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RESEARCH ARTICLE

Influence of confinement on the spreading of bacterial populations

Daniel B. Amchin 1, Jenna A. Ott 1, Tapomoy Bhattacharjee², Sujit S. Datta 1, Tapomoy Bhattacharjee³, Suji

- 1 Department of Chemical and Biological Engineering, Princeton University, Princeton, New Jersey, United States of America, 2 Andlinger Center for Energy and the Environment, Princeton University, Princeton, New Jersey, United States of America
- * ssdatta@princeton.edu

Abstract

The spreading of bacterial populations is central to processes in agriculture, the environment, and medicine. However, existing models of spreading typically focus on cells in unconfined settings—despite the fact that many bacteria inhabit complex and crowded environments, such as soils, sediments, and biological tissues/gels, in which solid obstacles confine the cells and thereby strongly regulate population spreading. Here, we develop an extended version of the classic Keller-Segel model of bacterial spreading via motility that also incorporates cellular growth and division, and explicitly considers the influence of confinement in promoting both cell-solid and cell-cell collisions. Numerical simulations of this extended model demonstrate how confinement fundamentally alters the dynamics and morphology of spreading bacterial populations, in good agreement with recent experimental results. In particular, with increasing confinement, we find that cell-cell collisions increasingly hinder the initial formation and the long-time propagation speed of chemotactic pulses. Moreover, also with increasing confinement, we find that cellular growth and division plays an increasingly dominant role in driving population spreading—eventually leading to a transition from chemotactic spreading to growth-driven spreading via a slower, jammed front. This work thus provides a theoretical foundation for further investigations of the influence of confinement on bacterial spreading. More broadly, these results help to provide a framework to predict and control the dynamics of bacterial populations in complex and crowded environments.

Author summary

The spreading of bacteria through their environments critically impacts our everyday lives; it can be harmful, underlying the progression of infections and spoilage of foods, or can be beneficial, enabling the delivery of therapeutics, sustaining plant growth, and remediating polluted terrain. In all these cases, bacteria typically inhabit crowded environments, such as soils, sediments, and biological tissues/gels, in which solid obstacles confine the cells and regulate their spreading. However, existing models of spreading typically focus on cells in unconfined settings, and thus are frequently not applicable to cells

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in more complex environments. Here, we address this gap in knowledge by extending the classic Keller-Segel model of bacterial spreading via motility to also incorporate cellular growth and division, and explicitly consider the influence of confinement. Through numerical simulations of this extended model, we show how confinement fundamentally alters the dynamics and morphology of spreading bacterial populations—in particular, driving a transition from chemotactic spreading of motile cells to growth-driven spreading via a slower, jammed front. These results provide a foundation for further investigations of the influence of confinement on bacterial spreading, both by yielding testable predictions for future experiments, and by providing guidelines to predict and control the dynamics of bacterial populations in complex and crowded environments.

1 Introduction

The ability of bacterial populations to spread through their surroundings plays a pivotal role in our everyday lives. It can be harmful, underlying the progression of infection in the body [1–8] and the spoilage of foods [9, 10]. In other cases, it can be beneficial, enabling bacteria to deliver drugs to hard-to-reach spots in the body [11, 12], move toward and protect plant roots in soil [13–17], and degrade environmental contaminants [18–22]. Therefore, the development of accurate models of spreading is critically important for the prediction and control of bacterial populations in medicine, food, agriculture, and the environment.

One common way in which many bacteria spread is through active motility using, for example, flagellar propulsion in liquids or pili-mediated propulsion on solids. Populations of cells can thereby direct their spreading via chemotaxis: as the cells continually consume a surrounding chemoattractant, such as a nutrient or oxygen, they collectively generate a local gradient that they, in turn, bias their motion along. This biased motion leads to the spectacular formation of a coherent pulse of cells that continually propagates, enabling populations to escape from harmful conditions and colonize new terrain [23–28]. Building on the seminal work of Keller and Segel in 1971, continuum-scale models have been developed that can successfully capture the key features of this chemotactic spreading in bulk liquids [26, 27, 29–33]; hence, such models form a cornerstone of theoretical studies of emergent phenomena and collective behavior in biology.

However, while such models and the lab studies that they describe typically focus on cells in unconfined environments, bacteria often inhabit more crowded settings—such as porous gels and tissues in the body, micro- and meso-porous foods, and soils, sediments, and subsurface formations in the environment—in which a solid matrix obstructs and confines the motion of cells. Depending on the degree of confinement, bacterial populations may still be able to spread via chemotaxis [28, 34], but with two notable differences, as revealed by recent experiments. First, as confinement increases, collisions with the solid matrix increasingly impede the spreading of individual cells [35-38]. Second, also as confinement increases, the amount of free space available for cells to move through decreases, promoting cell-cell collisions that further impede cellular spreading [28]. Indeed, if the cells are sufficiently confined and their local density is sufficiently high, cell-cell collisions dominate and they ultimately become jammed, unable to self-propel at all [39, 40]—abolishing chemotaxis entirely. In this case, the population instead spreads solely through cellular proliferation, referred hereafter as 'growth' for brevity, in which metabolically-active cells grow, divide, and push each other to new locations [41, 42]. Hence, confinement—which is an inherent feature of many bacterial habitats—is a strong regulator of population spreading.

Despite its widespread use, the classic Keller-Segel model of chemotactic spreading treats cells as being non-contacting, does not explicitly incorporate confinement, and does not consider population spreading via growth. Previous work took a useful first step toward extending this model by modifying the underlying transport parameters to incorporate the influence of cell-solid collisions [34]. Nevertheless, due to the limited understanding of single-cell motility in confinement at the time, this approach necessarily relied on *ad hoc* approximations; moreover, it did not incorporate cell-cell collisions/jamming or growth. While a previous study [43] has addressed some aspects of the role of growth in a slow-moving front, it did not examine the influence of confinement or crowding—which can influence the relative importance of motility versus growth in driving spreading. As a result, there remains a need for models that can more accurately describe the spreading of bacteria, both by motility and growth, in crowded and highly-confining environments.

Here, we present an extended version of the classic Keller-Segel model that takes a first step toward addressing these gaps in knowledge: it describes bacterial spreading via *both* motility and growth, and explicitly incorporates the influence of confinement on spreading by considering both cell-solid and cell-cell collisions, motivated by recent experimental observations. We identify key dimensionless parameters emerging from this extended model that describe bacterial spreading. Furthermore, by numerically solving the model, we show how confinement fundamentally alters the dynamics and morphology of spreading bacterial populations. In particular, with increasing confinement, we find that cell-cell collisions increasingly hinder the initial formation and the long-time propagation speed of chemotactic pulses. Moreover, also with increasing confinement, growth plays an increasingly dominant role in driving population spreading compared to cellular motility—eventually leading to a transition from chemotactic spreading to growth-driven spreading via a slower, jammed front. Thus, our work provides a foundation for future investigations of the influence of confinement, and yields quantitative principles that could guide the prediction and control of bacterial spreading in crowded and complex environments.

2 Methods

2.1 Classic Keller-Segel model, also incorporating growth

Two forms of the Keller-Segel model have been explored in the prior literature to describe two distinct biophysical processes: one describes cellular aggregation and pattern formation in response to chemoattractant produced by the cells themselves [44–46], while the other describes cellular spreading in response to an exogenous chemoattractant that is not produced, but just consumed, by the cells [26, 27, 29–33]. Here, we focus on the latter case. Before considering confinement, we first describe how chemotactic spreading is typically modeled using this form of the classic one-dimensional Keller-Segel model—which does not incorporate the influences of growth and confinement, but can successfully capture the key features of experiments on dilute populations of bacteria in bulk liquid [25–27, 29–34]. We also introduce growth into this model.

To directly connect the model to many experiments [24, 26, 28, 34], we consider a sole nutrient that also acts as the chemoattractant—as is conventionally done [24, 29, 32]—represented by the continuum variable c(x, t), where x is the position coordinate and t is time. The number density of bacteria, in turn, is given by the continuum variable b(x, t). Furthermore, given the experimental conditions, we assume that the cells do not excrete their own chemoattractant or other diffusible signals, as is sometimes the case in low-nutrient conditions and for specific strains. Recent extensions of this model have also considered the case in which

nutrient and attractant are separate chemical species, which leads to fundamentally different behavior that would be interesting to explore using our framework in future work [27, 47].

As the nutrient diffuses through space with thermal diffusivity D_c it is consumed by the cells at a rate $b\kappa g(c)$; here, κ is the maximum consumption rate per cell and the Monod function $g(c) \equiv c/(c+c_{\rm char})$, with the characteristic concentration $c_{\rm char}$, quantifies the reduction in consumption rate when nutrient is sparse [27, 34, 48–52]. Therefore, the nutrient dynamics are given by

$$\frac{\partial c}{\partial t} = D_c \nabla^2 c - b \kappa g(c). \tag{1}$$

The bacterial dynamics have two contributions: a motility-driven flux \vec{J}_m and cellular proliferation. The flux arises from the combination of the undirected spreading of cells, a diffusive process with an active diffusivity D_{b0} [53], and directed chemotaxis with a drift velocity $\vec{v}_c \equiv \chi_0 \nabla f(c)$ that quantifies the ability of the bacteria to logarithmically respond to the local nutrient gradient [29–31]. The well-established function $f(c) \equiv \log[(1+c/c_-)/(1+c/c_+)]$ quantifies the ability of the cells to sense nutrient with characteristic bounds c_- and c_+ [26, 27, 54–62], while the chemotactic coefficient χ_0 quantifies the ability of the cells to bias their motion in response to a sensed nutrient gradient. Therefore, the motility-driven flux $\vec{J}_m = -D_{b0}\nabla b + b\vec{v}_c$. Proliferation, on the other hand, is given by $b\gamma g(c)$, where γ is the maximal growth rate per cell and g(c) reflects the reduction in growth rate when nutrient is sparse—circumventing the need to introduce an ad hoc "carrying capacity" of a logistically-growing population, as is sometimes done. However, as we detail further in §2.3, the added feature of confinement does introduce a maximum cell density at which purely growth-driven spreading dominates. Here, in the absence of confinement, the bacterial dynamics are therefore given by

$$\frac{\partial b}{\partial t} = \underbrace{D_{b0} \nabla^2 b - \nabla \cdot (b \nu_c)}_{-\nabla \vec{J}_m} + b \gamma g(c). \tag{2}$$

Together, Eqs 1 and 2 represent the classic Keller-Segel model that describes the coupled dynamics of nutrient and bacteria, also including the added influence of cellular growth. In particular, they successfully capture the key features of chemotactic spreading in unconfined liquid, in which cells collectively generate a local gradient of nutrient that they in turn bias their motion along—leading to the formation of a coherent pulse of bacteria that continually propagates, sustained by its continued consumption of the surrounding attractant [23–26].

