Understanding the human innate immune system

*In-silico studies*

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Supplemented alkaline phosphatase supports the immune response in patients undergoing cardiac surgery: clinical and computational evidence

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Modeling the Human Innate Immune Response for Systemic Inflammation

Abstract

Alkaline phosphatase (AP) is an enzyme that exhibits anti-inflammatory effects by dephosphorylating inflammation triggering moieties (ITMs) like bacterial lipopolysaccharides and extracellular nucleotides. AP administration aims to prevent and treat peri- and post-surgical ischemia reperfusion injury in cardiothoracic surgery patients. Recent studies reported that intravenous bolus administration and continuous infusion of AP in patients undergoing coronary artery bypass grafting with cardiac valve surgery induce an increased release of liver-type “tissue non-specific alkaline phosphatase” (TNAP) into the bloodstream. The release of liver-type TNAP into circulation could be the body’s way of strengthening its defense against a massive ischemic insult. However, the underlying mechanism behind the induction of TNAP is still unclear. To obtain a deeper insight into the role of AP during surgery, we developed a mathematical model of systemic inflammation that clarifies the relation between supplemented AP and TNAP.
and describes a plausible induction mechanism of TNAP in patients undergoing cardiothoracic surgery. The model was validated against clinical data from patients treated with bovine Intestinal AP (bIAP treatment) or without AP (placebo treatment), in addition to standard care procedures. We performed additional in-silico experiments adding a secondary source of ITMs after surgery, as observed in some patients with complications, and predicted the response to different AP treatment regimens. Our results show a strong protective effect of supplemented AP for patients with complications. The model provides evidence of the existence of an induction mechanism of liver-type tissue non-specific alkaline phosphatase, triggered by the supplementation of AP in patients undergoing cardiac surgery. To the best of our knowledge this is the first time that a quantitative and validated numerical model of systemic inflammation under clinical treatment conditions is presented.

**Keywords:** alkaline phosphatase, innate immune response, cardiac surgery, ODE model, in-silico, clinical trial
1.1. Introduction

Alkaline Phosphatase (AP) is an enzyme originally known for its pivotal role in skeletal mineralization but also for its capability to reduce inflammation. AP is in fact capable to reduce inflammation in animals by dephosphorylating inflammation triggering moieties like bacterial lipopolysaccharides (LPS) and extracellular nucleotides. In addition, several studies demonstrated that AP has a key function in maintenance and restoration of physiological barriers in addition to this anti-inflammatory role of AP. In fact, many, if not all of these barriers may become hyper-permeable and or dysfunctional during such systemic ischemic and inflammation triggering insult. Extracellular nucleotides, like Adenosine Triphosphate (ATP) and Adenosine Diphosphate (ADP), having pivotal energy housekeeping functions, intracellularly act as ITM as soon they have leaked out of cells exposed to ischemic insults. LPS, a major component of the Gram negative bacterial outer membrane that is responsible for mediating septic shock. These inflammation triggering moieties (ITMs) are pro-inflammatory signals that may start local and systemic inflammatory responses in the innate immune system. Clinical trials involving the parenteral administration of AP to patients with severe sepsis, showed significant improvement in renal function.

Humans have four distinct AP isozymes: tissue-nonspecific AP (liver/bone/kidney type AP), which is the most predominant circulating form of isozyme, intestinal-, placental-type, and germ cell AP. The anti-inflammatory effects of AP have been confirmed in settings with intestinal-, placental- and liver-type AP.

Coronary artery bypass grafting (CABG) is one of the most common types of open-heart surgery which very often triggers a systemic inflammatory response, the clinical impact thereof is specific for the specific patient and depends on multiple factors like age, underlying diseases and other confounding factors. The average annual number of CABG procedures in Western practice is about 62.2 per 100 000, ranging from 29.3 procedures...
in Spain to 135.4 procedures in Belgium. According to the Society of Thoracic Surgeons National Database, CABG-mediated complications contribute to 1.8% in-hospital and 2.2% operative mortalities, but caused 24% post-operative atrial fibrillation incidences in 151,474 patients in 2015. We focus on the experiments by Kats et al., where we assume a systemic insult due to the amounts of ITMs introduced and generated peri- and post-cardiac surgery. Cardiac surgery invokes a vigorous systemic inflammatory response where massive amounts of ITMs are simultaneously generated from various sources in the body: a) CABG and valve surgery under CPB (Cardio Pulmonary Bypass) induces sheer stress on blood cells damaging them and releasing a massive amount of ITMs in the process, b) surgical area where tissue is damaged locally, and c) reperfusion damage by accumulated ITMs that have crossed the gut barrier during hypo-perfusion and become systemically available upon re-circulation. The body thereby deals with a massive amount of ITMs that enter the circulation and are transported into the tissue via blood flow, and circulate in the blood stream due to the effects of cardio pulmonary bypass grafting and reperfusion injury.

*De novo* synthesis and release in circulation of AP induced by AP prophylaxis could be the body’s way to improve its defense mechanism. A study in 2012 by Kats et al. demonstrated that intravenous bolus administration and continuous infusion of bovine intestinal Alkaline Phosphatase (bIAP, bRESCAP, and APPIRED studies by Alloksys Life Sciences), in patients undergoing CABG (with or without valve surgery) results in the release of endogenous tissue non-specific AP (TNAP), most likely liver-type AP. This release exhibits a *unique* feat that was not before observed in septic shock patients. Induction of liver-type non-specific AP supports the idea that AP contributes significantly to the immune response. Additional Phase III clinical trials are currently on the way to confirm the beneficial effects of AP previously reported in CABG and valve surgery.

If indeed excess AP or the release of additional liver-type TNAP is beneficial to the clinical outcome of patients undergoing major surgeries as well as for
individuals suffering from acute and chronic inflammation, then there is an urgent need to develop computational models that can reproduce and predict the dynamics of induced TNAP in circulation. The developed model could then pave way to better understand when and more importantly how much of this liver type TNAP is expressed and released back into circulation through in-silico experiments. We develop a new model of systemic inflammation based on existing models of the innate immune response to acute inflammation, with the purpose of describing, and gaining further insight on the dynamics of the innate immune system response through in-silico experiments. We thereby report, to the best of our knowledge, the first calibrated and validated mathematical model for systemic inflammation.

The human innate immune system (HIIS) is the body's first line of defense to an infection or trauma. This is commonly manifested in the form of acute inflammatory response, resulting from these and other oxidative stress conditions. Numerous studies were reported aiming to understand the acute inflammatory response based on the response of a single population of white blood cells to invading pathogens. Unlike in previous models that only deal with a general population of invading and invaded entities, Dunster distinguished between populations of white blood cells by incorporating activated macrophages, activated, apoptotic and necrotic neutrophil populations. More specific mathematical models of HIIS that further distinguish white blood cells into distinct populations have also been developed. For instance, Su et al. used a system of partial differential equations (PDE) that capture the spatial and temporal dynamics of the innate and adaptive immune response via the following stages: recognition, initiation, effector response and resolution of infection. This model of the human innate immune response was adapted by Pigozzo et al., who focused on the dynamics of LPS, neutrophils, pro-inflammatory and anti-inflammatory cytokines.

This paper focuses on how the concentrations of the innate immune response components evolve over time. Partial differential equations (PDEs)
provide ways to analyse both time and spatial dynamics of key aspects of HIIS. Since we are modelling a systemic insult, where a massive amount of ITMs are coming from various sources in the body and inflammation is not confined to a specific tissue or organ, we can assume that these moieties are present and distributed all throughout the organism. Thus, we regard the "tissue" as representative of the entire body. Given this assumption, the role of microscopic spatial effects for the dynamics of the system is negligible and we use ordinary differential equations (ODEs) to describe the dynamics of the immune response. However, we take into account spatial effects by modelling various compartments (liver, blood stream and tissue) and the transport of cells and molecules between them due to the inflammation in blood and tissue. This compartmentalization of the organism allows us to account for chemotaxis at a macroscopic level using the change in permeability of the endothelium during the different stages of inflammation to affect the transport of immune cells and molecules between blood and tissue.

We therefore construct the HIIS model from 46, 49, and 50 by introducing the following key differences: compartmentalization of the organism into liver, blood and tissue; introduction of the dual pathway to neutrophils death, necrosis being pro-inflammatory and apoptosis being anti-inflammatory; introduction of the anti-inflammatory action of AP and of the mechanism of AP induction; the dilution of cellular components in tissue typical of systemic inflammatory responses as opposed to the increased concentration of cellular components in a localized region typical of acute inflammatory responses.

1.2. **Clinical Trial Data**

Patients undergoing open-heart surgery were stratified according to a risk assessment score system called EuroSCORE (Type 1). EuroSCORE is a risk measure for severe complications (mortality) associated with this type of surgery. In addition to standard care treatment, patients undergoing cardiothoracic surgery were divided into two distinct categories based on
the type of treatments they received: a) placebo treatment (physiological buffer containing no AP) and b) bovine intestinal AP (bIAP) treatment in the same buffer.

In the APPIRED I study patients with a $2 < \text{EuroSCORE} \leq 6$, were initially given either placebo or 1000 IU of Bovine Intestinal AP (bIAP) followed by a continuous infusion during 36 hours of either placebo ($n=31$, with mean EuroSCORE $= 3.7 \pm 1.4$) or 5.6 IU per kg body weight per hour (total 9000 IU) ($n=32$, with mean EuroSCORE $= 3.6 \pm 1.2$).

In the APPIRED II study patients with a EuroSCORE of $\geq 5$, were initially given either placebo ($n=25$, with mean EuroSCORE $5.6 \pm 2.6$) or 1000 IU of bIAP followed by a continuous infusion during 8 hours (total 9,000 units) ($n=27$ with mean EuroSCORE $5.8 \pm 3.1$) (total 9,000 units).

Where in APPIRED I, 63 patients underwent CABG only, in APPIRED II a total of 52 patients were included that underwent CABG combined with valvular surgery under CPB. This type of combined surgery is associated with an increased risk.

Further details of the APPIRED clinical trials have been described by Kats et al. 28. The induction of endogenous alkaline phosphatase in this trial was described in 9. Primary endpoints were cytokine levels peri- and post-surgery next to clinical outcome.

