Patterns of bacterial motility in microfluidics-confining environments

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Understanding the motility behavior of bacteria in confining microenvironments, in which they search for available physical space and move in response to stimuli, is important for environmental, food industry, and biomedical applications. We studied the motility of five bacterial species with various sizes and flagellar architectures (Vibrio natriegens, Magnetococcus marinus, Pseudomonas putida, Vibrio fischeri, and Escherichia coli) in microfluidic environments presenting various levels of confinement and geometrical complexity, in the absence of external flow and concentration gradients. When the confinement is moderate, such as in quasi-open spaces with only one limiting wall, and in wide channels, the motility behavior of bacteria with complex flagellar architectures approximately follows the hydrodynamics-based predictions developed for simple monotrichous bacteria. Specifically, V. natriegens and V. fischeri moved parallel to the wall and P. putida and E. coli presented a stable movement parallel to the wall but with incidental wall escape events, while M. marinus exhibited frequent flipping between wall accumulator and wall escaper regimes. Conversely, in tighter confining environments, the motility is governed by the stochastic interactions between bacteria and the surrounding walls. In mesoscale regions, where the impacts of hydrodynamics and stochastic interactions overlap, these mechanisms can either push bacteria in the same directions in linear channels, leading to smooth bacterial movement, or they could be oppositional (e.g., in mesoscale-sized meandered channels), leading to chaotic movement and subsequent bacterial trapping. The study provides a methodological template for the design of microfluidic devices for single-cell genomic screening, bacterial entrapment for diagnostics, or biocomputation.

Significance

Understanding bacterial movement is crucial for health, agriculture, environment, and industry. Studying the motility of five bacterial species in microfluidic environments showed that bacterial motility behavior is the result of a “tug-of-war” between hydrodynamics and local nanomechanics. In less confining spaces, bacterial motility is governed by hydrodynamics and can be approximately predicted by modeling developed for the simplest species. Conversely, in tightly confining environments, movement is mainly controlled by the stochastic interactions between flagella and the surrounding walls. Intriguingly, in mesoscale-sized geometries, hydrodynamics and bacterial-wall interactions overlap, either “constructively,” leading to smooth movement in straight channels, or “destructively,” leading to trapping. Our study provides a methodological template for the development of devices for single-cell genomics, diagnostics, or biocomputation.

Many motile bacteria live in confining microenvironments (e.g., animal or plant tissue, soil, waste, granulated, and porous materials) and consequently are important to many applications like health [infectious diseases (1, 2), pharmaceuticals (3), and nutrition (4)], agriculture [veterinary (5) and crops (6)], environmental science [photosynthesis (7), biodegradation (8), and bioremediation (9)], and industrial activities [mining (10) and biofueling (11)]. Bacterial motility is essential in the search for available physical space as well as for enabling bacterial taxis in response to external stimuli, such as temperature (12), chemical gradients (13, 14), mechanical cues (15), or magnetic fields (16).

To thrive in environments with diverse geometrical and physical characteristics, from open spaces to constraining environments, motile bacteria have evolved a multitude of propelling mechanisms (17), with flagellum-driven being the most common (18, 19). Flagellum-based machinery features various numbers of flagella (20) and designs: monotrichous, lophotrichous, amphitrichous, or peritrichous. The mechanics of this machinery, coupled with cell morphology (21) (e.g., coccus, rod-like, or curved) translates into several motility modes (e.g., turn angle, run-and-tumble, or run-and-flick) (22), and various motility behaviors (e.g., swimming, tumbling, and swarming) (17, 23). Environmental factors (24, 25) (e.g., chemical composition, viscosity, temperature, pH, and the chemistry and the roughness of adjacent surfaces) also influence bacterial motility.

“Pure” bacterial motility, unbiased by chemotaxis or fluid flow, was reported near simple flat surfaces (26, 27) and in channels (28–30). Simulations of model bacteria in analogous conditions were also undertaken (31–37), but owing to the complexity of bacterial mechanisms (38), modeling from first principles did not provide sufficient understanding to accurately predict movement patterns of different species in complex, confined environments. Consequently, studies of the effects of bacterial geometry in confined geometries were limited to models of simple, monotrichous bacteria with an assumed rigid flagellum (32, 39).

Microfluidic devices (40, 41) are commonly used for the manipulation of individual or small populations of cells in micrometer-sized channels for medical diagnostics (42), drug screening (43), cell separation (44, 45), detection and sorting (46), and single-cell genomics (47). While microfluidic structures are used for the study of bacterial motility | microfluidic devices | space partitioning | wall escaper | wall accumulator

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the motility of mammalian cells (48, 49), and microorganisms [e.g., fungi (50, 51), algae (52), or bacteria (29, 53–56)], these studies typically focus on a single species.