2.2 Characteristic dimensionless parameters

Non-dimensionalizing Eqs 1 and 2 yields useful dimensionless parameters for characterizing population spreading. We rescale $\{c, b, t, x\}$ by the characteristic quantities $\{c_{\infty}, b_0, t_{c,0}, \zeta\}$, where c_{∞} is the initial nutrient concentration taken to be constant everywhere, b_0 is the maximal initial cell density, $t_{c,0} \equiv c_{\infty}/(b_0\kappa)$ is a characteristic time scale of nutrient consumption, and $\zeta_0 \equiv \sqrt{D_{b0}t_{c,0}}$ is the characteristic extent of cellular diffusion over the duration $t_{c,0}$. This process yields the non-dimensional equations

$$\frac{\partial \tilde{c}}{\partial \tilde{t}} = \delta_0 \tilde{\nabla}^2 \tilde{c} - \tilde{b} \tilde{g} \tag{3}$$

$$\frac{\partial \tilde{b}}{\partial \tilde{t}} = \tilde{\nabla}^2 \tilde{b} - \alpha_0 \tilde{\nabla} \cdot (\tilde{b} \tilde{\nabla} \tilde{f}) + \beta_0 \tilde{b} \tilde{g}, \tag{4}$$

where the tildes indicate non-dimensionalized variables. Three dimensionless parameters emerge:

- The diffusion parameter $\delta_0 \equiv D_c/D_{b0}$ compares the thermal diffusion of nutrient to the active diffusion of bacteria. When $\delta_0 \ll 1$, variations in nutrient are localized to the leading edge of the bacterial population, whereas when $\delta_0 \gg 1$, nutrient levels vary over large spatial extents.
- The directedness parameter $\alpha_0 \equiv \chi_0/D_{b0}$ compares the influence of chemotaxis to active diffusion in driving cellular spreading. When $\alpha_0 \ll 1$, diffusion dominates and cells do not appreciably direct their motion in response to a nutrient gradient, whereas when $\alpha_0 \gg 1$, motile cells strongly direct their motion in response to a gradient.
- The yield parameter $\beta_0 \equiv \gamma/(b_0\kappa/c_\infty)$ compares the rates of cell growth and nutrient consumption, γ and $t_{c,0}^{-1}$, respectively. It therefore quantifies the yield of new cells produced as a population consumes nutrient. When $\beta_0 \ll 1$, nutrient consumption is much faster than proliferation, whereas when $\beta_0 \gg 1$, many new cells are produced in the time required to consume the available nutrient.

The quantity $\Lambda_0 \equiv \alpha_0/(\beta_0 \delta_0) = \gamma^{-1} \cdot \chi_0/(D_c t_{c,0})$ therefore characterizes the interplay between chemotatic and growth-driven spreading of bacterial populations. In particular, $[\chi_0/(D_c t_{c,0})]^{-1}$ is a characteristic time scale needed to spread via chemotaxis over the nutrient diffusion length $\sqrt{D_c t_{c,0}}$, while γ^{-1} is the time scale over which cells grow. Previous studies in bulk liquid focused solely on chemotactic spreading, which is characterized by the limit $\Lambda_0 \gg 1$ [25, 26, 29, 31, 33]. Other studies of non-chemotactic cells focused solely on growth-driven spreading, characterized by the opposite limit $\Lambda_0 = 0$ [41, 42, 63–66]. However, experiments performed in semi-solid agar [34] as well as in defined packings of particles [28] indicate that confinement in such crowded media introduces new cell-cell and cell-medium interactions that are not incorporated in the classic Keller-Segel model. Hence, in this paper, we describe a first step toward incorporating these complexities, which not only tune Λ_0 over a broad range, but also fundamentally alter spreading dynamics—as described hereafter.

2.3 Keller-Segel model incorporating confinement

As a model system, we consider bacterial populations confined in media with close-packed, rigid, and immovable obstacles surrounding a free space that is sufficiently large for cells to move through. This form of confinement alters bacterial spreading dynamics in three ways:

- (i). Collisions with the surroundings impede cellular spreading [35–38], reducing the transport parameters D_{b0} and χ_0 , as quantified in recent experiments in 3D porous media [28, 38] as well as in semi-solid agar [34];
- (ii). The presence of surrounding obstacles reduces the free space available to cells to move through, increasing cellular crowding and promoting cell-cell collisions that further truncate the motility parameters, observed experimentally using *in situ* microscopy [28];
- (iii). When the number density of cells is sufficiently high, this reduction in free space causes the cells to be jammed; hence, they are able to spread only through proliferation, which pushes cells outward, as quantified in experiments using single cell visualization [39, 40].

Notably, (ii)-(iii) are absent from the classic Keller-Segel model, which treats cells as non-contacting, and require modifications beyond simply changing the transport parameters D_{b0} and χ_0 .

- (i). Impeded spreading of isolated cells. Bacterial spreading is typically modeled as a random walk with directed steps of characteristic length ℓ and characteristic duration τ that are punctuated by reorientation events [53]. Consequently, both transport parameters D_{b0} , which describes the unbiased component of the random walk, and χ_0 , which describes the biased component, are set by $\sim \ell^2/\tau$. In bulk liquid, the directed steps are known as runs, which extend along straight-line paths $\ell \sim 40 \, \mu \text{m}$ long, punctuated by rapid tumbles. In tight confinement, however, a cell collides with an obstacle and becomes transiently trapped well before it completes such a run. Therefore, as established in recent experiments [37, 38], runs are truncated by collisions with surrounding obstacles, and the directed steps of the random walk are instead set by the geometry of the available free space; thus, for isolated cells, $\ell \sim \ell_c$, the mean length of straight line *chords* [67] that fit in the free space [37]. Moreover, because the trapping process induced by collisions with obstacles occurs over a duration τ_t that is longer than that of the truncated runs, $au pprox au_{ ext{t}}$. As a result, for cells confined in tight media, both transport parameters D_{b0} and χ_0 are instead $\propto \ell_c^2/\tau_t$ —and because increasing confinement both decreases ℓ_c and increases τ_t [37], it concomitantly decreases both D_{b0} and χ_0 , as confirmed experimentally [28]. Within the context of prior work investigating diffusion in porous media [68], we note that while the volume fraction of free space (porosity) ϕ is known to influence diffusion, it alone does not determine the diffusion coefficient because the geometry of the free space plays a key role as well. Thus, in our model, ϕ influences cellular transport (active diffusion and chemotaxis) indirectly through its effect on the chord length ℓ_{\odot} which characterizes the length scale associated with straight paths that fit within the free space—and therefore determines the length scale over which cells can move in a directed manner.
- (ii). Crowding-induced collisions between cells. Confinement also reduces the free space available to cells. Our definition of the number density of bacteria b quantifies the number of cells per unit total volume of space, which includes the volume of surrounding obstacles; hence, the local density of cells is given by b/ϕ , where $\phi < 1$ is the volume fraction of free space that is reduced by the presence of obstacles. This increase in the local density of cells increases the propensity of neighboring cells to collide as they move, further truncating ℓ . Single-cell imaging in a porous medium confirms this expectation [28]: when the available free space is so tight that multiple cells cannot fit side-by-side, cells are necessarily restricted to end-on collisions between each other as they move, also inducing reorientations akin to those induced by collisions with surrounding obstacles. Therefore, as a first step toward incorporating this behavior into the model described in §2.1, we adopt a mean-field treatment of cell-cell interactions in which cells truncate each other's directed steps in a density dependent manner, inducing transient trapping events again of duration $\tau_{\rm t}$ akin to collisions with obstacles.

In particular, wherever the local density b/ϕ is larger than a threshold value b^*/ϕ such that the mean separation between the surfaces of neighboring cells, $\ell_{\rm cell}$, decreases below the mean chord length $\ell_{\rm c}$, we expect that cell-cell collisions truncate ℓ from $\ell_{\rm c}$ to $\ell_{\rm cell}$ (schematized in the middle inset of Fig 1). Because the diffusion and chemotactic coefficients both vary as $\propto \ell^2$, we therefore multiply both density-independent parameters D_{b0} and χ_0 that characterize isolated cells by the density-dependent correction factor $\mu_{\rm crowd}(b) = (\ell_{\rm cell}/\ell_{\rm c})^2$, where the cell separation is approximated as the mean value $\ell_{\rm cell} \equiv (3\phi/4\pi b)^{1/3} - d$; here, $d \approx 1$ µm is the characteristic size of a cell, and therefore $b^* \equiv 3\phi/(4\pi \ell_{\rm c} + d)^3$]. As b increases further, it eventually reaches the jamming density $b_{\rm jammed} \equiv 3\phi/(4\pi d^3)$ at which cells cannot move at all, and $\ell_{\rm cell} = 0$; in this case, both transport

