Bacterial Route Finding and Collective Escape in Mazes and Fractals

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Bacteria which grow not on the featureless agar plates of the microbiology lab but in the real world must navigate topologies which are nontrivially complex, such as mazes or fractals. We show that chemo-sensitive motile E. coli can efficiently explore nontrivial mazes in times much shorter than a no-memory (Markovian) walk would predict, and can collectively escape from a fractal topology. The strategies used by the bacteria include individual power-law probability distribution function exploration, the launching of chemotactic collective waves with preferential branching at maze nodes and defeating of fractal pumping, and bet hedging in case the more risky attempts to find food fail.

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1. INTRODUCTION

The two topologies of mazes and fractals pose quite different challenges to motile bacteria. A maze typically has well-defined entrance and exit ports [1]. However, within a maze there may be nodes leading to dead ends and internal loops which greatly lengthen the time it takes to exit from the maze if followed. While exit from a maze by a stochastic random walk is inevitable, the time to escape from a maze via a random walk can be quite long because the distance covered scales as the square root of time [2]. There are many different kinds of fractals; the ones we work with here are treelike fractals [3]. The key characteristic of a fractal is self-similarity: A fractal looks locally the same as you branch down into the fractal, hence there are no obvious physical clues as to how deep you are into the fractal, and this makes escape from a fractal more challenging than from a maze. We show that chemotactic bacteria get temporarily trapped in a fractal but eventually escape in a collective manner.

The maze we designed is shown in Fig. 1(a). It is a nontrivial, non-simply-connected maze [4]. A calculation
given in the Appendix A shows that we can expect a collection of identical random walkers to take anywhere from 150 h (10 walkers) to 20 h ($10^6$ walkers) for the first walker to emerge on average, where the walkers travel along the shortest path of length $L = 3 \times 10^3 \text{ μm}$, assuming an effective diffusion coefficient $D_b$ of $300 \text{ μm}^2/\text{sec}$ for each bacterium. Even a nonrandom walker, such as a wall follower, will take a long time to exit (or even fail) if the maze is not simply connected and contains internal loops. Figures 1(a) and 1(b) show the schematic of our maze, both the actual microfabricated design and broken down into its elements. As in any maze, there is an entrance and an exit, in our case the exit leads to a ring of microhabitats which are connected to an external food source by nanoslits [5].

The fractal tree we work with is presented in Fig. 1. It is a physical fractal of circular microhabitat “leaves,” with levels from 0 to $N = 6$. The leaves of the fractal tree are a series of microhabitats of radius $R_n = ζ^n \times 425 \text{ μm}$. The leaves are connected via long and narrow branches of width $w_m = ζ^m \times 70 \text{ μm}$ and length $l_m = ζ^m \times 2600 \text{ μm}$, hence the aspect ratio $σ$ is the same for every tunnel and preserves self-similarity, with $n \in [0,N]$, $m \in [0,N−1]$, and $ζ = 5/8$. The number of microhabitats of level $n > 1$ is given by (see Appendix B)

$$N = \frac{(3 - \sqrt{3})(2 + \sqrt{3})^n + (3 + \sqrt{3})(2 - \sqrt{3})^n}{2}. \quad (1)$$

For large $n \gg 1$, the value $N$ obeys the self-similar power-law growth $\sim γ^n$ with $γ = 2 + \sqrt{3}$, indicative of a scale-free “heavy-tail” network with fractal dimension $D$ given by

$$\lim_{n \rightarrow +\infty} N(n) \sim (ζ^n)^{-D} \Rightarrow D = -\ln γ/ \ln ζ. \quad (2)$$

For our fractal tree $D \approx 2.90$. The fractal dimension of the human lung $D_{\text{lung}} \approx 2.5$ [6], hence our fractal can be viewed as rather similar to a human lung. Although $E.\ coli$ do not typically infect a lung, other species of bacteria can form very serious resistant infections in the human lung [7].

![Diagram](image.png)

FIG. 1. (a) A non-simply-connected maze. Bacteria are introduced in the center. The shortest path is shown in blue hexagons. (b) Three nodes with high branching levels labeled A, B, and C are shown. (c) A fractal of dimension $D = 2.90$. Bacteria are introduced in the center hole.
II. METHODS

The chip topologies were microfabricated on a 100-mm silicon wafer using standard silicon etching techniques. We grew at 1000°C in an O₂ atmosphere a 0.2-μm-thick SiO₂ layer to obtain a hydrophilic surface after the etching step. We observed that bacteria left behind a surface modification of the SiO₂ which prevented stable rewetting of the chip [8,9]. To enable fresh reruns the chip surfaces were cleaned by a wet HF strip of the SiO₂ surface followed by thermal regrowth of the SiO₂ surface. The top of the etched chip was sealed by pressing a 20-μm-thick gas-permeable LUMOX polymer film LUMOX (Sarstedt AB, Nümbrecht, Germany) using pressurized air (10⁴ Pa). Since LUMOX is highly gas permeable the bacteria in the chip were in equilibrium with 21% O₂ levels at all times. All experiments were done at 20°C.

The chip was sterilized with an initial 70% ethanol rinse followed by 2 min of O₂ plasma sterilization in a Harrick PDC-001 plasma cleaner (Harrick Plasma, Ithaca, NY). The plasma treatment was also important in making the chip surface temporarily hydrophilic for enhanced wetting. The sterilized chip was wet with 20 g/L Luria-Bertani (LB) broth so the initial conditions were uniformly high food concentrations everywhere.

