Reduced order modeling and analysis of the human complement system

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Abstract

Complement is an important pathway in innate immunity, inflammation, and many disease processes. However, despite its importance, there are few validated mathematical models of complement activation. In this study, we developed an ensemble of experimentally validated reduced order complement models. We combined ordinary differential equations with logical rules to produce a compact yet predictive model of complement activation. The model, which described the lectin and alternative pathways, was an order of magnitude smaller than comparable models in the literature. We estimated an ensemble of model parameters from in vitro dynamic measurements of the C3a and C5a complement proteins. Subsequently, we validated the model on unseen C3a and C5a measurements not used for model training. Despite its small size, the model was surprisingly predictive. Global sensitivity and robustness analysis suggested complement was robust to any single therapeutic intervention. Only the simultaneous knockdown of both C3 and C5 consistently reduced C3a and C5a formation from all pathways. Taken together, we developed a validated mathematical model of complement activation that was computationally inexpensive, and could easily be incorporated into pre-existing or new pharmacokinetic models of immune system function. The model described experimental data, and predicted the need for multiple points of therapeutic intervention to fully disrupt complement activation.

Keywords: Complement, systems biology, reduced order modeling, biochemical engineering

Introduction

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Complement is an important pathway in innate immunity. It plays a significant role in inflammation, host defense as well as many disease processes. Complement was discovered in the late 1880s where it was found to 'complement' the bactericidal activity of natural antibodies (1). However, research over the past decade has suggested the importance of complement extends beyond innate immunity. For example, complement contributes to tissue homeostasis (2). It has also has been linked with several diseases including Alzheimers, Parkinson's, multiple sclerosis, schizophrenia, rheumatoid arthritis and sepsis (3, 4). Complement also plays positive and negative roles in cancer; attacking tumor cells with altered surface proteins in some cases, while potentially contributing to tumor growth in others (5, 6). Lastly, several other important biochemical systems are integrated with complement including the coagulation cascade, the autonomous nervous system and inflammation (6). Thus, complement is important in a variety of beneficial 13 and potentially harmful functions in the body. Despite its importance, there have been few approved complement specific therapeutics, largely because of safety concerns and 15 challenging pharmacokinetic constraints, however, progress is being made (7). 16

The complement cascade involves many soluble and cell surface proteins, receptors and regulators (8, 9). The outputs of complement are the Membrane Attack Complex (MAC), and the inflammatory mediator proteins C3a and C5a. The membrane attack complex, generated during the terminal phase of the response, forms transmembrane channels which disrupt the membrane integrity of targeted cells, leading to cell lysis and death. On the other hand, the C3a and C5a proteins act as a bridge between innate and adaptive immunity, and play an important role in regulating inflammation (5). Complement activation takes places through three pathways: the classical, the lectin and the alternate pathways. The classical pathway is triggered by antibody recognition of foreign antigens or other pathogens. A multimeric protein complex C1 binds antibody-antigen

complexes and undergoes a conformational change, leading to an activated form with proteolytic activity. The activated C1-complex cleaves soluble complement proteins C4 and C2 into C4a, C4b, C2a and C2b, respectively. The C4a and C2b fragments bind to form the C4bC2a protease, also known as the classical pathway C3 convertase (CP C3 30 convertase). The lectin pathway is initiated through the binding of L-ficolin or Mannose 31 Binding Lectin (MBL) to carbohydrates on the surfaces of bacterial pathogens. These 32 complexes, in combination with mannose-associated serine proteases 1 and 2 (MASP-33 1/2), also cleave C4 and C2, leading to additional CP C3 convertase. Thus, the classical 34 and lectin pathways, initiated by different cues on foreign surfaces, converge at the CP C3 35 convertase. On the other hand, the alternate pathway is activated by a 'tickover' mechanism in which complement protein C3 is spontaneously hydrolyzed to form an activated 37 intermediate C3w; C3w recruits factor B and factor D, leading to the formation of C3wBb. 38 C3wBb cleaves C3 into C3a and C3b, where the C3b fragment further recruits additional factor B and factor D to form C3bBb, the alternate C3 convertase (AP C3 convertase) (10). The role of classical and alternate C3 convertases is varied. First, AP C3 conver-41 tases mediate signal amplification. AP C3 convertases cleave C3 into C3a and C3b; the C3b fragment is then free to form additional alternate C3 convertases, thereby forming a positive feedback loop. Next, AP/CP C3 convertases link complement initiation with the terminal phase of the cascade through the formation of C5 convertases. Both classical and alternate C3 convertases can recruit C3b subunits to form the classical pathway C5 convertase (C4bC2aC3b, CP C5 convertase), and the alternate pathway C5 convertase 47 (C3bBbC3b, AP C5 convertase), respectively. Both C5 convertases cleave C5 into the C5a and C5b fragments. The C5b fragment, along with the complement proteins C6, C7, C8 and multiple C9s, form the membrane attack complex. On the other hand, both 50 C3a and C5a are important inflammatory signals involved in several responses (8, 9). 51 Thus, the complement cascade attacks invading pathogens, while acting as a beacon for adaptive immunity.

