# Dynamic Modeling of the Human Coagulation Cascade using Reduced Order Effective Kinetic Models

Adithya Sagar and Jeffrey D. Varner\*

School of Chemical and Biomolecular Engineering Cornell University, Ithaca NY 14853

Running Title: Modeling blood coagulation

To be submitted: Processes

\*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**

Keywords: Coagulation, Mathematical modeling, Systems Biology

#### Introduction

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The human coagulation system is an archetype proteolytic cascade regulated by both positive and negative feedback loops. The biology of coagulation is well studied [1-4]. Coagulation is mediated by a family serine proteases, called factors and a key group of blood cells, called platelets. The activation of coagulation factors and platelets, which are normally inactive in the circulation, requires a trigger event. Trigger events such as vessel injury expose procoagulant materials like collagen, fibronectin, von Willebrand factor (vWF) or tissue factor (TF). These materials drive platelet localization and activation, coagulation factor activation and ultimately clot formation [5]. Once coagulation has been activated, two converging pathways activate the master protease thrombin. The extrinsic cascade is generally believed to be the main mechanism of thrombinogenesis in the blood [1, 2, 6]. Thrombin generation consists of three phases, initiation, propagation and termination [7, 8]. Initially thrombin is produced upon cleavage of prothrombin by fluid phase activated factor X (FXa) [9]. Picomolar amounts of thrombin can then activate the cofactors factors V and VIII (fV and fVIII) and platelets, resulting in the formation of the tenase 15 and prothrombinase complexes on the surface of activated platelets. These complexes 16 amplify the early coagulation signal by activating downstream and upstream coagulation 17 factors. Termination occurs after prothrombin is consumed or activated thrombin is neu-18 tralized by inhibitors such as activated protein C (APC) or antithrombin III (ATIII). 19

Malfunctions in coagulation can have serious or potentially even fatal consequences. For example, aggressive clotting is involved with Coronary Artery Diseases (CADs), which collectively account for 38% of all deaths in North America [10]. Coagulation management during surgery can also be challenging, particularly because of the increasing clinical use of antithrombotic drugs [11]. Insufficient coagulation due to genetic disorders such as hemophilia can also result in recurrent bleeding. The coagulation factors VIII (fVIII) and IX (fIX) are deficient in Hemophilia A and B, respectively [12–14]. People with mild

hemophilia have 5-40% of the normal clotting factor levels while severe hemophiliacs have <1% [14]. Hemophilia can be controlled with regular infusions of the deficient clotting factors. However, clotting factor replacement sometimes leads to the formation of in-vivo fVIII and fIX inhibitors [15]. Activated Prothrombin Complex Concentrates (aPCCs) from 30 pooled plasma have been used to treat hemophilia. However, aPCCs can have safety 31 issues [16-18]. Alternatively, recombinant factor VIIa (rFVIIa) has been used to treat bleeding disorders [19, 20] including hemophilia with and without factor VIII/IX inhibitors 33 [21-24]. However, rFVIIa requires frequent administration (every 2-3 hr) because of its 34 short half-life in the circulation. Many questions also remain about its mechanism [9, 25? 35 ? ? -28] and effective dose range [15]. Thus, despite its wide adoption, the utility of 36 rFVIIa in trauma-associated hemorrhage remains controversial [?]. 37

Modeling approaches differ in their degree of mechanistic detail, and the choice of ap-38 proach is often determined by prior system knowledge [29]. The ability of ODE modeling to capture dynamics has made it one of the most common tools for studying signaling systems. However, dynamics and ODEs come at the expense of difficult (and often impossible) to solve model parameter identification problems. For example, Gadkar et al., showed that even with near-perfect information and high frequency sampling, it was often impossible to identify (to within a specified uncertainty) all the parameters in typical signal transduction models [30]. This reality highlights the perhaps under-appreciated role that experimental design could play in generating the best training and validation data sets for model identification [31]. Moreover, detailed ODE models require significant mechanistic knowledge of the underlying biology. However, as Bailey suggested more than a decade ago, achieving qualitative and quantitative understanding of complex biological systems should not require complete structural and parametric knowledge [32]. Since 50 Bailey's complex biology with no parameters hypothesis, Sethna and coworkers showed 51 that model behavior and predictive ability were sensitive to only a few parameter combina-

tions, a characteristic seemingly universal to multi-parameter signaling models referred to as sloppiness [33]. Thus, reasonable model predictions could be possible with limited parameter information. Taking advantage of this property of model behavior, we developed sloppy techniques for parameter identification using ensembles of deterministic models. 56 Pareto optimal ensemble techniques (POETs) incorporate principles of competing objec-57 tives into a multi-objective optimization framework, enabling the exploration of kinetic parameter space while accounting for uncertainty and potential conflicts in the experimental 59 training data [34]. We have proposed that the sloppy behavior of biological networks may 60 also be a source of cell-to-cell [?] or even patient-to-patient heterogeneity [35]. Recently, 61 Bayesian techniques of parameter identification have also been used to explore cell-tocell heterogeneity [36, 37]. Thus, a population of cells could be viewed as a dynamic 63 ensemble of networks as the operational biochemical pathways are often context-specific 64 [38]. 65