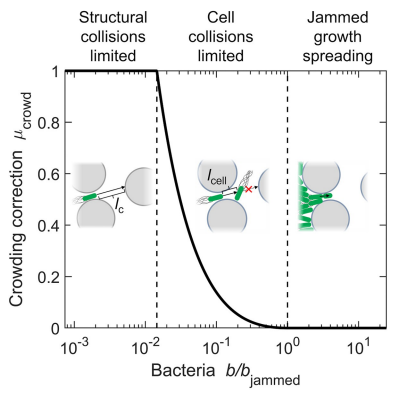


Fig 1. Summary of the cell density-dependent crowding correction μ_{crowd} , which we use in the model to incorporate the influence of confinement on cell-cell collisions. In particular, the cellular transport parameters are multiplied by μ_{crowd} , this case shown for the prototypical case of intermediate confinement. (Left) At low densities, spreading of cells (green) is impeded only by collisions with surrounding solid obstacles (grey), not with neighboring cells, so $\mu_{\text{crowd}} = 1$. This impeded spreading is quantified by the transport parameters D_{b0} and χ_0 , whose values are regulated by the characteristic chord length ℓ_c characterizing the amount of free space between obstacles. (Middle) When the local density of cells is so large that the characteristic separation between neighboring cells ℓ_{cell} is less than the characteristic chord length ℓ_c , cell-cell collisions further truncate the transport parameters. This effect is quantified by $\mu_{\text{crowd}} < 1$. (Right) At the maximal density $b = b_{\text{jammed}}$, the cells are jammed and have no free space to move. Therefore, $\mu_{\text{crowd}} = 0$, and the population spreads solely through growth and division of cells. Note that our definition of the number density of bacteria b is as the number of cells per unit total volume of space, which includes the volume of surrounding obstacles.

parameters are zero, and the bacterial population can only spread via growth. Therefore, in Eq 2, D_{b0} and χ_0 are replaced by the corrected values

$$D_b(b) = D_{b0} \times \mu_{\text{crowd}}(b) \tag{5}$$

$$\chi(b) = \chi_0 \times \mu_{\text{crowd}}(b) \tag{6}$$

where the crowding correction factor $\mu_{\text{crowd}}(b)$ is piecewise defined as

$$\mu_{\text{crowd}}(b) = \begin{cases} 1 & \text{when } b \leq b^* \\ \left[\frac{3\phi}{4\pi b} \right]^{1/3} - d \\ \ell_{\text{c}} & \end{cases} \text{when } b^* < b < b_{\text{jammed}} \end{cases}$$

$$0 & \text{when } b \geq b_{\text{jammed}}$$

$$(7)$$

as shown in Fig 1; the limits b^* and $b_{\rm jammed}$ are indicated by the left and right vertical dashed lines, respectively. This way of correcting the diffusive term in Eq 2 represents a simplifying approximation; strictly speaking, one cannot simply commute the divergence operator with the diffusion coefficient, given that D_b depends on b(x,t) via the crowding correction factor. However, as we describe further in S1 Text, this simplification does not appreciably influence our results and conclusions. We term cases with low cell density ($b < b^*$) the obstacle collisions limited regime described in (i) above; cases with intermediate cell density ($b^* \le b < b_{\rm jammed}$) the cell collisions limited regime; and cases with the highest possible density of cells ($b = b_{\rm jammed}$) the jammed growth spreading regime described in (iii) below. Because we take the cells and surrounding obstacles to be incompressible, b cannot exceed $b_{\rm jammed}$.

(iii). Jammed growth spreading. When they are jammed, cells form a contact network that holds them in place and prevents motion by active propulsion. However, these cells can continue to proliferate if supplied with nutrient; based on the experiments in [28], we assume that the maximal growth rate γ is not affected by confinement. Thus, in this case, their high body stiffness enables growing cells to push outward on their neighbors; the bacterial population can then be treated as an incompressible "fluid" in which the added stress due to cellular growth relaxes rapidly via spreading, as is conventionally done in models of growing immotile populations [41, 69–71] and supported by experiments [39, 40]. Because we treat the obstacles comprising the medium as being rigid and immovable, and the interstitial free space large enough for cells to move through without being deformed, this process leads to jammed growth spreading. We incorporate this behavior into the Keller-Segel model following previous work modeling the growth of immotile biofilms [42]. In particular, at each time step δt , we first identify the smallest x_i at which $b(x_i, t + \delta t)$ exceeds b_{jammed} ; we then set $b(x_i, t + \delta t) =$ b_{jammed} and instead relocate the newly-formed cells $\delta b(x_i) \equiv b(x_i, t + \delta t) - b_{\text{jammed}}$ to the nearest location $x_i > x_i$ at which $b(x_i, t) < b_{jammed}$. We then repeat this process for all successive positions $x > x_i$ such that at time $t + \delta t$, the upper limit on cell density b_{jammed} is globally satisfied.

2.4 Implementation of numerical simulations

To explore the influence of confinement, we perform numerical simulations of Eqs 1 and 2, modified as described in §2.3. Motivated by its simplicity and amenability to the addition of our discrete jammed expansion rule, we implement a forward Euler method to solve these equations; specifically, we discretize the spatial coordinate x using a forward difference form for first derivatives and a central difference form for second derivatives. The update equations for nutrient concentration and bacterial cell density, corresponding to Eqs 1 and 2 respectively, are then:

$$\begin{aligned} c_i^{n+1} &= c_i^n + \delta t \left[\frac{D_c}{\delta x^2} (c_{i-1}^n - 2c_i^n + c_{i+1}^n) - \kappa b_i^n g(c_i^n) \right] \\ b_i^{n+1} &= b_i^n + \delta t \left[\frac{D_{b0} \mu(b_i^n)}{\delta x^2} (b_{i-1}^n - 2b_i^n + b_{i+1}^n) - \frac{1}{\delta x} (b_{i+1}^n v_{c_{i+1}}^n - b_i^n v_{c_i}^n) + \gamma b_i^n g(c_i^n) \right] \end{aligned}$$

where time points advance in discrete steps of δt and are indexed by n, and spatial positions are separated by discrete steps of δx and are indexed by i. The spatial resolution δx is 10 μ m and the time step δt is 0.01 s; as shown in S3 Fig, these choices are sufficiently fine so that our results are not sensitive to the choice of resolution. We note that implicit methods (such as

backwards Euler) or semi-implicit methods would likely improve the efficiency of the numerical simulations—an important consideration for those seeking to extend our work e.g., to higher spatial dimensions.

To connect our results to an experimental system, we use input parameters and initial conditions that mimic the experiments described in [28], which explored the chemotactic spreading of E. coli populations in 3D porous media composed of densely-packed hydrogel particles. We use a Cartesian rectilinear coordinate system extending to a maximum distance of 1.75×10^4 µm, matching the length of the experimental system. Because our system is onedimensional, vectors (e.g. fluxes) oriented in the + or -x directions are represented by positive or negative quantities, respectively, with the vector notation suppressed. Both boundaries have no flux conditions. In these experiments, L-serine was considered to act as the primary nutrient and chemoattractant for cells. Because the hydrogel particles are polymer networks swollen in liquid, they are permeable to the nutrient, similar to many other naturally-occurring media such as biological gels and microporous clays/soils. Therefore, we take the nutrient diffusivity D_c to be equal to its value in bulk liquid, 800 μ m² s⁻¹ [72], and the nutrient is initially saturated at $c_{\infty} = 10$ mM throughout the simulation domain. For all the simulations, we use direct measurements of individual cells [27, 28, 34] to choose fixed values of the cellular parameters c_, c_+ , and γ given by 1 μ M, 30 μ M, and 0.69 h⁻¹, respectively; furthermore, as detailed in S2 Text, we use the data from experiments on spreading populations [28] to directly determine c_{char} and κ , given by 10 μ M and 1.3 × 10⁻¹² mM (cells/mL)⁻¹ s⁻¹, respectively.

Each experiment used a long 3D-printed cylinder of close-packed cells not containing hydrogel particles ($\phi=1$) as the initial inoculum, embedded within and surrounded by the 3D porous medium. The cells then continued to spread radially outward through the pore space. Thus, as the initial condition in all the simulations, we consider a Gaussian profile of b(x,t=0) centered at x=0 with a full width at half maximum of 100 μ m and a peak number density of $b_0=b_{\rm max}\equiv 3/(4\pi d^3)=2.4\times 10^{11}$ cells/mL, where $b_{\rm max}$ is defined as the number density of close-packed cells and is therefore the maximum possible value of b_0 —with the exception of the lower-density simulations that employ a lower value of b_0 , as detailed further in §3.1.3. Hence, for all simulations except those in §3.1.3, the initial inoculum has a maximal cell density $b_{\rm max}>b_{\rm jammed}$, where $b_{\rm jammed}$ instead corresponds to the maximal possible density of cells in confinement ($\phi<1$). For simplicity, wherever $b(x,t=0)>b_{\rm jammed}$, we still apply the jammed growth spreading rule described in §2.3(iii), but with $b_{\rm jammed}$ replaced by b(x,t=0).

The experiments tuned cellular confinement by using porous media with varying porosities ϕ and mean chord lengths ℓ_c [37], resulting in varying values of the transport parameters D_{b0} and χ_0 [28, 38]. In particular, as determined from the experiments, D_{b0} and χ_0 both decrease with increasing confinement as cellular mobility is increasingly hindered. Hence, in our simulations, we tune confinement by varying these parameters, using the values of D_{b0} obtained from single-cell imaging [28] and extracting χ_0 from experimental measurements of population spreading, as detailed in S2 Text. The confinement-dependent parameters are summarized in Table 1.

The corresponding dimensionless parameters characterizing the Keller-Segel model (§2.2) are also summarized in Table 1:

- The diffusion parameter $\delta_0 \equiv D_c/D_{b0}$ increases with confinement as cellular mobility is increasingly hindered. For all conditions tested here, however, δ_0 is always much greater than one, reflecting fast diffusion of nutrient; thus, we expect that nutrient levels vary over large spatial extents, as confirmed in the simulations that follow.
- For all conditions tested here, the directedness parameter $\alpha_0 \equiv \chi_0/D_{b0}$ is always much greater than one, indicating that motile cells strongly direct their motion in response to the nutrient

Table 1. Parameters used to describe bacteria in weak, intermediate, and strong confinement, as defined in the text. All parameters are defined in the text and their values are obtained from experiments as detailed in §2.4 and \$2 Text, with the exception of \bar{b} , which is determined directly from the simulation. We note that the values of α_0 are taken directly from the experiments in [28], which indicate that this parameter surprisingly decreases with increasing confinement. Thus, while we expect that both transport parameters χ_0 and D_{b0} are tuned by confinement in a similar way, with both proportional to \bar{l}_c^2/τ_t , it appears from the experiments that the ratio of the proportionality constants for each is also confinement-dependent. That is, experiments suggest that confinement more strongly hinders directed (quantified by χ_0) versus undirected (quantified by χ_0) spreading. While more work needs to be done to fully unravel why this is the case, in absence of a theoretical model for the confinement-dependence of α_0 , we directly use the experimental values in our work. Furthermore, in the absence of any experimental data assessing the influence of cell density on α_0 , we make the simplest possible assumption that cell-cell collisions hinder both χ_0 and χ_0 and χ_0 through the same crowding correction factor χ_0 , which quantifies the reduction in free space available to the cells. Thus, we take $\alpha=\alpha_0$. Future experiments could further probe this density dependence and motivate the introduction of additional extensions to our model.