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The complement cascade is regulated by plasma and host cell surface proteins which 54 balance host safety with effectiveness. The initiation of the classical pathway via complement protein C1 is controlled by the C1 Inhibitor (C1-Inh); C1-Inh irreversibly binds 56 to and deactivates the active subunits of C1, preventing chronic complement activation 57 (11). Regulation of upstream processes in the lectin and alternate pathways also oc-58 curs through the interaction of the C4 binding protein (C4BP) with C4b, and factor H with 59 C3b (12). Interestingly, both factor H and C4BP are capable of binding their respective 60 targets while in convertase complexes as well. At the host cell surface, membrane co-61 factor protein (MCP or CD46) can interact with C4b and C3b, which protects the host 62 cell from complement self-activation (13). Delay accelerating factor (DAF or CD55) also 63 recognizes and dissociates both C3 and C5 convertases on host cell surfaces (14). More 64 generally the well known inflammation regulator Carboxypeptidase-N has broad activity against the complement proteins C3a, C4a, and C5a, rendering them inactive by cleavage of carboxyl-terminal arginine and lysine residues (15). Although Carboxypeptidase-N 67 does not directly influence complement activation, it silences the important inflammatory signals produced by complement. Lastly, assembly of the MAC complex itself can be inhibited by vitronectin and clusterin in the plasma, and CD59 at the host surface (16, 17). Thus, there are many points of control which influence complement across the three activation pathways.

Developing quantitative mathematical models of complement could be crucial to fully understanding its role in the body. Traditionally, complement models have been formulated as systems of linear or non-linear ordinary differential equations (ODEs). For ex-75 ample, Hirayama et al., modeled the classical complement pathway as a system of linear ODEs (18), while Korotaevskiy and co-workers modeled the classical, lectin and alter-77 nate pathways as a system of non-linear ODEs (19). More recently, large mechanistic

models of sections of complement have also been proposed. For example, Liu et al., analyzed the formation of the classical and lectin C3 convertases, and the regulatory role of C4BP using a system of 45 non-linear ODEs with 85 parameters (20). Zewde and 81 co-workers constructed a detailed mechanistic model of the alternative pathway which 82 consisted of 107 ODEs and 74 kinetic parameters and delineated between the fluid, host 83 and pathogen surfaces (17). However, these previous studies involved large models with 84 little experimental validation. Thus, while these models are undoubtably important theo-85 retical tools, it is unclear if they can describe or quantitatively predict complement mea-86 surements. The central challenge of complement model identification is the estimation of 87 model parameters from experimental measurements. Unlike other important cascades, 88 such as coagulation where there are well developed experimental tools and publicly avail-89 able data sets, the data for complement is relatively sparse. Data sets with missing or 90 incomplete data, and limited dynamic data also make the identification of large mecha-91 nistic complement models difficult. Thus, reduced order approaches which describe the biology of complement using a limited number of species and parameters could be important for pharmacokinetic model development, and for our understanding of the varied role of complement in the body.

96 Results

In this study, we estimated an ensemble of experimentally validated reduced order com-97 plement models using multiobjective optimization. The modeling approach combined or-98 dinary differential equations with logical rules to produce a complement model with a 99 limited number of equations and parameters. The reduced order model, which described 100 the lectin and alternative pathways, consisted of 18 differential equations with 28 param-101 eters. Thus, the model was an order of magnitude smaller and included more pathways 102 than comparable models in the literature. We estimated an ensemble of model param-103 eters from in vitro time series measurements of the C3a and C5a complement proteins. Subsequently, we validated the model on unseen C3a and C5a measurements not used for model training. Despite its size, the model was surprisingly predictive. After validation, 106 we performed global sensitivity and robustness analysis to estimate which parameters 107 and species controlled model performance. Sensitivity analysis suggested CP C3 and C5 108 convertase parameters were critical, while robustness analyses suggested complement 109 was robust to any single therapeutic intervention; only the knockdown of both C3 and 110 C5 consistently reduced C3a and C5a formation for all cases. Taken together, we de-111 veloped a reduced order complement model that was computationally inexpensive, and 112 could easily be incorporated into pre-existing or new pharmacokinetic models of immune 113 system function. The model described experimental data, and predicted the need for 114 multiple points of intervention to disrupt complement activation. 115

Reduced order complement network. The complement model described the alternate and lectin pathways (Fig. 1). A trigger event initiated the lectin pathway (encoded as a logical rule), which activated the cleavage of C2 and C4 into C2a, C2b, C4a and C4b, respectively. Classical Pathway (CP) C3 convertase (C4aC2b) then catalyzed the cleavage of C3 into C3a and C3b. The alternate pathway was initiated through the spontaneous hydrolysis of C3 into C3a and C3b. The C3b fragments generated by hydrolysis (or by CP)

C3 convertase) could then form the alternate pathway (AP) C3 convertase (C3bBb). We did not consider C3w, nor the formation of the initial alternate C3 convertase (C3wBb). Rather, we assumed C3w was equivalent to C3b and only modeled the formation of the 124 main AP C3 convertase. Both the CP and AP C3 convertases catalyzed the cleavage of 125 C3 into C3a and C3b. A second C3b fragment could then bind with either the CP or AP 126 C3 convertase to form the CP or AP C5 convertase (C4bC2aC3b or C3bBbC3b). Both C5 127 convertases catalyzed the cleavage of C5 into the C5a and C5b fragments. In this study, 128 we simplified the model by assuming both factor B and factor D were in excess. However, 129 we did explicitly account for the action of two other control proteins, factor H and C4BP. 130 Lastly, we did not consider MAC formation, instead we stopped at C5a and C5b. Lectin 131 pathway activation, and C3/C5 convertase activity were modeled using a combination of 132 saturation kinetics and non-linear transfer functions, which resulted in a significant size re-133 duction of the model, while maintaining performance. Binding interactions were modeled 134 using mass-action kinetics, where we assumed all binding was irreversible. Thus, while 135 the reduced order complement model encoded significant biology, it was highly compact 136 consisting of only 18 differential equations and 28 model parameters. Next, we estimated 137 an ensemble of model parameters from time series measurements of the C3a and C5a complement proteins.

