

Dynamic Optimization with Particle Swarms (DOPS): A meta-heuristic for parameter estimation in biochemical models

Adithya Sagar, Christine Shoemaker[†] and Jeffrey D. Varner*

School of Chemical and Biomolecular Engineering

[†]School of Civil and Environmental Engineering

Cornell University, Ithaca NY 14853

Running Title: Parameter estimation in biochemical models

To be submitted: *PLoS ONE*

*Corresponding author:

Jeffrey D. Varner,

Associate Professor, School of Chemical and Biomolecular Engineering,

244 Olin Hall, Cornell University, Ithaca NY, 14853

Email: jdv27@cornell.edu

Phone: (607) 255 - 4258

Fax: (607) 255 - 9166

Abstract

Mathematical modeling is a powerful tool to analyze, and ultimately design biochemical networks. However, the estimation of biochemical model parameters is a significant challenge. Parameter estimation in biochemical models typically involves expensive function evaluations and noisy data, making it difficult to quickly obtain optimal solutions. Biochemical models often also have many local extrema which further complicates parameter estimation. Toward these challenges, we developed Dynamic Optimization with Particle Swarms (DOPS), a novel global meta-heuristic that combined features of multi-swarm particle swarm optimization with dynamically dimensioned search (DDS). DOPS uses a multi-swarm particle swarm optimization technique to generate candidate solution vectors, the best of which is greedily updated using dynamically dimensioned search. We tested the performance of DOPS on a model of human coagulation cascade. We performed $\mathcal{T} = 25$ trials with $\mathcal{N} = 4000$ function evaluations per trial, and compared the performance of DOPS with other commonly used meta-heuristics such as differential evolution (DE), simulated annealing (SA) and dynamically dimensioned search (DDS). We further tested the predictive power of the coagulation model parameters against data not used in training, and found good agreement between simulations and experimental measurements. Lastly, we tested the performance of DOPS on commonly used test functions for global optimization and on published biochemical parameter estimation benchmark problems. For the wide range of problems that we considered, DOPS outperformed other meta-heuristic approaches despite a limited number of function evaluations. Taken together, DOPS is a promising meta-heuristic approach for the estimation of biochemical model parameters in relatively few function evaluations.

Keywords: Parameter identification, Meta-heuristic optimization, Biochemical modeling

1 Introduction

2 Cells process nutrients and respond to changes in their environment using complex bio-
3 chemical networks. These networks contain thousands of components interconnected
4 through nonlinear enzyme catalyzed reactions. Mathematical modeling has evolved as a
5 powerful paradigm to analyze, and ultimately design these complex networks [1–5]. Math-
6 ematical modeling of biochemical networks is often an iterative process. First, models are
7 formulated from biochemical knowledge, and then model parameters are estimated using
8 experimental data [6–8]. Parameter estimation is typically framed as a non-linear opti-
9 mization problem wherein the residual (or objective function) between experimental data
10 and model simulations is minimized using an optimization strategy [9]. Optimal parame-
11 ters obtained from model training are then used to validate the model on unseen experi-
12 mental data. If validation fails, model construction and calibration are repeated iteratively
13 until satisfactory results are obtained.

14 Parameter estimation is a major challenge in the iterative development of biochemical
15 models. Although parameter estimation has been a well studied problem in engineering
16 for decades [10–13], the complex dynamics of large biological systems and noisy, often
17 incomplete experimental data pose a unique estimation challenge. Often optimization
18 problems involving biological systems are non-linear and multi-modal i.e. typical mod-
19 els often have multiple local minima or maxima [7, 9]. Non-linearity coupled with multi-
20 modality generally renders local optimization techniques such as pattern search [14],
21 Nelder-Mead simplex methods [15], steepest descent methods or Levenberg-Marquardt
22 [16] incapable of reliably obtaining optimal solutions as they generally terminate at local
23 minimum. Though deterministic global optimization techniques (for example algorithms
24 based on branch and bound) can handle non-linearity and multi-modality [17, 18], the
25 absence of derivative information, discontinuity of the objective functions, non-smooth
26 regions or the lack of knowledge about the objective function hampers these techniques.

Meta-heuristic stochastic optimization approaches like Genetic Algorithms (GAs) [ADI-REF], Simulated Annealing (SA) [19], Evolutionary Programming [ADI-REF] and population based searches like Differential Evolution (DE) [20] have shown promise on nonlinear multi-modal problems [21]. These techniques do not make any assumptions about the structure of objective function nor do they require *a priori* information about the objective function. Though they do not guarantee strong convergence, these approaches are effective in finding near optimal solutions. Mendes et al. [22] used simulated annealing to estimate rate constants for the irreversible inhibition of HIV proteinase, Modchang et al. [23] used a genetic algorithm to estimate parameters for a model of signal transduction [ADI-EXPAND], differential evolution approaches have also been effective on various biological problems [24–26]. Tashkova et al. compared different meta-heuristics for parameter estimation on a dynamic model of endocytosis and showed that DE was the most effective [27]. Banga and co-workers have also successfully applied scatter-search methods to estimate parameters on non-linear biological processes [28–30]. Hybrid approaches that combine a meta-heuristic with a local optimization search, wherein a near globally optimal solution obtained using a meta-heuristic is refined using a local search have also become popular. Villaverde et al. combined scatter search with local search methods for parameter estimation in a collection of systems biology models [31]. Fan et al. recently showed that population based meta-heuristics along with decomposition based methods can be also used to model gene circuits from mRNA data [32]. Despite these successes, a major drawback of most metaheuristic approaches is the large number of function evaluations required to explore the parameter space. Typically as models grow in size and complexity, evaluation of the objective function becomes computationally expensive. Thus performing a large number of function evaluations is not desirable (and perhaps not feasible).

In this study, we developed Dynamic Optimization with Particle Swarms (DOPS), a

novel meta-heuristic that combines the global search capability of multi-swarm particle swarm optimization and dynamically dimensioned search (DDS). The objective of DOPS is to obtain near optimal parameter estimates for large biochemical models within a relatively few function evaluations. DOPS uses a multi-swarm particle swarm optimization technique to generate candidate solution vectors which are then greedily updated using dynamically dimensioned search. We first considered a model of human coagulation cascade to test the performance of DOPS. Coagulation is a large, complex biochemical network involving strong positive feedback. We then tested the performance of DOPS on commonly used test functions for global optimization (Ackley and Rosenbrock) and published biochemical parameter estimation benchmark problems [31]. DOPS outperformed common meta-heuristic approaches like Differential Evolution (DE), Simulated Annealing (SA) and dynamically dimensioned search (DDS) on both the test functions and the coagulation model. DOPS also performed well on the benchmark problems where it outperformed enhanced scatter search (eSS) and recovered the nominal parameters with only 4000 function evaluations ([ADI-compared with what?]) across all the benchmark problems considered. Taken together, these studies suggest DOPS is a promising meta-heuristic approach for the estimation of biochemical model parameters in relatively few function evaluations.

Results

Optimization problem formulation. The problem of parameter estimation in a dynamic biological model consists of finding an optimal parameter vector which minimizes the difference between model simulations and \mathcal{E} experimental measurements. This difference is quantified by an objective function $K(\mathbf{p})$ which is typically the Euclidean norm of the simulation error subject to problem specific and parameter bounds constraints:

$$\begin{aligned} \text{minimize} \quad & K(\mathbf{p}) = \sum_{i=1}^{\mathcal{E}} (g_i(t_i, \mathbf{x}, \mathbf{p}, \mathbf{u}) - y_i)^2 \\ \text{subject to} \quad & \dot{\mathbf{x}} = \mathbf{f}(t, \mathbf{x}(t, \mathbf{p}), \mathbf{u}(t), \mathbf{p}) \\ & \mathbf{x}(t_0) = \mathbf{x}_0 \\ & \mathbf{c}(t, \mathbf{x}, \mathbf{p}, \mathbf{u}) \geq \mathbf{0} \\ & \mathbf{p}^L \leq \mathbf{p} \leq \mathbf{p}^U \end{aligned} \tag{1}$$

where t is time, $\mathbf{x}(t, \mathbf{p})$ is the state variable vector with an initial state \mathbf{x}_0 , $\mathbf{u}(t)$ is a model input vector, \mathbf{f} is the system of model equations (e.g., differential equations or algebraic constraints) and \mathbf{p} is the model parameter vector. The parameter search (or model simulations) can be subject to \mathbf{c} linear or non-linear constraints, and parameter bound constraints where \mathbf{p}^L and \mathbf{p}^U denote the lower and upper parameter bounds, respectively. The problem eventually is to find:

$$\mathbf{p}^* = \arg \min_{\mathbf{p}} K(\mathbf{p}) \tag{2}$$

Dynamic optimization with particle swarms (DOPS). DOPS is a novel meta-heuristic which combines multi-swarm particle swarm methods with dynamically dimensioned search (Fig. 1). The goal of DOPS is to estimate optimal or near optimal parameter vectors for high-dimensional biological models within a specified number of function evaluations. Toward this objective, DOPS begins by using a particle swarm search and then dynamically

switches, using an adaptive switching criteria, to the DDS search phase. We began the particle swarm phase by randomly initializing a swarm of \mathcal{K} -dimensional particles (represented as z_i), wherein each particle corresponded to a \mathcal{K} -dimensional parameter vector. After initialization, particles were randomly partitioned into k equal sized sub-swarms $\mathcal{S}_1, \dots, \mathcal{S}_k$. Thereafter within each sub-swarm \mathcal{S}_k , particles were updated according to the rule:

$$\mathbf{z}_{i,j} = \theta_{1,j-1} \mathbf{z}_{i,j-1} + \theta_2 \mathbf{r}_1 (\mathcal{L}_i - \mathbf{z}_{i,j-1}) + \theta_3 \mathbf{r}_2 (\mathcal{G}_k - \mathbf{z}_{i,j-1}) \quad (3)$$

where $(\theta_1, \theta_2, \theta_3)$ were adjustable parameters, \mathcal{L}_i denotes the best solution found by particle i within sub-swarm k for function evaluation $1 \rightarrow j-1$, and \mathcal{G}_k denotes the best solution found over all particles within sub-swarm \mathcal{S}_k . The quantities r_1 and r_2 denote uniform random vectors with the same dimension as the number of unknown model parameters ($\mathcal{K} \times 1$). Equation (3) is similar to the general particle swarm update rule, however, it does not contain velocity terms. In DOPS the parameter $\theta_{1,j-1}$ depends upon the function evaluations and is updated according to:

$$\theta_{1,j} = \frac{(\mathcal{N} - j) * (\mathbf{w}_{max} - \mathbf{w}_{min}))}{(\mathcal{N} - 1)} + \mathbf{w}_{min} \quad (4)$$

where \mathcal{N} represents the total number of function evaluations, \mathbf{w}_{max} and \mathbf{w}_{min} are the maximum and minimum inertia weights, respectively. While updating the particles, we made sure all dimensions of the solution represented by the particle were within bounds using a set of reflection boundary conditions:

After every \mathcal{M} function evaluations, particles were randomly redistributed to a new

```

if  $z_{i,j}^{old} < z_i^{min}$  then
   $z_{i,j}^{new} = z_{i,j}^{old} + (z_i^{min} - z_{i,j}^{old})$  if  $z_{i,j}^{new} > z_i^{max}$  then
     $z_{i,j}^{new} = z_i^{max}$ 
  end
end
if  $z_{i,j}^{old} > z_i^{max}$  then
   $z_{i,j}^{new} = z_{i,j}^{old} + (z_{i,j}^{old} - z_i^{max})$  if  $z_{i,j}^{new} < z_i^{min}$  then
     $z_{i,j}^{new} = z_i^{min}$ 
  end
end

```

106 sub-swarm, and updated according to Eqn. (3). This process continued till $\mathcal{F} * \mathbf{N}$ number
 107 of functions evaluations, where \mathcal{F} is the fraction of evaluations with the multi-swarms. The
 108 fraction of evaluations within the swam phase is based on the drop in error. If the error
 109 does not drop by 1% for a certain number of evaluations the search within swarm phase is
 110 terminated. At the end of these function evaluations, we froze all the solutions represented
 111 by various particles and chose the particle with best solution among $\mathcal{G}_1 \cdots \mathcal{G}_{NS}$ as the
 112 initial candidate vector \mathcal{G} for the remaining $(1 - \mathcal{F}) * \mathbf{N}$ number of function evaluations.
 113 This particle was then updated according to the rule:

$$\mathcal{G}_{new}(\mathbf{J}) = \begin{cases} \mathcal{G}(\mathbf{J}) + \mathbf{r}_{normal}(\mathbf{J})\sigma(\mathbf{J}), & \text{if } \mathcal{G}_{new}(\mathbf{J}) < \mathcal{G}(\mathbf{J}). \\ \mathcal{G}(\mathbf{J}), & \text{otherwise.} \end{cases} \quad (5)$$

114 where \mathbf{J} represents the set containing the specific dimensions being perturbed, r_{normal}
 115 denotes a normal random vector of the same dimensions as \mathcal{G} , and σ is the amplitude of
 116 perturbation:

$$\sigma = \mathbf{R}(\mathcal{MAX} - \mathcal{MIN}) \quad (6)$$

117 where \mathbf{R} is the scalar perturbation size parameter, \mathcal{MAX} and \mathcal{MIN} are $(\mathcal{K} \times 1)$ vec-
 118 tors that represent the maximum and minimum bounds on each dimension. The set \mathbf{J}
 119 was constructed using a probability function \mathcal{P}_j that represented the probability whether

120 a specific dimension j was perturbed or not. This function is a monotonically decreasing
 121 function that decreases with the number of function evaluations. \mathcal{P}_j can be any monoton-
 122 ically decreasing function, in our approach we used the following function:

$$\mathcal{P}_j = 1 - \log(j / ((1 - \mathcal{FR}) * \mathbf{N})) \quad (7)$$

123 Thus the number of dimensions of the candidate vector that are updated or perturbed
 124 decreases with the as the number of function evaluations increase. These updates are
 125 greedy in nature, so \mathcal{G}_{new} becomes the new solution vector only if it is better than the old
 126 one \mathcal{G} . The decrease of weight function in equation 4, reflection boundary conditions in
 127 equation 5, the update function described in equations 6 and 7, the selection probability in
 128 equation 8 are the means by which DDS ideas are incorporated into DOPS. The fraction
 129 of evaluations \mathcal{FR} within the swarm phase is based on a switching strategy wherein the
 130 switch from swarm phase to DDS phase happens when the error due to the best solution
 131 does not drop more than 1% of the original error, continuously for more than a prescribed
 132 number of function evaluations. This allows the solution to quickly jump out of local op-
 133 tima and avoid any convergence issues that are generally associated with swarm based
 134 searches.

Performance of DOPS on a model of the human coagulation cascade. We compared the performance of DOPS with simulated annealing (SA), differential evolution (DE), and dynamically dimensioned search (DDS) for a model of the human coagulation cascade. Coagulation is an archetype biochemical network that is highly interconnected, containing both positive and negative feedback (Fig. 2). In this study, we used the coagulation model of Luan et al [33] which modeled coagulation as a system of coupled non-linear ordinary differential equations. The Luan model contained 148 parameters and 92 species and has been validated using 21 published datasets.

The biochemistry underlying coagulation, though quite complex, has been well studied [34–40], and reliable experimental coagulation models have been developed [33, 41–43]. Coagulation is mediated by a family proteases in the circulation, called factors and a key group of blood cells, called platelets. The central process in coagulation is the conversion of prothrombin (fII), an inactive coagulation factor, to the master protease thrombin (FIIa). Thrombin generation involves three phases, initiation, amplification and termination [13,14]. Initiation requires a trigger event, for example vessel injury, which leads to the activation of factor VII (FVIIa). Two converging pathways, the extrinsic and intrinsic cascades, then process and amplify this initial coagulation signal. The extrinsic cascade is generally believed to be the main mechanism of thrombinogenesis in the blood [15–17]. Initially, thrombin is produced upon cleavage of prothrombin by fluid phase activated factor X (FXa), which itself has been activated by Tissue Factor/activated factor VII (TF/FVIIa) [10]. Picomolar amounts of thrombin then activate the cofactors factors V and VIII (fV and fVIII) and platelets, leading to the formation of the tenase and prothrombinase complexes on activated platelets. These complexes amplify the early coagulation signal by further activating FXa, and directly converting prothrombin to thrombin. There are several control points in the cascade that inhibit thrombin formation, and eventually terminate thrombin generation. Tissue Factor Pathway Inhibitor (TFPI) inhibits FXa formation catalyzed by

TF/FVIIa, while antithrombin III (ATIII) neutralizes several of the proteases generated during coagulation, including thrombin. Thrombin itself also inadvertently plays a role in its own inhibition; thrombin, through interaction with thrombomodulin, protein C and endothelial cell protein C receptor (EPCR), converts protein C to activated protein C (APC) which attenuates the coagulation response by proteolytic cleavage of fV/FVa and fVIII/FVIIIa. Termination occurs after either prothrombin is consumed, or thrombin formation is neutralized by inhibitors such as APC or ATIII.