The APPIRED II data was used in Sections 1.5 and 1.7 to validate the model. Data relative to APPIRED I was not used to validate the model due to the limited number of data points relative to AP. However, the secondary peak of ITMs observed in 16% of the patients in APPIRED I was used qualitatively to investigate the case of patients with complications in APPIRED II by adding a secondary source of ITMs in-silico in Section 1.8. This secondary source of ITMs is relative to the documented median peak IL6 concentration in septic shock patients in 53

Patients underwent CABG combined with valvular surgery. CABG pumps (heart-lung machine) were used during the surgery. The operations were
primary, therefore operated specifically for open heart surgery, and were planned prior to the actual surgery, hence non-emergent. An intra-aortic balloon pump was not used during the surgery, except for one patient who exhibited cardiogenic shock with multi-organ failure. The surgery lasted for an average of $4.7 \pm 1.4$ hours. The average perfusion time was $134 \pm 40$ minutes and average cross clamping time was $105 \pm 40$ minutes. Premedication such as relaxants, anaesthetics, antibiotics, and blood products such as red blood cells or platelets were given prior and during the CABG surgery. We summarize patients’ demographics, type and method of cardioplegia used, and patients’ medical history in Table 1, Table 2, and Table 3 of the Supporting Information respectively.

The data used was approved by the Ethics committee with IRB approval number M09-1965. The set-up of the study as well as appropriate consent procedures, have been reviewed and approved by the Institutional Review Board (METC). The central Independent Ethics Committee of The Netherlands (CCMO) has been informed. Approval from the METC of ZOL Genk was obtained in 6. March, 2012. The Belgium Competent Authority office: FAGG (Federaal Agentschap voor Geneesmiddelen en Gezondheidsproducten) was informed about the METC approval to initiate the study.

1.3. Biological Mechanisms of the Human Innate Immune System and Model Description

The HIIS model is constructed based on the biological mechanisms that occur in three separate compartments – blood, tissue and liver. The blood and tissue are separated by the endothelial lining that acts as a modulated barrier towards accessing blood circulation derived components like immune cells. Models of acute inflammation commonly neglect the dynamics of immune cells in the blood compartment under the assumption that it plays a minor role on the dynamics of the innate immune response, acting as a reservoir of immune cells. Systemic inflammation is considerably different and it is characterized by a dilution of the immune
cells in tissue caused by the delocalized inflammation. Additionally, AP is known to act on ITM both in blood and in tissue. For this reason, it is crucial to model both blood and tissue compartments to accurately capture the dynamic of resolution of a systemic inflammatory response.

**Blood Compartment**

Upon a systemic insult cytokines are released by both tissue cells and immune cells in tissue, with different rates. Cytokines then migrate first toward the endothelial barrier, with which they interact changing its permeability to recruit more immune cells into the inflamed tissue, and then in part migrate into the bloodstream. Since we only have data about the concentration of cytokines in blood, our model describes the dynamics of cytokines (IL-6 and IL-10) in the bloodstream. The rates of cytokines production used in our model represent the rate with which cytokines reach the bloodstream after being secreted by macrophages and necrotic neutrophils in tissue. These rates thus take into account not only the secretion rate by each cell type in response to the inflammatory state (presence of ITMs) but also of the mechanism of transport from tissue to the bloodstream.

In our model we assume that AP is the only component of the immune system that interacts with ITMs in the bloodstream, forming ITM-AP complexes that are later removed in the liver by Kupffer cells. Similarly to immune cells, the transport of AP and ITMs from the bloodstream into the tissue is controlled by a permeability factor.

**Tissue Compartment**

The innate immune response triggered in tissue by invasive cardiac surgery is shown in Fig 1. We assume that the presence of ITMs in tissue is mostly due to the migration into the tissue of ITMs released in bulk in the circulation through damaged blood cells and gut hypo-perfusion and later transported via the bloodstream. ITMs in tissue are also due to local tissue damage caused by the invasive surgery, but we consider this amount negligible compared to the other two sources of ITMs. An inflammatory response is
triggered as soon as ITMs activate resting macrophages ($M_R$) leading them to differentiate into “activated” macrophages residing in tissue (I). Activated macrophages ($M_A$) secrete pro-inflammatory cytokines ($CH$), which result in increasing the permeability of the endothelial barrier (II) via a series of intermediate stages. Consequently, resting neutrophils ($N_R$) in circulation are primed by circulating ITMs and then enter the tissue through the endothelial barrier via a process called ‘diapedesis’ (III). In the context of the computational model, resting neutrophils are only rendered active when they enter the tissue through the endothelial barrier. Activated neutrophils ($N_A$) phagocytose and/or release their granules to neutralize or antagonize inflammation (IV). If the inflammation is cleared, the neutrophils go into apoptotic or programmed death (V). Activated macrophages remove the apoptotic neutrophils ($ND_A$) by phagocytosis and in the process induce an anti-inflammatory effect as shown in Figure 1 by the green arrows (VI). If inflammation is too intense and not resolved rapidly, the neutrophils go into a necrotic ($ND_N$) state (designated by the red arrows in Figure 1), which releases additional ITMs in the tissue (VII). The presence of yet another batch of ITMs in the tissue induces ongoing inflammatory responses that causes tissue damage, which in turn perpetuates overall inflammatory response by macrophage activation and neutrophil influx into the local inflamed tissue areas (VIII).
Figure 1. Description of the innate immune response to inflammation. For details see the main text. This figure is taken from 7.

**Liver Compartment**

At the onset of surgery, a very high concentration of ITMs is released in the blood stream as a consequence of the damage to blood cells caused by the cardiac surgery bypass. In response to this massive ITM insult the liver releases all its stored AP (~5300 IU) into the bloodstream. After 2 – 4 hours the liver is able to supply newly synthesized AP again 54,55. Endogenous AP is naturally produced by the body and is highly expressed at physiological barriers like the gut, placenta, lungs, kidney glomerulus and the blood-brain barrier. Upon interaction with ITMs that are present in the bloodstream, endogenous AP is released from the apical membrane of specific physiological barriers expressing high levels of AP (like liver bile duct membrane, blood brain barrier, kidney and gut) and brought into circulation or gut lumen as ITM-AP conjugates. ITM-AP complexes are eventually removed from circulation by the liver Kupffer cells 24. We assume that due to its size AP can, under normal non-inflammatory conditions, enter the
tissue through the endothelial barrier fenestrae. Intravenously administered bovine AP (from here on noted as 'bIAP treatment') follows the same mechanism as endogenous AP, where it enters the tissue and detoxifies local ITMs through dephosphorylation. Note that the supplemented AP also detoxifies circulating ITMs directly and is removed from circulation by the Kupffer cells. In the case of an oxidative stress insult, such as that induced by cardiac surgery, ITM-AP is removed from circulation by Kupffer cells 24. This removal is observed in pre-clinical and clinical studies as a decrease in AP concentration in plasma. This decrease serves as a “distress signal” for the liver to release its stored AP at the bile ductal membrane barrier indirectly into the bloodstream (Pike et al., 2013). The release of liver AP into circulation implies that AP residing at blood-brain, kidney and gut-barriers may also be released, compromising the integrity of these barriers. This may result in clinical phenotypic conditions like kidney failure and cognitive impairment observed upon major surgery. The hypothesis central to the AP intervention is that by replenishing AP through either de novo synthesis or supplementation during surgery, the impairments can be circumvented by helping reduce the inflammation and preserving the integrity of such barriers. De novo synthesis, in the strictest sense, refers to the general production of an entity. In the context of our model, we use the term “de novo” synthesis as the continuous production of AP by the liver.

Our HIIS model takes into account the following key mechanisms: a) activation and inhibition of $M_R$, b) changes in endothelial permeability, c) phagocytosis of rest products of ITMs, d) phagocytosis of $NDA$ and $ND_N$, e) release of ITMs from necrotic cells, f) natural death of immune cells and degradation of molecular entities, g) production of $CH$ and $ACH$ h), induction of $D$, and finally i) delay in necrosis and cytokine production. The following key mechanisms are used to model the dynamics of AP: a) release of endogenous/stored AP from the liver bile canalicular membrane, b) de novo synthesis of AP in the liver, and c) administration of bIAP into the bloodstream. The model does not take into account the AP released from other physiological barriers, since the amount of AP present on these is
negligible compared to the AP released from the liver.

1.4. **Code Implementation and Repository**

We used Python 3.6.5 on a 3.30 GHz Intel® Core™ i7-5820K CPU with 16.0 GB RAM in all our simulations. Python libraries used were: numpy, pandas, scipy, joblib, SALib, and scikit-learn. The python codes and sample data have been uploaded to [https://github.com/avpresbitero/HIIS](https://github.com/avpresbitero/HIIS).

1.5. **Human innate immune system model with the induction mechanism of TNAP**

In the first sub-section we describe first the calibration process of the model with data from the bIAP branch of APPRED II under the assumption that supplemented bIAP stimulates the liver cells to produce additional TNAP. The calibrated model is then used to predict the dynamics of the immune response for the placebo branch. These predictions are validated using data from the placebo branch of APPRED II. In the second sub-section we show the dynamics of all cellular and molecular entities in the model and highlight the action of bIAP on the dynamics of systemic inflammation.

In this study we use the median for each branch of the APPRED II clinical trial to designate the values that best represent the population of patients undergoing cardiac surgery. Since we assume that the induction mechanism of alkaline phosphatase is inherent to all patients injected with bolus alkaline phosphatase, we did not cluster patients into different subgroups. Although clustering patients into subgroups based on their response would lead to a deeper understanding of the induction mechanism, the current dataset is not large enough to look into the individual trends of the patients’ blood parameters. Providing a personalized take on the modeling of the innate immune response and on the individual response to the supplemented AP, this endeavor is beyond the scope of the current research but will be investigated in future studies.
Calibration and Validation

Patients in the APPIRED studies were supplemented with a bolus of AP plus continuous infusion of AP and showed a surge of TNAP in the bloodstream (Figure 2A, bIAP Calibration). We calibrate the model parameters (summarized in 8.7 of the Supporting Information) using three datasets: AP, pro-inflammatory, and anti-inflammatory cytokine profiles of patients in the bIAP treatment experiment. We summarize the results in Figure 2.