Parameter	Weak confinement	Intermediate confinement	Strong confinement
φ	0.36	0.17	0.04
\bar{l}_{c} (μm)	4.6	3.1	2.4
$D_{b0} (\mu \text{m}^2 \text{s}^{-1)}$	2.3	0.93	0.42
$\chi_0 (\mu \text{m}^2 \text{s}^{-1})$	3700	94	16
$ar{b}/b_{ ext{max}}$	0.10	0.027	0.026
$\delta_0 \equiv D_c/D_{b0}$	340	860	1900
$\delta \equiv D_c/D_b(ar{b})$	2.8×10^4	1.2×10^4	4.3×10^{5}
$\alpha = \alpha_0 \equiv \chi_0/D_{b0}$	1600	100	38
$eta_0 \equiv \gamma/(b_0 \kappa/c_\infty)$	0.0063	$0.0063 (b_0 = b_{\text{max}}), 630 (b_0 = 10^{-5} b_{\text{max}})$	0.0063
$eta \equiv \gamma/(ar{b}\kappa/c_{\infty})$	0.06	0.23	0.25
$\Lambda_0 \equiv lpha_0/(eta_0 \delta_0)$	750	18 ($b_0 = b_{\text{max}}$), 1.8×10^{-4} ($b_0 = 10^{-5} b_{\text{max}}$)	1.3
$\Lambda \equiv lpha/(eta\delta)$	0.95	0.037	3.6×10^{-4}

gradient established through consumption. Intriguingly, the α_0 determined from the experimental parameters decreases with increasing confinement, indicating that confinement more strongly hinders directed versus undirected motion—consistent with previous reports that confinement fundamentally alters the mechanism by which cells perform chemotaxis [28]. Further investigating the determinants of α_0 in confinement will be a useful direction for future experiments.

• Because the maximal growth rate is not affected by confinement [28], the yield parameter $\beta_0 \equiv \gamma/(b_0\kappa/c_\infty)$ is independent of confinement for all of our simulations. For all simulations employing $b_0 = b_{\rm max}$, β_0 is much less than one, reflecting the fact that nutrient consumption by a maximally dense population is faster than cellular proliferation; conversely, for the lower-density simulations presented in Figs 4 and 5, β_0 is much greater than one, indicating the dominant role of proliferation in this case.

Therefore, for our simulations testing the influence of confinement on bacterial spreading, the parameter $\Lambda_0 \equiv \alpha_0/(\beta_0\delta_0)$ varies over a broad range, decreasing over nearly three orders of magnitude as confinement increases. We note that because the different parameters δ_0 , α_0 , β_0 do not incorporate the influence of density-dependent cellular crowding, we do not expect this transition to occur precisely at $\Lambda_0 \approx 1$. We therefore define a new version of this parameter, $\Lambda \equiv \alpha/(\beta\delta)$, where now $\delta \equiv D_c/D_b(\bar{b})$, $\alpha \equiv \chi(\bar{b})/D_b(\bar{b}) = \alpha_0$, and $\beta \equiv \gamma/(\bar{b}\kappa/c_\infty)$ (Table 1); \bar{b} is defined as the long-time mean cell density within each propagating pulse, and is directly calculated from each simulation as described further below. Thus, the newly-defined Λ explicitly incorporates density-dependent crowding. As summarized in Table 1, our simulations explore the transition from weak confinement ($\Lambda=0.95$) to strong confinement ($\Lambda=3.6\times 10^{-4}$); consistent with our expectation, this range reflects a transition from chemotactic to growth-driven spreading, as demonstrated directly by the simulations presented below.

3 Results

3.1 Intermediate confinement

As a prototypical starting case, we first examine bacterial spreading from a dense-packed Gaussian-shaped inoculum under intermediate confinement ($\Lambda=0.037$), shown by the initial profile for t=0 in Fig 2A. The simulation incorporates both motility and growth. The cells rapidly deplete nutrient locally via consumption (S1 Video) over a time scale $\sim c_{\infty}/(\kappa b_{\rm max}) \approx 30$ s, establishing a steep nutrient gradient at the leading edge of the population. This gradient extends over a large distance ahead of the population (Fig 2B and inset)—as expected from our calculation of the diffusion parameter $\delta_0\gg 1$. Cells at this leading edge then continue to grow outward as a jammed front with $b=b_{\rm jammed}$, shown by the flat region at t=1.8 h in Fig 2A. Eventually, a lower-density, coherent pulse of cells detaches from this jammed region (t=3.7 h), continues to propagate the nutrient gradient along with it, and thus continues to spread outward (t>3.7 h), as shown by the outward-moving peak in Fig 2A.

Indeed, this pulse spans the extent over which nutrient varies between the upper and lower bounds of sensing, c_+ and c_- (pluses and diamonds shown for the t = 11 h profiles, respectively)—reflecting the central role of chemotaxis in driving its propagation. The forward face of the pulse is also exposed to sufficient nutrient for cells to proliferate (with $c \ge c_{\text{char}}$, the characteristic Monod concentration, shown by the triangles on the t = 11 h profiles)—suggesting that cellular growth contributes to population spreading over long time scales, as well. The overall width of this pulse, $W \approx 200 \,\mu\text{m}$, is set by the length scale over which nutrient is depleted by consumption; at its rear, the nutrient concentration and nutrient gradient are both low, causing both growth and chemotaxis to be hindered. As a result, cells are shed at a nearconstant density $b_{\text{trailing}} \approx 0.02b/b_{\text{max}}$ (see 0.5 mm < x < 1.2 mm in Fig 2A). This coherent pulse of cells continues to move apparently without an appreciable change in shape, as suggested by the inset to Fig 2A, at a speed $v_{\text{pulse}} \approx 0.15$ mm/h. However, given the limited duration of the simulation, our results do not enable us to definitively conclude that the simulated pulse develops into a traveling wave with an unchanging shape; building on our simulation to explore longer length and time scales to test this possibility will be a useful direction for future work. The nutrient profile concomitantly propagates with the pulse, as shown in Fig 2B. Notably, similar spreading behavior was observed in experiments [28]; as shown in S4 Fig, in both simulation and experiment, we observe similarly-shaped bacterial pulses with comparable widths, trailing densities (compared to the peak densities), and final positions at $t \approx 11$ h.

3.1.1 Initial dynamics. To further characterize these spreading dynamics, we track the position x_l of the leading edge of the population over time t, as shown in Fig 3A. Specifically, motivated by a similar definition used in prior experiments [28], we define x_l as the position at which b falls below the threshold value $b/b_{\rm max} = 10^{-4}$. Initially, population spreading is hindered ($x_l \sim t^{\nu}$ with $\nu \ll 1$ e.g., red point), but as the coherent pulse forms and propagates, it eventually approaches constant speed spreading ($\nu \approx 1$ e.g., blue point). A similar transition from hindered to constant speed spreading was observed in experiments [28], although the underlying reason has thus far remained unclear. Here, we use our model to clarify the origin of this transition.

In particular, we examine the two different contributions to the motility-driven flux of cells —active diffusion and chemotaxis—for the population at early and late times (Fig 3B and 3C, respectively); for simplicity, we do not consider the added influence of growth, which only plays an appreciable role for long times $t \gg \gamma^{-1}$, until the next subsection. The magnitude of the active diffusive flux $-D_b(b)\nabla b = -D_{b,0}\mu_{\rm crowd}(b)\nabla b$ as it varies across the population is shown by the dashed lines in the bottom panels of Fig 3B and 3C, while the magnitude of the chemotactic flux $bv_c = b\chi_0\mu_{\rm crowd}(b)\nabla f(c)$ is shown by the dash-dotted lines instead. At early

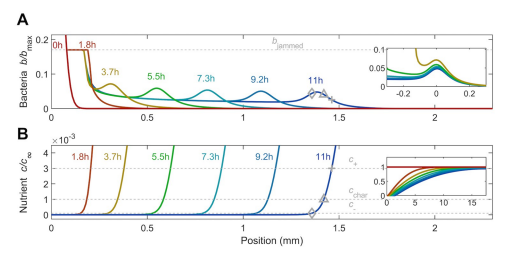


Fig 2. Results from a numerical simulation of population spreading in intermediate confinement. The simulation incorporates both motility and growth. (A) shows the dynamics of the cells while (B) shows the corresponding dynamics of the nutrient, quantified by the normalized density $b/b_{\rm max}$ and concentration c/c_{∞} , respectively. As noted in §2.4, the initial inoculum is composed purely of dense-packed cells with liquid between them ($\phi = 1$), with the entire inoculum surrounded by the obstacle-filled medium ($\phi = 0.17$); hence, the initial inoculum has $b = b_{\text{max}}$, which is larger than b_{jammed} , the jamming density of cells in confinement. Different colors indicate different times as listed. The dense inoculum initially centered about the origin spreads outward, first as a jammed front (jamming density shown by the dashed grey line in A), then detaching as a coherent lower-density pulse that propagates continually via chemotaxis. At long times, this pulse appears to approach an unchanging shape and speed, as suggested by the collapse of the profiles in the upper inset (showing the same data, but shifted horizontally to center the peaks). The cellular dynamics arise in response to consumption of the nutrient, which is initially saturated everywhere, but is rapidly depleted and forms a gradient that is propagated with the pulse (inset shows the same data but with both axes zoomed out). In B, the three dashed grey lines show the characteristic concentrations of sensing c_+ and c_- and the characteristic Monod concentration c_{char} ; the corresponding positions are shown by the pluses, diamonds, and triangles, respectively, in A-B. An animated form of this Figure is shown in S1 Video, and a sample MATLAB code to implement this simulation is provided in \$1 File.

times, the gradient in cell density is steep, as set by the sharp initial profile of cells and the limited extent of subsequent population spreading (Fig 3B, top). As a result, spreading is primarily due to active diffusion, which dominates over chemotaxis, as shown in the lower panel of Fig 3B. By contrast, as cells spread outward, the gradient in cell density becomes less steep. As a result, at late times, spreading is primarily due to chemotaxis, which dominates over active diffusion, as shown in the lower panel of Fig 3C. This behavior is also reflected by the bottom set of circles and squares in Fig 3D, which represent the maximal diffusive and chemotactic fluxes across the population (exemplified by the circles and squares in Fig 3B and 3C) over time. Initially, the diffusive flux dominates over the chemotactic flux; however, as the population continues to spread and consume nutrient, the diffusive flux decreases and the chemotactic flux increases, with both eventually approaching constant values at long times.

