Some Remarks on Input Choices for Biochemical Systems

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Control systems theory concerns the study of open dynamical systems that process time-dependent input signals (stimuli, ligands, controls, forcing functions, test signals) into output signals (responses, measurements, read-outs, reporters). Such input/output (i/o) systems may be studied by themselves, or as components (subsystems, modules) of larger systems. Two extreme approaches to modeling systems (or particular subsystems) are as follows [1, 2]:

- The black-box, purely input/output, or phenomenological description ignores all mechanistic detail and characterizes behavior solely in terms of behavioral or stimulus-response data. This data may be summarized for example by transfer functions (for linear systems and certain types of “bilinear” systems), Volterra, generating series, or Wiener expansions (for general nonlinear systems), statistical models that correlate inputs and outputs, neural network and other approximate models, or machine learning prediction formalisms.

- The state-space, internal, or mechanistic description, in which all relevant players (“state variables” such as proteins, mRNA, and metabolites) are specified, and the forms for all reactions and reaction constants are described.

In between these two extremes, one finds combinations (“gray box” view). Available mechanistic information is incorporated into the model, and this is supplemented by i/o data, which serves to impose constraints on (or, in a Bayesian approach, to provide prior distributions of) unknown parameters to be further identified. In engineering applications, black-box modeling is common, for example, in chemical process control, and state-space modeling is often used in mechanical and aerospace engineering. Many biological models involve combinations of both approaches.

The subfield of realization theory (“reverse-engineering”) deals with the question of how much of the internal system can be deduced from its input/output or black-box behavior. One may pose questions about the structure itself as well as about parameters, once that a concrete systems structure has been hypothesized. The basic results classify symmetry groups of possible internal realizations of a given behavior, typically through linear algebra, algebraic-geometric, and differential-geometric techniques, depending on the class of systems being studied, or provide conditions for parameter “identifiability” from i/o data. (The term is not to be confused with identification theory, the field that incorporates stochastic aspects so as to deal with noisy data. The underlying theoretical questions of what is ultimately achievable, even in ideal noise-free conditions, are easier to understand in a deterministic setting.)

Key concepts used in realization theory are observability (how much information about internal dynamic variables, such as chemical concentrations, can be ultimately obtained from time-varying stimulus/response measurements?) and reachability (or controllability) (what state configurations can be attained, from a given initial state by manipulating inputs, also called “open-loop” controls?). Minimal models explaining observed input/output behavior typically are characterized in terms of reachability and
observability. Also, identifiability of parameters can be reduced to an observability problem. Many tests exist for observability and controllability, for linear and more generally for large classes of nonlinear systems [1], although the theory of nonlinear controllability is still very incomplete.

There are many other important subfields of control theory, among them the study of disturbances and robustness (how does one quantify the effect of actuator and sensor disturbances, and the robustness to unstructured perturbations?), feedback (how does one design feedback laws or “closed-loop controls” so as to enhance desirable system characteristics such as stability and robustness to structural perturbations, noise, and other disturbances?), optimal control (among those open-loops controls which serve to drive one internal configuration of state variables to another, which ones are optimal in appropriate senses?), and the question of hybrid or discrete/continuous interfaces (how do digital sensors, actuators, limited-bandwidth communication channels, and feedback devices interact with systems described by analog real-valued signals?). Each of these subfields brings up its own set of challenges special to systems biology [3, 4].

Here, we make some elementary remarks regarding how the choice of inputs being applied to a system (pulses, steps, etc) affects parameter identifiability. To keep the exposition accessible, we work out some extremely simple explicit examples, instead of discussing general theoretical matters. Some of the material, such as that covering linear systems, is standard [1]. The nonlinear examples illustrate ideas from a paper in preparation.

Various Input Classes

We discuss next how the choice of input signals impacts the theoretical possibility of identifying system responses (black-box behavior) and/or internal structure (state-space view).

Normally, the analysis of responses to small signals (or small perturbations from steady states, leading to “weakly activated” systems) can be handled by linear control theory, which provides a rich toolkit for the analysis of biochemical networks. On the other hand, if relevant signals are large, or if nonlinear effects, such as receptor desensitization or dimerization, affect signaling at a comparable time-scale, then linear tools do not suffice and new techniques must be brought to bear. Nonlinear control systems analysis has undergone an exceptional period of development and maturation within the last 20 years, largely driven by application needs from aerospace, mechanical, and other engineering disciplines. The tools that have been developed reflect the requirements from these fields, hence the emphasis on studying certain classes of systems, such as Hamiltonian dynamics, which are less relevant to biochemical networks. Although many of the fundamental ideas are still applicable to biology, much further theoretical work is needed.