for the development of any dynamic model is the estimation of model parameters. We estimated an ensemble of complement model parameters using *in vitro* time-series data sets generated with and without zymosan, a lectin pathway activator (21). The residual between model simulations and experimental measurements was minimized using the Pareto Optimal Ensemble Technique (JuPOETs) [REFs] starting from a initial guess generated by the dynamic optimization with particle swarms (DOPS) routine [REF]. Unless otherwise specified, all initial conditions were assumed to be at their mean physiological

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values. While we had significant training data, the parameter estimation problem was underdetermined (we were not able to uniquely determine model parameters). Thus, instead of using the best-fit yet uncertain parameter set, we estimated an ensemble of probable parameter sets to quantify model uncertainty (N = 2100, see materials and methods). The complement model ensemble captured the behavior of both the alternate and lectin pathways (Fig. 2). To estimate alternate pathway model parameters, we used C3a and C5a measurements in the absence of zymosan (Fig. 2A and B). On the other hand, lectin pathway parameters were estimated from C3a and C5a measurements in the presence of 1mg/ml zymosan (Fig. 2C and D). The reduced order model reproduced a panel of alternate and lectin pathway data sets in the neighborhood of physiological factor and inhibitor concentrations. However, it was unclear whether the reduced order model could predict new data, without updating the model parameters. To address this question, we fixed the model parameters and simulated data sets not used for model training.

We tested the predictive power of the reduced order complement model with data not used during model training (Fig. 3). Six validation cases were considered, three for C3a and C5a, respectively. All model parameters and initial conditions were fixed for the validation simulations (with the exception of zymosan, and other experimentally mandated changes). The ensemble of reduced order models predicted the qualitative dynamics of C3a formation (Fig. 3, top), and C5a formation (Fig. 3, bottom) at three inducer concentrations. The rate of C3a formation and C3a peak time were directly proportional to initiator dose. Similarly, the C5a plateau and rate of formation were also directly proportional to initiator dose, with the lag time being indirectly proportional to initiator exposure for both C3a and C5a. However, there were shortcomings with model performance. First, while the overall C3a trend was captured (within the 99% confidence interval), the C3a dynamics were too fast with the exception of the low dose case. We believe the C3a time scale was related to our choice of training data, how we modeled the tickover mechanism, and

factor B and D limitation. We trained the model using either no or 1 mg/ml zymosan, but predicted cases in a different initiator range; comparing training to prediction, the model 175 performance e.g., the shape of the C3a trajectory was biased towards either high or very 176 low initiator doses. Next, tickover was modeled as a first-order generation processes 177 where C3wBb formation and activity was lumped into the AP C3 convertase. Thus, we 178 skipped an important upstream step which could influence AP C3 convertase formation 179 by attenuating the rate C3 cleavage into C3a and C3b. We also assumed both factor B 180 and factor D were not limiting, thereby artificially accelerating the rate of AP C3 conver-181 tase formation. The C5a predictions followed a similar trend as C3a; we captured the 182 long-time C5a behavior but over predicted the time scale of C5 cleavage. However, be-183 cause the C5a time scale depends strongly upon C3 convertase formation, we can likely 184 correct the C5 issues by fixing the rate of C3 cleavage. Despite these shortcomings, we 185 qualitatively predicted experimental measurements not used for model training typically 186 within the 99% confidence of the ensemble, for three inducer levels. Next, we used global 187 sensitivity and robustness analysis to determine which parameters and species controlled 188 the performance of the complement model. 189

Global analysis of the reduced order complement model. We conducted sensitivity analysis to estimate which parameters controlled the performance of the reduced order complement model. We calculated the total sensitivity of the C3a and C5a residual to changes in model parameters with and without zymosan (Fig. 4). In the absence of zymosan (where only the alternative pathway is active), the most sensitive parameter was the rate constant governing the assembly of the AP C3 convertase, as well as the rate constant controlling basal C3b formation via the tickover mechanism. The C5a trajectory was sensitive to the AP C5 convertase kinetic parameters (Fig. 4A). Interestingly, neither the rate nor the saturation constant governing AP C3 convertase activity were sensitive in the absence of zymosan. Thus, C3a formation in the alternative pathway was more heav-

