of BST and DEB approaches. Due to the large number of parameters to be estimated in the BST model, we applied powerful parameter identification techniques. Both models fitted equally well, but the BST model employs more parameters. The DEB model uses similarities of processes under growth and no-growth conditions and under aerobic and anaerobic conditions, which reduce the number of parameters. This paper discusses some future directions for the integration of knowledge from these two rich and promising areas, working top-down and bottom-up simultaneously. This middle-out approach is expected to bring new ideas and insights to both areas in terms of describing how living organisms operate. Keywords: metabolic networks, optimization, dynamic modeling, lactic acid bacteria
1 Introduction

Mathematical modelling of cellular and molecular processes is gaining increasing attention in recent years. The availability of high-throughput data and the development of new techniques and algorithms lead to the re-emergence of systems biology as a key area to understand organisms as a whole [12].

In this regard, new developments in molecular and cellular biology are shedding some light on how cells work in vivo and which molecular processes are present. It is now possible to understand part of the regulatory processes for some organisms and the networks of interactions, such as protein-protein and gene regulatory networks [2].

Although much effort is being made to analyse and integrate information from different techniques and levels of cellular and molecular organisation, this is still a challenging task. For example, too little is known about all interactions and the dynamics of all metabolites to allow for straightforward modelling. This includes dynamic models for metabolic networks, which represent the set of chemical processes and reactions occurring in a cell. Yet such models can have substantial impact in areas such as the food and pharmaceutical industries, biotechnology and medicine.

This type of modelling is a field in rapid progress and in which very different approaches are used. Depending on the level of detail considered, it is possible to use stochastic discrete models, for instance, to allow for collisions between molecules in microscopic systems with a finite number of particles. For systems with a high number of molecules, formalisms based on deterministic and continuous models are usually preferred. In this regard, metabolic network models typically consist of a set of coupled non-linear ordinary differential equations (ODE’s) where the state variables represent metabolite concentrations. The estimation of the high number of kinetic parameters of these equations from experimental data, typically multivariate time series, represents a major challenge; the parameter identification problem requires intensive search methods in high dimensional space due to the complex behaviour of the numerical error function [17, 1, 7]. Deviations from model predictions can originate from wrong parameter values, but also from an inadequate model structure. A complete understanding of the driving forces behind the processes in a cell is unknown and even the nature of some processes is not fully understood. The use of concentrations of metabolites in ODE’s presumes spatial homogeneity, while cells are actually not homogeneous at all. This contributes to the approximative nature of models for metabolic networks implying that the quantitative characterization of biochemical systems still remains an open problem.

Biochemical Systems Theory (BST) [22, 23, 36] provides a mathematical framework for modeling biological networks. Under BST, the time evolution of metabolite concentrations is modeled with systems of coupled non-linear ODE’s with a specific structure. The source and sink terms in the equations for fluxes of metabolites are products of power-laws of concentrations of metabolites. The rationale for this approach is based on the linear approximation of these terms by a first order Taylor series in logarithmic space.

Dynamic Energy Budget (DEB) theory [14] uses as primary focus the level of the individual, which can be a whale, a tree or a bacterial cell. Populations are treated as sets of interacting individuals and their dynamics typically involve transport of resources in the
environment (cq medium). Most experimental data in molecular biology relate (sometimes indirectly) to the population level, not to the sub-cellular level as such (including the example we will discuss here). DEB theory is based on several formalised assumptions [27], using mass and energy conservation explicitly and respecting stoichiometric constraints on production. These constraints are far from trivial because DEB theory allows for changes in the chemical composition of the individual. DEB theory uses several homeostasis concepts. Strong homeostasis means that metabolic pools, reserve(s) and structure(s), do not change in chemical composition; weak homeostasis means that the individual as a whole does not change in chemical composition during growth in constant environments (see [28], this volume). Contrary to BST models, DEB models do not follow the fate of any particular metabolite, but these homeostasis concepts restrict to possible dynamics of such metabolites considerably. DEB theory does not make use of the concept concentration of metabolites in an individual, as opposed to BST. Transport is linked to surface areas (membranes) and maintenance to (structural) volume, so changes in shape of the individual affect the kinetics of metabolites. Metabolic switches, such as cell division, are linked to the level of maturity. Maturity maintenance is proportional to the level of maturity, somatic maintenance is proportional to the amount of structure. Many aspects of the uptake and use of food (i.e. organisms) by animals are captured realistically and accurately by this standard DEB model. This model (the focus of most contributions to this theme-issue) has three state variables, one reserve, one structure and maturity, and 12 parameters. The explanation for the wide applicability of this model is that the complex food of animals couples the uptake of all substrates they require; animals are adapted to this coupling and acquired a relatively high level of homeostasis. DEB models for most other organisms, however, such as bacteria, fungi, algae and plants, have multiple reserves; the uptake of the various substrates and nutrients from the environment is mutually independent for them. As long as a single substrate or nutrient is limiting growth (see [21] and [16], in this volume), it frequently suffices to follow one reserve only, but such organisms remain multiple reserve systems. A large family of extending modules has been developed to add particular details, such as changes in shape, food selection, co-metabolism, interaction and adaptation.

The present paper briefly reviews the rationale behind BST and DEB models for microorganisms and compares their performance and accuracy in a case-study on the glycolysis in Lactococcus lactis.

2 Biochemical Systems Theory

A broad class of metabolic network models considers systems with a large number of interacting metabolites, which can approximatively be described through deterministic and continuous formalisms. In this context, changes in metabolite concentrations in a cell, here called \( X = \{X_i(t)\}, i = 1, \ldots, n \), are usually modelled with coupled non-linear differential equations:
\[ \frac{d}{dt} X_i = F_i(X,t; u(t), p) \]  

where \( u(t) \) are input signals (such as substrate or nutrient concentrations in the environment) and \( p \in \mathbb{R}^P \) are the parameters of the model. The functions \( F_i \) can be specified in several ways, giving rise to distinct methodologies and formalisms. One option is to use a bottom-up or mechanistic approach, where individual processes are gathered using, for example, mass action or Michaelis-Menten kinetics. Cooperativity or switch-like kinetics associated with regulatory events have been largely described in a phenomenological way by the Hill equation.