In this study, we constructed a reduced order effective model of thrombin generation 66 in normal and hemophilic blood. We used this model to understand how physiological 67 coagulation was altered by prothrombin levels, the activated protein C pathway and the level of ATIII. Previous models of coagulation have been formulated as systems of nonlinear ordinary differential equations, often using mass-action kinetics to describe the rates of biochemical conversions [REFHERE]. While this approach has proven highly effective, it does require significant experimental data for model identification and validation. For example, a previous coagulation model from our laboratory, which described 193 proteins and protein complexes interconnected by 301 biochemical interactions, contained 467 unknown parameters (301 kinetic parameters and 166 initial conditions) [REFHERE]. The key innovation of our approach is the seamless integration of simple effective rules encoding complex regulatory motifs with traditional kinetic pathway modeling. This inte-77 gration allows the description of complex regulatory interactions, such as time-dependent

- 79 allosteric regulation of enzyme activity, in the absence of specific mechanistic informa-
- 80 tion. The regulatory rules are easy to understand, easy to formulate and do not rely on
- overarching theoretical abstractions or restrictive assumptions. We tested our approach
- by modeling the time evolution of the human coagulation cascade.

### 83 Results

- Formulation and properties of reduced order effective models. Reduced order ef-
- 85 fective models are [FINISH ME].

Identification of reduced order coagulation model parameters using particle swarm optimization. A critical challenge for any dynamic model is the estimation of kinetic parameters. We estimated kinetic and control parameters simultaneously from in vitro coagulation data sets with and without the protein C pathway. The residual between model 89 simulations and experimental measurements was minimized using particle swarm opti-90 mization (PSO). A population of particles (N = 20) was initialized with randomized kinetic 91 and control parameters and allowed to search for parameter vectors that minimized the 92 residual. However, not all parameters were varied simultaneously. We partitioned the pa-93 rameter estimation problem into two subproblems based upon the biological organization 94 of the training data; (i) estimation of parameters associated with thrombin formation in the 95 absence of the protein C pathway and (ii) estimation of parameters associated with the 96 protein C pathway. Only those parameters associated with each subproblem were varied 97 during the optimization procedure for that subproblem e.g., thrombin parameters were not varied during the protein C subproblem. The PSO procedure was run for 20 generations for each subproblem, where each generation was 1200 iterations. The best particle from 100 each generation was used to generate the particle population for the next generation. We 101 rotated the subproblems, starting with subproblem 1 in the first generation. 102

The reduced order coagulation model captured the role of initial prothrombin abundance, and the decay of the thrombin signal following from ATIII activity (Fig. 3). However, we systematically under-predicted the thrombin peak and the strength of ATIII inhibition in this training data set. On the other hand, with fixed thrombin parameters, we captured peak thrombin values and the decay of the thrombin signal (at least for the 150% fII case) in the presence of both ATIII and the protein C pathway (Fig. 4). Lastly, we were unable to capture global differences in initiation time *across* separate data sets with a single ensemble of model parameters. These differences likely result from normal experimental variability. For example different thrombin generation experiments within our training data

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(at the same physiological factor levels) have significantly different initiation times (supplemental results). However, this also highlights a potential shortcoming of the initiation 113 module within the model. To capture the variability in initiation time across training data sets, we included a constant time-delay parameter  $(T_D)$  for each data group. The delay 115 parameter was constant within a data set, but allowed to vary across training data sets. 116 Introduction of the delay parameter allowed the model to simulate multiple training data 117 sets using a single ensemble of model parameters. Taken together, the model identifica-118 tion results suggested that our kinetic-rules based approach could reproduce a panel of 119 thrombin generation data sets conducted at physiological factor and inhibitors concentra-120 tions. However, it was unclear whether the reduced order model could predict new data, 121 without updating the model parameters. 122

