REPORT

The space between us: Modeling spatial heterogeneity in synthetic microbial consortia dynamics

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ABSTRACT A central endeavor in bioengineering concerns the construction of multistrain microbial consortia with desired properties. Typically, a gene network is partitioned between strains, and strains communicate via quorum sensing, allowing for complex behaviors. Yet a fundamental question of how emergent spatiotemporal patterning in multistrain microbial consortia affects consortial dynamics is not understood well. Here, we propose a computationally tractable and straightforward modeling framework that explicitly allows linking spatiotemporal patterning to consortial dynamics. We validate our model against previously published results and make predictions of how spatial heterogeneity impacts interstrain communication. By enabling the investigation of spatial patterns effects on microbial dynamics, our modeling framework informs experimentalists, helps advance the understanding of complex microbial systems, and supports the development of applications involving them.

INTRODUCTION

Since the discovery of quorum sensing in the early 1970s (1), understanding of microbial communication has grown tremendously. Multiple communication mechanisms such as acyl-homoserine lactones (AHLs) (2, 3) and autoinducing polypeptides (AIPs) (4, 5) have been uncovered, and their role in the formation of complex social behaviors such as biofilms and swarming motility have been deciphered (6–8). Recognizing that consortia of interacting microbial populations can perform more complicated behaviors than any one species individually, researchers have begun to leverage these systems for various biotechnology applications (9). For example, microbial consortia have been designed using natural or genetically engineered bacterial strains for use in biosensors (10), bioremediation (11, 12), and more efficient bio-production (13–15). With this wide level of utility, the impact of microbial consortia on industry and human health are likely to grow as more efficient tools are developed for their design and application.

Substantial tools exist for controlling a particular consortium’s behavior genetically using synthetic regulatory circuits (16–19), and tools for engineering a consortium’s spatial organization have also begun to be developed (20). For example, Romano et al. recently developed a novel optogenic method for regulating gene expression that allowed them to re-create highly complex images using bacterial lawns (21). Similarly, Alnahhas et al. demonstrated that the number of subpopulation bands in an extended microfluidic device

SIGNIFICANCE Multistrain microbial consortia are able to accomplish tasks difficult to engineer in any one strain. Thus, substantial research in bioengineering has focused on the construction of multistrain microbial consortia with desired properties. Yet how emergent spatiotemporal patterning affects consortial dynamics is still poorly understood. A better understanding of this effect can aid the development and deployment of microbial consortia for various biotechnology applications. Toward this goal, we propose a computationally tractable and straightforward modeling framework that explicitly allows linking spatiotemporal patterning to consortial dynamics. Furthermore, our framework is flexible and generalizes to a wide range of applications.
can be regulated through the seeding of the trap. Other methods for regulating the spatial organization of consortia including adhesion, cell-communication, and motility are also being developed (20, 22–26). With previous studies demonstrating that the spatial organization of a consortium can significantly alter its dynamics (18, 27), these experimental methods are enabling an additional layer of control for engineering microbial consortia.

What is lacking from the research are sufficiently flexible modeling frameworks for investigating the impact of complex spatiotemporal structure on microbial consortium dynamics. These tools have the potential to streamline the rational engineering of spatial patterns to control consortium behavior. Previous approaches to modeling the spatiotemporal dynamics of microbial consortia have traditionally relied on well-mixed compartment models (16, 17) or agent-based frameworks (28–30). However, the well-mixed compartment models do not allow for the explicit incorporation of strain patterns, and agent-based modeling frameworks are computationally expensive for larger systems (31). To address this gap in the literature, we borrow from previous research focused on modeling diffusion in heterogeneous media (32–34) and develop a piecewise-defined reaction-diffusion equation framework for modeling spatiotemporal consortium dynamics. Here, we show that this modeling framework allows for the explicit incorporation of consortium spatial organization while remaining computationally tractable for larger systems. We then use this framework to investigate the impact of spatial heterogeneity on consortium dynamics and, using our results, demonstrate the rational design of consortium dynamics through spatial organization. Importantly, our framework provides an explicit link between interstrain communication and spatiotemporal patterning.

Modeling framework

For diffusion in heterogeneous media, variations in the composition of the medium result in corresponding discontinuities in its physical properties that need to be accounted for when modeling molecular transport. Researchers traditionally model this phenomenon using a reaction-diffusion equation (32–34):

\[
\frac{\partial c}{\partial t} - \nabla \cdot (D(x)c(x, t)) = f(c, x, t),
\]

\[
t \geq 0, \quad x = (x, y, z) \in \Omega,
\]