As an alternative, approximate formalisms have been developed to describe biochemical processes when their underlying mechanisms are unknown. This corresponds to define function \( F \) as a sum of source and sink terms and approximate all these terms by the same function for all metabolites, whose terms only differ in parameter values. From these approximated formalisms, the most well-known are the power-law within the Biochemical Systems Theory [22, 23, 36], the linlog/log(linear) approximation [8, 9], and, more recently, the saturable and cooperative formalism [26]. The advantage of using structured formalisms, besides their mathematical clearness, is related with their ability to provide a system level analysis environment and the possibility to develop specifically tailored optimisation algorithms and methods.

Biochemical Systems Theory (BST) is a powerful framework for systems analysis of biochemical processes [23, 22, 36]. During the past decades, it has been successfully applied to a wide range of problems and has proven to be a mathematical formalism that can adequate describe and predict complex non-linear behaviour.

In BST each source or sink term \( V \) is approximated by a power-law applied to concentrations of metabolites, which derives from the first order Taylor series expansion in log-space. In this approximation the logarithm of each flux, \( \log V \), is a function of the logarithm of all the metabolite concentrations in the system, \( \log X_j \), with \( j = 1, \ldots, n \), in a neighbourhood of a given operating point \( (X_{10}, X_{20}, \ldots, X_{n0}, V_0) \):

\[
\log V = \log V_0 + \sum_{j=1}^{n} \frac{\partial \log V}{\partial \log X_j} \bigg|_{X_j=X_{j0}} (\log X_j - \log X_{j0})
\]

\[
V = V_0 \prod_{j=1}^{n} \left( \frac{X_j}{X_{j0}} \right)^{\frac{\partial \log V}{\partial \log X_j}} \bigg|_{X_j=X_{j0}}
\]

\[
= \gamma X_1^{f_1} X_2^{f_2} \cdots X_n^{f_n} = \gamma \prod_{j=1}^{n} X_j^{f_j}
\]

where

\[
f_j = \frac{\partial \log V}{\partial \log X_j} \bigg|_{X_j=X_{j0}} = \frac{X_j}{V} \frac{\partial V}{\partial X_j} \bigg|_{X_j=X_{j0}}
\]

\[
\gamma = V_0 X_1^{-f_1} X_2^{-f_2} \cdots X_n^{-f_n}
\]
The parameters $f_j$ represent the kinetic orders of the reaction, which may take any real number, and $\gamma$ is the rate constant of the reaction, which can take only positive values. It is important to note that these parameters refer to particular points $X_{j0}$ and $V_0$, which result from the Taylor series approximations. This means that the approximation is good near that operating point but loses accuracy if the system deviates too far from this region.

In practice, these parameters are inferred as a whole, lumping all the contributions of all the reference values. This means that one cannot, in practice, distinguish each component, estimating directly the $f_j$ and $\gamma$ parameters.

Under this formalism, the time evolution $dX_i/dt$ of metabolite $i$ concentration $X_i$ depends on $P$ fluxes $V_{ik}$ as defined above. This approach leads to a system of differential equations, usually called Generalized Mass Action (GMA):

$$
\frac{dX_i}{dt} = \sum_{k=1}^{P} V_{ik} = \sum_{k=1}^{P} \pm \gamma_{ik} \prod_{j=1}^{n} X_{j}^{f_{ijk}}
$$

(5)

The principle of mass action takes meeting frequencies (and so transformation rates) between two types of molecules proportional to the product of their concentrations, using a diffusion argument. The approach taken by BST can be seen as a variation on this principle.

The sources and sinks can be grouped together, i.e., the set of fluxes producing a metabolite and the set of fluxes consuming it can be aggregated in a power-product term. The dynamics of the system is thus represented in the form leading to a set of ODE’s that is called an S-system:

$$
\frac{dX_i}{dt} = \alpha_i \prod_{j=1}^{n} X_{j}^{g_{ij}} - \beta_i \prod_{j=1}^{n} X_{j}^{h_{ij}}
$$

(6)

where the $\alpha_i$’s and $\beta_i$’s are rate constants. Their dimensions vary according to the number of metabolites involved and depend on the reference values, as stated above. The parameters $g_{ij}$ and $h_{ij}$ represent the kinetic orders.

S-systems representation has several advantageous features. It is mathematically structured, which means that each equation has the same form. This fact can be explored in the effective simulation of the system [11]. S-systems also yield closed-form analytical steady-state solutions that can be enumerated [24] and algorithms tailored to their inference from experimental data were developed [32, 33]. However, mass conservation must be imposed by adding additional parameter constraints or restricting the solution space. If the topology of the network is known, the fluxes can be more adequate described using GMA, where mass conservation is directly implicit in the equations.

In BST, if a particular metabolite does not partake in a source or sink, it has kinetic order zero. The kinetic order is positive in the case of activation, negative in case of inhibition. The choice of the metabolites $X_j$ to include in each reaction is motivated by biochemical information. In fact, when the topology of the metabolic network is known, each reaction can be expressed as a function of the metabolites that influence it. BST is
a formalism that has proven to be flexible and can capture a rich dynamical behaviour of metabolic systems, as explored elsewhere [36].

3 DEB theory for subcellular organisation

To see the link with the standard DEB model (with which most contributions in this issue deal), it helps to realise that DEB models for dividing unicellular organisms represent both a simplification and an extension. Biomass is (formally) partitioned in one structure and a number of reserves that is typically equal to the types of substrates (cq nutrients) that are taken up independently from the environment; substrates are converted to reserves and mobilised reserves are used for all other metabolic needs (maintenance, growth, maturation). The metabolic pools (reserves, structure) are supposed to have a constant chemical composition (strong homeostasis). Cells growing in (chemically) constant environments do not change in composition (weak homeostasis). The weak homeostasis requirement fully specifies the mobilisation fluxes [27]. After subtraction of the reserve-specific somatic maintenance costs, the remaining fluxes are converted to structure using the merging rules of the kinetics of Synthesizing Units (SUs) [13], which accounts for fixed stoichiometries. The metabolites that are rejected by the growth-SUs return to the original reserves or are excreted into the environment (possibly in transformed form).

A direct consequence of strong homeostasis is that any particular chemical compound, such as an enzyme or metabolite, is a fixed fraction of structure and/or any of the reserves. The relative sizes of these pools depend on the growth rate of the cell. Changes in cells chemical composition as a function of the growth rate reveal the chemical composition of pools. Since rRNA per dry weight increases with the growth rate (in a particular way) in energy limited cultures, most rRNA belongs to the energy reserve [38]; compounds in reserve can have active metabolic functions. Since DNA per dry weight decreases with the growth rate (in eukaryotes), DNA belongs to the structure [14]. Strong homeostasis is obviously a simplified and idealized assumption, but many other model approaches make this assumption implicitly. Metabolic Control Analysis (MCA) [10], for instance, assumes that enzyme concentrations are parameters that can be manipulated by enhancing gene performance. Dilution by growth is typically not included by MCA, and a single metabolic network is considered, excluding the synthesis of participating enzymes. BST [36] does not assume a priori any constraints on the parameter space and the solutions, obtained by optimization procedures, should be further checked for consistency, as to represent thermodynamically meaningful systems.