Validation of the reduced order coagulation model. We tested the predictive power of the reduced order coagulation model with validation data sets not used during model 124 training. Two validation data sets were used, thrombin generation for various prothrombin 125 and ATIII concentrations with the protein C pathway, and thrombin generation in normal 126 versus hemophilic plasma in the presence of the protein C pathway. Lastly, we compared 127 the qualitative output of the model to rFVIIa addition in the presence of hemophilia. The 128 hemophilia case was an especially difficult test as it was taken from a different study 129 which used a plasma-based in vitro assay involving platelets instead of phospholipid vesi-130 cles (PCPS). All kinetic and control parameters were fixed for the validation simulations. 131 The only globally adjustable parameter  $T_D$ , was fixed within each validation data set but 132 allowed to vary between data sets. The reduced order model predicted the thrombin 133 generation profile for ratios of prothrombin and ATIII in the absence of the protein C path-134 way (Fig. 5). Simulations near the physiological range (fII,ATIII) = (100%, 100%) or 135 (125%,75%) tracked the measured thrombin values (Fig. 5B and C). On the other hand, 136 predictions for factors levels outside of the physiological range (fll,ATIII) = (50%, 150%)

or (150%, 50%), while qualitatively consistent with measured thrombin values, did show significant deviation from the measurements (Fig. 5A and D). Likewise, simulations of 139 thrombin generation in normal versus hemophilia (missing both fVIII and fIX) were con-140 sistent with measured thrombin values (Fig. 6). We modeled the dependence of thrombin amplification on factor levels using a product rule ( $\mathcal{Z} = fV \times fX \times fVIII \times fIX$ ), which 142 was then was integrated using a min integration rule into the control variable governing 143 amplification. Thus, in the absence of fVIII or fIX, the amplification control variable evalu-144 ated to zero, and the only thrombin produced was from initiation (Fig. 6B). However, the 145 decay of the thrombin signal was underpredicted in the normal case (Fig. 6A), while the 146 activated thrombin level was overpredicted in hemophilia simulations, although thrombin 147 generation was far less than normal (Fig. 6B). Taken together, the reduced order model 148 performed well in the physiological range of factors, even with unmodeled components 149 such as platelet activation in the hemophilia data set. Thus, our kinetic-rules framework 150 predicted the output of a physiologically important cascades such as coagulation, despite significant unmodeled components. 152

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The model ensemble predicted a direct correlation between thrombin generation and rFVIIa addition in hemophilia (Fig. 7). In the current proof of concept model, we cannot distinguish between different initiation sources, e.g., TF/FVIIa and rFVIIa, as we have only a single lumped initiation source (trigger). Thus, we simulated the addition of rFVIIa in hemophilia by removing fVIII and fIX from the model, and modulating the level of trigger. The baseline level of trigger is consistent with the previous hemophilia simulations, where the only thrombin being produced is from initiation mechanisms (Fig. 7,  $1 \times$  trigger). As we increased the strength of the trigger event, the thrombin peak time and the maximum value of thrombin increased (Fig. 7, 50× trigger). However, as the trigger strength increased, the thrombin generated quickly decayed. For example, for the highest trigger strength simulated (50×trigger), 95% of the thrombin was gone by 20 min after initiation.

We performed flux analysis to understand how the reduced order coagulation model 164 balanced initiation, amplification and inhibition of thrombin formation for normal coagula-165 tion and hemophilia. Analysis of the reaction flux through the reduced order network for 166 thrombin generation in normal, hemophilia and rFVIIa-treated hemophilia identified three 167 distinct operational modes (Fig. 8). The reduced order network includes four lumped re-168 actions, initiation, amplification, thrombin-induced APC generation and total thrombin in-169 hibition (including both APC and ATIII action). Directly after the addition of a trigger (e.g., 170 TF/FVIIa or rFVIIa), the lumped initiation flux was the largest for all three cases. However, 171 within a few minutes enough thrombin was generated by the initiation mechanism to in-172 duce the amplification stage. During amplification, thrombin catalyzes its own formation 173 and inhibition by generating activated protein C (APC), a potent inhibitor of the coagulation 174 cascade. For normal coagulation, amplification and thrombin inhibition are the dominate 175 reactions by 6 min after initiation (Fig. 8, left). After 10 min, the dominate reaction has 176 shifted to thrombin inhibition (both ATIII and APC action). In the current proof-of-principle 177 model, APC inhibits upstream of amplification thus APC activity will slow the rate of throm-178 bin formation directly from the initiation trigger. On the other hand, ATIII inhibits thrombin directly, as well as upstream coagulation factors. In hemophilia (missing both fVIII and fIX), the amplification reaction does not occur and the only thrombin produced is from initiation (Fig. 8, center). Initiation is quickly inhibited by APC, and the thrombin level 182 stabilizes and eventually decays because of ATIII activity. Lastly, when 50×rFVIIa is used 183 to induce thrombin formation in hemophilia (absence of fVIII/fIX), initiation mechanisms 184 dominate for up to 6 min following initiation (Fig. 8, right). However, similar to hemophilia 185 alone, without amplification the thrombin signal is quickly extinguished by the combined 186 action of ATIII and APC generated by thrombin. 187

