

A hybrid individual-based mathematical model to study bladder infections

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## EDITED BY

Esteban A. Hernandez-Vargas,  
University of Idaho, United States

## REVIEWED BY

Babita K. Verma,  
Johns Hopkins University, United States  
Florian Wagenlehner,  
University of Giessen, Germany

## \*CORRESPONDENCE

Ruth Bowness  
✉ rb2250@bath.ac.uk

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# A hybrid individual-based mathematical model to study bladder infections

Anas Lasri Doukkali<sup>1</sup>, Tommaso Lorenzi<sup>2</sup>, Benjamin J. Parcell<sup>3</sup>,  
Jennifer L. Rohn<sup>4</sup> and Ruth Bowness<sup>5\*</sup>

<sup>1</sup>Schools of Medicine and Mathematics and Statistics, University of St Andrews, St Andrews, United Kingdom, <sup>2</sup>Department of Mathematical Sciences, Politecnico di Torino, Torino, Italy, <sup>3</sup>Medical Microbiology, NHS Tayside, Ninewells Hospital and Medical School, Dundee, United Kingdom, <sup>4</sup>Division of Medicine, Department of Renal Medicine, Centre for Urological Biology, University College London, London, United Kingdom, <sup>5</sup>Department of Mathematical Sciences, Claverton Down, University of Bath, Bath, United Kingdom

**Introduction:** Bladder infections are common, affecting millions each year, and are often recurrent problems.

**Methods:** We have developed a spatial mathematical framework consisting of a hybrid individual-based model to simulate these infections in order to understand more about the bacterial mechanisms and immune dynamics. We integrate a varying bacterial replication rate and model bacterial shedding as an immune mechanism.

**Results:** We investigate the effect that varying the initial bacterial load has on infection outcome, where we find that higher bacterial burden leads to poorer outcomes, but also find that only a single bacterium is needed to establish infection in some cases. We also simulate an immunocompromised environment, confirming the intuitive result that bacterial spread typically progresses at a higher rate.

**Conclusions:** With future model developments, this framework is capable of providing new clinical insight into bladder infections.

## KEYWORDS

mathematical, model, individual-based, simulation, bladder, infection, *Escherichia coli*

## 1. Introduction

Urinary tract infections (UTIs) are infections affecting the urethra, bladder and kidneys. Lower UTIs affect the bladder and urethra, and upper UTIs affect the kidneys. Simple or uncomplicated UTIs are caused by a number of pathogens in people with a normal urinary tract and kidney function, and no predisposing co-morbidities. They can often be self-limiting or readily cleared with a short course of antibiotics. One in four uncomplicated UTIs patients, however, experience persistent or recurrent infection and/or treatment failure, and complicated UTIs carry an increased likelihood of such outcomes [1].

UTIs are some of the most common bacterial infections, with an infection toll of 150 million people each year worldwide [2]. UTIs are a significant cause of morbidity, particularly affecting females, older men and infant boys [1]. Ramifications of such infections can lead to pyelonephritis with sepsis, renal damage in young children, pre-term labor in pregnant women and complications that arise from frequent antimicrobial use (for example, antibiotic resistance). Patients who suffer from symptomatic UTI are normally treated with the use of antibiotics. Currently, the antibiotics most commonly used include Nitrofurantoin, Trimethoprim, Cotrimoxazole (a combination of Trimethoprim and Sulfamethoxazole), Fosfomycin or Nitrofurantoin. However, the rise we have seen in antibiotic resistance in addition to the highly recurrent profile of the infection means healthcare professionals may have to use alternative lines of treatment, which have greater risk of infection with *Clostridioides difficile*. Extended-spectrum beta-lactamase (ESBL)-producing Enterobacterales are increasing and these bacteria are often multi-drug resistant. With limited treatment options and a lack of new antibiotics being developed it is essential to focus on prevention of UTIs and gain more

understanding of the cellular and molecular dynamics of how uropathogens adhere, colonize and adapt to the nutritionally lacking environment of the bladder; elude the immune system; persevere and spread through the urinary tract [3–5].

UTIs are highly recurrent and may persist through different rounds of treatments. This recurrence is due to either relapse (symptomatic recurrent UTIs with the same organism following adequate therapy) or re-infection (recurrent UTIs with previously isolated bacteria after treatment and with a negative intervening urine culture, or a recurrent UTI caused by a second bacterial isolate). In the majority of cases however, recurrent UTIs are thought to represent reinfection with the same organism. UTIs are caused by both Gram-negative and Gram-positive bacteria, and in some cases by specific fungi, but the majority are caused by the bacterial species Uropathogenic *Escherichia coli* (UPEC) [1]. Although the causes of recurrence are multifactorial—with at least a dozen mechanisms, some acting in tandem—one key factor could be the ability of UPEC to establish quiescent intracellular reservoirs (QIRs). Thus, the establishment of these bacterial communities at deeper levels of the urothelium may allow them to evade immune cell interaction and other mechanisms such as bacterial shedding.

The bladder has a number of strategies to protect itself against microbial invaders. First and foremost, the urinary tract has anatomical barriers, alongside the flushing action of urination, which usually suffice to prevent pathogenic elements from entering the tract and establishing a persistent infection. The presence of virulence factors in some bacteria, however, can help them to bypass these barriers and successfully establish infection. Virulence factors are any set of characteristics that will help bacteria in eluding the primary line of defense, and hence increase the risk of successful infection [1]. A crucial step during the infection process of a number of species of UTI pathogens, including the most common uropathogen, *Escherichia coli* (*E. coli*), is the ability to permeate and invade the bladder epithelial barrier and seek refuge within the bladder epithelial cells (BECs). This is normally done by hijacking the capacity of the epithelial cells to regulate bladder volume [6]. Another important thing to note is, due to the presence of toxic compounds in urine, the bladder must maintain a tight epithelial barrier. Thus, any immune reaction has to carefully balance the response to microbial challenge with the need to maintain the structural integrity of the epithelial barrier [7]. This will sometimes lead to premature termination of the anti-inflammatory response, resulting in chronic or recurrent infections due to residual bacteria. The innate immune response refers to the non-specific defense mechanisms that are deployed by the immune system usually hours after a pathogen is established. This is particularly important in bladder infections, as the role of the adaptive immune system in UTI is less straightforward, and is still being clarified [8]. The innate immune system is composed of various recruited and resident cells that express a wide range of pattern recognition receptors (PRRs). These receptors allow for early recognition of the pathogen which will be translated to induce a quick and robust pro-inflammatory response [9]. As noted before, although these immune responses are important, they have to be controlled to avoid any substantial damage to the epithelial barrier. As a last line of defense, the bladder sheds cells from its internal walls to reduce bacterial load. These host shedding mechanisms are modulated to avoid adverse effects [10, 11]. Below we review the main innate immune cells that operate in the urinary tract and focus on their unique antimicrobial activities.

## 1.1. Bladder epithelial cells

These cells, lining the interior of the urinary tract, will deploy the first line of defense against the infection, with the secretion of factors, both pro-inflammatory cytokines and antibacterial agents. The first cytokines detected after infection are IL-1, IL-6, and IL-8 [6, 9, 12]. This group of cytokines is important for the recruitment of phagocytes into the infected bladder [13]. Additional factors secreted by epithelial cells hinder bacterial growth by eliminating growth factors present in urine. Following bacterial infection of BECs, the pathogens are then encapsulated in RAB27b+ fusiform vesicles with exocytic properties, meaning BECs will be able to expel intracellular bacteria back into the extracellular medium [14]. If any bacteria avoids expulsion, by escaping the RAB27b+ vacuoles, it is then met with a second wave of elimination in which such pathogens are recognized and captured by autophagy. If BECs become heavily infected, a preventive measure is the shedding of the superficial epithelial cell layer through the urine, as mentioned above [10]. This leaves the underlying tissue exposed to urine toxins and *de novo* infection of deeper layers, so the shedding of BECs is immediately followed by a shift of the urothelium to a highly active and proliferative phase to restore the previous loss *via* replacement by cells underneath that differentiate and take their place [15].

## 1.2. Neutrophils

Neutrophils are the first type of immune cells to be recruited to the bladder. Initially, they respond to CXC-chemokine ligand 1 (CXCL1) and other chemo-attractants produced by superficial epithelial cells, macrophages and mast cells [16]. In mouse experiments, neutrophils are detected in urine as early as 2 h post-infection, with their peak at approximately 6 h post-infection [17]. The number of neutrophils is proportional to that of bacteria in the urinary tract [6].

## 1.3. Macrophages

A substantial population of macrophages resides in the submucosa of the urinary tract, and more cells are recruited to these sites following infection. Once activated, macrophages produce important chemokines and cytokines that regulate the activity of these and other immune cells in the vicinity, hence, influencing the timing and the intensity of inflammatory responses during UTIs [18]. Communication between bladder macrophages leads to mobilization of neutrophils into the epithelium and their subsequent activation alongside neighboring immune cells. These responses ensure efficient bacterial clearance whilst minimizing unnecessary and damaging inflammation.

## 1.4. Mast cells

Mast cells are also resident immune cells located underneath the uroepithelium adjacent to the blood vessels and lymphatic vessels that traverse the mucosal region. These type of cells have a pivotal sentinel and key immunomodulatory role during UTIs, which is partly due to

their ability to release many pre-stored inflammatory mediators upon activation. These chemicals are kept within cytoplasmic granules and gradually secrete the chemical mediators after being released extracellularly [19–21]. Although it is currently unknown what causes the activation of bladder mast cells when the epithelium is still intact, it is believed that factors released from responsive epithelial cells such as ATP, LL-37 and IL-33 could contribute to this process. The number of mast cells in the bladder increases during bladder infection, which points to an important dynamic role for these cells in the fight against the infection. Although mast cells are considered pro-inflammatory immune cells, experiments in mice show that once the infection progresses, such cells adopt anti-inflammatory responses (by releasing cytokines, IL-10) [22]. This switch in function seems to happen in parallel with the breaking down of the epithelial barrier and could facilitate epithelial regeneration.