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ily influenced by the spontaneous hydrolysis of C3, rather than AP C3 convertase activity, in the absence of zymosan. In the presence of zymosan, the C3a residual was controlled by the formation and activity of the CP C3 convertase, as well as tickover and degradation parameters. On the other hand, the C5a residual was controlled by the formation and activity of CP C5 convertase, and tickover C3b formation in the presence of zymosan (Fig. 204 4B). The lectin initiation parameters were sensitive, but to a lesser extent than CP conver-205 tase kinetic parameters and tickover C3b formation. Thus, sensitivity analysis suggested 206 that CP C3/C5 convertase formation and activity dominated in the presence of zymosan, but tickover parameters and AP C5 convertase were more important without initiator. AP 208 C3 convertase assembly was important, but its activity was not. Next, we compared the 209 sensitivity results to current therapeutic approaches; pathways involving sensitive param-210 eters have been targeted for clinical intervention (Fig. 4C). In particular, the sensitivity analysis suggested AP/CP C5 convertase inhibitors, or interventions aimed at attenuat-212 ing C3 or C5 would most strongly influence complement performance. Thus, there was 213 at least a qualitative overlap between sensitivity and the potential of biochemical efficacy. 214 However, sensitivity coefficients quantify how changes in parameters e.g., rate or satura-215 tion constants affect model performance. To more closely simulate a clinical intervention 216 e.g., administration of an anti-complement inhibitor, we performed robustness analysis.

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Robustness analysis suggested there was no single intervention that inhibited complement activation in the presence of both initiation pathways (Fig. 5). Robustness coefficients quantify the response of a protein to a macroscopic structural or operational perturbation to a biochemical network. Here, we computed how the C3a and C5a trajectories responded to a decrease in the initial abundance of C3 and/or C5 with and without lectin initiator. We simulated the addition of different doses of anti-complement inhibitor cocktails by decreasing the initial concentration of C3, C5 or the combination of C3 and C5 by 50%, 90% and 99%. This would be conceptually analogous to the administration

of a C3 inhibitor e.g., Compstatin alone or combination with Eculizumab (Fig. 4C). The response of the complement model to different knock-down magnitudes was non-linear; a 227 90% knock-down had an order of magnitude more impact than a 50% knock-down. As ex-228 pected, a C5 knockdown had no effect on C3a formation for either the alternate (Fig. 5A) 229 or lectin pathways (Fig. 5B). However, C3a and to a greater extent C5a abundance de-230 creased with decreasing C3 concentration in the alternate pathway (Fig. 5A). This agreed 231 with the sensitivity results; changes in AP C3-convertase formation affected the down-232 stream dynamics of C5a formation. Thus, if we only considered the alternate pathway, C3 233 alone could be a reasonable target, especially given that C5a formation was surprisingly 234 robust to C5 levels in the alternate pathway. Yet, when both pathways were activated, 235 C5a levels were robust to the initial C3 concentration (Fig. 5B); even 1% of the nominal 236 C3 was able to generate enough AP/CP C5 convertase to maintain C5a formation. Thus, 237 the only reliable intervention that consistently reduced both C3a and C5a formation for all 238 cases was a knockdown of both C3 and C5. For example, a 90% decrease of both C3 239 and C5 reduced the formation of C5a by an order of magnitude, while C3a was reduced 240 to a lesser extent (Fig. 5B).

Discussion

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In this study, we developed an ensemble of experimentally validated reduced order com-243 plement models using multiobjective optimization. The modeling approach combined or-244 dinary differential equations with logical rules to produce a complement model with a lim-245 ited number of equations and parameters. The reduced order model, which described the 246 lectin and alternative pathways, consisted of 18 differential equations with 28 parameters. 247 Thus, the model was an order of magnitude smaller and included more pathways than 248 comparable mathematical models in the literature. We estimated an ensemble of model 249 parameters from in vitro time series measurements of the C3a and C5a complement proteins. Subsequently, we validated the model on unseen C3a and C5a measurements that were not used for model training. Despite its small size, the model was surprisingly predic-252 tive. After validation, we performed global sensitivity and robustness analysis to estimate 253 which parameters and species controlled model performance. These analyses suggested complement was robust to any single therapeutic intervention. The only intervention that 255 consistently reduced C3a and C5a formation for all cases was a knockdown of both C3 256 and C5. Taken together, we developed a reduced order complement model that was 257 computationally inexpensive, and could easily be incorporated into pre-existing or new 258 pharmacokinetic models of immune system function. The model described experimen-259 tal data, and predicted the need for multiple points of intervention to disrupt complement 260 activation. 261

Despite its importance, there has been a paucity of validated mathematical models of complement pathway activation. To our knowledge, this study is one of the first complement models that combined multiple initiation pathways with experimental validation of important complement products like C5a. However, there have been several theoretical models of components of the cascade in the literature. Liu and co-workers modeled the formation of C3a through the classical pathway using 45 non-linear ODEs (20). In

contrast, in this study we modeled lectin mediated C3a formation using only five ODEs. Though we did not model all the initiation interactions in detail, especially the cross-talk between the lectin and classical pathways, we successfully captured C3a dynamics with respect to different concentrations of lectin initiators. The model also captured the dynamics of C3a and C5a formed from the alternate pathway using only seven ODEs. The reduced order model predictions of C5a were qualitatively similar to the theoretical complement model of Zewde et al., which involved over 100 ODEs (17). However, we found that the C3a produced in the alternate pathway was nearly three orders of magnitude greater than the C5a generated. While this was in agreement with the experimental data (21), it differed from the theoretical predictions made by Zewde et al., who showed C3a was eight orders of magnitude greater than the C5a concentration (17). In our model, the time profile of both C3a and C5a generated changed with respect to the quantity of zymosan (the lectin pathway initiator). In particular, the C3a peak time was directly proportional to initiator, while the lag phase for generation was inversely proportional to the initiator concentration. Korotaevskiy et al. showed a similar trend using a theoretical model of complement, albeit for much shorter time scales (19). Thus, the reduced order complement model performed at least as well as existing larger mechanistic models, despite being significantly smaller.

