Quorum Sensing Desynchronization Leads to Bimodality and Patterned Behaviors

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Abstract

Quorum Sensing (QS) drives coordinated phenotypic outcomes among bacterial populations. Its role in mediating infectious disease has led to the elucidation of numerous autoinducers and their corresponding QS signaling pathways. Among them, the Lsr (LuxS-regulated) QS system is conserved in scores of bacteria, and its signal molecule, autoinducer-2 (AI-2), is synthesized as a product of 1-carbon metabolism. Lsr signal transduction processes, therefore, may help organize population scale activities in numerous bacterial consortia. Conceptions of how Lsr QS organizes population scale behaviors remain limited, however. Using mathematical simulations, we examined how desynchronized Lsr QS activation, arising from cell-to-cell population heterogeneity, could lead to bimodal Lsr signaling and fractional activation. This has been previously observed experimentally. Governing these processes are an asynchronous AI-2 uptake, where positive intracellular feedback in Lsr expression is combined with negative feedback between cells. The resulting activation patterns differ from that of the more widely studied LuxIR system, the topology of which consists of only positive feedback. To elucidate differences, both QS systems were simulated in 2D, where cell populations grow and signal each other via traditional growth and diffusion equations. Our results demonstrate that the LuxIR QS system produces an ‘outward wave’ of autoinduction, and the Lsr QS system yields dispersed autoinduction from spatially-localized secretion and uptake profiles. In both cases, our simulations mirror previously demonstrated experimental results. As a whole, these models inform QS observations and synthetic biology designs.

Author Summary

Bacterial behavior is responsive to a multitude of soluble molecular cues. Among them are self-secreted autoinducers that control quorum sensing (QS) processes. While new quorum sensing systems are constantly being discovered, several systems have been well defined in terms of their molecular and genetic topologies, each influencing a variety of resultant phenotypes. These quorum sensing systems include LuxIR homologs that use an array of species specific autoinducers and Lsr system homologs that share a single
autoinducer among numerous species. Here we suggest that the regulatory topology of these two systems mark them as opposites of a sort. Whereas the LuxIR system bears a strong positive intercellular feedback mechanism, the Lsr system bears strong negative intercellular feedback. In our simulations these differences are manifested in distinct patterns of signaling. This was readily visualized in the outward spread of autogenous LuxIR expression in a growing bacterial 2D ‘colony’ whereas a dispersed activity was produced by autogenous Lsr expression in an otherwise identical colony. Here, this dispersed activity is a reflection of bimodal Lsr expression. We show that this bimodality could arise from desynchronized Lsr driven autoinducer import (intercellular negative feedback). This may have consequences on the arrangement of downstream phenotypes.

Introduction

Quorum sensing (QS) is a bacterial response to self-secreted signaling molecules known as autoinducers. While QS has been observed among individual bacteria in experimentally manipulated settings [1–3], QS often informs the coordination of processes that are metabolically burdensome and ineffectual for individual cells, yet beneficial at multicellular or population scales (e.g. virulence factor production and biofilm formation) [4]. Coordination arises from accumulated self-secreted autoinducer acting as a shared pool of extracellular signal. This regulatory strategy can focus phenotypic outcomes, reducing the effect of noise and organizing population activity [5,6]. This coordination sometimes involves the entire population such as with LuxIR QS, the signaling of which is defined by positive feedback mediated through autoinducing acylated homoserine lactones (AHL) [7] (Fig 1A). Behaviors that are driven by or occupy an entire population may not always be favored, however, as illustrated by instances of bet hedging [8] and role diversification [9]. For example, subpopulations are known to emerge in graded environments such as at the transition between biofilm margin and bulk [10,11].

The modes by which bacteria perceive and transduce the autoinducer signal can influence the extent of the QS behavior. In certain circumstances, Lsr based QS signaling via autoinducer-2 (AI-2) [12] (Fig 1B) for example, appears to generate activated and unactivated subpopulations, producing a bimodal distribution of QS activity [13,14]. Despite a fairly detailed understanding of the Lsr signal transduction process and the prevalence of Lsr in gammaproteobacteria [15,16] and AI-2 in eubacteria [17], the mechanisms underlying this coordinated fractional activation have been unexplored. Moreover, while the signaling network topology of many QS systems have been elucidated (e.g., LuxIR and Lsr systems) and mathematical models have delineated and compared a variety of QS processes [18–23], nominal consideration [24,25] has been given to Lsr QS, the extracellular negative feedback of which is unique among QS topologies.

Indeed, Lsr expression produces an AI-2 importer and kinase that work in tandem to de-repress the system, creating positive intracellular feedback. This same process draws down extracellular AI-2, generating negative intercellular feedback [26–29]. This is in addition to negative intracellular feedback from induced LsrR and LsrFG. We hypothesized that these intertwined feedback loops could provide a rationale for observations of bimodal Lsr activation [13]. We further hypothesized that the disparate characteristics of LuxIR and Lsr activity could result in distinctive expression patterns among subpopulations. In order to explore these hypotheses, models of Lsr and LuxIR were developed and compared.

Our simulations suggest that bimodal expression of the Lsr system is a consequence of desynchronized Lsr activation and subsequent autoinducer recompartmentalization and...
catabolism, which creates a competition for autoinducer. Here, desynchronization was imposed by cell-to-cell variation of select parameters. Moreover, using a 2D finite difference agent based model of cell growth and QS signaling, we also recapitulated the expansion phase of LuxIR QS activation from a cell colony center seen in an earlier experimental study [30]. Next when LuxIR signaling mechanisms were replaced with Lsr processes, cell colonies produced dispersed pockets of Lsr autoinduction. The "speckled" expression pattern was attributed to imposed population heterogeneity. Differences in population scale signaling patterns and responses to population heterogeneity were directly related to network signaling topology.

Model

Lsr QS. Deterministic simulations were drawn from an ODE system comprised of both statistical mechanics-like and phenomenological Michaelis-Menten and Hill function terms. Parameters were chosen to be biologically realistic and to conform to Lsr expression and AI-2 activities seen in batch cultures [13,25,27,31,32]. The rationale behind the ODE design choices and parameter values are discussed further in S1 Text. Our intent was to examine topology more so than describe a specific set of experimental results. Illustrative solutions are represented in Fig 2 where shortly after 4 hours of culture Lsr mRNA expression increased abruptly.
while the balance of AI-2 rapidly shifted from extracellular to intracellular. Examined briefly in the S1 Fig, this system was sensitive to multiple parameters controlling AI-2 synthesis, AI-2 export, AI-2 import, AI-2 phosphorylation, and Lsr transcription. The parameter basal (S1E Fig) represents the rate of AI-2 import through low-affinity pathways in E. coli, possibly through ribose transporters [33] or the phosphotransferase system [34]. Along with the rate of AI-2 synthesis, baseline expression of Lsr ABC-type importer, and AI-2 export through TqsA, basal AI-2 import through low affinity transporters is known to play an important role in the balance of AI-2 in the uninduced Lsr system, controlling extracellular AI-2 accumulation and Lsr activation [31,33,35]. Each of these influences was represented by a separate term in the ODEs.