Many bacteria grow as rods that increase in length only; this causes a particular change in shape during growth that depend on the aspect ratio of the cell at division: the ratio of the width and the length of the cell. Kooijman [14] showed that rods represent a static mixture of V0- and V1-morphs. Surface area increases proportional to volume to the power zero (so it remains constant) in V0-morphs, and to the power 1 in V1-morphs; the caps of a rod act as V0-morph, the rest as V1-morph. Isomorphs would be \( V_2^3 \)-morphs in this nomenclature. Since transport (such as uptake) is linked to surface, and maintenance to (structural) volume, V1-morphs have no intrinsic size control; they continue growing as...
long as substrate allows. Cell volume increases exponentially in time at constant substrate
density. V0-morphs, on the other hand, rapidly balance uptake with maintenance and
growth rapidly declines during the cell cycle. The shape of the growth curve of cells is thus
very sensitive for the aspect ratio, which is exactly what has been found empirically [14,
Fig 4.11]. The growth curve of individual cocci (short rods) rapidly satiates before division.
At division, surface area increases rapidly. The total surface area in a population of cells
increases with the number of cells, so the population as super-individual acts as a V1-
morph. If the aspect ratio is very small (and caps of rods hardly matter), the individuals
act as V1-morphs as well, and so the organisation level of the individual completely drops
out and measurements at the population level (as typically done in molecular biology)
directly match the sub-cellular level. This is very different in the case the aspect ratio is
not small; the relationship between the levels of organisation is much more complicated.
Even if population growth is constant (e.g. chemostats in equilibrium), the growth rate of
individual cells jumps up and down during its cycle. If we are interested in sub-cellular
phenomena (such as the dynamics of metabolic networks), the cell cycle generally matters.

DEB theory deals with surface (for transport) and volume (for maintenance), not with
length directly. The quantifier for reserve mobilisation in the DEB standard model, the
energy conductance $\dot{v}$, has dimension length per time (see [28], this volume). This length is
in fact the ratio between volume and a surface area. A subtlety is worth mentioning here,
that is easier to see for isomorphs than for V1-morphs. Reserve mobilisation follows from
weak homeostasis [27]; a reasonable mechanism of this particular dynamics takes mobili-
sation rate proportional to the interface between reserve and structure [15, 14]. Reserve
and structure are segregated at the molecular level, an essential component of DEB theory.
Although the reserve density frequently features in DEB theory as a variable, it is defined
as a ratio of two amounts (reserve and structure), and is not conceived as a concentration.
The latter implies well-mixedness at the molecular level and this state is behind the law
of mass action. Since surface area and volume are proportional in V1-morphs, $\dot{v}/L$ for iso-
morphs should be replaced by $k_E$ for V1-morphs. Likewise $\{\dot{F}_m\}L^2$ for isomorphs should
be replaced by $[\dot{F}_m]L^3$ for V1-morphs, and $\{\dot{J}_{EAm}\}L^2$ by $[\dot{J}_{EAm}]L^3$.

Since the roles of surfaces and volumes have become identical for V1-morphs, it is no
longer handy to work with length, and we better remove length from the specification of
changes for state variables. This is simple, thanks to the strong homeostasis assumption.
E.g. the searching rate $[\dot{F}_m]L^3$ is replaced by $\dot{f}_mM_V$, where $\dot{f}_m$ is the mass-specific searching
rate (better called specific substrate affinity in this context); $[\dot{J}_{EAm}]L^3$ is replaced by
$\dot{j}_{EAm}M_V$, where $j_{EAm}$ is the mass-specific maximum assimilation. The surface-specific
maintenance costs (e.g. for osmotic work) absorb into the volume-specific maintenance
cost.

DEB theory states that a cell divides if maturity exceeds a threshold; maturity is reset
at division. Maturity represents information and has no mass or energy. Allocation of
reserve to maturity maintenance plus maturation is a fixed fraction of the mobilised flux,
and maturation maintenance is proportional to the maturity. Since maturity represents
a level of metabolic learning, only energy reserves matter. Maturity density (the ratio of
maturity and the amount of structure) typically varies little during a cell cycle. This is why
maturity and somatic maintenance costs can typically be added and taken proportional to
the amount of structure; cells then divide when the amount of structure exceeds a threshold value. Changes in maturity can be important to capture cell cycle phenomena, and how cell size at division depends on the nutritional status, but for many applications these are only details. The fraction allocated to maturation then becomes irrelevant.

Growth of a V1-morph on a single limiting substrate can then be captured by 2 state variables (reserve and structure) and 6 parameters (specific affinity $f_m$, specific maximum assimilation rate $j_{E Am}$, yield of reserve on substrate $y_{EX}$ and of structure on reserve $y_{VE}$, specific maintenance costs $j_{EM}$, reserve turnover rate $k_E$). The mass-specific maintenance costs $j_{EM}$ applies as long as the mobilisation rate allows. If not, the remaining costs are supplemented from structure by shrinking, which involves the parameter $j_{VM}$. If reserve is in fact internalised substrate (or nutrient), we simply have $y_{EX} = 1$.

The specific reserve mobilisation rate of a V1-morph works out as $j_{EC} = j_{EC}/M_V = j_{E Am}(e - \dot{r}/\dot{k}_E)$, where $\dot{r}$ is the specific growth rate $\frac{d}{dt} \ln M_V$ and $e$ the scaled reserve density $e = m_E/m_{Em} = m_E k_E/j_{E Am}$. The change in scaled reserve density $e$ and structure $M_V$ is given by

\[
\frac{d}{dt} e = (f - e)k_E \quad \text{with} \quad f = \frac{X}{X + K} \quad \text{and} \quad K = \frac{j_{E Am}}{y_{EX} f_m} \quad (7)
\]

\[
\frac{d}{dt} M_V = \dot{r} M_V \quad \text{with} \quad \dot{r} = \dot{k}_E \frac{m_E k_E - j_{EM} - y_{EV} j^M_V}{m_E + y_{EV}} \quad \text{and} \quad (8)
\]

This simplest formulation for the payment of maintenance costs, where reserve has absolute priority above structure as substrate for maintenance, is a special case of a more general formulation where priority is less extreme and can be set by a preference parameter, see [30].

