1 Introduction:

The dynamic response of any system can be modeled as a set of ordinary differential equations (ode) (Equation 1a) in the state variables (\mathbf{x}). The changes in the state variables are expressed as a function of the state variables, the system parameters (θ) and the inputs to the system (u).

$$\dot{\mathbf{x}} = g(\mathbf{x}, \theta, u) \tag{1a}$$

$$\mathbf{y} = h(\mathbf{x}, \mu, u) \tag{1b}$$

When not all states are observable, Equation (1a) is usually augmented with a system that defines the relationship between the observable output variables (\mathbf{y}) and the state variables (\mathbf{x}) as in Equation (1b). The parameters used to establish this relationship (Equation 1b), μ , may or may not include system parameters θ defined in Equation (1a).

Typically, a nonlinear programming formulation with the objective of minimizing the least square error between the measured (y_{kl}^*) and modeled (y_{kl}) outputs over the time course (l = 1, ..., d) for which experimental data is collected (Equation 2) (Raue et al., and other parameter estimation/identifiability papers), is used to estimate parameters θ and μ in Equation 1.

$$\chi^{2}(\theta, \mu) = \sum_{k=1}^{m} \sum_{l=1}^{d} \left(\frac{y_{kl}^{*} - y_{kl}}{\sigma_{kl}^{*}} \right)^{2}$$
 (2)

The difference between the data and the model estimate for each output, and at each time point, is weighted by the variance in the experimental data σ_{kl}^* for that corresponding variable and time point. The minimization problem is solved subject to constraints on the state and output variables expressed in Equation 1, as well as bounds on the estimated parameter values. The ability to determine a unique solution to the parameter estimation problem is however governed by the identifiability of the parameters in the model (McLean AND McAuley 2012).

The identifiability of parameters in nonlinear models can be classified into two categories: structural (or a priori) and practical (or posterior) identifiability. Any system (Equation 1) is said to be structurally identifiable if, for an input-output mapping defined by $\mathbf{y} = \Phi(\mu, u)$ for at least one input function u, any two

values of parameters μ_1 and μ_2 satisfy the relationship in Equation (3) below.

$$\Phi(\mu_1, u) = \Phi(\mu_2, u) \iff \mu_1 = \mu_2 \tag{3}$$

Accordingly, any system that has an infinite number of solutions to the parameter estimation problem for all input functions is said to structurally non-identifiable. Thus, the structural identifiability of parameters in a dynamic model helps establish the presence or absence of a relationship between the unobservable system states and the observable system outputs. Accordingly, the effect of model structure and parameterization on the ability to infer true parameter values from experimental data is determined by the structural identifiability of the parameter.

Experimental data from many physical systems is usually noisy, and when parameters are estimated on the basis of noisy data, the ability to estimate unique parameter values to satisfy Equation (3) is referred to as practical identifiability. The effect of the available experimental data on the ability to estimate unique parameter values is determined by the practical identifiability of the parameter. Accordingly, practical identifiability of a parameter is contingent upon the nature, quality and quantity of data available to estimate the parameter as opposed to the structure and parameterization of the model.

Thus, on the one hand, establishing the structural identifiability of parameters enables one to propose models that are not only appropriate representations of physical processes, but also are parameterized in such a way that the value of these parameters can be estimated. On the other hand, establishing practical identifiability of parameters in any model helps design experiments that are minimal, informative and useful for parameter estimation.

The dynamics of metabolic networks can also be represented by Equation (1) wherein the metabolite concentrations, \mathbf{x} , are state variables. The changes in the metabolite concentrations (\mathbf{x}) are expressed as a function of reaction rates dependent on enzyme kinetics, which are in turn expressed as nonlinear functions of the states (\mathbf{x}), kinetic parameters (θ) and other input variables (u).

Algorithms have been extensively developed for establishing structural identifiability of dynamic models of biological systems (IEEE Trans paper from 2007). Most of these algorithms use methods based on differential algebra (Glad and Ljung, 1994). More recent methods take a profile-likelihood-based approach (2012 Paper) to establish both structural and practical identifiability. However, these methods not only scale poorly

with increases in size of the modeled system, but also require dynamic time course data of the observable variables of the system. While computational burden due to poor scalability can be partly addressed with the current increases in computational power, the ability to obtain dynamic data for establishing identifiability of parameters in kinetic models of metabolism still remains a challenge.