This ODE model of Lsr activity was incorporated into an agent based simulation in a 500 x 500 μm 2D finite difference environment with an implied depth of 6 μm. Diagrammed in Fig 3A, space was discretized into 2x2 μm grid spaces and time into 0.0667 second steps. Also discretized was the net exchange of AI-2 between cells and the finite difference grid spaces in which they resided. Autoinducer diffusion was estimated by a central in concentration forward in time discretization, with the coefficient of diffusion estimated by the Wilke-Chang correlation [36].

Along with Lsr/AI-2 behavior, cell motility was also modeled at each time step. The distance traveled at each time step was drawn from a velocity distribution with an average of 20 μm per second and σ = 0.05 [37]. The direction (θ) at each step was uniformly distributed. Movement was constrained, however, by simulation boundaries that were modeled as impermeable. Cells with projected positions outside of the simulation boundary were retained within the 2D space by relocating them at twice the normal distance away from the boundary that they otherwise would have crossed (depicted in Fig 3B). Movement was further limited such that a cell could not enter a grid with three or more cells in order to control cell density.

That is, motility within a 2D colony was first modeled entirely as a consequence of cell growth and division. For cell division events in the colony interior, one daughter cell pushed adjacent cells outward towards the closest colony edge not abutting a simulation boundary. For cells at the colony boundary, duplicated cells randomly occupied free adjacent spaces. Each
time step also included the possibility of cell replication and changes in QS state, as indicated in Fig 3C. Simulations began with a cell number approximating a concentration of 0.03 OD$_{600}$. In swimming simulations, cells were initially randomly distributed. For simulations involving colony growth, a single colony was initiated randomly in the simulated 2D space. Cell division was accounted for by converting a population-wide Monod growth rate into a per-cell doubling time that was assumed to be normally distributed from cell to cell with $\sigma = 0.05$; time to division was marked by counters for each cell. Divided cells were assigned with the same characteristics as their parent, including state variable values, QS state, etc., except for a) any parameter varied in the simulated population, b) doubling time, and c) cell position.

QS state assignment was based on the level of Lsr transporter protein. Cells were considered QS ‘positive’, ‘activated’ or ‘induced’ if the Lsr transporter protein concentration exceeded 5 times their initial concentration, and QS ‘negative’ or ‘uninduced’ otherwise. This designation was for downstream analysis and did not otherwise influence model behaviors.

Represented in Fig 3D are trajectories for ten cells from the first two hours of simulations for colony growth (left) and unbiased movement (right). As previously described, rules for motility in a 2D colony consisted entirely of movement into neighboring spaces based on cell growth events. This heuristic yielded no substantial cell trajectories. For cells following swimming motility rules, trajectories reflected the free exploration of simulated space. Such trajectories did indeed appear random, as expected.

As noted, noise or heterogeneity was applied using a Monte Carlo method to assign basal or (other parameter values) from lognormal distributions as a reflection of cellular constituents’ natural variability [38–41]. The assigned parameter values were maintained throughout simulated time and were considered as average values for a particular cell over that time [42,43].

Fig 3. Schematic of agent based-finite difference simulation of Lsr activity. (A) Schematic of a representation of a cell within a grid element (bounded by dashed lines) moving a specific distance in a given direction (vector). The cell is exposed to the concentration of Al-2 within the grid element and conducts a net import or export according to a numerical solution for a set of ODE’s described in detail in S1 Text. (B) Multiple elements are depicted, each bearing a different shade of blue to reflect the concentration of Al-2 in each grid. Also represented is the treatment of motility at the impermeable simulation boundary (represented by solid line), where assignment of a cell motion across a boundary triggered a reflection in the opposite direction at twice the normal distance. (C) Cells also have a chance of replicating and changing their QS state in a given time step according to the values of state variables in the ODE’s. (D) Tracked motion for ten random cells from simulations of basic colony growth and unbiased movement ("swimming") for 200 minutes after initialization. Each line represents a single cell trajectory, and each segment represents an elapsed minute. Tracks are normalized to a (0,0) starting position to elucidate patterns of motion, and due to this scale ranges from -500 to 500 $\mu$m for each axis of each plot.

doi:10.1371/journal.pcbi.1004781.g003
**LuxIR QS.** LuxIR QS behavior was also simulated. Cells produced AHL at an average baseline rate ($\mu = 1 \, \mu M/min$); this value was also distributed over the entire population ($\sigma = 0.0225$, log-normal distributions) to the same degree as the parameter $\text{basal}$ in Lsr simulations. Once an intracellular AHL threshold of 2.9 $\mu M$ was exceeded, LuxIR expression was increased linearly so that a maximal rate of AHL synthesis ($10 \, \mu M/min$) was obtained after a thirty minute period. This artificially captured inherent system cooperativity and the time lag associated with transcription and translation. Diffusion across the membrane was modeled by an ODE with a conductivity of 0.6 between the intracellular and extracellular spaces driven by concentration difference [19]. Cells with LuxIR QS activity were considered to be ‘active’ or ‘induced’ when AHL production was five times the basal level.

LuxIR was only simulated in the context of colony growth (no swimming). As with Lsr simulations, newly divided cells shared the same characteristics as their parent, including state variable values, QS state (‘on’ or ‘off’), etc., with the exceptions of cell position (guided by colony growth dynamics alone), the uninduced rate of AHL production (which was varied between cells), and doubling time.

**Results**

**Lsr autoinduction with noise leads to fractional QS activation**

In order to simulate population scale phenomena, we employed a mix of two methodologies (a comprehensive set of ODEs simulating batch cultures and a 2D agent based approach, see Model section). First, ODEs were developed to reflect the trends of Lsr activation seen experimentally in batch cultures. To this, non-genetic heterogeneity was introduced as a type of noise and source of desynchronization. Specifically, the rate of AI-2 import through low affinity transporters (described by $\text{basal}$) among the cell population was varied log normally. This created a spectrum of Lsr-activation sensitivity to AI-2 and was posited as a means of generating bimodal Lsr activity. The distribution from which $\text{basal}$ values were drawn was varied in the standard deviation ($\sigma$) and mean ($\mu$) of its natural logarithm. This was done to find the variation necessary to reflect the previously reported bimodal activation [13], but also served to explore parameters to which fractional QS activation was sensitive and to interrogate underlying system mechanics.

