

Role of Fitted Reaction Rates in Predicting Thrombin Production

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Abstract

Background: The blood coagulation system is composed of a complex network of chemical reactions. This network can be modeled by a Hockin-Mann et al theoretical model.

Results: We have used a systematic approach to analyze how simulation fitted reaction rate constants in the Hockin-Mann model of the extrinsic (TF mediated) coagulation pathway affect rates of thrombin production. We first present algorithms based on numerical algebraic geometry to compute steady state solutions for different initial conditions. By applying these algorithms to general rank-deficiency systems, steady states which cannot be obtained by the time marching method are determined. Stability of these steady state solutions is studied using mathematical analysis and numerical simulations. We then perform both variance decomposition based on the sensitivity analysis and Morris design method to rank the significance of the 16 data-fitted reaction rates with respect to the total thrombin production. The sensitivity ranking of the reaction rates show agreement between both methods. Subsequently, the sparse grid probabilistic collocation method (SGPCM) is employed to quantify how uncertainties of these 16 reaction rates influence total thrombin production. Numerical results obtained by SGPCM are consistent with the results obtained from the classic Monte Carlo simulations. These methods can be applied to analysis of mathematical model with large scale parameters. Additionally, the numerical results indicate that SGPCM can achieve much faster convergence than classic Monte Carlo method with much lower

computational cost. Finally, we ranked the importance of different reaction rate constants using sensitivity analysis to identify critical threshold levels for factors VIII and IX for hemophilia therapy.

Conclusions: We present a general framework for analyzing reaction network represented by system of ordinary differential equations (ODEs). In these systems, there exists a large number of parameter sets. Especially, in Hockin-Mann model, among these 16 fitted reaction rate constants, the reaction rate k_{14} , which is related to factor IX, plays most important role in determining the rate of thrombin generation. Moreover, the sensitivity analysis provides threshold values for individual factors or provided reaction rate constants in controlling thrombin production. Identifying the threshold concentration of factor VIII provides clinical information to direct therapy for hemophilia patients experiencing bleeding disorder. In our simulation, we suggest that increasing the value of k_{14} and decreasing the concentration of factor X would provide a feasible therapy for patients of bleeding disorder.

Introduction

It is becoming increasingly vital to combine laboratory research with mathematical modeling and computer simulation to get additional insight into biological processes [1, 2, 11, 22, 36, 42, 44, 49, 55, 56, 58, 61, 64]. First, conceptual model is established by making some simplifying assumptions. Then, it is implemented mathematically and analyzed using analytical approaches or numerical methods. During the course of model development, experiments are used to estimate model parameters and to calibrate and validate the model. However, due to errors in experimental measurements, there always exists uncertainty in parameter estimation. This uncertainty includes model simplifications, computational errors coming from numerical schemes, and random local rules in Monte Carlo simulations. The situation could become even worse if certain parameter values may be impossible to measure using current experimental techniques, and to test a range of parameter values. Thus, identifying the sensitivity and uncertainty of the model parameters becomes essential for using models to make predictions and to explore biological hypotheses.

The Hockin-Mann's model [27] simulate blood coagulation reactions, one of the key processes controlling blood clot (or thrombus) formation. Blood clotting (or hemostasis) is critical to preventing hemorrhaging. Prior to the Hockin-Mann's model [27], Mann's group proposed in the form of system of ordinary differential equations [31,32] to describe the TF-initiated pathway reactions based on *in vitro* experiments. Despite wide applications of Hockin-Mann's model [31], structures, characteristics of steady state solutions of the model, sensitivity of the model, and uncertainty of parameter values have yet to be discussed. So far, only Luan *et al.* [42] applied sensitivity analysis to a network of coagulation reactions to identify fragile sites. Using

parameter values from the literature, reactions were identified where small changes in parameter values would have dramatic effects on thrombin generation and platelet activation. This analysis [42] identified reactions involving an interaction of factors IX and VIII as the most sensitive reactions to small fluctuations of relevant parameters.

In this paper, we introduce a new general framework for quantifying uncertainty, analyzing the parameters, and determining steady states of biological models (see Fig. 1), then demonstrate this approach by using a specific example (Hockin-Mann’s model). We first compute the steady state solution structures and characteristics of Hockin-Mann’s model [27] using numerical algebraic geometry method. We present a new numerical algebraic geometry algorithm to find all steady state solutions for arbitrary initial conditions. The steady state system of the model [27] is a singular sparse polynomial system. To deal with the singularity, we propose a new algorithm for a large number of variables. The steady state solutions are known to be dependent on the initial conditions and reaction rates. Exploring these solution structures and characteristics provides insights into understanding the inventory of the proteins and the associated biophysical and enzymatic processes involved in blood clotting. Moreover, analysis and control of threshold of factor VIII shows that the therapy of haemostasis controlling k_{14} and factor X can reduce the risk of bleeding disorder.

There are 16 reaction rates of the model [27] which were determined by fitting computational results with empirical data. Evaluation of uncertainties of these reaction rates has not been conducted yet. Moreover, sensitivity of these reaction rates has not been studied to the best of our knowledge. However, these parameters’ study would provide a feasible therapy (involving the most important reaction rates), and simplify the mathematical model. We next apply sensitivity analysis and uncertainty quantification to rank the importance of the reaction rates, and to quantify their interactions. Variance of these reaction rates on thrombin production and covariance of any pair of these reaction rates on thrombin production are decoupled and plotted in Figure 2. Among these fitted rate constants, we identified that the reaction rates (k_{14} and k_{15} , associated with factor IX, shown in Table 10) associated with factor IX play the most important roles in regulating the rate of thrombin production. Specifically at 20% variation in k_{14} and k_{15} can increase thrombin production by 200%.

Most of the therapeutic drugs treating hemophilia would vary one or two reaction rates. Using our sensitivity analysis results, we studied the effects of several hemophilia drugs (shown in Table 9) on the rate of thrombin production. Moreover, analysis of the threshold value of factor VIII (FVIII) could provide useful information to treat hemophilia, and reduce the risk of bleeding disorder: using normal levels of all coagulation network components (except for FVIII) the threshold FVIII concentration is 4.7% of normal

levels. We determined that the reaction rate k_{14} and the factor X (FX) concentration have dramatic effects on thrombin generation. By increasing the value of k_{14} and the concentration of FX, the threshold concentration of FVIII is reduced to 3.3 of normal. This technique could be valuable in identifying hemophilia carriers who are at risk of bleeding. FVIII levels in hemophilia carriers can vary widely from 10 - 90% of normal values. By determining the level of other factors and rate constants, one can identify whether an individual carrier has FVIII levels near their critical threshold and thus may be at risk of uncontrolled bleeding.

Biological Background

During the initial step of clot formation, resting platelets in blood are activated following adhesion to matrix proteins exposed from the damaged vessel wall. At the same time, blood borne factor VII binds with vessel wall tissue factor (TF) to form VIIa-TF complex (VIIa stands for activated factor VII) to initiate a complex set of blood coagulation reactions [46]. Coagulation reactions lead to the formation of thrombin from prothrombin. Thrombin is efficient in activating platelets and also converts fibrinogen in blood to fibrin which self-polymerizes, forming a fibrin network that is a major structural component of the clot [28, 35].

Platelets lead to pro-thrombotic changes upon activation. These include the release of their contents of alpha and dense granules that contain coagulation factors and platelet activators, modification of the platelet surface to promote surface dependent coagulation reactions, and the activation of the integrin receptor gpI-IaIIIb which can bind fibrin(ogen), von Willebrand Factor (vWF) and vitronectin, enabling these molecules to bind adjacent platelets and increase platelet to platelet adhesivity.

In parallel with platelet activation, VIIa-TF complex formed by factor VII and TF binding catalyzes the conversion of factor X to Xa directly or via a IXa-VIIIa intermediary step. Activated factor Xa combines with its protein cofactor, factor Va, on the surfaces of platelets to form prothrombinase (Va-Xa complex) which, in turn converts prothrombin (II) to thrombin (IIa) through an intermediate meizothrombin (mIIa). Transient mIIa is rapidly converted to IIa in the presence of factor Va and pro-coagulant lipids. Factor V is activated to its active form Va by both factor Xa and IIa. Thrombin activates platelets, converts fibrinogen to fibrin and activates factors V, VIII, and XI which provide a positive feedback loop propagating the production of thrombin. Thrombin is also a potent activator of resting platelets and thus mediates recruitment of resting platelets flowing nearby in the blood. The activated platelets provide a procoagulant surface that promotes coagulation enzyme activity. The formation and activity of VIIIa-IXa and Va-Xa complexes are dependent on the availability of phospholipid binding sites, which are only expressed on the

surfaces of activated platelets [1, 8, 27, 33–35].

The coagulation process is also regulated by inhibitors of coagulation reactions. TFPI binds VIIa-TF with factor Xa resulting in inhibition of the coagulation initiation reaction. ATIII inactivates several coagulation factors including thrombin, factor IXa and factor Xa. In addition, thrombin also initiates a negative feedback loop that possibly limits continued thrombin production by binding thrombomodulin on endothelial cells and activating Protein C. Activated Protein C inactivates factor Va and factor VIIIa which are required for continued thrombin generation. The activation of the PC anticoagulant pathway is initiated after thrombin generation begins, suggesting that it might provide a negative feedback mechanism to limit growth of a developing thrombus. Additionally, activated platelets release endothelial cell selective adhesion molecule (ESAM) which interferes with platelet to platelet adhesion. Presumably the release of anti thrombotic components at late stages of platelet activation might stop continued thrombus growth [12, 17, 27, 31, 32].

The Hockin-Mann’s model provided a good approximation of empirical data for blood clot formation. Subsequently, this model [31, 32] was improved by including blood anticoagulants tissue factor pathway inhibitor (TFPI) and antithrombin III (AT-III), and detailed descriptions of coagulation enzyme activities [27]. The new model (termed Hockin-Mann’s model thereafter) accurately predicted the nonlinear dependence of thrombin generation on the tissue factor and has been widely used to understand how various coagulation factors interact with each other and how variations of these factors affect thrombin production [8, 9, 12, 17, 29, 34, 66]. Lo *et al.* conducted kinetic Monte Carlo simulations [41] using the reaction network described in this model [27] to allow them to accurately simulate blood coagulation with low concentrations of blood zymogens and enzymes, which deterministic models often fail to predict. Specifically, rates of thrombin production were studied using low TF concentrations. Simulations revealed that 0.2pM TF was the critical concentration to cause 50% of reactions containing 3-fold diluted whole blood to generate 0.05 U/ml thrombin by 1 hour [41]. In addition to the other models [7, 18, 27, 30–34, 42, 66–68] have been proposed in recent years to integrate various aspects of coagulation reactions.

Results

Mathematical and computational framework for analyzing biological reaction networks

We present a general approach to compute the steady states and analyze importance of parameters, and provide diagram of different stages in the process in Figure 1 and Box 1.

Box 1 For a general reaction network model: how to study steady states and analyze the effects of parameters.

- 1 numerical algebraic geometry (NAG) introduces a novel paradigm of computing steady states. Especially, we present a new algorithm for the rank-deficient polynomial system which is derived from the reaction network model;
- 2 uncertainty quantification (UQ) and sensitivity analysis (SA) estimate and identify parameters in the mathematical model. In particular, we employ SGPCM, Morris method to analyze the effects of parameters;
- 3 Adjusting the most sensitive parameters, and comparing with the existence therapy, provides the threshold for these parameters for a given disease.

Through these numerical simulations, we are able to validate and calibrate the mathematical model, and provide some prediction for biological application. We use this approach to solve Hockin-Mann's model, and then provide an example to demonstrate general approach, and show outcome and benefits in specific biomedical application.

Analysis of steady state solutions

Using **Algorithm 2** to solve the steady state system of the blood coagulation model, we found 88 equilibrium (steady state solutions) for Hockin-Mann's model based on initial conditions from [27], and that all equilibriums are determined by linear and quadratic equations shown in Table 1. In particular, Thrombin IIa (x_{13}) and meizothrombin mIIa (x_{16}) are free variables, and linearly depend on the initial condition (linear equations), reaction rates, and values of other species such as TF=VIIa=X (x_{17}), TF=VIIa=Xa (x_{19}) as well as Xa=TFPI (x_{20}) (quadratic equations). Moreover, the constants shown in linear equations, such as 3.3383×10^{-9} and 2.33×10^{-10} , are determined by initial conditions. Quadratic equations account for effects of reaction rates k_i . Linear stability analysis is used to explore the steady state solution and study the matrix eigenvalue problems of the linearized system. Moreover, the nonlinear stability analysis is applied to a simplified steady state solution.

Linear stability

We consider the local stability of steady state solutions. Assuming that $x_i = x_i^s + \epsilon x_i^1 + O(\epsilon^2)$ for $i = 1, \dots, 34$, where x_i^s is the i -th component of steady state solutions we obtained using the method described in section **Methods**, and x_i^1 is the first order term by Taylor expansion at the steady states. By substituting x_i into Eq. (7) in Appendix B, and dropping $O(\epsilon^2)$ and higher order terms, the linearized system is obtained and

shown in Appendix D. The linearized system can be written as

$$\frac{d\mathbf{x}^1}{dt} = A(\mathbf{x}^s)\mathbf{x}^1,$$

which implies that $\mathbf{x}^1 = e^{A(\mathbf{x}^s)t}$. If all the eigenvalues of $A(\mathbf{x}^s)$ are non-positive, the steady state solution \mathbf{x}^s is stable. Since the matrix $A(\mathbf{x}^s)$ linearly depends on \mathbf{x}^s , we obtain the following formula

$$A(\mathbf{x}^s) = \sum_{i=1}^{34} A_i x_i^s,$$

where A_i is a real matrix and independent on x_i^s .

All the maximum eigenvalues of A_i are negative and shown in Table 2. Therefore all eigenvalues of $A(\mathbf{x}^s)$ are also negative for any \mathbf{x}^s with positive components. Thus, the linearized system corresponding to any steady state solution is stable. Then, we switch our attention to check the nonlinear stability of solutions shown in Table 1.