(Reaction-Diffusion Equation)

where \(c = c(x, t)\) is the local concentration of the diffusing molecule, \(D\) is the diffusion coefficient, \(t\) is time, \(\Omega\) is the domain, and \(f(c, x, t)\) describes the production and degradation of \(c\). To account for variations in transport properties in heterogeneous media, \(D\) is piecewise-defined according to the medium composition to have the appropriate value in the appropriate (user-controlled) location. Recognizing a natural correspondence between media heterogeneity and cell heterogeneity, we modify this modeling approach so that it can be used to study the spatiotemporal dynamics of microbial consortia. In our modified approach, the concentration of each signaling molecule in the consortium is modeled using a reaction-diffusion equation with components representing the local production and degradation described by a source term \(f(c, x, t)\). We incorporate the spatial organization of the consortium strains into the resulting partial differential equation (PDE) model by structuring the source term for each signaling molecule on \(\Omega\) as a piecewise function. This ensures that each signal is only generated in areas that contain a strain capable of producing it. Since each strain is uniquely defined by the signaling molecules that it produces, piecewise defining our system creates areas of production that mimic the spatial organization of the strains (see Fig. 1A). This allows the resulting system to incorporate the influence of spatial organization on the dynamics of the microbial consortium and allows for the investigation of any desired organization.

Evaluation of framework and results

As a first application of this framework, we used it to model a recent experimental study of the spatiotemporal dynamics of a well-known microbial consortium (16). There, Kim et al. compared the dynamics of four microbial consortia in a spatially extended microfluidic trap consisting of different types of feedback networks. Each consortium consisted of an activator and repressor microbial strain in an alternating columnar spatial arrangement (diagrammed in Fig. 1A). The two strains regulated each other’s gene expression in a dual-feedback mechanism to produce an oscillating fluorescent pattern (17). Through experimentation and modeling, Kim et al. found that the addition of a positive feedback loop resulted in globally coordinated oscillations. This was in spite of the fact that the length of the trap was significantly longer than the diffusion correlation length of the signaling molecules, thereby preventing direct communication between strains located far apart from each other (16). However, their model omitted the organization of the stripe spatial patterns. Stripe thickness could affect the ability of the strains to couple their behavior through diffusion. Thus, the influence of strain spatial organization on consortium dynamics warrants investigation. For simplicity, we only model two of the
To model the spatiotemporal dynamics of the consortia studied in Kim et al., $P_2N_1$ and $P_1N_1$ (diagrammed in Fig. 1 B and C), since they behave very similarly to the other two consortia. The $P_2N_1$ and $P_1N_1$ consortia also exhibited more robust columnar stripe patterns making them better suited to study in 1D.

Model derivation

To model the spatiotemporal dynamics of the $P_2N_1$ and $P_1N_1$ consortia using our framework, we constructed reaction-diffusion equations describing the dynamics of the consortium’s extracellular signaling molecules. Like previous research into microbial consortia (35), we developed a simplified model of the dynamics since a thorough description of the underlying gene circuit interactions has been previously shown to require 16 delay-differential equations (DDEs) (17). To do so, we suppressed the dynamics of the fluorescent reporters and considered only the dynamics of the extracellular signaling molecules. This simplification is valid because the fluorescent reporters are produced in proportion to the local concentration of the lactone-based signaling molecules (17). We also ignored the role of intermediaries in producing these signaling molecules by modeling their production using Langmuir-functions, which take into account the overall activator-repressor network interactions. These functions mirror the form of previously developed equations for modeling production in consortia (16, 17) and are provided below, where $a$ represents the local concentration of the activator N-butanoyl-L-homoserine lactone (C4-HSL, C4) and $r$ represents the local concentration of the repressor N-(3-hydroxytetradecanoyl)-DL-homoserine lactone (3-OHC14-HSL, C14). We chose a similar strategy to model their enzymatic degradation by AHL-lactonase, AiiA, since it is produced in response to the signaling lactones and a 7.5-min transcriptional delay, $\tau$ (16, 17).

With these interactions characterized, the source terms $f_a(x, t)$ and $f_r(x, t)$ are given by

$$f_a(x, t) = \frac{\eta_{a_0} + \eta_{a_1} \left( \frac{a_x}{K_a} \right)^{n_a}}{1 + \left( \frac{a_x}{K_a} \right)^{n_a} + \left( \frac{r_x}{K_r} \right)^{n_r}} - \left( \frac{r_x}{K_r} \right)^{n_r} - \gamma_a$$

$$f_r(x, t) = \frac{\eta_{r_0} + \eta_{r_1} \left( \frac{a_x}{K_a} \right)^{n_a}}{1 + \left( \frac{a_x}{K_a} \right)^{n_a} + \left( \frac{r_x}{K_r} \right)^{n_r}} - \left( \frac{r_x}{K_r} \right)^{n_r} - \gamma_r$$

where $a_x \equiv a(x, t - \tau)$ and $r_x \equiv r(x, t - \tau)$. A complete description of all the parameters and their corresponding values can be found below in Table 1, and a description of how the parameters values were determined can be found in the Methods section of the Supporting