Another key factor that hinders the initial population spreading is cellular crowding. To assess the influence of crowding, we compare the maximal diffusive and chemotactic fluxes across the population, but with or without the crowding correction factor $\mu_{\text{crowd}}(b)$ (corresponding to the "with crowding" and "no crowding" datasets, respectively, in Fig 3D). In both cases, the active diffusive flux dominates over the chemotactic flux initially, but chemotaxis eventually dominates as the population continues to spread and establish the nutrient gradient (e.g. top set of squares for $t \leq 0.1$ h). The spreading of the population remains hindered, however; due to the high initial density of cells, crowding continues to limit the chemotactic flux of

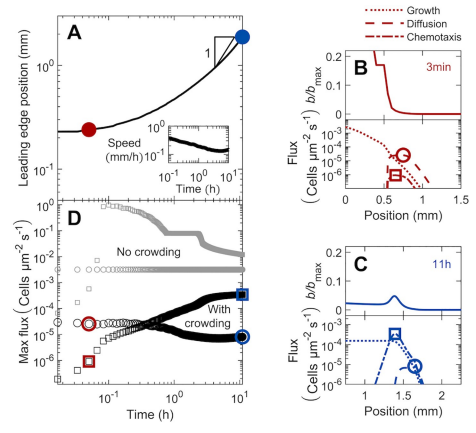


Fig 3. Population dynamics, morphology, and fluxes driving spreading for the simulation of bacteria in intermediate confinement (Fig 2). The simulation incorporates both motility and growth. (A) Increase in the position of the leading edge of the population is initially hindered (red), but approaches constant-speed motion indicated by the triangle at long times (blue). The corresponding instantaneous speed $\nu_{\rm pulse}$ is shown in the inset. (B) At a short time corresponding to the red point in A, the population expands as a jammed front (top panel). Lower panel shows that cellular growth and diffusion are the primary contributors to the expansion of this front. (C) At a long time corresponding to the blue point in A, the population spreads as a coherent pulse (top panel). Lower panel shows that chemotaxis is the primary contributor to pulse propagation. Positions of the maximal diffusive and chemotactic fluxes are indicated by the circles and squares, respectively, in B-C; note the slight upward kink in the diffusive flux in C indicated by the circle. (D) Variation of the maximal diffusive and chemotactic fluxes, indicated by the circles and squares in B-C, over time. The initial population dynamics are dominated by cellular diffusion (circles), while at longer times chemotaxis dominates (squares). To illustrate the role played by cellular collisions, we show the same data with and without the crowding correction $\mu_{\rm crowd}$ in the upper (grey) and lower (black) datasets; crowding hinders population spreading, as shown by the vertical offset in the curves, but plays a less appreciable role at long times, as shown by the curves approaching each other.

cells, only enabling a small fraction at the leading edge of the population to spread outward—as exemplified by the first two profiles in Fig 2A, the sharp decrease in both diffusive and chemotactic fluxes for x < 0.5 mm in Fig 3B, and the large difference between the two sets of squares in Fig 3D. Eventually, as this leading edge continues to spread, crowding in the forward face of the population becomes sufficiently low, enabling the coherent pulse of cells to detach from the population—as exemplified by the t = 3.7 h profile in Fig 2A and the "kink" in the top set of squares at $t \approx 3$ h in Fig 3D. Hindrance due to crowding continues to decrease over time, as shown by the diminishing difference between the two sets of squares in Fig 3D for t > 3 h, and eventually approaches a constant value.

Hence, population spreading is initially slow due to the time required for cellular consumption to establish a sufficiently strong nutrient gradient to drive chemotactic spreading. Cellular crowding near the initial inoculum then continues to hinder spreading until enough of the forward face of the population has spread outward—enabling cells to detach as a coherent pulse that continues to move outward, eventually approaching a constant speed.

3.1.2 Long-time behavior. Having established how the spreading population forms a moving pulse, we now seek to clarify the factors that continue to drive its propagation. As previously described (Fig 3), active diffusion plays a negligible role at these longer times. Instead, as noted previously when describing the t=11 h profiles in Fig 2, we expect that chemotaxis and growth are the principal contributors to population spreading. In particular, the outward-moving pulse spans the extent over which nutrient varies between the upper and lower bounds of nutrient sensing—reflecting the central role of chemotaxis in driving its propagation. The forward face of the pulse is also exposed to sufficient nutrient for cells to proliferate—suggesting that cellular growth contributes to spreading, as well. Indeed, the time scale over which this pulse propagates over its width $\sim W/v_{\rm pulse}=1.3$ h is comparable to the time scale of cellular proliferation, $\gamma^{-1}=1.4$ h, further indicating that growth may contribute to population spreading. However, the relative influence of chemotaxis versus growth in driving population spreading remains unclear.

Hence, we examine the long-time behavior of the pulse by considering a coordinate system that moves with the pulse, $\xi \equiv x - t v_{\rm pulse} + \xi_0$; ξ_0 is a constant shift factor chosen such that $\xi = 0$ is located at the rear of the pulse, at which $b \approx b_{\rm trailing}$. Here, both the bacterial and nutrient gradients are negligible, eliminating diffusive and chemotactic fluxes of cells, as shown in Fig 3C. Within a time increment dt, the moving pulse leaves behind $N_{\rm loss} \approx b_{\rm trailing} v_{\rm pulse} A dt$ cells, where A is the transverse cross sectional area. Simultaneously, growth generates $N_{\rm grown} \approx A dt \int_{\xi=0}^{\infty} b(\xi') \gamma g(\xi') d\xi'$ new cells within the pulse. Therefore, $N_{\rm loss} \approx N_{\rm grown}$ to preserve what we assume for simplicity to be a nearly-unchanging pulse of cells (Fig 2A, inset).

More generally, at locations further ahead ($\xi \ge 0$), $N_{\text{motile}} \approx J_m A dt$ cells also travel with the pulse through their motility-driven flux $J_m = -D_b \nabla b + b v_c$; here, $b, \nabla b, D_b$, and v_c are all ξ -dependent quantities. Thus, an unchanging profile of cells requires the more general flux balance $N_{\text{loss}} - N_{\text{motile}} \approx N_{\text{grown}}$, where now $N_{\text{loss}} \approx b v_{\text{pulse}} A dt$ and $N_{\text{grown}} \approx A dt \int_{\xi}^{\infty} b(\xi') \gamma g(\xi') d\xi'$; that is,

$$\underbrace{bv_{\text{pulse}}}_{\text{Loss}} + \underbrace{D_{\text{b}}\nabla b}_{\text{Diffusion}} - \underbrace{bv_{\text{c}}}_{\text{Chemotaxis}} \approx \underbrace{\int_{\xi}^{\infty} b\gamma g d\xi'}_{\text{Growth}}$$
(8)

where all quantities except for the constants v_{pulse} and γ are position-dependent. This equation quantifies the intuition that the cells that cannot keep up with the moving pulse through their motility must be replaced by growth so as to prevent a net loss of cells from the region ahead of ξ . Therefore, for a given position ξ , the right hand side of Eq 8 represents the additional contribution to the overall spreading of the pulse due to cellular growth at $\xi \geq 0$. We therefore term this quantity the *growth flux* and compare it to the chemotactic flux bv_c .

Both fluxes are shown for the final profile in Fig 3C; the growth flux is shown by the dotted line and the chemotactic flux is shown by the dash-dotted line, both plotted on a logarithmic scale. A version showing these fluxes on linear scales is given in §3.2. For this case of intermediate confinement, both fluxes are appreciable, with the maximal chemotactic flux $(3.4 \times 10^{-4} \text{ cells } \mu\text{m}^{-2}\text{s}^{-1})$ slightly larger than the maximal growth flux $(1.5 \times 10^{-4} \text{ cells } \mu\text{m}^{-2}\text{s}^{-1})$, indicating that chemotaxis plays a greater role in driving population spreading. To further quantify this behavior, we evaluate Eq 8 at two distinct positions: the rear of the pulse $(\xi=0)$ and the

peak of chemotactic flux, which we denote ξ_{peak} (indicated by the square in Fig 3C). At both locations, the gradient in cell density is approximately zero, eliminating diffusive flux and simplifying our analysis. The chemotactic flux is also approximately zero at the rear of the pulse ($x \approx 1.2$ mm in Fig 3C). Moreover, at both locations, the growth flux is approximately the same—reflecting the fact that only the forward face of the pulse is exposed to sufficient nutrient for cells to proliferate. Hence, equating both of these implementations of Eq 8 yields an expression for the long-time pulse speed:

$$v_{\rm pulse} \approx v_c(\xi_{\rm peak}) + v_{\rm pulse} \frac{b_{\rm trailing}}{b_{\rm peak}},$$
 (9)

where we have defined $b_{\rm peak} \equiv b(\xi_{\rm peak})$. Therefore, the ratio $b_{\rm trailing}/b_{\rm peak} = 40\%$ approximates the fraction of the overall pulse speed attributable to growth, while the remaining 60% is due to chemotaxis.