In Fig 4A we depict distributions of $\text{basal}$ with different standard deviations from which parameter values were assigned. In Fig 4B, the effect of broadening $\text{basal}$ distributions on Lsr autoinduction is represented. This is a plot of the fraction of activated cells over time, including the mean trajectory and its variance. The broadest $\text{basal}$ distribution ($\sigma = 0.025$) produced the earliest onset of activation and the lowest final number of activated cells. A distinct trend was observed so that as the distribution became narrower, the onset of activation was delayed and a greater fraction of activated cells was obtained. Without any variation ($\sigma = 0$), Lsr QS was activated almost synchronously throughout the cell population. These results suggest that variation in $\text{basal}$ coincides with and is required for the development of bimodal signaling.

Varying the parameter $K_{\text{synth}}$ (the rate of AI-2 synthesis) as a stand-in for cell-cell heterogeneity instead of $\text{basal}$ generated similarly shifting patterns of Lsr activity. In Fig 5A, increasing standard deviations of $K_{\text{synth}}$ ($\sigma$ values ranging from 0.005 to 0.025) led to a decreasing fraction of the population ultimately becoming activated. The same range of standard deviation was also applied to the parameter $V_{\text{ygG}}$ (the rate of AI-2 export through YdgG). In Fig 5B, while the produced effect on fractional autoinduction was less than that for $K_{\text{synth}}$, increasing standard deviations of $V_{\text{ygG}}$ led to decreasing fractions of the population becoming autoinduced nonetheless. Notably, the parameters, $K_{\text{synth}}$ and $V_{\text{ygG}}$ affect the rate of AI-2 accumulation inversely (synthesis and export, respectively) which further supports the idea that, in general,
desynchronized Lsr expression (tested here by introducing QS related heterogeneity using three independent parameters) leads to bimodal activity.

Results from an ODE system modeling two Lsr networks sharing the same extracellular Al-2 pool highlighted the character of this bimodal signaling (Fig 6). The parameter basal of the second network was equal to or less than that of the first network, creating a difference in time to activation between the two systems. Increasing differences between the networks’ basal values, eventually led to the second network remaining inactive. That is, as basal of cell 2 decreased from that of cell 1 (510 μM/min) increases in the concentration of phospho-Al-2 in cell 2 stagnated, reflecting a two cell system where the first cell (cell 1) became autoinduced whereas the second cell (cell 2) remained uninduced.

Results presented in Figs 4 through 6 suggest that the more desynchronized a population is in its time to activation, the lower the final fraction of cells expressing Lsr at high levels. For basal distributions in particular, the broader the variation, the earlier cells with the highest values of basal will begin Lsr expression and the earlier extracellular Al-2 will be depleted. Conversely, cells with lower basal values will thereby be increasingly prevented from autoinduction, especially as they themselves are increasingly less sensitive to AI-2. Overall, this conceptual model of desynchronized QS activation serves as a basis to explain bimodal distributions of Lsr activation [13], and further bolsters the link between increased desynchronization and lower fractional activation. The exact fraction should depend on the compound heterogeneous system characteristics that desynchronize the timing of Lsr expression. In addition to changing the variation of the basal distribution, its mean value was also varied (specifically the mean the natural logarithm, Fig 7A). As seen in Fig 7B, these shifts in mean (μ = 6.17–6.27) also impacted the timing and fraction of cells that ultimately became Lsr activated. Since decreasing basal led to slower Lsr autoinduction in simple ODE simulations (S1 Text, S1E Fig), delayed initial Lsr expression upon decreasing the mean of the basal distribution was expected. The trend for the fraction of cells ultimately Lsr autoinduced was non-monotonic, however. Initially, decreasing mean and slower activation led to diminishing fractional activity. For the lowest tested means (μ = 6.15–6.17), however, further decrease and
delay of initial Lsr de-repression led to slightly higher final fractional activity, producing a mixed trend, indicating that competing factors were at play in the tested range.

When starting from distributions with higher mean, decreasing quorum activation was associated with decreasing intercellular distance. Fig 8A depicts the median of nearest neighbor cell-cell distances from twenty simulations. The gradually decreasing distance resulted from growth dynamics, which are independent of Lsr activity. Markers on the line describing cell-cell distance indicate when autoinduction first begins for systems with different basal means, with lower basal means leading to later Lsr autoinduction (as described before) during periods of smaller cell-cell distances. As described in Fig 7, in tested range of basal means, lower values also led to decreasing fractional activation. Decreasing cell-cell distance and decreasing Lsr fractional activation could be mechanistically linked by the local diminution of AI-2 effecting more cells at these later times. This interpretation is buttressed in the Supporting Information (S1 Text and S2 Fig), where smaller integrated cell-cell distance resulting from different
motility schemes led to consistently earlier initial activation and a smaller fraction of the population ultimately becoming Lsr-induced.

Any such effect was constrained, however. From previous figures, when basal distributions with the lower mean values were applied, we observed increasing fractional Lsr expression. Fig 8B, which represents the dynamics of extracellular AI-2 concentration per QS activated cell, provides further insight and suggests a plausible explanation. As the initiation of Lsr expression (shown by upward spikes in plot) was further and further delayed with decreasing mean basal, cell and extracellular AI-2 concentrations were increasingly elevated by the time Lsr-induction had begun. Under such conditions, the period between initial Lsr induction and significant reductions in extracellular AI-2 pools increased (represented by horizontal bars on plot). Effectively more cells had to become activated before AI-2 was drawn down to negligible levels.

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Fig 6. Solution to dual ODE system where basal of the second cell was reduced relative to the first cell. Results are from a pair of Lsr ODE’s sharing the same extracellular AI-2 but with different values for basal. Cell 1’s parameter values were held constant. Over separate simulations, cell 2’s parameter, basal (μM/min) was varied at comparatively lower values (in legend). The trajectory of Al2-P accumulation for each of these cases is represented. Rapid Al2-P increases reflects Lsr autoinduction.

doi:10.1371/journal.pcbi.1004781.g006

Fig 7. Changes in the Lsr-induced fraction of cells from changes to the basal mean. (A) The distribution of assigned basal values given select μ for the natural logarithm. (B) Presented results were derived from finite difference agent based simulations of Lsr activity in bacteria swimming in an unbiased manner. The fraction of the population QS activated over time was influenced by the mean of the natural logarithm, μ, of the log-normal distribution for the parameter basal, for μ values ranging from 6.15 to 6.27, run in triplicate. Dark lines represent the average values, whereas the lighter surrounding shade represents the standard deviation.

doi:10.1371/journal.pcbi.1004781.g007
concentrations. Put differently, as local depletion of AI-2 was delayed, more cells became Lsr-induced than would have if AI-2 depletion had been more sudden.