Just as in the standard DEB model, we have 3 organising fluxes, assimilation (substrate $X$ is converted to reserve $E$ and products $P$ with rate $J_{X A}$), dissipation (reserve $E$ is converted to products $P$ at rate $J_{EM}$) and growth (reserve $E$ is converted to structure $V$ and products $P$).

For $n$ independent substrates we need $n + 1$ state variables ($n$ reserves $E_i$ and structure $V$), and $6n + 1$ parameters (specific substrate affinity $f_{m_i}$, maximum specific assimilation rate $j_{E_i Am}$, yield of reserve on substrate $y_{E_i X_i}$, specific maintenance cost $j_{E_i M}$, yield of structure on reserve $y_{VE_i}$, excretion fraction of rejected mobilized reserve $1 - \kappa_{E_i}$ and common mobilisation rate $k_E$) for the baseline model with a single structure.

4 Example: Lactic Acid Bacteria metabolism

We now compare BST and DEB models by fitting them to the same experimental data for *Lactococcus lactis*. This is a Gram-positive bacterium that plays an essential role in the manufacture of fermented dairy products, being the primary model organism for Lactic Acid Bacteria. One of the most studied pathways in *Lactococcus lactis* is glycolysis, which is the sequence of reactions that metabolize one glucose molecule to two molecules of
pyruvate with the concomitant net production of two molecules of ATP. Figure 1 represents a simplified version of glycolysis in \textit{L. lactis}, and also the subsequent pyruvate metabolism.

Figure 1: Pathways of glucose and pyruvate metabolism in \textit{Lactococcus lactis}. The reactions indicated are catalyzed by the following enzymes: 1. phosphoenolpyruvate:phosphotransferase system (PEP:PTS); 2. EMP enzymes; 3. lactate dehydrogenase; 4. pyruvate-formate lyase; 5. pyruvate dehydrogenase; 6. acetaldehyde dehydrogenase and alcohol dehydrogenase; 7. phosphotransacetylase and acetate kinase; 8. \(\alpha\)-acetolactate synthase; 9. \(\alpha\)-acetolactate decarboxylase; 10. 2,3-butanediol dehydrogenase. From [18].

Glycolysis constitutes an excellent test pathway since its metabolic network topology is known for many organisms and it is an ubiquitous process in living cells. However, the dynamical behavior of this system is not yet completely understood.

A model capturing all known details of glycolysis was developed by Teusink and colleagues [29] but it can only be used if a lot of detailed information is available. Contrary to DEB theory, Biochemical Systems Theory (BST) [36] does not assume any specific mechanism for changes in metabolite concentrations, given its approximative nature. Nevertheless the parameters have biochemical interpretation, at least locally, and translate directly to the kinetic orders and rate constants of the corresponding reactions.
In recent years, it is possible to accompany the time evolution of metabolite concentrations in vivo through Nuclear Magnetic Resonance (NMR). This technique allows inspecting directly how living cells metabolize substrates. In these experiments a pulse of labelled glucose is provided to a cell suspension and the labelling is traced throughout the pathway. This means that we only need to consider pathways related to the assimilation process. Such data might provide important insights about metabolic behaviour of living cells.

We analyse the results of 3 experiments with batch cultures of *L. lactis*: resting cells under aerobic or anaerobic conditions, which were prevented to grow due to absence of essential amino acids, as well as a growing culture under anaerobic conditions, where glucose is the limiting energy substrate.

In resting suspensions, meaning no-growth, metabolites were measured using NMR [20], while during growth end-products and glucose were determined in supernatant samples by HPLC. The details of the growth experiment are fully given in [19].

4.1 BST formulation

This section presents models for glycolysis in *L. lactis* using Biochemical Systems Theory. The systems of differential equations comply with the mathematical formalism of power-laws described and the choice of the state variables was made according to the availability of experimental data. This translates into a simplified model for glycolysis where some reactions are grouped together.

4.1.1 No-growth under aerobic and anaerobic conditions

Departing from the topology of the glycolytic pathway described above and represented in Figure 1, the BST specification collects the terms involved in each reaction using power-laws.

The fitted BST model relates the concentrations of glucose $G$, glucose-6-phosphate G6P, fructose-1,6-bisphosphate FPB, common pool of 3-phosphoglycerate PGA and phosphoenolpyruvate PEP, pyruvate $P$ and lactate $L$. In these experiments, the measured extracellular metabolites are $G$ and $L$, and the intracellular metabolites are G6P, FPB, PGA, PEP and $P$.

According to the structural analysis performed [34], showing that the reversible reaction between PGA and PEP is extremely fast, both pools were merged into a unique state variable, PGAPEG = PGA + PEP. In fact, these two metabolites can be considered at equilibrium, which inevitably hampers the correct identification of the parameters of this reversible reaction. This is also justified by the biochemical data that support the hypothesis of an invariant ratio between the concentrations of these two pools. By defining this ratio as $k_{45}$, i.e. $PGA = k_{45} PEP$, the equations to compute the individual concentrations from the common pool of PGAPEG become: $PGA = k_{45} PGAPEG/(1 + k_{45})$ and $PEP = PGAPEG/(1 + k_{45})$, respectively.

The model for glycolysis is described by the following system:
\[
\frac{d}{dt} G = -k(1 + \alpha t^\beta) G
\]
\[
\frac{d}{dt} G_{6P} = \beta_1 G^{h_{11}} G_{6P}^{h_{12}} PEP^{h_{25}} - \beta_2 G_{6P}^{h_{22}} ATP^{h_{2ATP}}
\]
\[
\frac{d}{dt} FPB = \beta_2 G_{6P}^{h_{22}} ATP^{h_{2ATP}} - \beta_3 FPB^{h_{33}} P_i^{h_{33}} Pi - NAD^{h_{3NAD}}
\]
\[
\frac{d}{dt} PGAPEP = 2\beta_3 FBP^{h_{33}} P_i^{h_{33}} Pi - NAD^{h_{3NAD}} - \beta_1 G^{h_{11}} G_{6P}^{h_{12}} PEP^{h_{25}} + \beta_5 FBP^{h_{513}} P_i^{h_{513}} P_i - \beta_52 PEP^{h_{525}}
\]
\[
\frac{d}{dt} P = \beta_1 G^{h_{11}} G_{6P}^{h_{12}} PEP^{h_{25}} + \beta_5 FBP^{h_{513}} P_i^{h_{513}} P_i + \beta_61 FBP^{h_{616}} NAD^{h_{61NAD}} - \beta_62 P^{h_{626}}
\]
\[
\frac{d}{dt} L = \beta_61 FBP^{h_{616}} NAD^{h_{61NAD}}
\]
\[
\frac{d}{dt} A = \beta_7 P^{h_{626}}
\]

The rationale behind the choice of this simplified model is directly linked to the known topology of the network. As an example, Figure 1 shows that the degradation of G6P (and the production of FBP) depends on the G6P concentration and the amount of ATP. Therefore, the structure of the rate law describing that particular reaction is based on the product of the intertwining metabolites, in this example \(\beta_2 G_{6P}^{h_{22}} ATP^{h_{2ATP}}\).