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Global analysis of the complement model suggested potentially important therapeutic targets. Complement malfunctions are implicated in a spectrum of diseases, however the development of complement specific therapeutics has been challenging (3, 22). Previously, we have shown that mathematical modeling and analysis can be useful tools to estimate therapeutically important mechanisms (23–26). In this study, we analyzed a validated ensemble of reduced order complement models to better understand the strengths and weaknesses of the cascade. In the presence of an initiator, C3a and C5a formation was sensitive to CP C3/C5 convertase assembly and activity, and to a lesser extent

lectin initiation parameters. Formation of the CP convertases can be inhibited by targeting upstream protease complexes like MASP-1,2 from the lectin pathway (or C1r, C1s from 295 classical pathway). For example, Omeros, a protease inhibitor that targets the MASP-2 296 complex, has been shown to inhibit the formation of downstream convertases (27). Lam-297 palizumab and Bikaciomab, which target factor B and factor D respectively, or naturally 298 occurring proteins such as Cobra Venom Factor (CVF), an analogue of C3b, could also 299 attenuate AP convertase formation (28-30). Removing supporting molecules could also 300 destabilize the convertases. For example, Novelmed Therapeutics developed the anti-301 body, NM9401 against propedin, a small protein that stabilizes alternate C3 convertase 302 (31). Lastly, convertase catalytic activity could be attenuated using small molecule pro-303 tease inhibitors. All of these approaches are consistent with the results of the sensitivity 304 analysis. On the other hand, robustness analysis suggested C3a and C5a generation 305 could only be significantly attenuated by modulating the free levels of C3 and C5. The 306 most commonly used anti-complement drug Eculizumab, targets the C5 protein (22). Sev-307 eral other antibodies targeting C5 are also being developed; for example, LFG316 targets 308 C5 in Age-Related Macular Degeneration (32), while Mubodina is used to treat Atypical 309 Hemolytic-Uremic Syndrome (aHUS) (33). Other agents such as Coversin (34) or the aptamer Zimura (35) could also be used to knockdown C5. The peptide inhibitor Compstatin and its derivatives are promising approaches for the inhibition of C3 (36). However, while the knockdown of C3 and C5 affect C3a and C5a levels downstream, the abundance, turnover rate and population variation of these proteins make them difficult targets 314 (37, 38). For example, the eculizumab dosage must be significantly adjusted during the 315 course of treatment for aHUS (39). A validated complement model, in combination with 316 personalized pharmacokinetic models of immune system function, could be an important 317 development for the field. 318

The performance of the complement model was impressive given its limited size. How-

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ever, there are several questions that should be explored further. A logical progression for this work would be to expand the network to include the classical pathway and the 321 formation of the membrane attack complex (MAC). However, time course measurements 322 of MAC abundance (and MAC formation dynamics) are scarce, making the inclusion of 323 MAC challenging. On the other hand, inclusion of classical pathway activation is straight-324 forward. Liu et al., have shown cross-talk between the activation of the classical and lectin 325 pathways through C reactive proteins (CRP) and L-ficolin (LF) under inflammation condi-326 tions (20). Thus, inclusion of these species, in addition to a lumped activation term for the 327 classical pathway should allow us to capture classical activation. Next, we should address 328 the C3a time scale issue. We believe the C3a time scale was related to our choice of train-329 ing data, how we modeled the tickover mechanism, and factor B and D limitation. Tickover 330 was modeled as a first-order generation processes where C3wBb formation and activity 331 was lumped into the AP C3 convertase. Thus, we skipped an important step which could 332 strongly influence AP C3 convertase formation by slowing down the rate C3 cleavage 333 into C3a and C3b. The model should be expanded to include the C3wBb intermediate, 334 where C3wBb catalyzes C3 cleavage at a slow rate compared to normal AP or CP C3 335 convertases. We also assumed both factor B and factor D were not limiting, thereby artificially accelerating the rate of AP C3 convertase formation. This shortcoming could be 337 addressed by including balances around factor B and D, and including these species in the appropriate kinetic rates. The C5a predictions also had an accelerated time scale. 339 However, because the C5a time scale depended strongly upon C3 convertase formation, 340 we can likely correct the C5 issues by fixing the rate of C3 cleavage. Lastly, we should 341 also consider including the C2-bypass pathway, which was not included in the model. 342 The C2-bypass mediates lectin pathway activation, without the involvement of MASP-1/2. 343 Thus, this pathway could be important for understanding the role of MASP-1/2 inhibitors 344 on complement activation.

