Stochastic Turing patterns in a synthetic bacterial population

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The origin of biological morphology and form is one of the deepest problems in science, underlying our understanding of development and the functioning of living systems. In 1952, Alan Turing showed that chemical morphogenesis could arise from a linear instability of a spatially uniform state, giving rise to periodic pattern formation in reaction–diffusion systems but only those with a rapidly diffusing inhibitor and a slowly diffusing activator. These conditions are disappointing to achieve in nature, and the role of Turing instabilities in biological pattern formation has been called into question. Recently, the theory was extended to include noisy activator–inhibitor birth and death processes. Surprisingly, this stochastic Turing theory predicts the existence of patterns over a wide range of parameters, in particular with no severe requirement on the ratio of activator–inhibitor diffusion coefficients. To explore whether this mechanism is viable in practice, we have genetically engineered a synthetic bacterial population in which the signaling molecules form a stochastic activator–inhibitor system. The synthetic pattern-forming gene circuit destabilizes an initially homogenous lawn of genetically engineered bacteria, producing disordered patterns with tunable features on a spatial scale much larger than that of a single cell. Spatial correlations of the experimental patterns agree quantitatively with the signature predicted by theory. These results show that Turing-type pattern-forming mechanisms, if driven by stochasticity, can potentially underlie a broad range of biological patterns. These findings provide the groundwork for a unified picture of biological morphogenesis, arising from a combination of stochastic gene expression and dynamical instabilities.

Turing patterns | biofilm | synthetic biology | signaling molecules | stochastic gene expression

A central question in biological systems, particularly in developmental biology, is how patterns emerge from an initially homogeneous state (1). In his seminal 1952 paper “The chemical basis of morphogenesis,” Alan Turing showed, through linear stability analysis, that stationary, periodic patterns can emerge from an initially uniform state in reaction–diffusion systems where an inhibitor morphogen diffuses sufficiently faster than an activator morphogen (2). However, the requirements for realizing robust pattern formation according to Turing’s mechanism are prohibitively difficult to realize in nature. Although Turing patterns were observed in a chemical system in 1990 (3), the general role of Turing instabilities in biological pattern formation has been called into question, despite a few rare examples (ref. 4 and references therein).

Recently, Turing’s theory was extended to include intrinsic noise arising from activator and inhibitor birth and death processes (5–8). According to the resulting stochastic Turing theory, demographic noise can induce persistent spatial pattern formation over a wide range of parameters, in particular, removing the requirement for the ratio of inhibitor–activator diffusion coefficients to be large. Moreover, stochastic Turing theory shows that the extreme sensitivity of pattern-forming systems to intrinsic noise stems from a giant amplification resulting from the nonorthogonality of eigenvectors of the linear stability operator about the spatially uniform steady state (8). This amplification means that the magnitude of spatial patterns arising from intrinsic noise is not limited by the noise amplitude itself, as one might have thought naively. These developments imply that intrinsic noise can drive large-amplitude stochastic Turing patterns for a much wider range of parameters than the classical, deterministic Turing theory. In particular, it is often the case in nature that the activator and inhibitor molecules do not have widely differing diffusion coefficients; nevertheless, stochastic Turing theory predicts that, even in this case, pattern formation can occur at a characteristic wavelength that has the same functional dependence on parameters as in the deterministic theory.

To explore how global spatial patterns emerge from local interactions in isogenic cell populations, we present here a synthetic bacterial population with collective interactions that can be controlled and well-characterized (an introduction to this perspective is in ref. 9), where patterning is driven by activator–inhibitor diffusion across an initially homogeneous lawn of cells. Synthetic systems can be forward-engineered to include relatively simple circuits that are loosely coupled to the larger natural

Significance

In 1952, Alan Turing proposed that biological morphogenesis could arise from a dynamical process in reaction systems with a rapidly diffusing inhibitor and a slowly diffusing activator. Turing’s conditions are disappointingly hard to achieve in nature, but recent stochastic extension of the theory predicts pattern formation without such strong conditions. We have forward-engineered bacterial populations with signaling molecules that form a stochastic activator–inhibitor system that does not satisfy the classic Turing conditions but exhibits disordered patterns with a defined length scale and spatial correlations that agree quantitatively with stochastic Turing theory. Our results suggest that Turing-type mechanisms, driven by gene expression or other source of stochasticity, may underlie a much broader range of patterns in nature than currently thought.


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system into which they are embedded. This makes it easier to design and control the molecular underpinnings of the biological pattern phenomenon (10) and even front propagation phenomenon (11). Previous pattern formation efforts in synthetic biology have focused on oscillations in time (12) or required either an initial template (13) or an expanding population of cells (14), neither of which show a Turing mechanism.