Not all the measured variables are considered in the model. In fact, ATP, NAD\(^+\), NADH and inorganic phosphate (\(P_i\)) are not state variables of this system (Eq. 9) due to the lack of information regarding the complete set of reactions they are involved in. In fact, these metabolites are ubiquitous in the cells and participate in many pathways, which hampers their detailed description. For this reason, they are considered as input variables.

The parameters were estimated using the method described in [34, 37] for the three experiments separately, using the method of least squares. Figure 2 shows the fits of the system using the parameter values as reported in Tables 1 and 2.

There are some noteworthy results regarding this system. The accumulation of PGA and PEP is a characteristic of starved cells. The feed-forward activation of FBP can explain this behavior [35] from a mathematical point of view. The accumulation of PEP allows the starved cells to assimilate glucose almost immediately, upon availability, since the main transport systems uses PEP, through direct phosphorylation of glucose to G6P, using the phosphoenolpyruvate:sugar-phosphotransferase system (PEP:PTS). This was shown to confer robustness to cells. Overall, this model is capable of describing the experimental data accurately.

### 4.1.2 Growth under anaerobic conditions

When modelling the growth experiment using the BST approach, some alterations must be made in the equations due to limitations of data availability. In the previous no-growth
Table 1: The BST estimated parameter values of model (9) for the glucose metabolism (Figure 1). The corresponding simulations are represented in Figure 2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Aerobic</th>
<th>Anaerobic</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>0.0530251</td>
<td>0.124738</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>0.0419958</td>
<td>0.134194</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$\beta$</td>
<td>2.68092</td>
<td>2.6674</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_1$</td>
<td>7.20321</td>
<td>5.86735</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$h_{11}$</td>
<td>0.997546</td>
<td>1.25193</td>
<td>-</td>
</tr>
<tr>
<td>$h_{12}$</td>
<td>-1.48643</td>
<td>-1.06961</td>
<td>-</td>
</tr>
<tr>
<td>$h_{25}$</td>
<td>0.38576</td>
<td>0.288687</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_2$</td>
<td>0.345889</td>
<td>0.379794</td>
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<td>$h_{22}$</td>
<td>1.54399</td>
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<td>-</td>
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<td>$h_{2\text{ATP}}$</td>
<td>1.51599</td>
<td>0.26203</td>
<td>-</td>
</tr>
<tr>
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<td>0.181082</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$h_{33}$</td>
<td>1.09298</td>
<td>1.02783</td>
<td>-</td>
</tr>
<tr>
<td>$h_{3\Pi}$</td>
<td>0.258372</td>
<td>-0.137778</td>
<td>-</td>
</tr>
<tr>
<td>$h_{3\text{NAD}}$</td>
<td>-0.0966562</td>
<td>0.174484</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_{32}$</td>
<td>0.134164</td>
<td>0.000447956</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$h_{525}$</td>
<td>0.0940446</td>
<td>0.103001</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_{51}$</td>
<td>0.862421</td>
<td>0.683548</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$h_{513}$</td>
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<td>0.854603</td>
<td>-</td>
</tr>
<tr>
<td>$h_{515}$</td>
<td>0.0382342</td>
<td>0.0921911</td>
<td>-</td>
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<td>$h_{51\Pi}$</td>
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<td>-0.279396</td>
<td>-</td>
</tr>
<tr>
<td>$\beta_{61}$</td>
<td>0.0324743</td>
<td>0.0306174</td>
<td>min$^{-1}$</td>
</tr>
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<td>$h_{613}$</td>
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</tr>
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<td>$h_{61\text{NAD}}$</td>
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<td>-</td>
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<td>$\beta_{62}$</td>
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<td>1.915812</td>
<td>min$^{-1}$</td>
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<td>$h_{626}$</td>
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<td>-</td>
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<tr>
<td>$\beta_7$</td>
<td>0.0386</td>
<td>0</td>
<td>min$^{-1}$</td>
</tr>
<tr>
<td>$k_{45}$</td>
<td>2.04035</td>
<td>1.54695</td>
<td>min$^{-1}$</td>
</tr>
</tbody>
</table>
Figure 2: Experimental time trajectories of external and internal metabolites, energy compounds and biomass dry-weight, with prediction by the BST formulation. The simulations of the predicted solution correspond to the integration of Equations 9 with parameters of Tables 1 and 2. (a) No-growth, aerobic conditions; (b) No-growth, anaerobic conditions; (c) Growth, anaerobic conditions.

experiments, time series were available for $P_i$, NAD$^+$, NADH and ATP, which allowed the inclusion of these variables as input functions in the model. However, in this experiment, these variables were not measured, which hampers inevitably the utilization of the same system of differential equations. For this reason, the model for growth should necessary be adjusted to exclude non-measured input variables.

The equations for this growth experiment relate the concentrations of extracellular metabolites glucose ($G$), acetate ($A$), lactate ($L$), ethanol ($E$), formate ($F$) and biomass ($V$):

$$\frac{d}{dt} G = -k(1 + \alpha t^\beta)G$$
$$\frac{d}{dt} A = \alpha_2 G^{g_{31}} A^{g_{42}}; \quad \frac{d}{dt} E = \alpha_4 G^{g_{41}} E^{g_{44}}; \quad \frac{d}{dt} F = \alpha_5 G^{g_{51}} F^{g_{55}}$$
$$\frac{d}{dt} L = \alpha_3 G^{g_{31}} L^{g_{33}}; \quad \frac{d}{dt} V = \alpha_6 G^{g_{61}} V^{g_{66}}$$

The rationale behind this representation is to consider, as previously, that glucose decay depends on its concentration and a time acceleration, and the other fluxes $dX_i/dt$ are functions of glucose concentration and their own concentration $X_i$.