In response to a massive insult, the liver releases all its stored AP into the bloodstream. The liver then takes roughly 2 hours to recover. We actually see this dynamics on the in-silico prediction of the model initially exhibited as a high concentration of AP at the onset of surgery. This is then followed by an immediate drop in AP concentration, corresponding to the time interval when the liver is still recuperating. Note that the effect of the AP bolus on the concentration disappears within 20 minutes after its supplementation as attributed to its short half-life and its interaction with ITMs. Then the liver begins supplying AP again at around 2 hours after surgery.

**bIAP Calibration**

**Placebo Validation**
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Figure 2. Innate immune response to systemic inflammation with the addition of the induction of TNAP by supplemented bIAP. The three plots (A, C, E) on the bIAP calibration column show the result of the calibration of the model parameters using data from the bIAP branch of APPIRED II. Data points are shown in red and correspond to the median value of the patients in this branch. The error bar shows the median absolute error. Blue lines correspond to the dynamics of the in silico model after calibration. (A) shows the dynamics of AP in blood (B) shows the dynamics of the pro-inflammatory cytokine represented in the model compared against IL6 data. (C) shows the dynamics of anti-inflammatory cytokines in the model against IL10 data. The three plots (B, D, F) on the placebo validation column show the validation of the model against data from the Placebo branch of APPIRED II. The model is able to predict the dynamics of placebo branch using the parameters calibrated with the data from the bIAP branch. The model predicts a protective effect of AP. As a consequence, the model predicts a greater concentration of pro-inflammatory cytokines in the placebo branch (D). See unit conversion of AP and cytokines from \( \frac{\text{molecules}}{\text{mm}^3} \) in Equations (31) and (32) (section 8.10 of the Supporting Information) respectively.

A continuous supply of bIAP was administrated into the patients for 8 hours, in addition to the initial concentration of 1000 IU bovine AP. It was observed that liver-type TNAP is induced in these patients as is shown by the overall concentration of AP in circulation in Figure 2 (A). This supports the
conjecture that, as a result of an ischemic condition, added AP serves as an indirect trigger for the liver to release more AP into the bloodstream. We therefore introduce an induction term \( \frac{r_{\text{inducepeak}}}{1 + \exp(r_{\text{induce}}(t - t_{\text{APdelay}}))} (AP_{\text{Stissue}} + AP_{\text{Sblood}}) \) in Equation (15) that is dependent on the concentration of bolus AP supplied into the system. The induction mechanism of AP is modelled as a reverse sigmoid function that is centered at 1 hour - corresponding to the lag of release of AP from the liver, having flushed all its contents, as the liver recuperates.

The rate at which AP is being used up by the system and the rate at which AP is replenished back into the bloodstream from the liver should be the same regardless of the treatment type. This is because the two groups of patients (bIAP and placebo branches) underwent the same type of cardiothoracic surgical procedure. Hence, we assume the same scale of insult, or the same amount of ITMs on both branches. We validate our model by using the parameter values that we have previously calibrated on the supplemented AP treatment branch to predict the AP profiles of patients in the placebo treatment branch. Our results are shown in Figure 2 (B Placebo Validation).

1.6. Dynamics of the HIIS with the induction mechanism of TNAP

**Dynamics of Macrophages**

In the case of a massive insult, such as cardiac surgery, where ITMs are simultaneously originating from numerous sources in the body, the entire population of resting macrophages immediately becomes activated. This is evident in Figure 3 (A) where we see, from simulated data, an immediate drop of resting macrophage population at the moment surgery is initiated (time = 0), which corresponds to an immediate rise of activated macrophage population as shown in Figure 3 (B).
A

Resting Macrophages

\[ \text{Concentration (cells/L)} \]

\[ \log_{10} \text{concentration} \]

Hours After Operation

B

Activated Macrophages

\[ \text{Concentration (cells/L)} \]

\[ \log_{10} \text{concentration} \]

Hours After Operation

**Figure 3.** (A) Resting Macrophages residing in tissue concentration drops within very short time after initiating ischemic insult upon a massive insult and induced by circulating ITM released under CPB/surgical conditions, which in our case is cardiac surgery. (B) Activated Macrophages concentration is driven to a maximum brought about by the immediate turnover of resting macrophage population to activated macrophages.

**Dynamics of Neutrophils**

In the context of our model, resting neutrophils become “activated” when they enter the tissue from the bloodstream via the endothelial barrier. The recruitment of neutrophils is proportional to the concentration of pro-inflammatory cytokines that increase the permeability of the endothelial barrier. This means that the larger the insult, the more resting neutrophils are recruited from the blood stream into the tissue.

For placebo patients the model predicts (Figure 4) an increased level of neutrophils necrosis in tissue in response to slightly higher concentration of ITM in tissue. The increased number of necrotic neutrophils leads to the production of additional ITMs, which acts as a positive feedback for inflammation. In AP-treated patients, the presence of additional AP prevents or reduces the necrosis of neutrophils. This could explain the lower number of adverse events reported for the AP branch of the clinical trial compared to the placebo branch.28

The model dynamics shown in Figure 4 supports the idea that AP is an anti-inflammatory mediator that plays an important and active role in the human innate immune system, even though the impact of AP observed with the current levels of ITMs appears to be confined to a small shift in the ratio of
apoptotic versus necrotic neutrophils.

Figure 4. Dynamics of (A) ITMs in Plasma, (B) ITMs in Tissue, (C) Resting Neutrophils, (D) Activated Neutrophils,
Activated neutrophils in the placebo treatment go into necrosis quicker than in the supplemented AP treatment. It seems indeed that the addition of bolus AP contributes to the human innate immune system as an anti-inflammatory mediator by reducing the amount of neutrophils that go into the necrosis pathway.

1.7. HIIS model without the induction mechanism of TNAP

In this section we present the model results under the assumption that supplemented biAP does not stimulate the liver cells to produce additional TNAP. In this case we use the previous model without the induction term introduced in section 0. We first attempted to calibrate the parameters of this alternative model on the supplemented biAP branch. However, we were not able to model the AP dynamics of the biAP branch without the induction term. For this reason we calibrated the model with data from the placebo branch of APPIRED II. The calibrated alternative model is then used to predict the dynamics of the immune response for the biAP branch. We compare these predictions with data from the biAP branch of APPIRED II and observe that the calibrated model fails to predict the AP dynamics observed in the clinical trial. Since the model without the induction mechanism cannot be validated we do not show the detailed dynamics of cellular and molecular entities as we did in section 3.1.2.

**Calibration and Validation**

Instead of calibrating the parameters using the AP treatment and validating using the placebo treatment, we now reverse the process and calibrate instead the parameters in the placebo treatment first and validate them using the AP patient data. The aim of which is to find out whether we could model the induced amount of endogenous AP in the supplemented branch without using the induction term.
Figure 5. Innate immune response to systemic inflammation without the induction term. The three plots (A, C, E) on the placebo calibration column summarize the result of the calibration of the model parameters using data from the placebo branch of APPIRED II. Data points are shown in red and correspond to the median
value of the patients in this branch. The error bar shows the median absolute error. Blue lines correspond to the dynamics of the in silico model after calibration. (A) shows the dynamics of AP in blood. (B) shows the dynamics of the pro-inflammatory cytokine represented in the model compared against IL6 data. (C) shows the dynamics of anti-inflammatory cytokines in the model against IL10 data. The three plots (B, D, F) on the bIAP validation column show the validation of the model against data from the bIAP branch of APPREDE II.

Without the induction term, we are not able to reproduce the Alkaline Phosphatase profile in the AP-treated patients. See unit conversion of AP and cytokines from $\text{mm}^3$ in Equations Equations (31) and (32) (section 8.10 of the Supporting Information) respectively.

Using the calibrated parameters from the placebo study, we predict the dynamics of the bIAP study without the induction term. As shown in Figure 5B, the model without the induction term is not able to reproduce the AP profile of bIAP treatment patients. This clearly shows that an additional mechanism is missing and that the missing term has to account for additional production of TNAP to be released from the liver (the major source of stored AP) into the bloodstream. See Supporting Information section 8.9 for details on the dynamics of the various compartments of the HIIS without the induction mechanism of TNAP.

1.8. Predicting the innate immune response for patients having excess ITMs using different AP treatment regimens

The validation in section 3.1 shows that the model with an induction mechanism for TNAP is able to predict the dynamics of the human innate immune system response in patients with systemic inflammation. Under the conditions of reported in APPREDE II patients in the placebo treatment branch have been able to resolve inflammation almost as effectively as patients in the bIAP branch, the only measurable difference being different levels of plasma AP and a reduced number of adverse events in the bIAP branch. Supplementation of AP under these conditions has an impact on the amount of necrotic neutrophils but did not drastically change the dynamics of immune cells from that of the placebo treatment group. However, since in the APPREDE I study 16% of patients show an excess amount of ITMs, the
source of which is unknown, we explore the impact of supplemented AP in a system stressed by an additional source of ITMs after surgery. We model this scenario by adding a secondary source of insult in-silico and predicting how the human innate immune system would respond in different AP regimens using our model. The amount of this secondary source of ITMs is set to a reasonable value as indicated in. We perform two sets of in-silico experiments: in the first set we predict how the immune cells respond to an excess amount of ITMs; in the second set we predict how different AP regimens (i.e. different concentrations of bolus AP) affect the body's response to an additional source of ITMs as the one observed in APPIRED I.

**In-Silico Experiment #1: Innate immune system dynamics for patients with excess ITMs**

The model predicts a protective effect due to supplemented AP when a patient is challenged by a second source of ITMs immediately after or during the surgery. Figure 6 and 7 show the predicted dynamics for bIAP and placebo branches in case of a source of additional ITMs. Apoptotic neutrophils in supplemented branch have a concentration higher than in the placebo branch. On the other hand, necrotic neutrophils and ITMs in placebo branch are higher than in the supplemented treatment branch. The model predicts a more intense inflammation in the placebo branch as shown in the pro-inflammatory cytokines plot (Figure 7C). This is confirmed by (Figure 7D) which shows more anti-inflammatory cytokines in the supplemented than in the placebo branch.
Figure 6. (A) Activated Neutrophils, (B) Apoptotic Neutrophils, (C) Necrotic Neutrophils, and (D) ITMs in Tissue in Supplemented and Placebo branches for patients with excess ITMs. The model predicts higher concentrations of apoptotic neutrophils in the supplemented than in the placebo branch, but higher concentrations of necrotic neutrophils in the placebo than in the supplemented branch.
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Figure 7. (A) Alkaline Phosphatase, (B) ITMs in Plasma (C) Pro-inflammatory, and (D) Anti-inflammatory Cytokine Concentrations in Supplemented and Placebo branches for patients with excess ITMs. The placebo branch produces more pro-inflammaotry cytokines and lesser anti-inflammatory cytokines, suggesting a more intense inflammation in the placebo branch compared to the supplemented branch. See unit conversion of AP and cytokines from $\text{mmol} \cdot \text{m}^{-3}$ in Equations (31) and (32) (section 8.10 of the Supporting Information) respectively.

