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# Dynamic mode structure of active turbulence

Richard J. Henshaw,<sup>1</sup> Olivia G. Martin,<sup>1,2</sup> and Jeffrey S. Guasto<sup>1,\*</sup>

<sup>1</sup>Department of Mechanical Engineering, Tufts University, Medford, MA 02155

<sup>2</sup>Department of Mechanical Engineering, Stanford University, Stanford, CA 94305

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Dense suspensions of swimming bacteria exhibit chaotic flow patterns that promote the mixing and transport of resources and signalling chemicals within cell colonies. While the importance of active turbulence is widely recognized, the structure and dynamics of the resulting collective flows are the subject of intense investigation. Here, we combine microfluidic experiments with proper orthogonal decomposition (POD) analysis to quantify the dynamical flow structure of this model active matter system under a variety of conditions. In isotropic quasi-2D turbulence, the modal analysis encompasses the most energetic spatio-temporal flow structures across a range of suspension activity levels and benchmarks the potential for low-dimensional order representations of active turbulence. In confined geometries, POD analysis illustrates the role of boundary interactions for the transition to bacterial turbulence, and it quantifies the evolution of coherent active structures in externally applied flows. Beyond establishing the physical flow structures underpinning the complex dynamics of bacterial turbulence, the low-dimensional representation afforded by this modal analysis will facilitate data-driven modelling of active turbulence.

## I. INTRODUCTION

The collective motion of concentrated suspensions of swimming bacteria and self-propelled actin filaments are typical of a broad class of active systems, ranging from flocking birds and schooling fish down to micron-sized synthetic active colloids [1–5]. Such active suspensions regulate fundamental ecological and biological processes for example in the gut microbiome of the human gastrointestinal tract [6], as well as the mitotic spindle structure during cell division [7]. At high concentrations, swimming bacteria form emergent jet-like flow structures and eddies [8–12], which are correlated over length scales that are one order of magnitude larger than the individual cells and enhance the colony mobility [13]. These chaotic flow patterns – termed as active turbulence [14] – are reminiscent of classical, inertial turbulence [15] and exhibit a range of novel physical properties, such as: enhanced material transport [13, 16], boundaryinduced pattern formation [17, 18], and reduced viscosity and super-fluid-like behavior [19, 20]. Collective flows are highly dependent on the physical properties of their constituent agents, where variations in cell size, swimming speed, suspension density, nutrient availability, media/substrate, and viscosity can alter the nature of the cell motion [11, 21–24]. Apart from traditional, bulk statistical measures and velocity spectra [3, 4, 11, 12], relatively little experimental progress has been made in understanding the underlying flow structure of active turbulence, especially in the presence of boundaries and ambient flows. A host of theoretical and computational models have been proposed to explain these phenomena [4, 25–28], as well as the onset of active turbulence under confinement [29], and recent work using machine learning approaches to uncover the equations of motion for active turbulence are promising [30]. However, the vast experimental parameter space and necessity for big data sets to characterize these chaotic systems complicate data-driven discovery through simulations. Thus, establishing new approaches to characterize the dynamic structure of active turbulence and provide compact reduced-order representations are essential, for example to understand the movement ecology of collective microbial motion [31] and to design and engineer new classes of active materials [32].

Inspired by approaches from inertial turbulence, we apply proper orthogonal decomposition (POD) to analyse the spatio-temporal dynamics of self-generated bacterial turbulence flows. POD – also known as PCA or the Karhunen-Loève procedure [33–35] – computes an orthogonal basis set that optimally represents the fluid velocity field, an approach that has been instrumental for example in the development of reduced-order flow models [34, 36, 37]. In POD, a velocity field  $\mathbf{u}(\mathbf{x},t)$  at N discrete times is represented by a linear combination of N orthonormal spatial modes,  $\phi_n(\mathbf{x})$ , as

$$\mathbf{u}(\mathbf{x},t) = \sum_{n=1}^{N} A_n(t)\phi_n(\mathbf{x}), \tag{1}$$

where  $A_n(t) = a_n(t)\sqrt{\lambda_n}$ ,  $a_n(t)$  is a normalized temporal coefficient, and  $\lambda_n$  is the spatio-temporal average kinetic energy for a given mode [37]. This process is achieved through singular value decomposition (SVD; see Appendix. A 2) [38], which ranks the spatial modes,  $\phi_n(\mathbf{x})$ , by decreasing energy,  $\lambda_n$ . Despite the striking similarities between inertial and active turbulence, POD techniques have seen limited applications in soft matter