In this paper, we propose a methodology to establish practical identifiability for parameters in kinetic models of metabolism. We present a computer algebra-based method that can facilitate experimental design for estimating parameters separately for each individual reaction within a metabolic network based on available steady state experimental data. This enables us to address the twin issues of scalability and data availability. We illustrate the utility of this method by applying it for a small network of gluconeogenesis in *E. coli* and demonstrating our ability to propose experiments that will facilitate parameter estimation for a kinetic model of this network. We also demonstrate the scalability of the proposed methodology to facilitate experimental design by applying it to a relatively larger metabolic network of the human red blood cell hepatocyte.

Outline:

- Importance of parameter values for estimating in vivo response using kinetic models of metabolism and consequently for metabolic network design using kinetic models of metabolism
- The need for parameter identifiability to determine unique and true parameter values from observed data
- Types and purpose of identifiability for parameters
- Methods for structural identifiability and existing methods for practical identifiability
- Lack of methods for practical identifiability and consequently experimental design (not covered?)
- work done in this paper for practical identifiability
- scalability of computer algebra-based methods for structural identifiability (using CRNT to reduce networks to make structural identifiability scalable) (move to discussion may be in discussion)

2 Methods:

2.1 Kinetic model of gluconeogenesis in E. coli:

The proposed model for acetate consumption through gluconeogenesis and its corresponding kinetic model is used as a case study to illustrate the utility of identifiability analysis for the design of experiments for estimating parameters in kinetic models of metabolism. The kinetic model is described below.

$$\frac{d}{dt}pep = v_1 - v_2 - v_4 \tag{4}$$

$$\frac{d}{dt}fdp = v_2 - v_3 \tag{5}$$

$$\frac{d}{dt}E = v_{e,max} \left(\frac{1}{1 + \left(\frac{fdp}{K_f^{fdp}} \right)^{n_e}} \right) - dE$$
 (6)

The kinetic expressions for fluxes v_1 through v_4 are given below. The consumption of acetate through v_1 and conversion of pep through v_2 are expressed in Equations (7) and (8) respectively using Michaelis-Menten kinetics. The acetate flux through v_1 is also governed by the quantity of available enzyme E.

$$v_1 = k_1^{cat} E \frac{acetate}{acetate + K_1^{acetate}}$$
 (7)

$$v_2 = V_2^{max} \frac{pep}{pep + K_2^{pep}} \tag{8}$$

$$v_{3} = V_{3}^{max} \frac{\tilde{fdp} \left(1 + \tilde{fdp}\right)^{3}}{\left(1 + \tilde{fdp}\right)^{4} + L_{3} \left(1 + \frac{pep}{K_{3}^{pep}}\right)^{-4}}$$
(9)

The allosterically regulated flux v_3 for the consumption of fdp is expressed in Equation (9) using the Monod-Wyman-Changeux (MWC) model for allosterically regulated enzymes, where \tilde{fdp} refers to the ratio of fdp with respect to its allosteric binding constant K_3^{fdp} . The added flux v_4 for the export of pep is expressed as a linear equation dependent on pep in Equation (10).

$$v_4 = k_4^{cat}.pep (10)$$

2.2 A method for practical identifiability of kinetic models of metabolism:

In this section, we show how practical identifiability of kinetic parameters in a dynamic model of metabolism can be established using the algebraic relationship between fluxes that form the network. In a kinetic model, the value of every flux v_i is expressed using one of the many available enzyme kinetic formulations. Without loss of generality, all of these kinetic formulations can be expressed as nonlinear algebraic equations (Equation 11). The fluxes are expressed as a function of the metabolite concentrations x and the kinetic parameters θ .

$$v = f(\mathbf{x}, \theta) = \frac{N(\mathbf{x}, \theta)}{D(\mathbf{x}, \theta)}$$
(11)

In Equation 11, $N(\mathbf{x}, \theta)$ is the numerator of f, and $D(\mathbf{x}, \theta)$ is the denominator of f.