To estimate coagulation model parameters, we used data sets from TF-VIIa initiated coagulation in the absence of anticoagulants. The objective function was a linear combination of two error functions that used data sets representing coagulation initiated with different concentrations of TF-VIIa (5pM, 5nM) [41]. We restricted the number of function evaluations to 4000 for each algorithm we tested, and performed 25 trials of each experiment. DOPS converges faster and has a lower final error compared to the other algorithms (Fig. 3). Within the first 1000 function evaluations of DOPS there is a very rapid drop in error. Approximately between 500-1000 function evaluations a switch to dynamically dimensioned search phase happens (this switch varies from trial to trial since the switch is based on the error from the swarm phase). Overall at the end of 4000 function evaluations DOPS minimizes the error (final objective error is 5.5916×10^4) to a much greater extent than any of the other algorithms. Using the parameters obtained at the end of 4000 function evaluations we examined the 'fits' between models predictions and experimental data (Fig.4). The solid lines represent the mean value of prediction over 25 trials and the shaded region represents the 99% confidence interval. Subsequently we used these optimal parameters to make model predictions that was compared against completely 'unseen' or untrained experimental data where coagulation was initiated with 500pM, 50pM, 10pM concentrations of TF-VIIa respectively (Fig.5).

Performance of DOPS on benchmark problems and test functions. Villaverde and co-workers recently published a set of benchmark biochemical problems to evaluate parameter estimation methods [31]. From a computational cost perspective problems they categorized the problems as most expensive, intermediate and least expensive. We evaluated the performance of our algorithms on a problem from the most expensive and least expensive categories. The first problem (most expensive) is a genome wide kinetic model of *Saccharomyces cerevisiae* with 261 reactions, 262 variables and 1759 parameters (henceforth referred to as problem B1). The reactions were modeled using modular rate law and generalized form of Michaelis-Menten kinetics. The second problem (henceforth referred to as problem B4, least expensive) is a metabolic model of Chinese Hamster Ovary (CHO) with 35 metabolites, 32 reactions and 117 parameters. In both cases pseudo time series data was generated by the authors. For problem B1, the time series data consisted of 44 observables and for problem B4 the data corresponded to 13 different metabolites. For a detailed description about the model architectures please refer to the work of Villaverde et al. [31]

We fixed the number of function evaluations at 4000 for DOPS and trained both the models against the pseudo experimental data. In both cases we found good fits (Fig. 8 and Fig. 9) to the problems within 4000 evaluations. We recaptured the 'nominal' parameters' in both cases within 4000 evaluations (Fig. 7). The final objective function value (Table 1) is an order smaller than nominal error for B1 possibly due to overfitting against the noise that was added to the synthetic data.

Having obtained good fits on the coagulation problem and the benchmarks we proceeded to compare the performance of DOPS with other algorithms on commonly used test functions for global optimization. We used a 300 dimensional Rastrigin function and 300 dimensional Ackley function. Both Ackley and Rastrigin have multiple local minima and maxima and attain a global minimum value of 0. We tested the performance of the 4

different heuristic approaches on these two functions. In each experiment we again fixed the number of function evaluations at 4000 and ran 25 experiments. In both cases (Fig. 6) we see that the error convergence rate for DOPS is much faster as compared to the other three heuristics and it finds the global minimum of 0 in both cases. In (Table 1) we summarize the results obtained on the coagulation model, the benchmark problems and the test functions.

Discussion

Our study presents a novel approach for high-dimensional parameter estimation in complex biological systems with relatively few function evaluations. In this approach we combined a variant of a well known meta heuristic particle swarm optimization with Dynamically Dimensioned Search (DDS). We tested our approach on an ODE model of coagulation with 148 parameters and 92 species. Coagulation is an ideal system to test our approach since the biology is well known and complex, with multiple feed back loops that are tightly regulated. We used experimental data under different conditions to obtain optimal parameters and used these parameters to make predictions against unseen experimental data. We obtained good fits and made sufficiently accurate enough predictions using parameters obtained from 4000 function evaluations. Further, we also used high-dimensional forms of commonly used test functions of global optimization and showed that we were able to find the global minimum for 300 dimensional Ackley and Rastrigin functions faster than other meta-heuristics. We also considered two recently published benchmark problems to test parameter optimization approaches and showed that we were able to retrieve the nominal parameter vector within 4000 function evaluations. Meta-heuristic approaches are generally effective in finding close to optimum solutions of complex, multimodal functions. In addition, they generally obviate the need for any *a priori* knowledge (like function derivative). However they take an exorbitant number of objective function evaluations to come close to an optimum. When the objective function evaluations tend to become expensive it is infeasible to take up a large number of evaluations. As the dimensionality of parameter space increases, the search region gets widened and thus the problem becomes more challenging. In addition, most of these approaches require optimization of 'algorithm parameters' before the actual optimization and also involve computationally expensive update operations. Tolson and Shoemaker, through DDS, showed that randomly perturbing a subset of dimensions in high dimensional parameter space is

an effective way to obtain near optimal solutions with few function evaluations. Though their approach is based on a single solution, the decision vector carries forward from one iteration to the next. Hence it behaves similar to a population based search. In our approach, we tried to eliminate the probability of starting from a bad region of search using a variant of particle swarm optimization. Particle Swarm Optimization (PSO) is a population based meta heuristic which does not have any complex operations like recombination, mutation or selection that are associated with other population based meta-heuristics like Differential Evolution (DE) or Genetic Algorithm (GA). Several particle swarm variants have been proposed to improve the search ability and rate of convergence, that involve different neighborhood structures, multi-swarms or adaptive parameters. Multi-swarm PSO with small particle neighborhoods have been shown to better in searching on complex multi-modal solutions [44]. Multi swarm methods, in addition, avoid rapid convergence to a local optimum or stable point and are able to generate diverse solutions. Generation of diverse solutions in the early stage gives a better exploratory capability and thus converge of upon multiple optima.

We utilized this capacity of multi swarms to generate diverse candidate solutions which can be used as initial solutions for a DDS like search. Thus the solution vectors obtained at the end of multi swarm search phase have a better propensity to not get stuck in bad regions in DDS phase as opposed to starting with a totally random initial solution. Given a fixed number of function evaluations, we have been able to show that we were able to obtain better solutions for the coagulation model and the test functions faster than other commonly known meta-heuristics and also DDS alone. The computational cost was prohibitively high for parameter estimation on this problem using a standard PSO. Thus we used SA, DE and DDS for the purpose of comparison. Choosing the number of function evaluations is largely a function of cost and complexity of the objective function. Traditionally the stopping condition for a parameter optimization problem can be the number

of function evaluations, percentage of initial error achieved or an absolute error threshold. However in case of complex, expensive functions where we desire a value within a certain period of time, the number of evaluations are used as a stopping criterion. In our current study we used a value of 4000 which we based upon the time taken (approximately 8-10 seconds) for a single objective function evaluation in the coagulation case. We used the same value of 4000 for benchmarks published by Villaverde and co-workers [31]. In this work by Villaverde et al. they used a population based search enhanced Scatter Search (eSS) to estimate the biochemical model parameters. Quite surprisingly we took a couple of orders lesser number of function evaluations 10^3 to obtain the optimal parameter vector as compared to the enhanced Scatter Search (eSS) with a local optimizer, which took around the order of 10^5 number of evaluations. The amount of CPU time taken (on an Intel Xeon processor 2.4 GHz) is lesser as compared to eSS on a similar architecture.

A surprisingly remarkable aspect about our algorithm has been that we did not 'pre optimize' any parameters of the algorithm to suit a specific problem. In the swarm phase this includes the number of particles, number of sub-swarms, acceleration constants or the number of generations after which the particles are redistributed and the neighborhood perturbation parameter in DDS phase. We used the same parameters for all the problems. The same rule was applied to the rest of the meta-heuristics barring Simulated Annealing. For SA, we optimized the cooling schedule for the coagulation model. Thus, in this approach any overhead that usually comes with additional function evaluations to pre optimize parameters was avoided.