Materials and Methods

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

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

where \mathcal{R} denotes the number of reactions and \mathcal{M} denotes the number of proteins in 350 the model. The quantity τ_i denotes a time scale parameter for species i which captures 351 unmodeled effects. For the current study, τ scaled with the level of initiator (z) for C5a 352 and C5b; $\tau_i=z/z^*$ for i = C5a, C5b where z^* was 1mg/ml, τ_i = 1 for all other species. The quantity $r_j(\mathbf{x}, \epsilon, \mathbf{k})$ denotes the rate of reaction j. Typically, reaction j is a non-linear function of biochemical and enzyme species abundance, as well as unknown model parameters \mathbf{k} ($\mathcal{K} \times 1$). The quantity σ_{ij} denotes the stoichiometric coefficient for species i in reaction j. If $\sigma_{ij} > 0$, species i is produced by reaction j. Conversely, if $\sigma_{ij} < 0$, species iis consumed by reaction j, while $\sigma_{ij}=0$ indicates species i is not connected with reaction j. Species balances were subject to the initial conditions $\mathbf{x}(t_o) = \mathbf{x}_o$. 359 Rate processes were written as the product of a kinetic term (\bar{r}_i) and a control term 360 (v_i) in the complement model. The kinetic term for the formation of C4a, C4b, C2a and 361 C2b, lectin pathway activation, and C3 and C5 convertase activity was given by: 362

$$\bar{r}_j = k_j^{max} \epsilon_i \left(\frac{x_s^{\eta}}{K_{js}^{\eta} + x_s^{\eta}} \right) \tag{2}$$

where k_j^{max} denotes the maximum rate for reaction j, ϵ_i denotes the abundance of the enzyme catalyzing reaction j, η denotes a cooperativity parameter, and K_{js} denotes the saturation constant for species s in reaction j. We used mass action kinetics to model

protein-protein binding interactions within the network:

$$\bar{r}_j = k_j^{max} \prod_{s \in m_j^-} x_s^{-\sigma_{sj}} \tag{3}$$

where k_j^{max} denotes the maximum rate for reaction j, σ_{sj} denotes the stoichiometric coefficient for species s in reaction j, and $s \in m_j$ denotes the set of *reactants* for reaction j. We assumed all binding interactions were irreversible.

The control terms $0 \le v_j \le 1$ depended upon the combination of factors which influenced rate process j. For each rate, we used a rule-based approach to select from
competing control factors. If rate j was influenced by $1, \ldots, m$ factors, we modeled this relationship as $v_j = \mathcal{I}_j \left(f_{1j} \left(\cdot \right), \ldots, f_{mj} \left(\cdot \right) \right)$ where $0 \le f_{ij} \left(\cdot \right) \le 1$ denotes a regulatory transfer
function quantifying the influence of factor i on rate j. The function $\mathcal{I}_j \left(\cdot \right)$ is an integration
rule which maps the output of regulatory transfer functions into a control variable. Each
regulatory transfer function took the form:

$$f_{ij}\left(\mathcal{Z}_{i}, k_{ij}, \eta_{ij}\right) = k_{ij}^{\eta_{ij}} \mathcal{Z}_{i}^{\eta_{ij}} / \left(1 + k_{ij}^{\eta_{ij}} \mathcal{Z}_{i}^{\eta_{ij}}\right) \tag{4}$$

where \mathcal{Z}_i denotes the abundance of factor i, k_{ij} denotes a gain parameter, and η_{ij} denotes a cooperativity parameter. In this study, we used $\mathcal{I}_j \in \{min, max\}$ (40). If a process has no modifying factors, $v_j = 1$. The model equations were implemented in Julia and solved using the CVODE routine of the Sundials package (41, 42). The model code and parameter ensemble is freely available under an MIT software license and can be downloaded from http://www.varnerlab.org.

Estimating an ensemble of complement model parameters. We estimated a single initial parameter set using the Dynamic Optimization with Particle Swarms (DOPS) technique (43). DOPS is a novel hybrid meta-heuristic which combines a multi-swarm particle

swarm method with the dynamically dimensioned search approach of Shoemaker and colleagues (44). DOPS minimized the squared residual between simulated and C3a and 387 C5a measurements with and without zymosan as a single objective. The best fit set esti-388 mated by DOPS served as the starting point for multiobjective ensemble generation using 389 Pareto Optimal Ensemble Technique in the Julia programming language (JuPOETs) (45). 390 JuPOETs is a multiobjective approach which integrates simulated annealing with Pareto 391 optimality to estimate model ensembles on or near the optimal tradeoff surface between 392 competing training objectives. JuPOETs minimized training objectives of the form: 393

$$O_j(\mathbf{k}) = \sum_{i=1}^{T_j} \left(\hat{\mathcal{M}}_{ij} - \hat{y}_{ij}(\mathbf{k}) \right)^2 + \left(\frac{\mathcal{M}'_{ij} - \max y_{ij}}{\mathcal{M}'_{ij}} \right)^2$$
 (5)

subject to the model equations, initial conditions and parameter bounds $\mathcal{L} \leq \mathbf{k} \leq \mathcal{U}.$ The 394 first term in the objective function measured the shape difference between the simulations and measurements. The symbol $\hat{\mathcal{M}}_{ij}$ denotes a scaled experimental observation (from training set j) while the symbol \hat{y}_{ij} denotes the scaled simulation output (from training set j). The quantity i denotes the sampled time-index and \mathcal{T}_j denotes the number of time points for experiment *j*. The scaled measurement is given by:

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$$\hat{\mathcal{M}}_{ij} = \frac{\mathcal{M}_{ij} - \min_{i} \mathcal{M}_{ij}}{\max_{i} \mathcal{M}_{ij} - \min_{i} \mathcal{M}_{ij}}$$
(6)