Nonlinear stability

Among these 88 steady states, we analyze one steady state, which is “close” to results obtained by solving Hockin-Mann’s model [27] using the time marching method. Specifically, this steady state solution is obtained by the following manner: combining with the time marching result from [27], the free variables in Table 1 are set to zero except the thrombin (x_{13}), meizothrombin (x_{16}) and TF=VIIa=X (x_{17}). Thus, a simplified steady state solution is obtained in Table 3. The difference between the solution computed by the time marching method and the steady state solution that we choose to analyze is smaller than 10^{-7} in L_1 norm. Relative differences of the nonzero components are shown in Table 4. By solving the polynomial system with respect to variables x_{13}, x_{16} and x_{17} with Bertini [4], we get one real positive solution as follows

$$x_{13} = 2.99990 \times 10^{-8}, x_{16} = 2.32829 \times 10^{-14}, x_{17} = 1.03445 \times 10^{-12}.$$

Then x_{19} and x_{20} can be obtained by substituting x_{16} and x_{17} into the linear equations in Table 3. This solution shows that total thrombin concentration is related with the complex TF=VIIa=X (x_{17}). In particular, equation $k_{18}x_{17} + k_{25}x_{17} - k_{19}x_{16}x_{13} = 0$ reveals the fact that TF=VIIa=X links with thrombin and meizothrombin through IXa=VIIIa in No. 11 and 15 chemical equations. Moreover, TF=VIIa=X also affects the amount of TF=VIIa=Xa (x_{19}) and Xa=TFPI (x_{20}) together with meizothrombin (linear equations).

Nonlinear stability analysis is accomplished by the following steps: applying random perturbations (a sequence ϵ_n in Table 5) to the steady state solution gives initial conditions for the initial value problem

shown in Appendix B; Runge-Kutta method is used to solve this initial value problem numerically. The results, summarized in Table 5, demonstrate that the steady state solution is non-asymptotically stable. In this case, the system moves away in a small region from the steady state solution after introducing small disturbances. Therefore, this chemical reaction network depends sensitively on the initial condition and the reaction rates.

Steady states for different initial conditions

Since initial conditions determine linear equations by generating an augmented system and play an important role in computing steady states, we consider different initial conditions from [34], which respond in a threshold manner to changes in the availability of particular surface binding sites for enzymes and zymogens. Using the **Algorithm 2**, we obtained steady state solutions and show them in Table 6. The free variables in Table 6 are set to zero, and then a specified steady state solution is obtained in Table 7. By solving the polynomial system in Table 7, we get two positive real solutions as follows:

$$x_{13} = 4.99656 \times 10^{-7}, x_{16} = 4.12563 \times 10^{-13}, x_{17} = 3.43569 \times 10^{-10}; \quad (1)$$

$$x_{13} = 4.46036 \times 10^{-7}, x_{16} = 7.25906 \times 10^{-11}, x_{17} = 5.39634 \times 10^{-8}. \quad (2)$$

In these two steady state solutions, (1) can be obtained by a time marching method based on the given initial conditions, while (2) can not be obtained by the time marching approach. However, (2) is also stable since it can come back to itself by giving small perturbation to (2) by time-marching. In many biological systems, there exists multiple equilibriums even for a given initial condition. Moreover, a small perturbation to an equilibrium can approach the equilibrium by time-marching indicates that the equilibrium is stable. Therefore, our method is competitive to find multiple steady state solutions.

Comparing Tables 3 and 7, polynomial equations have different coefficients. Therefore, these steady state solutions show a very sharp transition in predicting thrombin production as the initial conditions vary. Changes of these initial conditions induce an increase of thrombin concentration (x_{13}) by 10 times. Moreover, meizothrombin concentration (x_{16}) is also increased by 10 or even 1000 times. Our results are consistent with that of [34] and [41]. In fact, initial TF density is known to change the total thrombin significantly when a vessel is injured. A threshold dependence on TF concentration is found in the experiments by Veer and Mann [63]. Additionally, numerical study for low TF density was carried out in [41] by using Kinetic Monte Carlo simulation. However, the steady state solutions depends not only on initial condition, but also on reaction rates. It is obvious that the quadratic equation is related with rate constants k_i . Understanding

how variation of reaction rates affects steady or dynamic solutions is difficult to discern using linear or nonlinear stability analysis. We therefore perform sensitivity analysis and uncertainty quantification to the system and show their effects.

Sensitivity analysis and uncertainty quantification of reaction rates and initial conditions and their impacts on steady state solutions

In Hockin-Mann’s model, there are 16 reaction rates (shown in Table 8) that are derived through fitting computational results to empirical data. In order to investigate the uncertainties associated with these 16 reaction rates and depict their impacts on model predictions, the sparse grid probabilistic collocation method (SGPCM) [39, 40, 65] is utilized to generate the collocation points on the feasible parameter space of these 16 reaction rates, which are assumed to be random inputs with multivariate uniform distribution.

Sensitivity analysis of 16 data-fitted reaction rates

In this subsection, we apply two different stochastic sensitivity analysis methods, Morris method [45] based on Monte Carlo (MC) simulation and sparse grid collocation based on SGPCM approach, to rank the importance of the reaction rates on the total thrombin and study how the variation (uncertainty) in the total thrombin can be attributed to variations in the reaction rates as the inputs, i.e., systematically changing reaction rates in the model to determine the effects of such changes on the total thrombin concentration.

In applied statistics, the Morris method [45] for global sensitivity analysis is very efficient compared to other methods. We used the “Morris one at a time” (MOAT) module based on MC simulation performed by PSUADE [62], which is a software toolkit to facilitate the uncertainty analysis and design exploration. Figure 3 and Figure 4 show that the reaction rates k_{14} , k_{15} and k_{11} have more significant impact on the total thrombin synthesis compared with other fitted reaction rates. This result is consistent with the current understanding of coagulation cascade. k_{14} is the rate to form complex TF=VIIa=IX using TF=VIIa and IX. TF=VIIa=IX is subsequently disassociated to TF=VIIa and IXa with rate k_{15} . Moreover, k_{11} can control the production of TF=VIIa and Xa from TF=VIIa=Xa. The formation of the extrinsic factor Xa plays a key role in the total thrombin production, since prothrombinase catalyzes the conversion of zymogen prothrombin to thrombin.

Next, we used sparse grid probabilistic collocation methods [39, 40, 65], which can also result in sensitivity information, to rank sensitivities of these reaction rates and compare the results with that obtained by the Morris method [45]. Total variance of reaction rates on catalyzing thrombin was decomposed using analysis

of variance (ANOVA) technique as

$$v_{total} = \sum v_i + \sum v_{ij} + \sum v_{ijk} + \dots,$$

where v_i is the variance of reaction rate i , v_{ij} is the covariance of rates i and j , and v_{ijk} is the covariance of rates i , j and k , etc.

This variance decomposition indicates the amount of information each reaction rate (v_i) contributes to the total thrombin production and determines how much of the variance of each reaction rate can be explained by the other reaction rates (v_{ij}). We use a network graph to present ranks of the importance of each reaction rate with respect to the total thrombin and also illustrate the interactions between 16 data-fitted reaction rates. See Figure 2 in which the radius of disks corresponding to reaction rate i is proportional to v_i/v_{total} , representing the normalized variance associated with reaction rate i . The lines between any pair of reaction rates i and j is proportional to v_{ij}/v_{total} , representing the correlation between that pair. The thicker the line is, the larger the correlation is. Figure 2 also helps us to avoid the mistake of ignoring reactions which have small variances associated with themselves, but large correlation with other reactions. It is obvious that k_{14} and k_{15} play major roles in generating thrombin. We note that only two-way interactions are shown in Figure 2, but higher order interactions can also be displayed using the same approach. We also note that, when the output is switched, the sensitivity is also switched. We applied variance decomposition and plotted the sensitivity analysis in Figure 5 when output changed to IXa=VIIIa and IXa=ATIII. Figure 5 shows that k_4 and k_{14} play important roles in generating IXa=VIIIa (x_{33}) and IXa=ATIII (x_{30}) respectively.

We conclude that the results obtained by the Morris design based on the MC simulations are consistent with the results based on SGPCM (Figure 2). Both of them show that the variances of k_{14} , k_{15} and k_{11} are larger, and that they are dominant reaction rates in generating the total thrombin. We remark that Morris method is a classical method on sensitivity analysis, it depends on MC method which has a lower convergence than SGPCM. In addition, the sparse grid collocation can also provide the effects between two reaction rates.

Sensitivity analysis of initial conditions

From pervious study, we showed that initial concentrations of factors also contribute to steady states of the blood coagulation model. Here we performed sensitivity analysis of 10 non-zero initial concentrations of zymogen factors using SGPCM. These non-zero initial concentrations of zymogen factors are assumed to vary in 80%-120% of their normal values, and follow a uniformly distribution. The effects of initial

conditions on the total thrombin synthesis is shown in Figure 6. Factor X (x_{15}) plays an important role in generating total thrombin and its cofactor Xa interacts with TFPI (x_{18}), which inhibits the thrombin production. Blood coagulation simulation is dependent on various model components such as reaction rates, and initial concentrations. Sensitivity analysis of these reaction components can give a guidance to experimental variability in increasing and decreasing thrombin production.

Validation of uncertainty quantification method

The SGPCM approach can achieve much faster convergence than a classic MC method and it handles a larger number of uncertain parameters with relatively lower computational cost, when comparing sparse grids with full tensor products [39]. Here we utilized the MC simulations to verify accuracy of the SGPCM approach, which we employed to do uncertainty quantification. In Figure 7, we plotted the difference of the mean and variance of the total thrombin obtained by SGPCM (545 points) and MC (1000 points). The results (mean and variance of the total thrombin) obtained by the SGPCM approach is consistent with that of MC simulations. But the SGPCM is more efficient, and achieves faster convergence than MC. We used three different levels of sparse grid points (184 points, 1228 points and 6430 points) associated with 16 data-fitted reaction rates and treated the mean and variance of 6430 points as the “real solution”. The absolute errors of the mean and variance for 34 variables are shown in Table 12 (Appendix E) which verifies the convergence of the SGPCM approach.

Propagation of uncertainty of reaction rates

Propagation of uncertainty of reaction rates in model solutions is studied in this section by using initial conditions described in [27]. The 16 data-fitted reaction rates are input variables and steady state concentrations of different factors are output variables. Runge-Kutta method is used to compute the time marching system on the sampling space of input variables. Figure 8 shows the mean and error bars of steady state solutions whose magnitudes are greater than 10^{-10} μM . Error bars show how the data are spread and encompass the lowest and highest values by using formula $mean \pm standard\ deviation (SD)$, where SD is the average or typical difference between the data points and their mean. It is obvious that the steady state solutions we found are lying inside the intervals of Figure 8. The results of the mean and error bar for time marching simulations of undiluted whole blood triggered with 25 pM TF has been shown in Figure 9. Since it shows that thrombin production and consumption reach equilibrium by 700 seconds in Hockin-Mann’s model, we studied the simulation over 700 seconds in time. Total thrombin is defined as thrombin (IIa)

and meizothrombin (mIIa) with meizothrombin multiplied by a factor 1.2 because of meizothrombin's 20% greater activity toward substrates [27]. Figure 9 plots the mean and error bar of the total thrombin with total 545 sparse grid points for 16 data-fitted reaction rates. This prediction confidence interval of total thrombin production is provided with the model as the quantification of the uncertainty.

Prediction of hemophilia therapy targeting factors VIII & IX

The sensitivity analysis provides threshold values for individual factors or reaction rate constants in controlling thrombin production. One application of our study is to examine women who are carriers for hemophilia. Hemophilia is a group of hereditary genetic disorders that impair the body's ability to control blood clotting or coagulation, which is used to stop bleeding when a blood vessel is broken. There are two types of hemophilia diseases: one is factor VIII deficiency; and the other one is factor IX deficiency. These carriers contain a normal and defective gene for factor VIII or factor IX, and can present with a wide range of factor VIII or factor IX concentration levels in blood; as wide as 0-20% of their normal values respectively. By knowing the threshold concentration of factor VIII or factor IX given the concentration level of other factors, one can predict the level of thrombin production in case of bleeding, and subsequently determine if a carrier is at risk of bleeding disorder using our approach.

In this section, we show how factors VIII and IX quantitatively control the total thrombin concentration to identify individuals at risk of bleeding disorder for hemophiliac patients. We assume that the amounts of factors VIII and IX vary in ranges of 0-20% of their normal values. We also assume that the threshold of the total thrombin is 10% of its normal value. Below this level, people would have bleeding disorder. Other factors are assumed to be at their normal levels of concentration. We use uniformly distributed sampling points on factors VIII and IX, each factor having 100 points. The total thrombin is computed at each pair of sample points of factors VIII and IX by solving the time dependent system (see Appendix B). The total thrombin is compared to the assumed threshold value of total thrombin. Figure 10 shows that the threshold concentration of factor VIII is 4.7 % of its normal value, and that the threshold is increased when the concentration of factor IX increases. The histogram in Figure 11 shows the probability of the total thrombin above the threshold is 75.4% when factor VIII and factor IX vary. In this case, the risk of bleeding disorder is high (24.6%). The threshold of factor VIII is 4.7% of the normal value.