Simulated Lsr and LuxIR expression in colony growth

Lsr processes were further evaluated in the context of colony growth. Most models of colony growth attempt to capture the growth and motility dynamics involved in pattern formation found on variable solid or semi-solid media substrates [44–46]. The interest here laid in the likelihood that any coordination conferred by Lsr dynamics [23] is likely to depend upon the spatial organization of the cells involved [47,48]. To avoid the need to consider complex phenotypes associated with colony growth [45] only initial colony development was modeled using space filling constraints as described previously. Both Lsr and LuxIR signaling were studied. Originally identified in *V. fischeri* [49], LuxIR signaling has strong positive intercellular feedback [50], and is topologically distinct from Lsr signaling. As described in the Model section, LuxIR activation was modeled heuristically, incorporating essential system features including a threshold of autoinducer required for activation, a time delay, a basal rate of autoinducer production, and a maximal rate of autoinducer production.

A set of select images from a representative simulation for LuxIR signaling in colony growth is presented in Fig 9A. AHL concentration was initialized well below the activation threshold. Along with the selected rate of basal AHL production, this led to a couple hours delay before AHL accumulated above the QS activation threshold, accounting for the absence of QS signaling (yellow) early in initial colony growth (panels i-iii). Once initiated, however, LuxIR activity (red) spread quickly outward from its point of origin near the center of colony growth (due to positive feedback processes of cell growth and autoinduced AHL production), engulfing inactive neighbors in an outward wave of high AHL concentration, and coordinating population expression across an expanding front (panels iv-vi). The outward propagation of activity is evident in S1 Video. This activation pattern approximately recapitulated the initial expansion phase of activation seen using LuxIR engineered cells as noted above [23,30]. As in previous simulations [23,51], LuxIR initiation was biased towards simulated non-permeable boundaries.
This is reported in Fig 9B, which plots positions of the colony center versus that of the initial LuxIR activation for different simulations. For each type of position, the original ‘x’ and ‘y’ values for the colony center are mapped onto the abscissa axis, while original ‘x’ and ‘y’ values for the autoinduction center are mapped onto the ordinate axis. Original ‘x’ and ‘y’ positions from twenty simulations are marked by orange circles and blue squares, respectively. The dark line represents a linear fit to the data compared against a dashed, light gray 1:1::colony center::autoinduction center unskewed line.

doi:10.1371/journal.pcbi.1004781.g009

Fig 10. Lsr dynamics coupled with gliding during colony growth produces dispersed autoinduction. Images are from a representative simulation of Lsr induction in a growing bacterial colony. Lsr induced cells are in red, while uninduced cells are yellow. Over the course of the simulation, cells divided, duplicating the QS state and constituent concentration of their parent cells. As a result of division processes, cells pushed neighboring cells outward if necessary in the direction of the shortest path to unoccupied space.

doi:10.1371/journal.pcbi.1004781.g010
scattered throughout the remainder of simulated colony growth (panels iv-v). Underlying this expression pattern are the dynamics governing fractional activation as described earlier. Cells moved outward to available spaces upon cell division, and produced AI-2, which accumulated fastest near the colony center. The cells most sensitive to AI-2 became Lsr-induced, leading to the uptake of nearby AI-2. Proximal cells with less sensitivity were prevented from activation. This result provides a visual depiction of the bimodal phenotype arising from Lsr based AI-2 recompartmentalization and subsequent catalysis that comprises both positive and negative feedback [26,27]. Moreover, this emphasizes the importance of spatial information in QS of all types [23,48].

Additional analysis attributes the differences in induction patterns to the different mechanisms by which QS systems deal with heterogeneity. This is shown in Fig 11 where the assigned values of the baseline AHL synthesis rate (LuxIR, Fig 11A) or the basal AI-2 uptake rate (Lsr, Fig 11B) are provided in histogram form. The vertical light blue lines represent the average and standard deviation of these rates. The vertical red lines correspond to the average and standard deviation for parameter values of the first cells to autoinduce. For both Lsr and LuxIR simulations the first cells to autoinduce had parameter values drawn from the higher end of each distribution. For LuxIR QS, however, the first cells to activate were not always those with the highest rate of baseline AHL synthesis, indicated by the slightly more central position of the average value along the distribution and the wider standard deviation compared to Lsr QS represented in Fig 11B. For Lsr QS, first activators had basal values only from the distribution tail (and carried a smaller standard deviation). In LuxIR/AHL activity the more centralized relative position and greater distribution implies that heterogeneity associated with the baseline rate of AHL production was somewhat blunted. On the other hand, the distribution of Lsr/AI-2 expression over the colony was strongly sensitive to the imposed cell heterogeneity. Quantification of the spatial heterogeneity of LuxIR and Lsr signaling in colony growth is presented in S3 Fig.

Despite the dissimilarities between colony growth and traditional experimental biofilm models [52], the development of dispersed Lsr autoinduction modeled here may nonetheless

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**Fig 11.** Comparison of parameter value bias among first induced cells in simulations of LuxIR and Lsr colony growth. Results from finite difference agent based simulations of LuxIR or Lsr activity in bacteria growing in a colony. Histograms of assigned parameter values (blue; event count of 10,000), with the corresponding average and standard deviation overlaid (light blue). Overlaid in red is the average parameter value of the first autoinduced cell and the corresponding standard deviation (n = 20). (A) In LuxIR simulations, where the base rate of AHL synthesis was varied, some of the first activators had parameter values less than the median indicating a smoothing effect by LuxIR on heterogeneity. (B) In Lsr simulations, where the basal rate of AI-2 import was varied, nearly all the first autoinduced cells bared the highest basal values.

doi:10.1371/journal.pcbi.1004781.g011
capture Lsr expression patterns in early adhering bacterial microcolonies [53]. As Lsr activity informs macrostructure development in *E. coli* biofilms [54] and may be important for biofilm development for other bacteria [55], whether patchiness in Lsr activity plays a role in the early development of amorphous patterning in biofilms remains of interest. These or similar simulations may be helpful in understanding Lsr signaling in biofilms, which would inherently involve spatial dynamics [56].

**Discussion**

Our simulation results of bimodal autoinduction within a clonally homogenous cell population are consistent with experimental observations showing that the accessibility of QS activatable cells to autoinducer can define the fraction of the total cell population that generates QS phenotypes [13,57]. In Fig 2, we show results for solutions to an ODE system modeling Lsr signaling among cells in a batch culture. In Figs 4 and beyond (with the exception of Fig 6) we applied the same ODE systems into 2D agent based models to more readily visualize effects of cell movement and autoinducer diffusion. This was done by modeling the system as a collection of discrete cells having identical growth, AI-2 synthesis, and import parameters, while also explicitly accounting for diffusion and motility. Using these agent based models in Figs 4, 5 and 7 we demonstrated that heterogeneity within an otherwise clonal population, embodied here by a distribution in the basal AI-2 import rate among cells, led to a subpopulation of cells taking in AI-2 and conducting QS activity while other cells remained unactivated. The net result was fractional Lsr autoinduction at the population scale.