Under this scaling, the lowest measured concentration become zero while the highest equaled one, where a similar scaling was defined for the simulation output. The secondterm in the objective function quantified the absolute error in the estimated concentration scale, where the absolute measured concentration (denoted by \mathcal{M}'_{ij}) was compared with the largest simulated value. In this study, we minimized two training objectives, the total C3a and C5a residual w/o zymosan (O₁) and the total C3a and C5a residual for 1 mg/ml zymosan (O_2). JuPOETs identified an ensemble of N \simeq 2100 parameter sets which were used for model simulations and uncertainty quantification subsequently. JuPOETs is open source, available under an MIT software license. The JuPOETs source code is freely available from the JuPOETs GitHub repository at https://github.com/varnerlab/POETs.jl. The objective functions used in this study are available in the GitHub model repository available from http://varnerlab.org.

Sensitivity and robustness analysis. We conducted global sensitivity and robustness analysis to estimate which parameters and species controlled the performance of the reduced order model. We computed the total variance-based sensitivity index of each parameter relative to the training residual for the C3a/C5a alternate and C3a/C5a lectin objectives using the Sobol method (46). sThe sampling bounds for each parameter were established from the minimum and maximum value for that parameter in the parameter ensemble. We used the sampling method of Saltelli *et al.* to compute a family of N (2d + 2) parameter sets which obeyed our parameter ranges, where N was the number of trials per parameters, and d was the number of parameters in the model (47). In our case, N = 400 and d = 28, so the total sensitivity indices were computed using 23,200 model evaluations. The variance-based sensitivity analysis was conducted using the SALib module encoded in the Python programming language (48).

Robustness coefficients quantify the response of a marker to a structural or operational perturbation to the network architecture. Robustness coefficients were calculated as shown previously (49). Log-transformed robustness coefficients denoted by $\hat{\alpha}$ (i, j, t_o, t_f) were defined as:

$$\hat{\alpha}\left(i, j, t_o, t_f\right) = \log_{10}\left[\left(\int_{t_o}^{t_f} x_i\left(t\right) dt\right)^{-1} \left(\int_{t_o}^{t_f} x_i^{(j)}\left(t\right) dt\right)\right] \tag{7}$$

Here, t_o and t_f denote the initial and final simulation time, while i and j denote the indices

for the marker and the perturbation, respectively. A value of $\hat{\alpha}(i,j,t_o,t_f)>0$, indicates increased marker abundance, while $\hat{\alpha}(i,j,t_o,t_f)<0$ indicates decreased marker abundance following perturbation j. If $\hat{\alpha}(i,j,t_o,t_f)\sim0$, perturbation j did not influence the abundance of marker i. In this study, we perturbed the initial condition of C3 or C5 or a combination of C3 and C5 by 50%, 90% and 99% and measured the area under the curve (AUC) of C3a or C5a with and without lectin initiator. We computed the robustness coefficients for a subset of the parameter ensemble (N = 65) and reported the mean robustness value.

437 Competing interests

The authors declare that they have no competing interests.

Author's contributions

J.V directed the study. A.S developed the reduced order complement model and the parameter ensemble. A.S, W.D and M.M analyzed the model ensemble, and generated figures for the manuscript. The manuscript was prepared and edited for publication by A.S, W.D, M.M and J.V.

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Fig. 1: Simplified schematic of the human complement system. The complement cascade is activated through three pathways: the classical, the lectin, and the alternate pathways. Complement initiation results in the formation of classical or alternative C3 convertases, which amplify the initial complement response and signal to the adaptive immune system by cleaving C3 into C3a and C3b. C3 convertases further react to form C5 convertases which catalyze the cleavage of the C5 complement protein to C5a and C5b. C5b is critical to the formation of the membrane attack complex (MAC), while C5a recruits an adaptive immune response.

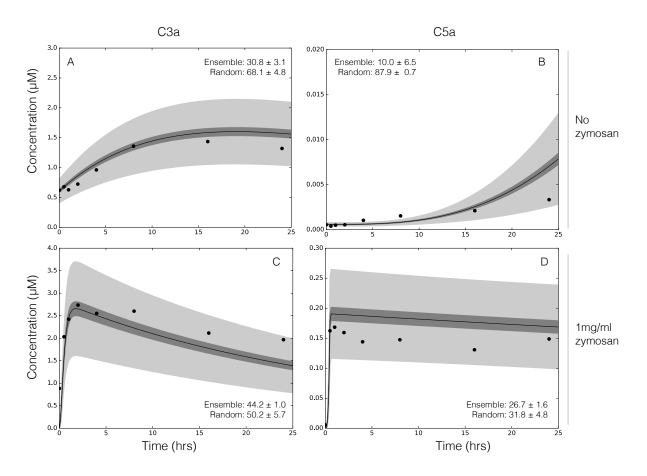


Fig. 2: Reduced order complement model training. An ensemble of model parameters were estimated using multiobjective optimization from C3a and C5a measurements with and without zymosan (21). The model was trained using C3a and C5a data generated from the alternative pathway (**A–B**) and lectin pathway initiated with 1 mg/ml zymosan (**C–D**). The solid black lines show the simulated mean value of C3a or C5a for the ensemble, while the dark shaded region denotes the 99% confidence interval of mean. The light shaded region denotes the 99% confidence interval of the simulated C3a and C5a concentration. All initial conditions were assumed to be at their physiological serum levels unless otherwise noted.