Small-signal analysis: equivalence of step, impulse, and pulse responses

When probing systems with signals of small magnitude, the full power of the well-developed linear control theory can be taken advantage of. To illustrate the problems that arise in this context, we use a very simple example, that of a single species $X$ whose concentration $x(t)$ is described by an ordinary differential equation:

$$\frac{dx}{dt}(t) = -ax(t) + bu(t)$$

in which the variable $u(t)$ measures the, generally time varying, concentration of an external input $U$, and $a, b$ are unknown positive constants. (From now on, we will drop the time arguments $t$ when clear from the context.)

For example, $X$ might represent the concentration of bound receptors, and $U$ the concentration of a ligand which is present only in very small quantities. In this case, $a$ is the degradation rate of $X$, due
to internalization, ligand unbinding, and other effects, and \( b \) represents the concentration of unoccupied receptors, which we consider as constant by assuming that their abundance is large compared to the ligand. (If the concentration is not constant, then a linear model is not appropriate.) Generally, variables such as \( x \) and \( u \) represent not absolute concentrations but rather perturbations from a steady state.

We assume that the reporter being used provides a measurement \( y(t) = cx(t) \) equal to the product of the concentration of \( X \) and some (unknown) positive constant \( c \). Mechanistically, the form of \( y \) might arise from an additional differential equation:

\[
\varepsilon \frac{dy}{dt} = -y + cx
\]

which models the formation and decay of the reporter \( Y \) (we took a unit decay to simplify) and \( 0 < \varepsilon \ll 1 \) indicates that this process happens at a much faster time-scale than the production of \( x \). Using a quasi-steady state approximation (singular perturbation), we set \( \varepsilon = 0 \) and consider \( y = cx \) as the output. Another way in which an output of this type could arise is as a model of unknown measurement device calibration.

For any input \( u(t) \) for \( t > 0 \), and \( x(0) = 0 \), we have, for \( t > 0 \):

\[
y(t) = (k * u)(t) = \int_0^t k(t - r)u(r) \, dr
\]

(star denotes convolution), where

\[
k(r) = ce^{-\alpha t}b.
\]

The function \( k(t) \) is called the impulse-response of the system, because a unit impulse \( u = \delta_0 \) at time 0 (or, in practice, a very narrow pulse with unit area) elicits the response \( y(t) = k(t) \). The function \( K(t) \) is called the step-response, because for any step input \( u(t) \equiv u_0 \) for \( t > 0 \), the output is \( y(t) = K(t)u_0 \) for \( t > 0 \).

Knowledge of \( K \) is equivalent to knowledge of \( k \), because \( k = K' \) and \( K = \int_0^t k \). Therefore, knowing the time-series response to any single step input is sufficient to determine \( K(t) \) (just divide: \( K(t) = y(t)/u_0 \)), which in turn (convolution formula) determines the response to any other possible input, for instance pulses.

In fact, knowing the response to basically any single input is theoretically enough for finding \( k \). A proof using Laplace transforms, is clear from the fact that \( \hat{y}(s) = W(s)\hat{u}(s) \), where \( W(s) \) is the frequency response of the system, that is to say, the Laplace transform of \( k \). Thus, as long as \( \hat{u}(s) \) does not vanish, one can solve for \( W = \hat{y}/\hat{u} \), and \( k \) can then be obtained by Laplace inversion. In practice, of course, one does not perform just a single input/output experiment, since measurements will be noisy, nor does one obtain \( k \) as \( K' \), since differentiation of an estimated step function is numerically difficult. Several measurements are averaged, through a least-squares regression or another technique. This is discussed at length in textbooks in identification theory. It is easier, however, to explain the theory in the noise-free case.