The performance of our approach seems impressive given that it performed well on different complex systems with no pre optimization of algorithm parameters being required. We comfortably outperformed existing, widely used meta-heuristics and were also able to find minima of high dimensional global optimization test functions. Thus this approach may be well suited to large scale global optimization. In addition, surprisingly, we were

able to obtain optimal parameter vectors for two different large scale systems biology models with a couple of orders fewer number of function evaluations as compared to enhanced Scatter Search (eSS). However it is quite possible that highly optimized versions of common meta heuristics may outperform us on these systems. This aspect is currently beyond the scope of this study. Our approach can also be combined with local derivative based searches to improve upon the accuracy of the solutions. In addition, the current implementation of the algorithm is designed to switch only once from the swarm phase to the DDS. In the DDS phase, the search uses only one candidate vector although there is a provision to start with a population of candidate vectors. Incorporating a more intelligent switching strategy that can do switch from swarm phase to DDS phase multiple times and having a population of candidate vectors are some aspects of the algorithm that can be studied further.

Acknowledgements

This study was supported by an award from the Army Research Office (ARO #59155-LS).

References

1. Assmus HE, Herwig R, Cho KH, Wolkenhauer O (2006) Dynamics of biological systems: role of systems biology in medical research. *Expert Review of Molecular Diagnostics* .
2. van Riel NAW (2006) Dynamic modelling and analysis of biochemical networks: mechanism-based models and model-based experiments. *Briefings in Bioinformatics* 7: 364–374.

KEY: Riel:2006aa
ANNOTATION: 10.1093/bib/bbl040
3. Jaqaman K, Danuser G (2006) Linking data to models: data regression. *Nat Rev Mol Cell Biol* 7: 813–819.

KEY: Jaqaman:2006aa
ANNOTATION: 10.1038/nrm2030
4. Kitano H (2002) Systems biology: a brief overview. *Science* 295: 1662–1664.
5. Hood L, Heath JR, Phelps ME, Lin B (2004) Systems biology and new technologies enable predictive and preventative medicine. *Science* 306: 640–643.
6. Aldridge BB, Burke JM, Lauffenburger DA, Sorger PK (2006) Physicochemical modelling of cell signalling pathways. *Nat Cell Biol* 8: 1195–1203.

KEY: Aldridge:2006aa
ANNOTATION: 10.1038/ncb1497
7. Banga JR (2008) Optimization in computational systems biology. *BMC systems biology* 2: 47.
8. Ashyraliyev M, Fomekong-Nanfack Y, Kaandorp JA, Blom JG (2009) Systems biology: parameter estimation for biochemical models. *Febs Journal* 276: 886–902.

9. Moles CG, Mendes P, Banga JR (2003) Parameter estimation in biochemical pathways: a comparison of global optimization methods. *Genome research* 13: 2467–2474.
10. Nieman R, Fisher D, Seborg D (1971) A review of process identification and parameter estimation techniques†. *International Journal of Control* 13: 209–264.
11. Beck JV, Arnold KJ (1977) Parameter estimation in engineering and science. James Beck.
12. Young P (1981) Parameter estimation for continuous-time models—a survey. *Automatica* 17: 23–39.
13. Beck JV, Woodbury KA (1998) Inverse problems and parameter estimation: integration of measurements and analysis. *Measurement Science and Technology* 9: 839.
14. Hooke R, Jeeves TA (1961) “direct search” solution of numerical and statistical problems. *Journal of the ACM (JACM)* 8: 212–229.
15. Nelder JA, Mead R (1965) A simplex method for function minimization. *The computer journal* 7: 308–313.
16. Moré JJ (1978) The levenberg-marquardt algorithm: implementation and theory. In: *Numerical analysis*, Springer. pp. 105–116.
17. Esposito WR, Floudas CA (2000) Deterministic global optimization in nonlinear optimal control problems. *Journal of Global Optimization* 17: 97–126.
18. Horst R, Tuy H (2013) *Global optimization: Deterministic approaches*. Springer Science & Business Media.
19. Kirkpatrick S, Gelatt CD, Vecchi MP, et al. (1983) Optimization by simulated annealing. *science* 220: 671–680.
20. Storn R, Price K (1997) Differential evolution—a simple and efficient heuristic for global optimization over continuous spaces. *Journal of global optimization* 11: 341–359.
21. Sun J, Garibaldi JM, Hodgman C (2012) Parameter estimation using metaheuristics in

systems biology: a comprehensive review. *Computational Biology and Bioinformatics*,
IEEE/ACM Transactions on 9: 185–202.

22. Mendes P, Kell D (1998) Non-linear optimization of biochemical pathways: applications to metabolic engineering and parameter estimation. *Bioinformatics* 14: 869–883.

23. Modchang C, Triampo W, Lenbury Y (2008) Mathematical modeling and application of genetic algorithm to parameter estimation in signal transduction: Trafficking and promiscuous coupling of g-protein coupled receptors. *Computers in Biology and Medicine* 38: 574–582.

24. Tsai KY, Wang FS (2005) Evolutionary optimization with data collocation for reverse engineering of biological networks. *Bioinformatics* 21: 1180–1188.

25. Wang FS, Su TL, Jang HJ (2001) Hybrid differential evolution for problems of kinetic parameter estimation and dynamic optimization of an ethanol fermentation process. *Industrial & engineering chemistry research* 40: 2876–2885.

26. Noman N, Iba H (2007) Inferring gene regulatory networks using differential evolution with local search heuristics. *IEEE/ACM Transactions on Computational Biology and Bioinformatics (TCBB)* 4: 634–647.

27. Tashkova K, Korošec P, Šilc J, Todorovski L, Džeroski S (2011) Parameter estimation with bio-inspired meta-heuristic optimization: modeling the dynamics of endocytosis. *BMC systems biology* 5: 159.

28. Villaverde AF, Egea JA, Banga JR (2012) A cooperative strategy for parameter estimation in large scale systems biology models. *BMC systems biology* 6: 75.

29. Rodríguez-Fernández M, Egea JA, Banga JR (2006) Novel metaheuristic for parameter estimation in nonlinear dynamic biological systems. *BMC bioinformatics* 7: 483.

30. Egea JA, Rodríguez-Fernández M, Banga JR, Martí R (2007) Scatter search for chemical and bio-process optimization. *Journal of Global Optimization* 37: 481–503.

- 386 31. Villaverde AF, Henriques D, Smallbone K, Bongard S, Schmid J, et al. (2015)
387 Biopredyn-bench: a suite of benchmark problems for dynamic modelling in systems
388 biology. BMC systems biology 9: 8.
- 389 32. Fan M, Kuwahara H, Wang X, Wang S, Gao X (2015) Parameter estimation methods
390 for gene circuit modeling from time-series mrna data: a comparative study. Briefings
391 in bioinformatics : bbv015.
- 392 33. Luan D, Zai M, Varner JD (2007) Computationally derived points of fragility of a hu-
393 man cascade are consistent with current therapeutic strategies. PLoS computational
394 biology 3: e142.
- 395 34. Mann KG, Butenas S, Brummel K (2003) The dynamics of thrombin formation. Arte-
396 riosclerosis, thrombosis, and vascular biology 23: 17–25.
- 397 35. Mann K, Brummel K, Butenas S (2003) What is all that thrombin for? Journal of
398 Thrombosis and Haemostasis 1: 1504–1514.
- 399 36. Mann KG (2003) Thrombin formation. CHEST Journal 124: 4S–10S.
- 400 37. Vogler EA, Siedlecki CA (2009) Contact activation of blood-plasma coagulation. Bio-
401 materials 30: 1857–1869.
- 402 38. Diamond SL (2013) Systems biology of coagulation. Journal of Thrombosis and
403 Haemostasis 11: 224–232.
- 404 39. Fogelson AL, Tania N (2005) Coagulation under flow: the influence of flow-mediated
405 transport on the initiation and inhibition of coagulation. Pathophysiology of haemosta-
406 sis and thrombosis 34: 91–108.
- 407 40. Anand M, Rajagopal K, Rajagopal K (2003) A model incorporating some of the me-
408 chanical and biochemical factors underlying clot formation and dissolution in flowing
409 blood: review article. Journal of Theoretical Medicine 5: 183–218.
- 410 41. Hockin MF, Jones KC, Everse SJ, Mann KG (2002) A model for the stoichiometric
411 regulation of blood coagulation. Journal of Biological Chemistry 277: 18322–18333.