**Experimental Results**

**Synthetic Biology of a Bacterial Community.** In our synthetic gene network design, which was guided by computational modeling (SI Appendix, section 1), we used two artificial diffusible morphogens: the small molecule N-(3-oxododecanoyl) homoserine lactone, denoted here as A3OC12HSL, and the small molecule N-butanoyl-l-homoserine lactone, denoted here as IC4HSL, from the *Pseudomonas aeruginosa* las and rhl quorum sensing pathways, respectively, in *P. aeruginosa* (15). A3OC12HSL serves as an activator of both its own synthesis and that of IC4HSL, while IC4HSL serves as an inhibitor of both signals (Fig. 1 A and B and SI Appendix, section 1). A3OC12HSL activates its own synthesis and synthesis of IC4HSL by binding regulatory protein LasR to form a complex that activates the hybrid promoter P_{las-OR1}. This promoter regulates expression of LasI, an A3OC12HSL synthase, and RhlI, an IC4HSL synthase. To increase the sensitivity of A3OC12HSL self-activation, LasR is regulated by a second copy of P_{las-OR1}. IC4HSL inhibits synthesis of A3OC12HSL and itself by forming a complex with the regulatory protein RhlR. This complex activates expression of lambda repressor CI, which in turn, represses transcription of LasI, RhlI, LasR, and RhlR. Pattern formation in our system can be modulated by altering the concentration of isopropyl β-D-1-thiogalactopyranoside (IPTG), a small molecule inducer that binds LacI and alleviates repression of P_{Rhl-lacO}–GFP and red fluorescent protein (RFP) are expressed from the rhl and las hybrid promoters, respectively, to aid in experimental observation (SI Appendix, section 2).

In our experimental setup, the A3OC12HSL activator diffuses more slowly than the IC4HSL inhibitor (SI Appendix, section 3). The estimated diffusion coefficient for A3OC12HSL is 83 μm²/s, and for IC4HSL, it is 1,810 μm²/s. The experimentally determined ratio of diffusion rates in our system of 21.6 is much higher than the value of 1.5 predicted by Wilke–Chang correlation in water (16), likely due to partitioning of A3OC12HSL in the cell membrane, which slows its diffusion from cell to cell (17). The slower diffusion rate of A3OC12HSL coupled with positive feedback regulating its synthesis allows A3OC12HSL to aggregate in local domains, leading to formation of visible red fluorescent spots (cellular lawn illustration is shown in Fig. 1C). Within these red domains, both A3OC12HSL and IC4HSL are found in high concentrations, but because A3OC12HSL competitively binds RhlR (SI Appendix, section 4 and Fig. S8), GFP is attenuated (18). The faster diffusion rate of IC4HSL allows it to diffuse into regions outside of the red fluorescent domains. Here, IC4HSL is free to bind RhlR, activating GFP expression. Collectively, these processes lead to green regions between red spots.

**Experimental Patterns and Controls.** To study pattern-forming behavior, engineered cells are first cultured in liquid media, and then, they are concentrated and plated on a petri dish to form a global cellular lawn. Cells express fluorescing proteins according to the profiles above and produce a global multicellular pattern. In control experiments, we show that our patterns are not simply a result of the outward growth of clusters of differently colored cells (Fig. 2 B and C and SI Appendix, section 5). Also, by performing an experiment with cells that harbor independent bistable green/red toggle switches, we test whether observable patterns would emerge if individual cells autonomously made cell fate decisions at some point after plating (SI Appendix, section 5). The fluorescence fields after 24 h of incubation at 30 °C in both control experiments are uniform, showing no emergence of patterns.

Next, we examine how changes in the strengths of localized interactions lead to different global outcomes in our pattern-forming gene circuit. In our system, IPTG can be used to modulate the inhibitory efficiency of IC4HSL in individual cells by affecting CI expression from P_{Rhl-lacO}. Specifically, IPTG relieves LacI repression of CI and GFP reporter. The increased range of CI ultimately increases inhibition of both morphogens, which is expected to decrease activator spot sizes, while causing...
the system at the communication signals and protein levels as follows:

\[
\frac{\partial U}{\partial t} = \alpha_u I_u - \gamma_u U + D_u \nabla^2 U \tag{1}
\]

\[
\frac{\partial V}{\partial t} = \alpha_v I_v - \gamma_v V + D_v \nabla^2 V \tag{2}
\]

\[
\frac{\partial I_u}{\partial t} = \alpha_{iu} F_1(X_1, C) - \gamma_{iu} I_u \tag{3}
\]

\[
\frac{\partial I_v}{\partial t} = \alpha_{iv} F_1(X_1, C) - \gamma_{iv} I_v \tag{4}
\]

\[
\frac{\partial C}{\partial t} = \alpha C_2(X_2, I) - \gamma C, \tag{5}
\]

where \( U \) and \( V \) are the concentrations of the two diffusible morphogens A3OC12HSL and IC4HSL, respectively; \( I_u \) and \( I_v \) are the concentrations of corresponding acyl homoserine lactone (AHL) synthases, respectively; and \( C \) refers to CI.