In-Silico Experiment #2: Treating Patients with excess ITMs with Various Alkaline Phosphatase Regimen

Phase II and Phase IIIa clinical trials have observed an increased concentration of TNAP in circulation in the AP treatment group compared to the placebo group given an AP protocol. For instance in APPIRED I, a bolus of 1000 IU of bovine AP was first injected to patients undergoing cardiac
surgery followed by a continuous infusion of 5.6 IU/L per kg body weight for 8 hours. In this section we predict the dynamics of the innate immune response under the conditions described in section 3.3.1 given two different AP supplementation regimens for which we increase the supplemented AP to twice and thrice the original protocol respectively. We then compare the results with the protocol tested in APPIRED II study.

The model predicts an increasing protective effect the higher the concentration of supplemented AP by showing an increasing neutralizing effect on ITMs both in plasma and in tissue (Figure 8A and B). As the concentration of supplemented AP increases, more and more activated neutrophils are inclined to go into apoptosis rather than necrosis (Figure 8C and D). Pro-inflammatory profiles show that increasing supplemented AP decreases the amount of pro-inflammatory cytokines, while increasing the population of anti-inflammatory cytokines, indicating a better resolution of the systemic inflammation.
1.9. Summary and Conclusion

The induction mechanism of liver-type Tissue Non-specific Alkaline Phosphatase (TNAP) into circulation observed in patients undergoing coronary artery bypass grafting with cardiac valve replacement could be the body's way of strengthening its defense mechanism against such a massive insult, making TNAP a de facto key player in the human innate immune system. Hence, directing the attention to the role of Alkaline Phosphatase in
systemic inflammation and understanding its role in supporting an appropriate innate immune response is of utmost immunological importance. Computational modelling makes it possible to mimic and understand such intricate details and mechanisms of the human innate immune system and offers a predictive power for experimental outcomes through *in-silico* experiments.

To our knowledge, we have developed the first mathematical model of systemic inflammation that is calibrated and validated on clinical data. We show that the amount of additional endogenous TNAP released in circulation in the supplemented branch is proportional to the total concentration of bolus AP being supplied. We also provide a plausible mathematical function that describes this mechanism. Our model predicts a protective effect of AP in the supplemented branch of APPIRED II, evidence of which can be concluded from the dynamics of the neutrophils. This effect, minimal under the conditions of APPIRED II, becomes obvious when patients exhibit an excess of ITMs after surgery (as observed in 16% of patients of APPIRED I study). We present a scenario where we mimic excess ITMs as seen in some patients by adding a secondary source of insult and see how the model reacts to different AP regimens by varying doses of AP supplied to patients. We show that additional AP has indeed a protective effect and this effect is more prominent in patients with excess ITMs. In this case *in-silico* experiments predict that the amount of apoptotic neutrophils in the supplemented AP branch is much higher than in the placebo branch. Additionally, the amount of pro-inflammatory cytokines predicted for the supplemented AP branch is lower than in the placebo branch, giving further evidence that supplemented AP reduces the intensity of systemic inflammation. As expected, the model predicts more anti-inflammatory cytokines in the supplemented branch than in the placebo branch. In other words, the dynamics predicted by the *in-silico* model show that resolution of inflammation is faster and more efficient with increasing concentration of AP. Hence, our findings suggest that AP indeed plays an important role in mitigating inflammation especially in systemic inflammation and that this
protective effect can be modified through variation in AP protocol.

Our model for systemic inflammation is in fact similar to mechanistic models of physiological process, more specifically that of pharmacokinetic/pharmacodynamic (PKPD) models, that are used to develop insights on the dynamics and magnitude of the effect of a drug through quantitative analysis. These models are used to describe the dynamics of the physiological variables in different states. In our case, we model and validate the dynamics of bolus AP together with the human innate immune response for patients undergoing cardiac surgery in two treatment arms respectively: with or without AP supplementation. After validation, the model is used to understand and predict the effect of experimental perturbations, such as variations in AP regime, an approach that is referred to as “forward engineering.” A synergy, henceforth, is created between systems biology and PKPD through iterations between computational/mathematical modeling and experimentation 56. Helmlinger et al. have provided a comprehensive review of drug-disease modeling in the pharmaceutical industry 57. A robust application of systems pharmacology, or the application of systems biology in order to understand how drugs affect the human body, is detailed by Gadkar et al. in 58.

The current model has three main limitations: the limited biological knowledge regarding the mechanism that regulates the induction of endogenous AP production triggered by supplemented AP, the potential bias introduced by the modeling approach, and the bias introduced by studying the median dynamics rather than attempting to cluster patients in groups with different dynamics.

1) The goal of this model is to prove the existence of AP induction and propose a possible mechanism describing these dynamics. We did validate the existence of the induction of AP, yet the available experimental data is not sufficient to unequivocally unravel the underlying mechanism.

2) The other limitation relates to the modeling approach used. Ordinary
differential equations do not take into account the spatial properties of the innate immune response during systemic inflammation. The spatial effects are limited to the compartmentalization of the body into blood, liver and tissue while microscopic spatial effects of the cellular dynamics are neglected. It is possible that modeling the spatial properties of the system would result in a more accurate description of the dynamics, especially for the first 3 hours during which there is a major displacement of cells between blood and tissue. For that, we would need a high resolution spatial-temporal clinical data.

3) Given the data at our disposal we decided to study the systemic inflammation via the median dynamics of the two branches of the APPIRED II clinical trial. The large variability in the dataset suggests that clustering patients in subgroups with different dynamics might provide a more accurate prediction of model parameters. However, we believe that the approach used in this paper is sufficient to prove the existence of the induction mechanism of AP and to provide a preliminary description of its dynamics.

The physiological relevance of this study is that to the best of our knowledge this is the first mathematical model describing systemic inflammation. Additionally, this is the first model describing the role of Alkaline Phosphatase in the resolution of inflammation after invasive cardiac surgery, laying the foundations to understand systemic inflammatory response syndrome. The main clinically relevant result is the evidence of the existence of an induction mechanism triggered by supplemented AP. This model provides a starting point to investigate the amount of endogenous AP induced in the body, and consequently, the optimal amount of supplemented AP to be administered during and after invasive cardiac surgery.

Using the proposed model, we have shown in-silico the dynamics of systemic inflammation within a period of 36 hours using different AP regimens. This information is being taken into account in the planning of a clinical trial phase III b. Data from the new multi-center clinical trial will be
used to further refine the model. This model and its future iterations will be useful to predict the dynamics of neutrophils during systemic inflammation (namely the balance between apoptotic and necrotic neutrophils and the subsequent resolution of inflammation) and act as a tool to optimize the administration of anti-inflammatory drugs (not necessarily AP) in clinical trials dealing with systemic inflammatory response syndromes.

We perform a global sensitivity test (see Supporting Information Section 8.8) where we vary our input parameters within intervals that correspond to the values found in literature.

The model provides evidence of the existence of an induction mechanism of liver-type tissue non-specific alkaline phosphatase, triggered by the supplementation of AP in patients undergoing cardiac surgery. We show that the AP branch of the clinical trial can only be explained using a mechanism that induces a release in circulation of liver TNAP that is proportional to the amount of supplemented AP. We provide a possible mathematical description of this induction mechanism. The model is validated using novel clinical AP, pro-inflammatory and anti-inflammatory cytokine profiles of placebo- and bIAP-treated patients. This is the first time that liver-type tissue nonspecific alkaline phosphatase has been modelled together with the human innate immune system. To date, there are no other existing published clinical trials that tackle the exact mechanisms of liver-type TNAP induction, let alone, a model that describes systemic inflammation. To the best of our knowledge this is the first numerical model of a complex innate immune response that is quantitatively validated with clinical data. Our work paves the way to a deeper understanding of the immunological mechanisms underpinning this important innate immune response to oxidative stress mediated inflammation.

1.10. **Supporting Information**

All references of the supporting information for this chapter are in Section 1.10.10.
1.10.1. Model Terms and Biological Mechanisms

Activation and Inhibition of Resting Macrophages

As deduced from the prevalence of complications following cardiothoracic surgery, i.e. in kidney function and brain temporary cognitive impairment, the mere presence of inflammation triggering moieties (ITMs) as by-products of simultaneous insults, both local or transported into the tissue, is enough to activate resting macrophages ($M_R$) at a rate of $\phi_{M_R|ITM}$. The activation of $M_R$ is inhibited by the presence of anti-inflammatory cytokines ($A_C$) at a rate of $\theta_{A_C}$ and consequently the concentration of $M_A$ is reduced accordingly. This is motivated by the biological mechanism where increasing concentration of apoptotic neutrophils cleared by activated macrophages induces a higher concentration of $A_C$. In other words, the higher $A_C$ concentration is, the slower the activation process of resting macrophages becomes.

$$-\phi_{M_R|ITM}M_RITM_{issue} \frac{1}{1 + \theta_{A_C}A_C}$$

Induction of Endothelial Permeability

We model diapedesis and the movement of primed cellular entities from the blood into the tissue compartment by a permeability factor of the endothelial membrane, to account for the macroscopic spatial dynamics (movement of cells between blood and tissue) involved in a systemic inflammatory response.