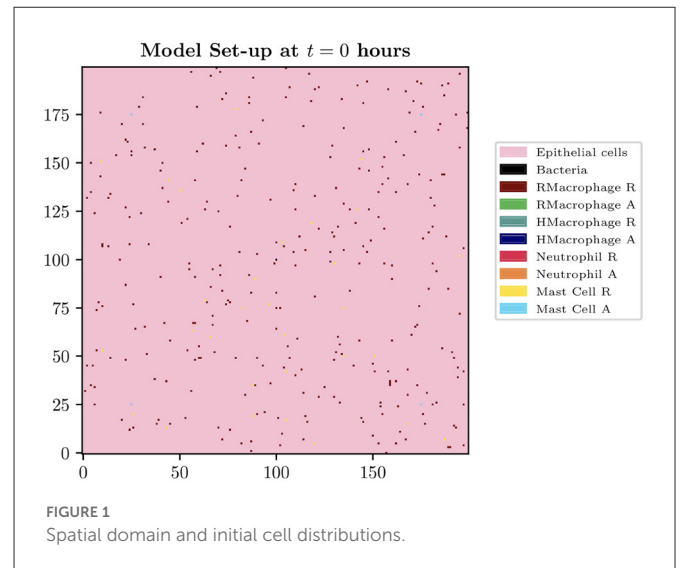
Although previous mathematical frameworks have been developed to model the role of bladder hydrodynamics in bacterial elimination in lower UTIs [23], we are not aware of any studies that describe spatio-temporal models of the bladder. We have built an individual-based mathematical model capable of simulating discrete bacteria and immune cells, as well as the diffusion of relevant cytokine molecules, which crucially considers a spatial domain. This spatial model is capable of investigating the potential for the bacteria to evade bacterial shedding and treatment by establishing QIRs and hence can assess to what extent this mechanism is responsible for infection recurrence. In Section 2 we describe the mathematical model, with all “agent” rules explained and differential equations outlined. Section 3 explores the results from model simulations before we make some concluding comments and outline future research in Section 4.

## 2. Methods

We have developed a simple individual-based mathematical model, *UTImodel*, to simulate an infection in a section of the bladder. Individual-based models simulate individual elements or ‘agents’ on a computational grid. The grid describes the environment, here the bladder, and the model includes rules to describe agent-agent and agent-environment interactions. These interactions between agents and between some agents and their environment can be quite complex; not only can these interactions change with time but so can the strategies used to decide what action will be employed at any particular time. These actions are generally constituted by the exchange of information which as a result, agents can either update their internal state or take other actions [24]. Individual-based modeling, in contrast to many other modeling techniques, allows for heterogeneity in population and environment [25, 26]. Our model is a hybrid model; as well as discrete agents (bacteria and immune cells), it also contains a generic chemokine molecule that acts as a chemoattractant, directing the immune cells to the site of infection. This generic chemokine is modeled *via* a Partial Differential Equation (PDE).

### 2.1. Model environment

The spatial domain is formed by the innermost bladder layer, the urothelium, where the two-dimensional model domain is of size



$L_g \times L_g$ , where  $L_g = 200$  for the simulations presented in this paper. The size of each grid cell within our environment is constructed in order to contain the largest agent within the biological system to be described, the macrophage. The diameter of a human macrophage is approximately  $20 \mu\text{m}$ , hence each grid cell will be equivalent to a  $400 \mu\text{m}^2$  section of the bladder. We therefore model over a  $16 \text{mm}^2$  section of the superficial urothelium. We choose this small section of the bladder to investigate as we want to focus on the beginning of a bladder infection and believe this size is sufficient to study the initial growth and immune response. Each grid cell will represent a location instance within our model which can adopt the following states: empty, occupied by immune cell, bacteria or blood vessel. Although blood vessels do not appear on the urothelium in reality, we assume that these are the spatial locations of the blood vessels at deeper levels. Future iterations of the model will study a three-dimensional space where we can simulate the representation of the vasculature more realistically. On our 2-D spatial domain, we fix grid cells to be locations to represent cross-sections of 4 blood vessels, with locations [(175, 25), (25, 175), (175, 175), (25, 25)]. We place bacteria in the center of the domain. See Figure 1 to see the spatial domain and initial cell distributions for our simulations.

### 2.2. Agents and rules

For our simple model, we define five types of agents: bacteria, resident macrophages ( $LY6C^-$ ), helper macrophages ( $LY6C^+$ ), neutrophils and mast cells. We have chosen not to include BECs as agents within this version of the model, in order for computational simplicity. This will be included in future work. The interaction between such agents will be set by specific rules defined within our model. Functions for all these interactions are stored in separate classes in the code created for each specific agent.

#### 2.2.1. Bacteria

In our model, we simulate UPEC bacteria. Initially,  $I_B$  bacteria are seeded at the center of the grid. We place bacteria in the center of the

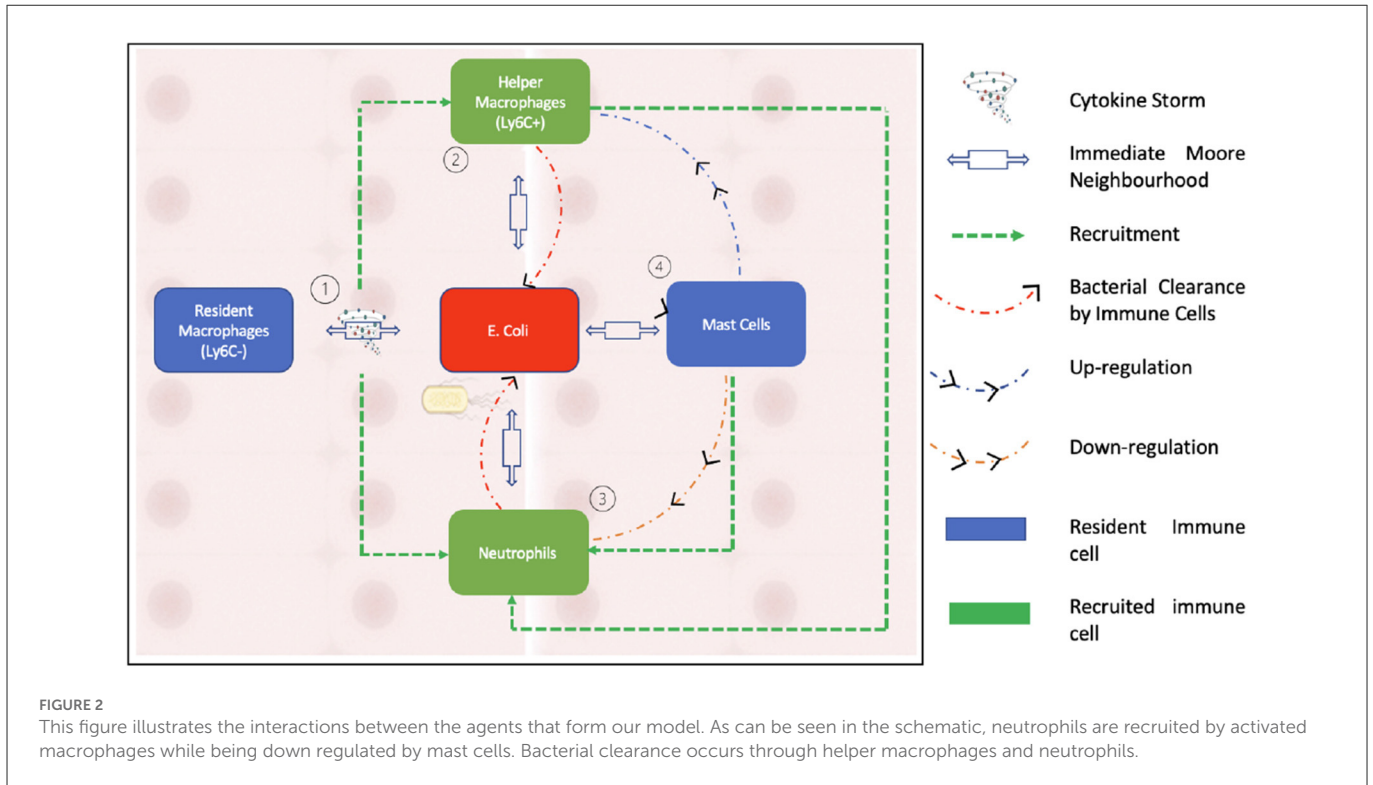
TABLE 1 Table of model parameter values.