We used the bacterial strains W3110 for chemotactic experiments and isogenic W3110 cheA resulting from P1 transduction of a cheA deletion carrying strain (RP9535) that has lost its chemotaxis ability. To capture bacteria motion into the mazes, both strains express green fluorescent proteins (GFP) under the Plac promoter, which intrinsically express GFP. The bacteria were cultured in LB broth at 30°C for 24 h to reach saturation, then 20 μL of the culture transferred into a fresh 2-mL LB broth tube for 3 h at 30°C to maintain motility under constant shaking until mid-log phase (0.6 OD). Then 0.5 μL of the mid-log bacterial growth was taken directly from the culture tube and inoculated at the center of the chip, and immediately covered with a 3-mm-diameter cover glass to prevent evaporation.

Images were taken in a Nikon 90i upright microscope (Nikon Instruments Inc., Melville, NY) with an Andor Neo 5.5 sCMOS camera (Andor Inc., Concord, MA) using a Nikon 4X Plan APO λ objective. A 3 x 3 stitch field was used, with 0.5-sec exposure times per field, so each stitched field took approximately 5 sec to accumulate. Stitched fields were taken every 2 or 5 min. Images were stitched together using NIKON ELEMENTS software [10] and later analyzed with ImageJ software [11] and MATLAB software (MATLAB and Statistics Toolbox Release 2019b, The MathWorks, Inc., Natick, MA). Stitching of high-magnification images allowed us to detect individual bacteria while still observing the global dynamics of the bacteria.

III. RESULTS

A. Mazes

We first show the maze escape experiments. In a microfabricated maze with one inlet and one outlet but a large internal array of paths, it is virtually impossible to have zero pressure difference between the input and output ports, and thus there is typically a very gentle fluid flow through the device. We use 40-nm-diam red fluorescent beads in an attempt to track the flow, but the velocity of the flow, estimated to be under 1 μm/sec in the narrow channels and under 0.01 μm/sec in the open areas, is low and overwhelmed by diffusion of the beads, while larger beads simply sink and stick to the surfaces and are immobile.

Motile bacteria are sensitive to net biases in flow, called rheotaxis [12], and may also simply be unable to swim against too strong a flow velocity. As we show, depending on the sign of the flow into or out of the maze entrance, the collective bacteria population escape dynamics are quite different, which we attribute here to changes in the bacterial population dynamics rather than rheotaxis. Since the flow velocities are so low that they are difficult to directly measure, we assume a net flow to be outbound from the entrance to the exit if individual bacteria when inoculated immediately enter the maze. Figure 2 shows a series of images of a run with single bacterial resolution. Supplemental Material video smovie-1 [13] shows the full movie. Visualization of individual bacteria at the movie length scale shown is difficult to detect by eye, but easily captured using ImageJ software.

For an outflow situation, it should be no surprise that bacterial flow out of the maze along the shortest path even if they are not chemotactic, or even not motile due to pure advection of the medium. Since we are able to track the bacteria individually in the case of low number density it is also possible to measure the number density ρ(x, t) of the number of bacteria along the shortest path through the maze.

We compute at time t the number of bacteria S(x, t) which have gotten to a distance x or greater for both chemotactic and nonchemotactic bacterial strains, a form of a survival function [14]:

\[ S(x, t) = 1 - \frac{\int_{L_{ent}}^{L_{exit}} \rho(x')dx'}{\int_{L_{ent}}^{L_{exit}} \rho(x')dx'}, \tag{3} \]

where \( L_{ent} \rightarrow L_{exit} \) is the path of length \( L = L_{exit} - L_{ent} \) from entrance to exit. Figure 2 shows \( S(x, t) \) as a function of distance along the shortest path out of the maze versus time at 270 and 360 min for chemotactic bacteria and nonchemotactic bacteria. Figure 2 shows that the chemosensitive and chemoinsensitive bacteria have very different survival functions \( S(x, t) \), with the chemosensitive bacteria
presenting a power-law decay for $S(x, t)$, while chemoinsensitive bacteria have an exponential survival curve.

We are most interested in the number distribution of the bacteria in the maze versus time. Since we see power-law distributions and not exponentials, this indicates some form of chemotaxis is occurring. One possibility is a biased random walk, which in the limit of all bacteria having the same bias still yields exponentials in contradiction to the data, but could yield rapid passage. However, we do not explicitly introduce known powerful chemoattractants such as $L$ aspartate, and whatever gradient exists must be quite gentle since our path is $3 \times 10^4$-$\mu$m long. Thus, we would guess that the results of Nagy et al. [15] for weak chemoattractants such as $L$ lysine over gradients on the order of 100 mM/1000 $\mu$m to be UPPER values for what biased chemotaxis in LB would yield since our gradient is about $1/20$ their gradient. In the work of Nagy et al. it takes about 4 h for $E. coli$ to segregate over $1/20$ our length scale. With weak-biased chemotaxis a weak-biased random walk should take least 80 h. We believe a homogeneous-biased walk does not explain our result.

Power-law probability distributions are indicative of a non-Markov (persistent) walk where the bacteria do not randomly choose step sizes after each step, but rather show a persistent distribution in step sizes $g(l_p)$, where $l_p$ is the persistent step size of a given bacterium, which is maintained from step to step. Assuming then a power-law distribution of $g(l_p, t)$, we map this step size over to an effective diffusion constant distribution $g(D, t)$. The probability $P(x, t)$ then of finding a bacterium at position $x$ at time $t$ becomes, assuming a Gaussian kernel for $P_D(x, t)$ for a given diffusion constant,

$$ P(x, t) = \int_{D_{\text{min}}}^{D_{\text{max}}} g(D, t)P_D(x, t) dD. \quad (4) $$

If we assume that $g(D, t) \sim D^{-\gamma(t)}$, the spatial-dependence part of the survival function becomes

$$ S(x, t) \sim x^{1-2\gamma(t)}. \quad (5) $$

Since we see a power-law survival distribution as is predicted by Eq. (5), the conclusion seems to be that in a maze in outflow conditions the bacteria lock into a power-law distribution of step sizes in a non-Markovian random flight model of bacterial searching since each bacterium chooses stochastically a different step size with each iteration [16,17]. Here, we have long steppers and short steppers which are fixed phenotypes. Note that this distribution
$g(l_p, t)$ is rather like the power-law distribution of relaxation times seen in proteins frozen in at low temperatures [18]. The work of the Emonet group analyzing the motion of chemosensitive bacteria and spatial self-organization is related to what we see here [19].