Pro-inflammatory cytokines ($C_H$) indirectly result in opening up the endothelial barrier. This allows the migration of monocytes and circulating primed neutrophils, as well as plasma resident molecules like albumin and alkaline phosphatase, from the bloodstream into the tissue. We model endothelial permeability through Equation (2), where $\zeta$ could either be $M_R, N_R, ITM$, or AP. $P_{\zeta}^{\max}$ and $P_{\zeta}^{\min}$ are the maximum and minimum endothelial permeability respectively, $C_H$ designates the concentration of...
pro-inflammatory cytokines, while \( K_{eqCH} \) is the concentration of \( CH \) that gives 50% of the maximum effect on permeability. The term \( f_{dilution} \) is simply a dilution factor that represents the diffusion of immune cells from the bloodstream (5 liters) into the tissue (75 liters, assuming an 80 kg person)

\[
f_{dilution} = \frac{p_{\max} - p_{\min}}{CH + K_{eqCH} + p_{\min}}
\]

Equation (2) is modelled using a Hill type equation to make the permeability of the endothelial barrier non-linearly dependent on the \( CH \) concentrations. These permeability factors are limited by a multiplier that depends on the represented entity.

**Interaction of cells with Inflammation Triggering Moieties**

Equation (3) models the interaction of cells with ITMs by \( M_A, N_A, G, \) and \( AP \), where the \( \lambda \) coefficients correspond to the rates at which the ITMs are being detoxified by dephosphorylation in the tissue.

\[
ITM_{tissue}(\lambda_{ITM|M_A|MA} + \lambda_{ITM|N_A|NA} + \lambda_{ITM|AP|Etissue}AP_{Etissue} + \lambda_{ITM|AP|Stissue}AP_{Stissue})
\]

**Phagocytosis of Apoptotic and Necrotic Neutrophils**

Activated neutrophils either go into apoptosis or necrosis depending on the scale of insult in the body. \( M_A \) then proceeds to dispose of these neutrophils through phagocytosis. This process is modelled by Equation (4) where \( \eta \) represents either apoptotic or necrotic state of neutrophils, and \( \lambda_{M_A|ND_\eta} \) is the rate of phagocytosis.

\[
\lambda_{M_A|ND_\eta}M_A ND_\eta
\]

**Induction of Inflammation Triggering Moieties**

Neutrophils undergoing necrosis cause all of their cellular contents, with
endogenous-ITM characteristics, to spill out into the surrounding tissue, thereby triggering a vigorous pro-inflammatory response. We model the induction of ITMs through Equation (5) where $\alpha_{ND} N$ is the rates of induction of ITM due to necrotic neutrophils ($ND_N$).

$$\alpha_{ND} N$$

### Natural Decay of Species

HIIS key players can disappear from the system due to natural species decay, modelled in Equation (6) where $\mu_{\psi}$ represents the decay rate of species $\psi$.

$$-\mu_{\psi} \psi$$

### Production of Pro-inflammatory and Anti-inflammatory Cytokines

Equation (7) models the production of CH, such as interleukin (IL)-1, and tumor necrosis factor (TNF), where the term $\beta_{MAITM} M \ ITM_{tissue} + \beta_{NAITM} N \ ITM_{tissue}$ describes the production of CH when $M_A$ and $N_A$ phagocytose ITM. This mechanism is regulated by $ACH$. The larger the $ACH$ concentration is, the slower the increase in CH. The term $1 - CH/CH^{max}$ models the growth factor of CH. Hence, CH is no longer produced when it reaches a maximum concentration of $CH^{max}$ in the tissue.

$$\frac{(\beta_{MAITM} M \ ITM_{tissue} + \beta_{NAITM} N \ ITM_{tissue})(1 - \frac{CH}{CH^{max}})}{1 + \theta_{ACH} ACH}$$

ACH, such as IL4, IL10, and IFN-alpha, are actively produced by $M_A$, as well as when $M_A$ phagocytoses $ND_A$. This is modelled by the term $$(\beta_{ND} M_A N \ ITM_{tissue} + \alpha_{ACHM} M_A)$$ in Equation (8). The production of ACH is limited by the maximum concentration of ACH allowed in the tissue ($ACH^{max}$) through a growth factor represented by the term $1 -$.
\[ \frac{ACH}{ACH^{\text{max}}} = (\beta_{ND_A|M_A}M_A ND_A + \alpha_{ACH|M_A}M_A)(1 - \frac{ACH}{ACH^{\text{max}}}) \] (8)

**Necrosis and Cytokines Delay**

Neutrophils naturally die through a programmed death process called "apoptosis." However, if the stimulus is too strong, neutrophils go into the necrotic state, which is a turbulent cell death that triggers the release of tremendous amounts of ITMs in the surrounding tissue. It remains unclear how much of this activated neutrophil concentration goes into apoptosis, or conversely into necrosis. We simplify these mechanisms in our model by assuming that necrosis takes place when activated neutrophils phagocytose ITMs. Therefore, the larger the insult, the more ITMs there are in the body and the higher the concentration of necrotic neutrophils becomes. Apoptosis, on the other hand is simply modelled by the decay of activated neutrophils and is inversely proportional to the concentration of ITMs in the system. These mechanisms are modelled by Equation (9), where the term on the left corresponds to apoptosis, and the term on the right to necrosis with temporal delay \( t_{NDN\text{delay}} \) of 5 hours.

\[ -\mu_{ND_A} \frac{N_A}{1 + ITM_{\text{tissue}}} - \frac{1}{1 + \exp^{-\tau_{ND}(t-t_{NDN\text{delay}})}} \lambda_{ITM|N_A,N_A ITM_{tissue}} \] (9)

It is also of great importance to model the delay of the cytokine response. This was observed in patients by cytokines increment in the plasma after about 20 minutes. We model this delay mechanism as a step function given by Equation (10) where \( b \) is the rate which we set to maximum to account for the immediate increase in cytokine population as soon as the time reaches \( t_{CH\text{delay}} \).

\[ \frac{1}{1 + \exp^{-b(t-t_{CH\text{delay}})}} \] (10)
Release of Endogenous Alkaline Phosphatase from the Liver

Alkaline phosphatase in the bloodstream is used up in numerous ways. The body then establishes homeostasis by releasing a considerable amount of liver-derived AP (about 50% of total plasma concentration of AP) into the blood circulation through a process described by Pike et al., (2013). Note that the largest amount of AP in the body is stored at the bile canalicular membrane. We model this supply of AP through a Gaussian function that is centred around zero $\mathcal{A}_{\mathcal{E}_{\mathcal{E}}\mathcal{M}_{\mathcal{M}}} + \mathcal{A}_{\mathcal{S}_{\mathcal{S}}\mathcal{S}_{\mathcal{S}}\mathcal{M}_{\mathcal{M}}}$ concentration, which served as the body’s “distress signal” to fire up the liver into releasing its contents (indirectly) into the blood stream. ‘w’ is the width of the Gaussian function that sets the AP “distress signal” concentration and $r_{\text{distress}}$ is the rate at which AP is released into circulation by the liver.

$$r_{\text{distress}} \left( \exp \left( \frac{(\mathcal{A}_{\mathcal{E}_{\mathcal{E}}\mathcal{M}_{\mathcal{M}}} + \mathcal{A}_{\mathcal{S}_{\mathcal{S}}\mathcal{S}_{\mathcal{S}}\mathcal{M}_{\mathcal{M}}})^2}{2w^2} \right) \right)$$  \quad (11)

Administration of Bolus Alkaline Phosphatase

In APPIRE II, a total amount of 10,000 units AP (1000 IU bolus and 9000 IU infusion) was administered over a period of 8 hours. AP infusion is modelled by a step function given by Equation (12) where $b$ is set to maximum to represent the sudden decline in bIAP concentration as soon as the infusion has been terminated at $t_{\text{stop}}$.

$$r_{\text{inject}} \left( \frac{1}{1 + \exp[b(t-t_{\text{stop}})]} \right)$$  \quad (12)

Induction of Alkaline Phosphatase

The intricate details pertaining to the underlying mechanisms to how alkaline phosphatase is induced upon a systemic insult are still unknown and not found anywhere in literature. We propose a hypothesis that the induced AP is only proportional to the amount of supplemented AP that is in the system. We model this induction mechanism as the inverse of a sigmoid function where the amplitude is dependent on parameter $r_{\text{induce peak}}$ and the
total concentration of supplemented AP that is in the system \((AP_{\text{tissue}} + AP_{\text{blood}})\). Note that the induction mechanism has a delay of \(t_{\text{APdelay}}\), which refers to the time it takes for the liver to be able to supply AP again, having emptied out 1000 IU at the onset of surgery.

\[
\frac{r_{\text{inducepeak}}}{1+\exp^{r_{\text{induce}}(t-t_{\text{APdelay}})}}(AP_{\text{tissue}} + AP_{\text{blood}})
\]  

### 1.10.2. The Human Innate Immune System Model (HIIS)

The HIIS model is summarized in equations (14)-(27). Equations (14)-(19) model the biological mechanisms in the blood compartment while equations (20)-(27) correspond to the tissue compartment. The parameters and initial values that we used in the model are those from literature, as well as those indicated in the cardiac surgery clinical study.