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TABLE 1 Parameter Values and Descriptions

| Parameter | Description | Value   |
|-----------|-------------|---------|
| $\eta_{ai}$ | Basal production rate of activator signaling molecule from promoter. | 985.56 nM min$^{-1}$ |
| $\eta_{ai}$ | Maximal production rate of activator signaling molecule from promoter in response to activator signal. | 17,991.82/0 nM min$^{-1}$ |
| $\eta_{ri}$ | Basal production rate of repressor signaling molecule from promoter. | 74.46 nM min$^{-1}$ |
| $\eta_{ri}$ | Maximal production rate of repressor signaling molecule from promoter in response to activator signal. | 46,495.80 nM min$^{-1}$ |
| $K_a$ | $EC_{50}$ of activator signaling molecule for production. | 5937 nM |
| $K_r$ | $IC_{50}$ of repressor signaling molecule for repression. | 10 nM |
| $n_a$ | Langmuir coefficient of production for activator signaling molecule. | 4 |
| $n_r$ | Langmuir coefficient of repression for repressor signaling molecule. | 2 |
| $D$ | Diffusion coefficient signaling molecules. | 4080 $\mu$m$^2$ min$^{-1}$ |
| $d_l$ | Maximal rate of internal enzymatic degradation of signaling molecules. | 2257 min$^{-1}$ |
| $K_e$ | $EC_{50}$ of repressor signaling molecule for enzymatic degradation. | 1000 nM |
| $n_e$ | Langmuir coefficient of internal enzymatic degradation for repressor signaling molecule. | 4 |
| $\gamma$ | Inherent rate of degradation of signaling molecules caused by dilution from both media flow and cell growth. | 0.128 min$^{-1}$ |
| $\tau$ | Time delay for transcription and translation. | 7.5 min |

To differentiate between the $P_2N_1$ and $P_1N_1$ consortia, $\eta_{ai}$ is set to zero for the $P_1N_1$ consortium to account for the absence of the auto-activation, and the $(\frac{\omega}{K_i})^{n_a}$ in the denominator of the production term is removed. We thus arrived at the following two-equation system, partial delay-differential equation model of the $P_2N_1$ and $P_1N_1$ consortium spatiotemporal dynamics:

$$\begin{align*}
\frac{\partial a}{\partial t} - D\nabla^2 a &= f_a(x, t) \\
\frac{\partial r}{\partial t} - D\nabla^2 r &= f_r(x, t)
\end{align*}$$

The strain spatial organizations are included in this model in a user-defined fashion through the piecewise definition of $f_a(x, t)$ and $f_r(x, t)$. Specifically, we set the production rate terms in $f_a(x, t)$ and $f_r(x, t)$ to be zero in areas that did not include an activator or repressor strain, respectively.

Model validation

To validate our simplified model of the $P_2N_1$ and $P_1N_1$ consortia, we investigated its ability to accurately capture the spatiotemporal dynamics recently observed in Kim et al. The main result of this paper was that strains containing a second positive feedback loop synchronized their dynamics across a spatially extended microfluidic trap. Thus, we validated our model by testing whether it could reproduce this key behavior. For this test, we simulated the $P_2N_1$ and $P_1N_1$ consortia in one dimension. A 1D model was chosen since diffusion and variations in strain composition in the vertical direction are assumed to be negligible in the spatially extended trap (16). We therefore modeled the $L = 2$ mm long trap (e.g., device size in Kim et al. (16)) using the 1D spatial domain, $\Omega = [0, L]$. The consortia were implemented by dividing the spatial domain into 100 stripes of equal width alternating between activator and repressor strains. The consortium was then divided into two equally spaced halves oscillating with an initial phase difference of $|\Delta \varphi_0| = 25\%$ of the oscillation period, $\varphi$. We chose this value because it resulted in synchrony for $P_2N_1$ and asynchrony for $P_1N_1$ in Kim et al. Both the $P_2N_1$ and $P_1N_1$ consortia were then analyzed for whether synchronization between the two halves occurred. To determine the presence of synchronization, we calculated the phase difference of the activator concentration oscillations for the middle spatial points of the left and right subpopulations, $\Delta \varphi$ (detailed explanation provided in the Methods section of the Supporting material). A percent phase difference, $|\Delta \varphi|$ was then calculated as follows:

$$|\Delta \varphi| = \frac{\Delta \varphi}{\varphi} \times 100$$

We assumed a phase difference of $|\Delta \varphi| < 2.5\%$ between the two halves corresponded to synchronization across the trap. Since $\varphi$ depends on the basal production rate, $\eta_{ai}$, we chose to measure the phase difference after the 10th oscillation instead of after a specified period of time had passed.