The evolutionary advantage to such a power law might be that there exist chemotactic “pioneers” which explore regions rapidly tumbling, and conservatives which explore spaces more slowly, a theme which is now becoming evident from agar plate work [20,21] and now reveals itself in a more realistic maze environment with outflow.

The bacterial population dynamics in a maze for inflow conditions are very different but perhaps connected to the outflow situations we just presented. Figure 3 shows a series of stitched images of a population of bacteria introduced to the center of the maze with what we believe was a net inward speed from the exit to the entrance in the narrow channels of under 1 μm/sec (which corresponds to about 0.03 μm/sec in the open hexagonal habitats). For a period of approximately 6 h the bacteria basically do not (possibly cannot individually) enter the maze but rather build in density by reproduction until suddenly a collective surge of bacteria enters the maze, conquering the inward flow. Once the collective entry occurs a solitary wave of bacteria penetrates the maze and after a period of several hours collectively exits the maze, in a time far shorter than a simple random walk. Note that the maze fills out in a nonergodic manner, but not the sole path shown for outflow conditions. The full movie can be found in Supplemental Material video smovie-2 [13].

The average speed of the wave moving through the maze is approximately 1 μm/sec in the open habitat hexagons moving toward the exit. Note also that by no means do all of the bacteria track with the main wave front: There are bacteria which lead the main front; there are also bacteria which do not continue to move with the wave front through the maze but drop out, as shown in Fig. 3. These front-leading bacteria are presumably different from the bacteria which entered the maze before the wave formed. The wave also leaves behind “bet hedgers” bacteria no longer traveling with the wave front. Although passive chemical waves can navigate mazes [22], they show little of this

FIG. 3. Stitched images of bacteria entering a maze with a net collective inward flow at selected times. We have superposed the maze as red hexagons, the shortest path is shown with blue hexagons, in every other time slice.
complexity. The biological origins of this complexity, which we assume provides a fitness advantage, we do not know.

It is clear from Fig. 3 that the wave-propagating bacterial front does not uniformly fill the array but rather that some areas are populated more than others. If the collective wave-propagation front is merely due to classic bacterial Patlak-Keller-Segel (PKS) chemotactic response [23,24], at every maze path junction the wave should bifurcate equally into all exiting channels since initially there is no chemical clue which way to go. However, the wave seems to preferentially choose the path which has the shortest path to the next node. Figure 4 shows graphically an example of this: At node C in the maze an incoming solitary wave is faced exits to five outgoing channels, only one of which leads to the shortest path to the next major node. Although all five nodes are entered, the majority of the wave population entering the node chooses the correct shortest path. Exactly how this preferred selection of the “correct path” occurs we do not know, but it may be connected to the leading edge pioneer bacteria.

B. Fractals

Fractals pose a different challenge to bacteria but we see strategies related to the maze solutions. Since the fractal had no exit the flow issues of the maze work are not present, but we see both strategies observed in the maze experiment used in a fractal to good effect. Figure 5 presents six time frames of bacterial density vs time after bacterial inoculation in our fractal. A video of this can be seen in the Supplemental Material video smovie-3 [13].
the chemosensitive bacteria, that is the fractal serves as a pump to chemotactic swimmers to the smallest length scale of the fractal.

We measure the probability distribution of this initial trapping by measuring the local bacterial density \( \rho(r, t) \) at radius \( r \) from the inoculation center occupied by bacteria. This trapping only occurs for chemosensitive bacteria. Simple random walkers, even in a fractal landscape, should show a Gaussian distribution for \( \rho(r, t) \), which we verify both by running nonchemosensitive bacteria (strain RP9535 Δ cheA) as seen in Fig. 5, and by running a simulation on MATLAB (see Supplemental Material smovie-4, smovie-5, and Appendix D [13]).

For chemosensitive bacteria after inoculation the bacterial density distribution is initially a power-law distribution in space like the bacteria moving into a maze as we discussed above, as seen in Fig. 5(a). Because of the termination of fractal at the smallest scale, the leading bacteria become trapped and the distribution becomes inverted, with a probability distribution increasing with radius, rather than decreasing with distance, as seen in Fig. 5(b). This initial fractal trapping of bacteria shows no collective dynamics and can be compared to the outflow bacterial dynamics of the maze followed by pileup at the fractal cutoff scale due to fractal pumping.

After the fractal trapping stage, counterpropagating collective waves are launched throughout the fractal, one set is launched from the inoculation port side outward, the other set is launched from the fractal pumped termini of the smallest scale fractal leaves inward. These outgoing and ingoing waves collide, bifurcating and scattering as we show in Fig. 4 at fractal levels, with the scattered waves moving into all arms of the fractal tree. The synchronization of the two counterpropagating waves and the bifurcation of the colliding waves into regions of low bacterial density would seem to strongly argue for a high degree of bacterial communication during this collective dynamics. Although these synchronized waves are best appreciated in the Supplemental Material video smovie-3 [13], Fig. 6 shows some stages of the collective dynamics.

The net effect of these counterpropagating collective waves is to defeat the fractal-pumping phenomena and end with a uniformly filled (ergodic) fractal tree.