The solution was obtained through the use of several optimization algorithms implemented in the Systems Biology Toolbox 2 (SBTOOLBOX2) for MATLAB [25]. In partic-
Table 2: The BST parameter values of growth model (Equation 10). The simulations of the corresponding model are represented in Figure 2.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Growth</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k$</td>
<td>0.036084</td>
<td>h(^{-1})</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>0.0012835</td>
<td>h(^{-\beta})</td>
</tr>
<tr>
<td>$\beta$</td>
<td>6.0178</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha_2$</td>
<td>1.0054</td>
<td>h(^{-1}) mM(^{-1})</td>
</tr>
<tr>
<td>$g_{21}$</td>
<td>1.0265</td>
<td>-</td>
</tr>
<tr>
<td>$g_{22}$</td>
<td>1.3567</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha_3$</td>
<td>0.04975</td>
<td>h(^{-1}) mM(^{-1})</td>
</tr>
<tr>
<td>$g_{31}$</td>
<td>0.53391</td>
<td>-</td>
</tr>
<tr>
<td>$g_{33}$</td>
<td>1.2574</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha_4$</td>
<td>0.16885</td>
<td>h(^{-1}) mM(^{-1})</td>
</tr>
<tr>
<td>$g_{41}$</td>
<td>0.46223</td>
<td>-</td>
</tr>
<tr>
<td>$g_{44}$</td>
<td>0.77257</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha_5$</td>
<td>0.13177</td>
<td>h(^{-1}) mM(^{-1})</td>
</tr>
<tr>
<td>$g_{51}$</td>
<td>0.57726</td>
<td>-</td>
</tr>
<tr>
<td>$g_{55}$</td>
<td>1.1216</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha_6$</td>
<td>0.1931</td>
<td>h(^{-1}) mM(^{-1})</td>
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<tr>
<td>$g_{61}$</td>
<td>0.36009</td>
<td>-</td>
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<td>$g_{66}$</td>
<td>1.0102</td>
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</tr>
<tr>
<td>$G(0)$</td>
<td>50.51</td>
<td>mM</td>
</tr>
<tr>
<td>$A(0)$</td>
<td>2.91648e-006</td>
<td>mM</td>
</tr>
<tr>
<td>$L(0)$</td>
<td>1.77998</td>
<td>mM</td>
</tr>
<tr>
<td>$E(0)$</td>
<td>4.36213e-008</td>
<td>mM</td>
</tr>
<tr>
<td>$F(0)$</td>
<td>0.0157067</td>
<td>mM</td>
</tr>
<tr>
<td>$V(0)$</td>
<td>0.0203951</td>
<td>g DW/L</td>
</tr>
</tbody>
</table>

Since some of the initial conditions are not measured, namely for Acetate, Ethanol and Formate, the algorithms also estimate them. This avoids numeric problems of assuming zero concentrations, which would not allow the current system to progress. Furthermore, it is expected that cells have minimum quantities of these metabolites in vivo, which makes this procedure more realistic too. Therefore, the obtained solution for the parameters also estimates the initial values $A(0)$, $E(0)$ and $F(0)$ along with $G(0)$, $L(0)$ and $V(0)$ to allow for more flexibility in the optimization step and translating the initial uncertainty associated with these state variables.

The estimated parameters and initial conditions are specified in Table 2. The simulation of this solution is presented in Figure 2c.

Alternative models with different structure where also analyzed and tested. For example, the decay of glucose can be argued to depend on the total amount of biomass $V$, $dG/dt = -k(1 + \alpha t^\beta)GV^{\alpha_0}$. However, this model led to an estimated low value for
$\alpha_0 = 0.00666$ and a less accurate fitting. For this reason, and in order to simplify as much as possible the system, this value was put to zero, without affecting the overall behaviour of the simulation. Another interesting characteristic is that all kinetic orders $g_{ii}$ are positive, which can be argued to be counterintuitive. The end product should inhibit its own production, which would correspond to $g_{ii} < 0$. This behaviour was, however, not observed using these equations. One possible explanation might be the fact that some other missing state variables, if included, could change the sign of these parameters.

### 4.2 DEB formulation

We applied DEB theory for V1-morphs with 1 reserve and 1 structure and one extra route for aerobic glucose uptake. Contrary to the BST formulation, we here capture all three experiments simultaneously by imposing the constraint that comparable parameter values are equal in all three experiments. This leads to a substantial reduction of the number of free parameters. As stated before dry weight (total biomass) has contributions from reserve and structure.

DEB theory assumes explicitly that substrate uptake only depends on substrate availability and amount of structure, not on the amount of reserve, so also not on the nutritional conditions.

The experimental no-growth conditions concern growth limitations of (mineral) substrates that are not measured. Glucose and its products relate to reserves that are non-limiting. In the growth-condition, these limiting nutrients have been added to the medium *ad libitum*. In this case, glucose is the limiting substrate, and the non-limiting reserves hardly matter. This latter follows from the dynamics of synthesising units for growth on multiple complementary reserves.

We further assume that one uptake route for glucose is not active anaerobically or, at least, has a negligible contribution. In fact, there are three known routes for glucose uptake in *L. lactis* and they are all active under anaerobiosis [6]. Our results suggest that two of these routes have the same kinetic properties and can be lumped in the model.

We also observe that the population has an adaptation period in the aerobic case, which is negligible in the two anaerobic situations. The DEB theory has no explanation for this, and it might be related to the pre-treatment of the population.

We include this process using an *ad hoc* formulation where inhibition of substrate uptake decays exponentially.