To make progress toward a more general understanding of the motility of individual bacterial cells in confining microenvironments, as well as to assess the extent to which the behavior of bacteria with complex architectures can be assimilated with that of the more predictable monotrichous bacteria, the present work investigated the movement of five species (i.e., *Vibrio natriegens*, *Magnetococcus marinus*, *Pseudomonas putida*, *Vibrio fischeri*, and *Escherichia coli*) in microfluidic geometries with various levels of confinement and geometrical complexity.

**Results and Discussion**

The modulation of motility behavior by confinement was assessed by observing, through three-dimensional (3D) imaging, the movement of individual bacteria in microfluidic geometries with various levels of confinement and linear channels (top row), angled, and meandered channels (bottom row); and 5) SEM image of a bacterium (here, *E. coli*) in a channel.

**Motility in Large Chambers.**

**Impact of the distance between horizontal planes.** To minimize the possible coupling of the impact of horizontal planes, the designs of microfluidic chambers, made of polydimethylsiloxane (PDMS), had to find a compromise between their height and fabrication and operation issues. From the design perspective, it was found that a height of 6 μm (Fig. 1A and SI Appendix, Table S1) allowed, conservatively, the unencumbered bacterial motility. Furthermore, preliminary experiments comparing motility in both types of microfluidic structures presented evidence (Movie S1) of the coupling of the impact on both horizontal planes on bacterial motility for those with 4 μm heights. Consequently, 6 μm–tall microfluidic structures were used for all further experiments. A detailed discussion is presented in SI Appendix.

**Spatial distribution of bacteria.** The bacterial species studied presented different motility behaviors with respect to proximity of vertical walls and corners (Fig. 2A–C). First, *V. fischeri*, *V. natriegens*, and *E. coli* moved at small distances from vertical walls. Second, *M. marinus* presented an uneven, broken density near vertical walls, due to the frequent “ping-pong”–like collisions and reflections (Movie S1). Third, *P. putida* presented an even spatial distribution throughout the chamber.

Fig. 1. Microfluidics chip for testing microbial motility. (A) Scanning electron micrographs (SEM) images of bacteria studied with various architectures and dimensions (full details in SI Appendix, Table S1). (B) Graphical projection of the fit of the total bacterial length (body plus flagella) positioned at 45° versus the height of the microfluidic structures for 6 μm and 4 μm heights. (C) Sequential, from left to right, zoom-in images of the experimental device: 1) the bacterial suspension is introduced from the side of the chip attached to the cover slide; 2) the overall architecture of the chip; 3) zoom-in of one lane of experimental structures (sequence of angled channels separated by plazas); 4) detailed image of the experimental structures used in this study (i.e., plazas) and linear channels (top row), angled, and meandered channels (bottom row); and 5) SEM image of a bacterium (here, *E. coli*) in a channel.
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Fig. 2. Motility in plazas with 6 μm high ceilings. (A) Density maps of bacterial locations. Color code (bottom): “min” and “max” represent no and the highest presence of bacteria, respectively. (B) Spatial distribution of bacteria obtained by superimposing and averaging the data from all four quarters of the density maps in A. Color code (bottom): frequency of bacterial presence, with red for the highest and dark green for the minimum probability. (C) Characteristic long2D projections of bacterial trajectories. (D) 3D bacterial trajectories. By rows, from top to bottom, are the following: V. natriegens (average count of bacterial positions in each frame, n = 14/frame); M. marinus (n = 12/frame); P. putida (n = 15/frame); V. fischeri (n = 15/frame); and E. coli (n = 13/frame). Movie S1 presents the bacterial movement in plazas, with representative trajectories (similar to C).

for P. putida and M. marinus, which frequently fluctuated between z-planes (SI Appendix, Figs. S7 and S8). Statistical analysis of the bacterial positions (SI Appendix, Fig. S9) showed that V. natriegens, V. fischeri, and E. coli moved preferentially in a parallel plane to the horizontal walls and that P. putida and M. marinus presented a rather uniform distribution of positions on the vertical axis.

Theoretical classification of bacterial motility behavior. For bacteria that are propelled by a flagellum or flagellar bundle behind the cell, the fluid flow generated by swimming has a dipolar structure: the fluid is pushed backward by the flagellum and pulled forward by the cell body. This flow has been shown to attract swimmers to solid walls, causing them to remain close to the wall for long time periods despite rotational Brownian motion (57). A separate effect of swimming near surfaces is that hydrodynamic interactions between the wall and rotating flagellum and between the wall and counter-rotating cell body, respectively, lead to bacteria swimming in circular orbits when they are close to a wall (58).