Furthermore, this analysis also identifies other factors that might lower the threshold concentration of factor VIII so that above threshold level of thrombin is produced to prevent from bleeding disorder. Based on the sensitivity analysis, we noticed that the threshold concentration of factor VIII would be lowered by

increasing concentrations of the pro-coagulant factors that lead to produce total thrombin. Figures 2 and 6 show that the total thrombin concentration is activated by raising the value of reaction rate k_{14} and the initial concentration of factor X (x_{15}). Figure 12 shows that the total thrombin production is increased from 0.145 to 0.17 as k_{14} increases by 10% of the normal value. Increasing the value of reaction rate k_{14} is to activate factor IX to its active form IXa. This study is consistent with the study of the local measures [26], which are used with a pre-operative dose of raising the concentration level of factor IXa. In [26], factor IXa is increased 10% by using local measures of tranexamic acid solution, and excellent haemostasis is achieved for severe hemophiliac patients. In order to lower the threshold value of factor VIII in Figure 10, we let the rate of k_{14} and the initial concentration of factor X (x_{15}) vary in the range of 100-200% of their normal values. We again take 100 uniformly distributed sampling points in each of their parameter value spaces. The total thrombin is computed on the factor VIII and IX sampling space by time marching method. Figure 13 shows that the threshold of factor VIII is decreased to 3.3% by raising the value of reaction rate k_{14} and the initial concentration of factor X. Figure 14 shows the probability of bleeding disorder is also decreased to 17.7%. Thus it is possible by altering the activity the level of reaction rate k_{14} , and the initial concentration of factor X, one could reduce the level required of factor VIII to maintain normal hemostasis. Moreover, Factor IX is an important drug of haemostasis (see Table 9 for a sampling of current trials involving factor IX). Most trials involve the important factor IX. The lack of factor IX can produce bleeding episodes that cause damage of the bone, muscles, joints, and tissues. Control mechanisms involving factor IXa are common strategies in current anticoagulation conical therapies and clinical trials [42].

Discussion

This paper presents a general mathematical and computational framework to identify effective drug therapy and predict its performance with confidence interval (Figure 1). First, an algorithm is developed to solve the steady state system with rank-deficiency derived from a mathematical model and establish a quantitative understanding of effects of simulation fitted reaction rates and initial conditions on the total thrombin. Second, we have developed a new technique to quickly solve the steady state system, as well as analyze the chemical reaction model using stochastic approaches. Finally, effective drug therapy is identified by comparing the importance ranking of the corresponding reaction rates obtained through sensitivity analysis study. We employ the numerical model to predict the performance of the most effective drug therapy with confidence interval of corresponding thrombin production using SGPCM. Further analysis will be required to assess the effects of all parameters, such as reaction rates, using compressive sensing and optimization

under uncertainty. Optimal control on the parameters for manipulating the total thrombin will be derived in our future works.

We have analyzed Hockin-Mann’s model that describes the tissue factor mediated pathway of blood coagulation. A new algorithm based on numerical algebraic geometry is proposed to compute the steady state system with rank-deficiency and is applied to the Hockin-Mann’s model. This algorithm is the first one we are aware of that utilizes methods of numerical algebraic geometry to solve rank deficient systems. It can be used to analyze other dynamic systems with rank-deficiency. Moreover, our results show that the steady states depend not only on initial conditions but also on reaction rates. This algorithm enables us to find all solutions, among which the time marching method would not be able to solve the dynamical system with multiple equilibriums for given initial condition.

We used methods of sensitivity analysis and uncertainty quantification to explore the effects of 16 reaction rates whose values are obtained by fitting data and initial conditions. Both variance decomposition based sensitivity analysis and Morris design method were performed to rank the significance of the 16 uncertain reaction rates with respect to the total thrombin. Ranks of the importance of each reaction rate with respect to total thrombin concentration and the interactions between pair reactions are illustrated using a network graph (See Figure 5). Good agreement on the sensitivity ranking of the reaction rates is observed using both methods. SGPCM is employed to study the effect of the 16 uncertain reaction rates on the total thrombin concentration. The numerical results are consistent with the results obtained from classic Monte Carlo simulations. Additionally, the numerical results indicate that SGPCM can achieve much faster convergence than classic Monte Carlo method with much lower computational cost. (Figure 7)

Computational studies support the following three main conclusions:

- the steady states depend on the initial concentrations of factors and chemical reaction rates;
- our methods compute all steady states, which couldn’t be founded by time marching method;
- among 16 fitted rate constants, the reaction rate k_{14} plays an important role in generating the total thrombin;
- the threshold concentration of factor VIII is controlled by k_{14} and factor X. This provides clinical guidance for the examination of bleeding disorder.

All “important” reaction rates, factors are listed in Box 2.

Box 2 Important reaction rates and factors

- k_{14} and k_{15} : are the most important reaction rates in generating IXa=VIIIa and IXa=ATIII respectively, eventually play important roles in generating the total thrombin;
- **factors VIII and IX**: deficiency causes two types of hemophilia diseases;
- k_{14} and **factor X**: control the threshold of factor VIII, and reduce the risk of bleeding disorder.

Although previous studies showed that experimental coagulation initiation at different levels of TF effects the total thrombin production. Our analysis investigates how altering both the initial concentrations of pro-coagulant factors and values of reaction rate constants can effect thrombin production, and shows that k_{14} and k_{15} play a major role in generating the total thrombin since they directly control generation of the complex TF=VIIa=IX and thereafter control the generation of factor IXa. On the other hand, rates k_{37} and k_9 , which consume TF=VIIa complex and factor X, prevent the total thrombin production. Moreover, k_4 activates, and k_{37} inhibits the role of k_{14} . In fact, larger values of k_4 yield more generation of TF=VIIa, which speeds up the thrombin production, and then activates the role of k_{14} . While k_{37} inhibits the thrombin production by consuming TF=VIIa to generate the complex TF=VIIa=Xa=TFPI, and then gives negative feedback to k_{14} . Furthermore, blocking k_{14} and k_{15} results in significant sensitivity shifts. Figure 15 shows that their affinities are predicted to be less important when values of k_{14} and k_{15} are reduced to 10% of their normal values. Conversely, the sensitivity of k_{23} related with factor VIIIa is found to increase. Our analysis also shows that the factor VIII plays an important role in the examination of bleeding disorder for women who are hemophilia carriers, and can vary within a wide range, namely, 20-80% of the normal value. Analysis of the threshold concentration of factor VIII can be used to predict if a carrier is at high risk of bleeding disorder. The threshold level for this particular factor in a patient would depend upon the activity rate and the concentration of all coagulation factors. The current common practice for hemophilia treatment is to replace the missing factor with exogenous material. If it is a human derived factor VIII, this carries the risk of spreading disease (as in the case of HIV and the high number of hemophiliacs who developed AIDS). One could provide recombinant proteins, but these are very expensive. Also recombinant proteins can induce the patient to generate an immune response to the foreign proteins and block their effectiveness. Thus, identifying the threshold value of factor VIII enables the use of small molecule inhibitors which are less likely to be immunogenic and should be cheaper than production of recombinant proteins. The sensitivity analysis could provide important clinical information to direct therapy for bleeding and thrombotic disorders.

Furthermore, the sensitivity analysis can identify new targets for hemorrhagic disorders. One approach

to treat coagulation factor deficiencies is to replace the missing factor with exogenous material. Prophylactic use of exogenous material to prevent disease is very expensive and exogenous material is often just provided during the bleeding episodes but not for prophylactic purpose. Furthermore, the exogenous material may induce an immune response antagonizing the effect of replacement therapy. Sensitivity analysis may help identify other elements of the coagulation network where changes in activities may reduce the threshold concentration of the missing factors. Thus small molecule inhibitors or activators of the other elements (which are less likely to induce an immune response) can reduce the effects of the factor deficiency. This potentially may provide new therapeutic targets to treat coagulation deficiencies, and it also may be possible to individualize the treatment for a specific patients based upon their level of coagulation components based on the guidance of the sensitivity analysis.

Methods

Overview of methods

Over the last decade, numerical algebraic geometry (see [37] and [57] for a review), which grew out of continuation methods for finding all isolated solutions of systems of nonlinear multivariate polynomials, became efficient and was applied to variety of problems. To approximate all isolated solutions of polynomial systems, numerical path following techniques have been proven reliable and efficient during the past two decades. In the 90's, homotopy methods were developed to exploit special structures of the polynomial system, in particular its sparsity. In numerical algebraic geometry, we apply and integrate homotopy continuation methods to describe solution components of polynomial systems. Moreover, we have been successful in the past in dealing with these types of problems, zebrafish model [23], necrotic core tumor growth model [24] and three dimensional tumor growth model [25]. In [24], we found multiple steady state solutions of the zebrafish model, which is hard to accomplish using classical numerical PDE algorithms. We applied homotopy continuation to free boundary problems discussed in [24,25], and showed the efficiency of our algorithms. These studies provide the benefits of numerical algebraic geometry in studying nonlinear systems. All the computations are based on Bertini [4], which is a software package in the field of numerical algebraic geometry that numerically compute all solutions of polynomial systems over \mathbb{C}^n . However, Bertini cannot be directly applied to a “deficient steady state system”, which is defined as if some equations of a system can be expressed by a linear combination of other equations in this system. In other words, the Jacobian associated with the system is rank deficient at any given point. If the deficient system is also the steady state system derived from a differential equations system, then we call it as a “deficient steady state system”.

There is rich literature on methods to study biochemical reaction networks, such as reaction network structures [16], Michaelis-Menten kinetics, linear kinetics and the rational parameterizations [60]. Specifically, [42] studies the fragility and robustness of the model and discusses the computationally derived points, correlation between the model simulation and experimental data, and computation of the Overall State Sensitivity Coefficients (OSSCs); [16] discusses the relationship between reaction network structure and the possibility of multiple equilibria based on computer algebra software; [60] uses a parameterization and transfers steady state system into a L algebraic equation. One common difficulty encountered by these methods is to deal with rank-deficient systems. To overcome this difficulty, we propose a new algorithm, which can compute the steady states for rank-deficient systems and analyze stability of the solutions. Our results cannot be derived from any existing mathematical theories of biochemical reaction networks. The steady state solutions depend upon the initial condition and reaction rates. In Hockin-Mann’s coagulation model, 16 reaction rate constants are obtained by data-fitting. Uncertainties associated with the TF pathway of blood coagulation due to uncertainty associated with these reaction rate constants remain unknown. We explore the significance of these data-fitted reaction rates by using uncertainty quantification and sensitivity analysis.

Sensitivity analysis is a powerful tool in ranking the importance of the random inputs and quantifying their interactions. The sensitivity approach in [50] uses a few sample points in a large parameter domain to rank the random inputs, which is a computationally efficient technique for a linear model. However, when the system has strong nonlinearities, more sample points covering the whole parameter domain have to be used. In sensitivity analysis, a common approach is changing One-At-a-Time (OAT), to see what effect this produces on the output. Sensitivity methods that use OAT design are sensitive approaches where the impact of changing the values of each factor is evaluated in turn. Randomly moved OAT as inputs is the most used screening method in sensitivity analysis. Many screening methods have been developed, e.g., Morris’s method [10, 45, 53], Cotter’s method [15], Andres’ iterated fractional factorial method [3], Bettonvil’s sequential bifurcation [6], and variance-based sensitivity analysis [53]. By using a sensitivity analysis framework to explore in the space of reaction rate parameters, we study the output statistics to understand how input uncertainties propagate through the Hockin-Mann’s coagulation model and how Mann’s model responds to variation in values of these simulation fitted reaction rates. The Morris method is utilized to show effects of reaction rates on generating the total thrombin. The total variance of total thrombin is decomposed by using Analysis Of Variance technique (ANOVA). We remark that the sensitivity analysis performed in the present paper can be extended to inverse modeling to calibrate model parameters

to improve simulations. The relationships between the output responses and input parameters (response curves or surfaces) can also be used to develop reduced-order models for the subset of processes, which can then be used to perform more extensive uncertainty quantification of the complete model. Recently the probabilistic collocation method (PCM) has been applied for sensitivity analysis and can efficiently identify the sensitive parameters [51, 52]. In [38], the Multi-Element-PCM based sensitivity analysis method is developed, which is efficient and can achieve fast convergence compared to standard Monte Carlo (MC) based sensitivity analysis methods for cases with 20-100 random parameters. Moreover, PCM can be also used to speedup the convergence for uncertainty quantification introduced by Tatang and McRae [59]. Recently Xiu and Hesthaven [65] have used Lagrange polynomial interpolation to construct high-order stochastic collocation methods. The properties of PCM were extensively studied in the past 10 years. In [5, 47, 48], the errors of integrating or interpolating functions with Sobolev regularity were analyzed for Smolyak constructions based on one-dimensional nested Clenshaw-Curtis rules. In [48], the degree of exactness of the Smolyak quadrature using Clenshaw-Curtis and Gaussian one-dimensional rules was investigated. In [65], the efficiency of Clenshaw-Curtis-based sparse grid stochastic collocation was demonstrated in comparison to other stochastic methods on an elliptic problem. In 2003, Gerstner and Griebel [20] introduced the dimension-adaptive tensor product quadrature method. In [19], sparse grid collocation schemes were applied to solving stochastic natural convection problems. In [21, 43], an adaptive hierarchical sparse grid collocation algorithm has been developed. In this study, we apply the Sparse Grid Probabilistic Collocation Method (SGPCM) to study the uncertainty quantification process of Mann’s model in a faster convergence rate. We validate our method by comparing results obtained by SGPCM and MC.