The used DEB model reads:

\[
\frac{d}{dt} G = (1 - \exp(-\dot{h}t)) V(\ddot{J}_{GAm} f + \dddot{J}_{GAm} f_a) \quad \text{with } f = \frac{G}{K + G} \quad \text{and } f_a = \frac{G}{K_a + G}
\]

\[
\dot{h} = \infty \quad \text{and } \ddot{J}_{GAm} = 0
\]

\[
\frac{d}{dt} FBP = -y_{FBP,G} \frac{d}{dt} G - \dot{k}_{FBP} FBP
\]

\[
\frac{d}{dt} PEP = -y_{PEP,G} \frac{d}{dt} G - \dot{k}_{PEP} PEP
\]

\[
\frac{d}{dt} A = -y_{AG} \frac{d}{dt} G; \quad \frac{d}{dt} E = -y_{EG} \frac{d}{dt} G; \quad \frac{d}{dt} F = -y_{FG} \frac{d}{dt} G
\]

(11)
\[
\frac{d}{dt} L = -y_{LG} \frac{d}{dt} G + y_{L,PEP} \dot{k}_{PEP} \text{PEP}; \quad PGA = y_{PGA,PEP} \text{PEP}
\]

\[
\frac{d}{dt} V = \dot{r} V \quad \text{with} \quad \dot{r} = \dot{k}_E \frac{e - l_d}{e + g}
\]

\[
\frac{d}{dt} e = (f_a - e) \dot{k}_E; \quad W = d_V V(1 + \omega_d e)
\]

where \( V \) is (structural) biovolume, \( G \) is glucose, \( L \) lactate, \( A \) acetate, \( E \) ethanol, \( F \) formate, \( \text{FBP} \) fructose-1,6-bisphosphate, \( \text{PEP} \) phosphoenolpyruvate, \( \text{PGA} \) 3-phosphoglycerate, \( e \) scaled reserve density, \( f \) and \( f_a \) scaled functional responses, \( V \) biomass dry weight, \( \dot{r} \) the specific growth rate, where \( \dot{r} = 0 \) for the two no-growth experiments.

Notice that the growth experiment is in anaerobic conditions, so in this case, a single uptake route is used \( (f = 0) \). In the no-growth experiments, we have \( r = 0 \), and \( V \) and \( e \) are constant. Compounds \( A, E, F, L, \text{PEP} \) and \( \text{FBP} \) are treated as assimilation products, while \( A, E, F, \) and \( L \) are excreted into the medium. Some \( L \) is also formed from \( \text{PEP} \) under aerobic conditions. \( \text{PGA} \) is taken to be proportional to \( \text{PEP} \), implying that they are interconverted.

The intra-cellular products \( \text{FBP} \) and \( \text{PEP} \) appear to decay slowly. DEB theory has no direct explanation of this; it doubtlessly links to other components of the metabolism, and perhaps to the problem that maintenance continues, while energy substrates are absent. Too little is known for a more satisfying inclusion of this process, so we use an \textit{ad hoc} first order decay, which does not capture the \( \text{FBP} \) decay under anaerobic conditions very well.

The parameters have the following interpretation: \([\dot{J}_{GAm}]\) and \([\dot{J}_{GAm}^a]\) are the aerobic and anaerobic volume-specific maximum glucose uptake rates, \( K \) and \( K_a \) the aerobic and anaerobic half-saturation constants, \( \dot{h} \) the decay rate of (aerobic) inhibition, \( y_{*1,*2} \) the yield of \(*_1\) on \(*_2\) (which might be different under aerobic and anaerobic conditions), \( \dot{k}_s \) the decay rate of internal metabolite \(*\), \( \dot{k}_E \) the specific conductance, \( g \) the energy investment ratio, \( l_d = \dot{k}_M g / \dot{k}_E \) where \( \dot{k}_M \) is the maintenance rate coefficient, \( d_V \) the specific density of structural biomass, \( \omega_d \) the proportionality constant for the contribution of the scaled reserve density to dry weight.

We fitted all 16 curves of the 3 experiments simultaneously because the parameters of anaerobic glucose uptake occur in all experiments, and the anaerobic yields of product occur in the 2 anaerobic experiments. We used a simplex method to estimate the parameters, as coded in DEBtool (\url{http://www.bio.vu.nl/thb/deb/deblab/}). The results are presented in Table 3 and Figure 3.

Notice that \( y_{L,PEP}, y_{FBP,G} \) and \( y_{PEP,G} \) need correction for the ratio of bacterial and medium volumes before they make biochemical sense. This is because the data represent concentrations in homogenised medium plus biomass, while \( L \) and \( G \) occur outside the cells, and \( \text{PEP} \) and \( \text{FBP} \) inside the cells. The yield coefficients directly relate to the metabolic scheme of Figure 1 and reveals the relative use of alternative pathways. Contrary to the BST model, these yield coefficients are constant, independent of metabolite concentrations. Since aerobic and anaerobic pathways are qualitatively different, the yield coefficients may differ between these conditions.
Table 3: The DEB parameter values of model (11) that belong to figure 2. The upper panel refers to the two no-growth experiments; the lower panel to the extra parameters for the anaerobic growth experiment.

<table>
<thead>
<tr>
<th>symbol</th>
<th>aerobic sd</th>
<th>anaerobic sd</th>
<th>unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>$V$</td>
<td>2.41 0</td>
<td>2.48 0</td>
<td>ml</td>
</tr>
<tr>
<td>$G(0)$</td>
<td>40 0</td>
<td>40 0</td>
<td>mM</td>
</tr>
<tr>
<td>$K$</td>
<td>24.74 38.84</td>
<td>23.86 7.15</td>
<td>mM</td>
</tr>
<tr>
<td>$[\dot{J}_{GAM}]$</td>
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<td>321.6 50</td>
<td>mM ml$^{-1}$h$^{-1}$</td>
</tr>
<tr>
<td>$y_{L,G}$</td>
<td>1.48 0.03476</td>
<td>1.83 0.0058</td>
<td>mol mol$^{-1}$</td>
</tr>
<tr>
<td>$y_{A,G}$</td>
<td>0.2374 0.0203</td>
<td>0 0</td>
<td>mol mol$^{-1}$</td>
</tr>
<tr>
<td>$\dot{h}$</td>
<td>6.45 5.47</td>
<td>$\infty$ 0</td>
<td>h$^{-1}$</td>
</tr>
<tr>
<td>$y_{PGA,PEP}$</td>
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<td>$y_{L,PEP}$</td>
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<td>0 0</td>
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<tr>
<td>$FBP(0)$</td>
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</tr>
<tr>
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<td>8.286 2.135</td>
<td>1.181 0.191</td>
<td>mol mol$^{-1}$</td>
</tr>
<tr>
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<td>mM</td>
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<tr>
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<td>0.0532 0.0356</td>
<td>mol mol$^{-1}$</td>
</tr>
<tr>
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<tr>
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<td>1 0</td>
<td>ml, -</td>
</tr>
<tr>
<td>$G(0), g$</td>
<td>49.72 1.25</td>
<td>0.093 1.66</td>
<td>mM, -</td>
</tr>
<tr>
<td>$d_v, \omega_d$</td>
<td>2.04 25.81</td>
<td>5.08 88.63</td>
<td>g ml$^{-1}$, -</td>
</tr>
<tr>
<td>$k_E, l_d$</td>
<td>0.9663 2.77</td>
<td>0.0542 1.527</td>
<td>h$^{-1}$, -</td>
</tr>
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<td>$y_{A,G}^a, y_{E,G}^a$</td>
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<td>0.0663 0.047</td>
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<tr>
<td>$y_{F,G}$</td>
<td>0.0729 0.0464</td>
<td></td>
<td>mol mol$^{-1}$</td>
</tr>
</tbody>
</table>
5 Discussion and conclusion

The models obtained using BST and DEB theory serve different purposes and objectives since they are based on different approaches and assumptions.