Sensitivity analysis of the ensemble of reduced order coagulation models. We conducted a variance based global sensitivity analysis to better understand which pa-

rameters were controlling the performance of the reduced order coagulation model. In particular, we calculated the sensitivity of two performance metrics, the time to maximum thrombin and the thrombin exposure (area under the thrombin curve) to changes in the model parameters (Fig. ZZ).

#### Discussion

Controlling hemophilia and more generally hemorrhage has been a driving force to under-195 stand the molecular basis of blood coagulation. Replacement therapy improves bleeding 196 times for most hemophilia patients. However, repeated fVIII/fIX administration can lead 197 to neutralizing inhibitors. Alternatively, rFVIIa has been used as a universal hemostatic 198 agent to initiate clotting with and without fVIII or fIX inhibitors [19]. Despite its wide adop-199 tion, the mechanism and the utility of rFVIIa in trauma-associated hemorrhage remains 200 controversial [?]. In this study, we formulated a model of the human coagulation cascade 201 and studied thrombin formation in normal and rFVIIa treated and untreated hemophilic plasma. The model described 193 proteins or protein complexes connected by 301 interactions. The coagulation network architecture was based on literature and a previous 204 mathematical model from our laboratory [? ]. Simulations of thrombin dynamics were independently validated using blood drawn from patients with coronary artery disease 206 (reported here) and previous TF/FVIIa and FVIIa in-vitro studies [3]. While the human 207 coagulation cascade is perhaps an ideal model system to develop network analysis tools, 208 not all the model parameters were identifiable given the current training data. Instead, an 209 ensemble of models (N = 437) was estimated using nine cell-based coagulation training 210 sets [?]. We used this family of consistent coagulation models in all simulations instead 211 of a single best-fit but uncertain model. Additionally, using sensitivity analysis on the 212 ensemble of models, we identified structurally sensitive components of the coagulation 213 architecture as a function of condition. We also demonstrated that the ensemble robustly 214 constrained model predictions of independent validation sets, despite having many poorly 215 constrained parameters. Thus, the model ensemble displayed sloppy behavior similar to 216 that observed by Sethna and coworkers for other signal transduction networks [? ]. By 217 studying a family of models, we perhaps partially addressed the uncertainty stemming 218 from the many poorly characterized model parameters. However, many other factors could influence our results. For example, missing structural interactions or biophysical factors such as blood flow could play a large role. Thus, while the results presented here

may be a valuable first step, more studies are required.

#### **Materials and Methods**

Formulation and solution of the model equations. We used ordinary differential equations (ODEs) to model the time evolution of proteins  $(x_i)$  in our reduced order coagulation model:

$$\frac{dx_i}{dt} = \sum_{j=1}^{\mathcal{R}} \sigma_{ij} r_j \left( \mathbf{x}, \epsilon, \mathbf{k} \right) \qquad i = 1, 2, \dots, \mathcal{M}$$
 (1)

where  $\mathcal{R}$  denotes the number of reactions,  $\mathcal{M}$  denotes the number of protein species in the model. The quantity  $r_j\left(\mathbf{x},\epsilon,\mathbf{k}\right)$  denotes the rate of reaction j. Typically, reaction jis a non-linear function of biochemical species abundance, as well as unknown kinetic parameters  $\mathbf{k}$  ( $\mathcal{K} \times 1$ ). The quantity  $\sigma_{ij}$  denotes the stoichiometric coefficient for species i in reaction i. If i is i on metabolite i is produced by reaction i. Conversely, if i on metabolite i is consumed by reaction i, while i is indicates metabolite i is not connected with reaction i. Lastly, i denotes the scaled enzyme degradation constant. The system material balances were subject to the initial conditions i is i on i o

Each reaction rate was written as the product of two terms, a kinetic term  $(\bar{r}_j)$  and a regulatory term  $(v_j)$ :

$$r_i(\mathbf{x}, \epsilon, \mathbf{k}) = \bar{r}_i v_i \tag{2}$$

We used multiple saturation kinetics to model the reaction term  $\bar{r}_i$ :

$$\bar{r}_j = k_j^{max} \epsilon_i \left( \prod_{s \in m_j^-} \frac{x_s}{K_{js} + x_s} \right) \tag{3}$$

where  $k_j^{max}$  denotes the maximum rate for reaction j,  $\epsilon_i$  denotes the scaled enzyme activity which catalyzes reaction j, and  $K_{js}$  denotes the saturation constant for species s in reaction j. The product in Eqn. (3) was carried out over the set of *reactants* for reaction j (denoted as  $m_j^-$ ).