Sparse grid probabilistic collocation method

In this paper, we applied the uncertainty quantification method - SGPCM to achieve faster convergence in the uncertainty quantification process, compared with the classic MC simulations. The general procedure of the PCM approach is similar to that of the MC method except that different sampling points and corresponding weights are selected. The procedure of the PCM consists of the following of three main steps:

1. Generate N_c collocation points in probability space of random parameters as independent random inputs based on a quadrature formula;
2. Solve a deterministic problem at each collocation point;

3. Estimate the solution statistics using the corresponding quadrature rule,

$$\langle u(\mathbf{x}, t) \rangle = \int_{\Gamma} u(\mathbf{x}, t, \xi) \rho(\xi) d\xi \approx \sum_{k=1}^{N_c} u(\mathbf{x}, t, \xi_k) w_k, \quad (3)$$

$$\sigma(u)(\mathbf{x}, t) = \sqrt{\int_{\Gamma} (u(\mathbf{x}, t, \xi) - \langle u \rangle)^2 \rho(\xi) d\xi} \approx \sqrt{\sum_{k=1}^{N_c} u^2(\mathbf{x}, t, \xi_k) w_k - \langle u \rangle^2}, \quad (4)$$

where $\rho(\xi)$ is the probability density function (PDF) of the random variable ξ , N_c is the number of quadrature points, $\{\xi_k\}$ is the set of quadrature points, and $\{w_k\}$ is the corresponding set of weights, which is the combination of quadrature weights in each random dimension. In the second step of the PCM, as for the MC method, any existing code can be used. Extensive reviews on the construction of quadrature formulas may be found in [13, 14]. The Smolyak formula [54] is used to construct the collocation point set, which is a linear combination of tensor product formulas and the resulting point set has a significantly smaller number of points than the full tensor product set. Recently, Lagrange polynomial interpolation has been used [39, 40, 65] to construct high order stochastic collocation methods based on sparse grids using the Smolyak formula [54]. Such sparse grids do not depend as strong on the dimensionality of the random space as those more suitable for applications with high-dimensional random inputs. Detailed descriptions on building the collocation point set can be found in [39, 40, 65].

Global sensitivity analysis based on Polynomial Chaos Expansion (PCE)

Global sensitivity analysis can be obtained from PCEs in order to identify the dominant sources of uncertain parameters and their attributions. Generally, PCEs of a second-order stochastic processes with d number of independent random variables can be expressed as

$$g(x_1, \dots, x_d) = \sum_{k=0}^{K-1} c_k \Psi_k(\mathbf{x}),$$

where $\mathbf{x} = (x_1, \dots, x_d)$ is a set of independent random variables, $\Psi_k(\mathbf{x})$ are multidimensional orthogonal polynomials with regard to the inner product, and c_k are the deterministic polynomial chaos coefficients of g . Then the global sensitivity analysis based on variance decomposition is employed. Here the total variance is defined as

$$Var[g(\mathbf{x})] = \sum_{k>0} c_k^2 \|\Psi_k(\mathbf{x})\|^2,$$

where $\|\Psi_k(\mathbf{x})\|$ can be pre-computed, and c_k is calculated using the sparse grid probabilistic collocation method and the orthogonal property of polynomial chaos. Then the first-order (or main) effect sensitivity

indices of S_i are

$$S_i = \frac{\text{Var}[\mathbb{E}(g(\mathbf{x}|x_i))]}{\text{Var}[g(\mathbf{x})]} = \frac{\sum_{k \in \mathbb{I}_i} c_k^2 \|\Psi_k(\mathbf{x})\|^2}{\sum_{k > 0} c_k^2 \|\Psi_k(\mathbf{x})\|^2},$$

where \mathbb{I}_i is the set of bases with only x_i involved. S_i is the uncertainty contribution that is due to i -th parameter only. Similarly, the joint sensitivity indices can be written as

$$S_{ij} = \frac{\text{Var}[\mathbb{E}(g(\mathbf{x}|x_i, x_j))]}{\text{Var}[g(\mathbf{x})]} - S_i - S_j = \frac{\sum_{k \in \mathbb{I}_{ij}} c_k^2 \|\Psi_k(\mathbf{x})\|^2}{\sum_{k > 0} c_k^2 \|\Psi_k(\mathbf{x})\|^2},$$

where \mathbb{I}_{ij} is the set of bases with only x_i and x_j involved. S_{ij} is the uncertainty contribution that is due to (i, j) parameter pair.

Method for solving rank-deficient polynomial system

We develop a numerical algorithm for studying the steady state system derived from the tissue factor-mediated coagulation pathway model introduced in [27] in this section. The coagulation cascade formulated in [27] is listed in Table 10 of Appendix A for the self-completeness of the paper. From this coagulation cascade, we get a mathematical model consisting of 34 ordinary differential equations (see Appendix B) with 42 rate constants which describe 27 independent equilibrium expressions listed in Table 10. Table 11 of Appendix A lists the notations to represent species in Table 10 used by ODEs in Appendix B. The corresponding steady state system, which is obtained by setting t -derivatives to be 0 for the ODE system, is a polynomial system with 34 variables. Unfortunately, this system cannot be solved directly by polynomial solvers such as Bertini [4] because of its rank-deficiency. In order to deal with the rank-deficiency, let us start with considering the rank of the polynomial system. From [57], the rank of a polynomial system equals to that of the Jacobian associated with the system at any random generic point. For our model, the rank of the steady state system is 24, which suggests that the system is not of full rank. An additional 10 linearly independent equations need to be added to the system in order to generate an augmented polynomial system. Here we introduce an idea of symbolic computation to determine these additional linear equations. Let $C(\mathbf{x}, \mathbf{k})$ denote monomial components of the steady state polynomial system, i.e., the multiplication of the rate constants and reactants,

$$C(\mathbf{x}, \mathbf{k}) = \left(k_2 x_1 x_2, k_1 x_3, k_4 x_1 x_4, k_3 x_5, k_5 x_5 x_2, k_6 x_6 x_2, k_7 x_7 x_2, k_9 x_5 x_8, k_8 x_9, k_{12} x_5 x_6, k_{11} x_{10}, k_{14} x_5 x_{11}, \right. \\ k_{13} x_{12}, k_{16} x_6 x_{14}, k_{17} x_7 x_{15}, k_{19} x_{16} x_{13}, k_{18} x_{17}, k_{21} x_{17} x_8, k_{20} x_{18}, k_{24} x_{16}, k_{23} x_{19} x_{20}, k_{25} x_{18}, k_{25} x_{17}, \\ k_{26} x_7 x_{21}, k_{28} x_6 x_{22}, k_{27} x_{23}, k_{30} x_{23} x_{14}, k_{29} x_{24}, k_{32} x_{25} x_{23}, k_{34} x_6 x_{26}, k_{33} x_{27}, k_{36} x_{10} x_{26}, k_{35} x_{28}, \\ \left. k_{37} x_5 x_{27}, k_{38} x_6 x_{29}, k_{39} x_{25} x_{29}, k_{40} x_{13} x_{29}, k_{41} x_7 x_{29}, k_{42} x_5 x_{29}, k_{10} x_9, k_{15} x_{12}, k_{22} x_{18}, k_{31} x_{24} \right)^T,$$

then the steady state polynomial system $f(\mathbf{x})$ can be expressed as

$$f(\mathbf{x}) = A \times C(\mathbf{x}, \mathbf{k}),$$

where A is a real valued matrix. Obviously, the rank of A is the same as the rank of the system $f(\mathbf{x})$. Since $\text{rank}(A) = 24$, there exists 24 linearly independent rows of A so that the other 10 rows of A can be represented by them (see Appendix C). Moreover, each row of A , say i -th row $A(i, :)$, is corresponding to i -th equation of $f(\mathbf{x})$ or $\frac{dx_i}{dt}$. For example, we have

$$A(24, :) = -A(21, :) - A(22, :) - A(23, :),$$

which means

$$\begin{aligned} \frac{dx_{24}}{dt} &= -\frac{dx_{21}}{dt} - \frac{dx_{22}}{dt} - \frac{dx_{23}}{dt} \\ \Rightarrow x_{24} + x_{21} + x_{22} + x_{23} &= \text{Const}, \end{aligned} \quad (5)$$

where Const is a constant determined by the initial condition. In this case, we can generate additional linear equations by the following algorithm.

Algorithm 1: Algorithm for generating linear equations

Input : The coefficient matrix $A_{n \times m}$.

Output: Linear equations generated.

for $i = 1 : n$ **do**

 determine if the row $A(i, :)$ is linearly independent upon B ?

if true then

 | add $A(i, :)$ into B and add i into \mathcal{I}_B .

else

 | write $A(i, :)$ as a linear combination of the rows of B and insert the coefficients of B into \mathcal{C} .

 | add i into \mathcal{I}_C .

end

end

for $i = 1 : \text{size of } \mathcal{I}_C$ **do**

 | print $\frac{dx_{\mathcal{I}_C i}}{dt} = \sum_j \mathcal{C}(i, j) \frac{dx_{\mathcal{I}_B j}}{dt}$

end

For Mann's model, 10 linear equations can be generated and denoted as $L(x_1, x_2, \dots, x_{34})$. Therefore, the augmented polynomial system becomes

$$F(x_1, x_2, \dots, x_{34}) = \begin{pmatrix} f(x_1, x_2, \dots, x_{34}) \\ L(x_1, x_2, \dots, x_{34}) \end{pmatrix}. \quad (6)$$

Now we switch our attention to the computation of steady state solutions to system (6). Noticing that there are some simple polynomial equations such as f_{11}, f_{18}, f_{24} . We propose the following algorithm to simplify and to solve this rank-deficient polynomial system (RDPS).

Algorithm 2: Algorithm for solving RDPS(f)

Input : A set $f = \{f_1, \dots, f_N\}$ of N polynomials, the linear system L .
Output: A set $g = \{f_1, \dots, f_M\}$ of M simplified polynomials. A zero solution set $S = \{x_i | x_i = 0\}$.
Set $S = \{\emptyset\}$.
Call RDPS_iterative(f, L, S).

Algorithm 3: RDPS_iterative(f, L, S)

Input : A polynomial set $f = \{f_1, \dots, f_{N'}\}$, linear system L and solution set S
Output: true (success) or false
Set display_solution=true
if S does not satisfy L **then**
| **return** false
end
for $i = 1 : N'$ **do**
| **if** f_i is single monomial **then**
| | display_solution=false
| | remove f_i from f , assume $f_i = k_{s_i} x_{i_1} \cdots x_{i_m}$
| | **for** $j = 1 : m$ **do**
| | | add x_{i_j} into S , evaluate f as f' and L as L'
| | | **if** RDPS_iterative(f', L', S) **then**
| | | | **return** true
| | | **else**
| | | | remove x_{i_j} from S
| | | **end**
| | **end**
| **end**
end
if display_solution **then**
| output f and S
| **return** true
end
return false

Author's contributions

WH, AS and ZX developed algorithms for steady states; WH, GL and ZX performed sensitivity and uncertainty to the model; WH, GL, ZX and ER analyze the therapy for bleeding disorder; AS and MA supervised the project. All authors read and approved the final manuscript.

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Tables

Table 1: Steady state solutions of Hockin-Mann's model [27] with initial conditions specified in [27]. See Table 11 in Appendix A for definition of variables.

Constant solutions	$x_1 = x_3 = x_5 = x_6 = x_7 = x_8 = x_9 = x_{10} = x_{12} = x_{18} = x_{23} = x_{24} = 0,$ $x_{25} = x_{27} = x_{28} = x_{29} = 0, x_{26} = 8.33 \times 10^{-10}, x_{30} = 5.33 \times 10^{-9}, x_{34} = 2.5 \times 10^{-11}$
Free variables	$x_4, x_{11}, x_{13}, x_{14}, x_{15}, x_{16}, x_{17}, x_{21}, x_{33}$
Linear expression	$x_2 = 3.3383 \times 10^{-9} - x_4, \quad x_{19} = x_{20} = 2.33 \times 10^{-10} - x_{15} - x_{16} - x_{17}$ $x_{22} = 6.67 \times 10^{-9} - x_{21}, \quad x_{31} = 4.67 \times 10^{-7} - x_{14} - x_{33}$ $x_{32} = 3 \times 10^{-8} - x_{11} - x_{13} - x_{17}$
Other equations	$k_{18}x_{17} + k_{25}x_{17} - k_{19}x_{16}x_{13} = 0, k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20} - k_{19}x_{16}x_{13} = 0$

Table 2: Max eigenvalues for A_i

$\max(\rho(A_1))$	-4.3e-19	$\max(\rho(A_7))$	-7.1e-06	$\max(\rho(A_{13}))$	-4.9e-07
$\max(\rho(A_{18}))$	-1.1e-04	$\max(\rho(A_{24}))$	-1.1e-04	$\max(\rho(A_{30}))$	-1.1e-04
$\max(\rho(A_2))$	-1.1e-04	$\max(\rho(A_8))$	-1.1e-04	$\max(\rho(A_{14}))$	-1.1e-04
$\max(\rho(A_{19}))$	-1.1e-04	$\max(\rho(A_{25}))$	-7.1e-06	$\max(\rho(A_{31}))$	-1.1e-04
$\max(\rho(A_3))$	-1.1e-04	$\max(\rho(A_9))$	-1.1e-04	$\max(\rho(A_{15}))$	-1.1e-04
$\max(\rho(A_{20}))$	-1.1e-04	$\max(\rho(A_{26}))$	-1.8e-20	$\max(\rho(A_{32}))$	-1.1e-04
$\max(\rho(A_4))$	-4.3e-19	$\max(\rho(A_{10}))$	-3.6e-04	$\max(\rho(A_{16}))$	-1.1e-04
$\max(\rho(A_{21}))$	-1.1e-04	$\max(\rho(A_{27}))$	-6.4e-06	$\max(\rho(A_{33}))$	-1.1e-04
$\max(\rho(A_5))$	-2.3e-07	$\max(\rho(A_{11}))$	-1.1e-04	$\max(\rho(A_{17}))$	-1.1e-04
$\max(\rho(A_{22}))$	-1.1e-04	$\max(\rho(A_{28}))$	-1.1e-04	$\max(\rho(A_{34}))$	-1.1e-04
$\max(\rho(A_6))$	-1.5e-06	$\max(\rho(A_{12}))$	-1.1e-04		
$\max(\rho(A_{23}))$	-1.1e-04	$\max(\rho(A_{29}))$	-4.9e-07		