The presence of the autoinducing positive feedback loop in the $P_2N_1$ consortium was cited in Kim et al. as the reason $P_2N_1$ oscillations became coordinated, whereas $P_1N_1$ oscillations did not. Thus, we originally only varied the value of $\eta_{ai}$ between our $P_2N_1$ and $P_1N_1$ simulations, setting it to zero in the $P_1N_1$ simulation to account for an absence of the feedback loop.
We failed to see any significant difference in coordination between the \( P_2N_1 \) and \( P_1N_1 \) consortia (Fig. 2 C). With this finding, it was clear that the difference in the activator basal production rate between the \( P_2N_1 \) and \( P_1N_1 \) consortia needed to be incorporated into our model. In Chen et al., it was reported that the activator basal production rate in the \( P_1N_1 \) consortium is approximately nine times greater than in the \( P_2N_1 \) consortium, and this difference was even included in the model used in Kim et al. Thus, to investigate the role changing this basal production rate had on \( P_1N_1 \) behavior, we re-ran the \( P_1N_1 \) synchronization tests for different values of \( \eta_{a0} \). Specifically, we scaled \( \eta_{a0} \) by a basal multiplier, \( b_{am} \), that varied from 0.1 to 10. The results of these tests are provided in Fig. 2 C and show that the coordination of oscillations is strongly influenced by the value of the activator basal production rate, \( \eta_{a0} \). Kymographs illustrating the level of coordination in the \( P_2N_1 \) and \( P_1N_1 \) consortia are provided in Fig. 2 A and B, respectively.

Investigating the role of spatial heterogeneity

Having validated our model against the experimentally observed difference between the \( P_2N_1 \) and \( P_1N_1 \) consortia, provided the difference in activator basal production rates is incorporated, we proceeded to test our model's ability to capture the influence of spatial heterogeneity on consortium dynamics. This is the primary focus of this work and will provide new insight into heterogeneous systems. First, we investigated the impact changing stripe width had on the coordination of oscillations in the \( P_2N_1 \) consortium. This was done by repeating the aforementioned synchronization simulations including kymograph of activator concentration during the 8th to 11th oscillations of the \( P_2N_1 \) (A) and \( P_1N_1 \) (B) consortia (basal multiplier of 0.7). (C) Graph showing the relationship between phase difference and the value of the basal multiplier for the \( P_1N_1 \) consortium at two different initial phase differences, \( |\Delta \phi_0| = 25\% \) and \( 20\% \).

![Biophysical Reports 2, 100085, December 14, 2022](https://example.com/image.png)
test for the $P_2N_1$ consortium with stripe widths varying from 10 to 200 μm. The results of these simulations are provided in Fig. 3 B and show that coordination breaks down as stripe width increases. In fact, we observe a significant breakdown in synchrony for stripe widths larger than 90 μm. This shows quantitative agreement with previous experimental studies of the same consortia in Gupta et al. There, they found that separation distances larger than 100 μm between activator and repressor populations led to a loss of oscillatory behavior. This demonstrates that our modeling framework results are consistent with experimental results from different studies (16, 18).

Interestingly, a stripe width of 70 μm is optimal for synchrony. We intuit this result as follows. For stripe widths larger than 70 μm, the diffusion correlation length of the quorum sensing molecules is not great enough to allow for interstrain communication. That is, the coupling is too weak. On the other hand, for stripe widths less than 70 μm, chemical signals can diffuse across several stripes, flooding the trap with chemical signal. When active, the activator promoters are maximally productive, leading to an effective basal production rate that corresponds to the right-hand side of Fig. 2 C. In effect, the spatial heterogeneity provides an engineering methodology to control the effective basal production of the strains.

With this result in mind, we hypothesized that the coordination of the consortium could be controlled by only varying its spatial organization. To test this hypothesis, we simulated two $P_2N_1$ consortia side by side in a 4-mm-long trap. Both consortia contained alternating activator and repressor stripes whose widths increased linearly. However, the first consortium contained small-width stripes (0.56–47.06 μm), and the second consortium contained large-width stripes (19.05–266.67 μm). The results of this test are provided in a kymograph showing the concentration of the activator across the consortium and time. As anticipated by the results provided in Fig. 3 B, the left half of the consortium oscillated in a coordinated manner, and the right half of the consortium, containing larger stripes, had its oscillations become decoupled.

**DISCUSSION**

We developed a new modeling framework to investigate the impact of spatial heterogeneity on consortial dynamics. We verified our framework results against the modeling and experimental results found in Kim et al. (16). We also observed quantitative agreement between our stripe width investigation study and experimental findings from Gupta et al. (18). Although our model is much simpler than what is used by Kim et al. and coarse-grains over many subcellular processes, the qualitative behaviors of the two models coincide. Our model gains tractability at the expense of some fidelity to reality. But it is exactly in these scenarios that mathematical modeling can provide unexpected qualitative predictions regarding the biological system.