We note that bimodal expression arising from clonal origins is a common phenomenon, sometimes associated with pattern formation and differentiation in multicellular organisms [58,59], arising as a consequence of nonlinear responses to heterogeneity or asymmetry [60,61]. For the Lsr system, results indicate that bimodal expression arises when the intercellular negative feedback associated with Lsr activated AI-2 uptake and catabolism is desynchronized across a population (here represented by varied rates of AI-2 uptake and subsequent Lsr mediated gene expression).

Such bimodal expression may represent role diversification within the population. Bacterial diversification is frequently framed within the context of bet hedging [8] or as a graded response to environmental conditions, an example being the different conditions at the margins of a biofilm compared to those within the bulk [10]. While Lsr signaling does influence biofilm development in *E. coli* [62], whether population diversification through Lsr QS represents bet-hedging or specialization within the context of a population-wide transition to a sessile lifestyle remains of interest. As Lsr homologs are found in numerous bacteria [15], the heterogeneous influence of Lsr and Lsr-related systems on emergent phenotypes is likely to vary from species to species, and may only be effective in specific settings and circumstances.

By transitioning simulations to a 2D agent based model, the consequence of differences between Lsr and LuxIR mediated QS activation were made clear (Figs 9–11). For LuxIR signaling in colony growth simulations, the colony expanded its footprint as cells doubled and at the same time produced AHL that promoted increased AHL synthesis via LuxIR expression. The net phenotype was a sudden outward expansion of LuxIR expression initiated in colony centers, the regions of longest cell residence and highest accumulation of autoinducer. Instead, as governed by an autoinducer requiring active transport, Lsr system activity generated dispersed autoinduction, creating patterns similar to that of phase repulsive coupling in epileptic astrocyte cultures [63]. These distinctive spatiotemporal patterns are attributed to intracellular positive feedback processes also producing intercellular negative feedback in Lsr signaling, whereas LuxIR QS dynamics generate only positive feedback (recall Fig 1). Overall, while autoinducer
clearly has a coordinating and organizing effect on cell and Lsr activity, autoinduced Lsr expression is only experienced by a fraction of the cell population, and in a manner that appears to favor dispersion of that expression in space.

Ancillary interpretations also stem from our results; though speculative, they are worthy of discussion. For instance, since desynchronization appears to result in bimodal signaling due to induced AI-2 import and catalysis, the phenomenon would seem to be sensitive to the induced rate of AI-2 import, with slower import decreasing negative intercellular feedback. This intuition bears out insofar as when the induced rate of AI-2 import is lower a greater desynchronization (via increased basal heterogeneity) is required to achieve fractional Lsr induction. In all other simulations presented here, the induced rate was orders of magnitude greater than the basal rate. Using an induced rate less than 6 times the basal rate (by altering one parameter), significantly greater basal heterogeneity was required to produce a modest non-activating population (S4 Fig). Interestingly, heterogeneity for a glucose importer was reported by FACS to be of the same order of magnitude [39]. Moreover, tracking of heterogeneous expression within a population reveals that for the bulk of cells expression levels relative to the overall distribution are qualitatively stable over the time scales modeled here [42], with the exception of outlying high or low expressers which may be possibly attributable to regulatory mechanisms [40]. Moreover, while this study simplistically focuses on the heterogeneity of a single molecular species, the cells are composed of numerous components. We have not explored whether heterogeneity associated with multiple components produces different results.

Among other larger ideas, that bimodal expression evaluated here arises from a mechanism creating both positive intracellular and negative intercellular feedback indicates that Lsr expression may represent an exception or twist to the idea that bimodality is primarily a consequence of positive feedback regulation [60,64,65]. That is, while positive feedback clearly remains an integral feature of Lsr activity, this study may be seen to suggest that in examining conditions for bimodality, extracellular interactions should be considered for cases where inducer depletion results from induction, as nontrivial patterns may arise [63].

Finally, as previously mentioned, Lsr system homologs are plentiful among gammaproteobacteria, while AI-2 production occurs among eubacteria more broadly. That being the case, results here lend support the idea that in some circumstances the Lsr system could represent an intraspecies quorum sensing mechanism that uses a “universal” signaling molecule. This would seem to have ramifications for QS cheating and competition more broadly, which this model could be used to interrogate in future studies.

Supporting Information

S1 Fig. System sensitivity to parameter changes that manifested in shifting time to Lsr autoinduction. Solutions to ODE’s modeling Lsr activity, where rapidly increasing phosphorylated AI-2 concentration indicates Lsr autoinduction. Shifts in the time to activation were associated with reported parameter value changes. Changes to $k_2$ (transcription Hill parameter; A), $k_{phos}$ (phosphorylation; B), $K_{synth}$ (AI-2 synthesis; C), $V_{ydG}$ (AI-2 export; D), and basal (low affinity AI-2 import; E) are presented.

(TIF)

S2 Fig. Measures of the difference between different modes of motility when coupled with Lsr/AI-2 dynamics. Results were derived from finite difference agent based simulations of Lsr activity. (A) The fraction of the population that was QS activated over time, in Lsr simulations of different motility, with average values set in the darker lines and the standard deviation represented by lighter surrounding shades (n = 20). (B) The median minimum cell-cell distance for populations influenced by different combinations of motility and AI-2 uptake. Dark lines
are average value and the surrounding lighter shades reflect the corresponding standard deviation (n = 20). For example, cells undergoing colony growth had a predefined, regular distance between them, thus a single value prevailed across the entire time course and variability was zero.

S3 Fig. Local heterogeneity of Lsr versus LuxIR QS activation in colony growth. Results were derived from finite difference agent based simulations of LuxIR activity or Lsr activity using a median basal of 487.8 and a coefficient of variance of 0.052 (σ = 0.0225) for bacteria growing in a colony. (A) The dark lines represent the average local heterogeneity of 20 simulations, while the lighter, surrounding shades represent the standard deviation of those values. Also noted are the percentage of QS activity at the plateau of heterogeneity for Lsr simulation (represented in blue) and the percentage QS activity at the peak of heterogeneity for LuxIR simulations (represented in red). This is relevant, since the measure of local heterogeneity used is sensitive to the fraction of QS activation. This is seen in (B) for measures of colonies wherein QS state was assigned for each cell with a probability reflecting the percent QS activation. For enhanced context, the blue dot represents the heterogeneity for Lsr at plateau, whereas the red dot represents the heterogeneity for LuxIR at 50% activation.