**Blood Compartment**

\[
\frac{dITM_{\text{blood}}}{dt} = -\mu_{\text{ITM}}ITM_{\text{blood}}
\]

\[
-ITM_{\text{blood}}(\lambda_{\text{ITM}|AP_{\text{b}}}AP_{\text{b}} + \lambda_{\text{ITM}|AP_{\text{s}}}AP_{\text{s}})
\]

\[
-\left([p_{\text{max}} - p_{\text{min}}] \frac{CH}{CH + K_{eqCH}} + p_{\text{min}} \right)ITM_{\text{blood}} \left(1 - \frac{ITM_{\text{tissue}}}{ITM_{\text{max}}} \right)
\]

\[
+ \frac{t_{\text{ITMpeak}}}{1 + \exp^{t_{\text{ITM}}(t-t_{\text{ITM}})}} \left(1 - \frac{ITM_{\text{blood}}}{ITM_{\text{max}}} \right)
\]
\[
\frac{dAP_{Eblood}}{dt} = -\mu_{AP_e} AP_{Eblood} - \left[[P_{AP}^{\text{max}} - P_{AP}^{\text{min}}] \frac{CH}{CH + K_{eqCH}} + P_{AP}^{\text{min}} \right] AP_{Eblood} - \lambda_{ITM}\|AP_e ITM_{blood} AP_{Eblood} + \frac{1}{1 + \exp^{-\tau_{distress}}\left(\exp\left(\frac{(AP_{Eblood} + AP_{Sblood})^2}{2\omega^2}\right)\right)}
\]

\[+ \frac{r_{\text{distress}}}{1 + \exp^{-\tau_{AP}(t-t_{AP\text{delay}})}} \left(\exp\left(\frac{(AP_{Eblood} + AP_{Sblood})^2}{2\omega^2}\right)\right) (15)\]

\[
\frac{dAP_{Sblood}}{dt} = -\mu_{AP_s} AP_{Sblood} - \text{dilution} \cdot \left[[P_{AP}^{\text{max}} - P_{AP}^{\text{min}}] \frac{CH}{CH + K_{eqCH}} + P_{AP}^{\text{min}} \right] AP_{Sblood} + \frac{r_{\text{distress}}}{1 + \exp^{-\tau_{distress}}(t-t_{AP\text{delay}})} \left(\exp\left(\frac{(AP_{Eblood} + AP_{Sblood})^2}{2\omega^2}\right)\right) (16)\]

\[
\frac{dN_R}{dt} = -\mu_{N_R} N_R - \left[P_{max} \frac{CH}{CH + K_{eqCH}} \left(1 - \frac{N_A}{N_R^{\text{max}}}\right)\right] N_R + r_{\text{Nhomo}} (1 - \frac{N_R}{N_R^{\text{max}}}) N_R (17)\]

\[
\frac{dCH}{dt} = -\mu_{CH} CH + \frac{1}{1 + \exp^{-\beta(t-t_{CH\text{delay}})}} \left[\beta_{M_A|ITM} M_A ITM_{tissue} + \beta_{N_A|ITM} N_A ITM_{tissue}\right] (1 - \frac{CH}{CH^{\text{max}}}) (1 + \theta_{ACH} ACH) (18)\]

\[
\frac{dACH}{dt} = -\mu_{ACH} ACH + (\beta_{N,D_A|M_A} M_A N_D + \alpha_{ACH|M_A} M_A) (1 - \frac{ACH}{ACH^{\text{max}}}) (19)\]

**Tissue Compartment**

\[
\frac{dITM_{tissue}}{dt} = -\mu_{ITM} ITM_{tissue} - ITM_{tissue} \left(\lambda_{ITM}\|M_A M_A + \lambda_{ITM}\|N_A N_A\right) + \lambda_{ITM}\|AP_{Itissue} AP_{Itissue} + \lambda_{ITM}\|AP_{Stissue} AP_{Stissue} \left(\lambda_{ITM}\|AP_{Itissue} + \lambda_{ITM}\|AP_{Stissue}\right) + \alpha_{N,D_N} N_D + \text{dilution} \cdot \left[[P_{max} - P_{min}] \frac{CH}{CH + K_{eqCH}} + P_{min} \right] ITM_{blood}(1 - \frac{ITM_{tissue}}{ITM_{max}}) (20)\]
\[
\frac{dM_R}{dt} = -\mu M_R M_R - \frac{\phi_{M_R|TM} M_R ITM_{tissue}}{1 + \theta_{ACH} ACH}
\]

\[
+ f_{dilution}\left( (p^\text{max}_{M_R} - p^\text{min}_{M_R}) \frac{CH}{CH + K_{eqCH}} + p^\text{min}_{M_R} \right) [M^\text{max} - (M_R + M_A)]
\]

\[
\frac{dM_A}{dt} = -\mu M_A M_A + \frac{\phi_{M_A|TM} M_A ITM_{tissue}}{1 + \theta_{ACH} ACH}
\]

\[
\frac{dN_A}{dt} = -\mu_{NDA} N_A 1 + ITM_{tissue} - \frac{1}{1 + \exp^{-r_{NDN}(t - t_{NDN delay})}} \frac{\lambda_{ITM|N_A} N_A ITM_{tissue}}{N_R}
\]

\[
+ f_{dilution}[P^\text{max}_{N_R} \frac{CH}{CH + K_{eqCH}}] (1 - \frac{N_A}{N_R}) N_R
\]

\[
\frac{dNDA}{dt} = \mu_{NDA} \frac{N_A}{1 + ITM_{tissue}} - \lambda_{NDA|M_A} M_A NDA
\]

\[
\frac{dNDN}{dt} = \frac{1}{1 + \exp^{-r_{NDN}(t - t_{NDN delay})}} \frac{\lambda_{ITM|N_A} N_A ITM_{tissue}}{N_R} - \lambda_{NDN|M_A} M_A NDN
\]

\[
\frac{dA_P_{tissue}}{dt} = -\mu_{A_P} A_P_{tissue} + f_{dilution}\left( (p^\text{max}_{A_P} - p^\text{min}_{A_P}) \frac{CH}{CH + K_{eqCH}} \right)
\]

\[
+ [p^\text{min}_{A_P}] A_P_{blood} - \lambda_{ITM|A_P} A_P ITM_{tissue} A_P_{tissue}
\]

\[
\frac{dA_P_{blood}}{dt} = -\mu_{A_P} A_P_{blood} + f_{dilution}\left( (p^\text{max}_{A_P} - p^\text{min}_{A_P}) \frac{CH}{CH + K_{eqCH}} \right)
\]

\[
+ [p^\text{min}_{A_P}] A_P_{tissue} - \lambda_{ITM|A_P} A_P ITM_{tissue} A_P_{tissue}
\]
Table 1. Summary of Terms and Description Used in the Human Innate Immune System Model. Note that the values indicated for Immune cells (entries 1-14) are initial values.

<table>
<thead>
<tr>
<th>Category</th>
<th>Equation Terms</th>
<th>Description</th>
<th>Value</th>
<th>Unit</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$ITM_{blood}$</td>
<td>concentration of Inflammation Triggering Moeties (ITMs) in the bloodstream</td>
<td>$3.5 \times 10^7$</td>
<td>molecules/mm$^3$</td>
<td>60</td>
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<td>2</td>
<td>$AP_{eblood}$</td>
<td>concentration of endogenous Alkaline Phosphatase (AP) in the bloodstream</td>
<td>4000</td>
<td>IU</td>
<td>61</td>
</tr>
<tr>
<td>3</td>
<td>$AP_{sblood}$</td>
<td>concentration of supplemented AP in the bloodstream</td>
<td>1000</td>
<td>IU</td>
<td>experiment*</td>
</tr>
<tr>
<td>4</td>
<td>$N_R$</td>
<td>concentration of resting neutrophils</td>
<td>$1.2 \times 10^5$</td>
<td>cells/mm$^3$</td>
<td>62</td>
</tr>
<tr>
<td>5</td>
<td>$CH$</td>
<td>concentration of pro-inflammatory cytokines</td>
<td>0</td>
<td>molecules/mm$^3$</td>
<td>50</td>
</tr>
<tr>
<td>6</td>
<td>$ACH$</td>
<td>concentration of anti-inflammatory cytokines</td>
<td>0</td>
<td>molecules/mm$^3$</td>
<td>50</td>
</tr>
<tr>
<td>7</td>
<td>$ITM_{tissue}$</td>
<td>concentration of ITMs in the tissue</td>
<td>$3.5 \times 10^5$</td>
<td>molecules/mm$^3$</td>
<td>50</td>
</tr>
</tbody>
</table>
Chapter 1 Modeling the Human Innate Immune Response for Systemic Inflammation

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<tr>
<td>8</td>
<td>$M_R$</td>
<td>concentration of resting macrophages</td>
<td>$1.5 \times 10^3$</td>
<td>cells mm$^{-3}$</td>
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<td>9</td>
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<td>concentration of activated macrophages</td>
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<td>cells mm$^{-3}$</td>
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<td>cells mm$^{-3}$</td>
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<td>12</td>
<td>$ND_N$</td>
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<td>cells mm$^{-3}$</td>
<td>50</td>
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<td>13</td>
<td>$AP_{Etissue}$</td>
<td>concentration of endogenous AP in the tissue</td>
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<td>IU</td>
<td>experiment*</td>
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<td>14</td>
<td>$AP_{Stissue}$</td>
<td>concentration of supplemented AP in the tissue</td>
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<td>IU</td>
<td>experiment*</td>
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<tr>
<td>15</td>
<td>$\mu_{ITM}$</td>
<td>Decay rate of ITMs</td>
<td>$4.4 \times 10^{-6}$</td>
<td>1/min</td>
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<tr>
<td>16</td>
<td>$\mu_{AP_e}$</td>
<td>Decay rate of endogenous AP</td>
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<td>17</td>
<td>$\mu_{MR}$</td>
<td>Decay rate of resting macrophages</td>
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<td>1/min</td>
<td>62</td>
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<tr>
<td>18</td>
<td>$\mu_{ND_A}$</td>
<td>Decay rate of necrotic neutrophils</td>
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<td>-</td>
<td>calibrated</td>
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</tr>
<tr>
<td>20</td>
<td>$\mu_{AP_s}$</td>
<td>Decay rate of supplemented AP</td>
<td>0.1</td>
<td>1/min</td>
<td></td>
</tr>
<tr>
<td>21</td>
<td>$\lambda_{ITM</td>
<td>AP_E}$</td>
<td>neutralization rate of ITMs by endogenous AP</td>
<td>$3.9 \times 10^{-9}$</td>
<td>mm$^3$/molecules.min</td>
</tr>
<tr>
<td>22</td>
<td>$\lambda_{ITM</td>
<td>AP_s}$</td>
<td>neutralization rate of ITMs by supplemented AP</td>
<td>$2.6 \times 10^{-9}$</td>
<td>mm$^3$/molecules.min</td>
</tr>
<tr>
<td>23</td>
<td>$\lambda_{ITM</td>
<td>M_A}$</td>
<td>phagocytosis rate of ITMs by activated macrophages</td>
<td>$7 \times 10^{-4}$</td>
<td>mm$^3$/cells.min</td>
</tr>
<tr>
<td>24</td>
<td>$\lambda_{ITM</td>
<td>N_A}$</td>
<td>phagocytosis rate of ITMs by activated neutrophils</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>25</td>
<td>$\lambda_{ITM</td>
<td>AP_E\text{tiss}}$</td>
<td>phagocytosis rate of ITMs by endogenous AP in tissue</td>
<td>$3.9 \times 10^{-9}$</td>
<td>mm$^3$/molecules.min</td>
</tr>
<tr>
<td>26</td>
<td>$\lambda_{ITM</td>
<td>AP_s\text{tiss}}$</td>
<td>phagocytosis rate of ITMs by supplemented AP in tissue</td>
<td>$2.6 \times 10^{-9}$</td>
<td>mm$^3$/molecules.min</td>
</tr>
<tr>
<td>27</td>
<td>$\lambda_{ND_A</td>
<td>M_A}$</td>
<td>rate at which necrotic neutrophils are engulfed by macrophages</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