Table 3: Steady state solution by setting free variables be zeros in Table 1

zero solutions	$x_1 = x_3 = x_4 = x_5 = x_6 = x_7 = x_8 = x_9 = x_{10} = x_{11} = x_{12} = x_{14} = 0,$ $x_{15} = x_{18} = x_{21} = x_{23} = x_{24} = x_{25} = x_{27} = x_{28} = x_{29} = x_{32} = x_{33} = 0$
constant solutions	$x_2 = 3.3383 \times 10^{-9}, x_{22} = 6.67 \times 10^{-9}, x_{26} = 8.33 \times 10^{-10},$ $x_{30} = 5.33 \times 10^{-9}, x_{31} = 4.67 \times 10^{-7}, x_{34} = 2.5 \times 10^{-11}$
polynomial system	$3 \times 10^{-8} - x_{13} - x_{17} = 0$ $k_{18}x_{17} + k_{25}x_{17} - k_{19}x_{16}x_{13} = 0$ $k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20} - k_{19}x_{16}x_{13} = 0$
linear equation	$x_{19} = x_{20} = 2.33 \times 10^{-10} - x_{16} - x_{17}$

Table 4: The relative difference of nonzero components of the steady state solution shown in 3 compared with the solution computed by time marching

x_i	relative difference	x_i	relative difference
x_2	4.67582e-08	x_{22}	7.70551e-09
x_{13}	1.84436e-08	x_{26}	3.94895e-09
x_{16}	7.40565e-08	x_{30}	8.09857e-08
x_{17}	4.58548e-08	x_{31}	6.09728e-08
x_{19}	7.49401e-08	x_{34}	4.63122e-08
x_{20}	7.49401e-08		

Table 5: Nonlinear stability of the steady state solution shown in Table 3

perturbation ϵ_n	$\ \tilde{x} - x_s\ $
$O(10^{-6})$	1.67733×10^{-5}
$O(10^{-7})$	1.33397×10^{-6}
$O(10^{-8})$	3.59671×10^{-7}

Table 6: Steady state solutions using initial conditions from [34]

Constant solutions	$x_1 = x_3 = x_5 = x_6 = x_7 = x_8 = x_9 = x_{10} = x_{12} = x_{18} = x_{23} = x_{24} = x_{25} = 0,$ $x_{27} = x_{28} = x_{29} = 0, x_{26} = 4.66 \times 10^{-9}, x_{30} = 3.25 \times 10^{-8}, x_{34} = 3.1 \times 10^{-10}$
Free variables	$x_4, x_{11}, x_{13}, x_{14}, x_{15}, x_{16}, x_{17}, x_{21}, x_{33}$
Linear expression	$x_2 = 6.45 \times 10^{-8} - x_4, \quad x_{19} = x_{20} = 4.31 \times 10^{-9} - x_{15} - x_{16} - x_{17}$ $x_{22} = 3.33 \times 10^{-8} - x_{21}, \quad x_{31} = 5.32 \times 10^{-6} - x_{14} - x_{33}$ $x_{32} = 5 \times 10^{-7} - x_{11} - x_{13} - x_{17}$
Other equations	$k_{18}x_{17} + k_{25}x_{17} - k_{19}x_{16}x_{13} = 0,$ $k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20} - k_{19}x_{16}x_{13} = 0$

Table 7: Steady state solutions by setting free variables in Table 6 to zeros

zero solutions	$x_1 = x_3 = x_5 = x_6 = x_7 = x_8 = x_9 = x_{10} = x_{12} = x_{18} = x_{23} = x_{24} = 0,$ $x_{25} = x_{27} = x_{28} = x_{29} = 0 = x_4 = x_{11} = x_{14} = x_{15} = x_{21} = x_{32} = x_{33} = 0$
constant solutions	$x_2 = 6.45 \times 10^{-8}, x_{22} = 3.33 \times 10^{-8}, x_{26} = 4.66 \times 10^{-9}, x_{30} = 3.25 \times 10^{-8},$ $x_{31} = 5.32 \times 10^{-6}, x_{34} = 3.1 \times 10^{-10}$
polynomial system	$5 \times 10^{-7} - x_{13} - x_{17} = 0$ $k_{18}x_{17} + k_{25}x_{17} - k_{19}x_{16}x_{13} = 0$ $k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20} - k_{19}x_{16}x_{13} = 0$
linear equation	$x_{19} = x_{20} = 4.31 \times 10^{-9} - x_{16} - x_{17}$

Table 8: Listing of reaction rates analyzed in Hockin-Mann's model. These values are obtained by fitting computational results to empirical data.

Rate no.	Lower bound	Upper bound
1	6.29×10^{-4}	3.1×10^{-3}
2	1.6×10^5	3.2×10^6
3	8.76×10^{-4}	3.1×10^{-3}
4	1.6×10^5	2.3×10^7
8	69 nm(PCPS)	4.35 nm(PCPS)
9	69 nm(PCPS)	4.35 nm(PCPS)
11	17	21
12	2×10^7	2.4×10^7
13	2.1	2.7
14	9×10^6	1.1×10^7
15	1.6	2
23	2×10^4	2.4×10^4
34	8×10^5	1×10^6
35	1×10^{-4}	1.2×10^{-4}
36	2.9×10^8	3.5×10^8
37	4.5×10^7	5.5×10^7

Table 9: Clinical trials information for factor IX, was assembled from <http://ClinicalTrials.gov> using the search terms factor IX, where both open and closed trials were accepted.

Trial Identifier	Condition	Intervention	Purpose
NCT00581126	Hemophilia B	Recombinant Factor IX Coagulation	To assess efficacy and safety of BeneFix for prophylaxis in "Short-term" therapy and on demand therapy for all bleeding episodes of subjects with hemophilia B
NCT00004801	Hemophilia B Factor IX Deficiency	monoclonal factor IX replacement therapy	Assess the safety and long-term efficacy of monoclonal factor IX concentrate in patients with factor IX deficiency.
NCT00000582	Blood Coagulation Disorders Hematologic Diseases Hemorrhagic Disorders	Factor IX	To test the efficacy of prothrombin complex concentrates (Factor IX) in the treatment of hemophiliac patients who had inhibitors to Factor VIII.
NCT01159210	Prothrombin Complex Factor Deficiency	Prothrombin complex concentrate	To assess the efficacy and safety of Prothromplex Total as a treatment for the immediate reversal of oral anticoagulant therapy with vitamin K antagonists in patients with acquired deficiency of prothrombin complex coagulation factors (II, VII, IX, X).

Figures

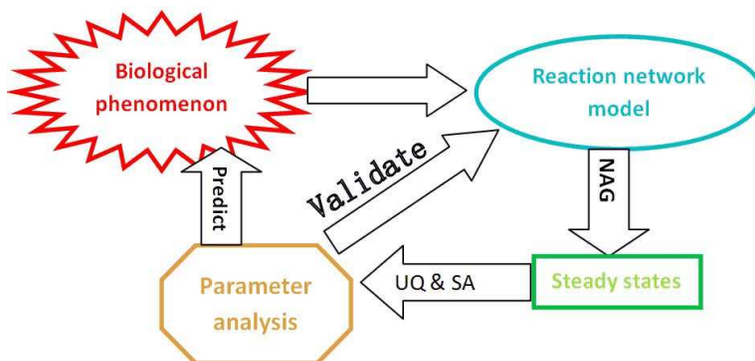


Figure 1: The general approach diagram of different stages in the analysis process: Numerical algebraic geometry (NAG) provides a numerical approach to compute all the steady states; Ranking importance of parameters, which is performed by uncertainty quantification (UQ) and sensitivity analysis (SA), validates the mathematical model and gives some medical prediction to the biological application.

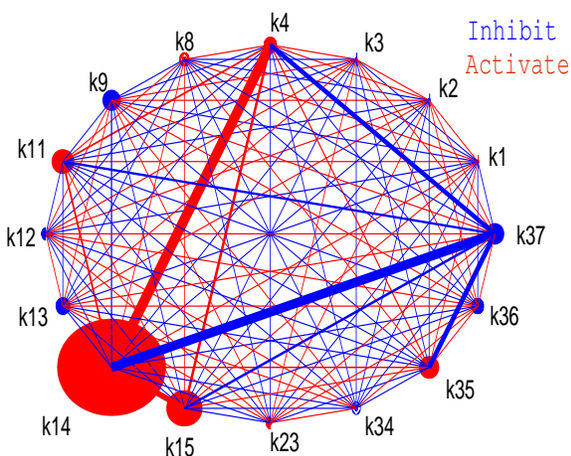


Figure 2: Network graph of importance of reaction rates and the interaction between pair reactions with respect to total thrombin. The radius of circles corresponding to reaction rate measures the rank of sensitivity while the width of lines of any pair of reaction rates represents rank of their co-sensitivity. The red color indicates that they play an activate role in generating thrombin production while the blue color means that they inhibit the generation of thrombin production. This figure shows that k_{14} , k_{15} and k_{11} are the most sensitive reaction rates in activating thrombin while k_{37} and k_9 inhibit thrombin production. Variance of k_4 and k_{14} can play a positive role in generation of thrombin production, however k_{37} and k_{14} inhibit generation of thrombin production.

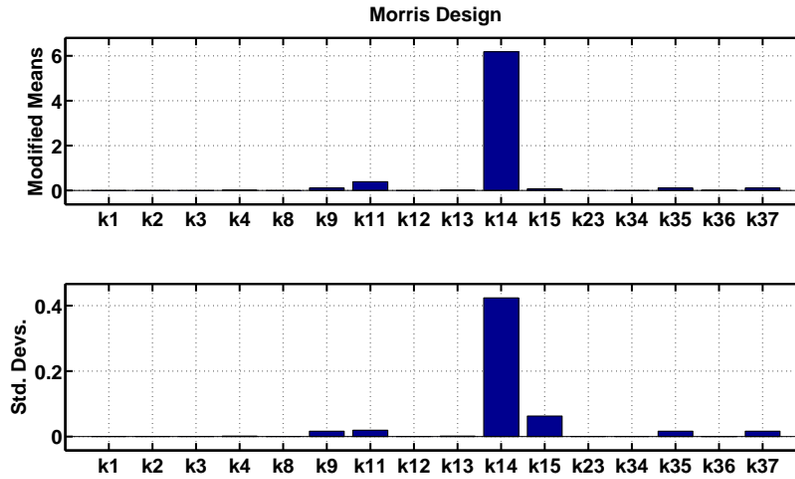


Figure 3: Modified means and standard deviations based sensitivity ranking of reaction rates with respect to the total thrombin using “Morris one at a time” method. Horizontal axis shows reaction rates. Vertical axis shows modified means and standard deviations respectively. This figure indicates that k_{14} has the largest modified mean and standard deviation. Therefore k_{14} is the most sensitive to thrombin production.

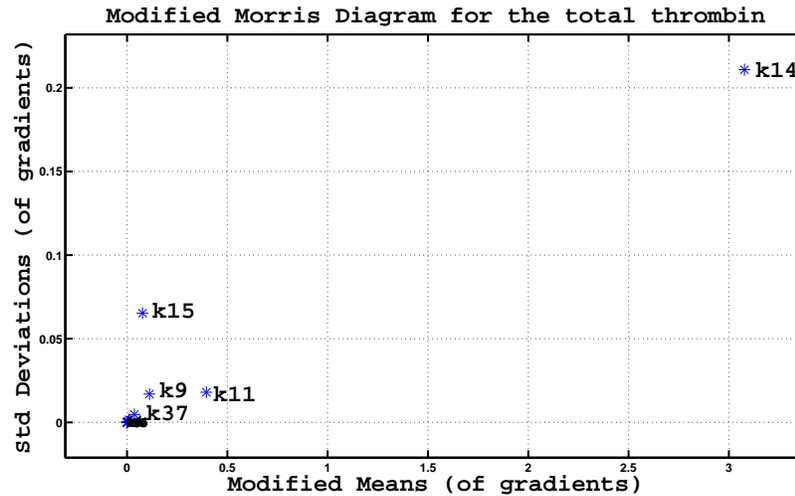


Figure 4: Modified Morris diagram on sensitivity ranking of reaction rates with respect to the total thrombin. Ranks of sensitivity for individual reaction rates are determined by their distance from the origin. From this figure, the five most sensitive reaction rates are k_{14} , k_{15} , k_{11} , k_9 and k_{37} . The other reaction rates cluster together due to the same magnitude of sensitivity.

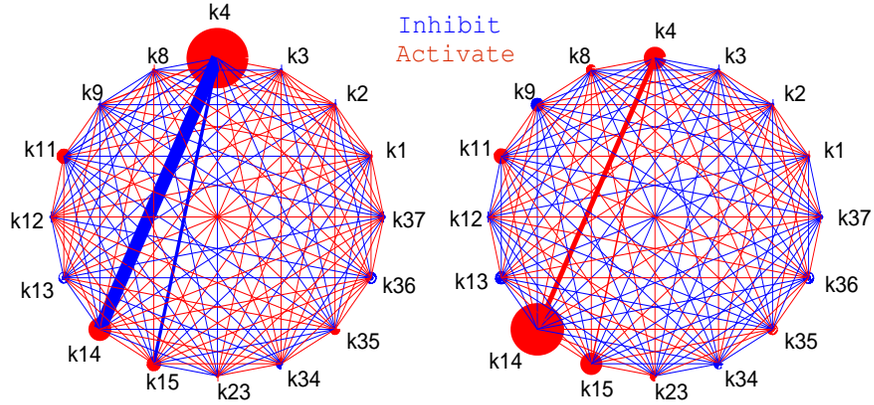


Figure 5: Network graph of importance of reaction rates and the interaction between reaction rates with respect to IXa=VIIIa (x_{33}) and IXa=ATIII (x_{30}) respectively. The same legend is used with Figure 1. This figure shows that the rank of sensitivity varies with respect to different output variables.