A relevant difference is that the BST approach tries to follow each chemical compound individually, while the DEB approach works with a reduced number of generalized compounds.

In both formalisms the models obtained are nonsegregated [3]. We consider that the population of cells is homogeneous, i.e., the population is lumped into one biophase interacting with the external environment, and, therefore, can be viewed as one species in solution. The cell concentration is thus described by one variable alone. Both formalisms relate measurements at the population level directly to the sub-cellular level.

The BST model required 80 parameters to describe the results of three experiments (17 curves), while the DEB model used 41 parameters. The number of DEB parameters can be reduced further by assuming $\dot{k}_{PEP}^a = 0$, $FBP(0) = FBP_a(0)$, $PEP(0) = PEP_a(0)$, $K = K_a$. This would hardly affect the goodness of fit. The overkill in the number of BST parameters makes their estimation a challenge. In fact, this formalism can lead to over-parameterized models, which can be tackled by detecting structural identifiability problems. DEB models usually provide better convergence in the optimization step since a simpler error surface is expected.

In BST (using Generalized Mass Action formalism) each flux is modelled individually and, consequently, each rate kinetics description is approximated through a power-law that
depends on the terms involved in that particular reaction. Mass conservation is imposed by specifying equal terms for common fluxes in the equations. In DEB each flux has a simple mechanism that is based on SU-dynamics and (weak) homeostasis. Mass is conserved explicitly.

Regarding the models inferred using the two formalisms, there are issues worth mentioning. The BST model for the growth experiment lets substrate (glucose) disappear as an autonomous process and biomass follows this disappearance; this is a consequence of the absence of different levels of organisation in this formalism. The DEB model for the growth experiment quantifies the uptake of glucose by biomass; the change in total uptake follows from the increase in biomass and uptake as function of substrate concentration; this formulation seems closer to reality. However, both the BST and the DEB model have a problem to capture the sigmoidal decay of glucose in the aerobic no-growth experiment. In both models there was the need to introduce a time variable in the equations, to account for the slow start of glucose uptake by the aerobic non-growing cells. It can simply be interpreted as an acceleration of this initial reaction. In fact, the time derivative of glucose uptake increases in the first period of the experiment, and that was identified independently by both approaches. Biochemically, one possible explanation for this phenomenon is the progressive energization of the cells, which would imply an increasing flux of glucose assimilation. The complete explanation of this behavior is, however, still lacking experimental evidence.

In principle, the detailed BST description provides tools for experimental design, since each parameter has a corresponding biochemical meaning. For example, if an enzyme that affects a particular sink or source term is over-expressed, the kinetic order would become more negative or more positive and/or the rate constant would be affected. Likewise, a knock-down of a gene that codes for such an enzyme would set the kinetic order of the substrate that is catalysed to zero; the rate constant could be affected as well.

The DEB parameters have phenotypic and genotypic aspects; changes in the activity of a particular gene might potentially affect several DEB parameters simultaneously. Such changes should be worked out in comparative experiments, from which parameters are estimated. Extensions of DEB models involving gene expression have been applied successfully to model adaptation and diauxic growth [5, 4].

In DEB equations the production of each chemical species is related with the rate of glucose decay, through biochemical relations with specific yield factors. This allows the decoupling of all the intermediate reactions since there are no cascades, as in the BST model. The main advantage of this procedure is that the model obtained is more parsimonious - the fittings use a less number of parameters - while maintaining the accuracy in describing the experimental data. Furthermore, it was possible, in this setting, to perform simultaneous inference of the aerobic independent experiments (growth and no-growth), obtaining a conjugated model that approximates the three time series. Even if the inherent biochemical processes are completely distinct, mathematical modeling allows to understand them as a whole, which is an interesting feature. However, DEB models can hardly provide answers about the outcome when some alteration of a particular enzyme in the pathway takes place. This prediction procedure might be useful to model e.g. mutant strains, for metabolic engineering purposes.
The detailed description provided by BST might hamper the generalization to other experimental settings, if all the input functions are to be included. In fact, the given model for no-growth experiments cannot be used, in this particular situation, to model the growth data. This is partially due to the lack of information regarding all the metabolites present in the model and also the non-availability of inorganic phosphate $P_i$, redox balance (NAD$^+$ and NADH) and ATP data. For this reason, a reduction of the number of state variables and input functions should be performed prior to any estimation and simulation procedure. Furthermore, one possible drawback of this approach, given its approximative nature, is that it lacks growth as a feedback mechanism.

In conclusion, one can classify the obtained DEB models as nonsegregated and unstructured since the compartments are modelled, not compounds [3]. BST, on the other hand, can provide structured models since each individual reaction and intracellular process can be accounted for.

The fit of the DEB formulation is generally very good, but PEP and PGA start to increase later than predicted and FBP does not decay anaerobically according to a first order process. The reason remains unknown. We did not fit ATP and inorganic phosphate $P_i$ because their trajectories depend on variables that have not been measured; it is unclear why $P_i$ first jumps down and then recovers (it is assimilated in glycolysis and then released).

Another aspect worth mentioning is the FBP decay in the anaerobic non-growing cells experiment, which is not correctly fitted in any of the presented formalisms. In fact, by observing both results, there is a slow down of the flux after some minutes that is not explained by either model. Using BST it was argued that this observation might be due to a structural factor missing in the model and not an optimization problem. For example, the FBP decay can be inhibited by NADH, which might slow down the process. Some other unknown inhibition can be at play, or at least, relations that are not accounted for.

These common and model-independent observations are very interesting since they presumably represent intrinsic properties of the system, giving further insights about the dynamical and topological behavior of the metabolic pathway.

This comparative study between Biochemical Systems Theory (BST) and Dynamic Energy Budget (DEB), each with its own properties and rationale, illustrates current efforts to model metabolic networks. The key question when opting for any formalism is to clearly define the ultimate purpose of a mathematical model. The future perspective of this field can be anticipated to be the strengthening of multidisciplinary efforts, thus promoting strong collaborations between several areas of research.

6 Acknowledgments

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