We model the hybrid promoters using the following Hill functions:

\[
\frac{I}{I_{\text{sat}} + [C]^n} + \frac{I}{I_{\text{sat}} + [C]^n} + \frac{I}{I_{\text{sat}} + [C]^n} + \frac{I}{I_{\text{sat}} + [C]^n}
\]

The field of CI and GFP reporter to be more strongly expressed. Our data show that mean GFP levels increase sigmoidally with inducer concentration, while the overall area of red spots decreases (Fig. 3 A and C). In addition to offering further support that our gene circuit gives rise to emergent patterns, these results show how pattern formation characteristics can potentially be tuned to fit future application needs.

**Theoretical Results**

Having established that our system forms emergent patterns, we proceeded to study the mechanisms driving these patterns. We formulated deterministic and stochastic models and analyzed our data to assess agreement with the theory of stochastic Turing patterns.

**Deterministic Model.** We first developed a detailed deterministic reaction–diffusion model (SI Appendix, section 7). The model explicitly describes chemical reactions for the LasI and RhlI synthases, regulatory protein CI, and synthesis and diffusion of the morphogens A3OC12HSL and IC4HSL. As the overall system involves a large number of reactions with rate constants that span multiple timescales, we made two commonly used simplifying assumptions. First, we assume that operator states of a promoter fluctuate much faster than protein degradation rates. Second, we assume that mRNA half-life is much shorter than protein half-life. These assumptions allow us to eliminate operator fluctuation and mRNA kinetics and model
patterns are formed but still within the regime where stochastic Turing patterns occur (5–8). Indeed, gene expression in microbes is inherently noisy due to the small volume of cells and the fact that many reactants are present in low numbers, suggesting that stochastic Turing patterns could be present in our system (24).

Noise in stochastic Turing patterns expands the range of parameters in which patterns form, in contrast to the usual expectation that noise serves as a destabilizing agent. The patterns observed in stochastic Turing systems correspond to the slowest decaying mode of the fluctuations. Similar noise stabilization phenomena can be observed in other systems that are out of equilibrium. For example, in predator–prey systems, fluctuations can drive temporal oscillations of populations (25, 26). Noise-driven stabilization has also been recently discovered in the clustering of molecules on biological membranes (27, 28) and in models that exhibit Turing-like pattern formation (7). In particular, whereas spatial symmetry breaking and pattern formation via the original Turing design require two morphogens with diffusion rates that differ by a large factor on the order of 10 or 100 (1), the requirements to form stochastic Turing patterns are less stringent. For example, in a pattern-forming plankton–herbivore ecosystem, the noise associated with discrete random birth and death processes reduces the required ratio of diffusion constants for pattern formation from a threshold of 27.8 for normal Turing patterns to a threshold of 2.48 for stochastic Turing patterns (5–8).

To determine whether noise in the chemical reactions underlying gene expression and morphogen diffusion in our system can cause the emergence of patterns over a wider range of parameters than a deterministic model, we constructed a stochastic spatiotemporal model using the same biochemical reactions, diffusion, and rate constants used in our deterministic model (SI Appendix, section 8). This model captures stochastic effects in the production and degradation of the proteins and morphogens in our system but approximates diffusion as deterministic. Simulations of the stochastic model generically produce patterns with large variability in spot size, shape, intensity, and intervals, which are similar to the patterns observed in our experiments and different from those predicted for the deterministic model (Fig. 3F). We have compared the experimental patterns with stochastic simulations in both real space and in 2D Fourier transform (2DFT) space (SI Appendix, section 8). Neither the experimental 2DFT nor the simulated 2DFT contain pronounced peaks that would be present in a deterministic honeycomb Turing pattern. Moreover, as the IPTG concentration is increased, both experimental and simulated patterns become more regular (Fig. 3G).