**IV. ANALYSIS**

There are four fundamental phenomena seen here as we challenge bacteria with complex topologies: (1) individual bacterial exploration with a time-invariant non-Gaussian probability distribution; (2) collective waves which solve
mazes by path selection at maze nodes; (3) bet hedging for a fraction of the collective wave bacteria; (4) escape for bacteria from fractal pumping by collective wave scattering.

The collective waves we see are not due to a Fisher-Kolmogorov population nonlinearity [26] because they are only observed for chemotactic bacteria, but rather are connected to PKS chemotactic waves. These waves are not true solitons since they collapse and scatter when they collide with each other. In the maze we constructed with loops this collapse of colliding waves serves to facilitate killing the endless looping in the maze: A wave that enters a loop can collide with another wave and the resultant collapse stops the entry process. However, the scattering process also seems to direct the incident wave into shortest path exit channels.

That the PKS equations by no means represent the full range of bacterial approaches to exploring a nontrivial maze can be seen from the outflow maze experiments, where no obvious collective waves form yet the maze is rapidly solved with a non-Gaussian probability distribution. For the fractal a different phenomenon occurs: Fractal pumping traps bacteria and collective waves untrap the bacteria. The challenge in a fractal is different from a maze; unlike a maze a fractal is self-similar, there are no topological clues in a fractal as to where a walker is within the structure. Furthermore, scale invariance predicts that the habitat area $S_i$ and channel lengths $l_i$ connecting areas as you proceed down the fractal must satisfy the following scaling laws for a systems of habitats of size $\{S_n\}$ connected via tunnel of width $\{w_n\}$ and length $\{l_n\}$ with the scale decreasing at a constant factor $\zeta$ between consecutive levels:

$$\frac{S_{n+1}}{S_n} = \zeta^2, \quad \frac{w_{n+1}}{w_n} = \frac{l_{n+1}}{l_n} = \zeta. \quad (6)$$

In such a train of decreasing areas $S_n$ we expect that for chemosensitive bacteria there will be a continuous cascade of population collapse because the sign of $\kappa$ in the PKS equations drives bacterial aggregation. In our fractal device $\zeta = 5/8$. Chemotactic signaling driven collapse favors bacteria moving from large areas $S_{\text{large}}$ to smaller adjacent...
areas $\zeta S_{\text{large}}$ because for the movement of a given bacteria from large to small the effective density of the bacteria increases $\rho$, and so does the effective concentration of the chemoattractant $c$. The actual time dependent nonlinear dynamics of this process is extremely complex (see Appendix C), but in the limit of assuming that the food concentration $f$ is fixed and ignoring bacterial growth and death processes, the bacterial density difference $\Delta \rho = \rho_{n+1} - \rho_n$ is roughly given by

$$\partial_t \Delta \rho \sim \kappa \alpha \left( \frac{1}{\zeta} - 1 \right) \Delta \rho \Delta c,$$  \hspace{1cm} (7)

where we have lumped all fixed parameters in $\alpha$, and $\Delta c$ is the (time and bacterial density dependent) chemoattractant concentration difference $c_{n+1} - c_n$. Equation (7) although an approximation, cannot be integrated analytically but it demonstrates the qualitative phenomena of fractal pumping: There is no pumping if $\zeta = 1$ (see Appendix D), while decreasing $\zeta$ results in more rapid population collapse. Figure 6(b) shows a fit of Eq. (7) of the bacterial density along one sector of the fractal at $t = 240$ min after inoculation. This pumping is not stable but decays due to backward wave propagation.

In the absence of backward propagating collective waves a fractal is then a trap for chemosensing bacteria and there is no escape from the smallest structures of area $\zeta^{12} S_n$. This makes the fractal structures extremely dangerous both to the individual infected with bacteria and to the bacteria themselves since they will increase in numbers to very high densities driven by fractal pumping. The only way out is to launch collective waves in both directions; when two of the waves collide this results in the scattering of both waves, as is seen in Fig. 6, and the eventual ergodic filling of the entire fractal.

V. CONCLUSIONS

Natural complex topologies form challenging existential puzzles for bacteria. Billions of years of evolution have shaped the response of bacteria to these puzzles, whose solutions can be found in how bacteria such as the common *E. coli* respond collectively and individually to challenges. We pose challenges to bacteria in forms of mazes and fractal spaces, and show the unexpected and clever, if we can say that word about bacteria, way in which they are able to path find their way through a non-simply-connected maze, and escape from the trap of fractal pumping.

There are controversial aspects of this work we freely admit. Modeling of the collective dynamics is difficult. Unfortunately the PKS equations are highly nonlinear and cannot be solved analytically. Assumption of fixed food concentration such as we have done is the simplification needed to get the solution of fractal pumping in a nice analytic form, which we expect to be due to geometric reasons (or at least insensitive to the food change).

We believe that you need both geometry and also chemotactic sensing to fully understand our results.

Even modeling of the individual bacterial behavior is challenging. While using the random-walk results provides a useful benchmark, we find that the biased random walk is informative for the presumed outflow case, but it fails miserably for the inflow case which is really driven by collective dynamics, launching wave against the flow.

A third possibility to help understand our results is modification of the environment itself by the bacteria in a semipermanent way by surface modification. In fact, we think something like this could be happening in the maze (this is called in the mathematics of maze solving literature the “Hansel and Gretel” mode of solving a maze, where an agent leaves a trail behind which can be followed later.

There are many puzzles to be continued as we explore these ancient organisms. Although we know a great deal about the basic biological physics of chemotaxis [27], it is clear that much remains to be done to clarify more complex collective aspects.