**Steady state measurements do not provide enough information about transient behavior**

Steady-state experiments measure the steady state output \( y_{ss} \) of the reporter variable as a function of the magnitude of a step input. With the above example, \( y_{ss} = cx_{ss} = \frac{cb}{a}u_0 \). In other words, the only information about the three parameters \( a, b, c \) that can be obtained from such experiments is the combination \( \gamma = bc/a \) (called the steady state gain of the system). For example, the two systems (i) \( dx/dt = -x + u \) with \( y = x \) and (ii) \( dx/dt = -2x + u \) with \( y = 2x \) have the same steady-state gain, \( \gamma = 1 \). The i/o behaviors of these two systems are different, however: when using a step input \( u(t) \equiv 1 \), and initial condition
In either case, we have gone from a “gray box” description to a complete mechanistic description. Partial imprecision in a biochemical assay provides the range “less gray” level of description than total ignorance of the parameters. Now it is possible to also determine multivariable inputs and outputs, in which case both can be measured and only if the parameters from a homologous protein, for example). That is, the invariants provide a “prior belief” about observability, the following theorem is true [1]: two triples are equivalent if and only if
t1
K(0) = K′′(0) = −cab are known, and a = −K′′(0)/K′(0) can be estimated. One says that the two parameter triples (a1, b1, c1) and (a2, b2, c2) are equivalent if they cannot be distinguished by any i/o experiment. Thus, we showed that two triples are equivalent if and only if b1c1 = b2c2 and a1 = a2. Another way to say state this fact is through a one-parameter group of symmetries: (a, b, c) is equivalent to (a, Tb, cT−1) for all T ≠ 0. (These symmetries amount to rescaling the concentration x(t): in different units, one has a new variable z(t) = Tx(t), which satisfies the differential equation dz/dt = −az + Tbu, and the reporter is now and y = cT−1x.)

A far-reaching generalization of this result is available for linear systems of arbitrary dimension n (number of species), not merely n = 1 as in our example: systems dx/dt = Ax + bu, y = cx, where now A is an n × n matrix, b is a column n-vector, and c is row n-vector. Under genericity conditions (“reachability and observability”), the following theorem is true [11]: two triples (Ai, bi, ci), i = 1, 2 are i/o equivalent if and only if (TAiT−1, Tbi, cTi−1) = (A2, b2, c2) for some invertible matrix T. (The theorem is also true for multivariable inputs and outputs, in which case both B and C are matrices.)

Continuing with our example, we may think of our system as a “gray box” in the sense that we do not know the complete mechanism (all the parameters) but neither do we have complete ignorance, since we are able to measure a and the product bc. Now suppose that a new experimental design allows us not merely to measure y(t) but also do directly measure the parameter c (for example, the binding of X to species Y may be characterized by means of a biochemical experiment). Now it is possible to also determine b, since b = K′(0)/c. Or perhaps, instead of a biochemical experiment, a new probe is devised, which allows one to directly measure x(t). Then, knowing that dx/dt = −ax + bu, and from knowledge of x(t) and x′(t) at two values t = ti, one can determine both a and b, and hence, c = K′(0)/b as well. In either case, we have gone from a “gray box” description to a complete mechanistic description. Partial knowledge of b or c can also be understood in these terms: suppose that we know that K′(0) = 1 and that an imprecise biochemical assay provides the range c ∈ [0.01, 0.1]. Then, we know that b ∈ [10, 100], a “less gray” level of description than total ignorance of b.

These ideas are particularly useful in a Bayesian context. The i/o knowledge given by the value of a and the product bc, combined with physically realistic ranges for the unknown b and c, can be used to provide priors for parameter estimation through further experiments (or to evaluate the reliability of using parameters from a homologous protein, for example). That is, the invariants provide a “prior belief” about the parameters π = (a, b, c), summarized as a probability P(π). After an experiment e, this probability may then be updated to obtain a posterior distribution P(π|e) through an application of Bayes Rule:

\[ P(\pi|e) = \frac{P(e|\pi)P(\pi)}{P(e)}, \]

where P(e|π) is the likelihood of the experimental results under the assumed parameters π.
Step inputs are not sufficient for nonlinear identification

There are strong theoretical justifications for the need for rich input classes in the context of reverse engineering biological systems. We use a simplified, and hence somewhat unrealistic, example in order to explicitly illustrate this point, but the phenomenon scales to larger systems.

Consider the following system, which describes the time-evolution of the concentrations $x(t)$ and $z(t)$ of two chemical species $X$ and $Z$:

$$
\begin{align*}
\frac{dx}{dt} &= -\lambda x + u^2 \\
\frac{dz}{dt} &= -\lambda z + u
\end{align*}
$$

with initial concentrations $x(0) = 0$ and $z(0) = 0$, and where $u(t)$ is the (possibly time-varying) concentration of an input (for instance a ligand) $U$ that helps promote formation of $X$ and $Z$. Our goal is to estimate the unknown parameter $\lambda$, the degradation rate of $X$ and $Z$. Note that as long as no input $U$ is applied ($u(t) = 0$), the concentrations of $X$ and $Z$ remain at steady state (zero).