- 412 42. Chatterjee MS, Denney WS, Jing H, Diamond SL (2010) Systems biology of coagu-
413 lation initiation: kinetics of thrombin generation in resting and activated human blood.
414 PLoS computational biology .
- 415 43. Mann KG, Brummel-Ziedins K, Orfeo T, Butenas S (2006) Models of blood coagula-
416 tion. *Blood Cells, Molecules, and Diseases* 36: 108–117.
- 417 44. Zhao SZ, Liang JJ, Suganthan PN, Tasgetiren MF (2008) Dynamic multi-swarm par-
418 ticle swarm optimizer with local search for large scale global optimization. In: *Evo-*
419 *lutionary Computation, 2008. CEC 2008.(IEEE World Congress on Computational*
420 *Intelligence)*. IEEE Congress on. IEEE, pp. 3845–3852.

Table 1: Table with optimization settings and results for the coagulation problem, the benchmarks and test functions using DOPS. For each problem the bounds on the parameter vector, the total number of function evaluations, the best initial objective value and the best final objective value are specified. Nominal objective value represents the objective value using the true parameter vector or the nominal parameter vector. The CPU time is the time taken for the problem on a 2.4GHz Intel Xeon Architecture running Matlab 2014b.

	Coagulation	B1	B4	Ackley 300D	Rastrigin 300D
Lower Bound	0.001.pnom	5.pnom	5.pnom	30	5.12
Upper Bound	1000.pnom	0.2.pnom	0.2.pnom	-15	-5.12
CPU Time	10.0835 hours	38.308 hours	6.2 minutes	2.8 seconds	2.6 seconds
Function Evaluations	4000	4000	4000	4000	4000
Initial Objective Value	1.11837e+07	1.4275e+07	1.8536e+07	21.12	99.985
Final Objective Value	5.5916e+04	3.5348e+04	38.1306	0	0
Nominal Objective Value	4.7785e+06	1.0986e+06	39.0676	0	0

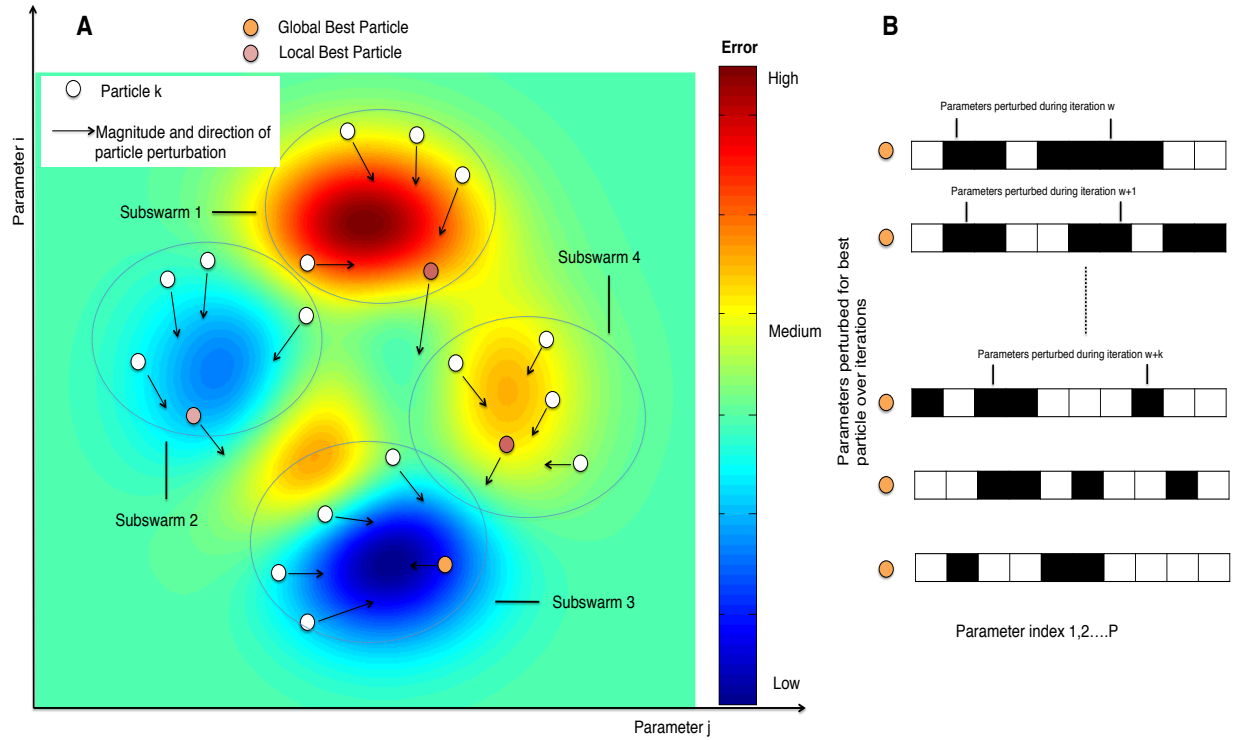


Fig. 1: Dynamic Optimization with Particle Swarms. **A:** Each particle represents an N dimensional parameter vector. Particles are given randomly generated initial solutions and grouped into different sub-swarms. Within each swarm the magnitude and direction of the movement a particle is influenced by the position of the best particle and also by its own experience. After every g number of function evaluations the particles are mixed and randomly assigned to different swarms. When the error due to the global best particle (best particle amongst all the sub-swarms - orange color) does not drop over a certain number of function evaluations, the swarm search is stopped and the search switches to a Dynamically Dimensioned Search with global best particle as the initial solution vector or candidate vector. **B:** The candidate vector performs a greedy global search for the remaining number of function evaluations. The search neighborhood is dynamically adjusted by varying the number of dimensions that are perturbed (in black) in each evaluation step. The probability that a dimension is perturbed decreases as the number of function evaluations increase. Thus as the evaluations increase the optimality of the solution is preserved.