<sup>\*</sup> To whom correspondence should be addressed.
Email: Jeffrey.Guasto@tufts.edu
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analyzed the data; and R.J.H., O.G.M., and J.S.G. wrote the
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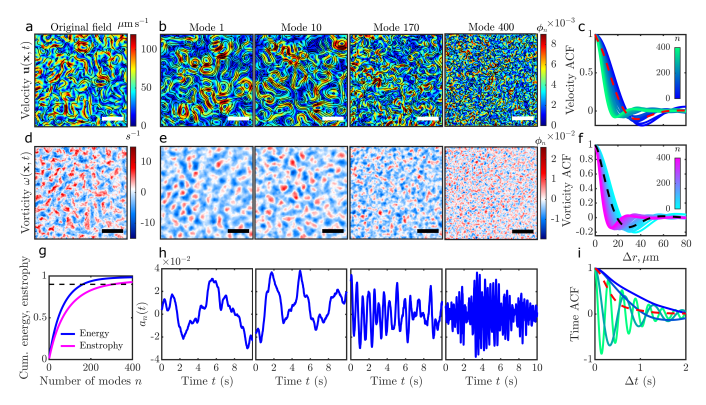


FIG. 1. POD reveals dominant spatio-temporal structures of bacterial turbulence. (a) Instantaneous measured velocity field,  $\mathbf{u}(\mathbf{x},t)$ , obtained from the collective motion of a dense suspension of swimming bacteria. (b) Sample velocity field POD modes,  $\phi_n(\mathbf{x})$ , organized by decreasing kinetic energy. (c) Spatial autocorrelation function (ACF) of the first 400 POD velocity field modes compared to the original flow field (dashed red), showing a decrease in characteristic length scale as the mode number increases. (d) Instantaneous vorticity field,  $\omega = \nabla \times \mathbf{u}$ , corresponding to (a). (e) Sample vorticity field POD modes organized by decreasing enstrophy. (f) ACF of POD vorticity field modes compared to the original vorticity field (dashed black). (g) Normalized cumulative mode energy (blue) and enstrophy (magenta),  $C(n) = \sum_{j=1}^{n} \lambda_j / \sum_{j=1}^{N} \lambda_j$ , for the first n modes. 162 and 305 modes are required to capture 90% (black dashed) of the cumulative energy and enstrophy, respectively. (h) Temporal amplitudes,  $a_n(t)$ , of the velocity field modes in (b). (i) Time ACF for the mode amplitudes shown in (h) compared to the velocity field time ACF (dashed red). Colors correspond to (c). Scale bars are 100  $\mu$ m.

and low Reynolds number flows [39–42].

In this work, we use microfluidic experiments to measure the chaotic self-generated flow fields of bacterial turbulence under a variety of conditions, which we then decompose using modal analysis to quantify their flow structure and dynamics. Bacterial turbulence is firstly investigated far from lateral boundaries (quasi-2D), where we determine the effects of varying cell activity [4, 12] on the spatio-temporal mode structure and benchmark the efficacy of data compression for active turbulence. Subsequently, we use POD to elucidate the role of boundary interactions in the transition to bacterial turbulence in confined geometries [18, 29], and to quantify the evolution of coherent active structures in externally applied flows [43]. Not only does POD analysis unveil the constituent flow structures underpinning the chaotic bacterial motion in each of these physical systems, but this framework also provides an accurate and efficient representation of active turbulence with remarkably few spatial modes.