Detailed hydrodynamic modeling of monotrichous bacteria showed that the geometrical parameters of the cell (length and width) and of the helical flagellum (length, helical amplitude, and wavelength) determine the motility behavior near a single flat surface (32). Based on this modeling framework, correlated with the experimental observations from the present study, three classes of behavior were observed, depending on the geometry of the bacterium. “Wall accumulators” descend to the walls and exhibit a strong propensity for swimming in the closest vicinity to the wall (with a separation of tens of nanometers between the bacterium and the surface), where steric interactions are likely, thus making difficult the precise prediction of motility behavior even for the simplest monotrichous bacteria. When bacteria swim at distances further than this from the wall but at a nearly constant separation, exhibiting the characteristic circular orbits predicted by simpler analysis, they are classified as “stable swimmers parallel to the wall.” It was observed (26) that dynamical interactions are negligible before collisions with the walls, but once bacteria swim on parallel planes a few micrometers away from surfaces, hydrodynamic forces...
maintain long residence times in this region. Finally, when hydrodynamic interactions result in bacterial movement away from surfaces, they are classified as “wall escapers.” The demarcation between these classes is approximate, due to the inherent stochasticity of bacterial motility.

Two key geometrical parameters determining whether a particular bacterium is an accumulator, escaper, or moving parallel to the wall are 1) the cell body aspect ratio and 2) the length of the flagellum. Higher aspect ratios (more rod-like) and shorter flagella encourage escape from walls (Fig. 3). For geometries at the boundary between parallel motion and escapers, it is possible for a bacterium to exhibit either stable motion close to the wall or escape depending on the angle of approach to the wall. It is useful to first determine the behavior of bacteria near a single wall because this is indicative of motility in more complex environments. For example, simulations showed that parallel–stable swimmers and escapers had different characteristics when placed between parallel walls (35) and in corners of rectangular channels (39). However, the variability of characteristic bacterial dimensions adds to the inherent stochasticity of movement. This in turn makes the demarcation between motility classes approximate. Details of the modeling used in Fig. 3 are given in SI Appendix, and the characteristic dimensions of bacteria are presented in SI Appendix, Table S2.

While these theoretical studies were based on a model with a single, polar flagellum, it was demonstrated that such models accurately reproduce the experimentally observed radius of curvature of near-wall tracks for E. coli, which swim with several flagella (31). Therefore, it is expected that this classification serves as a useful conceptual background for the characterization of motility behavior in relation to a solid surface, even though most of the species in the current study are architecturally more complex than the monotrichous model (here, V. natriegens). Indeed, the propensity to move near surfaces was observed experimentally for several nonmonotrichous bacterial species, for instance (extensively) for E. coli (26, 27, 30, 31, 57, 59), but also for Serratia marcescens (29) and Pseudomonas aeruginosa (60).

Comparison of experimental and theoretically predicted behavior. By comparison with monotrichous model bacteria of equivalent dimensions, M. marinus is predicted to be a wall accumulator, but it is actually near the boundary between accumulators and escapers (Fig. 3). All other species are expected to maintain stable motion parallel to and near the walls (Fig. 3), although variability within populations is sufficient for some individuals to be classified as escapers. There are elements that correlate well with the predicted motility behavior of simple bacteria with that of the more complex geometries studied as well as explanations for the deviations from this general “motility landscape” (Fig. 3):

1) Our experiments showed that M. marinus did not exhibit stable motion parallel to the wall but rather a “ping-pong”–like movement, with abrupt approaches to the walls alternating with equally abrupt breakouts. Recently, a model of the movement of a polar biflagellate bacterium (61), based on M. marinus, showed that such wall escaping (scattering) behavior could occur for certain arrangements of the two flagella. Additionally, it was recently reported that M. marinus swims with one flagellar bundle in front of the cell body and one behind (62), a mode of motility that is fundamentally different from the monotrichous model.

2) The density maps, probability maps, long trajectories, represented as two-dimensional (2D) projections and in 3D (Fig. 2 A–D, respectively) for P. putida and E. coli, showed characteristics of both escapers, more apparent for P. putida, and movement parallel to the wall, more apparent for E. coli. The persistent circular orbits indicate motion close to the horizontal walls, and for E. coli, the long trajectories along the vertical walls also highlight boundary accumulation. In contrast, the long, relatively straight trajectories through the middle of the chamber and frequent transitions between z-planes represent wall escaping behaviors. These seemingly contradictory observations are, in fact, consistent with the variability found in the measured cell shapes and flagella lengths. While the average values for both P. putida and E. coli lie within the movement parallel to the wall regime (Fig. 3), the spread of parameters extends considerably into the wall escape region.