The control term  $v_j$  depended upon the combination of factors which influenced the activity of enzyme i. For each enzyme, we used a rule-based approach to select from competing control factors (Fig. 2). If an enzyme was activated by m metabolites, we modeled this activation as:

$$v_j = \max\left(f_{1j}\left(\mathcal{Z}\right), \dots, f_{mj}\left(\mathcal{Z}\right)\right) \tag{4}$$

where  $0 \le f_{ij}(\mathcal{Z}) \le 1$  was a regulatory transfer function that calculated the influence of metabolite i on the activity of enzyme j. Conversely, if enzyme activity was inhibited by a m metabolites, we modeling this inhibition as:

$$v_{j} = 1 - \max\left(f_{1j}\left(\mathcal{Z}\right), \dots, f_{mj}\left(\mathcal{Z}\right)\right) \tag{5}$$

Lastly, if an enzyme had both m activating and n inhibitory factors, we modeled the regulatory term as:

$$v_i = \min\left(u_i, d_i\right) \tag{6}$$

251 where:

$$u_{j} = \max_{j^{+}} \left( f_{1j} \left( \mathcal{Z} \right), \dots, f_{mj} \left( \mathcal{Z} \right) \right) \tag{7}$$

$$d_{j} = 1 - \max_{j^{-}} \left( f_{1j} \left( \mathcal{Z} \right), \dots, f_{nj} \left( \mathcal{Z} \right) \right)$$
 (8)

The quantities  $j^+$  and  $j^-$  denoted the sets of activating and inhibitory factors for enzyme j.

If an enzyme had no allosteric factors, we set  $v_j=1$ . There are many possible functional forms for  $0 \le f_{ij}(\mathcal{Z}) \le 1$ . However, in this study, each individual transfer function took the form:

$$f_i(\mathbf{x}) = \frac{\kappa_{ij}^{\eta} \mathcal{Z}_j^{\eta}}{1 + \kappa_{ij}^{\eta} \mathcal{Z}_j^{\eta}} \tag{9}$$

where  $\mathcal{Z}_j$  denotes the abundance of the j factor (e.g., metabolite abundance), and  $\kappa_{ij}$  and  $\eta$  are control parameters. The  $\kappa_{ij}$  parameter was species gain parameter, while  $\eta$  was a cooperativity parameter (similar to a Hill coefficient). The model equations were encoded using the Python programming language and solved using the ODEINT routine of the SciPy module [39].

Estimation of model parameters from experimental data. Model parameters were estimated by minimizing the difference between simulations and experimental thrombin measurements (squared residual):

$$\min_{\mathbf{k}} \sum_{\tau=1}^{\mathcal{T}} \sum_{j=1}^{\mathcal{S}} \left( \frac{\hat{x}_j(\tau) - x_j(\tau, \mathbf{k})}{\omega_j(\tau)} \right)^2$$
 (10)

where  $\hat{x}_{j}\left( au\right)$  denotes the measured value of species j at time au,  $x_{j}\left( au,\mathbf{k}\right)$  denotes the 264 simulated value for species j at time  $\tau$ , and  $\omega_{j}\left( au\right)$  denotes the experimental measure-265 ment variance for species j at time  $\tau$ . The outer summation is respect to time, while 266 the inner summation is with respect to state. We minimized the model residual using 267 Particle swarm optimization (PSO) [40]. PSO uses a swarming metaheuristic to explore 268 parameter spaces. A strength of PSO is its ability to find the global minimum, even in the 269 presence of potentially many local minima, by communicating the local error landscape 270 experienced by each particle collectively to the swarm. Thus, PSO acts both as a local 271 and a global search algorithm. For each iteration, particles in the swarm compute their 272 local error by evaluating the model equations using their specific parameter vector realization. From each of these local points, a globally best error is identified. Both the local 274 and global error are then used to update the parameter estimates of each particle using 276 the rules:

$$\Delta_i = \theta_1 \Delta_i + \theta_2 \mathbf{r}_1 \left( \mathcal{L}_i - \mathbf{k}_i \right) + \theta_3 \mathbf{r}_2 \left( \mathcal{G} - \mathbf{k}_i \right)$$
(11)