The results demonstrate that our new modeling framework is able to effectively incorporate the effect of spatial organization on microbial consortium dynamics. By allowing for the explicit incorporation of stripe patterns, our work complements and extends previous research into modeling spatiotemporal dynamics of microbial consortia (16, 18, 22, 28–30, 36, 37). For example, Kim et al. modeled spatiotemporal dynamics using well-mixed compartments containing both activator and repressor strains. However, this approach prevents the incorporation of well-defined
spatial patterns into the consortia, which, in turn, limits any investigation into their effect on consortium dynamics. Our piecewise-defined diffusion equation framework allowed us to investigate this impact in the present study. Our modeling framework is thus more faithful to the robust single-strain stripes observed in Alnahhas et al. Compartment models force all cells in a given compartment to experience the same level of chemical signal. Our piecewise framework allows for spatial variations within and between single-strain stripes.

As expected, we found that small-width stripes allowed for more coordination of consortium dynamics due to better communication between stripes over shorter distances (18, 27). Interestingly, our heterogeneous stripe test demonstrates consortium behavior can be controlled through the consortium's spatial organization alone. Combined with a rise in experimental methods for controlling spatial organizations in consortia (20, 23, 24, 38), this opens an interesting avenue for regulating consortium dynamics in the future (36).

An unexpected finding from our work was the strong influence of the basal multiplier on consortium coordination. From the results of Kim et al., it was expected that the presence or absence of the positive feedback loop would be sufficient to control coordination. Our simulation results indicate that the basal production rate had a stronger impact. Since the results of the basal multiplier test mirror that of the stripe width investigation (see Figs. 2C and 3B), we hypothesize that the observed behavior was due to variations in communication strength at various multiplier values. For lower values, the production rate of signaling molecule is lower, which results in weaker signaling between stripes. The opposite holds for higher production rates. The fact that Kim et al. included differences in basal production rates into their model may have obfuscated the actual impact of the positive feedback loop. However, since our model substantially simplifies the underlying gene circuits in exchange for computational efficiency and understandability, the basal multiplier results could be an artifact of this simplification.

Separate from this simplified model is the underlying framework of using piecewise-defined reaction-diffusion equations to incorporate spatial patterns into studies on consortium dynamics. Our ability to study various spatial patterns using this framework demonstrates its usefulness for modeling the spatiotemporal dynamics of microbial consortia. One strength of this framework is its flexibility. By varying the spatial points where the production terms are nonzero, a researcher can precisely control the spatial organization they are studying, including heterogeneous arrangements. Besides flexibility, another benefit of our model is its computational tractability. One strategy that previous studies have used to explicitly incorporate consortium spatial patterns is agent-based modeling (28–30). Although the fine-grained nature of these frameworks provides the greatest level of control when investigating the impact of spatial patterns, the complexity of these systems makes the study of larger consortia computationally infeasible (29, 31). Our continuum-based framework complements these studies and fills a gap in the literature by allowing for incorporation of spatial organizations into studies on the dynamics of large, spatially extended consortia.

Although we did not investigate other geometries or consortia in this study, we are confident the flexibility and computational tractability of our modeling framework will enable the study of more complicated systems. For example, our model could be used to investigate the effects of square size on consortium dynamics for the checkerboard pattern recently developed by Perkins et al. Besides allowing for a better understanding of these systems, our heterogeneous stripe results show that our modeling framework could potentially be used to engineer consortium behavior by finding spatial organizations that modulate the communication between consortium subpopulations in a desired fashion. Specifically, this can be done by making the distance between subpopulations either smaller or larger to modify the signaling strength between the subpopulations. We used this principle here to design a consortium that oscillated synchronously on the half that contained small stripes and asynchronously on the half that contained large stripes. Combined with a growing number of experimental approaches for designing consortia with specific spatial arrangements, including optogenic, seeding, and other approaches (20, 22–24), modulating consortium dynamics through spatial organization is becoming more practical. Since the underlying numerical scheme used to solve our reaction-diffusion equation model has routinely been applied in higher dimensions (39–42), our framework is well suited to interrogate the effects of the more complex spatial organizations implemented by these experimental methods (23, 43–45). Through enabling these investigations into the impact of spatial organization on microbial consortium dynamics, we hope that our modeling framework helps advance the understanding of these systems and supports the development of new applications involving them.

**CODE AVAILABILITY**

The input MATLAB scripts used in our simulations and representative examples from this work are available for download from GitHub: https://github.com/sdryan/StripeHeterogeneity.
SUPPORTING MATERIAL

Supporting Material can be found online at https://doi.org/10.1016/j.bpr.2022.100085.

AUTHOR CONTRIBUTIONS

R.G., B.R.K., and S.D.R. conceived the problem and built the mathematical model. R.G. and S.D.R. designed the numerical approach and conducted the numerical simulations. B.R.K. provided background in modeling microbial consortia. R.G. prepared the manuscript after careful discussion with B.R.K. and S.D.R. R.G., B.R.K., and S.D.R. edited the paper.