3) Density and probability maps, as well as 2D projections and 3D bacterial trajectories (Fig. 2 A–D, respectively), are consistent with the placement of V. natriegens and V. fisheri deep in the movement parallel to the walls, according to the theoretical predictions in Fig. 3. Both species showed circular trajectories (more prominent in V. natriegens) and high densities around the perimeter of the chamber. Interestingly, V. natriegens was often observed swimming parallel to the vertical walls but at distances of around 3.5 μm from the wall (Fig. 2B) rather than keeping almost in contact with the wall. This type of parallel motion was found in simulations of boundary accumulators in corners of channels (39).
Motility patterns. The longest trajectories of bacterial motility in plazas had characteristics that were the most species specific (Fig. 2C and Movie S1, top row). V. natriegens, E. coli, and V. fischeri presented, to various degrees, two classes of trajectories: 1) movement along the vertical and horizontal walls and, when detached, 2) circular motions, until again attacking to the walls. M. marinus exhibited a “ping-pong”–like motility pattern, generally following relatively straight paths until it approached and scattered off a vertical wall, resulting in a statistically higher density localized near the walls (due to frequent collisions). There was little discernible movement along the vertical or horizontal walls of the plaza, and no complete circular orbits were observed. Two classes of behavior were present in the longest trajectories of P. putida. Some were relatively straight, spanning from one side of the chamber to the other, whereas other trajectories were circular and persisted for many overlapping cycles. Long trajectories around the perimeter of the chamber, as observed for V. natriegens, E. coli, and even V. fischeri, were uncommon for P. putida.

Circular motion. The circular motion of bacteria near surfaces was previously reported for E. coli both at air–liquid (27) and solid–liquid interfaces (58, 63) and for P. putida at solid–liquid interfaces (33, 64). Counterintuitively, despite their very different flagellar arrangements (Fig. 1 and SI Appendix, Figs. S1 and Table S1), circular patterns were also observed here for P. putida, to a lesser extent for E. coli, and for V. natriegens (Fig. 2 C and D). Theoretically, the hydrodynamic interactions between a flat surface and a bacterium swimming on a parallel plane to it are indeed able to explain this curved pattern of trajectories (58, 60).

Motility in Tightly Confining Geometries. Motility in linear channels. Following the experiments in plazas with high and low ceilings and to avoid (to the extent possible) the impact on motility from more than two vertical walls, further experiments used only microfluidic channels with a 6 μm distance between the horizontal planes.

Overall motility characteristics; sinusoidal movement. When laterally confined in wider channels (e.g., 6 to 8 μm), V. natriegens and E. coli showed the strongest propensity for moving along walls (Fig. 4A and SI Appendix, Fig. S10 for 3D trajectories), correlating well with their motility behavior in plazas (Fig. 2A and B) and their movement parallel to the vertical (Fig. 2C and D) and horizontal walls (SI Appendix, Fig. S9).

P. putida exhibited an apparent sinusoidal movement, especially in larger channels (Fig. 4A). A Fast Fourier Transform (FFT) analysis of the trajectories (SI Appendix, Figs. S11 and S12B) indicated that V. natriegens, V. fischeri, and, to a much lesser extent, E. coli also present sinusoidal movement characteristics, with wavelengths increasing roughly proportionally with an increase in channel widths (SI Appendix, Fig. S12B). It was demonstrated (39, 65) that monochirous wall escapers (with this behavior being predicted, partially, for P. putida in Fig. 3) move in distorted helical paths in channels of large rectangular transversal section. This upwards correlation between motility wavelengths and available volume for movement is similar to the larger radii of the circular movement in plazas with higher ceilings than in those with low ceilings (Fig. 2C and SI Appendix, Fig. S5C). M. marinus also exhibited sinusoidal-like behavior, but the FFT analysis showed that this movement is only the result of frequent collisions to, and bouncing from, the walls.

In narrower channels (i.e., 3 to 6 μm), the tighter confinement increasingly forced bacteria to move along the channel axis (except for M. marinus) rather than exhibited their motility behavior observed in open spaces (plazas). Moreover, in tighter (but still larger than the lateral size of the cell) channels, bacterial movement appeared to benefit from both hydrodynamics and steric interaction with the walls, which synergistically push bacteria in the same direction due to the lateral-only confinement of straight channels (66).