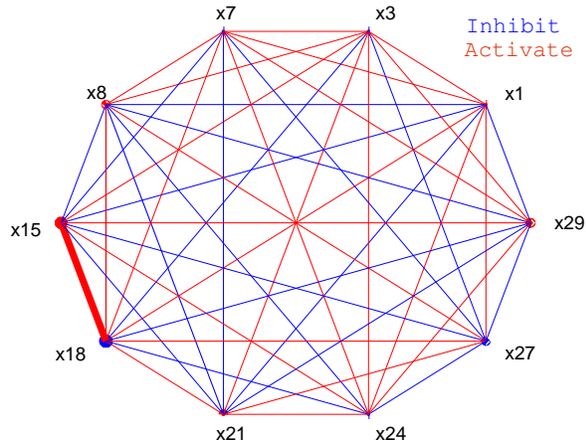


Figure 6: Network graph of importance of initial condition and the interaction between pair components with respect to total thrombin. See the legend in Figure 3. This figure shows that factor X (x_{15}) is the most sensitive reaction component in activating thrombin while TFPI (x_{18}) inhibits thrombin production. Moreover, their variance can play a positive role in generation of thrombin production.

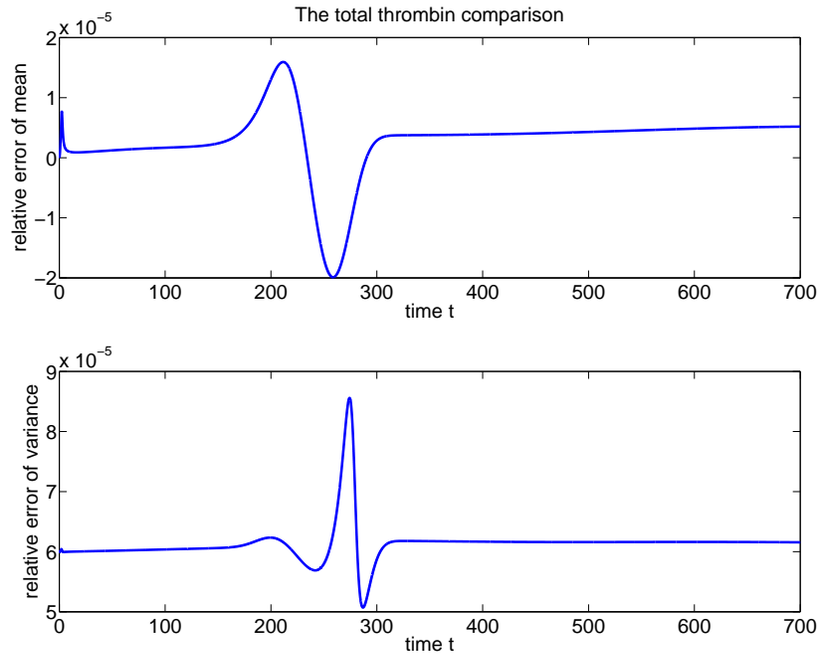


Figure 7: The Difference of mean and variance of the total thrombin obtained by SGPCM (545 points) and MC (1000 points). The figure shows that the results of two methods are same under small tolerance.

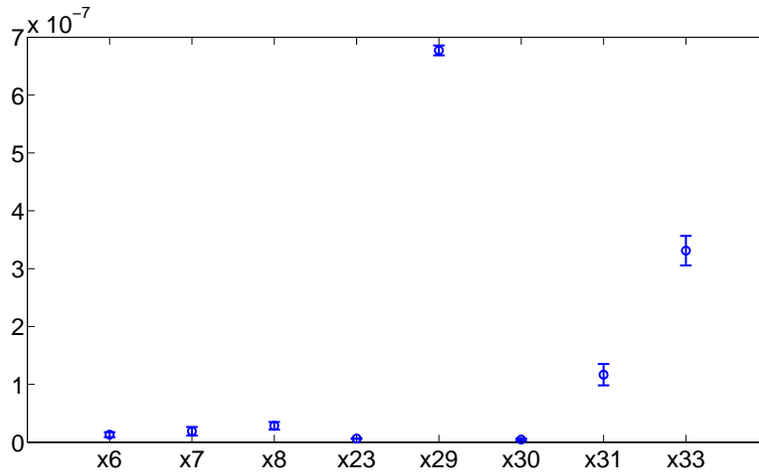


Figure 8: Mean and error bar of steady state solutions whose magnitude are greater than 10^{-10} . Horizontal axis is reaction rates. This figure shows confidence intervals of each reaction rates.

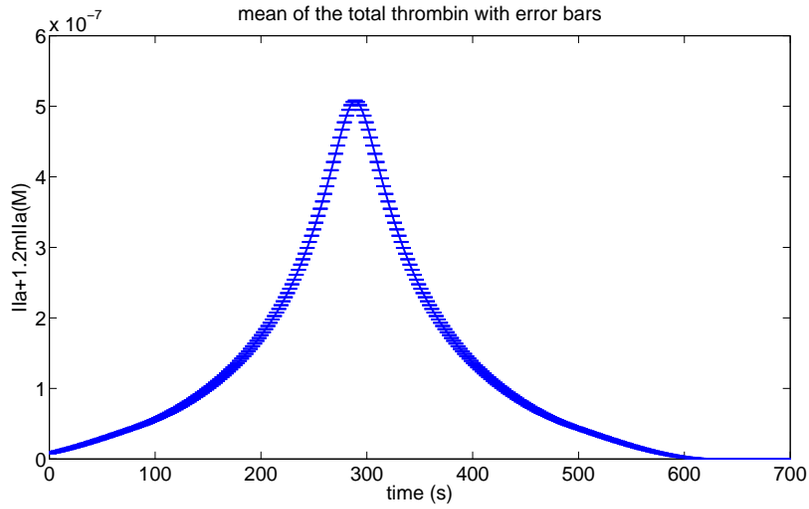


Figure 9: The mean and error bar of the total thrombin along the solution by time marching. Error bars show the confidence intervals of reaction rates along the curve.

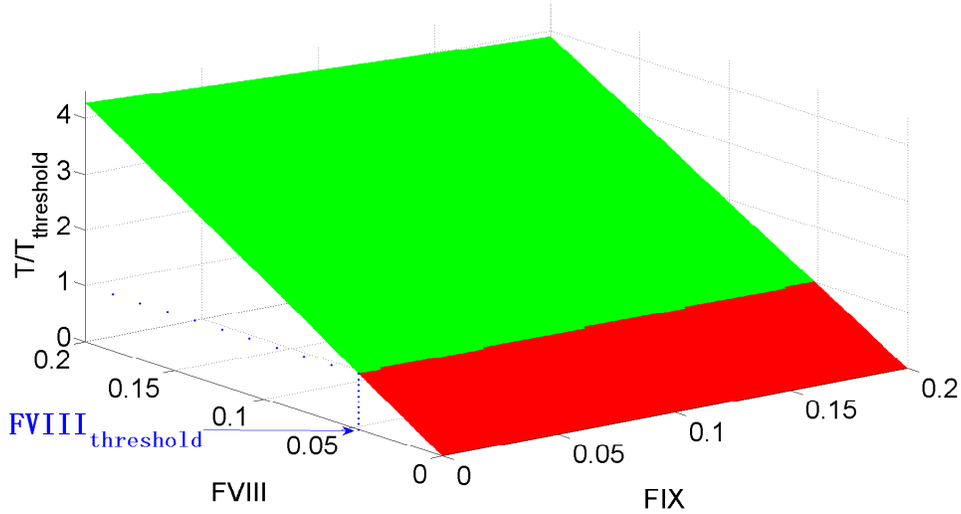


Figure 10: The total thrombin *v.s.* factors VIII (FVIII) and IX (FIX). $T_{threshold}$ stands for the threshold value of thrombin. The red portion of the plane means the total thrombin (T) is smaller than $T_{threshold}$ while the green portion means the total thrombin is greater than $T_{threshold}$. This figure shows that the threshold of factor VIII is 4.7 % of normal value, and that the total thrombin is greater than $T_{threshold}$ when factor VIII is greater than $FVIII_{threshold}$ regardless of factor IX concentration when it's in the assume range.

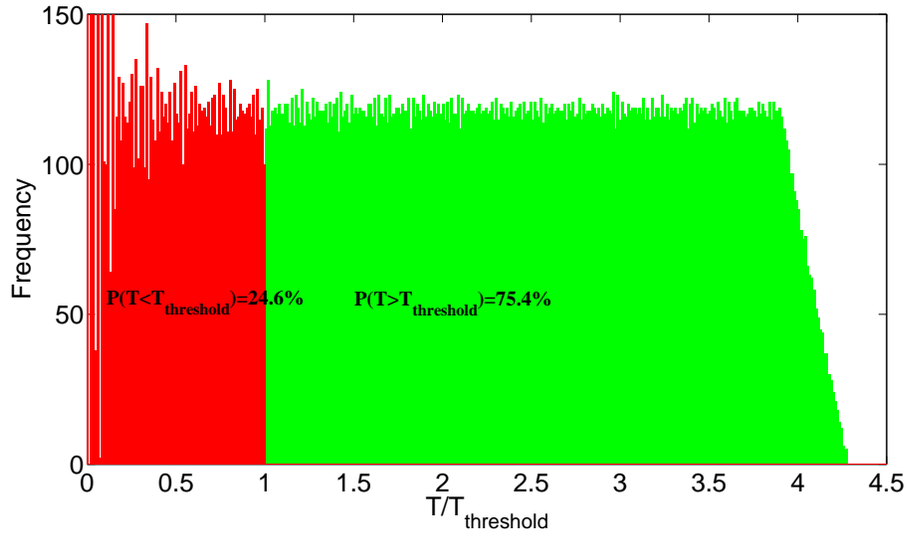


Figure 11: Histogram of the total thrombin. The probability that the total thrombin is greater than $T_{threshold}$ is 75.4%.

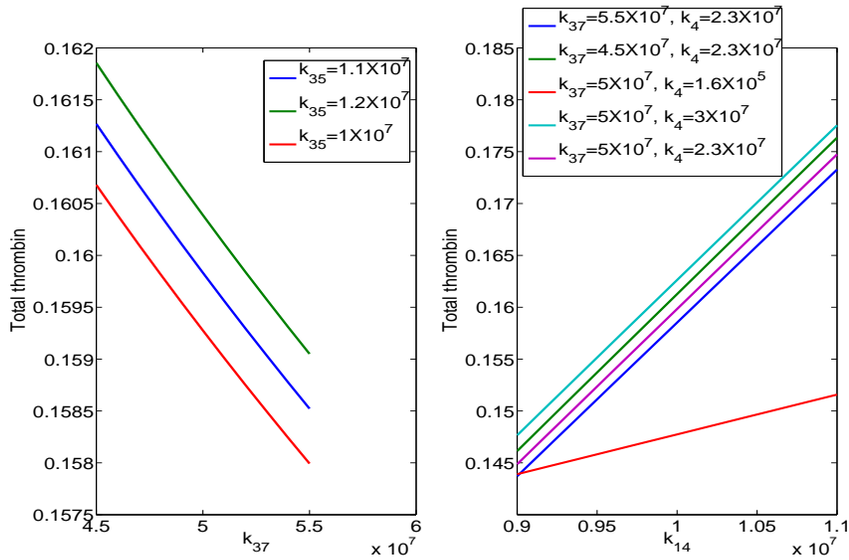


Figure 12: The total thrombin *v.s.* reaction rates k_{37} and k_{14} . k_{37} inhibits the generation of total thrombin as it increases. k_{35} plays a negative role in the effect of k_{37} . k_{14} activates the generation of total thrombin as it increases. k_{37} and k_4 play negative and positive roles in the effect of k_{14} respectively.

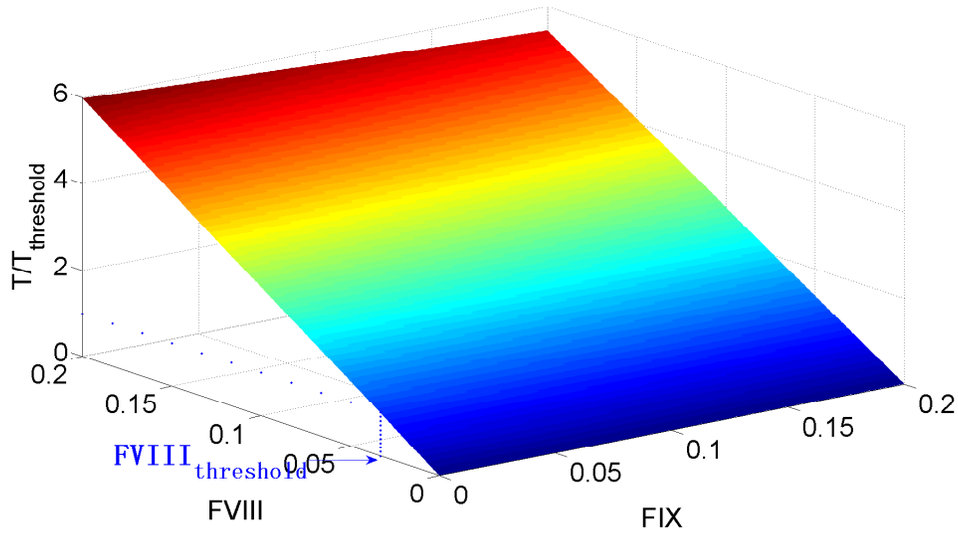


Figure 13: The mean of the total thrombin of sampling points of k_{14} and factor X, which vary from 100%-200% of their normal values. The color means the variance of the total thrombin. This figure shows that the threshold of factor VIII is decreased to 3.3% of normal value.