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APPENDIX A: SEARCH TIMES IN MAZES

Finding the exit from a maze for bacteria can be quite formidable in terms of time if a simple random-walk algorithm is used because typically a very circuitous path of long length must be followed to the exit. If we assume that bacteria simply do a random walk over the shortest path of length $x_{\text{final}}$, it is straightforward to calculate the mean first-arrival time of $N$ bacteria at a position $x_{\text{final}}$, ignoring the effects of dead ends and branching decisions. The probability density $\rho(x,t)$ of a bacteria to arrive at position $x$ after time $t$ for a random (Markov process) walker is given by a Gaussian distribution:

$$\rho(x,t) = \frac{1}{\sqrt{\pi D_b t}} e^{-x^2/4D_b t}, \hspace{1cm} (A1)$$

where $D_b$ is the effective diffusion coefficient of a swimming bacteria. Hence, the probability $P_L(t)$ for a bacteria to reach the exit of the maze at time $t$ is

$$P_L(t) = \int_0^\infty dx \rho(x,t) = \text{erfc} \left( \frac{L}{\sqrt{4D_b t}} \right), \hspace{1cm} (A2)$$
where erfc is the error function. Therefore, the expected time $T$ for 1 bacterium out of $N$ total bacteria to reach the exit can be estimated to be

$$T = \left[ \text{erfc}\left(\frac{1}{\sqrt{N}}\right) \right]^{-2} \frac{L^2}{4D_b}, \quad (A3)$$

where erfc$^2$ is the inverse error function. Assuming for $D_b$, a diffusion coefficient of $300 \mu m^2/\text{sec}$ [28], this yields formidably long first-arrival times for bacteria. Increased numbers does not help much, the mean first-arrival times for bacteria. Increased steps (see Fig. 7). A leaf is a single habitat with a single tunnel, connected to another habitat that we add it as follows:

**Step 0.**—Create a single habitat.

**Step 1.**—Add $\alpha$ leaves to the habitat.

**Step $n \geq 2.$**—For every habitat, given its linking number $\rho$: (1) If $\rho = 1$, add $\alpha - 1$ leaves to it. (2) If $\rho > 1$, add $\rho$ leaves to it.

It should be noted that the physical size of the leaves is decreasing with steps. The difference in length scale between leaves of consecutive steps is a constant scale factor $\zeta < 1$.

2. Calculating the fractal dimension

We follow Ref. [29] to calculate the fractal dimension. In short, given the number of structures $N(n)$ (the number of leaves at step $n$) at the characteristic length scale $\lambda(n) \sim \zeta^n$ (the length scale of the leaves at step $n$), the fractal dimension $D$ can be estimated by

$$D = -\lim_{n \to \infty} \frac{\ln N(n)}{\ln \lambda(n)}. \quad (B1)$$

If this limit is well defined, our geometry has a definite fractal dimension.

Note that the number of added leaves is equal to the number of added habitats at every step $n > 0$. For $n = 0$, the number of added habitats is simply 1. Define the number of habitats added at step $n$ to be $N(n)$, then

$$N(0) = 1, \quad N(1) = \alpha,$$

$$N(2) = \alpha(\alpha - 1) + \alpha = \alpha^2. \quad (B2)$$

Given the rules generating our fractal, any habitat added at step $n > 0$ will be added $(\alpha - 1)$ leaves at step $n + 1$, $\alpha$ leaves at step $n + 2$, $2\alpha$ leaves at step $n + 3, ..., 2^k \alpha$ leaves at step $n + k + 2, ...$. Therefore, for every step $n > 1$ one has the recursion relation

$$N(n) = (\alpha - 1)N(n - 1) + \sum_{j=0}^{n-2} 2^{n-2-j} \alpha N(j). \quad (B3)$$

Consider a shift by 1 in the recursion relation

$$N(n + 1) = (\alpha - 1)N(n) + \sum_{j=0}^{n-1} 2^{n-1-j} \alpha N(j). \quad (B4)$$

Therefore,

$$N(n + 1) - 2N(n) = \left( (\alpha - 1)N(n) + \sum_{j=0}^{n-1} 2^{n-1-j} \alpha N(j) \right)$$

$$- \left( 2(\alpha - 1)N(n - 1) + \sum_{j=0}^{n-2} 2^{n-1-j} \alpha N(j) \right)$$

$$= (\alpha - 1)N(n) + (2 - \alpha)N(n - 1). \quad (B5)$$

The recursion relation now becomes much more local:

$$N(n + 1) = (\alpha + 1)N(n) + (2 - \alpha)N(n - 1). \quad (B6)$$

Since Eq. (B3) holds only when $n > 0$, we can rewrite Eq. (B6) as followed when $n > 2$:

$$N(n) = (\alpha + 1)N(n - 1) + (2 - \alpha)N(n - 2). \quad (B7)$$

From the initial values (B2), one solves the above recursion relation and arrives at a complicated expression of the form...

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**APPENDIX B: FRACTAL ENVIRONMENT**

1. Generating our fractal

Our fractal is built from habitats and tunnels at many different scales (see Fig. 7). A leaf is a single habitat with a single tunnel, connected to another habitat that we add it on. A linking number of a habitat is the number of tunnel connected to it. Our $\alpha$-fold fractal is generated step by step as follows:

**Step 0.**—Create a single habitat.

**Step 1.**—Add $\alpha$ leaves to the habitat.

**Step $n \geq 2.$**—For every habitat, given its linking number $\rho$: (1) If $\rho = 1$, add $\alpha - 1$ leaves to it. (2) If $\rho > 1$, add $\rho$ leaves to it.

It should be noted that the physical size of the leaves is decreasing with steps. The difference in length scale between leaves of consecutive steps is a constant scale factor $\zeta < 1$.