Velocities in channels. Analysis of the velocities in straight channels appeared to further substantiate the synergy between hydrodynamics–driven and steric interactions–driven motility mechanisms. Indeed, while M. marinus exhibited a moderate decrease in average velocity with the decrease of the width of the channel, including compared with that in the plazas, due to an increase in collisions with the walls, all other species did not show any notable and systemic velocity variation with channel widths (SI Appendix, Fig. S13). Furthermore, the double histograms of the velocity in channels (Fig. 4B, for rectangular 6 × 6 μm channels; full analysis in SI Appendix, Fig. S14) revealed that V. natriegens and E. coli presented a distinctive bimodal distribution of velocities at the walls, with one velocity higher and one lower than the overall velocity. This bimodal distribution, for the species with the lowest ratios of the cell body and of the flagella (a1/a2 and L/a, respectively, Fig. 3 and SI Appendix, Table S2), could be the result of separate instances of cell adhesion to the wall and movement acceleration due to the steric interaction of flagella with the walls. In this context, it was reported (67) that the interaction between the walls and the flagella of E. coli translates into a “thrusting aid” for those bacteria running smoothly along solid surfaces. It was also reported, for E. coli (30, 68), B. subtilis (69), and S. marcescens (29), that bacteria exhibited higher velocities in narrower channels (which eventually decreases significantly in even narrower channels, due to the severe mechanical constraints applied to the cells), which is supported by the bimodal distribution of velocities observed for E. coli (and V. natriegens) here.

Straight versus U-turn movements. In straight channels, bacterial motility was expected to be increasingly driven by steric interactions, to the detriment of hydrodynamics, with a decrease in channel widths. This increased impact of the steric interactions can explain the species-specific proportion of U-turns (Fig. 4C). First, the species with the lowest ratio of flagellum/length/cell body (i.e., V. natriegens and E. coli) (Fig. 3) had the lowest overall proportion of U-turns, with an apparent decrease of U-turns with the channel width for the larger E. coli (Fig. 4C, Bottom). Conversely, the species clustered at higher characteristic values of L/a and a1/a2 ratios (i.e., P. putida and V. fischeri) (Fig. 3) have a considerably higher proportion of U-turns than V. natriegens and E. coli, and there was even a considerably higher proportion for V. fischeri (Fig. 4D, fourth from the top). Second, M. marinus, with its characteristic frequent collisions and rebounds from the walls, had a low ratio of U-turns, with the notable exception of the 2 μm–wide channels. This unique behavior can be explained by the extreme steric interactions of M. marinus with both walls in channels with 2 μm widths, (i.e., as large as the cell body) (Fig. 4D, second from the top), resulting in the bacterial cell being “pinned” by both vertical walls then “flipped” in the 6 μm–tall vertical plane of the channel, followed by the movement in the opposite direction. Third, P. putida, experiencing intermittent wall contact, exhibited a similar ratio of U-turns as V. natriegens and E. coli. Fourth, V. fischeri, which swim the closest to the wall (Fig. 2B), had the highest ratio of U-turns.

In summary, these results demonstrate that, when a strong and complex coupling exists between the interaction by parallel walls placed at distances similar to the dimensions of bacteria, their motility is primarily governed by the local steric interactions between the walls and the flagella and, in extreme confinement, the cell body. Consequently, the increase in confinement with narrower
Fig. 4. Bacterial motility in linear channels. (A, Left) Density maps of the movement patterns of bacteria in channels with different widths. (A, Right) Bacterial trajectories in 8 μm wide channels, moving from one direction (green) or from an opposite one (red). (B) Double histograms of velocity (y-axis) versus normalized distance from the center of the linear channel (channel wall on the extreme right) for 6 μm × 6 μm channels (the full analysis is presented in SI Appendix, Fig. S14). E. coli and V. natriegens present a specific bimodal distribution of velocities near the wall. (C) Influence of the channel width on the fraction of U-turns. By rows, from top to bottom, are the following: V. natriegens (average count of bacteria each frame, n = 20/frame); M. marinus (n = 10/ frame); P. putida (n = 19/frame); V. fischeri (n = 18/frame); and E. coli (n = 22/frame). (D) Graphical representation of the top view of a bacterium with their average dimensions, in linear channels. The thick and dotted lines represent the minimum and maximum channel widths. Movie S2 presents bacterial movement in channels, with representative trajectories (similar to A, Right).
channels leads to a decrease in hydrodynamics-based propulsion, and the dilution, or outright disappearance of the classes of motility behavior observed in open spaces.

**Motility in channels with angled exits.** In the structures with angled exits (Fig. 1C, lower right of the fourth image from the left), all bacterial species had a large preference for moving in straight trajectories along the middle axis of the channel, as qualitatively suggested by the density maps (Fig. 5A), by representative trajectories (Fig. 5B and Movie S3), and by representative bacterial 3D trajectories (SI Appendix, Fig. S15). Even for the smallest exit angle (i.e., 30°), the probability of movement in a straight line instead of exiting laterally (estimated as the ratio between bacterial movement straight and the total number that arrived at that intersection) ranges from 72% (for *P. putida*) to 58% (for *M. marinus*). While the general trend for all bacteria was that the exiting probability decreased with increasing exit angle, there were some species-specific details (Fig. 5C). First, *V. natriegens*, *E. coli*, and *P. putida* had a clearly decreasing exiting probability with an increase in exit angle, while for *V. fischeri* this trend was less visible, and *M. marinus* exhibited a rather indifferent relationship between exit probabilities and exit angles, following an abrupt drop at angles higher than 30°. Second, all species other than *M. marinus* had a relatively higher exiting probability at 90° angles.

