

Investigation of spatial dynamics of fungal-host-interactions of *Candida albicans* and innate immune cells

<u>Sandra Timme^{1,2}</u>, Teresa Lehnert^{1,2}, Johannes Pollmächer^{1,2}, Kerstin Hünniger^{2,3}, Oliver Kurzai^{2,3}, Marc Thilo Figge^{1,2}

¹ Applied Systems Biology, Leibniz Institute for Natural Product Research and Infection Biology – Hans-Knöll-Institute, Jena ² Friedrich Schiller University Jena

³ Septomics Research Center, Leibniz Institute for Natural Product Research and Infection Biology – Hans-Knöll-Institute, Jena

Introduction

An increasing clinical problem that is associated with high mortality rates are systemic infections caused by *Candida albicans*. This opportunistic human fungal pathogen is a human comensal that resides on mucosal surfaces and can enter the bloodstream via the intestinal gut or medical devices, such as catheters. In immunocompromised patients the immune system is not able to clear the infection which allows *C. albicans* to disseminate through the whole body and cause sepsis.

Whole blood infection assay with human blood¹:



Bottom-up approach

In order to investigate the infection process of *C. albicans* different aspects of the system are of particular interest. To capture non - spatial rates, like phagocytosis rates or killing rates as well as spatial aspects, such as diffusion constants, within time we developed a bottom-up approach². This approach incooperates different modeling techniques with increasing complexity building on one another.





² Lehnert and Timme et al.(2015),Frontiers in Microbiology 6(608)

Agent-based model (ABM)

The agent-based modeling approach is a spatial, rule-based, stochastic modeling technique, which simulates the action and interaction of agents (cells) within a defined environment. This enables the possibility to investigate spatial aspects of biological systems.





Parameter estimation approach

Adaptive regular grid search:

0	0	0	0	0	0	0	0

parameter

For the calibration of the model onto experimental data, system parameters have

to be estimated. The quality of a simulation (x^{sim}) compared to the experimental data (x^{dat}) is determined by the Least-Squares error (LSE):

$$E[\vec{p}] = \sum_{c} \epsilon_c \frac{1}{2} \sum_{k} (x_{k,c}^{dat} - x_{k,c}^{sim}[\vec{p}])^2.$$

$$\tag{1}$$

The estimation of system parameters requires a scanning of the parameter space. Since agent-based modeling is associated with high computational load, there is need for a search technique which keeps computation time whithin limits. Therefore the *adaptive regular grid search* was chosen, which systematically scannes the parameter space and performes a refinded search in regions with low LSE.

Parameter estimation results



The application of the *adaptive* regular grid search for the parameter estimation of the diffusion coefficients for PMN (D_G) and for monocytes (D_M) results in a landscape that has a valley within the search space.

- ullet not sensitve to variations in D_M
- ullet very sensitive to variations in D_G
- $(D_G^{min}, D_M^{min}) = (425, 275) \, \mu m^2 / min:$





Predictions on *C. albicans* immune escape mechanisms

Immune escape mechanisms have been observed for different pathogens in biological systems. However, due to the high cell density it is very hard to proof immune escape of *C. albicans* in human blood. We hypothesise that mechanisms of immune escape of *C. albicans* in human blood are correlated with the occurance of restistant *C. albicans* cells during the infection process.





fits the experimental data very wellstandard deviation due to the stochasticity of the model







I.

The preliminary results show, that experimental and simulated data are in good agreement. Therefore we suggest to consider the process of immune escape of C. albicans also in human blood.

Outlook

investigate other immune cells and pathogens
include cytokine dynamics with PDE (hybrid ABM)
distinguish hyper- and hypo-inflammation in immune dysregulation of sepsis patients

Acknowledgement

This work was financially supported by the excellence graduate school Jena School for Microbial Communication (JSMC) and the CRC/TR124 FungiNet, Project B4, that are both funded by the Deutsche Forschungsgemeinschaft (DFG). Further financial support is provided by the Center for Sepsis Control and Care (CSCC), Project Quantim, that is founded by Federal Ministry of Education and Research (BMBF).

Friedrich-Schiller-Universität Jena

Contact: sandra.timme@hki-jena.de