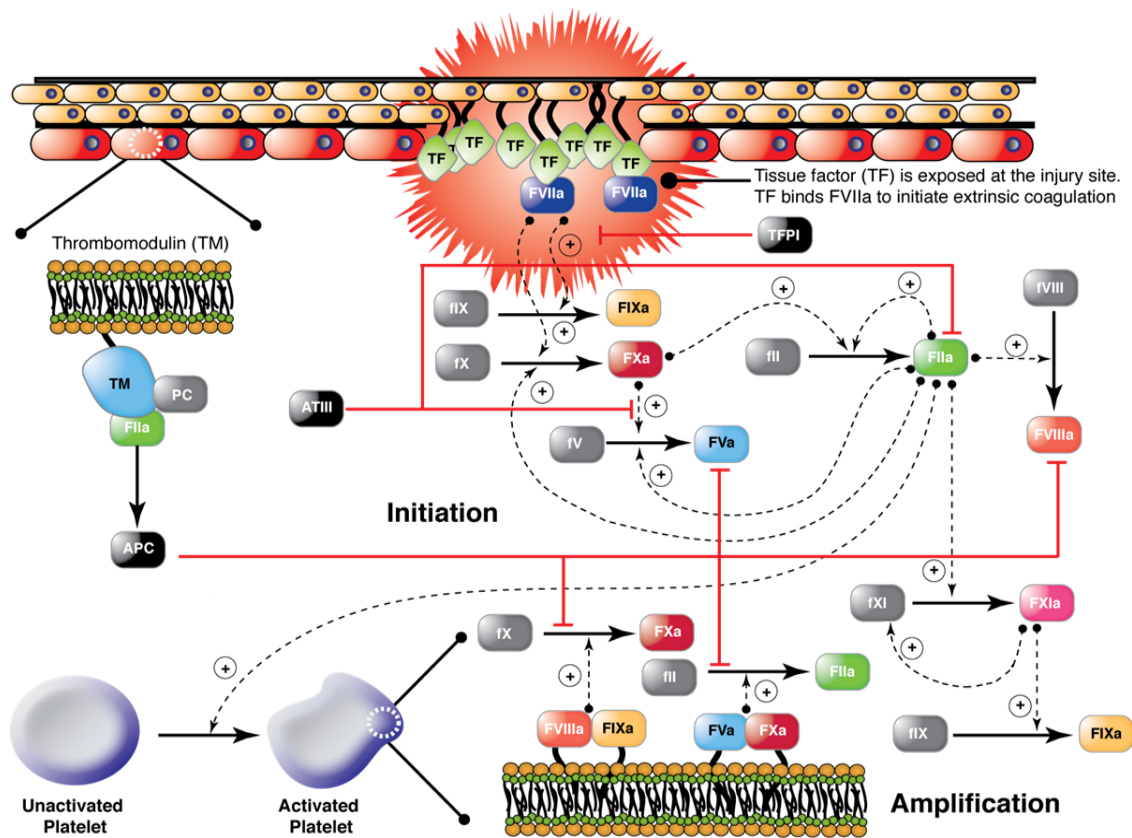


Fig. 2: Schematic of the extrinsic and intrinsic coagulation cascade[33]. Inactive zymogens upstream (grey) are activated by exposure to tissue factor (TF) following vessel injury. Tissue factor and activated factor VIIa (FVIIa) form a complex that activates factor X (fX) and IX (fIX). FXa activates downstream factors including factor VIII (fVIII) and fIX. Factor V (fV) is primarily activated by thrombin (FIIa). In addition, we included a secondary fV activation route involving FXa. FXa and FVa form a complex (prothrombinase) on activated platelets that converts prothrombin (fII) to FIIa. FIXa and FVIIIa can also form a complex (tenase) on activated platelets which catalyzes FXa formation. Thrombin also activates upstream coagulation factors, forming a strong positive feedback ensuring rapid activation. Tissue factor pathway inhibitor (TFPI) down-regulates FXa formation and activity by sequestering free FXa and TF-FVIIa in a FXa-dependent manner. Antithrombin III (ATIII) inhibits all proteases. Thrombin inhibits itself binding the surface protein thrombomodulin (TM). The Ila-TM complex catalyzes the conversion of protein C (PC) to activated protein C (APC), which attenuates the coagulation response by the proteolytic cleavage of fV/FVa and fVIII/FVIIIa.

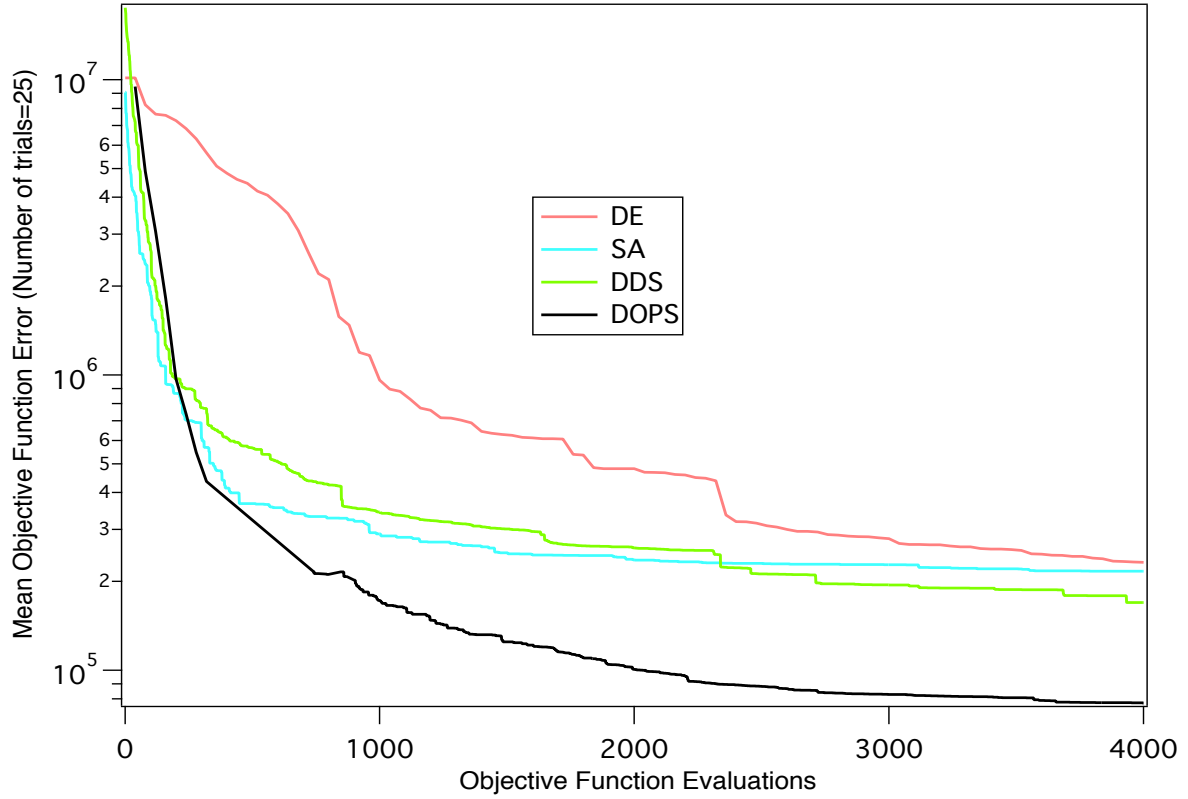


Fig. 3: Error convergence rates of the 5 different algorithms on the coagulation model. The objective error is the mean over $N=25$ trials. DOPS, DDS and SA have the steepest drop in error during first 300 function evaluations. Thereafter the error drop in DDS and SA remains nearly constant whereas DOPS continues to drops further. At the end of 4000 function evaluations DOPS attains the lowest error. The next best estimate using DDS is nearly 3 times greater than the lowest error using DDS.

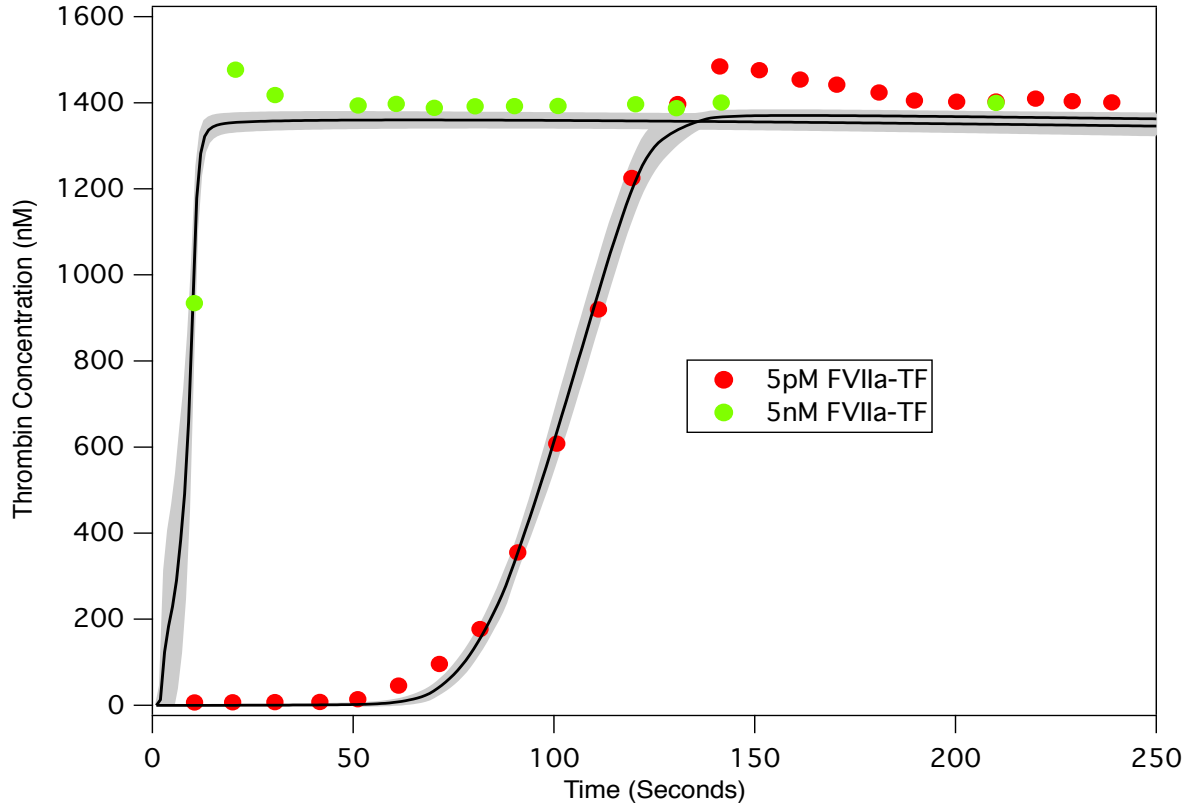


Fig. 4: Model fits on experimental data using DDSMLSPSO. The model parameters were estimated using DOPS. Solid black lines indicate the simulated mean thrombin concentration using parameter vectors from $N=25$ trials. The grey shaded region represents the 99% confidence estimate of the mean simulated thrombin concentration. The experimental data is reproduced from the synthetic plasma assays of Mann and co-workers. Thrombin generation is initiated by adding Factor VIIa-TF (5nM - Red and 5pM - Green) to synthetic plasma containing 200 $\mu\text{mol/L}$ of phospholipid vesicles (PCPS) and a mixture of coagulation factors (II,V,VII,VIII,IX,X and XI) at their mean plasma concentrations.