$$\mathbf{k}_i = \mathbf{k}_i + \mathbf{\Delta}_i \tag{12}$$

where  $(\theta_1,\theta_2,\theta_3)$  are adjustable parameters,  $\mathcal{L}_i$  denotes local best solution found by particle i, and  $\mathcal{G}$  denotes the best solution found over the entire population of particles. The quantities  $r_1$  and  $r_2$  denote uniform random vectors with the same dimension as the number of unknown model parameters ( $\mathcal{K} \times 1$ ). In thus study, we used  $(\theta_1,\theta_2,\theta_3)=(1.0,0.05564,0.02886)$ . The quality of parameter estimates was measured using two criteria, goodness of fit (model residual) and angle between the estimated parameter vector  $\mathbf{k}_j$  and the best parameter set  $\mathbf{k}^*$ :

$$\alpha_j = \cos^{-1}\left(\frac{\mathbf{k}_j \cdot \mathbf{k}^*}{\|\mathbf{k}_i\| \|\mathbf{k}^*\|}\right) \tag{13}$$

If the candidate parameter set  $k_j$  were perfect, the residual between the model and synthetic data and the angle between  $k_j$  and the true parameter set  $k^*$  would be equal to zero. The particle swarm optimization routine was implemented in the Python programming language. All plots were made using the Matplotlib module of Python [41].

## **Acknowledgements**

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

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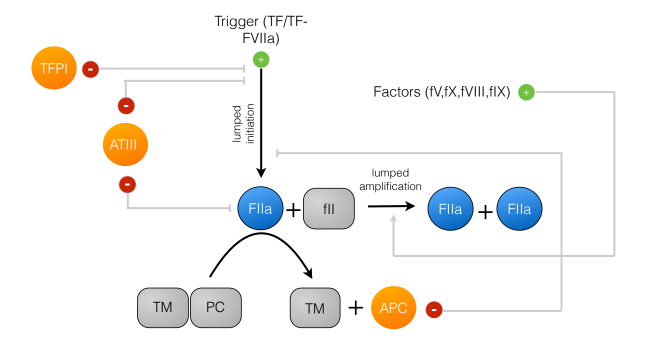
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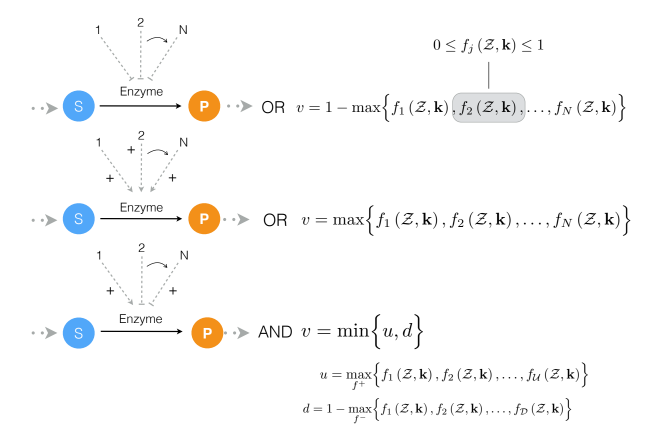
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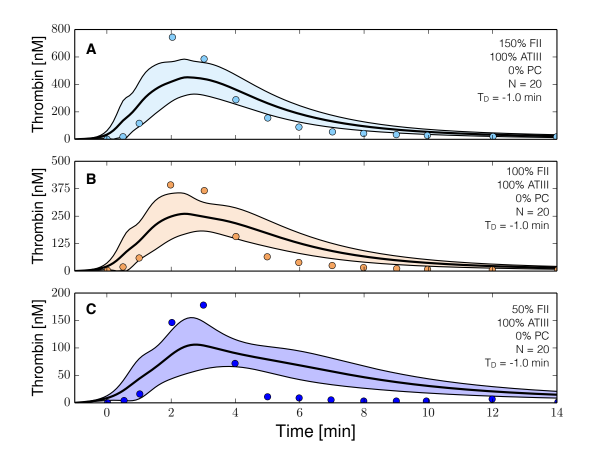
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**Fig. 1:** Proof of concept cell-free metabolic networks considered in this study. Substrate S is converted to products  $P_1$  and  $P_2$  through a series of chemical conversions catalyzed by enzyme(s)  $E_j$ . The activity of the pathway enzymes is subject to both positive and negative allosteric regulation.



**Fig. 2:** Schematic of rule based effective control laws. Traditional enzyme kinetic expressions e.g., Michaelis-Menten or multiple saturation kinetics are multiplied by an enzyme activity control variable  $0 \le v_j \le 1$ . Control variables are functions of many possible regulatory factors encoded by arbitrary functions of the form  $0 \le f_j(\mathcal{Z}) \le 1$ . At each simulation time step, the  $v_j$  variables are calculated by evaluating integration rules such as the max or min of the set of factors  $f_1, \ldots$  influencing the activity of enzyme  $E_j$ .