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DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental information

The space between us: Modeling spatial heterogeneity in synthetic microbial consortia dynamics

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1. Methods

**Determination of Parameter Values**

In our model, there were 14 parameter values that needed to be determined: \( \eta_{a0}, \eta_{a1}, \eta_{r0}, \eta_{r1}, K_a, K_r, n_a, n_r, D, d_e, K_e, n_e, \gamma, \tau \). These parameter values and their description are provided below in Table 1. The parameters \( K_a, n_a, n_r, d_e, n_e, \) and \( \tau \) were taken directly from Kim et al. Specifically, they were assigned the value of the biological parameter corresponding to their function in the model: \( K_a \) was assigned the EC50 value for \( P_{\text{Rhl/lac-s}} \) activation by C4, \( n_a \) was assigned the Langmuir-coefficient value for C4 activation of \( P_{\text{Rhl/lac-s}} \), \( d_e \) was assigned the value of the catalytic constant for C4 and C14 degradation by AiiA, \( n_e \) was assigned the Langmuir-coefficient value for \( P_{\text{cin}} \), activation by C14, and \( \tau \) was assigned the value of the time delay for transcription and translation. The values for \( K_e \) and \( K_r \) were chosen to give good qualitative agreement with the experimental results, and the remaining parameter values were calculated from parameters reported in Kim et al. These calculations are provided below.

| Parameter | Description | Value | Source |
|-----------|-------------|-------|--------|
| \( \eta_{a0} \) | Basal production rate of activator signaling molecule from promoter. | 985.56 nM min\(^{-1} \) | Calculated. |
| \( \eta_{a1} \) | Maximal production rate of activator signaling molecule from promoter in response to activator signal. | 17991.82/0 nM min\(^{-1} \) | Calculated |
| \( \eta_{r0} \) | Basal production rate of repressor signaling molecule from promoter. | 74.46 nM min\(^{-1} \) | Calculated |
| \( \eta_{r1} \) | Maximal production rate of repressor signaling molecule from promoter in response to activator signal. | 46495.80 nM min\(^{-1} \) | Calculated |
| \( K_a \) | EC50 of activator signaling molecule for production. | 5937 nM | Kim et al. |
| \( K_r \) | IC50 of repressor signaling molecule for repression. | 10 nM | |
| \( n_a \) | Hill coefficient of production for activator signaling molecule. | 4 | Kim et al. |
| \( n_r \) | Hill coefficient of repression for repressor signaling molecule. | 2 | Kim et al. |
| \( D \) | Diffusion coefficient signaling molecules. | 4080 \( \mu \)m\(^2\) min\(^{-1} \) | Calculated |
| \( d_e \) | Maximal rate of internal enzymatic degradation of signaling molecules. | 2257 min\(^{-1} \) | Kim et al. |
| \( K_e \) | EC50 of repressor signaling molecule for enzymatic degradation. | 1000 nM | |
| \( n_e \) | Hill coefficient of internal enzymatic degradation for repressor signaling molecule. | 4 | Kim et al. |
| \( \gamma \) | Inherent rate of degradation of signaling molecules caused by dilution from both media flow and cell growth. | 0.128 min\(^{-1} \) | Calculated |
| \( \tau \) | Time delay for transcription and translation. | 7.5 min | Kim et al. |

For diffusion, we made the simplifying assumption that that diffusion was the same for both
extracellular signaling molecules C4 and C14. Thus, we took the diffusion as the average of the reported diffusion values from Kim et al., $D_H$ for the activator (C4) and $D_I$ for the repressor (C14):

$$D = \frac{D_H + D_I}{2}$$  \hspace{1cm} (1)

$$= \frac{4800 \, \mu m^2 \, min^{-1} + 3360 \, \mu m^2 \, min^{-1}}{2}$$  \hspace{1cm} (2)

$$D = 4080 \, \mu m^2 \, min^{-1}$$  \hspace{1cm} (3)

The parameter value for non-enzymatic degradation, $\gamma$, was calculated by summing the contributions of dilution from cell-growth, $d$, and media flow, $\mu_e$:

$$\gamma = d + \mu_e$$  \hspace{1cm} (4)

$$= \ln(2)/25 \, min^{-1} + 0.1 \, min^{-1}$$  \hspace{1cm} (5)

$$\gamma = 0.128 \, min^{-1}$$  \hspace{1cm} (6)

The remaining parameters were the production rates for the extracellular signaling lactones: $\eta_{a_0}$, $\eta_{a_1}$, $\eta_{p_0}$, and $\eta_{p_1}$. These lactones upregulated or downregulated $P_{rhl/lac-s}$ which in turn produced the enzymes Rhll and Cinl which are synthases that catalyze the production of C4 and C14, respectively. Thus, to estimate production rates for C4 and C14 from the reported production rates for Rhll and Cinl in Kim et al., we made the simplifying assumption that each enzyme lasts 1 min on average before being degraded by ClpXP. With production rates for Rhll and Cinl of $16 \, min^{-1}$ and $2 \, min^{-1}$ this implies that each Rhll produces $16$ C4 on average and each Cinl produces $2$ C14 on average. Using this assumption, we can then calculate the production rates for C4 and C14 as follows. Note that in the following calculations, $S_R$ and $S_C$ were indeterminate scalings for the parameters that were estimated in Kim et al.. Additionally, $\eta_{R_x}$ and $\eta_{C_x}$ stand for the production rates of Rhll and Cinl from the promoters $P_{rhl/lac-s}$ and $P_{rhl/lac-w}$, respectively.