This analysis also provides a way to extend a previous scaling estimate [27] of the long-time pulse speed $\nu_{\rm pulse}$, which did not incorporate the influence of confinement in regulating spreading. First, we note that the chemotactic velocity scales as $\nu_c(\xi_{\rm peak}) \sim \chi(b_{\rm peak})/W$, where W is the pulse width. Next, we relate the mean number density of cells $\bar{b} \equiv W^{-1} \int_0^\infty b d\xi'$ to $\nu_{\rm pulse}$ through a flux balance of cells at long times, when the shape of the pulse is unchanging over time. In particular, as described earlier, the rate at which cells are left behind the pulse, $b_{\rm trailing}\nu_{\rm pulse}A$, is balanced by the rate at which growth generates new cells in the pulse, $A\int_0^\infty b\gamma g d\xi' = A\bar{b}W\gamma\bar{g}$, where we have defined the cell-weighted mean $\bar{g} \equiv \int_0^\infty g(c(\xi'))bd\xi'/\int_0^\infty bd\xi' = \int_0^\infty g(c(\xi'))bd\xi'/(\bar{b}W)$. This flux balance yields $W = b_{\rm trailing}\nu_{\rm pulse}/(\bar{b}\gamma\bar{g})$, and therefore, $\nu_c(\xi_{\rm peak}) \sim \chi(b_{\rm peak})\bar{b}\gamma\bar{g}/(b_{\rm trailing}\nu_{\rm pulse})$. Substituting this expression into Eq. 9,

$$u_{\rm pulse} pprox \chi \Big(b_{\rm peak}\Big) rac{ar{b} \gamma ar{g}(c)}{b_{
m trailing}
u_{
m pulse}} +
u_{
m pulse} rac{b_{
m trailing}}{b_{
m peak}}.$$
(10)

Multiplying both left and right hand sides by v_{pulse} , grouping terms to solve for v_{pulse}^2 , and multiplying the resulting solution by $\frac{b_{\text{peak}}b_{\text{trailing}}}{b_{\text{peak}}b_{\text{trailing}}}$ then yields our ultimate scaling estimate:

$$v_{\rm pulse}^2 \approx \chi \Big(b_{\rm peak}\Big) \gamma \bar{g}(c) \frac{\bar{b}b_{\rm peak}}{b_{\rm trailing}(b_{\rm peak} - b_{\rm trailing})}.$$
 (11)

This estimate thus extends a previous calculation [27] by explicitly incorporating the influence of confinement. To evaluate the accuracy of this estimate, we use the long-time simulation data to directly determine all the parameters on the right hand side of Eq 11 and thereby obtain $\nu_{\rm pulse}$. We find reasonable agreement between the predicted (via Eq 11) and simulated speeds to within a factor of two: the predicted value is 0.08 mm/h, while the simulation yields 0.15 mm/h. This agreement also extends to the case of weak confinement, discussed further in §3.2, for which the predicted value is 0.3 mm/h, while the simulation yields 0.8 mm/h, within a factor of 2.5. Hence, Eq 11 provides a straightforward way to approximately relate the long-time shape of a pulse to its propagation speed, even in confinement.

Finally, we note that the fluxes associated with chemotaxis and growth also determine the overall shape of the spreading population; for simplicity, we neglect the diffusive flux, given that it is at least one order of magnitude smaller than the chemotactic and growth fluxes (Fig 3C). In particular, as quantified in Eq.8, the cellular profile $b(\xi)$ is given by the sum of the chemotactic and growth fluxes, scaled by the constant v_{pulse} . Our results confirm this expectation:

as shown in Fig 3C, the location of the bacterial pulse nearly coincides with the peak in the chemotactic flux, while the steady increase in growth flux from the leading edge to the rear coincides with the additional asymmetry in the bacterial profile arising from the trail of cells shed from the moving pulse. Taken together, these results therefore demonstrate that the interplay between chemotaxis and growth determines both the long-time speed and shape of the spreading population.

3.1.3 Influence of initial cell density. Our analysis thus far considered a dense initial inoculum, for which cellular crowding hinders the formation and detachment of a pulse; at much longer times, this less-crowded pulse no longer resembles the initial inoculum, but instead is shaped by the interplay of chemotaxis and growth (Fig 3). We therefore expect that for a lower-density inoculum, a similar pulse also emerges at long times, but with initial dynamics that are limited instead by the time required for cellular consumption to establish a sufficiently strong nutrient gradient. To test this expectation, we repeat the simulation shown in Fig 2, but using an initial peak number density of cells that is 10^5 times smaller ($b_0 = 10^{-5}b_{\rm max}$)—shown in S2 Video.

In the previously-considered case of a dense inoculum, the cells deplete nutrient rapidly via consumption, and the population subsequently spreads from its leading edge as a growing jammed front (first two curves in Fig 2). By contrast, with a more dilute inoculum, nutrient depletion takes much longer. Instead, the population continually grows and spreads as a whole (red to blue curves in Fig 4A inset), not just at its leading edge, without appreciably depleting nutrient. It eventually reaches a maximal density $b' = b_0 e^{\gamma t'}$ for which the time scale of subsequent nutrient depletion $t_{\rm dep} \sim c_{\infty} / (\kappa b')$ is comparable to the time scale of subsequent growth $t_{\rm g} \sim \gamma^{-1}$; equating these time scales yields $t' = \gamma^{-1} \ln \left(\frac{c_{\infty} \gamma}{b_0 \kappa} \right)$. Therefore, we expect that nutrient is fully depleted at the initial inoculum after $t' + t_{\rm dep} \sim \gamma^{-1} \left[\ln \left(\frac{c_{\infty} \gamma}{b_0 \kappa} \right) + 1 \right] \approx 11$ h. The simulation results are consistent with this estimate, which neglects spatial variation in nutrient availability through the entire population and thus serves as a lower bound, showing that nutrient is fully depleted at the initial inoculum after ~ 14 h (red to blue curves in Fig 4B inset). The nutrient gradient again extends over a large distance ahead of the population, as expected from our calculation of the diffusion parameter $\delta_0 \gg 1$.

Unlike the case of a dense inoculum, the population does not subsequently spread as a jammed front. Instead, once the nutrient gradient is sufficiently strong, a coherent pulse of cells again detaches *without* the prior formation of a jammed front, continues to propagate the nutrient gradient with it, and continues to spread outward (t > 15 h in Fig 4). Consistent with our expectation, this pulse is noticeably similar to that which arises in the dense inoculum case: it has a nearly-identical shape and also appears to move without an appreciable change in shape, eventually reaching approximately a similar constant speed $v_{\rm pulse} \approx 0.1$ mm/h (compare late-time profiles in Figs 2 and 4). Evaluating the speed by instead tracking the position of the peak, instead of the leading edge, also yields a comparable value of $v_{\rm pulse} \approx 0.16$ mm/h.

To further characterize the population spreading dynamics, we again plot the leading edge position x_l as a function of time t. As in the case of a dense inoculum, $x_l \sim t^{\nu}$ with $\nu \ll 1$ at early times, transitioning to $\nu \approx 1$ at later times (Fig 5A); however, these seemingly similar dynamics reflect fundamentally different underlying processes at early times. With a more dilute inoculum, slower nutrient depletion causes the diffusive flux to initially dominate over chemotaxis (Fig 5B) without any influence of cellular crowding—indicated by the overlap of the early-time points with/without the crowding correction in Fig 5D. As cells continue to grow and consume nutrient, they eventually establish a sufficiently strong gradient and spread as a coherent pulse via chemotaxis—as indicated by the dominant role of the chemotactic flux

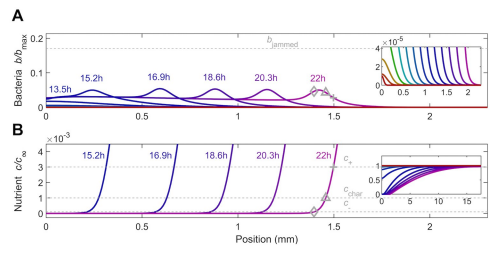


Fig 4. Results from a numerical simulation of population spreading in intermediate confinement starting from a more dilute inoculum. The simulation incorporates both motility and growth. (A) shows the dynamics of the cells while (B) shows the corresponding dynamics of the nutrient, quantified by the normalized density $b/b_{\rm max}$ and concentration c/c_{∞} , respectively. Different colors indicate different times as listed. The dilute inoculum (jamming density shown by the dashed grey line in A) initially centered about the origin first grows exponentially and spreads diffusively until nutrient is locally depleted (upper inset shows the same data, but zoomed in to the vertical axis); only then does a coherent pulse detach and propagate continually via chemotaxis in response to the nutrient gradient (lower inset shows the same data but with both axes zoomed out). Even though the short-time behavior is different from the case of a more dense inoculum shown in Fig 2, the long-time behavior of this pulse is identical. To facilitate comparison with Fig 2, in B, the three dashed grey lines again show the characteristic concentrations of sensing c_+ and c_- and the characteristic Monod concentration $c_{\rm char}$; the corresponding positions are shown by the pluses, diamonds, and triangles, respectively, in A-B. An animated form of this Figure is shown in S2 Video.

at long times (Fig 5C and 5D). At these later times, the different contributions to the bacterial flux are nearly identical to those that drive pulse propagation in the case of a dense inoculum (compare Figs 5C and 3C). Indeed, the fractions of the overall pulse speed attributable to chemotaxis and growth, as quantified by Eq 9, are \approx 58% and 42%, respectively—nearly identical to the case of a dense inoculum. Hence, while the initial dynamics of population spreading are sensitive to the initial cell density—consistent with experiments [28]—the properties of the pulse that forms and continues to drive spreading at long times are not, instead being set solely by the interplay between chemotaxis and growth.

3.2 Influence of confinement

For the case of intermediate confinement explored thus far, we have established that chemotaxis and growth both drive population spreading at long times. How does this behavior change with confinement? As quantified in Figs 3D and 5, confinement-induced crowding limits the chemotactic flux; therefore, we expect that with reduced or increased confinement, chemotaxis or growth plays a more dominant role in driving spreading, respectively. To test this expectation, we perform the same simulation with a dense inoculum as in Figs 2 and 3, but with different values of the confinement-dependent parameters as summarized in Table 1. In particular, our simulations explore $\Lambda = 0.95$, 0.037, and 3.6×10^{-4} , representing weak, intermediate, and strong confinement (top, middle, and bottom rows in Figs 6 and 7), respectively—also shown in S3 Video.