(see Table 2)
<p>|   | Rate of Engulfment of Neutrophils | ( \lambda_{ND_A|M_A} ) | rate at which apoptotic neutrophils are engulfed by macrophages | - | - | calibrated (see Table 2) |
|---|---|---|---|---|---|---|
| 28 | Permeability Constants | ( K_{eqCH} ) | concentration of pro-inflammatory cytokines that give 50% of the maximum effect of permeability | - | - | calibrated (see Table 2) |
| 29 | | ( p_{AP}^{\text{max}} ) | maximum permeability of AP | - | - | calibrated (see Table 2) |
| 30 | | ( p_{AP}^{\text{min}} ) | minimum permeability of AP | - | - | calibrated (see Table 2) |
| 31 | | ( p_{N_R}^{\text{max}} ) | maximum permeability of resting neutrophils | - | - | calibrated (see Table 2) |
| 32 | Maximum Concentration | ( ITM_{max} ) | maximum concentration of ITMs | ( 3.6 \times 10^7 ) | molecules/mm(^3) | 53 |
| 33 | | ( N_{R}^{\text{max}} ) | maximum concentration of resting neutrophils | ( 2.5 \times 10^3 ) | cells/mm(^3) | 50 |</p>
<table>
<thead>
<tr>
<th></th>
<th>of Immune Cells</th>
<th>$ACH^{max}$</th>
<th>maximum concentration of anti-inflammatory cytokines</th>
<th>$5.8 \times 10^6$</th>
<th>$\frac{molecules}{mm^3}$</th>
<th>65</th>
</tr>
</thead>
<tbody>
<tr>
<td>36</td>
<td>$M^{max}$</td>
<td>maximum concentration of Monocytes</td>
<td>$1.5 \times 10^2$</td>
<td>$\frac{cells}{mm^3}$</td>
<td>50</td>
<td></td>
</tr>
<tr>
<td>37</td>
<td>$t_{ITM}$</td>
<td>delay in generating ITMs at the onset of surgery</td>
<td>10</td>
<td>$min$</td>
<td></td>
<td></td>
</tr>
<tr>
<td>38</td>
<td>$t_{AP\text{delay}}$</td>
<td>delay at which AP is supplied into the bloodstream</td>
<td>60</td>
<td>$min$</td>
<td>correspond with Ruud Brands</td>
<td></td>
</tr>
</tbody>
</table>
| 39 | $t_{\text{suppstop}}$ | time at which bolus AP that is injected into the bloodstream is stopped | 480               | $min$                  | experiment*
| 40 | $t_{CH\text{delay}}$ | delay of cytokine recruitment                    | 20               | $min$                  | correspond with Ruud Brands |
### Chapter 1: Modeling the Human Innate Immune Response for Systemic Inflammation

<table>
<thead>
<tr>
<th></th>
<th>Parameter</th>
<th>Description</th>
<th>Value</th>
<th>Unit</th>
<th>Correspondence</th>
</tr>
</thead>
<tbody>
<tr>
<td>41</td>
<td>$t_{NDNdelay}$</td>
<td>delay of necrosis</td>
<td>300</td>
<td>min</td>
<td>correspondence with Ruud Brands</td>
</tr>
<tr>
<td>42</td>
<td>$r_{ITM}$</td>
<td>rate of increase of ITMs at the onset of surgery</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td>43</td>
<td>$r_{AP}$</td>
<td>rate of increase at which AP is supplied into the bloodstream</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td>44</td>
<td>$r_{induce}$</td>
<td>rate of induction of AP in the supplemented branch, where patients are supplied initially with 1000 IU of AP, and continuously with bolus AP</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td>45</td>
<td>$r_{NDN}$</td>
<td>rate at which activated neutrophils become necrotic</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td>46</td>
<td>$b$</td>
<td>set to maximum to signify an immediate rise/fall of concentration</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>---</td>
<td>--------------------------</td>
<td>-----------------------------------------------------------------------------------------------</td>
<td>------</td>
<td>------</td>
<td>----------------------------</td>
</tr>
<tr>
<td>47</td>
<td>Maximum</td>
<td>$r_{ITMpeak}$ peak concentration of ITMs</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td></td>
<td>Peak Saturation of</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Sigmoid Function</td>
<td>$r_{distress}$ peak concentration of AP that is supplied into the bloodstream to establish</td>
<td>-</td>
<td>-</td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>homeostasis</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>49</td>
<td></td>
<td>$r_{inducepeak}$ peak concentration of induced AP</td>
<td>-</td>
<td>-</td>
<td>calibrated, (see Table 2)</td>
</tr>
<tr>
<td>50</td>
<td></td>
<td>$r_{inject}$ peak concentration of injected bolus AP</td>
<td>-</td>
<td>-</td>
<td>experiment*</td>
</tr>
<tr>
<td>51</td>
<td>Pro-inflammatory</td>
<td>$\beta_{M_{ITM}}$ rate of pro-inflammatory cytokine production when activated macrophages</td>
<td></td>
<td></td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td></td>
<td>Cytokine Production</td>
<td>phagocytose ITMs</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>52</td>
<td></td>
<td>$\beta_{N_{ITM}}$ rate of pro-inflammatory cytokine production when activated neutrophils</td>
<td></td>
<td></td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>neutralize ITMs</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>53</td>
<td></td>
<td>$\theta_{ACH}$ rate at which anti-inflammatory cytokines inhibit pro-inflammatory cytokine</td>
<td></td>
<td></td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>production</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
### Anti-inflammatory Cytokine Production

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
<th>Value</th>
<th>Units</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \beta_{ND</td>
<td>MA} )</td>
<td>rate at which anti-inflammatory cytokines are produced when activated macrophages engulf apoptotic neutrophils</td>
<td></td>
<td></td>
</tr>
<tr>
<td>( \alpha_{ACH</td>
<td>MA} )</td>
<td>rate at which anti-inflammatory cytokines are produced by activated macrophages</td>
<td>( 1.3 \times 10^{-3} )</td>
<td>( 1/min )</td>
</tr>
<tr>
<td>( f_{dilution} )</td>
<td>dilution factor that accounts for the movement of immune cells from the bloodstream into the tissue</td>
<td>( 1/16 )</td>
<td></td>
<td>66</td>
</tr>
<tr>
<td>( \alpha_{NDN} )</td>
<td>rate of production of ITMs by necrotic neutrophils</td>
<td></td>
<td></td>
<td>calibrated (see Table 2)</td>
</tr>
<tr>
<td>( \phi_{M_R</td>
<td>ITM} )</td>
<td>rate at which resting macrophages are activated by ITMs</td>
<td>( 7.3 \times 10^{-5} )</td>
<td>( \frac{mm^3}{molecules.min} )</td>
</tr>
<tr>
<td>( r_{Nhomeo} )</td>
<td>rate of homeostasis of resting neutrophils in the bloodstream</td>
<td></td>
<td></td>
<td>calibrated (see Table 2)</td>
</tr>
</tbody>
</table>
Chapter 1 Modeling the Human Innate Immune Response for Systemic Inflammation

| 60 | Homeostasis of Neutrophils | $w$ | defines how long the concentration of neutrophils are replenished back into the bloodstream to establish homeostasis | - | - | calibrated (see Table 2) |

*concentration used in the clinical study
### 1.10.3. Parameter Estimation

A cost function quantifies how closely the output of the model agrees with the experimental data using a set of parameters $\theta$. This cost function is normally designated by the root mean of square error as shown in Equation (28):

$$
\chi^2(\theta) = \sum_{k=1}^{m} \left( \frac{\sum_{i=1}^{d} (\overline{y}_{kl}^i - \overline{y}_k(\theta, t_i))^2}{d_k} \right)
$$

(28)

where $y_{kl}^i$ corresponds to the $d$ data points for each observable $k$ (Alkaline Phosphatase, anti- and pro-inflammatory cytokines) at time points $t_i$. $d_k$ is the number of data points for observable $k$, and $y_k(\theta, t_i)$ is the model output for observable $k$ at time $t_i$ using the parameters $\theta$. We also normalized the experimental data and the model output for observable $k$ by dividing them with the maximum experimental value and model output value respectively. In this way, we assign equal contributions from each dataset to the objective function especially since they have different range of values.

$$
\overline{y}_{kl}^i = \frac{y_{kl}^i}{\max(y_{kl}^i)}, y_k(\theta, t_i) = \frac{y_k(\theta, t_i)}{\max(y_k(\theta))}
$$

(29)

The set of parameters $\theta$ can be estimated numerically by minimizing $\chi^2(\theta)$:

$$
\theta = \arg\min[\chi^2(\theta)]
$$

(30)

### 1.10.4. Parameter Optimization

We estimate model parameters by minimizing the cost function in Equation (30). Here we used the truncated Newton method, also known as Newton-Conjugate Gradient method, which solves large nonlinear optimization problems given a set constraints or boundaries for each parameter in $\theta$.

The NCG method is an iterative algorithm that searches for optimal parameters corresponding to the minimal cost function in the direction of
\( p_k^N \) as summarized in Equation (31)

\[
p_k^N = -\nabla^2 f_k^{-1} \nabla f_k
\]  

(31)

where \( p_k^N \) is the search direction, \( \nabla^2 f_k \) is the second derivative of the cost function (Hessian matrix), and \( \nabla f_k \) is the gradient of the cost function at an optimization step \( k \). However, the Hessian matrix \( \nabla^2 f_k \) may not always be positive definite and the step direction \( p_k^N \) may not always be in the descending direction. The NCG method avoids this issue using the conjugate gradient method, which terminates the iteration steps if a negative curvature of the Hessian matrix is encountered. Additionally, the NCG method is a Hessian-free method because it does not require explicit knowledge of the Hessian matrix. Rather, it only needs to calculate for the matrix-vector products between the Hessian matrix \( \nabla^2 f_k \) and the search vector \( p \), which can be solved via the finite-difference technique. A more detailed discussion of the conjugate gradient method and the finite-difference techniques can be found in 67.