To further test the hypothesis that we are observing stochastic Turing patterns, we measured the power spectrum for both of our fluorescent reporters. Theory predicts that the power spectrum will have a power law tail as a function of wavenumber, k, for large wavenumbers, with an exponent characteristic of the noise source (7, 25). The exponent values are −2 and −4 for stochastic Turing patterns and deterministic Turing patterns with additive noise, respectively, and can be interpreted simply as follows. The −2 arises, because at small frequency or wavenumber, the random variable (i.e., concentration) is simply diffusing and therefore, follows the behavior of a random walk, which has a power spectrum that exhibits a −2 power law. The −4 arises, because for a system that is executing deterministic damped periodic motion but driven by additive white noise, the response of the random variable is a Lorentzian, with an asymptotic behavior for the power spectrum that exhibits a −4 exponent.

For the GFP channel, we observe a power law tail with an exponent of −2.3 ± 0.4 (Fig. 4B and SI Appendix, section 9A). For the RFP channel, we also observe a power law tail with an exponent of −3.9 ± 0.4 (SI Appendix, section 9A). To better understand the implications of these tails, we examined our detailed stochastic model of the genetic systems and also developed a reduced stochastic model that explicitly includes only the morphogens (SI Appendix, section 9B). Both models
predict that our experimental parameters will produce a stochastic pattern with a power law tail of −2 for both the activator and the inhibitor at asymptotically large wavenumbers (Fig 4C and SI Appendix, section 9C). However, in the range of parameters likely to correspond to the experiments (Fig. 4A and SI Appendix, section 9C), the detailed stochastic model predicts that the exponent of the power law tail for the activator will be −4 over a large range of intermediate wavenumbers before it eventually undergoes a cross-over to a power law with an exponent −2 at high wavenumbers (SI Appendix, Fig. S24). This behavior once again agrees with our experimental data and supports our identification of stochastic Turing patterns. In summary, spectral analysis of the patterns of activator and inhibitor is consistent with a model in which fluctuations in the amount of signaling morphogens drive stochastic Turing patterns.

Our analysis of the stochastic Turing model predicts that stochastic patterns form over a wide range of parameters (SI Appendix, section 8). Indeed, our stochastic model predicts that stochastic Turing patterns are possible at the measured ratio of diffusion rates for A3OC12HSL and IC4HSL (Figs. 3 and 4A). In addition, to determine the sensitivity of the stochastic model to the parameters chosen, we individually varied parameters from 0.5× their nominal value to 1.5× their nominal value while keeping all other parameters fixed at their best estimated value. For each set of parameters, we calculate the analytical power spectrum and the eigenvalues of the Jacobian (linear stability matrix) of the stochastic model evaluated at a fixed point found numerically. Based on this analysis (SI Appendix, section 8), we classify each set of parameters as producing an unstable homogeneous state at wavenumber $k = 0$, a stable homogeneous state, a stochastic Turing pattern, or a deterministic Turing pattern. Specifically, we classify a set of parameters as producing a pattern if they produce a peak in the calculated power spectrum at a nonzero wavenumber. To distinguish between stochastic Turing patterns and deterministic Turing patterns, we examine the eigenvalues of the corresponding Jacobian. If the real part of all of the eigenvalues is negative for all wavenumbers, then the pattern must be due to stochasticity. If there is any range of wavenumbers that have corresponding positive real parts of their eigenvalues, then the pattern is produced by the traditional Turing mechanism. The results of this analysis are shown in SI Appendix, Fig. S22 and illustrate the significant ranges for each parameter that can lead to stochastic Turing patterns. Indeed, the estimated parameter values yield stochastic Turing patterns and variation of $D_a$, $D_r$, and IPTG, and several other parameters never produce deterministic patterns; therefore, our results are very insensitive to estimation error of these important parameters. Overall, varying the parameters one at a time, 68% of the values yield stochastic Turing patterns.

To quantify the way in which stochasticity changes the pattern-forming regime of parameter space, we simultaneously varied all model parameters and performed the classification used above. Specifically, we used Latin hypercube sampling to randomly generate 500 parameter sets, where all of the parameters were allowed to vary between 0.5× and 1.5× their nominal value. For this analysis, we found that 24.8% of parameters produced unstable fixed points, 43.2% produced stable homogeneous states, 13.2% produced stochastic Turing patterns, and 18.8% produced Turing patterns. Thus, over this arbitrarily large range of parameters, pattern formation occurs only 18.8% of the time in the absence of stochasticity but 32% of the time when stochasticity is included. By including stochasticity, the range in which patterns can form has been increased by 70%.