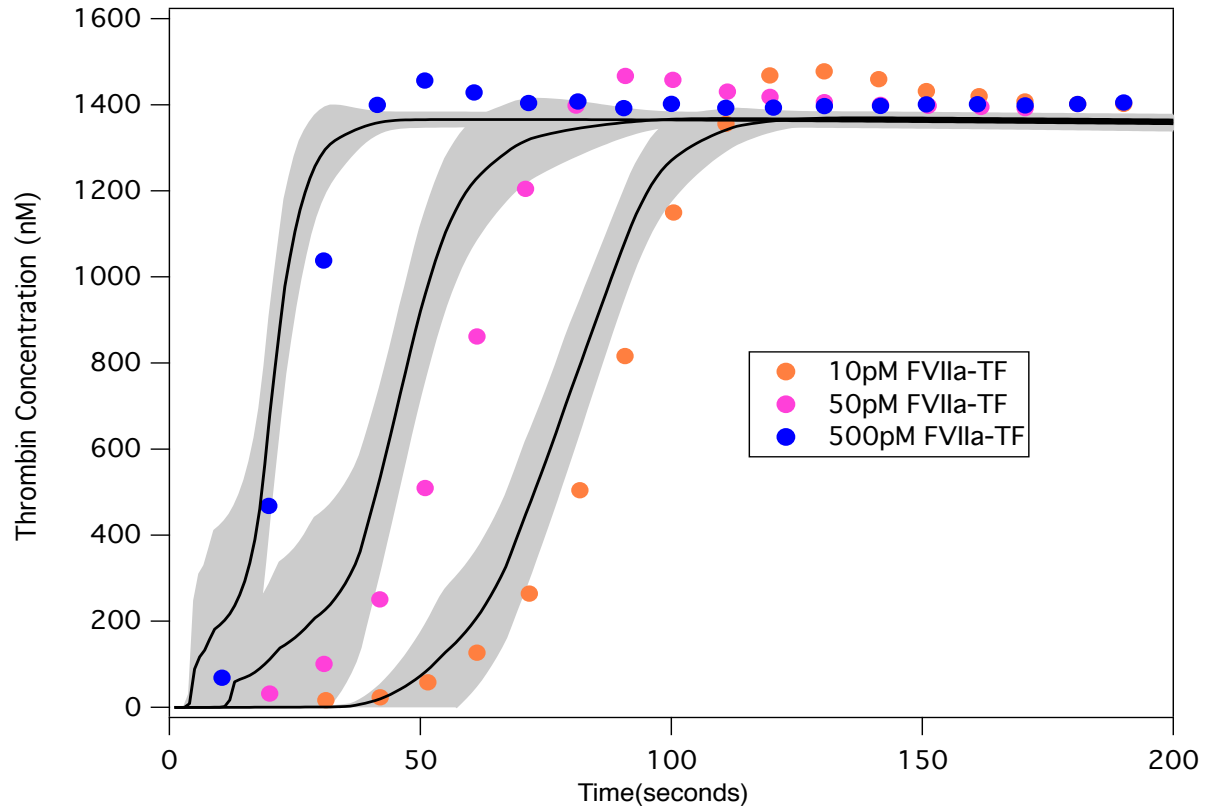


Fig. 5: Model predictions on unseen experimental data using parameters obtained from DOPS. The parameter estimates that were obtained using DOPS were tested against data that was not used in the model training. Solid black lines indicate the simulated mean thrombin concentration using parameter vectors from $N=25$ trials. The grey shaded region represents the 99% confidence estimate of the mean simulated thrombin concentration. The experimental data is reproduced from the synthetic plasma assays of Mann and co-workers. Thrombin generation is initiated by adding Factor VIIa-TF (500pM - Blue, 50pM - Pink and 10pM - Orange respectively) to synthetic plasma containing 200 $\mu\text{mol/L}$ of phospholipid vesicles (PCPS) and a mixture of coagulation factors (II,V,VII,VIII,IX,X and XI) at their mean plasma concentrations.

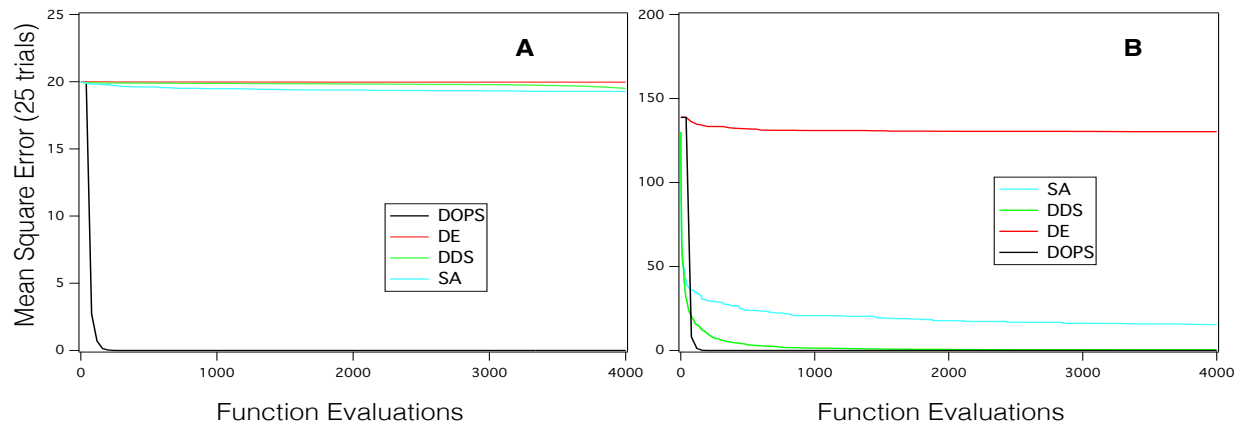


Fig. 6: Error convergence on 300 dimensional Ackley function and Rastrigin function. **(A)** Error convergence of 4 different meta-heuristics on 300 dimensional Ackley function **(B)** Error convergence of 4 different meta-heuristics on 300 dimensional Rastrigin function

