Variational Kinetics

Author: Masoud Ahookhosh, Systems Biochemistry Group, Luxembourg Centre for Systems Biomedicine

Reviewers: Ronan Fleming, Sylvain Arreckx

INTRODUCTION

During this tutorial, you will learn how to investigate steady states and moiety conserved steady states of biochemical reaction systems [1].

Let us consider a biochemical network with m molecular species and n reversiable elementary reactions. We define forward and reverse stoichiometry matrices $(F,R\in Z_+^{m\times n})$ respectively, where F_{ij} denotes the stoichiometry of the i^{th} molecular species in the j^{th} forward reaction and R_{ij} denotes the the stoichiometry of the i^{th} molecular species in the j^{th} reverse reaction. We assume that every reaction conservs mass, that is, there exists at least one positive vector $l \in R_{++}^m$ satisfying $(R-F)^T l = 0$. The matrix N = R - F represents net reaction stoichiometry and may viewed as the incidence matrix of a directed hypergraph. We are assume that there are less molecular species than there are net reactions, i.e., m < n. We assume the cardinality of each row of F and R is at least one and the cardinality of each column of N is at least two. We also assume that $\operatorname{rank}([F,R]) = m$, which is a requirement for kinetic consistency.

Steady state nonlinear system

Let $c \in R_{++}^m$ denote a variable vector of species concentrations. Assuming constant ningegative elementary kinetic parameters $k_f, k_r \in R_+^n$, we assume elementary reaction kinetics for forward and reverse elementary reaction rates as $s(k_f, c) := \exp(\ln(k_f) + F^T \ln(c))$ and $r(k_r, c) := \exp(\ln(k_r) + R^T \ln(c))$, respectively, where $\exp(\cdot)$ and $\ln(\cdot)$ denote the respective componentwise functions. Then, the deterministic dynamical equation for time evolution of molecular species concentration is given

$$\frac{dc}{dt} = N(s(k_f, c) - r(k_r, c)) = N\left(\exp(\ln(k_f) + F^T \ln(c)) - \exp\left(\ln(k_r) + R^T \ln(c)\right)\right) = f(x) := [N, -N] \exp(k + [F, R]^T x) =: -f(c),$$

where $[\cdot,\cdot]$ stands for horizental cancatenation operator. A vector c^* is a steady state if and only if it satisfies $f(c^*) = 0$, leading to the nonlinear system

$$f(x) = 0.$$

Moiety conserved steady state nonlinear system

Let us emphasize that a vector c^* is a steady state of the biochemical system if and only if

$$s(k_f, c^*) - r(k_r, c^*) \in \mathcal{N}(N),$$

where $\mathcal{N}(N)$ denotes the null space of N. Therefore, the set of steady states $\Omega = \left\{c \in R_{++}^m, f(c) = 0\right\}$ is unchanged if we replace the matrix N with \overline{N} with the same kernel. Suppose that $\overline{N} \in Z^{r \times n}$ is the submatrix of N whose rows are linearly independent, then $\operatorname{rank}(\overline{N}) = \operatorname{rank}(N) := r$. If one replaces N by \overline{N} and considers the logarithmic scale, by setting $x = \ln(c) \in R^m$ and $k = [\ln(k_f)^T, \ln(k_r)^T]^T \in R^{2n}$, then we have

$$\overline{f}(x) := [\overline{N}, -\overline{N}] \exp(k + [F, R]^T x).$$

Let also $L \in R^{m-r,m}$ denote a basis for the left nullspace of N, which implies $N^TL = 0$. We also have $\operatorname{rank}(L) = m - r$. WE say that the system satisfies moiety conservation if for any initial concentration $c_0 \in R^m_{++}$, it holds

$$L c = L \exp(x) = l_0$$

where $l_0 \in R_{++}^m$. Therefore, the problem of finding the moiety conserved steady state of a biochemical reaction network is equivalent to solving the nonlinear system of equations

$$h(x) := \begin{pmatrix} \overline{f}(x) \\ L \exp(x) - l_0 \end{pmatrix} = 0.$$

We introduce an interface to software that enables the computation of the elementary modes, or extreme pathways, given a network and user-defined reaction bounds.

METHODOLOGY

In order to solve above-mentioned nonlinear system, we here address three classes of methods, i.e.,

- 1. Levenberg-Marquardt methods [1,2,5,6,9.10],
- 2. DC programming methods [3],
- 3. derivative-free methods for duplomonotone mappings [4],

where each class of solvers are described shortly as follows:

- 1. The Levenberg-Marquardt methods are standard techniques used to solve nonlinear systems that each of which is a combination of the gradient descent and the Gauss-Newton methods. Therefore, knowing the first-order information (function values and gradients) of the mapping is enough to proceed the algorithm. We here consider two classes of Levenberg-Marquardt methods, namely locally convergent Levenberg-Marquardt methods (LLM_YF, LLM_FY, LLM_F, LLM) and globally convergent Levenberg-Marquardt methods (GLM_YF, GLM_FY, LevMar, LMLS, LMTR), see [1] and [2], respectively.
- 2. In DC programming methods, one needs to rewrite the nonlinear system as a minimization of a difference of two convex function. Then, the DC subproblem is the minimization of a convex function, which is constructed by keeping the first function and linearlizing the second function. We here address two DC programming methods, namely, DCA and BDCA, see [3] for detailed information.