Discussion

Alternative Hypotheses. Now, we consider alternative hypotheses to our claim that the theory of stochastic Turing patterns explains our experimental observations. We consider the duration and dynamics of our pattern formation experiments. One may expect to observe early events in Turing pattern formation, such as splitting of clusters or increases in intercluster distances. These processes may be, in fact, be taking place but may be difficult to observe due to weak reporter expression in the earlier stages. In addition, we must consider the limited duration of our experiments and the possibility that, theoretically, longer observations may result in different patterns if nonlinear processes eventually began to dominate dynamics. Indeed, we do not feed fresh nutrients to sustain the system for extremely long durations. However, as confirmed by analysis of the dynamics in SI Appendix, Fig. S12, cluster size growth and spacing between clusters appear to be stabilizing toward the end of the experiment. In addition, domains are neither created nor destroyed in the later time periods. Essentially, it appears that the patterns are close to stabilizing within the 32-h observation period.

Another alternative hypothesis is that cell growth dynamics primarily drive the observed pattern formation. Our control experiments with mixtures of red and green cell populations (SI Appendix, Fig. S9) along with our bistable switch control (SI Appendix, Fig. S10) suggest that cell growth does not explain our patterns. Moreover, our ability to tune pattern characteristics offers support for the fact that our patterns are not a simple consequence of natural biofilm growth morphologies but rather, are driven by our genetic circuit. However, growth may indeed impact regularity and may likely explain the fact that our
Summary of Evidence for Stochastic Turing Patterns. We summarize our evidence for showing stochastic Turing patterns and not showing amplification of random noise as follows. Our control experiments with mixtures of red and green cells (Fig. 2B and SI Appendix, Fig. S9) along with a bistable switch (SI Appendix, Fig. S10) did not produce the patterns that we observe with our genetic circuit (Fig. 2A). Our ability to tune pattern characteristics offers further support that pattern formation is driven by our genetic circuit. In addition to our experimental controls, we identify patterns in the stochastic model but not in the deterministic model of our system for the experimentally observed ratio of diffusion rates (Fig. 3F). These model patterns resemble the experimentally observed patterns in real space, exhibit no peaks in the 2DFT (SI Appendix, Figs. S13 and S20), and recapitulate the observed trend with IFTP variation. Analyses of our experimental data are also in accord with the theory of stochastic Turing patterns. The exponents in the tails of the experimental radial power spectra agree with theoretical predictions (Fig. 4B and SI Appendix, Fig. S13). In addition, although spatial regularity is weak, we observe a radial spectral peak for our experimental patterns (Fig. 4B and SI Appendix, Fig. S13), indicating a characteristic length scale. Furthermore, exploration of the large parameter space of the stochastic model indicates that the experimental parameters are most likely to be in the regime where only stochastic patterns can form (SI Appendix, section S). Collectively, this body of evidence suggests that our experiments indeed exhibit stochastic Turing pattern formation.

Materials and Methods

Strains and Conditions. Our patterning system was constructed using two plasmids that correspond to the upper and lower portions of the circuit diagram in Fig. 1B: pFNK512 and pFNK806 in Fig. 2A and pFNK512 and pFNK804/ΔlacO1ac in Fig. 3. The two-color bistable toggle switch plasmid pTOG-1 was constructed from plasmid pIK-E107. All plasmids were constructed using standard cloning and DNA recombination techniques. Plasmid construction details are described here and in SI Appendix, section 2. Escherichia coli strain MG1655 was used for all experiments.

Procedure. Cells harboring appropriate plasmids were initially grown in LB liquid media with corresponding antibiotics at 30 °C until OD 600 nm was reached 0.1 – 0.3. Cells were then concentrated and resuspended in M9 media with appropriate antibiotics (29); 0.5 mL of concentrated cell solutions OD600nm = 2.0 were poured onto a 2% M9 agar plate (60 × 15-mm petri dish) to form a cellular lawn. Plates were incubated at 30 °C, and fluorescence images were captured periodically. To examine the single-cell fluorescence evolution of toggle switch cell populations, we performed flow cytometry at the beginning of the experiment (0 h) and the end of the experiment (24 h).

Data Analysis. Fluorescence density plots, power spectrum of green fluorescence, averaged green fluorescence, total area of red spots, collectivity metric, and Moran’s I were all computed by analyzing the experimental time-lapse microscopy data with custom Matlab software.

Mathematical Modeling. The patterning system was simulated by numerically integrating differential equations using in-house-developed C software. We also developed stochastic spatiotemporal models using a hybrid stochastic simulation algorithm (30). SI Appendix has details about the models and the simulation environments.

Note Added in Proof. After the completion and acceptance of this work, an independent observation of stochastic Turing patterns in the cyanobacteria colonies of Anabaena sp. was reported (31).

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