**Fig. 3:** Reduced order coagulation model training simulations. Reduced order coagulation model parameters were estimated using particle swarm optimization (PSO) with and without the protein C pathway as a function of prothrombin. Solid lines denote the simulated mean value of the thrombin profile for N = 20 independent particles, points denote experimental data. The shaded region denotes the 99% confidence estimate of the mean simulated thrombin value (uncertainty in the model simulation). (A,B,C) training results for 150%, 100% and 50% of physiological prothrombin levels in the absence of the protein C pathway. Thrombin generation was initiated using 5 pmol/L FVIIa-TF in the presence of 200  $\mu$ mol/L of phospholipid vesicles (PCPS). All factors and control proteins were at their physiological concentration unless others denoted. The experimental training data was reproduced from the study of Butenas et al. [42].

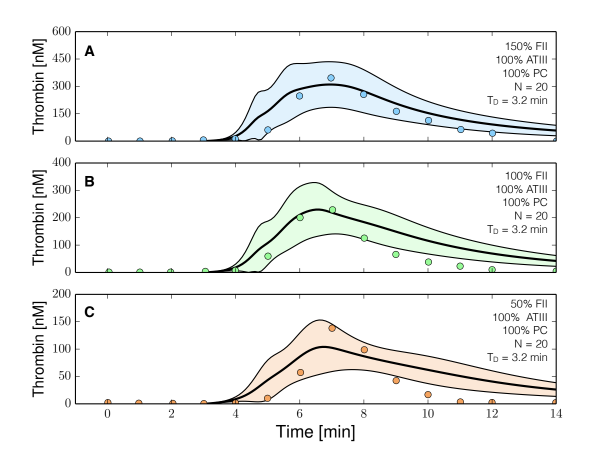
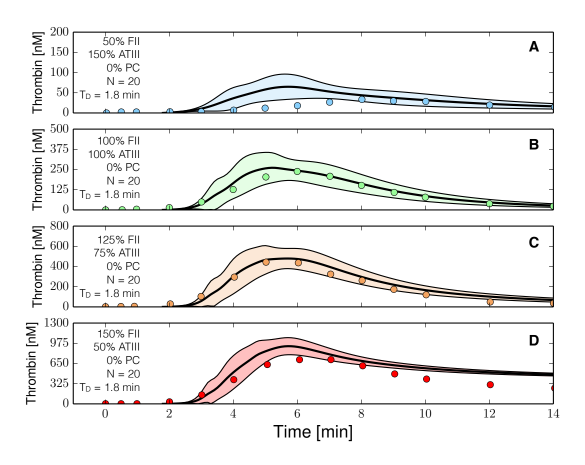
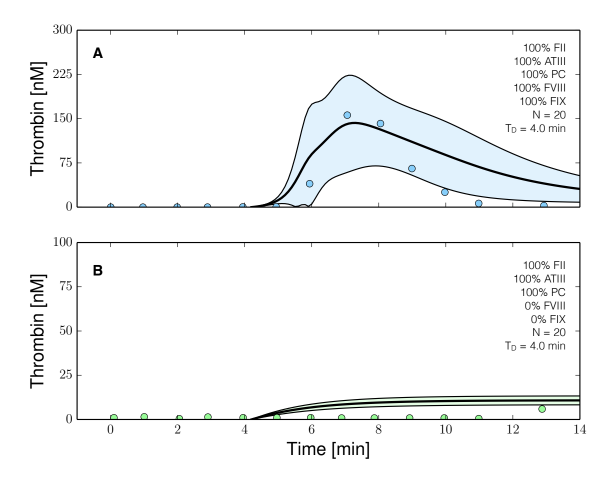