$$\eta_{a_0} = 16 \cdot \eta_{R_0}$$  \hspace{1cm} (7)

$$= 16 \cdot 20.13 \cdot S_R \, nM \, min^{-1}$$  \hspace{1cm} (8)

$$= 16 \cdot 20.13 \cdot 3.06 \, nM \, min^{-1}$$  \hspace{1cm} (9)

$$\eta_{a_0} = 985.56 \, nM \, min^{-1}$$  \hspace{1cm} (10)
\[ \eta_{a_1} = 16 \cdot \eta_R \]  
\[ = 16 \cdot 367.48 \cdot S_R \text{ nM min}^{-1} \]  
\[ = 16 \cdot 367.48 \cdot 3.06 \text{ nM min}^{-1} \]  
\[ \eta_{a_1} = 17991.82 \text{ nM min}^{-1} \]  

\[ \eta_{r_0} = 2 \cdot \eta_C \]  
\[ = 2 \cdot 1 \cdot S_C \text{ nM min}^{-1} \]  
\[ = 2 \cdot 1 \cdot 37.23 \text{ nM min}^{-1} \]  
\[ \eta_{r_0} = 74.46 \text{ nM min}^{-1} \]  

\[ \eta_{r_1} = 2 \cdot \eta_C \]  
\[ = 2 \cdot 624.44 \cdot S_C \text{ nM min}^{-1} \]  
\[ = 2 \cdot 624.44 \cdot 37.23 \text{ nM min}^{-1} \]  
\[ \eta_{r_1} = 46495.80 \text{ nM min}^{-1} \]  

**Non-Dimensionalization**

Our model was non-dimensionalized as follows. The activator and repressor concentrations were scaled by \( K_a \), time was scaled by \( \tau \), and space was scaled by \( \sqrt{D \cdot \tau} \). After introducing these scalings, the following system with non-dimensionalized parameters was obtained:

\[
\begin{align*}
\frac{\partial a}{\partial t} &= \frac{\eta_{a_{s_0}} + \eta_{a_{s_1}} a_T^{n_a}}{1 + a_T^{n_a} + \left( \frac{r_e}{K_{es}} \right)^{n_e}} + D_s \frac{\partial^2 a}{\partial x^2} - \frac{d_{es_a} \left( \frac{r_e}{K_{es}} \right)^{n_e}}{1 + \left( \frac{r_e}{K_{es}} \right)^{n_e}} - \gamma_s a \\
\frac{\partial r}{\partial t} &= \frac{\eta_{r_{s_0}} + \eta_{r_{s_1}} a_T^{n_r}}{1 + a_T^{n_r}} + D_s \frac{\partial^2 r}{\partial x^2} - \frac{d_{es_r} \left( \frac{r_e}{K_{es}} \right)^{n_e}}{1 + \left( \frac{r_e}{K_{es}} \right)^{n_e}} - \gamma_s r
\end{align*}
\]

The parameters in the above system are defined as follows:

\[ \eta_{a_{s_0}} = \frac{\eta_{a_0} \cdot \tau}{K_a} \]  

(24)
\[ \eta_{a1} = \frac{\eta_{a1} \cdot \tau}{K_a} \]  \hspace{1cm} (25)

\[ \eta_{r0} = \frac{\eta_{r0} \cdot \tau}{K_a} \]  \hspace{1cm} (26)

\[ \eta_{r1} = \frac{\eta_{r1} \cdot \tau}{K_a} \]  \hspace{1cm} (27)

\[ K_{rs} = \frac{K_r}{K_a} \]  \hspace{1cm} (28)

\[ D_s = 1 \]  \hspace{1cm} (29)

\[ d_{es} = d_e \cdot \tau \]  \hspace{1cm} (30)

\[ K_{es} = \frac{K_e}{K_a} \]  \hspace{1cm} (31)

\[ \gamma_s = \gamma \cdot \tau \]  \hspace{1cm} (32)