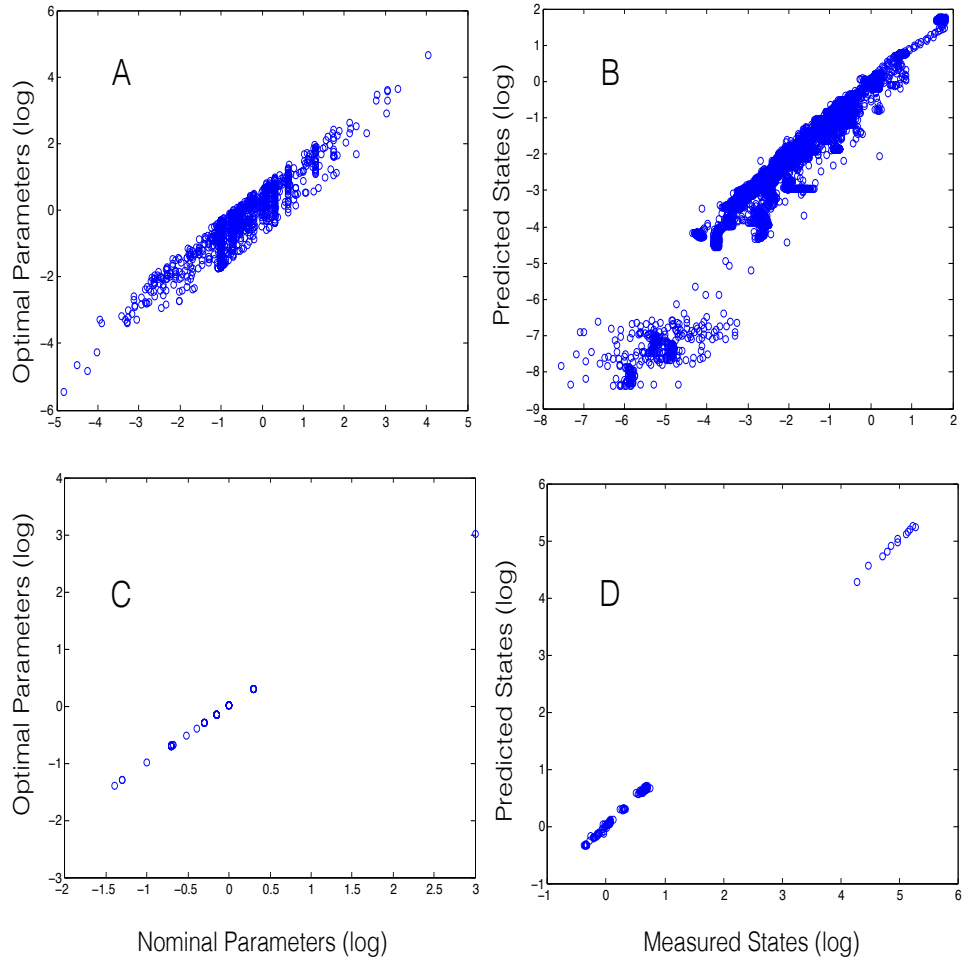


Fig. 7: (A) Difference between nominal and optimal parameters for problem B1: Genome wide kinetic model of *S.cerevisiae* with 1759 unknown parameters. **(B)** Difference between experimental (measured) data and data simulated with optimal parameters for problem B1: Genome wide kinetic model of *S.cerevisiae* with 1759 unknown parameters. **(C)** Difference between nominal and optimal parameters for problem B4: Metabolic model of Chinese Hamster Ovary Cells (CHO) cells with 117 parameters. **(D)** Difference between experimental (measured) data and data simulated with optimal parameters for problem B4: Metabolic model of Chinese Hamster Ovary Cells (CHO) cells with 117 parameters.

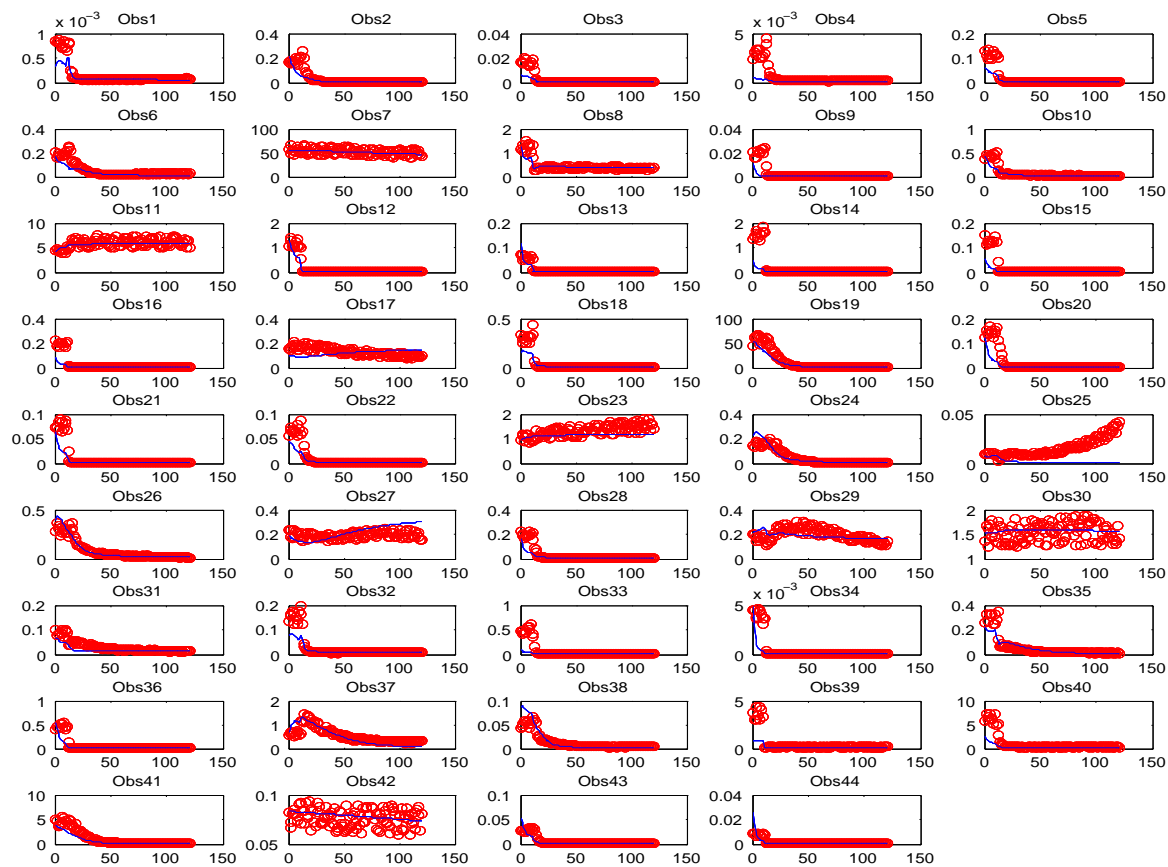


Fig. 8: (Data fits for Problem B1) Pseudo-experimental data (red circles) vs. optimal solution obtained using DOPS (solid blue lines) for the 44 observed states. X axis: time [s]; Y axis: metabolite concentrations [mM].

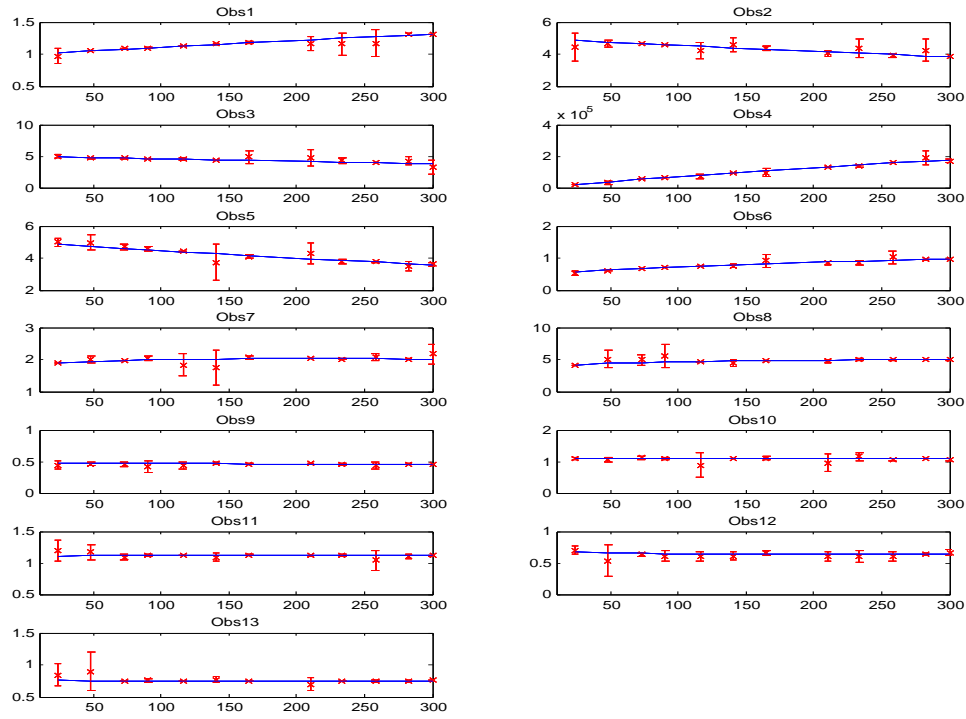


Fig. 9: (Data fits for Problem B4) Pseudo-experimental data (red x) vs. optimal solution obtained using DOPS (solid blue lines) for the 13 observed states. X axis: time [s]; Y axis: metabolite concentrations [mM].