Table 2. Summary of calibrated model parameters. Note that the calibration range excludes 0.

<table>
<thead>
<tr>
<th>Model Parameters</th>
<th>Unit</th>
<th>Calibrated Supplemented AP Experiment</th>
<th>Calibration Range</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 ( \beta_{MA</td>
<td>ITM} )</td>
<td>( \text{mm}^3 \text{ cells.min}^{-1} )</td>
<td>( 7.8 \times 10^{-4} )</td>
<td>(0,1]</td>
</tr>
<tr>
<td>2 ( \beta_{NA</td>
<td>ITM} )</td>
<td>( \text{mm}^3 \text{ cells.min}^{-1} )</td>
<td>( 4.8 \times 10^{-2} )</td>
<td>(0,1]</td>
</tr>
<tr>
<td>3 ( \theta_{ACH} )</td>
<td>( \text{mm}^3 \text{ molecules}^{-1} )</td>
<td>( 1 \times 10^{-10} )</td>
<td>(0,1]</td>
<td>constrained 53</td>
</tr>
<tr>
<td>4 ( \beta_{ND</td>
<td>MA} )</td>
<td>( \text{molecules.mm}^{-3} \text{ cells}^2 \text{ min}^{-1} )</td>
<td>( 9 \times 10^4 )</td>
<td>(0,10^4]</td>
</tr>
<tr>
<td>5 ( \lambda_{ITM</td>
<td>NA} )</td>
<td>( \text{mm}^3 \text{ cells.min}^{-1} )</td>
<td>( 1 \times 10^{-6} )</td>
<td>(0,10]</td>
</tr>
<tr>
<td>6 ( \alpha_{ND} )</td>
<td>( \text{molecules} \text{ cells.min}^{-1} )</td>
<td>( 1 \times 10^4 )</td>
<td>([10^2,10^4])</td>
<td>constrained 59</td>
</tr>
<tr>
<td>7 ( p_{\text{max}_AP} )</td>
<td>( 1/\text{min} )</td>
<td>( 2 \times 10^{-3} )</td>
<td>(0,1]</td>
<td>constrained (data)</td>
</tr>
</tbody>
</table>
Although we propose a model of the human innate immune response that aims at describing the dynamics of fourteen (14) immune cells and molecules that are located either in the bloodstream or in the tissue, clinical data is only available for AP, pro-inflammatory, and anti-inflammatory cytokines. Given the noise in the experimental data, the assurance of model parameters being estimated unambiguously becomes challenging. More often than not, experimental data is insufficient especially with respect to the size of the model, hence estimated parameters can be non-identifiable.

Parameter identifiability analysis addresses a crucial step in estimating...
parameters. This type of analysis tries to answer whether it is possible to recover a unique set of model parameters given a dataset. One tool that was developed to evaluate which parameters are identifiable, and hence consequently infer the feasibility of the model, was introduced by Rau et al. in 71 through a technique called **profile likelihood**. This type of approach aims at “profiling” a single parameter by fixing its value across a range of values, while fitting the remaining parameters using the objective function in Equation (28). The minimum value of the objective function constitutes the likelihood profile for the fixed parameter within the computed confidence interval derived using a threshold in the likelihood. Due to the intense computational requirements of a detailed parameter identifiability study, we performed a preliminary study (data not shown) focusing on the sensitive parameters identified in the global sensitivity analysis (see Section 1.10.7). This preliminary study shows that all sensitive parameters are either identifiable or practically non-identifiable parameters. Additionally for all practically non-identifiable parameters the issue is related to the lower boundary, which cannot be lower than 0, while the upper boundary always crosses the threshold in the likelihood.

### 1.10.6. Difference between Experimental Data and Model Output

We show the normalized residuals between experimental data and results of the model simulation for Alkaline Phosphatase, pro-inflammatory, and anti-inflammatory cytokines by calculating the difference between experimental data and model output divided by the experimental data. Normalized residual values are at maximum between 0-5 hours after surgery time for cytokines (see A-B). Considering the large standard deviation observed within the 0-5 hours post-surgery timeframe on the cytokines dataset that in fact spans 7 orders of magnitude in units cells/m³, our results still fall within this expected range. Note that patients’ blood parameters usually stabilize to normal concentrations within this time period after surgery.
Normalized residuals for AP are only at maximum at the onset of surgery. This is as expected because right at the start of surgery, patients are injected with bovine AP concentration of 1000 IU. Bovine AP has a short half life and could disappear within 20 minutes. However, the earliest recorded AP concentration from the experimental data is 30 minutes after the start of operation. Therefore, we do not see a trace of this added bovine AP on the data, giving a residual that is close to the initial concentration of AP given to the patient.

Figure 9. Normalized Residuals (Experimental Data–Predicted Value/Experimental Data) for A) Pro-inflammatory Cytokines, B) Anti-inflammatory Cytokines, and C) Alkaline Phosphatase.

1.10.7. Sensitivity Analysis

In order to obtain a deeper understanding of the model dynamics, it is important to quantify how the variability of model parameters affects the predictions of the model. For this reason, we perform a global sensitivity
analysis for three of the most significant model outputs: Alkaline Phosphatase, pro-inflammatory cytokines, and anti-inflammatory cytokines for which we have the experimental data. The strength of variance-based measures of sensitivity lies on the fact that they measure sensitivity across the entire input space and could deal with nonlinear responses and measure the effect of interactions in non-additive systems. We chose the Fourier amplitude sensitivity test (FAST) due to its computational efficiency.

The ranges of parameter values were varied in a range of biologically justified values given in Table 2. The 22 parameters of the model were then sampled using a periodic sampling approach with the number of parameter samples $N = 5000$ to ensure an adequate coverage of the multidimensional parameter space. We analyzed the sensitivity of the model by plotting the sensitivity indices with 95% confidence of the FAST method (see Figure 10).

We show that concentrations of pro-inflammatory cytokines are most sensitive to $\theta_{\text{ACH}}$, $\beta_{\text{MA}|\text{ITM}}$, $\beta_{\text{ND}|\text{MA}}$, and $\tau_{\text{NDN}}$ (see Figure 10A) in various time intervals. $\tau_{\text{NDN}}$ only becomes sensitive 30 minutes after the onset of surgery. This is due to the delay in the recruitment of neutrophils into the tissue. The sensitivity indices for $\beta_{\text{MA}|\text{ITM}}$, $\theta_{\text{ACH}}$, on the other hand peak right after the onset of surgery and decrease beyond this time interval. These parameters are sensitive because they are incorporated in the model Equation (18) which regulates the concentration of pro-inflammatory cytokines. The interactions between activated neutrophils and ITMs also trigger the recruitment of pro-inflammatory cytokines. Hence, the term $\tau_{\text{NDN}}$, which controls the activation rate of resting neutrophils in Equation (23), turns out to be sensitive as well.

$\beta_{\text{ND}|\text{MA}}$, $\beta_{\text{MA}|\text{ITM}}$, $\theta_{\text{ACH}}$, $\lambda_{\text{NDN}|\text{MA}}$, $K_{\text{eqCH}}$, and $P_{\text{NR}}^{\text{max}}$ are the most sensitive parameters among those investigated with respect to the concentration of anti-inflammatory cytokines as shown in Figure 10B. The first three parameters are the same for both pro-inflammatory cytokines and anti-inflammatory cytokines because these two immune cells depend on each other. However, for anti-inflammatory cytokines the sensitivity is more
pronounced as the anti-inflammatory response continues for several hours after surgery.

Finally, the parameters shown to be most sensitive for the production of endogenous AP in the bloodstream are $w$, $r_{AP}$, $r_{distress}$ and $p_{AP}^{min}$. These parameters have direct effect on the concentration of endogenous AP as can be seen in Equation (15) of the model. The sensitivity of $p_{AP}^{min}$ to endogenous AP shows an initial peak within the first hour, then it goes down and rises 40 minutes after surgery. Note that bolus AP is continuously injected up until 8 hours after surgery. Therefore, the effect of adjusting the minimum permeability of AP ($p_{AP}^{min}$) that allows the transfer from the bloodstream into the tissue is also minimal within this time period.
1.10.8. Human innate immune system model without the induction mechanism of TNAP

HIIS model without the induction term predicts exactly the same dynamics for ITMs in plasma, ITMs in tissue and neutrophils in both branches as shown in Figure 11. Hence, without the added induced AP, immune cells in the supplemented branch exhibit the same dynamics as in placebo branch.
Figure 11. Dynamics of (A) ITMs in Plasma, (B) ITMs in Tissue, (C) Resting Neutrophils, (D) Activated Neutrophils, (E) Apoptotic Neutrophils and (F) Necrotic Neutrophils in the tissue. Dynamics of immune cells.
are similar for supplemented bIAP and placebo branches.

1.10.9. Conversion of Concentrations

**Alkaline Phosphatase**

\[ AP \left( \frac{IU}{L} \right) = AP \left( \frac{molecules}{mm^3} \right) \times \frac{1\,\text{mole}}{6.02 \times 10^{23}\,\text{molecules}} \times \frac{1\,\text{mole}}{\text{Molecular Weight}_{AP}\,(g)} \times \frac{\text{Activity}_{AP}(IU)}{1 \times 10^{-3} \,g} \times \frac{1 \times 10^6 mm^3}{1\,L} \]  

(32)

where \( \text{Molecular Weight}_{AP} = 160\,kg/mole \)

**Cytokines**

\[ \text{cytokine}(\frac{pg}{ml}) = \text{cytokine}(\frac{molecules}{mm^3}) \times \frac{1\,\text{mole}}{6.02 \times 10^{23}\,\text{molecules}} \times \frac{1\,\text{mole}}{\text{Molecular Weight}_{cytokine}\,(g)} \times \frac{1 \times 10^{12}pg}{1\,g} \times \frac{1 \times 10^6 mm^3}{1\,L} \times \frac{1\,L}{1 \times 10^3 \,ml} \]  

(33)

where \( \text{Molecular Weight}_{cytokine} = 21\,kg/mole \)

1.10.10. Supplementary Information References


6. Price CP. Multiple Forms of Human Serum Alkaline Phosphatase: Detection and