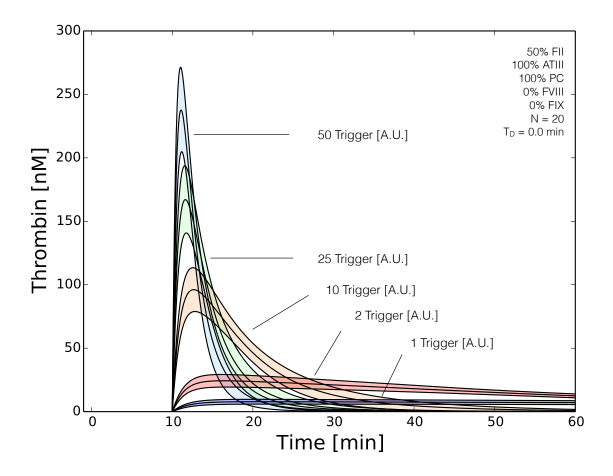
Fig. 4: Reduced order coagulation model training simulations. Reduced order coagulation model parameters were estimated using particle swarm optimization (PSO) with and without the protein C pathway as a function of prothrombin. Solid lines denote the simulated mean value of the thrombin profile for N = 20 independent particles, points denote experimental data. The shaded region denotes the 99% confidence estimate of the mean simulated thrombin value (uncertainty in the model simulation). (A,B,C) training results for 150%, 100% and 50% of physiological prothrombin levels in the presence of the protein C pathway. Only APC pathway parameters were allowed to vary in the simulations on the right. Thrombin generation was initiated using 5 pmol/L FVIIa-TF in the presence of 200  $\mu$ mol/L of phospholipid vesicles (PCPS). All factors and control proteins were at their physiological concentration unless others denoted. The experimental training data was reproduced from the study of Butenas et al. [42].



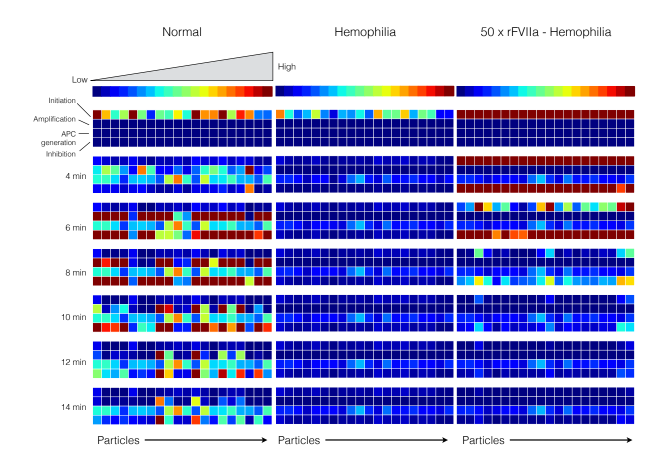
**Fig. 5:** Reduced order coagulation model predictions versus experimental data for normal coagulation. The reduced order coagulation model parameter estimates were tested against data not used during model training. Simulations of different levels of prothrombin and ATIII were compared with experimental data in the absence of the protein C pathway. Solid lines denote the simulated mean value of the thrombin profile for N = 20 independent particles, points denote experimental data. The shaded region denotes the 99% confidence estimate of the mean simulated thrombin value (uncertainty in the model simulation). (A,B,C,D) prediction results for (FII,ATIII): (50%,150%), (100%, 100%), (125%, 75%) and (150%, 50%) of physiological prothrombin and ATIII levels in the absence of the protein C pathway. Thrombin generation was initiated using 5 pmol/L FVIIa-TF in the presence of 200  $\mu$ mol/L of phospholipid vesicles (PCPS). All factors and control proteins were at their physiological concentration unless others denoted. The experimental validation data was reproduced from the study of Butenas et al. [42]



**Fig. 6:** Reduced order coagulation model predictions versus experimental data with and without FVIII and FIX. The reduced order coagulation model parameter estimates were tested against data not used during model training. Simulations of normal thrombin formation with ATIII and the protein C pathway were compared with thrombin formation in the absence of fVIII and fIX. Solid lines denote the simulated mean value of the thrombin profile for N = 20 independent particles, points denote experimental data. The shaded region denotes the 99% confidence estimate of the mean simulated thrombin value (uncertainty in the model simulation). (A,B) prediction results for normal thrombin generation and thrombin generation in hemophilia. All factors and control proteins were at their physiological concentration unless others noted. The experimental validation data was reproduced from the study of Allen et al. [17].



**Fig. 7:** Reduced order coagulation model predictions of rFVIIa administration. Simulations of thrombin formation in the presence of ATIII and the protein C pathway were conducted for a range of trigger values (1x - 50x) in the absence of fVIII and fIX. Solid lines denote the simulated mean value of the thrombin profile for N = 20 independent particles. The shaded region denotes the 99% confidence estimate of the mean simulated thrombin value (uncertainty in the model simulation). All factors and control proteins were at their physiological concentration unless others noted.



**Fig. 8:** Reaction flux distribution as a function of time for thrombin generation under normal (left), hemophilia (center) and rFVIIa treated hemophilia (right). Reaction flux was calculated for each particle at T=0,4,6,8,10,12,14 min after the initiation of coagulation. Blue colors denote low flux values red colors denote high flux values.