S4 Fig. Slower induced import results in less negative intercellular feedback and fewer induced cells. Results were drawn from finite difference agent based simulations of Lsr activity using a reduced rate of induced AI-2 import. The fraction of the population that was QS activated over time given changes to the variation for the distribution of the basal rate of AI-2 import. Variation was shifted over a range from 0.0 to 0.35.

S5 Fig. Comparison of solution for population with a single basal value versus a population with a unimodally distributed value of basal. Juxtaposition of the solution for extracellular AI-2 for a simulation of cells with a single basal value versus the average solution of extracellular AI-2 for a simulation of cells with a log normal distribution of the parameter basal

S6 Fig. Comparison of results from single versus multiple finite difference elements to define environment. The average trajectory of AI2-P for cells with the same parameter sets in simulations where the environment was defined as either a single finite difference element or by the standard array of elements as defined in the methods. Modeling with a single finite difference element eliminates spatial noise as a source of difference between cells. The addition of noise through the full implementation of finite difference elements, adds spatially associated noise to the simulation. This did not result in a significant change in the average trajectory of AI2-P.

S7 Fig. Congruence of solution from finite difference-agent based modeling versus implicit solution of pure ODE’s. A AI2-P trajectory from implicit numerical methods and the average AI2-P concentration from the finite difference-agent based approach. Here, cells from the finite-difference-agent based solution all held the same parameter values as that from the pure ODE solution. In the pure ODE approach, cells were modeled as a dependent variable. Ideally, the two solutions would bear identical traces. B The rate to activation was assessed by fitting the function, f(t), from 12–152 minutes to a first order linear regression, g(t). The first time point at which f(t)-g(t)>2g(t) was considered the point of activation. The time to activation for
each value of basal was calculated and the bearing on the solution by the modeling and numerical method used was evaluated by direct comparison along the primary axis and according to the ratio of activation times for the finite difference-agent based solution to the pure ODE solution on the secondary axis.

S1 Figure Source File. Excel file of data used for figures.

S1 Table. Table of parameters and parameter values.

S1 Text. Supporting information text.

S1 Video. Video of LuxIR QS dynamics coupled with gliding during colony growth. Video of a representative simulation for LuxIR/AHL dynamics coupled with colony growth. QS active cells are in red, whereas inactive cells are in yellow. The complete simulated time of the video is 400 minutes.

S2 Video. Video of Lsr QS dynamics coupled with gliding during colony growth. Video of a representative simulation for Lsr/AI-2 dynamics coupled with colony growth. QS active cells are in red, whereas inactive cells are in yellow. The complete simulated time of the video is 400 minutes.

S1 XML File. SBML format XML file of Lsr system model.

Acknowledgments
We would like to thank Dr. Jonathan Alberts for generously providing his Sim2D framework. We also like to thank Dr. Amin Zargar and Dr. Jessica Terrell for useful discussions about the manuscript.

Author Contributions
Conceived and designed the experiments: DNQ CYT HCW WEB. Performed the experiments: DNQ. Analyzed the data: DNQ. Contributed reagents/materials/analysis tools: DNQ. Wrote the paper: DNQ CYT WEB.

References
1. Boedicker JQ, Vincent ME, Ismagilov RF. Microfluidic Confinement of Single Cells of Bacteria in Small Volumes Initiates High-Density Behavior of Quorum Sensing and Growth and Reveals Its Variability. Angew Chem Int Ed. 2009; 48(32): 5908–11.
2. Carnes EC, Lopez DM, Donegan NP, Cheung A, Gresham H, Timmins GS, Brinker CJ. Confinement-induced quorum sensing of individual Staphylococcus aureus bacteria. Nat Chem Biol. 2009; 6(1): 41–4. doi:10.1038/nchembio.264 PMID: 19935660
3. Hagen SJ, Son M, Weiss JT, Young JH. Bacterium in a box: sensing of quorum and environment by the LuxI/LuxR gene regulatory circuit. J Biol Phys. 2010; 36(3):317–27. doi: 10.1007/s10866-010-9186-4 PMID: 21629592
4. Darch SE, West SA, Winzer K, Diggle SP. Density-dependent fitness benefits in quorum-sensing bacterial populations. Proc Natl Acad Sci. 2012; 109(21):8259–63. doi: 10.1073/pnas.1118131109 PMID: 22566647
5. Tabareau N, Slotine J-J, Pham Q-C. How synchronization protects from noise. PLoS Comput Biol. 2010; 6(1):e1000637. doi: 10.1371/journal.pcbi.1000637 PMID: 2090826

6. Tanouchi Y, Tu D, Kim J, You L. Noise reduction by diffusional dissipation in a minimal quorum sensing motif. PLoS Comput Biol. 2008; 4(8):e1000167. doi: 10.1371/journal.pcbi.1000167 PMID: 18769706

7. Fuqua WC, Winans SC, Greenberg EP. Quorum sensing in bacteria: the LuxR-Lux family of cell density-responsive transcriptional regulators. J Bacteriol. 1994; 176(2):269. PMID: 8288518

8. Veening JW, Smits WK, Kuipers OP. Bistability, epigenetics, and bet-hedging in bacteria. Annu Rev Microbiol. 2008; 62:193–210. doi: 10.1146/annurev.micro.62.081307.163002 PMID: 18537474

9. Stoodley P, Sauer K, Davies D, Costerton JW. Biofilms as complex differentiated communities. Annu Rev Microbiol. 2002; 56(1):187–209.

10. de Beer D, Stoodley P. Relation between the structure of an aerobic biofilm and transport phenomena. Water Sci Technol. 1995; 32(8):11–8.

11. Ley RE, Harris JK, Wilcox J, Spear JR, Millers SR, Bebout BM, Maresca JA, Bryant DA, Sogin ML, Pace NR. Unexpected diversity and complexity of the Guerrero Negro hypersaline microbial mat. Appl Environ Microbiol. 2006; 72(5):3685–95. PMID: 16672518

12. Taga ME, Semmelhack JL, Bassler BL. The LuxS–Ley RE, Harris JK, Wilcox J, Spear JR, Millers SR, Bebout BM, Maresca JA, Bryant DA, Sogin ML, Pace NR. Unexpected diversity and complexity of the Guerrero Negro hypersaline microbial mat. Appl Environ Microbiol. 2006; 72(5):3685–95. PMID: 16672518

13. Tsao C-Y, Hooshangi S, Wu H-C, Valdes JJ, Bentley WE. Autonomous induction of recombinant protein by minimally rewiring native quorum sensing regulon of E. coli. Metab Eng. 2010; 12(3):291–7. doi: 10.1016/j.ymben.2010.01.002 PMID: 20060924

14. Servinsky MD, Terrell JL, Wu H-C, Quan DN, Tsao C-Y, Allen P, Byrd CM, Sund C, Zargar A, Bentley WE. Directed Assembly of a Bacterial Quorum. ISME J. 2016; 10(1):158–69. doi: 10.1038/ismej.2015.89 PMID: 26048256

15. Quan DN, Bentley WE. Gene network homology in prokaryotes using a similarity search approach: Queries of quorum sensing signal transduction. PLoS Comput Biol. 2012; 8(8):e1002637. doi: 10.1371/journal.pcbi.1002637 PMID: 22916001

16. Pereira CS, de Regt AK, Brito PH, Miller ST, Xavier KB. Identification of functional LsrB-like autoinducer-2 receptor. J Bacteriol. 2009; 191(22):6975–87. doi: 10.1128/JB.00379-09 PMID: 19749048

17. Winzer K, Hardie KR, Burgess N, Doherty N, Kirke D, Holden MT, Linforth R, Cornell KA, Taylor AJ, Hill PJ. LuxS: its role in central metabolism and the in vitro synthesis of 4-hydroxy-5-methyl-3(2H)-furanone. Microbiology. 2002; 148(4):909–22.