**Numerical Scheme and Synchronization Test Procedures**

All simulations were performed in MATLAB 2020a using custom scripts. To carry out the synchronization test for the \( P_2N_1 \) and \( P_1N_1 \) consortia, the non-dimensionalized model, Equation 23, was implemented numerically using the Crank-Nicholson scheme in one spatial dimension. We chose to discretize space by constructing our spatial mesh out of 5,000 points with a non-dimensional spatial step of \( \Delta x = \frac{2000 \mu m}{\sqrt{D \cdot \tau(5000)}} \) and a time step of \( \Delta t = 0.0001 \). Here \( D \) is the signal diffusion strength and \( \tau \) is the delay time. The spatial step was chosen so that the scaled length of the spatial domain was 2,000 \( \mu m \) long, equivalent to the length of the microfluidic trap used in Kim et al. Taking into account the design of the trap, Dirichlet (open) boundary conditions were used at the endpoints: \( x = 0 \) and \( x = 2,000 \mu m \). Additionally, we defined the spatial domain, \( \Omega \), to be piecewise by initially breaking it into 100 stripes of equal width \( (s_1, s_2, \ldots, s_{100}) \). The activator production terms \( (\eta_{a0}, \eta_{a1}) \) were then defined to be zero for the even numbered stripes while the repressor production terms \( (\eta_{r0}, \eta_{r1}) \) were defined to be zero for the odd numbered stripes. This
created 100, 20 µm wide stripes of alternating activator and repressor producing areas, thereby allowing us to incorporate the columnar spatial patterns observed in Kim et al.. For the \( P_1N_1 \) consortia, the production term \( \eta_{a_1} \) was set to 0 everywhere to account for the absence of the positive feedback loop.

Using this scheme, we carried out the synchronization test by first simulating the consortium under study (\( P_2N_1 \) or \( P_1N_1 \)) with initial values of \( a_{0,x} = 0 \) and \( r_{0,x} = 0 \) at all spatial points. This resulted in synchronous oscillation of the consortium across the trap. We then extracted the period of this oscillation post-transience, \( \phi \), and recorded the concentration of \( a \) and \( r \) for the middle spatial point at time \( 0.25\phi \). These concentrations were then used to initialize a second simulation. Specifically, in the second simulation, the first 2500 spatial points (the left half of the consortium) were initialized with concentrations \( a_{0,x} = 0 \) and \( r_{0,x} = 0 \). The next 2500 spatial points (the right half of the consortium) were initialized with the activator and repressor concentrations extracted from the previous simulation: \( a_{0,x} = a_{0,25\phi,2500} \) and \( r_{0,x} = r_{0,25\phi,2500} \). This resulted in the left and right halves of the consortium oscillating with the initial phase difference \( |\Delta \phi| = 25\% \).

To evaluate whether synchrony occurred, we tracked the activator concentration for the spatial points corresponding to the middle of the left and right sub-populations: \( x = 1250 \) and \( x = 3750 \). We then calculated the time difference, \( \Delta t_{\phi} \), between the minima corresponding to the 10th oscillation for these spatial points. This value was then used to calculate the final phase difference using:

\[
|\Delta \phi| = \frac{\Delta t_{\phi}}{\phi} \times 100
\]  

This process was repeated for both the \( P_2N_1 \) and \( P_1N_1 \) consortia. In this study, we assumed a phase difference of 2.5% of the period of oscillation, \( |\Delta \phi| < 0.025\phi \) corresponded to synchronous oscillation across the trap.

This synchronization test was repeated with slight variations to study the effect of changing the basal production rate and stripe width on the synchronization of the \( P_1N_1 \) and \( P_2N_1 \) consortia, respectively. To study the effect of changing the basal multiplier on the synchronization of the \( P_1N_1 \) consortium, we repeated the \( P_1N_1 \) simulations for different values of the basal production rate \( \eta_{a_0} \). Specifically, we scaled \( \eta_{a_0} \) by a basal multiplier, \( b_m \), and evaluated synchrony for values of \( b_m \) ranging from 0.1-10. To evaluate the effect of changing the stripe width on the synchronization of the \( P_2N_1 \) consortium, we repeated the \( P_2N_1 \) simulations and evaluated synchrony for stripe widths ranging from 200 µm to 10 µm. We took care to choose stripe widths that provided an even number of stripes. This allowed us to properly partition the consortium into two equal halves.
**Heterogeneous Stripes Test**

To evaluate the effect of heterogeneous stripes on coordination of the $P_2N_1$ consortium, we again divided the consortium into two halves. However, we modified our numerical scheme so that both halves consisted of stripes whose width increased according to a linear gradient. Specifically, if the half consisted of $n_x$ spatial points and would contain $k$ stripes, we first divided $n_x$ by the sum of the first $k$ numbers. This gave us a common difference, $d$, which we used to define the number of spatial points in the $k$-th stripe, $n_k$: $n_k = kd$. For this study, we used 84 stripes in the left half of the consortium and 14 stripes in the right. These numbers were chosen so that the linear gradient could be properly implemented in each half of the consortium with a manageable number of spatial points equally divided between the halves. This resulted in $n_x = 7,140$ equally spaced spatial points. For this test, we also chose to simulate a larger 4,000 $\mu$m consortium which changed our spatial step to $\Delta x = \frac{4000 \, \mu m}{\sqrt{D \cdot \tau(n_x)}}$. All other aspects of our numerical scheme were left unchanged. The numerically implemented system was then simulated with initial values of $a_{0,x} = 0$ and $r_{0,x} = 0$ at all spatial points.
Supporting References

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