This species-specific motility behavior in angled channels appeared to be the result of bacterial movement being driven by both local hydrodynamics and by steric interactions with the walls. First, the decrease of exit probabilities with exit angles for all species, but especially for *V. natriegens* and *E. coli*, resembles the lower frequency of turning by large angles in open spaces (SI Appendix, Fig. S3). The deflection angles in open spaces (SI Appendix, Figs. S3 and S4) are near-instantaneous measurements, and while longer integration times would lead to larger apparent values, this could also incorporate other sudden changes of direction, thus obscuring the inherent propensity of bacteria for sideways movement. With this qualification, it is reasonable to expect a connection between the propensity to escape laterally at set angles in angled channels (Fig. 5C) and the deflection angles in open spaces. However, this similarity had notable limitations (e.g., all species studied had negligible probabilities of deflection angles at much lower angles than those for bacteria in the angle range of 10° to 30°). Second, the wide spread of deflection angles in plazas (SI Appendix, Figs. S3 and S16) for *P. putida* could justify its relatively wide spread of exit probabilities in angled channels, *E. coli*, which had a narrow distribution of deflection angles, had a considerably larger and wider distribution of exit probabilities in angled channels than *P. putida*. Similarly, while both *V. natriegens* and *V. fischeri* exhibited a monotone decrease of frequency with increasing deflection angles in plazas (SI Appendix, Figs. S3 and S16), this behavior translated into a monotone decrease of exit probabilities in angled channels only for the former, whereas the latter did not show any obvious correlation between exit probabilities and respective escape angles. Finally, *M. marinus* had a monotone decrease of frequency with increasing deflection angles (after 10°) but an approximately flat relationship between the exit probabilities and escape angles (after 30°).

These observations suggest that, in addition to species-specific hydrodynamics-driven spread of deflection angles in open spaces (plazas), another mechanism was also responsible for determining the exit probabilities in angled channels. Indeed, the species that exhibited a notable departure from the expected extrapolation of behavior in open spaces is also the species whose dimensions exceed the clearance in the angled channels (i.e., *E. coli* and *V. fischeri*) (Fig. 5D). Conversely, the species whose dimensions did not surpass the clearance in the angled channels (i.e., *V. natriegens* and *P. putida*) are also those which exhibited a reasonable extrapolation of deflection angles in open spaces to a monotonical decrease of exit probabilities with escape angles. The frequent collisions and bouncing of *M. marinus* had the effect of leveling the exit probabilities regardless of the escape angles (except for 30°, for which there is enough turning space and therefore a higher exit probability, Fig. 5D).

It must be also noted that the confinement at the intersection between central and lateral channels did not fully correlate with the respective exit angle. For instance, the 150° exit offered the largest volume available for movement at the intersection between the axial and lateral channels (highlighted in SI Appendix, Fig. S2), thus making the comparison with U-turns (at 180°) in tight linear channels, with no variation of widths, inconsistent. Finally, the relatively higher escape probabilities for 90° angles could be the result of smaller free volume at the intersection of the axial channel, with steric interactions biasing bacteria toward lateral exits.

In conclusion, bacterial motility studies in angled channels revealed that when the level of confinement is low, due to the large volume at cross-intersection in relation to smaller bacterial sizes, the movement is mostly driven by hydrodynamics, as an extension of the behavior observed in open spaces. Conversely, when the confinement is tight, due to larger bacterial sizes, the local steric interactions between flagella and the walls contribute substantially to the motility behavior.

**Motility in meandered channels.** The trapping of bacteria in purposefully designed microfluidics structures is of special interest to various applications [e.g., single-cell genomics (70) and accelerated evolution (71)], and therefore, the responsible mechanisms were studied (72, 73).

The meandered system comprised three channels, each with a different gap between the edge of the “teeth” (i.e., 5 μm [left], 10 μm [middle], and 15 μm [right]) (Fig. 1C, lower right, fourth image from left). The tightly confined, 5 μm–wide meandered channels made the motility of all species more complex (Fig. 6 and SI Appendix, Fig. S17 and Movie S4). The elastic-like collisions of *M. marinus* resulted in frequent trapings and, consequently, a considerably lower overall “success rate” (defined as the ratio of bacterial entries versus exits, at steady state) than the rest of the bacterial species (Fig. 6C). In addition, the 90°-angled corners appeared to operate as traps for *E. coli* and to a lesser extent for *V. natriegens* (bright spots in the density maps in Fig. 6D; the higher retention time for *E. coli*, SI Appendix, Tables S3 and S4). Intuitively, all species exhibited to a certain extent for *M. marinus*, *P. putida* and to a lesser extent for *V. natriegens* (SI Appendix, Figs. S3 and S4), the trapping effect of the 90°-angled corners more effective, to a near-total extent for *V. natriegens* and *E. coli*, and to a lower relative extent for *V. fischeri*, *M. marinus*, and *P. putida*. As expected, the highest passage “success rate” was provided by the meandered channels with the largest distance between walls (Fig. 6C). Again, *M. marinus* and *P. putida* executed more zigzagged trajectories than *V. natriegens*, *V. fischeri*, and *E. coli* (Fig. 6B).