Symbol	Parameter description	Values	Units	Source
<b>Bacteria</b>				
$I_b$	Minimal infectious dose	1–1,000	Unitless	E
$B_{sp}$	Bacterial replication rate	0.37–0.97	Per hour	[29], [10]
$N_{db}$	Neighborhood replication depth	2	Unitless	E
$S_B$	Bacterial load for shedding	6,000	Unitless	E
$P_{Bp}$	Prob bacteria penetrates bladder wall	0.156	Unitless	E
<b>Resident macrophages</b>				
$I_{rM}$	Initial No. resident macrophages	500	Unitless	E
$rM_{ls}$	Resident macrophages lifespan	30–32	Hours	[36]
$rM_{RhM}$	Resident macrophages recruitment prob.	0.04–0.08	Unitless	E
$rM_{RN}$	Resident macrophages movement rate	0.17–0.26	Per hour	[37]
$rM_{Ap}$	Resident macrophages activation prob.	0.261	Unitless	E
<b>Helper macrophages</b>				
$hM_{ls}$	Helper macrophage lifespan	30–32	Hours	[36]
$hM_{RhM}$	Helper macrophages recruitment prob.	0.007–0.031	Unitless	E
$hM_{RN}$	Helper macrophages movement rate	0.13–0.18	Per hour	E
$hM_{Ap}$	Helper macrophages activation prob.	0.129	Unitless	E
$hM_{pp}$	Helper macrophages killing prob.	0.12–0.13	Unitless	E
<b>Neutrophils</b>				
$N_{ls}$	Neutrophil lifespan	30–32	Hours	[38]
$N_{RN}$	Neutrophil movement rate	0.12–0.23	Per hour	E
$N_{RhM}$	Neutrophil recruitment prob.	0.07–0.14	Unitless	E
$N_{Ap}$	Neutrophil activation prob.	0.148	Unitless	E
$N_{Php}$	Neutrophil killing prob.	0.11–0.17	Unitless	E
<b>Mast cells</b>				
$I_{MC}$	Initial No. mast cells	50	Unitless	E
$MC_{ls}$	Mast cell lifespan	30–35	Hours	E
$MC_{RN}$	Mast cell movement rate	0.14–0.26	Per hour	E
$MC_{urM}$	Mast cell recruit. prob.	0.003–0.033	Unitless	E
$MC_{Ap}$	Mast cell activation prob.	0.261	Unitless	E

Where the source is given as E, this means it is estimated.

domain as our computational environment represents a section of a initially healthy bladder, which is then seeded with bacteria, in order to investigate bacterial growth without the infection approaching the computational boundary. This set-up is also optimal for using experimental results to verify model output, which is our intention in future work. Initial bacterial placement can easily be changed to either random allocation or to be located at the boundaries, which again is intended to be studied in future work. The bacteria are able to replicate according to a growth rate of  $B_{sp}$ . See Table 1 for these parameter values, with corresponding references to the experimental data these are based on. Bacteria can be found in either a replicating or “resting” state, where bacteria are deemed to be resting when there is no space on the computational grid for them to replicate into. At each time point, the neighborhoods are

checked and when empty spaces in the proximity of the bacteria are available, the bacteria are able to switch back to a replicating state and spread across the domain. Bacterial replication within our model is assumed to have a neighborhood depth of  $N_{db} = 2$ . Hence, if bacteria are able to replicate, an empty location within the specified depth is chosen and populated with a new bacterium. If all locations are occupied, the replication state is switched temporarily to resting. The neighborhood type is alternated between Moore and von Neumann at each time step, in order to more accurately model bacterial cluster shapes. A Moore neighborhood consists of the eight cells that surround the cell of interest, whereas a von Neumann neighborhood contains the four adjacent grid cells to the cell of interest, i.e. above, below, right and left. Again, in future model developments a 3-D version of this domain will be studied to simulate



the environment more realistically, where bacterial growth will be less restricted.

### 2.2.2. Macrophages

Macrophages are represented as both resident and recruited discrete agents on the grid. Only one macrophage can occupy each grid cell and, independently of their nature (resident or recruited), all macrophages have the following attributes: position, age and state (resting or activated). Such characteristics are updated at every model time step.  $I_{rM}$  resident (initially resting) macrophages are seeded onto the grid at the start of the simulation. Macrophages are randomly assigned an age  $rM_{Is}$  or  $hM_{Is}$  for resident or helper macrophages, respectively. Once a macrophage reaches its assigned age, it is removed from the grid. If a resident macrophage is in the Moore neighborhood (of order 2) of a bacterium, they are activated with probability  $rM_{Ap}$ . Once activated,  $Ly6C^+$ /helper macrophages are recruited into the domain with probability  $rM_{RhM}$ , and placed in the neighborhood of the activated macrophage. As well as recruiting helper macrophages, activated macrophages also secrete a generic chemokine molecule, the dynamics of which are described in Section 2.3. Macrophages move along the gradient of this chemoattractant, directing the immune cells to the site of infection. When no chemokine is present in the spatial domain, the macrophages follow a random walk. The movement rates are described by  $rM_{RN}$  and  $hM_{RN}$  for resident and helper macrophages, respectively.  $Ly6C^+$ /helper macrophages are able to kill bacteria: once a bacterium is in the immediate neighborhood (Moore neighborhood of order 1) of an activated macrophage, this will kill said bacterium with probability  $hM_{phP}$ .

### 2.2.3. Neutrophils

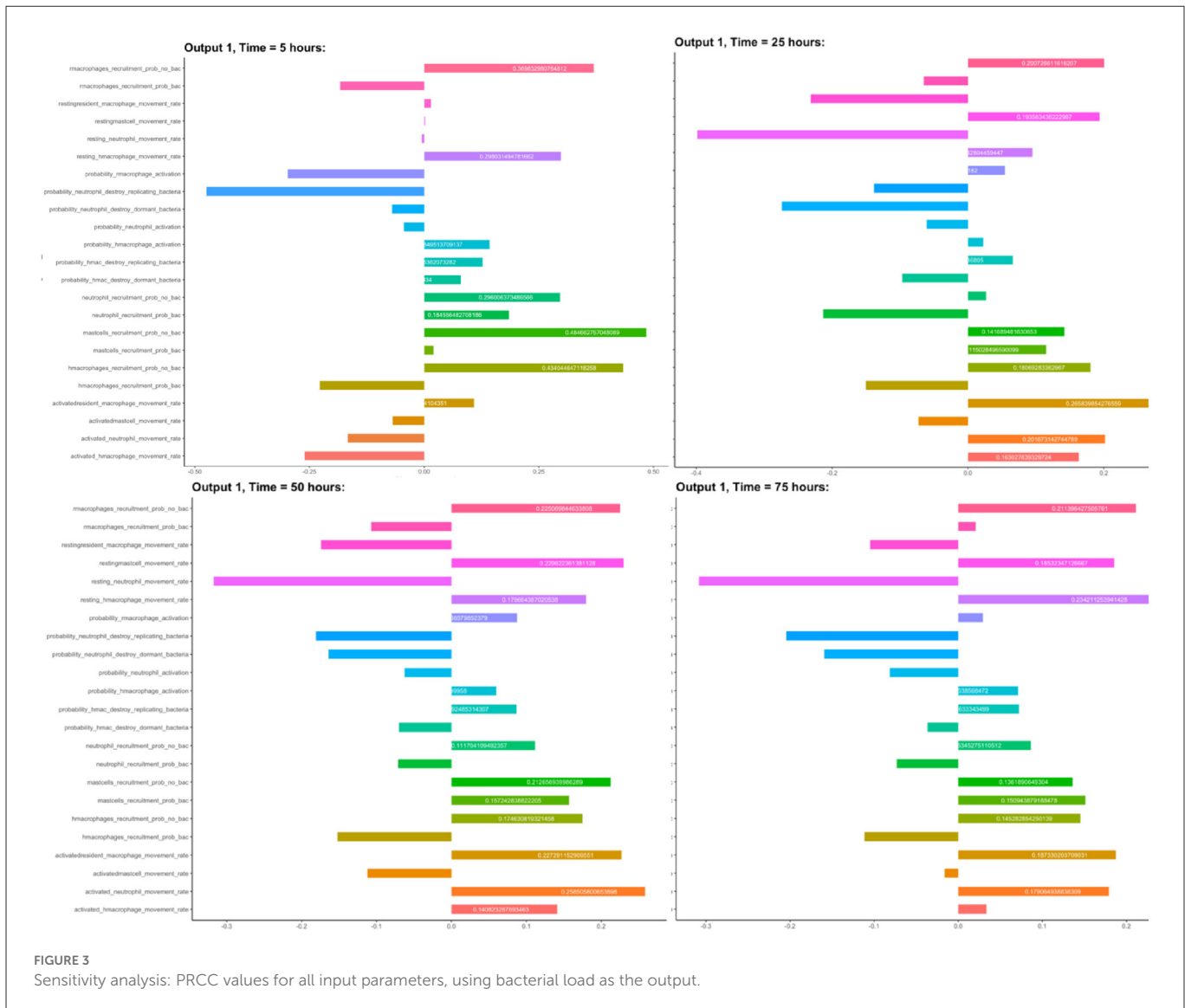
The lifespan of the neutrophils are set in the model as  $N_{Is}$ . They are recruited by activated macrophages, similarly to helper macrophages, with probability  $N_{RhM}$ . The neutrophils move chemotactically in the same manner as all immune cells in the system, up the gradient of the chemoattractant, or as a random walk when no chemokine is present. They move at rate  $N_{RN}$ . Neutrophils can kill bacteria in their immediate neighborhood with probability  $N_{phP}$ . Neutrophils are down-regulated by mast cells (see Section 2.2.4 for details).

### 2.2.4. Mast cells

There are  $I_{MC}$  mast cells initially seeded on the grid. Their lifespan is set to  $MC_{Is}$ , they move at rate  $MC_{RN}$  and are recruited through the blood vessels with probability  $MC_{urM}$ . They are activated with probability  $MC_{Ap}$ . Mast cells are responsible for up-regulating helper macrophages, aiding with the clearance of bacterial infection. When there is a bacterium in the immediate neighborhood (Moore neighborhood of order 1) of a mast cell and no helper macrophages are present within this same neighborhood, the mast cell recruits a helper macrophage to a random location within this neighborhood.

Mast cells are also known to down-regulate the recruitment of neutrophils [22, 27] in order to limit excessive damage to the epithelial tissue [28]. When a mast cell is in an immediate neighborhood of three or more neutrophils, it removes one (at random) from the grid.