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FIG. 7. The building of our fractal with $a$-fold, step-by-step from left to right. Here the parameter $a = 3$. 

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\[ \mathcal{N}(n) = c_+ r^n_+ + c_- r^n_. \]
\[ c_\pm = -\frac{\alpha [a - 5 \pm \sqrt{(a - 1)^2 + 8}]}{2(a - 2) \sqrt{(a - 1)^2 + 8}}, \]
\[ r_\pm = \frac{\alpha + 1 \pm \sqrt{(a - 1)^2 + 8}}{2}. \]

Plug this into (B1), one gets
\[ D = -\frac{\ln r_+}{\ln \zeta} = \frac{\ln 2 - \ln [\alpha + 1 \pm \sqrt{(a - 1)^2 + 8}]}{\ln \zeta}. \]

For \( \zeta = 1/2 \) and \( \alpha = 3 \), \( D \approx 1.90 \). For \( \zeta = 5/8 \) and \( \alpha = 3 \), the dimensionality is close to that of a human lung \( D \approx 2.80 \).

**APPENDIX C: GEOMETRICAL INFLUENCE ON BACTERIAL COLLECTIVE DYNAMICS**

The amplification of bacterial collapse in a multiscale environment can be explained theoretically with a nonlinear model of *E. coli* chemotaxis signaling network, which will be described in this section.

Consider a simple geometry that captures the multiscale nature—a system of habitats of size \( \{ S^n \} \) connected via tunnel of width \( \{ w_n \} \) and length \( \{ l_n \} \) with the scale decreasing at a constant factor between consecutive levels.

We end this multiscale series at level \( N \) (see Fig. 8):
\[ S_0 \rightarrow w_{1,1} \rightarrow S_1 \rightarrow w_{2,1} \rightarrow S_2 \rightarrow \ldots \]
\[ S_n \rightarrow w_{n+1,1} \rightarrow \ldots \rightarrow w_N \rightarrow S_N. \]

For level 0 being the largest structure, the factor \( \zeta \) has to be smaller than unity. The relation between consecutive levels’ geometrical values, as the length scale goes for a constant scale factor \( \zeta \):
\[ S_{n+1}^2 = S_n = \zeta^{2(N-n)}S_N, \]
\[ w_{n+1} = \zeta \Rightarrow w_n = \zeta^{n-N}w_N, \]
\[ l_{n+1} = \zeta \Rightarrow l_n = \zeta^{n-N}l_N. \]

The basic starting point for understanding collective bacteria dynamics is the Patlak-Keller-Segel equations [30,31], which in general are of the form
\[ \partial_t b = \chi_b(b,c,f) + D_b \nabla^2 b - \kappa_{bc} \nabla (b \nabla c), \]
\[ \partial_t c = \chi_c(b,c,f) + D_c \nabla^2 c, \]
\[ \partial_t f = -\chi_f(b,c,f) + D_f \nabla^2 f. \]

The densities of bacteria, chemoattractants, and food are given by the field \( b, c, \) and \( f \); the reproduction rate \( \chi_b \) of bacteria, the production rate \( \chi_c \) of chemoattractants, and the consumption rate \( \chi_f \) of food in general depends on both \( b, c, \) and \( f \); the diffusivity coefficients of bacteria, chemoattractants, and food are \( D_b, D_c, D_f \); the chemotaxis signaling can be represented by the sensitivity of bacteria to the chemo gradients, which has the sensitivity coefficient \( \kappa_{bc} \).

For simplification, we will look at the limit where there is no bacterial production and food concentration stays fixed:
\[ \chi_b \to 0, \quad f \to f_* = \text{const.} \]

Now, for the explicit form of \( \chi_c \), we will use the model in Brenner et al. [25]:
\[ \chi_c(b,c,f) = \beta f b. \]

Equations (C3) and (C4) become
\[ \partial_t b = D_b \nabla^2 b - \kappa_{bc} \nabla (b \nabla c), \]
\[ \partial_t c = \beta f_b + D_c \nabla^2 c. \]

### 1. Bacteria collapse due to chemotaxis

Consider the geometry given in Fig. 8 with \( N = 1 \) (a small habitat size \( S_1 \) connects to a big one size \( S_0 = \zeta^{-1}S_1 \) via a long and narrow tunnel \( w_1 \times l_1 \), using the mean-field approximation inside the habitats and across the tunnels with the bacteria density \( \{ b_0, b_1 \} \) and chemo concentration \( \{ c_0, c_1 \} \), the dynamics becomes
\[ \partial_t (S_0 b_0) = D_b \frac{w_1}{l_1} (b_1 - b_0) - \kappa_{bc} \frac{w_1}{l_1} b_0 + \frac{b_1}{l_1} (c_1 - c_0), \]
\[ \partial_t (S_1 b_1) = D_b \frac{w_1}{l_1} (b_1 - b_0) - \kappa_{bc} \frac{w_1}{l_1} b_1 + \frac{b_1}{l_1} (c_0 - c_1), \]
\[ \partial_t (S_0 c_0) = S_0 \beta f_b b_0 + D_c \frac{w_1}{l_1} (c_1 - c_0), \]
\[ \partial_t (S_1 c_1) = S_1 \beta f_b b_1 + D_c \frac{w_1}{l_1} (c_0 - c_1). \]

**FIG. 8.** A simple multiscale geometry that can be used to demonstrate the amplification of bacterial collapse and fractal attraction.
To demonstrate how chemotaxis signaling leads to the bacterial collapse phenomenon, drop diffusion terms from the equations. Define

$$\Delta_b = b_1 - b_0, \quad \Delta_c = c_1 - c_0. \quad (C12)$$

Since there is no bacteria growth,

$$b = \frac{S_1 b_1 + S_0 b_0}{S_1 + S_0} = \frac{b_0 + \frac{\kappa_c}{\mu} k_1}{1 + \frac{\kappa_c}{\mu} k_1} = \text{const}. \quad (C13)$$