In all cases, the cells first rapidly deplete nutrient locally via consumption, generating a nutrient gradient that again extends over a large distance and drives subsequent spreading at

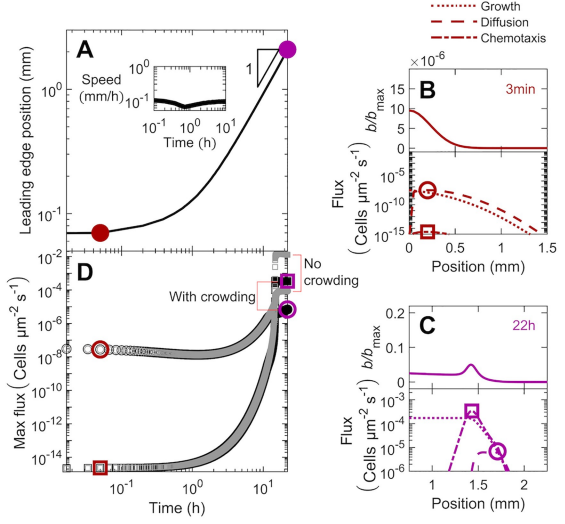


Fig 5. Population dynamics, morphology, and fluxes driving spreading for the simulation of bacteria in intermediate confinement, but from a more dilute initial inoculum (Fig 4). The simulation incorporates both motility and growth. (A) Increase in the position of the leading edge of the population is initially hindered (red), but approaches constant-speed motion indicated by the triangle at long times (purple). The corresponding speed $v_{\rm pulse}$ is shown in the inset. (B) At a short time corresponding to the red point in A, the population grows exponentially (top panel) and spreads primarily through growth and diffusion (lower panel). (C) At a long time corresponding to the purple point in A, the population spreads as a coherent pulse (top panel). Lower panel shows that chemotaxis is the primary contributor to pulse propagation. Positions of the maximal diffusive and chemotactic fluxes are indicated by the circles and squares, respectively, in B-C; note the slight upward kink in the diffusive flux in C indicated by the circle. (D) Variation of the maximal diffusive and chemotactic fluxes, indicated by the circles and squares in B-C, over time. The initial population dynamics are dominated by cellular diffusion (circles), while at longer times chemotaxis dominates (squares). To illustrate the role played by cellular collisions, we show the same data with (black) and without (grey) the crowding correction $\mu_{\rm crowd}$ in the datasets indicated by the red lines; the data are identical except at long times, when crowding slightly hinders population spreading.

the leading edge of the population. However, consistent with our expectation, and with experimental observations [28], the nature of this spreading is strongly confinement-dependent.

3.2.1 Weak confinement. In the case of weak confinement, cells detach and spread as a lower-density, coherent, propagating pulse *without* first growing outward as a jammed front (Fig 6A), unlike the case of intermediate confinement (Fig 6B). This pulse is notably sharper and faster, with the long-time pulse speed and peak height ≈ 5.4 and 3.9 times larger than in

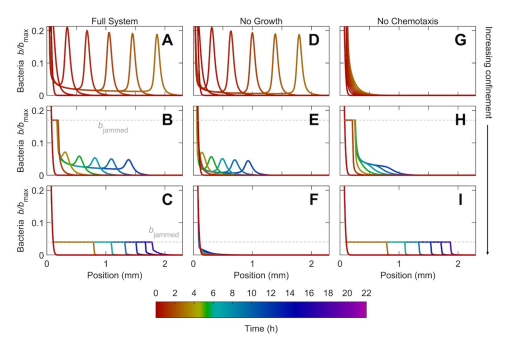


Fig 6. Increasing confinement causes a transition from fast chemotactic pulse propagation to slower jammed growth expansion. Panels show results from numerical simulations of population spreading from the same dense inoculum initially centered about the origin in weak, intermediate, and strong confinement, shown by top, middle, and bottom rows respectively. First column shows the results of the full model, while second and third columns show the same simulations with growth or chemotaxis omitted, respectively. We only show the normalized cellular density $b/b_{\rm max}$ for clarity. As noted in §2.4, the initial inoculum is composed purely of dense-packed cells with liquid in between them ($\phi = 1$), surrounded by the obstacle-filled medium ($\phi < 1$); hence, the initial inoculum has $b = b_{\rm max}$, which is larger than $b_{\rm jammed}$, the jamming density of cells in confinement. Different colors indicate different times as listed in the color scale. In weak confinement, a coherent pulse rapidly detaches and continually propagates; this pulse is driven primarily by chemotaxis, and thus, omitting growth barely changes the dynamics while omitting chemotaxis abolishes the propagation altogether. Conversely, in strong confinement, the population spreads slowly as a jammed front, driven primarily by growth. In intermediate confinement, both growth and chemotaxis drive population spreading. The dashed grey line shows the jamming density, which varies depending on confinement (and is larger than the vertical scale in the top row). An animated form of this Figure, along with the nutrient profiles, is shown in S3 Video.

intermediate confinement (also compare Panels A–B and D–E in Fig 7)—reflecting the dominant role of chemotaxis in driving spreading, as expected from the larger value of Λ . Quantification of the different fluxes driving spreading corroborates this expectation (Fig 7D); indeed, following our previous analysis summarized by Eqs 8 and 9, we find that \approx 93% of the overall pulse speed is attributable to chemotaxis in the case of weak confinement.

As a final confirmation of this point, we re-run the simulations, but with either growth or chemotaxis removed—shown by the second and third columns of Fig 6, respectively—thereby isolating the contributions of chemotactic and growth-driven spreading. In the prototypical case of intermediate confinement, both chemotactic and growth-driven spreading play appreciable roles; compare Panels E and H to B in Fig 6, as well as the different curves in Fig 7B and 7E. However, in the case of weak confinement, chemotactic spreading dominates, as expected; the simulation without growth (Fig 6D) is nearly identical to that incorporating all factors (Fig 6A), while the simulation without chemotaxis (Fig 6G) yields a population that barely spreads—also seen by comparing the different curves in Fig 7A and 7D. Therefore, chemotactic propagation dominates under lesser confinement, enabling the population to spread faster as a sharp, coherent pulse.

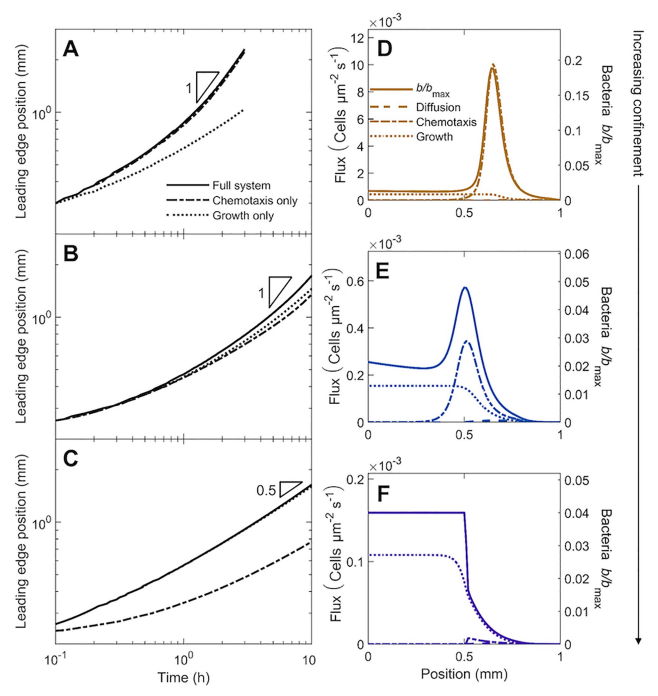


Fig 7. Population dynamics, morphology, and fluxes driving spreading for simulations of bacteria in weak, intermediate, and strong confinement (Fig 6) as shown by the top, middle, and bottom rows, respectively. (A-B) Increase in the position of the leading edge of the population is initially hindered, but approaches constant-speed motion indicated by the triangle at long times. In strong confinement (C), however, the long-time behavior approaches diffusive-like scaling instead. (D-E) At long times, the population spreads as a coherent pulse (solid line) driven primarily by chemotaxis in weak confinement, and by both chemotaxis and growth in intermediate confinement. (F) In strong confinement, however, the population spreads slowly as a jammed front, driven primarily by growth.

3.2.2 Strong confinement. Population spreading is markedly different in strong confinement. In this case, cells do not form a coherent pulse at all; instead, they continually grow outward as a jammed front (Fig 6C), unlike the case of intermediate confinement (Fig 6B). Notably, this front does not have a well-defined speed at long times, in stark contrast to the cases of weaker confinement explored previously. Instead, the leading edge position progresses as $x_l \sim t^{\nu}$ with $\nu \approx 0.5$ at long times, as shown by the solid curve in Fig 7C—and thus, the population spreads less effectively. This diffusive scaling of x_l is at odds with the prediction of the classic Fisher-KPP model, commonly used to describe growth-driven spreading, that the population spreads at a constant speed as a traveling wave [27, 47]. Instead, our finding is consistent with the results of agent-based simulations of a growing population of jammed, incompressible cells [41], which also found $v \approx 0.5$ in the limit of fast nutrient consumption. In this case, front propagation via growth of the jammed population lags behind nutrient diffusion—leading to the diffusive scaling of x_l observed in our simulations as well as those of [41]. This difference with the prediction of the classic Fisher-KPP model suggests that the logistic form of growth used therein does not adequately describe jammed growth spreading. We are not aware of any experiments testing this prediction; performing such a study would be a valuable direction for future research.

This dominant role of growth in driving spreading in the case of strong confinement is expected from the smaller value of Λ ; it is also corroborated by quantification of the different fluxes driving spreading (Fig 7F). Removing growth or chemotaxis from the simulation provides a final confirmation of this point; the simulation without growth (Fig 6F) yields a population that barely spreads, while that without chemotaxis (Fig 6I) is nearly identical to that incorporating all factors (Fig 6C)—also seen by comparing the different curves in Fig 7C and 7F. Hence, growth-driven spreading dominates under stronger confinement, enabling the population to spread diffusively as a jammed front.