Let $\theta \in \mathbb{R}^P$, and given a set of measurements for the metabolite concentrations \mathbf{x}_j and the reaction fluxes \mathbf{v}_j for each experiment j=1,2,...,n, if all parameters are identifiable, theoretically, the P unknown parameters in θ can be estimated by solving $P \subset 1,2,...,n$ nonlinear algebraic equations shown in Equation 12.

$$v_k = \frac{N(\mathbf{x}_k, \theta)}{D(\mathbf{x}_k, \theta)} \qquad \forall k \in 1, 2, ..., k, ..., P$$
(12)

Here, K = 1, 2, ..., k, ..., P is the index set of P experiments for which metabolite concentration (\mathbf{x}_K) and flux (\mathbf{v}_K) measurements are available for each metabolite and flux within the metabolic network. Parameter $\theta_p \in \theta$ is said to be identifiable if

However, if for any index l, \mathbf{x}_l and \mathbf{v}_l do not satisfy the condition that $D(\mathbf{x}_l, \theta) \neq 0$, i.e., at least one flux v_i is not identifiable based on experiment l, then the lth experiment cannot be used for parameter estimation and should be discarded. Consequently, an additional experiment $m \in 1, 2, ..., n, m \neq l$ is needed to estimate all p parameters in θ . This analysis can be performed for each flux in a metabolic network independent of all the other fluxes. This enables this method to be scalable to even genome-scale models. The following section demonstrates this methodology for one of the fluxes in the gluconeogenic model described earlier.

2.3 Experimental design for parameter estimation in kinetic models of metabolism:

The identifiability of each parameter based on each experiment indexed as j = 1, ..., n

2.4 Identifiability of parameters in a kinetic model of gluconeogenesis:

2.5 Data for establishing parameter identifiability in kinetic model of gluconeogenesis:

Steady state metabolomics and fluxomics data can be gathered under different physiological conditions by either perturbing the expression levels for different enzymes within a metabolic network, or by changing the substrate concentrations under which the cells grow. The aforementioned model of gluconeogenesis has three different fluxes $(v_1, v_2 \text{ and } v_3)$ whose enzyme expression parameters $(V_1^{max}, V_2^{max} \text{ and } V_3^{max})$ can be perturbed to simulate the repression and over expression of the corresponding enzymes. Furthermore, the acetate concentration on which the cell is grown can also be perturbed to measure cellular response to changes in the substrate concentration. We use the in silico metabolomics and fluxomics data generated from these perturbation experiments to demonstrate parameter identification with our methodology.

If any data generated from a perturbation experiment i results in nonidentifiability, we eliminate experiment i from the list of experiments that need to be performed for parameter estimation.

We use flux v_2 to demonstrate the identifiability analysis method described in the previous section. Flux v_2 has two parameters, V_2^{max} and K_2^{pep} that need to be estimated from experimental data. Here, we assume that at least two different sets of experimental data for the concentrations and fluxes are available. Accordingly, we label these dataset as pep^1 , v_2^1 and pep^2 , v_2^2 respectively. Subsequently, these experimental datasets can be included in the model to form two simultaneous nonlinear algebraic equations in the parameters V_2^{max} and K_2^{pep} (Equation 13).

$$V_2^{max} = \frac{v_2^1 v_2^2 (pep^1 - pep^2)}{v_2^2 pep^1 - v_2^1 pep^2}$$
 (13a)

$$K_2^{pep} = \frac{pep^1(v_2^1 pep^2 - v_2^2 pep^2)}{v_2^2 pep^1 - v_2^1 pep^2}$$
(13b)

Table 1: Table showing the perturbed values of all fluxes used for parameter estimation.

Designation	Perturbed Fluxes	Perturbed Values
P1	v_1	2
P2	v_2	0.2
Р3	v_3	0.5

3 Results:

- 3.1 Getting closed-form expressions for each flux in Kotte model:
- 3.2 Experiments that can establish identifiability for each flux in Kotte model:
- 3.3 Combinations of experiments that will enable identification of all model parameters:

3.4 Expanding methodology to RBC model:

Outline:

- parameter estimation is a well developed field typically using minimization of least square error to estimate model parameters from available experimental data
- if parameters are structurally identifiable, it does not guarantee practical identifiability from noisy experimental data
- identifiability dependent on whether given datasets (outputs) for estimation can sufficiently distinguish between different parameter values

Sections:

- datasets required for parameter estimation in kinetic models of metabolism (methods?)
- identifiability in kotte model scalability, number of experiments required, requirements for time course data(? in the intro)

• identifiability in large rbc model

References

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