We assume that the only available measurement is the (time-varying) concentration $y(t)$ of a reporter $Y$, which we take as a function of the instantaneous concentrations of $X$ and $Z$ according to the formula:

$$
y = x + u(a - z)
$$

were $a$ is some known positive constant. (One could view $y$ as obtained through a quasi-steady state approximation applied to

$$
\varepsilon \frac{dy}{dt} = -y + x + u(a - z)
$$

with $0 < \varepsilon \ll 1$, which describes a reaction in which a certain substance $A$ dimerizes with $U$ in order to promote formation of $y$, but $A$ is sequestered by $Z$ with 1-1 stoichiometry, so the free amount of $A$ equals $a - z = A_{\text{total}} - A_{\text{bound}}$, because the concentration of $A_{\text{bound}}$ equals the concentration of $Z$.)

If we use a step function input, $u(t) \equiv u_0$ for $t > 0$, then $x(t) \equiv z(t)u_0$, and therefore $y(t) \equiv au_0$, independently of the actual value of $\lambda$. Thus, with step inputs, no matter how many experiments (different values of $u_0$) are carried out, it is impossible to obtain any information about $\lambda$.

On the other hand, suppose that we use a unit ramp: $u(t) = t$. Then $y'''(t) = -\lambda(\lambda t - 2)e^{-\lambda t}$, and so

$$
\lim_{t \to 0} y'''(t) = 2\lambda
$$

and therefore $\lambda$ can be identified by differentiation. (Of course, differentiation will not be used in practice, nor is it needed since one may use other techniques for estimating $\lambda$, but we are just proving theoretical identifiability.) In summary, a single ramp input is enough to identify $\lambda$, but no amount of even complete transient time-measurements of step responses will achieve this result.

Instead of ramps, we could also use pulses. Suppose that we consider a unit pulse: $u(t) = 1$ for $0 \leq t \leq 1$ and $u(t) = 0$ for $t > 1$. Now, $x(t) = z(t) = \frac{1}{\lambda}(1 - e^{-\lambda t})$ for $t \in [0, 1]$ and $x(t) = z(t) = \frac{1}{\lambda}(e^{\lambda} - 1)e^{-\lambda t}$ for $t \geq 1$, so $y(t) \equiv x(t) + 0(A - z(t)) = x(t)$ for $t \geq 1$, which means that

$$
\lim_{t \to 1} y'(t) = e^{-\lambda} - 1,
$$

and therefore $\lambda$ can be calculated as:

$$
\lambda = -\ln \left(1 + \lim_{t \to 1} y'(t)\right).
$$
Once again, a non-step signal is successful in identifying the parameter $\lambda$, but steps are not enough.

Similar examples can be given with more parameters and more complicated systems. Also, we took the same degradation rates for $X$ and $Z$, because we are considering a noise-free ideal situation. If these degradation rates would not necessarily be equal but are close, then $y(t)$ will depend on $\lambda$, even in the step case. However, the dependence will be through a very small multiplier coefficient, indicating a very small sensitivity and therefore very high estimator variance (Cramer-Rao bounds), thus rendering estimation practically impossible. It is possible to prove (paper in preparation) that for bilinear systems (those in which the only possible products are between inputs and states), pulses provide as much theoretical information as any possible time-varying input signals.

Remarks

We have only discussed parameter identification, as opposed to structure identification. For small-signal (linear) analysis, the two problems are the same (at least if an upper bound on the number of species in the system is known), since zero entries in the differential equations indicate non-existent interactions. For nonlinear systems, the problem of identification of structure is far more difficult. It is the subject (at least for the ideal noise-free case) of nonlinear realization theory \[2\]. The parameter estimation problem is a special case of a nonlinear observability problem (thinking of unknown parameters as constant states), and a rich theory allows the use of Lie-algebra and differential-algebra techniques for determining observability \[1\].

There are results available characterizing the number of experiments required for identifying a given number of parameters in a system \[5\]. Another relevant theoretical mathematical fact is that generic “random” single inputs are enough for identification \[2\], and this fact is very useful in the context of identification of certain biochemical systems \[6\].

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