Figure 2 summarizes the interactions of the agents within our model.



### 2.3. Chemokine dynamics

A generic chemokine is secreted two-dimensionally by resident macrophages upon activation. This chemokine serves as a chemoattractant, directing all immune cells in the system to the site of infection. The spatiotemporal evolution of the chemokine concentration,  $C(x, t)$ , is described by the following PDE:

$$\frac{\partial C(x, t)}{\partial t} = \nabla \cdot (D(x)\nabla C(x, t)) + s(x, t) - (u(x, t) + d(x, t))C(x, t), \quad (1)$$

Which is subject to the initial condition  $C(x, 0) = 0$ . Here,  $D$  denotes the spatially-dependent diffusion coefficient,  $s$  denotes the source of chemokine from the activated macrophages,  $u$  denotes the uptake of chemokine by the host cells, and  $d$  denotes the extracellular decay.

### 2.4. Uncertainty and sensitivity analysis

Model parameter values were guided by available experimental data. There is always some uncertainty surrounding parametrisation of models such as these. To mitigate this, we carried out uncertainty and sensitivity analysis (UA/SA) for all parameters shown in Table 1 to have a range of values. This was conducted in three stages:

1. Once we searched the experimental literature to find suitable ranges for parameter estimates, we generated our sample space using Latin Hypercube Sampling (LHS). The output of this first stage is a data matrix consisting of all the data samples to be fed into our model. Here we introduced a testing framework using YAML (Yet Another Markup Language), which is a data serialization language that allows us to develop a framework using python, C++ and Rstudio. We use Rstudio to generate our LHS samples which are then imported through python. Python is then responsible for taking in a data frame and outputting  $N_s$  YAML files, each containing a separate parameter set which, through a newly developed python module, we are able to feed to the C++ model.

- 2. The second stage includes sorting simulation output and plotting simulation results. We simply analyse the model data here. By plotting various outputs, we can visualize the results from the whole parameter space.
- 3. We then carry out the sensitivity analysis using Rstudio. We utilize the global SA technique Partial Rank Correlation Coefficient (PRCC), which works by establishing nonlinear but monotonic relationships between model inputs and outputs.

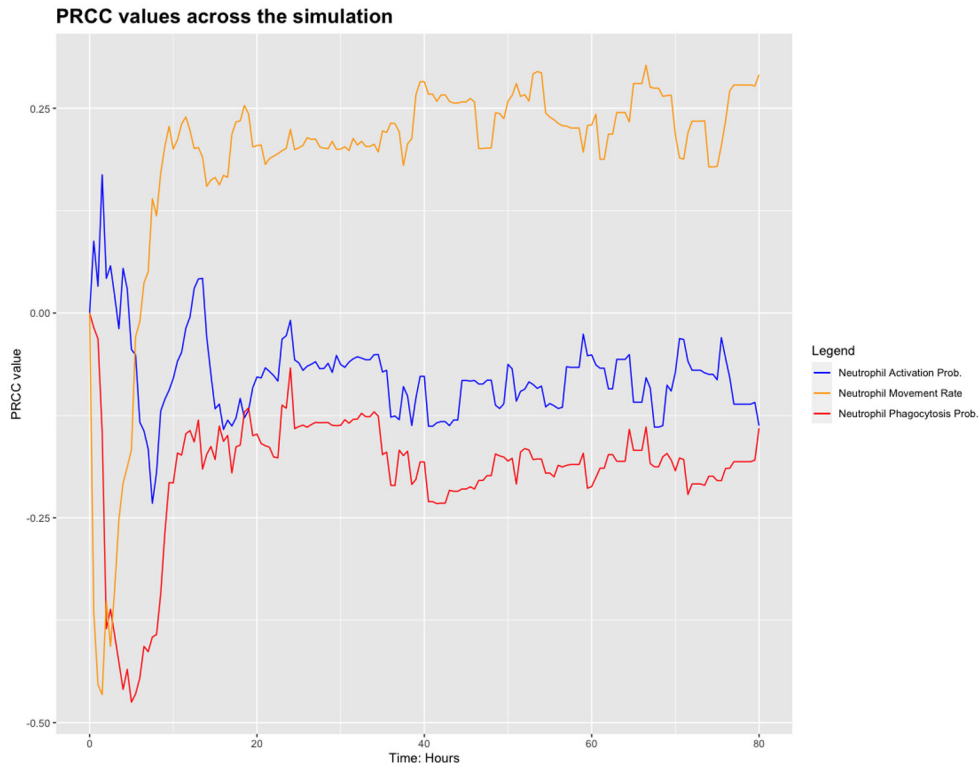


FIGURE 4 PRCC values for neutrophil parameters.