This complex and species-specific behavior can be explained in view of previous findings as follows. In channels with large gaps between comb teeth, all bacteria can negotiate the passage, their movement being driven mostly by hydrodynamics, with only occasional interference of the local steric interactions between the flagella and the walls. Conversely, in channels with tight confinement, bacteria are also capable of successfully negotiating the channels, this time “channeled” by the local steric interaction between flagella and the 90°-angled walls. Finally, in the channels in the mesoscale region (i.e., 10 μm distance between the “comb teeth”), the mechanism based on hydrodynamics and that based on the local steric interaction do not operate synergistically,
Fig. 5. Bacterial motility in channels with angled exits. (A) Density maps of the movement patterns of bacteria in straight parallel channels, connected by side channels at angles ranging from 0° to 150°. All channels are nominally 4 μm wide. (B) Bacterial trajectories, either from top (green) or from the opposite direction (red). (C) Frequencies of bacteria moving at different exit angles. By rows, from top to bottom, are the following: *V. natriegens* (average count of bacteria each frame, \( n = 10 \) frame); *M. marinus* \( (n = 8 \) frame); *P. putida* \( (n = 10 \) frame); *V. fischeri* \( (n = 13 \) frame); and *E. coli* \( (n = 11 \) frame). (D) Graphical representation of the top view of bacterium with average dimensions in the angled channels (few representative angles). The areas in light brown represent spaces that exceed the dimensions of the respective bacteria in the respective position. Movie S3 presents bacterial movement in angled networks and representative trajectories (similar to A, Right).
increasing the chaotic character of motility and making the overall forward advancement difficult. This is particularly obvious for *V. natriegens* and *E. coli*, which swim parallel to walls (Fig. 3) at a ~2 μm distance from the walls (Fig. 2B), thus leading to frequent U-turns. Additionally, these two species are those with the lowest ratios of the cell body and of the flagella (Fig. 3).
and SI Appendix, Table S2). The more compact architectures of *V. natriegens* and *E. coli* could explain the near-perfect trapping by frequent circular movements in very confined spaces leading to long retention times. Conversely, but for different reasons, *V. fischeri* (a species swimming the closest to the walls, Fig. 2B) and *P. putida* (a species with opportunistic distribution in free volumes) can avoid, to a larger extent than *V. natriegens* and *E. coli*, being trapped in the meandered channels. Tellingly, these two species are also those with the highest ratios of the cell body and of the whole bacterium (Fig. 3 and SI Appendix, Table S2). This dichotomy of behavior for species swimming parallel to the walls suggests that the steric interactions-driven movement in tight confinement is also modulated by bacterial shape and not only by size (presented schematically in Fig. 6D). Indeed, *V. natriegens* and *E. coli* are both very effectively trapped in mesoscale-sized meandered channels, and while *P. putida*, a much shorter species (SI Appendix, Table S2), appeared to have some success, *V. fischeri*, the largest of the species swimming parallel to the walls, had the best success rate. Finally, *M. marinus* was also found to exit mesoscale meandered channels more, but its frequent collision-and-rebound on the walls led to slightly lower trapping efficiencies.

To elucidate whether the trapping effect is permanent or transient, the average duration for successfully traversing the meandered channels was quantified (SI Appendix, Table S3). Within the experimental time window (4 to 5 min), *V. natriegens* and *E. coli* were unable to successfully traverse the middle-meandered channels. Although *M. marinus* had a shorter retention time due to its high velocity, the distance that it needed to travel in order to be able to exit the meandered channel was longer. Representative 3D trajectories in meandered channels are presented in SI Appendix, Fig. S17. The color-coded trajectories for U-turns, successful passages, and trapped bacteria are presented in SI Appendix, Fig. S18.

To conclude, in complex geometries, such as meandered channels, hydrodynamics-driven motility is prevalent in wider channels, and the local steric interactions–based mechanism governs bacterial motility in narrow channels. However, in the mesoscale region, these two mechanisms do not act in synergy, resulting in trapping bacteria, with high efficiency for species swimming parallel to the walls, finely modulated by their characteristic shape ratios.