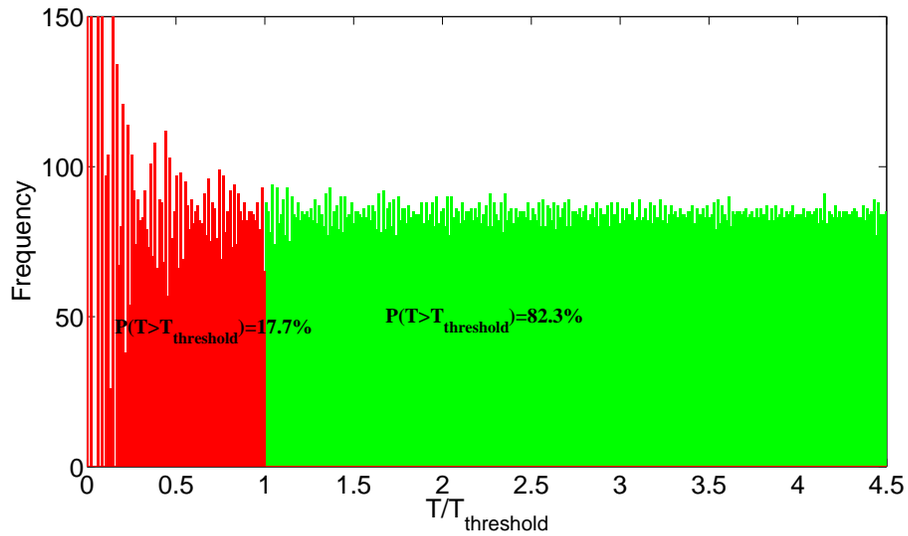


Figure 14: Histogram of the total thrombin as reaction rate k_{14} and factor X is introduced. The probability that the total thrombin is greater than $T_{\text{threshold}}$ is increased to 82.3%.

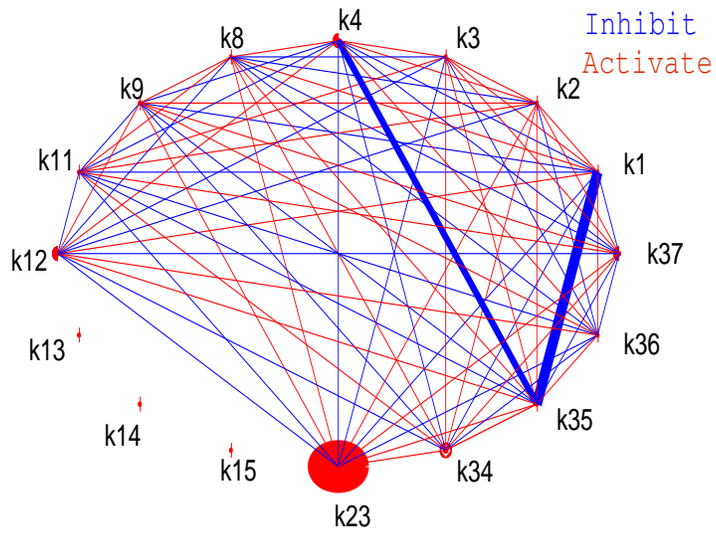


Figure 15: Network graph of importance of reaction rates and the interaction between pair components with respect to total thrombin when blocking k_{14} and k_{15} (their values are reduced by 90%). See the legend in Figure 1. This figure shows that the sensitivity shifts, i.e., reaction rate k_{23} becomes the most sensitive reaction rates in activating thrombin when k_{14} and k_{25} are blocked. Moreover, k_{13} plays no roles since chemical reaction with k_{13} is combined with k_{14} and k_{25} .

APPENDIX A

Chemical equations

Table 10: Chemical expressions for the coagulation cascade

Line	Chemical expressions
1	$TF + VII < 1 - 2 > TF=VII$
2	$TF + VIIa < 3 - 4 > TF=VIIa$
3	$TF=VIIa + VII -5 > TF=VIIa + VIIa$
4	$Xa + VII -6 > Xa + VIIa$
5	$IIa + VII -7 > IIa + VIIa$
6	$TF=VIIa + X < 8 - 9 > TF=VIIa=X -10 > TF=VIIa=Xa$
7	$TF=VIIa +Xa < 11 - 12 > TF=VIIa=Xa$
8	$TF=VIIa + IX < 13 - 14 > TF=VIIa=IX -15 > TF=VIIa + IXa$
9	$Xa + II -16 > Xa +IIa$
10	$IIa + VIII -17 > IIa + VIIIa$
11	$VIIIa + IXa < 18 - 19 > IXa=VIIIa$
12	$IXa=VIIIa + X < 20 - 21 > IXa=VIIIa=X -22 > IXa=VIIIa + Xa$
13	$VIIIa < 23 - 24 > VIIIa_1 \cdot L + VIIIa_2$
14	$IXa=VIIIa=X -25 > VIIIa_1 \cdot L + VIIIa_2 + X +IXa$
15	$IXa=VIIIa -25 > VIIIa_1 \cdot L + VIIIa_2 +IXa$
16	$IIa+ V -26 > IIa +Va$
17	$Xa + Va < 27 - 28 > Xa=Va$
18	$Xa=Va + II < 29 - 30 > Xa=Va=II -31 > Xa=Va + mIIa$
19	$mIIa+ Xa=Va -32 > IIa + Xa=Va$
20	$Xa + TFPI < 33 - 34 > Xa=TFPI$
21	$TF=VIIa=Xa + TFPI < 35 - 36 > TF=VIIa=Xa=TFPI$
22	$TF=VIIa + Xa=TFPI -37 > TF=VIIa=Xa=TFPI$
23	$Xa+ATIII -38 > Xa=ATIII$
24	$mIIa+ ATIII -39 > mIIa=ATIII$
25	$IXa + ATIII -40 > IXa=ATIII$
26	$IIa + ATIII -41 > IIa=ATIII$
27	$TF=VIIa + ATIII -42 > TF=VIIa=ATIII$

Table 11: Variables and Species

x_1	TF	x_{13}	IIa	x_{25}	IXa
x_2	IXa=VIIIa=X	x_{14}	Xa=Va=II	x_{26}	Xa=ATIII
x_3	VII	x_{15}	X	x_{27}	II
x_4	VIIIa ₁ · L	x_{16}	mIIa	x_{28}	mIIa=ATIII
x_5	TF=VII	x_{17}	TF=VIIa=X	x_{29}	VIII
x_6	VIIIa ₂	x_{18}	TFPI	x_{30}	IXa=ATIII
x_7	VIIa	x_{19}	TF=VIIa=Xa	x_{31}	VIIIa
x_8	V	x_{20}	Xa=TFPI	x_{32}	IIa=ATIII
x_9	TF=VIIa	x_{21}	IX	x_{33}	IXa=VIIIa
x_{10}	Va	x_{22}	TF=VIIa=Xa=TFPI	x_{34}	TF=VIIa=ATIII
x_{11}	Xa	x_{23}	TF=VIIa=IX		
x_{12}	Xa=Va	x_{24}	ATIII		

APPENDIX B

Model equations

$$\begin{aligned}
\frac{dx_1}{dt} &= -k_2x_1x_2 + k_1x_3 - k_4x_1x_4 + k_3x_5, & \frac{dx_2}{dt} &= -k_2x_1x_2 + k_1x_3 - k_5x_5x_2 - k_6x_6x_2 - k_7x_7x_2 \\
\frac{dx_3}{dt} &= k_2x_1x_2 - k_1x_3, & \frac{dx_4}{dt} &= -k_4x_1x_4 + k_3x_5 + k_5x_5x_2 + k_6x_6x_2 + k_7x_7x_2 \\
\frac{dx_5}{dt} &= k_4x_1x_4 - k_3x_5 - k_9x_5x_8 + k_8x_9 - k_{12}x_5x_6 + k_{11}x_{10} - k_{14}x_5x_{11} + k_{13}x_{12} - k_{37}x_5x_{27} \\
&\quad - k_{42}x_5x_{29} + k_{15}x_{12} \\
\frac{dx_6}{dt} &= -k_{12}x_5x_6 + k_{11}x_{10} - k_{28}x_6x_{22} + k_{27}x_{23} - k_{34}x_6x_{26} + k_{33}x_{27} - k_{38}x_6x_{29} + k_{22}x_{18} \\
\frac{dx_7}{dt} &= k_{16}x_6x_{14} + k_{32}x_{25}x_{23} - k_{41}x_7x_{29}, & \frac{dx_8}{dt} &= -k_9x_5x_8 + k_8x_9 - k_{21}x_{17}x_8 + k_{20}x_{18} + k_{25}x_{19} \\
\frac{dx_9}{dt} &= k_9x_5x_8 - k_8x_9 - k_{10}x_9, & \frac{dx_{10}}{dt} &= k_{12}x_5x_6 - k_{11}x_{10} - k_{36}x_{10}x_{26} + k_{35}x_{28} + k_{10}x_9 \\
\frac{dx_{11}}{dt} &= -k_{14}x_5x_{11} + k_{13}x_{12}, & \frac{dx_{12}}{dt} &= k_{14}x_5x_{11} - k_{13}x_{12} - k_{15}x_{12} \\
\frac{dx_{13}}{dt} &= -k_{19}x_{16}x_{13} + k_{18}x_{17} + k_{25}x_{18} + k_{25}x_{17} - k_{40}x_{13}x_{29} + k_{15}x_{12} \\
\frac{dx_{14}}{dt} &= -k_{16}x_6x_{14} - k_{30}x_{23}x_{14} + k_{29}x_{24} \\
\frac{dx_{15}}{dt} &= -k_{17}x_7x_{15}, & \frac{dx_{16}}{dt} &= k_{17}x_7x_{15} - k_{19}x_{16}x_{13} + k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20} \\
\frac{dx_{17}}{dt} &= k_{19}x_{16}x_{13} - k_{18}x_{17} - k_{21}x_{17}x_8 + k_{20}x_{18} - k_{25}x_{17} + k_{22}x_{18} \\
\frac{dx_{18}}{dt} &= k_{21}x_{17}x_8 - k_{20}x_{18} - k_{25}x_{18} - k_{22}x_{18} \\
\frac{dx_{19}}{dt} &= \frac{dx_{20}}{dt} = k_{24}x_{16} - k_{23}x_{19}x_{20} + k_{25}x_{18} + k_{25}x_{17} \\
\frac{dx_{21}}{dt} &= -k_{26}x_7x_{21}, & \frac{dx_{22}}{dt} &= k_{26}x_7x_{21} - k_{28}x_6x_{22} + k_{27}x_{23} \\
\frac{dx_{23}}{dt} &= k_{28}x_6x_{22} - k_{27}x_{23} - k_{30}x_{23}x_{14} + k_{29}x_{24} + k_{31}x_{24} \\
\frac{dx_{24}}{dt} &= k_{30}x_{23}x_{14} - k_{29}x_{24} - k_{31}x_{24}, & \frac{dx_{25}}{dt} &= -k_{32}x_{25}x_{23} - k_{39}x_{25}x_{29} + k_{31}x_{24} \\
\frac{dx_{26}}{dt} &= -k_{34}x_6x_{26} + k_{33}x_{27} - k_{36}x_{10}x_{26} + k_{35}x_{28} \\
\frac{dx_{27}}{dt} &= k_{34}x_6x_{26} - k_{33}x_{27} - k_{37}x_5x_{27}, & \frac{dx_{28}}{dt} &= k_{36}x_{10}x_{26} - k_{35}x_{28} + k_{37}x_5x_{27} \\
\frac{dx_{29}}{dt} &= -k_{38}x_6x_{29} - k_{39}x_{25}x_{29} - k_{40}x_{13}x_{29} - k_{41}x_7x_{29} - k_{42}x_5x_{29}, & \frac{dx_{30}}{dt} &= k_{38}x_6x_{29} \\
\frac{dx_{31}}{dt} &= k_{39}x_{25}x_{29}, & \frac{dx_{32}}{dt} &= k_{40}x_{13}x_{29}, & \frac{dx_{33}}{dt} &= k_{41}x_7x_{29}, & \frac{dx_{34}}{dt} &= k_{42}x_5x_{29}, \tag{7}
\end{aligned}$$

where $k_i, i = 1, 2, \dots, 42$ are constants, and $x_i, i = 1, 2, \dots, 34$ are variables representing the species (see Table 11). The steady state system can be obtained by taking $t \rightarrow \infty$, i.e., $\frac{dx_i}{dt} = 0, \forall i$.

APPENDIX C

$$f_1 = -k_4x_1x_4 + k_3x_5$$

$$f_2 = -k_5x_5x_2 - k_6x_6x_2 - k_7x_7x_2$$

$$f_3 = k_2x_1x_2 - k_1x_3$$

$$f_4 = -k_{12}x_5x_6 - k_{42}x_5x_{29} + k_4x_1x_4 - k_3x_5 + k_8x_9 - k_9x_5x_8 + k_{11}x_{10} - k_{37}x_5x_{27}$$

$$f_5 = -k_{12}x_5x_6 - k_{34}x_6x_{26} + k_{11}x_{10} + k_{33}x_{27} - k_{38}x_6x_{29} + k_{22}x_{18}$$

$$f_6 = k_{16}x_6x_{14} + k_{32}x_{25}x_{23} - k_{41}x_7x_{29}$$

$$f_7 = -k_9x_5x_8 + k_8x_9 - k_{21}x_{17}x_8 + k_{20}x_{18} + k_{25}x_{18}$$

$$f_8 = k_9x_5x_8 - k_8x_9 - k_{10}x_9$$

$$f_9 = k_{12}x_5x_6 - k_{11}x_{10} - k_{36}x_{10}x_{26} + k_{35}x_{28} + k_{10}x_9$$

$$f_{10} = -k_{14}x_5x_{11} + k_{13}x_{12}$$

$$f_{11} = -k_{15}x_{12}$$

$$f_{12} = -k_{19}x_{16}x_{13} + k_{18}x_{17} + k_{25}x_{18} + k_{25}x_{17} - k_{40}x_{13}x_{29}$$

$$f_{13} = -k_{16}x_6x_{14} - k_{30}x_{23}x_{14} + k_{29}x_{24}$$

$$f_{14} = -k_{17}x_7x_{15}$$

$$f_{15} = -k_{19}x_{16}x_{13} + k_{18}x_{17} - k_{24}x_{16} + k_{23}x_{19}x_{20}$$

$$f_{16} = k_{19}x_{16}x_{13} - k_{18}x_{17} - k_{21}x_{17}x_8 + k_{20}x_{18} - k_{25}x_{17} + k_{22}x_{18}$$

$$f_{17} = k_{21}x_{17}x_8 - k_{20}x_{18} - k_{25}x_{18} - k_{22}x_{18}$$

$$f_{18} = -k_{26}x_7x_{21}$$

$$f_{19} = -k_{28}x_6x_{22} + k_{27}x_{23}$$

$$f_{20} = k_{29}x_{24} - k_{30}x_{23}x_{14} + k_{31}x_{24}$$

$$f_{21} = -k_{32}x_{25}x_{23} + k_{31}x_{24}$$

$$f_{22} = -k_{34}x_6x_{26} + k_{33}x_{27} - k_{36}x_{10}x_{26} + k_{35}x_{28}$$

$$f_{23} = k_{34}x_6x_{26} - k_{33}x_{27} - k_{37}x_5x_{27}$$

$$f_{24} = k_{39}x_{25}x_{29},$$

The next 10 rows can be expressed as:

$$A_4 = A_1 - A_2$$

$$A_{19} = -A_{15} - A_{16} - A_{17} - A_{18}$$

$$A_{20} = -A_{15} - A_{16} - A_{17} - A_{18}$$

$$A_{24} = -A_{21} - A_{22} - A_{23}$$

$$A_{28} = -A_{26} - A_{27}$$

$$A_{29} = A_1 + A_3 + A_5 + A_6 + A_7 + A_8 + 2A_9 + 2A_{10} + A_{11} + 2A_{12} + A_{13} + A_{14} + A_{17} + 2A_{18} \\ - 2A_{21} - 2A_{22} - A_{23} + A_{25} - 2A_{26} - A_{27}$$

$$A_{30} = -A_6 - A_8 - A_9 - A_{10} - A_{18} + A_{21} + A_{22} + A_{26}$$

$$A_{32} = -A_{11} - A_{12} - A_{13} - A_{17}$$

$$A_{33} = -A_7 - A_{14} + A_{21} + A_{22} + A_{23} - A_{25} - A_{31}$$

$$A_{34} = -A_1 - A_3 - A_5 - A_9 - A_{10} - A_{12} + A_{26} + A_{27},$$

where A_i , $i = 1, 2, \dots, 34$ represents the i -th row of A .