Early stage: Infection Progression Profiles,  $I_B = 1$

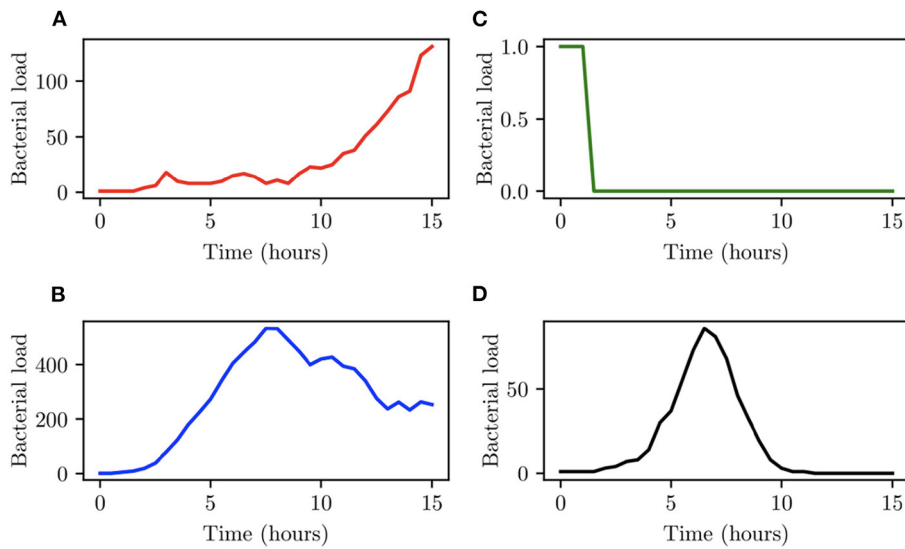
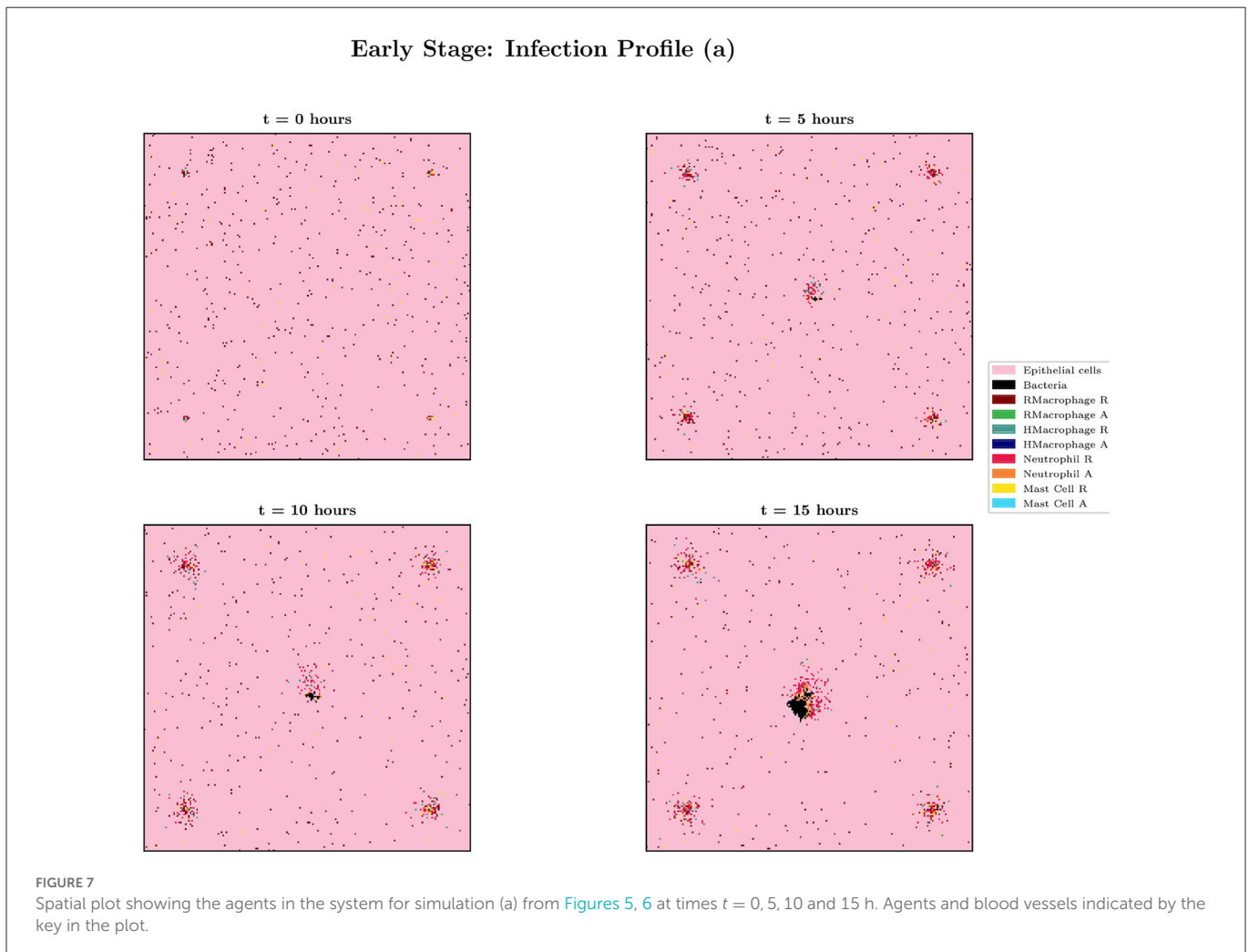
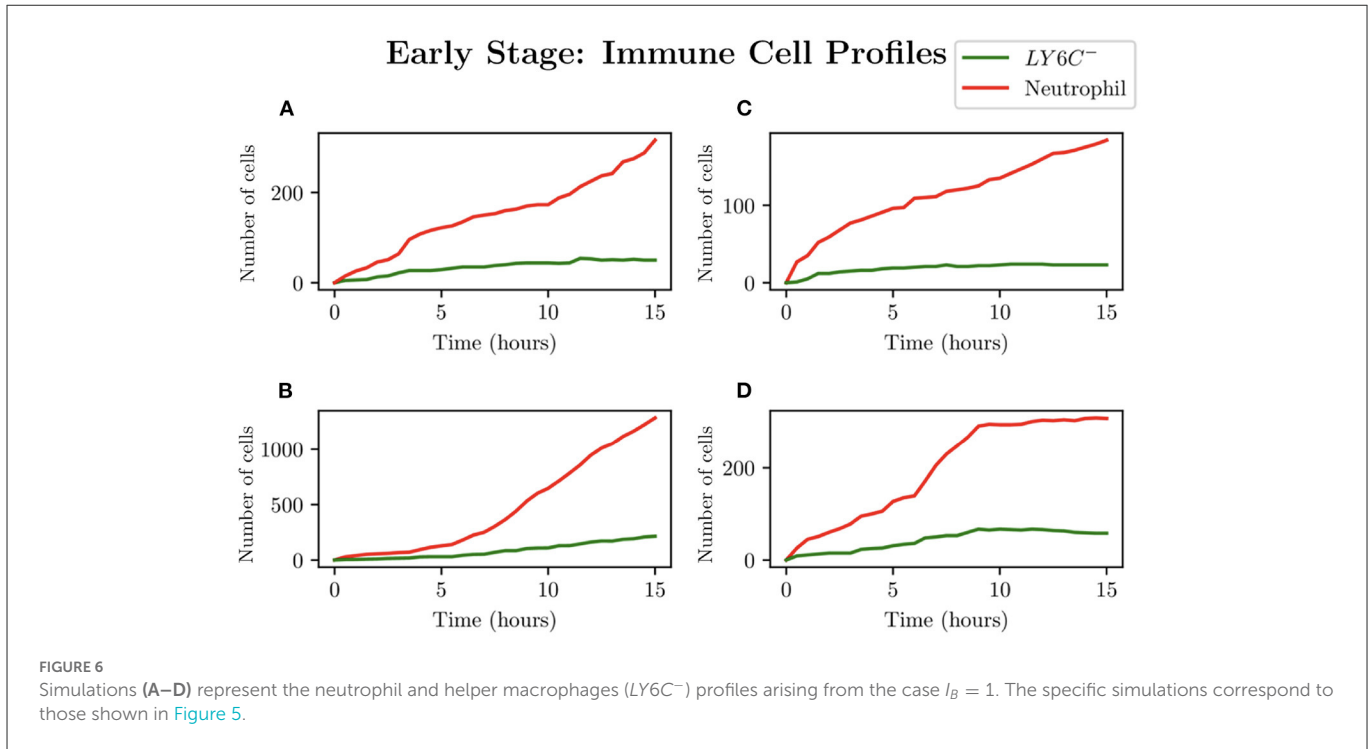
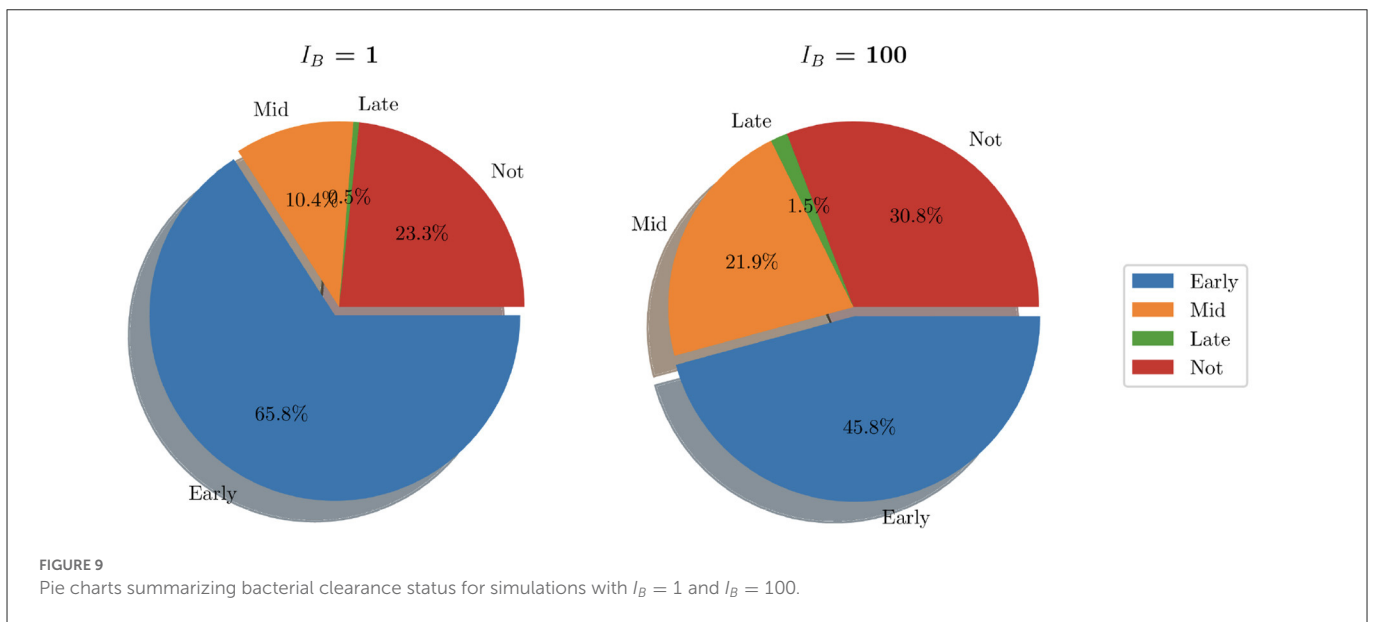
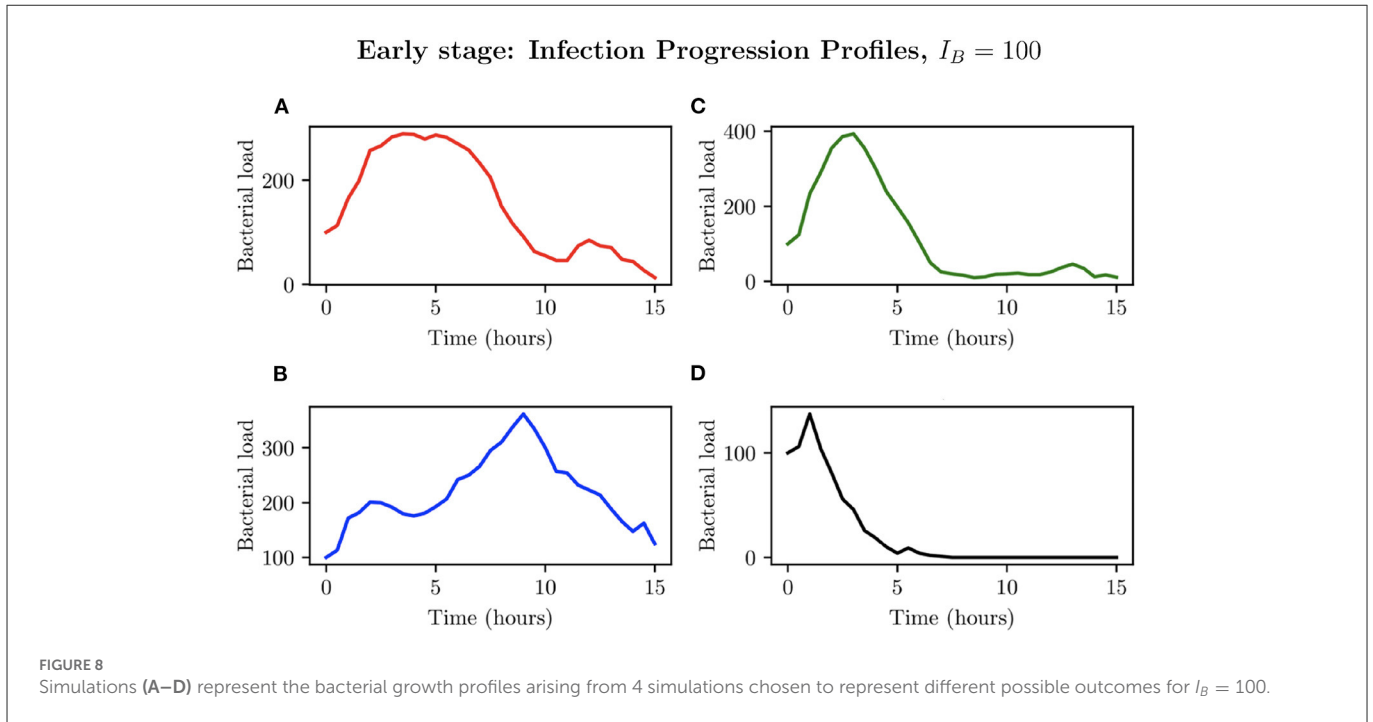


FIGURE 5 Simulations (A–D) represent 4 different bacterial growth profiles arising with  $I_B = 1$ .