3. Derivative-free methods are a class of methods that only needs function values to minimize a nonlinear least-squares problem. We here consider three derivative-free methods, namely, BDF, CSDF, and DBDF, see Algorithms 1-3 in [4] for more details.

MATERIALS

• Please ensure that the COBRA Toolbox has been properly installed and initialised.

PROCEDURE

Computing steady states of biochemical systems

The mandatory inputs for computing steady states are a <u>model</u> involving F and R, the name of a <u>solver</u> to solve the nonlinear system, an initial point x_0 , and <u>parameters</u> for the considered solvers. We first need to load data from a ".mat" file involve F, R, and F (kinetic vector). For example, for "Ecoli core" model, we have

```
global CBTDIR
tutorialPath = fileparts(which('tutorial_variationalKinetics.mlx'));
load([tutorialpath filesep 'Ecoli_core_data.mat']);
```

Then, we need to make a struture "model" by

```
model.F = F;
model.R = R;
```

and specify a solver by

```
solver = 'LMTR';
```

and detemine the parameters for the selected algorithm

```
parms.MaxNumIter = 1000;
parms.adaptive = 1;
parms.kin = kin;
```

otherwise, the selected algorithm will be run by the default parameters assigned in the codes. We finally need to run the function "optimizeVKmodel.m" like

```
output = optimizeVKmodel(model, solver, x0, parms);
```

Let us emphasize that all the slovers (LLM_YF, LLM_FY, LLM_F, LLM,GLM_YF, GLM_FY, LevMar, LMLS, LMTR, DCA, BDCA, BDF, CSDF, DBDF) can be used to find steady states of biochemical systems; however, based on our experiments, "LMTR" and "LMLS" perform better than the others. If you are not familiar with the solvers, we may suggest to use solvers with the dafault values for parameters.

Computing moiety conserved steady state of biochemical systems

The mandatory inputs for computing moiety conserved steady states are a <u>model</u> involving F, R, L, and l_0 , the name of a <u>solver</u> to solve the nonlinear system, an <u>initial point</u> x_0 , and <u>parameters</u> for the considered solvers. We first need to load data from a ".mat" file involve F, R, L, l_0 , and kin (kinetic vector). For example, for "Ecoli core" model, we have

Then, we need to make a struture "model" by

```
model.F = F;
model.R = R;
model.L = L;
model.10 = 10;
```

and specify a solver by

```
solver = 'LMLS';
```

and detemine the parameters for the selected algorithm

```
parms.MaxNumIter = 1000;
parms.adaptive = 1;
parms.kin = kin;
```

otherwise, the selected algorithm will be run by the default parameters assigned in the codes. We finally need to run the function "optimizeVKmodel.m" like

```
output = optimizeVKmodel(model, solver, x0, parms);
```

Let us emphasize that among above-mentioned solvers one can use the Levenberg-Marquardt slovers (LLM_YF, LLM_FY, LLM_GLM_YF, GLM_FY, LevMar, LMLS, LMTR) to find moiety conserved steady states of biochemical systems; however, based on our experiments, "LMTR" and "LMLS" perform better than the others. If you are not familiar with the solvers, we may suggest to use solvers with the dafault values for parameters.

Optional inputs

The function can have some optional inputs for slover, x_0 , and the parameters corresponding to the selected solver. Therefore, we here explain the most important optional inputs in the following with respect to the selected solver.

Parameters for all solvers:

- MaxNumIter: is the maximum number of iterations:
- MaxNumMapEval: is the maximum number of function evaluations;
- MaxNumGmapEval: is the maximum number of gradient evaluations;

- TimeLimit: is the maximum time limit;
- epsilon: is the accuracy parameter;
- kin: is a kinetic parameter in R^(2n);
- x_opt: is the optimizer (if available);
- psi_opt: is the optimum (if available);
- flag_x_error: is a flag to specify if the relative error of iteration points is needed (1) or not (0);
- flag_psi_error: is a flag to specify if the relative error of merit function is needed (1) or not (0);
- flag_time: is a flag to specify if saving time in each iteration is needed (1) or not (0);

Parameters for Levenberg-Marquardt solvers:

- solver: is one of the solver:
- 1. LLM_YF: the locally convergent Levenberg-Marquardt method of Yamashita and Fukushima [10];
- 2. LLM_FY: the locally convergent Levenberg-Marquardt method of Fan and Yuan [5];
- 3. LLM_F: the locally convergent Levenberg-Marquardt method of Fischer [6];
- 4. LLM: the locally convergent Levenberg-Marquardt method of Ahookhosh, Artacho, Fleming, and Phan [1];
- 5. GLM_YF: the globally convergent Levenberg-Marquardt method of Yamashita and Fukushima [10];
- 6. GLM_FY: the globally convergent Levenberg-Marquardt method of Fan and Yuan [5];
- 7. LevMar: the globally convergent Levenberg-Marquardt method of Ipsen, Kelley, and Pope [9];
- 8. LMLS: the globally convergent Levenberg-Marquardt method of Ahookhosh, Artacho, Fleming, and Phan [2];
- 9. LMTR: the globally convergent Levenberg-Marquardt method of Ahookhosh, Artacho, Fleming, and Phan [2];
- adaptive: is a flag to specify lambda should be updated adaptively (1) or not (0);
- eta: is a constant for Levenberg-Marquardt parameter;
- Stopping_Crit: is a stopping criterion;
- 1. 1: stop if the norm of gradients is less or equal than epsilon;
- 2. 2: stop if the norm of rhe mapping is less or equal than epsilon:
- 3. 3: stop if maximum number of iterations is reached;
- 4. 4: stop if maximum number of function evaluations is reached;
- 5. 5: stop if maximum number of gradient evaluations is reached;
- 6. 6: stop if time limit is reached;
- 7. 7: stop if ||grad||<=max(epsilon,epsilon^2*ngradx0)
- 8. 8: stop if ||nhxk||<=max(epsilon,epsilon^2*nhx0)
- 9. 9: stop if ||hxk||<=epsilon or maximum number of iterations is reached

Parameters for DC programming solvers:

• solver: is one of the solver;

- 1. DCA: DC programming algorithm of Artacho, Fleming, and Phan (Algorithm 1) [3];
- 2. BDCA: DC programming algorithm of Artacho, Fleming, and Phan (Algorithm 2 and 3) [3];
- alpha: is a constant for the line search;
- beta: is the backtarcking constant;
- lambda_bar: starting step-size for the line search;
- rho: is the strong convexity parameter;
- flag line search: is a flag determines either "Armijo" or "Quadratic interpolation" should be used;
- Stopping Crit: is a stopping criterion;
- 1. 1: stop if the norm of rhe mapping is less or equal than epsilon;
- 2. 2: stop if maximum number of iterations is reached;
- 3. 3: stop if maximum number of function evaluations is reached;
- 4. 4: stop if time limit is reached;
- 5. 5: stop if ||fxk||<=epsilon or maximum number of iterations is reached

Parameters for Derivative-free solvers:

- solver: is one of the solver;
- 1. BDF: backtracking derivative-free algorithm of Artacho and Fleming [4];
- 2. CSDF: constant step derivative-free algorithm of Artacho and Fleming [4];
- DBDF: double backtracking derivative-free algorithm of Artacho and Fleming [4];
- alpha: is a constant with alpha < 2 sigma;
- beta: is the backtarcking constant;
- lambda_min: lower bound of the step-size;
- lambda_max: upper bound of the step-size;
- Stopping Crit: is a stopping criterion;
- 1. 1: stop if the norm of rhe mapping is less or equal than epsilon;
- 2. 2: stop if maximum number of iterations is reached;
- 3. 3: stop if maximum number of function evaluations is reached;
- 4. 4: stop if time limit is reached;
- 5. 5: stop if ||fxk||<=epsilon or maximum number of iterations is reached;

For a complete list of optional inputs and their definition, you can also run the following command.

help optimizeVKmodel

Output

The output of optimizeVKmodel.m is a structure "output" involving the fields

- x best: is the best approximation of the optimizer;
- psi_best: is the best approximation of the optimum;
- T: is the running time;
- Niter: is the total number of iterations;
- Nmap: is total number of mapping evaluations;
- Ngmap: is total number of mapping gradient evaluations (if gradient used in the algorithm);
- merit_func: is an array including all merit function values;
- x_error: is the relative error of iteration points;
- psi_error: is the relative error of merit function;
- Time: is the running time of all iterations;
- Status: is the reason of termination;

TIMING

Running, the code is dependent on the size of models and the solver selected, which may take long from less than 1 second to few hours.

ANTICIPATED RESULTS

Finding steady states or moiety conserved steady state with one of the above-mentioned solvers (e.g., solver = 'LMTR') leads to the following results.

output = optimizeVKmodel(model, solver, x0, parms)

TROUBLESHOOTING

In order to compute moiety conserved steady states, one should not use DC programming algorithms (DCA and BDCA) or derivative-free algorithms (BDF, CSDF, DBDF) because the current version of these codes are designed to deal with steady state of biochemical systems.

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