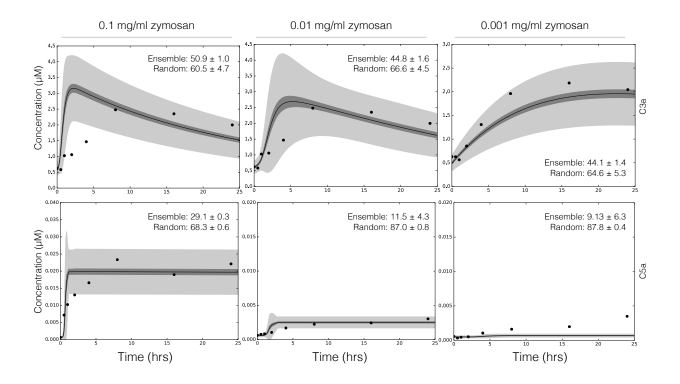


Fig. 3: Reduced order complement model predictions. Simulations of C3a and C5a generated in the lectin pathway using 0.1 mg/ml, 0.01 mg/ml, and 0.001 mg/ml zymosan were compared with the corresponding experimental measurements. The solid black lines show the simulated mean value of C3a or C5a for the ensemble, while the dark shaded region denotes the 99% confidence interval of mean. The light shaded region denotes the 99% confidence interval of the simulated C3a and C5a concentration. All initial conditions were assumed to be at their physiological serum levels unless otherwise noted.

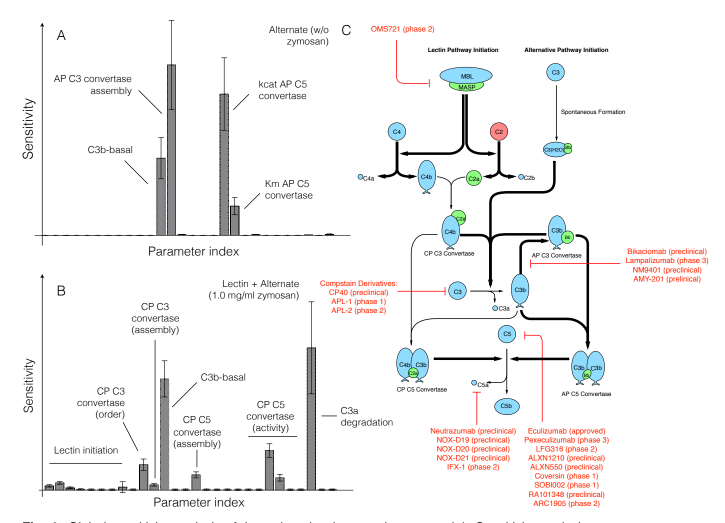


Fig. 4: Global sensitivity analysis of the reduced order complement model. Sensitivity analysis was conducted on the two objectives used for model training. **A:** Sensitivity of the C3a and C5a residual w/o zymosan. **B:** Sensitivity of the C3a and C5a residual with 1 mg/ml zymosan. The bars denote the mean total sensitivity index for each parameter, while the error bars denote the 95% confidence interval. **C:** Pathways controlled by the sensitivity parameters. Bold black lines indicate the pathway involves one or more sensitive parameters, while the red lines show current therapeutics targets. Current complement therapeutics were taken from the review of Morgan and Harris (22).

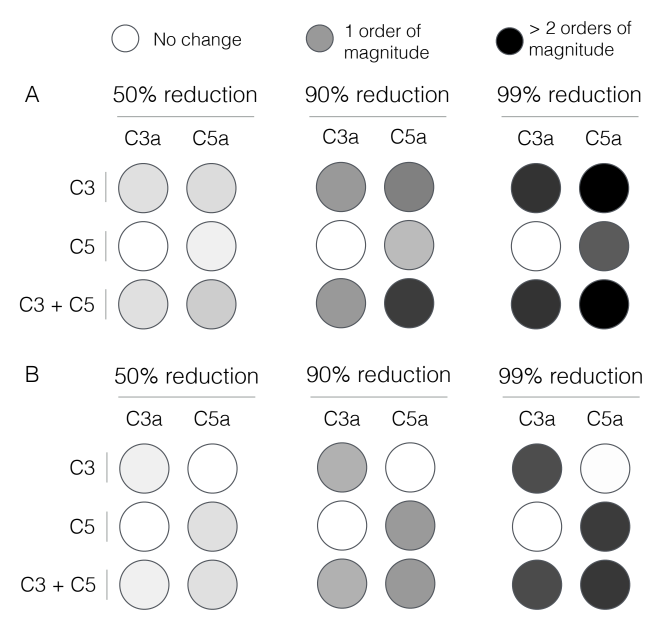


Fig. 5: Robustness analysis of the complement model. Robustness coefficients were calculated for a 50%, 90% and 99% reduction in C3, C5, or C3 and C5 initial conditions. **A:** Mean robustness index for C3a and C5a generated from the alternate pathway (w/o zymosan). **B:** Mean robustness index for C3a and C5a generated from the lectin and alternate pathway (1 mg/ml zymosan). The color describes the degree of reduction of C3a or C5a following the network perturbation. Robustness coefficients were calculated using all parameter sets with Pareto rank less than five (N = 65).