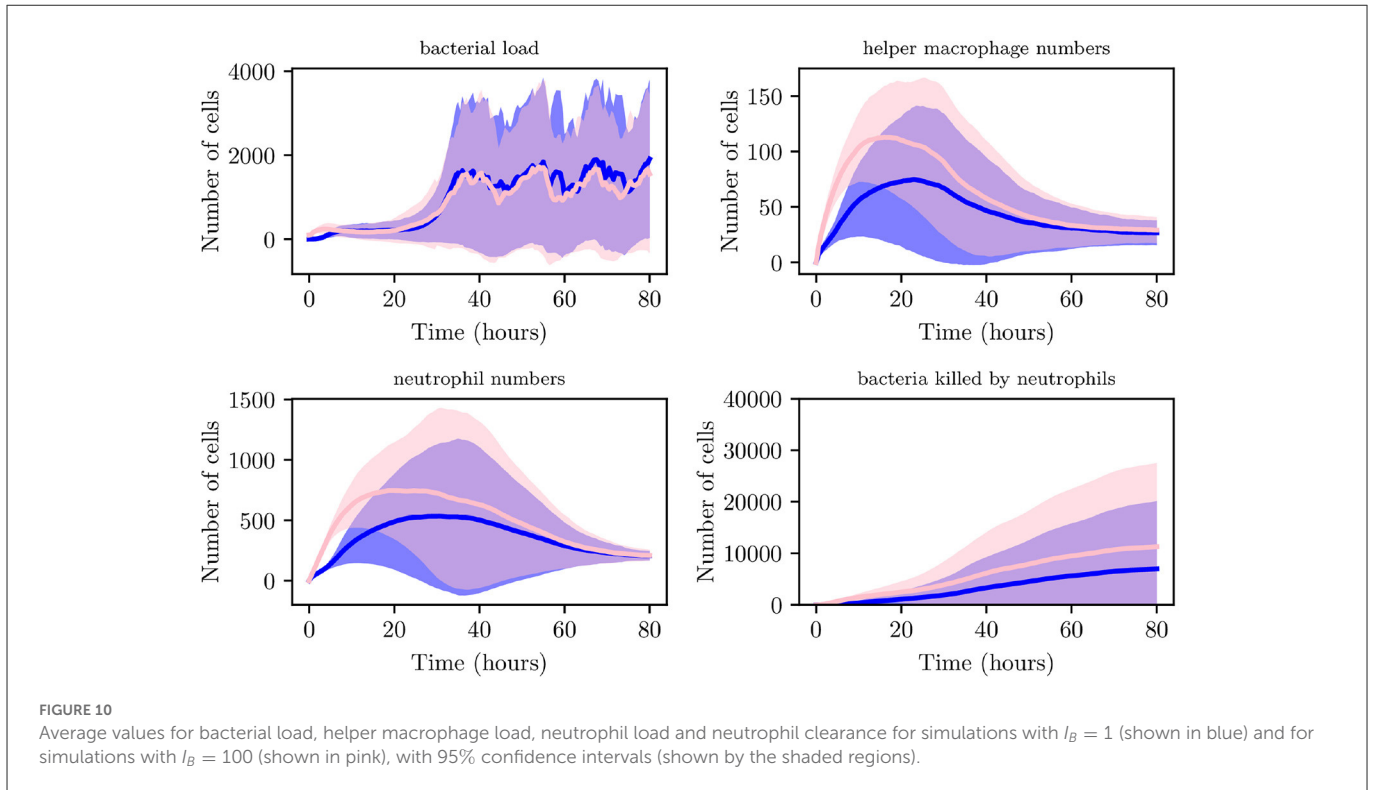
We chose to analyse PRCC, using bacterial load as the output of interest. Figure 3 shows the results from this sensitivity analysis, which suggest that the parameters related to the neutrophils in the model are significant. We investigate this further by plotting the PRCC values for these parameters through 80 h of a simulation, shown in Figure 4.

We chose to analyse PRCC, using bacterial load as the output of interest. Figure 3 shows the results from this sensitivity analysis, where we show the parameters with high sensitivity. Also notice in this figure that some parameters appear more than once in different forms depending on their state. For example, the probability of recruitment of macrophages is investigated separately; when bacteria are present and when bacteria are not present.

We have identified these parameters as the sensitive ones in the model but that their sensitivity only appears to be significant near the start of the simulation. Future work will include investigation of these parameters with regard the outcome of simulations. For the work presented here, however, we set these parameters as outlined in Table 1.

### 3. Results

Our model is able to capture a number of possible outcomes. Individual simulations may resolve in a variety of ways: immune cells may be able to completely eradicate bacterial infection



with the combination of immune agents and periodic bacterial clearance through shedding of the bladder wall; bacteria may colonize the modeled bladder section leading to established cystitis; or bacteria may persist in small numbers, perhaps evading the shedding mechanism by penetrating the epithelial barrier. These later simulations may be thought of as non-symptomatic cases.

### 3.1. Initial bacterial load

Here we investigate the effect that the initial bacteria load,  $I_B$ , can have on the infection dynamics. We first fix all model parameters, with the exception of  $I_B$ . Subsequently, we generate 200 simulations through each variation of  $I_B$ .

As we simulate 80 h, we will refer to an infection as being in the early stages when  $t \in [0, 15)$  hours. We refer to  $t \in [15, 30)$  hours as the middle stage of infection, which in the literature is identified as the stage when recruited immune cells are heavily involved in bacterial clearing. Finally, we refer to  $t \in [30, 80)$  hours as the late stage of the infection. Bacterial shedding occurs within the model when  $N_B > 6000$ , which usually occurs during the late stages of infection.

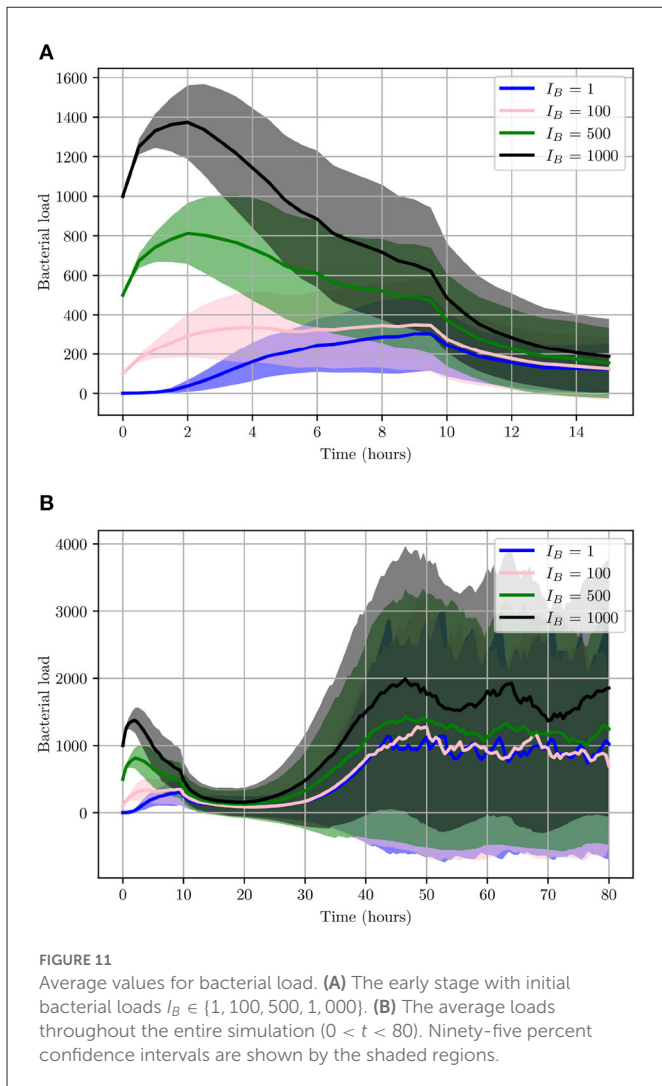
Throughout this analysis, we will use the following notation  $N_B^{t=y}$  to refer to the bacterial load at time  $y$  hours.

#### 3.1.1. Case where $I_B = 1$

When choosing  $I_B = 1$ , there is a large amount of variation in the infection outcomes. Progression profiles of UPEC can be observed, each resulting in a different final outcome. Figure 5 shows the number of bacteria from  $t = 0$  to  $t = 15$ , for 4 simulations that highlight the extremes of infection progression outcomes. In Figure 5A, bacterial

replication within the initial hours is slow; however, UPEC manages to evade the efforts of clearance by the immune cells and is able to replicate at higher rates toward the end of the early stage, reaching a final bacterial load of  $N_B^{t=15} = 120$ . In Figure 5B however, the infection progresses initially at higher speeds reaching a bacterial load of  $N_B^{t=6} = 420$ . Following this, the immune response manages to achieve good partial clearance outpacing replication at  $t = 7.5$  h. Toward the end of the early stage, the bacterial load is  $N_B^{t=15} = 218$ . In Figure 5C, bacteria is almost instantly cleared prior to being able to replicate, where immune cells successfully kill the bacteria. As immune cells are initially randomly placed within our grid, it is possible for an individual simulation to have immune cells within the immediate vicinity of  $I_B$ , which, in addition to having a low initial bacterial load ( $I_B = 1$ ), leads to fast and total clearance of bacteria. In Figure 5D, the infection progresses quickly reaching  $N_B^{t=6} = 100$ . The bacterial increase however, although steady throughout the early stage of the infection, halts and the immune system manages to successfully clear all bacteria with  $N_B^{t=15} = 0$ . To gain a better insight into the immune clearance process we also present Figure 6 which describes the immune cell dynamics for the same 4 simulations as shown in Figure 5. As expected, we see here higher recruitment of both macrophages and neutrophils in simulations with higher bacterial loads. Figure 7 shows spatial plots of the simulation shown in Figures 5A, 6A at timepoints  $t = 0, 5, 10, 15$  h. We see here as the infection progresses, there is a large influx of immune cells through the blood vessels that are subsequently directed by the chemokine signal to the site of infection at the center of the domain. We see that, at  $t = 15$  h, the immune response has not been successful in containing the infection spread.