APPENDIX D

The linearized system of differential equation system shown in Appendix B:

$$\begin{aligned}
x^1_{1t} &= -k_2x^s_1x^1_2 - k_2x^1_1x^s_2 + k_1x^1_3 - k_4x^s_1x^1_4 - k_4x^1_1x^s_4 + k_3x^1_5 \\
x^1_{2t} &= -k_2x^s_1x^1_2 - k_2x^1_1x^s_2 + k_1x^1_3 - k_5x^s_5x^1_2 - k_5x^1_5x^s_2 - k_6x^s_6x^1_2 - k_6x^1_6x^s_2 \\
&\quad -k_7x^s_7x^1_2 - k_7x^1_7x^s_2 \\
x^1_{3t} &= k_2x^s_1x^1_2 + k_2x^1_1x^s_2 - k_1x^1_3 \\
x^1_{4t} &= -k_4x^s_1x^1_4 - k_4x^1_1x^s_4 + k_3x^1_5 + k_5x^s_5x^1_2 + k_5x^1_5x^s_2 + k_6x^s_6x^1_2 + k_6x^1_6x^s_2 \\
&\quad +k_7x^s_7x^1_2 + k_7x^1_7x^s_2 \\
x^1_{5t} &= k_8x^1_9 + k_4x^s_1x^1_4 + k_4x^1_1x^s_4 + k_{13}x^1_{12} + k_{11}x^1_{10} - k_3x^1_5 - k_9x^s_5x^1_8 - k_9x^1_5x^s_8 \\
&\quad -k_{12}x^s_5x^1_6 - k_{12}x^1_5x^s_6 - k_{14}x^s_5x^1_{11} - k_{14}x^1_5x^s_{11} - k_{37}x^s_5x^1_{27} - k_{37}x^1_5x^s_{27} \\
&\quad -k_{42}x^s_5x^1_{29} - k_{42}x^1_5x^s_{29} + k_{15}x^1_{12} \\
x^1_{6t} &= k_{33}x^1_{27} + k_{27}x^1_{23} + k_{22}x^1_{18} + k_{11}x^1_{10} - k_{12}x^s_5x^1_6 - k_{12}x^1_5x^s_6 - k_{28}x^s_6x^1_{22} \\
&\quad -k_{28}x^1_6x^s_{22} - k_{34}x^s_6x^1_{26} - k_{34}x^1_6x^s_{26} - k_{38}x^s_6x^1_{29} - k_{38}x^1_6x^s_{29} \\
x^1_{7t} &= k_{16}x^s_6x^1_{14} + k_{16}x^1_6x^s_{14} + k_{32}x^s_{25}x^1_{23} + k_{32}x^1_{25}x^s_{23} - k_{41}x^s_7x^1_{29} - k_{41}x^1_7x^s_{29} \\
x^1_{8t} &= -k_9x^s_5x^1_8 - k_9x^1_5x^s_8 + k_8x^1_9 - k_{21}x^s_{17}x^1_8 - k_{21}x^1_{17}x^s_8 + k_{20}x^1_{18} + k_{25}x^1_{18} \\
x^1_{9t} &= k_9x^s_5x^1_8 + k_9x^1_5x^s_8 - k_8x^1_9 - k_{10}x^1_9 \\
x^1_{10t} &= k_{12}x^s_5x^1_6 + k_{12}x^1_5x^s_6 - k_{11}x^1_{10} - k_{36}x^s_{10}x^1_{26} - k_{36}x^1_{10}x^s_{26} + k_{35}x^1_{28} + k_{10}x^1_9 \\
x^1_{11t} &= -k_{14}x^s_5x^1_{11} - k_{14}x^1_5x^s_{11} + k_{13}x^1_{12} \\
x^1_{12t} &= k_{14}x^s_5x^1_{11} + k_{14}x^1_5x^s_{11} - k_{13}x^1_{12} - k_{15}x^1_{12} \\
x^1_{13t} &= -k_{19}x^s_{16}x^1_{13} - k_{19}x^1_{16}x^s_{13} + k_{18}x^1_{17} + k_{25}x^1_{18} + k_{25}x^1_{17} - k_{40}x^s_{13}x^1_{29} - k_{40}x^1_{13}x^s_{29} \\
&\quad +k_{15}x^1_{12} \\
x^1_{14t} &= -k_{16}x^s_6x^1_{14} - k_{16}x^1_6x^s_{14} - k_{30}x^s_{23}x^1_{14} - k_{30}x^1_{23}x^s_{14} + k_{29}x^1_{24} \\
x^1_{15t} &= -k_{17}(x^s_7x^1_{15} + x^1_7x^s_{15}) \\
x^1_{16t} &= k_{17}x^s_7x^1_{15} + k_{17}x^1_7x^s_{15} - k_{19}x^s_{16}x^1_{13} - k_{19}x^1_{16}x^s_{13} + k_{18}x^1_{17} - k_{24}x^1_{16} + k_{23}x^s_{19}x^1_{20} \\
&\quad +k_{23}x^1_{19}x^s_{20} \\
x^1_{17t} &= k_{19}x^s_{16}x^1_{13} + k_{19}x^1_{16}x^s_{13} - k_{18}x^1_{17} - k_{21}x^s_{17}x^1_8 - k_{21}x^1_{17}x^s_8 + k_{20}x^1_{18} - k_{25}x^1_{17} \\
&\quad +k_{22}x^1_{18}
\end{aligned}$$

$$\begin{aligned}
x^1_{18t} &= k_{21}x^s_{17}x^1_8 + k_{21}x^1_{17}x^s_8 - k_{20}x^1_{18} - k_{25}x^1_{18} - k_{22}x^1_{18} \\
x^1_{19t} &= k_{24}x^1_{16} - k_{23}x^s_{19}x^1_{20} - k_{23}x^1_{19}x^s_{20} + k_{25}x^1_{18} + k_{25}x^1_{17} \\
x^1_{20t} &= k_{24}x^1_{16} - k_{23}x^s_{19}x^1_{20} - k_{23}x^1_{19}x^s_{20} + k_{25}x^1_{18} + k_{25}x^1_{17} \\
x^1_{21t} &= -k_{26}(x^s_7x^1_{21} + x^1_7x^s_{21}) \\
x^1_{22t} &= k_{26}x^s_7x^1_{21} + k_{26}x^1_7x^s_{21} - k_{28}x^s_6x^1_{22} - k_{28}x^1_6x^s_{22} + k_{27}x^1_{23} \\
x^1_{23t} &= k_{28}x^s_6x^1_{22} + k_{28}x^1_6x^s_{22} - k_{27}x^1_{23} - k_{30}x^s_{23}x^1_{14} - k_{30}x^1_{23}x^s_{14} + k_{29}x^1_{24} + k_{31}x^1_{24} \\
x^1_{24t} &= k_{30}x^s_{23}x^1_{14} + k_{30}x^1_{23}x^s_{14} - k_{29}x^1_{24} - k_{31}x^1_{24} \\
x^1_{25t} &= -k_{32}x^s_{25}x^1_{23} - k_{32}x^1_{25}x^s_{23} - k_{39}x^s_{25}x^1_{29} - k_{39}x^1_{25}x^s_{29} + k_{31}x^1_{24} \\
x^1_{26t} &= -k_{34}x^s_6x^1_{26} - k_{34}x^1_6x^s_{26} + k_{33}x^1_{27} - k_{36}x^s_{10}x^1_{26} - k_{36}x^1_{10}x^s_{26} + k_{35}x^1_{28} \\
x^1_{27t} &= k_{34}x^s_6x^1_{26} + k_{34}x^1_6x^s_{26} - k_{33}x^1_{27} - k_{37}x^s_5x^1_{27} - k_{37}x^1_5x^s_{27} \\
x^1_{28t} &= k_{36}x^s_{10}x^1_{26} + k_{36}x^1_{10}x^s_{26} - k_{35}x^1_{28} + k_{37}x^s_5x^1_{27} + k_{37}x^1_5x^s_{27} \\
x^1_{29t} &= -k_{42}x^s_5x^1_{29} - k_{42}x^1_5x^s_{29} - k_{38}x^s_6x^1_{29} - k_{38}x^1_6x^s_{29} - k_{41}x^s_7x^1_{29} \\
&\quad - k_{41}x^1_7x^s_{29} - k_{40}x^s_{13}x^1_{29} - k_{40}x^1_{13}x^s_{29} - k_{39}x^s_{25}x^1_{29} - k_{39}x^1_{25}x^s_{29} \\
x^1_{30t} &= k_{38}(x^s_6x^1_{29} + x^1_6x^s_{29}) \\
x^1_{31t} &= k_{39}(x^s_{25}x^1_{29} + x^1_{25}x^s_{29}) \\
x^1_{32t} &= k_{40}(x^s_{13}x^1_{29} + x^1_{13}x^s_{29}) \\
x^1_{33t} &= k_{41}(x^s_7x^1_{29} + x^1_7x^s_{29}) \\
x^1_{34t} &= k_{42}(x^s_5x^1_{29} + x^1_5x^s_{29})
\end{aligned}$$

Appendix E

Table 12: Convergence study of SGPCM

Abs. Err.				
No.	Mean		Variance	
	184	1228	184	1228
1	2.285678e-004	6.276008e-005	1.414116e+000	5.270631e-001
2	3.528170e-003	9.790568e-004	1.421438e+000	5.282797e-001
3	1.117445e-002	3.058945e-003	1.451133e+000	5.334394e-001
4	1.737843e-004	4.822458e-005	1.421425e+000	5.282775e-001
5	8.828347e-003	2.389501e-003	1.486561e+000	5.391973e-001
6	1.988338e-002	5.317003e-003	1.566493e+000	5.520586e-001
7	4.584148e-003	1.255587e-003	1.427882e+000	5.293661e-001
8	2.048073e-004	5.578918e-005	1.454606e+000	5.339573e-001
9	3.396093e-003	9.241881e-004	1.424554e+000	5.287619e-001
10	3.256174e-003	8.869390e-004	1.423624e+000	5.286163e-001
11	8.412776e-006	2.279950e-006	1.480436e+000	5.382099e-001
12	8.808674e-003	2.384262e-003	1.486394e+000	5.391702e-001
13	8.277039e-003	2.243433e-003	1.479954e+000	5.381284e-001
14	3.103763e-003	8.465279e-004	1.443678e+000	5.320069e-001
15	7.814273e-003	2.193114e-003	1.426201e+000	5.286388e-001
16	1.328375e-002	3.739625e-003	1.415327e+000	5.270061e-001
17	9.094976e-003	2.458848e-003	1.490370e+000	5.397825e-001
18	8.172212e-003	2.212621e-003	1.483100e+000	5.386074e-001
19	2.363583e-003	6.446736e-004	1.460426e+000	5.347973e-001
20	2.363583e-003	6.446736e-004	1.460426e+000	5.347973e-001
21	7.814273e-003	2.193114e-003	1.426201e+000	5.286388e-001
22	7.593356e-003	2.131857e-003	1.425644e+000	5.288239e-001
23	2.706446e-003	7.513140e-004	1.429067e+000	5.297737e-001
24	1.078126e-002	2.958615e-003	1.431272e+000	5.298548e-001
25	9.639244e-003	2.646268e-003	1.426693e+000	5.291310e-001
26	3.609443e-004	9.730916e-005	1.492187e+000	5.400877e-001
27	1.019074e-002	2.748554e-003	1.499708e+000	5.413532e-001
28	2.549383e-003	6.949484e-004	1.420171e+000	5.280515e-001
29	5.662490e-004	1.541640e-004	1.448082e+000	5.327725e-001
30	9.960178e-003	2.677646e-003	1.508239e+000	5.426681e-001
31	2.958200e-003	8.078085e-004	1.445202e+000	5.322928e-001
32	7.845463e-003	2.126566e-003	1.478513e+000	5.378746e-001
33	2.120963e-003	5.769229e-004	1.427220e+000	5.293214e-001
34	8.380451e-003	2.269408e-003	1.483615e+000	5.387117e-001