18. Drees B, Reiger M, Jung K, Bischols IB. A modular view of the diversity of cell-density-encoding schemes in bacterial quorum-sensing systems. Biophys J. 2014; 107(1):266–77. doi: 10.1016/j.bpj.2014.05.031 PMID: 24983660

19. Dockery JD, Keener JP. A mathematical model for quorum sensing in Pseudomonas aeruginosa. Bull Math Biol. 2001; 63(1):95–116. PMID: 11146885

20. Klapper I, Dockery JD. Mathematical description of microbial biofilms. SIAM Rev. 2010; 52(2):221–65.

21. Hunter G, Vasquez FG, Keener J. A mathematical model and quantitative comparison of the small RNA circuit in the Vibrio harveyi and Vibrio cholerae quorum sensing systems. Phys Biol. 2013; 10(4):046007. doi: 10.1088/1478-3975/10/4/046007 PMID: 23820088

22. Pérez-Velázquez J, Quiñones B, Hense BA, Kuttler C. A mathematical model to investigate quorum sensing regulation and its heterogeneity in Pseudomonas syringae on leaves. Ecol Complex. 2015; 21:128–41.

23. Melke P, Sahlin P, Levchenko A, Jonsson H. A cell-based model for quorum sensing in heterogeneous bacterial colonies. PLoS Comput Biol. 2010; 6(6):e1000819. doi: 10.1371/journal.pcbi.1000819 PMID: 20585545

24. Barrios AFG, Zuo R, Hashimoto Y, Yang L, Bentley WE, Wood TK. Autoinducer 2 controls biofilm formation in Escherichia coli through a novel motility quorum-sensing regulator (MqsR, B3022). J Bacteriol. 2006; 188(1):305–316. PMID: 16352947

25. Hooshangi S, Bentley WE. LsrR quorum sensing ‘switch’ is revealed by a bottom-up approach. PLoS Comput Biol. 2011; 7(9):e1002172. doi: 10.1371/journal.pcbi.1002172 PMID: 21982072

26. Wang L, Hashimoto Y, Tsao C-Y, Valdes JJ, Bentley WE. Cyclic AMP (cAMP) and cAMP receptor protein influence both synthesis and uptake of extracellular autoinducer 2 in Escherichia coli. J Bacteriol. 2005; 187(6):2066–76. PMID: 15743955

27. Xavier KB, Bassler BL. Regulation of uptake and processing of the quorum-sensing autoinducer AI-2 in Escherichia coli. J Bacteriol. 2005; 187(1):238–48. PMID: 15601708
28. Marques JC, Lamosa P, Russell C, Ventura R, Maycock C, Semmelhack MF, Miller ST, Xavier KB. Processing the interspecies quorum-sensing signal autoinducer-2 (AI-2) characterization of phospho-(S)-4, 5-dihydroxy-2,3-pentanedione isomerization by LsrG protein. J Biol Chem. 2011; 286(20):18331–43. doi: 10.1074/jbc.M111.230227 PMID: 21454635

29. Marques JC, Oh IK, Ly DC, Lamosa P, Ventura MR, Miller ST, Xavier KB. LsrF, a coenzyme A-dependent thiolase, catalyzes the terminal step in processing the quorum sensing signal autoinducer-2. Proc Natl Acad Sci. 2014; 111(39):14235–40. doi: 10.1073/pnas.1408691111 PMID: 25225400

30. Danino T, Mondragón-Palomino O, Tsimring L, Hasty J. A synchronized quorum of genetic clocks. Nature. 2010; 463(7279):326–30. doi: 10.1038/nature08753 PMID: 20090747

31. Wang L, Hashimoto Y, Tsao C-Y, Valdes JJ, Bentley WE. Cyclic AMP (cAMP) and cAMP receptor protein influence both synthesis and uptake of extracellular autoinducer 2 in Escherichia coli. J Bacteriol. 2005; 187(6):2066–76. PMID: 15743955

32. Wang L, Li J, March JC, Valdes JJ, Bentley WE. LuxS-dependent gene regulation in Escherichia coli K-12 revealed by genomic expression profiling. J Bacteriol. 2005; 187(24):8350–60. PMID: 16321939

33. Shao H, James D, Lamont RJ, Demuth DR. Differential interaction of Aggregatibacter (Actinobacillus) actinomycetemcomitans LsrB and RbsB proteins with autoinducer 2. J Bacteriol. 2009; 189(15):5559–65.

34. Pereira CS, Santos AJ, Bejerano-Sagie M, Correia PB, Marques JC, Xavier KB. Phosphoenolpyruvate phosphotransferase system regulates detection and processing of the quorum sensing signal autoinducer-2. Mol Microbiol. 2012; 84(1):93–104. doi: 10.1111/j.1365-2958.2012.08010.x PMID: 22384939

35. Herzberg M, Kaye IK, Peti W, Wood TK. YdgG (TqsA) controls biofilm formation in Escherichia coli K-12 through autoinducer 2 transport. J Bacteriol. 2006 188(2):587–98. PMID: 16385049

36. Wilke C, Chang P. Correlation of diffusion coefficients in dilute solutions. AIChE J. 1955; 1(2):284–270.

37. Boehm A, Kaiser M, Li H, Spangler C, Kasper CA, Ackermann M, Kaeve M, Sourjik V, Roth V, Jenal U. Second messenger-mediated adjustment of bacterial swimming velocity. Cell. 2010; 141(1):107–16. doi: 10.1016/j.cell.2010.01.018 PMID: 20303158

38. Limpert E, Stahel WA, Abbt M. Log-normal distributions across the sciences: Keys and clues on the charms of statistics, and how mechanical models resembling gambling machines offer a link to a handy way to characterize log-normal distributions, which can provide deeper insight into variability and probability—normal or log-normal: That is the question. BioScience. 2001; 51(5):341–52.