While analysis of these individual simulations for the first 15 h is useful, we also look at the aggregate behavior resulting from our simulations with  $I_B = 1$ . With a total of 200 simulations, run for a



total of 80 h, clearance was achieved in 76.7% of instances. Within these, the vast majority (65.8%) was due to early clearance, i.e. before 15 h, with mid and late clearance accounting collectively for the remaining 10.9% of clearance cases.

Thus, our model indicates that low initial bacteria loads usually lead to total bacterial clearance within the initial stages of infection, with the majority of instances achieving total clearance within 6 h. However we see in this work that when only a single bacterium can lead to significant infection spread, as in 23.3% of simulations, clearance was never achieved. Here the bacteria were able to further colonize bladder epithelial tissue, bypassing immune cell action and bacterial shedding.

### 3.1.2. Case where $I_B = 100$

We now investigate an initial bacterial load of  $I_B = 100$ , by examining the different possible infection progression profiles. As can be seen in Figure 8, total clearance here does not occur at any time before  $t = 6$  h, with the simulations in Figures 8A–D showing early periods of considerable bacterial growth before halting due to immune cell intervention. Simulation (Figure 8A) presents an individual infection simulation where the bacterial load reaches a maximum of  $N_B^{t=4.5} = 280$ , after which, the immune

response achieves successful total clearance. Simulation (Figure 8B) on the other hand presents a relatively steady bacterial load until  $t = 9.50$  h, reaching  $N_B^{t=9.5} = 327$ , after which the immune cells are only able to achieve partial clearance of the bacteria, with  $N_B^{t=15} = 107$ . In simulations (Figures 8C, D) we see the bacterial load decline occurring earlier in the simulations, with successful immune responses to the infection.

In looking at the aggregate of the 200 simulations where  $I_B = 100$ , we can see that bacteria persist in 30.8% of simulations (as opposed to 23.3% when  $I_B = 1$ ). Early clearance accounted for 45.8% of simulations while mid stage clearance accounted for 21.9% of instances. Late clearance was achieved in 3 simulations (1.5%). See Figure 9 for pie charts summarizing this.

Although an increased  $I_B$  causes recruitment of immune cells at earlier times to the site of infection in comparison with lower  $I_B$ , bacteria are able to replicate and reach higher loads, typically having reached a maximum within the initial 5 h. As can be seen in Figures 8B, C instances attained bacterial loads  $N_B > 300$ . Reaching these stable numbers allows the bacterial population to persist despite attempted clearing by neutrophils and helper macrophages. Through analysis of spatial plots we can see that earlier immune cell intervention in simulations with  $I_B = 100$ , as opposed to simulations with  $I_B = 1$  is due to the closer proximity of the boundary of the bacterial population to recruited immune cells. As can be seen from Figure 10, the aggregate profiles and patterns of bacterial growth in the  $I_B = 1$  simulations are similar to those with  $I_B = 100$ . The differences in the profiles of helper macrophages, neutrophil and bacterial clearance are due to the higher intensity of immune action due to the increased initial bacterial count.

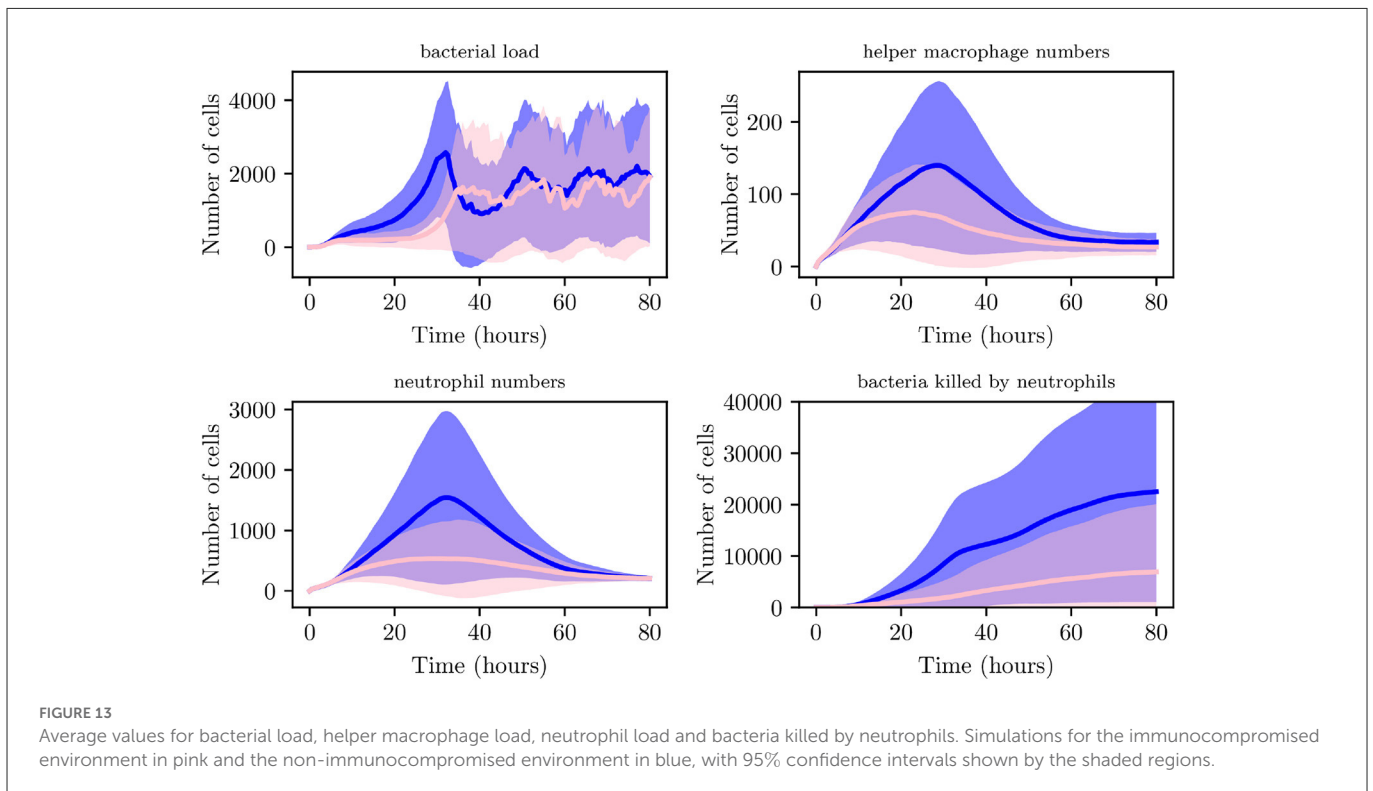
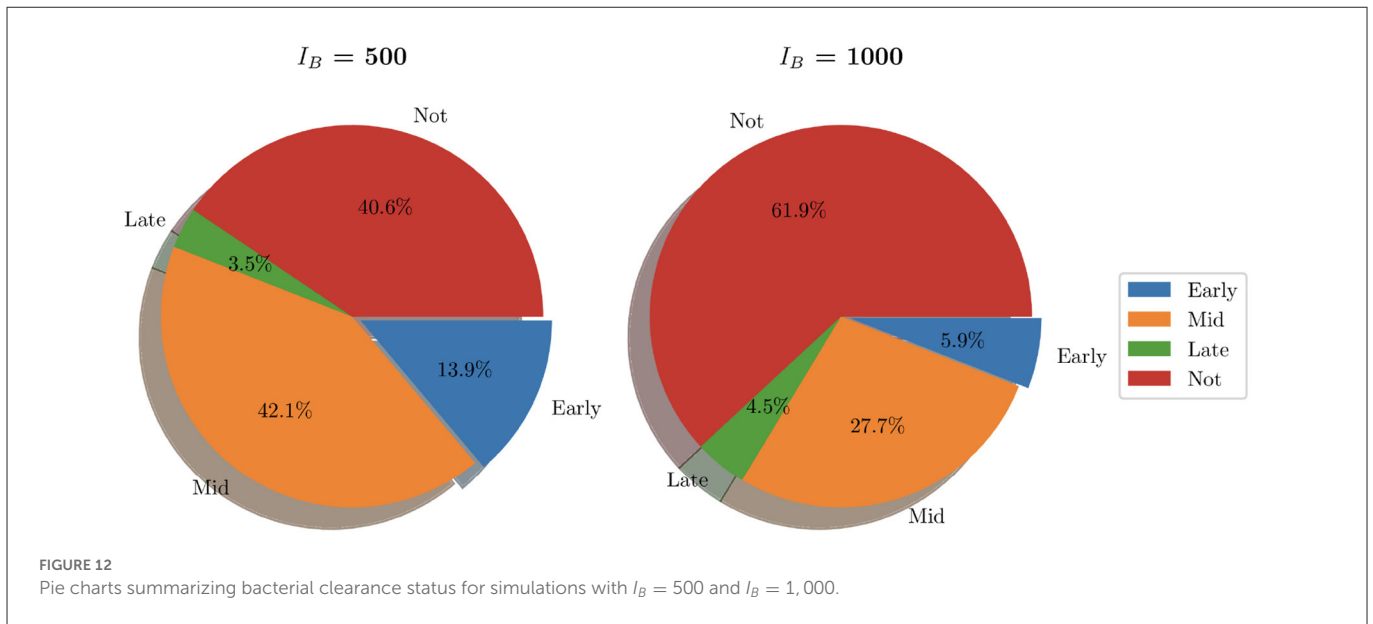
## 3.2. Bacterial replication rate