**Perspectives and Future Work.** The present study, in which we studied a wide range of bacterial motility behavior, provides insights in several areas of applications, as well as suggesting further research. *Fundamentals of bacterial motility mechanics in microenvironments.* It was previously shown that a fundamental understanding of the mechanics of the movement of monoflagellated (32, 33, 39) and even biflagellated (61) bacteria in simple geometries, such as the proximity to a surface, can accurately predict motility patterns of bacteria. However, the current study, which described motility patterns of more complex bacterial architectures and in more complex geometries, revealed the limits of this understanding, which would be critical for designing microdevices manipulating bacteria for biosensing, drug delivery, cell sorting, or bioocomputation. Further theoretical directions suggested by our study, perhaps coupled with long-term monitoring (74), include analyzing the impact of population variation on cell behavior, investigating the extent to which more complicated bacterial geometries and flagellar arrangements can be represented by more advanced mechanical models, and the need to conduct systematic validation studies. Studies of this type, using artificial microfluidics systems mimicking their natural counterparts, recently carried out for bacteria (75) and fungi (76) or for specifically investigating stochastic processes in bacteria (77), are motivated by the abundance of microbial habitats comprising linear and meandered channels and spaces with different angled turns (SI Appendix, Fig. S20).

**Motility of magnetic bacteria in biological networks.** Chemically or magnetically guided self-propelled bacteria were used for non-systemic delivery of drugs and cargoes in tumor therapy (78–81). The targeted physiological regions (e.g., deep enteric tissues, hypoxic tumors, tissue granules, and arterioles) (78, 82) are essentially impenetrable to probing devices, but they can be accessed, in principle, by robust bacteria operating as autonomous microrobots moving in the natural microfluidic vascular system (78). The description of bacterial motility, in particular that of *M. marinus*, in PDMS microfluidic channels mimicking the microvascular system surrounding the tumor (e.g., micrometer-range sizes and relevant mechanical elasticity) can lead to the optimization of the operation of these microrobots outside clinical settings, which are expensive to operate and unable to provide reproducible observations at the microscale and in real time.

**Bacterial cell sorting.** The efficient characterization, sorting, or selection of individual bacterial cells in small volumes are achieved in various microfluidics-based applications, such as those derived from the classical flow cytometry (83–85) to the more recent single-cell analysis (70). In fact, microfluidic devices have been increasingly used for assessing bacterial chemotaxis (86–89), motility (29, 30, 90, 91), and for bacterial cell sorting (46, 72, 91–93), and our results can offer insights for the design of these devices. For instance, the characterization of bacteria as wall accumulators, or wall escapers, can suggest entirely different geometries for microfluidic structures for bacterial cell sorting. Similarly, microfluidic channels can be designed so as to increase retention time (e.g., by having helical profiles) or to amplify the differences in mechanical responses to flow in microfluidics-based flow cytometry.

**Network-based bio computation.** Microfluidics-based approaches to computation of problems intractable to electronic computers have been proposed for clique problem (94) and subset sum problem (95). These biological computers require the independent exploration of microfluidic networks encoding a mathematical problem by autonomous agents such as beads (94), cytoskeletal filaments (95), or microorganisms (96). The precision of the microfluidics-based computation is determined by the capacity of biological agents, such as bacteria, to faithfully follow the movement rules embedded in the logic junctions they visit (97). Consequently, the selection of bacterial candidates and the designs of computational microfluidic networks will require the removal or at least minimization of errors, such as U-turns in narrow channels, as well as optimization of the angles of logic gates channels.

**Conclusion**

We here provided a comprehensive account of the motility of individual bacterial cells, belonging to five species with considerably varied dimensions and morphologies, in microfluidic networks and with various levels of confinement and complexity. For less-confining geometries, such as facing one limiting wall, the motility behavior of the five species studied can be assimilated, with qualifications, to that of monotrichous bacteria with similar dimensions. However, when increasing confinement complexity, as for instance in straight channels with various widths, in networks with exits at various angles, and meandered channels, the classification as swimming parallel to the walls for *V. natriegens*, *E. coli*, *V. fischeri*, and *P. putida* and as escapers, partially, for *E. coli*, *P. putida*, and *M. marinus* is increasingly inaccurate, as a result of the increase of the impact of local steric interaction of species-specific morphology with the tightly confining geometry. The study can be also used as a methodological template for the optimization of the design of microfluidic devices with specific functions (e.g., motility-based cell selection for single-cell genomic screening, detection of rare cells, bacterial entrapment devices for diagnostics, or bioocomputation).

**Materials and Methods**

All experimental, modeling, and simulation data analysis protocols are presented in SI Appendix.

**Data Availability.** All study data are included in the article and/or supporting information.
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