From Eqs. (C10) and (C11), after some algebraic manipulations,

$$\Delta_b = \frac{1}{\beta f_s} \partial_t \Delta_c, \quad \frac{\kappa_c}{\mu} k_1 \partial_t \Delta_c = \frac{2b_0 + 1 - \frac{\kappa_c}{\mu} k_1}{1 + \frac{\kappa_c}{\mu} k_1} \partial_t \Delta_c \Delta_c. \quad (C14)$$

Linear order gives instability

$$\partial_t \Delta_c \approx (1 + \zeta) \frac{2b_0 + 1 - \zeta}{1 + \zeta} \partial_t \Delta_c. \quad \Rightarrow \Delta_b \approx e \sqrt{(1 + \zeta) \frac{2b_0 + 1 - \zeta}{1 + \zeta} \partial_t \Delta_c}. \quad (C15)$$

Note that we arrive at the same behavior found by Park et al. [32]—an exponential collapse.

The $\pm$ indicates that the collapse seems to be able to happen both ways—into the small habitat or into the large one, in which the later is not physical and has not been observed experimentally. Indeed, to get rid of the unphysical solution from a theoretical point of view, one has to keep the full nonlinear order

$$\partial_t \Delta_c \approx (1 + \zeta) \frac{2b_0 + 1 - \zeta}{1 + \zeta} \partial_t \Delta_c \Delta_c. \quad (C16)$$

From here then on, to get the positive or negative sign correctly, without loss of generality we consider $\zeta \leq 1$ as similar considerations can be done with $\zeta \geq 1$ with sign flips.

For $\Delta_b \sim \partial_t \Delta_c$ (C14) negative $< 0$, the difference $\Delta_c$ cannot grow arbitrarily large negatively—it is bounded from below:

$$\partial_t \Delta_c \geq -2 \frac{1 + \zeta}{1 - \zeta} \beta f_s b. \quad \Rightarrow \Delta_b \geq \Delta_b \bigg|_{\text{min}} \geq -2 \frac{1 + \zeta}{1 - \zeta} b. \quad (C17)$$

Hence, collapse will not happen in that direction, and even fluctuation of density due to bacteria randomly walking around can already surpass such a limit (make the bacteria go back) when the number of bacteria in the system (as the average density) is low enough:

$$-\Delta_b \bigg|_{\text{min}} \leq \frac{1}{h S_0} + \frac{1}{h S_1} \Rightarrow b \geq \frac{2(1 - \zeta)}{h S_1}. \quad (C18)$$

Note that $h$ is the depth of the environment. The above estimation is done by letting one single bacterium going from $S_1$ to $S_0$ cause a density change high enough to surpass $\Delta_b \big|_{\text{min}}$. For $\zeta \ll 1$, just a few bacteria getting inside $S_1$ might already make the density $b_1$ much larger than that of $b_0$.

For $\Delta_b \sim \partial_t \Delta_c$ (C14) positive $> 0$, the difference $\Delta_c$ can grow arbitrarily large positively—it is unbounded from above. At large time $t$ behavior,

$$\partial_t \Delta_c \approx \frac{1 + \zeta}{1 - \zeta} \beta f_s b \Rightarrow \partial_t \Delta_c \approx (1 + \zeta) \frac{\kappa_c}{\mu} k_1 \beta f_s \Delta_c. \quad (C19)$$

Note that the solution here is singular—the bacteria density can grow to infinity in finite time, and the smaller $S_1$ the stronger the collapse. However, physically, there should be a cutoff on how much the density can be due to the volume $\Omega$ of a bacterium:

$$b \big|_{\text{max}} \lesssim \frac{1}{\Omega}. \quad (C20)$$

This concludes our theoretical analysis on how bacteria collapse $n$ into smallest structures.

2. A physical model for fractal pumping

Consider the geometry given in Fig. 8 with $N \gg 1$, using the mean-field approximation inside the habitats and across the tunnels with the bacteria density $\{b_n\}$ and chemo concentration $\{c_n\}$, the dynamics becomes

$$\partial_t (b_n b_n) = D_b \frac{w_n}{I_n} (b_n b_n - b_n) + D_b \frac{w_{n+1}}{I_{n+1}} (b_{n+1} b_n - b_n) - \kappa_c \frac{w_n}{I_n} (b_{n-1} + b_n) \frac{c_{n-1} - c_n}{2} - \kappa_c \frac{w_n}{I_{n+1}} (b_n + b_{n+1}) \frac{c_{n+1} - c_n}{2}, \quad (C21)$$

$$\partial_t (c_n c_n) = S_c \beta f_s b_n + D_c \frac{w_n}{I_n} (c_n - c_n) + D_c \frac{w_{n+1}}{I_{n+1}} (c_n - c_n). \quad (C22)$$

Since the physical cutoff comes from the smallest structures, having the length scale comparable to that of a bacteria, we will keep $S_N, w_N,$ and $I_N$ fixed while the total number $N$ of levels can be changed:

$$S_N = S_+, \quad w_N = w_+, \quad I_N = I_+. \quad (C23)$$
The above system of ordinary differential equations is very hard to deal with, analytically. To go further, we will use the continuous representation for \( n \) so that the system becomes 2 partial differential equations:

\[
\begin{align*}
\partial_t b &= \sigma \partial_n \partial_n b = \sigma D_b \partial_n^2 b - \sigma \kappa_b \partial_n (b \partial_n c), \quad \text{(C28)} \\
\partial_t c &= \zeta_2^{-2(n-N)} \partial_n \partial_n c = \zeta_2^{-2(n-N)} \beta f_c b + \sigma D_c \partial_n^2 c. \quad \text{(C29)}
\end{align*}
\]

At stationary distribution of bacteria, Eq. (C28) leads to

\[
\begin{align*}
\partial_t b &= 0 \Rightarrow D_b \partial_n^2 b = \kappa_b \partial_n (b \partial_n c), \\
&\Rightarrow b \sim e^{(\kappa_b/D_c) c}. \quad \text{(C30)}
\end{align*}
\]

Bacteria density depends exponentially on chemo concentration is a feature of the nonlinearity of bacteria-chemo interaction.