Studies have shown that UPEC presents different replication profiles [29]. At early stages of the infection, *E. coli* replicates with an average doubling time of 20–45 min [30]. As the infection progresses however, research has shown that UPEC adopts a relatively lower replication rate, estimated with a doubling time of 40–60 min [29]. We now include this to our model and investigate the effects on bacterial clearance and immune presence. In order to do this, we include two parameters describing the replication rate.  $B_{earlysp}$  describes the replication rate during the earlier stages of the model (when  $t < 10$  h). On the other hand,  $B_{latesp}$  is then used after 10 h. Hence, we set

$$B_{sp} = \begin{cases} B_{earlysp}, & t < 10 \text{ h} \\ B_{latesp}, & t \geq 10 \text{ h} \end{cases} \quad (2)$$

Here we analyse the output resulting from 1,000 individual simulations of our model, allowing us to create and analyse a virtual population as in Verma et al. [31, 33] and Niederer et al. [32]. We allow UPEC to adopt two replication profiles as detailed above, whilst also varying the initial bacterial load,  $I_B \in \{1, 100, 500, 1,000\}$ , with 200 simulations performed for each case.

This will allow us to investigate the effects  $I_B$  can have on the progression of the infection. As can be seen from Figure 11, during the initial 2 h, lower  $I_B$  present a relatively slower growth rate while high  $I_B$  will initially present with a much higher growth rate. These differences however are less noticeable when looking at  $2 < t < 9$



h, as the bacterial count of high  $I_B$  begins to decline due to clearance by neutrophils and helper macrophages that have arrived at the site of infection. While in contrast, a low  $I_B$  allows bacteria to maintain a steady growth within the interval  $2 < t < 6$  h. Once  $9 < t < 14$  h, all simulations converge closely. The final bacterial load of the early stage,  $N_B^{t=15}$ , matches in order of initial bacterial count except for  $I_B = 1$  which has managed to overtake  $I_B = 100$ . Although the trend appearing from the early stages of infection seems to indicate possible bacterial clearance, once we look at the later stages of the simulation we realize the importance bacterial shedding plays, where we see the periodic nature of the bacterial profiles.

As illustrated in Figure 11, although the growth of UPEC during the early stages can be slow and heavily hindered by neutrophils, at later stages bacterial growth increases significantly. Consequently, bacterial shedding can be a crucial mechanism to control infection within the superficial urothelium and prevent full colonization of the bladder wall.

Similarly to Figure 9, we show the pie charts of these simulations where we have taken the following initial bacterial loads:  $I_B = 500$  and  $I_B = 1,000$ . In contrast to the earlier analysis shown in Figure 9 where we looked at relatively low initial bacterial loads, Figure 12 presents the clearance in the case of high initial bacterial loads. As we

can see, with higher initial bacterial loads, clearance is not achieved in 40.6% of instances (when  $I_B = 500$ ) and 61.9% when ( $I_B = 1,000$ ). In addition to this, total bacterial clearance is now achieved generally during the middle stages of infection rather than early stages as was shown with  $I_B = 1$  and  $I_B = 100$ . This signals toward the ability of higher  $I_B$  to establish more severe infections.

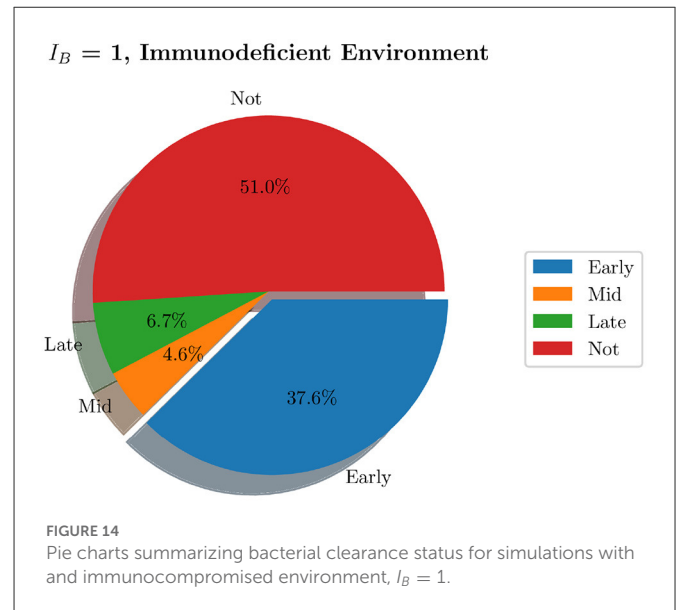
### 3.2.1. Simulating an immunocompromised environment

We now generate 200 simulations with  $I_B = 1$ , however, we change the number of resident immune cells to reflect an immunocompromised environment. We set  $I_{rM} = 300$ , and  $I_{MC} = 30$  (as opposed to all previous simulations where  $I_{rM} = 500$  and  $I_{MC} = 50$ ). As can be seen in Figure 13, in an immunocompromised environment with low initial loads of resident immune cells, our model indicates that bacteria are able to establish larger infections. Furthermore, an aggregate analysis of the 200 simulations (see Figure 14) indicates that our model does not achieve clearance in 51% of instances, in contrast with the non-immunocompromised environment (also  $I_B = 1$ ), only 23.3% of simulations do not achieve clearance. Our model produced late clearance in 6.7% of instances, which allows more time for the bacteria to penetrate the umbrella cells and potentially form more QIRs. This in turn would lead to greater bacterial loads and growth post shedding, thus increasing the likelihood of recurrence.

## 4. Discussion

We have developed a spatial mathematical model to investigate the pathogen-host response in a typical bladder infection. In a 2-dimensional  $16 \text{ mm}^2$  section of the bladder, we use our hybrid individual-based model to simulate Uropathogenic *E. coli* as individual elements, as well as immune cells. In particular, we model resident and helper macrophages, neutrophils and mast cells. In addition to these discrete elements, we also model a generic chemokine in order to act as a chemoattractant. This molecule directs the immune cells to the site of infection, diffusing into the spatial domain via a PDE.

In this paper, we have outlined the basic framework for the model. We include a gradual decrease in the speed of bacteria, as evidenced in the literature, and also model bacterial shedding as a mechanism for reducing bacterial burden in the bladder. We investigate several initial bacterial loads and show that even with one bacterium at the start of the simulation it is possible for infection to spread with colonization of the bladder epithelial tissue. This finding is consistent with recent work in the porcine model of UTI, in which an initial inoculum of  $\leq 10$  bacteria was sufficient to seed a productive infection *in vivo* [34]. Not surprisingly, as the initial bacterial burden increases, the time taken for the infection to clear increases. The number of simulations not cleared within the 80 h of the simulated infection also increases as the initial bacterial load increases. Interestingly, the average number of bacteria present in the cases where the infection is not cleared, is not too dissimilar for  $I_B = 1,000$ , and we only see a big difference when  $I_B = 1,000$ . At this initial bacterial load, the average number of



bacteria during the late stage of infection is significantly larger than for smaller initial bacterial loads. In Section 3.2.1 we investigate an immunocompromised environment with smaller numbers of immune cells acting against the infection. Here we see, as expected, that for immunocompromised simulations, more cases resulted in infection persisting at the end of the simulation than in the non-immunocompromised simulations. These results are consistent with clinical scenarios. For example, UTI occurs in up to 72% of immunocompromised donor kidney recipients, usually within the first few months post-transplant [35].

We have shown that the modeling framework that we have developed is able to simulate various scenarios in order to analyse the resulting impact of particular parameters, helping to aid understanding of these infections. Although we have only modeled the first 80 h of an infection course, as we keep a count in the model of the bacteria that manage to penetrate into the epithelial bladder wall and evade bacterial shedding, in future work we will run longer simulations to investigate the growth of these bacteria and their role in recurrent infections. We also plan to integrate a Pharmacokinetic/Pharmacodynamic (PK/PD) model of commonly used antibiotics in order to simulate treatment effects. To our knowledge, integrating a PK/PD model into a spatial model has not been done before to study bladder infections, and the spatio-temporal framework we present here is the crucial starting point. We are also in the process of completing parallel wet-lab experiments to compare to our model simulations. This analysis will be shown in future publications. We acknowledge the limitations of the model parameterisation of our framework and also plan to conduct a number of simulations to more rigorously test the effects of varying the neutrophil parameters, as these were highlighted in the sensitivity analysis as the most sensitive quantities. We also plan to include the BECs as agents within the model in future iterations, as well as expand the model to three dimensions in order to more realistically simulate the environment. With the planned future developments, this model will be capable of providing new clinical insight into bladder infections.

## Data availability statement

The original contributions presented in the study are publicly available. This data can be found at: <https://github.com/Ruth-Bowness-Group/UTImodel>, <https://doi.org/10.5281/zenodo.7293870>.

## Author contributions

RB provided funding for the work. RB and ALD contributed to model conception and wrote the first draft of the manuscript. ALD, RB, TL, JR, and BP contributed to the design of the study. ALD created the computational code and performed analyses. TL, JR, and BP wrote sections of the manuscript. All authors contributed to manuscript revision, read, and approved the submitted version.

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