4 Discussion

Ever since the discovery of bacteria over 300 years ago, lab studies of their spreading have typically focused on cells in unconfined environments such as in liquid cultures or near flat surfaces. However, in many real-world settings, bacteria must navigate complex and highlyconfining environments. Thus, motivated by experimental observations of bacterial motility [28, 34–38] and growth [39, 40] in confined settings, in this paper, we have presented an extended version of the classic Keller-Segel model that incorporates the influence of confinement on bacterial spreading through both motility and growth. Versions of the Keller-Segel model describing cellular aggregation and pattern formation in response to cell-generated chemoattractant have in some cases considered cell density-dependent motility [46], but do not explicitly consider confinement, and do not also incorporate cellular growth. Moreover, to our knowledge, there is no version of the Keller-Segel model of bacterial spreading in response to external chemoattractant that treats the density- and confinement-dependence of motility in an experimentally-motivated manner, and also incorporates cellular growth. The model described here provides a first step toward filling these gaps in knowledge, and in doing so, enabled us to examine the confinement-dependent interplay between motility-mediated and growth-mediated spreading.

In particular, our extended model treats cellular collisions with rigid surrounding obstacles, cellular collisions with each other, and growth-mediated spreading of jammed populations of cells. As such, it helps to bridge the classic Keller-Segel model of chemotactic spreading—which does not treat these effects and is therefore only appropriate to describe the spreading of dilute populations in unconfined settings—and models of growth-driven spreading (e.g., [41,

42, 71])—which do not treat motility-based spreading and are therefore only appropriate to describe the spreading of highly-concentrated/confined and non-motile populations. Indeed, non-dimensionalizing our extended model revealed the parameter Λ that quantifies the confinement-mediated transition between chemotactic spreading (in weak confinement with $\Lambda \gtrsim 1$) and growth-driven spreading (in stronger confinement with $\Lambda < 1$). Our analysis also provided a straightforward way to estimate, in general, the relative contributions of chemotaxis and growth to the speed with which a population spreads.

While our prior experiments [28] motivated and helped to parameterize and validate the model used in this study, our prior work did not provide a full computational analysis of the vastly different confinement-dependent spreading behaviors encoded by the model, and how they can jointly influence bacterial population dynamics. Accomplishing this task was the central goal of the present manuscript. To this end, numerical simulations of the model enabled us to examine the implications of the confinement-mediated transition in behaviors for the full dynamics of bacterial spreading. As expected, in weak confinement, a dense inoculum of bacteria rapidly depletes nutrient locally, causing a coherent pulse of cells to detach and continually propagate outward via chemotaxis—as predicted by the classic Keller-Segel model [26, 27, 29–33]. However, with increasing confinement, cellular crowding increasingly hinders both the initial formation of this pulse as well as its long-time propagation speed. Moreover, with increasing confinement, growth plays an increasingly dominant role in driving population spreading—eventually leading to a transition from fast chemotactic spreading to slow, growth-driven spreading of a jammed front [41]. Therefore, confinement is a key regulator of population spreading.

While chemotactic pulse propagation is well-characterized in unconfined settings [23–26], and conversely, jammed growth expansion has been investigated in some highly-confined settings [39, 40], the interplay between these two behaviors has scarcely been studied. Hence, we anticipate that our numerical characterization of this confinement-mediated transition from chemotactic- to growth-driven spreading will help guide future experimental investigations of confined populations. Moreover, because our model describes spreading over large length and time scales, we expect it could help more accurately describe the spreading dynamics of bacteria in processes ranging from infections, drug delivery, agriculture, and bioremediation. To this end, it would be interesting to extend our one-dimensional simulations to higher dimensions, which could result in additional rich dynamics e.g., as recently explored in [73], and to media with spatially-varying confinement.

Our model represents a first step toward capturing the essential biophysical processes underlying these complex dynamics, and necessarily involved some simplifying assumptions and approximations. For example, based on recent experiments [28], we treated the influence of cell-cell collisions using a mean-field approach in which the transport parameters D_{b0} and χ_0 are truncated in a cell density-dependent manner; incorporating more sophisticated collective dynamics [74–77] will be an important extension of our work. Similarly, we described jammed growth expansion by treating the population as an incompressible "fluid", similar to other models of growing immotile populations [41, 69, 70] and motivated by some experiments [39, 40], implemented in a discrete manner. An alternate continuum description of growth could e.g., track the local growth velocity defined from the spatial gradient of a pressure field within the growing population that originates from cell growth. For the purposes of this paper, in which our central goal was to characterize the dynamics of population spreading, we needed to only track the motion of the outer boundary of the jammed region—which is readily accomplished using our discrete representation of growth expansion. Developing a more detailed treatment of these growth dynamics using either continuum or discrete approaches, such as by incorporating cellular deformations [41] and possible changes in cellular behavior that may result [78], will be a useful direction for future work. Furthermore, a simplifying

assumption made in our model is that the solid obstacles that induce confinement are rigid and immovable; incorporating deformations of the surrounding medium will likely give rise to even more complex dynamics that will be interesting to study. Finally, while our model assumes that nutrient diffusion is unimpeded by the solid medium—which is likely to be the case in highly-permeable media such as biological gels and microporous clays/soils—incorporating hindered nutrient diffusion that may arise in other media will likely result in more complex dynamics that future extensions of our work could explore.

Supporting information

S1 Fig. Simulations corresponding to Fig 6A-6C, but using the full diffusive term (Eq 12) in our model; we observe nearly identical results to those presented in the main text. (TIFF)

S2 Fig. Visualization of terms (i) and (ii) of Eq 12 corresponding to the cell density profiles in Fig 3B and 3C. (A) Top shows the same cell profile as in Fig 3B; bottom shows corresponding diffusion terms. This example of an early time point shows that term (ii), as used in the main text, is larger than term (i), the neglected term. (B) Top shows the same cell profile as in Fig 3C; bottom shows corresponding diffusion terms. This example of a late time point shows that term (ii), as used in the main text, is typically larger than term (i), the neglected term. We note that the magnitude of the diffusive flux during this later time is much smaller than that of chemotaxis, as shown in Fig 3C; thus, the minor impact on diffusion of including term (i) does not change the finding that diffusion is negligible relative to chemotaxis. (TIF)

S3 Fig. To assess the sensitivity of our results to numerical discretization, we repeat the simulation shown in Fig 2, which has spatial resolution of $dx = 10 \mu m$, with varying values of dx; the time step dt is correspondingly varied as $dt = 0.01 \text{ s} \times (dx/10 \mu m)^2$. As shown in the figure, the final pulse velocity v_{pulse} obtained from the simulations is not strongly sensitive to the choice of numerical discretization. (TIFF)

S4 Fig. Experimental cellular signal of traveling front under intermediate confinement from [28]. Experiments begin with a dense packed cylindrical inoculum of *E. coli* embedded within a porous media with mean pore size 1.7 μ m. A pulse forms and propagates outward; the dataset shows the final time point of 10.75 h. The experiment used confocal microscopy of cells constitutively expressing green fluorescent protein; we take the fluorescence data thereby obtained from the mid-plane of the bacterial cylinder and normalize it by the brightest region of the initial inoculum. This normalized cellular signal is then converted to cell density by multiplying with $b_{\text{max}} = 0.95 \times 10^{12}$ cells/mL. Arrow indicates location identified as trailing behind the pulse, x_{trailing} , and the corresponding cellular density is $b_{\text{trailing}} = 1.5 \times 10^9$ cells/mL. (TIFF)

S1 Video. Numerical simulations of bacterial spreading from a dense inoculum in intermediate confinement. Video corresponds to Fig 2. Top shows the dynamics of the cells while bottom shows the corresponding dynamics of the nutrient, quantified by the normalized density $b/b_{\rm max}$ and concentration c/c_{∞} , respectively. Lower inset shows the same nutrient data, but with both axes zoomed out). (MP4)

S2 Video. Numerical simulations of bacterial spreading from a dilute inoculum in intermediate confinement. Video corresponds to Fig 4. Top shows the dynamics of the cells while

bottom shows the corresponding dynamics of the nutrient, quantified by the normalized density $b/b_{\rm max}$ and concentration c/c_{∞} , respectively. Top inset shows the same cellular data, but with the vertical axis zoomed in. Lower inset shows the same nutrient data, but with both axes zoomed out).

(MP4)

S3 Video. Numerical simulations of bacterial spreading in weak, intermediate, and strong confinement, also with growth or chemotaxis removed. Video corresponds to Fig 6, but also includes the nutrient profiles in orange (same vertical scales as in Figs 2 and 4). Different rows show different degrees of confinement, while different columns show full simulations or simulations with growth or chemotaxis removed. The simulations shown in different rows progress over different durations of time.

(MP4)

S1 File. Sample MATLAB code to implement simulations. This sample code corresponds to Fig 2.

(ZIP)

S2 File. Raw data underlying all Figures. This zip file contains the raw data used to make all the Figures.

(ZIP)

S1 Text. Further details of the model. This text presents additional details on how diffusion is incorporated in our model.

(PDF)

S2 Text. Determining parameters from experimental data. This text presents additional details on how the values of model parameters were extracted from experimental data. (PDF)

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Author Contributions

Conceptualization: Daniel B. Amchin, Tapomoy Bhattacharjee, Sujit S. Datta.

Data curation: Daniel B. Amchin, Sujit S. Datta. **Formal analysis:** Daniel B. Amchin, Sujit S. Datta.

Funding acquisition: Sujit S. Datta. **Investigation:** Daniel B. Amchin.

Methodology: Daniel B. Amchin, Sujit S. Datta.

Project administration: Sujit S. Datta.

Resources: Sujit S. Datta.

Software: Daniel B. Amchin, Jenna A. Ott.

Supervision: Sujit S. Datta. **Validation:** Daniel B. Amchin.

Visualization: Daniel B. Amchin.

Writing – original draft: Daniel B. Amchin, Sujit S. Datta.

Writing – review & editing: Daniel B. Amchin, Sujit S. Datta.

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