To get some intuition, note that the physical role of the chemo to the random-walking self-propelled E. coli are similar to that of the energy potential (profile \( V \)) to particles (with density \( \rho \)) at thermal equilibrium (at temperature \( T \)):

\[
\begin{align*}
b &\sim e^{\kappa_b/D_c} \quad \Leftrightarrow \quad \rho \sim e^{-V/k_BT} \\
c &\sim -V. \quad \text{(C31)}
\end{align*}
\]

Hence, we will quantify the bacterial attraction with the chemo density \( c \).

Equation (C29) can be rewritten as

\[
\begin{align*}
\partial_t c &= \beta f_c b + \zeta_2^{-2(n-N)} \frac{\sigma}{S} D_c \partial_n^2 c. \quad \text{(C32)}
\end{align*}
\]

In the fast chemoattractant diffusion limit \( D_c \gg D_b \), for linear terms one can ignore the time-variation contribution \( \sim \partial_t c \) compared to that of the spatial-variation contribution \( \sim \partial_n^2 c \) and do the approximation

\[
D_c \rightarrow +\infty \Rightarrow 0 \approx \beta f_c b + \zeta_2^{-2(n-N)} \frac{\sigma}{S} D_c \partial_n^2 c. \quad \text{(C33)}
\]

Physically speaking, the chemoattractant field is treated as background. It should be noted that this approximation is also made in Brenner et al. [25].

Since bacterial collapse happens most strongly at the smallest scale structures, let us use the maximum possible bacteria density there \( b(N) = B \). Define

\[
\Delta(n) = c(n) - c(N). \quad \text{(C34)}
\]

The boundary conditions for \( \Delta \) at the smallest structure (level \( N \)):

\[
\Delta(N) = 0, \quad \partial_n \Delta(N) = 0. \quad \text{(C35)}
\]

Bacteria density can now be written as

\[
b = B e^{(\kappa_b/D_c) \Delta}. \quad \text{(C36)}
\]

Equation (C33) becomes

\[
0 = e^{(\kappa_b/D_c) \Delta} + \zeta_2^{-2(n-N)} \frac{\sigma}{S} \frac{D_c}{\beta f_c B} \partial_n^2 \Delta. \quad \text{(C37)}
\]

Redefine

\[
\tilde{\Delta} = \frac{\kappa_b}{D_b} \Delta, \quad \tilde{\zeta} = \zeta_2^{-2} \sqrt{(D_b / D_c)(\kappa_b / \beta f_c B)(\sigma/S)}. \quad \text{(C38)}
\]

One then arrives at

\[
0 = e^{\tilde{\Delta}} + \tilde{\zeta} \tilde{\nu} \partial_n^2 \tilde{\Delta}. \quad \text{(C39)}
\]

The general solution of this equation has the form

\[
\tilde{\Delta} = \ln \left[ \frac{\delta_1}{2} \sech \left( \frac{\sqrt{\delta_1}}{2} (\delta_2 + \tilde{n}) \right) \right] - \ln(\tilde{\zeta}) \tilde{n}. \quad \text{(C40)}
\]

From Eq. (C35), the boundary conditions are

\[
\tilde{\Delta}(\tilde{n} = 0) = 0, \quad \partial_n \tilde{\Delta}(\tilde{n} = 0) = 0. \quad \text{(C41)}
\]

The matching gives

\[
\delta_1 = 2 + \ln^2(\tilde{\zeta}), \quad \delta_2 = -\frac{\ln[1 + \ln^2(\tilde{\zeta})] + \ln(\tilde{\zeta}) \sqrt{2 + \ln^2(\tilde{\zeta})}}{\sqrt{2 + \ln^2(\tilde{\zeta})}}. \quad \text{(C42)}
\]

It is quite a complicated expression. In the limit of a large number of fractal orders \( N \gg 1 \) and drastic change of scale \( \zeta \ll 1 \), as \( n \rightarrow N \) one has the approximation \( \tilde{\Delta} \sim \tilde{n}^2 \).
To get bacteria distribution $b$ from $\Delta$, use Eq. (C36). The function can then be used to compare with results observed from experiments [see Figs. 9 and 6(b)].

**APPENDIX D: NONCHEMOSENSITIVE BACTERIA AND NO FRACTALS**

We test the role that chemotaxis plays in the role of fractal pumping by inoculating a fractal with a strain of bacteria which while motile has no chemotaxis sensitivity, strain RP9535 $\Delta$ cheA. Supplemental Material video smovie-4 [13] shows a full video of the bacterial movement, while Fig. 10 shows a snapshot of a simulation at 250 min. Without chemotaxis there is no fractal pumping.

The role of the fractal structure in driving bacteria can be tested using a simple branching with $\zeta = 1$. An example is shown in the Supplemental Material video smovie-6 [13]. This structure has branches which both simply terminate in dead ends and branches connected via nanoslits to food sources. There is no strong collective collapse into the dead ends and no back propagation away from food sources.

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