39. Nikolic N, Barner T, Ackermann M. Analysis of fluorescent reporters indicates heterogeneity in glucose uptake and utilization in clonal bacterial populations. BMC Microbiol. 2013; 13(1):258.

40. Lu C, King RD. An investigation into the population abundance distribution of mRNAs, proteins, and metabolites in biological systems. Bioinformatics. 2009; 25(16):2020–7. doi: 10.1093/bioinformatics/btp360 PMID: 19535531

41. Long T, Tu KC, Wang Y, Mehta P, Ong N, Bassler BL, Wingreen NS. Quantifying the integration of quorum-sensing signals with single-cell resolution. PLoS Biol. 2009; 7(3):e1000068.

42. Young JW, Locke JC, Altinok A, Rosenfeld N, Bacarian T, Swain PS, Mjolsness E, Elowitz MB. Measuring single-cell gene expression dynamics in bacteria using fluorescence time-lapse microscopy. Nat Protoc. 2012; 7(1):80–8.

43. Eijlander RT, Kuipers OP. Live-cell imaging tool optimization to study gene expression levels and spatiotemporal modulations of bacterial gene expression. Mol Biotechnol. 2008; 40(3):192–8. doi: 10.1007/s12033-008-9072-1 PMID: 18437261

44. Pipe LZ, Grimson MJ. Spatial-temporal modelling of bacterial colony growth on solid media. Mol Biosyst. 2008; 4(3):192–8. doi: 10.1039/b708241j PMID: 18437261

45. Bonachela JA, Nadell CD, Xavier JB, Levin SA. Universality in bacterial colonies. J Stat Phys 2011; 144(2):303–15.

46. Shapiro JA. The significances of bacterial colony patterns. Bioessays. 1995; 17(7):597–607. PMID: 7646482

47. Alberghini S, Polone E, Corich V, Carlot M, Seno F, Trovato A, Squartini A. Consequences of relative cellular positioning on quorum sensing and bacterial cell-to-cell communication. FEMS Microbiol Lett. 2009; 292(2):149–61. doi: 10.1111/j.1574-6968.2008.01478.x PMID: 19187204

48. Dilanj GE, Langebrake JB, De Leenheer P, Hagen SJ. Quorum activation at a distance: spatiotemporal patterns of gene regulation from diffusion of an autoinducer signal. J Am Chem Soc. 2012; 134(12):5618–26. doi: 10.1021/ja211593q PMID: 22372494

49. Devine JH, Shadel GS, Baldwin TO. Identification of the operator of the lux regulon from the Vibrio fischeri strain ATCC7744. Proc Natl Acad Sci. 1989; 86(15):5688–92. PMID: 2762291
50. Eberhard A, Longin T, Widrig CA, Stranick SJ. Synthesis of the lux gene autoinducer in Vibrio fischeri is positively autoregulated. Arch Microbiol. 1991; 155(3):294–7.

51. Trovato A, Seno F, Zanardo M, Alberghini S, Tondello A, Squartini A. Quorum vs. diffusion sensing: a quantitative analysis of the relevance of absorbing or reflecting boundaries. FEMS Microbiol Lett. 2014; 352(2):198–203. doi: 10.1111/1574-6968.12394 PMID: 24484313

52. Mikkelson H, Duck Z, Lilley K, Welch M. Interrelationships between colonies, biofilms, and planktonic cells of Pseudomonas aeruginosa. J Bacteriol. 2007; 189(6):2411–6. PMID: 17220232

53. Mondes RD, O’Toole GA. The developmental model of microbial biofilms: ten years of a paradigm up for review. Trends Microbiol. 2009; 17(2):73–87. doi: 10.1016/j.tim.2008.11.001 PMID: 19162483

54. Li J, Attila C, Wang L, Wood TK, Valdes JJ, Bentley WE. Quorum sensing in Escherichia coli is signaled by Al-2/LsrR: effects on small RNA and biofilm architecture. J Bacteriol. 2007; 189(16):6011–20. PMID: 17557827

55. Auger S, Kim E, Aymerich S, Gohar M. Autoinducer 2 affects biofilm formation by Bacillus cereus. Appl Environ Microbiol. 2006; 72(1):937–941. PMID: 16391139

56. Gu H, Hou S, Yongyat C, De Tore S, Ren D. Patterned biofilm formation reveals a mechanism for structural heterogeneity in bacterial biofilms. Langmuir. 2013; 29(35):11145–53. doi: 10.1021/la402608z PMID: 23919925

57. Luo X, Wu H-C, Tsao C-Y, Cheng Y, Betz J, Payne GF, Rubloff GW, Bentley WE. Biofabrication of stratified biofilm mimics for observation and control of bacterial signaling. Biomaterials. 2012; 33 (20):5136–43. doi: 10.1016/j.biomaterials.2012.03.037 PMID: 22507453

58. Albert R, Othmer HG. The topology of the regulatory interactions predicts the expression pattern of the segment polarity genes in Drosophila melanogaster. J Theor Biol. 2003; 223(1):1–18. PMID: 12782112

59. Lawrence PA, Casal J, Struhl G. Hedgehog and engrailed: pattern formation and polarity in the Drosophila abdomen. Development. 1999; 126(11):2431–9. PMID: 10226002

60. Birtwistle MR, Rauch J, Kiyatkin A, Aksamitiene E, Dobrzyński M, Hoek JB, Kolch W, Oggunnaie BA, Khodolenko BN. Emergence of bimodal cell population responses from the interplay between analog single-cell signaling and protein expression noise. BMC Syst Biol. 2012; 6(1):109.

61. Shalek AK, Satija R, Adiconis X, Gertner RS, Gaublomme JT, Raychowdhury R, Schwartz S, Yosef N, Malboeuf C, Lu D. Single-cell transcriptomics reveals bimodality in expression and splicing in immune cells. Nature. 2013. 498(7453), 236–240 doi: 10.1038/nature12172 PMID: 23685454

62. Li J, Attila C, Wang L, Wood TK, Valdes JJ, Bentley WE. Quorum sensing in Escherichia coli is signaled by Al-2/LsrR: effects on small RNA and biofilm architecture. J Bacteriol. 2007; 189(16):6011–20. PMID: 17557827

63. Balázs G, Cornell-Bell A, Neiman AB, Moss F. Synchronization of hyperexcitable systems with phase-repetitive coupling. Phys Rev E. 2001; 64(4):041912.

64. Kim KH, Sauro HM. In search of noise-induced bimodality. BMC Biol. 2012; 10(1):89.

65. Mitrophanov AY, Groisman EA. Positive feedback in cellular control systems. BioEssays News Rev Mol Cell Dev Biol. 2008; 30(6):542–55.