# II. RESULTS AND DISCUSSION

#### A. Modal analysis of quasi-2D bacterial turbulence

In unbounded domains, concentrated suspensions of motile bacteria exhibit complex spatio-temporal flow patterns. Dense suspensions of motile B. subtilis bacteria were loaded into microfluidic chambers (26  $\mu$ m deep) with widely separated side walls (2 mm). The active suspensions were imaged in the center of the channel, far from the boundaries, using video microscopy for  $\approx 1 \text{ min}$ (N = 6300 frames). Time-dependent velocity fields,  $\mathbf{u}(\mathbf{x},t)$ , characterizing the bacterial motion (Fig 1a) were measured using particle image velocimetry (PIV; see Appendices. A 1,A 3) [18]. The resulting chaotic flows exhibit vortex structures ( $\approx 40 \,\mu\mathrm{m}$ ) and mean flow speeds  $(\langle |\mathbf{u}| \rangle \approx 50 \,\mu\mathrm{m\,s^{-1}})$  that are significantly larger than the respective length  $(4 \, \mu \text{m})$  and swimming speed of individual cells ( $\approx 15 \,\mu\mathrm{m\,s^{-1}}$ ), consistent with previous observations [11, 12].

Energy-ranked velocity field modes from POD anal-

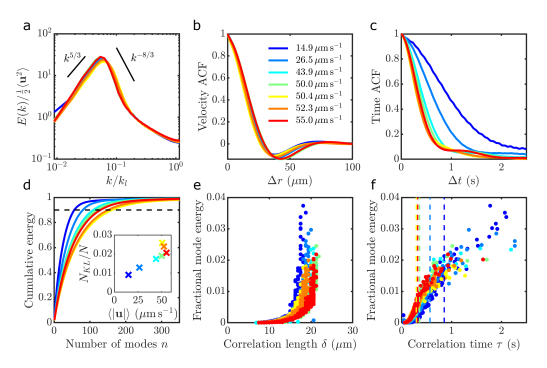


FIG. 2. Variation in bacterial turbulence structure with swimming activity. (a) Normalized kinetic energy spectra as a function of normalized wave number exhibit self-similarity and consistent power-law behavior [4] across a wide range of bacterial activity levels,  $\langle |\mathbf{u}| \rangle$ . The characteristic wave number is  $k_l = 2\pi/l$ , where  $l = 4\,\mu\mathrm{m}$  is the typical cell length. (b) Spatial velocity ACF remains constant across various cell activities. (c) Temporal velocity ACF broadens significantly as bacterial swimming activity decreases. (d) Cumulative kinetic energy contained within POD modes illustrates the relative increase in energy for low order modes with decreasing cell activity. Black dashed line corresponds to the 90% energy KL dimension. See also Appendix B Fig. 6. (e) Fractional mode energy  $\lambda_n/\Sigma\lambda_n$  as a function of the correlation length,  $\delta$ , of individual POD modes ( $n \leq 400$ ; 1/e decay of velocity ACF; Fig. 1c) show that , generally, the most energetic modes typically correspond to the largest spatial scales, which remain constant across bacterial activities. (f) Fractional mode energy  $\lambda_n/\Sigma\lambda_n$  as a function of the correlation time,  $\tau$ , of individual POD mode amplitudes ( $n \leq 400$ ; 1/e decay of time ACF; Fig. 1i). The most energetic modes have the largest  $\tau$ , which increase with decreasing activity. Vertical dashed lines indicate  $\tau$  for time ACFs from (c).

ysis reveal the hierarchy of flow structures underlying bacterial turbulence, whereby the highest energy modes are generally associated with the largest structures. The dominant flow structures are contained within the first several modes (Fig. 1b, n = 1, 10), followed by a transition to smaller structures (Fig. 1b, n = 170) and finally higher-order modes, which capture minor perturbations to the flow field (Fig. 1b, n = 400) as well as measurement noise. The modes appear similar across different realizations of the experiment (Appendix B Fig. 6a-b), but the precise mode structure varies due to the random nature of bacterial turbulence. The autocorrelation function (ACF; Appendix A3) of the spatial modes (Fig. 1c) illustrates an ordering generally from large to small flow structures with decreasing mode energy (increasing n). Furthermore, the highest energy modes encompass the dominant spatial scales of the original velocity field (Fig. 1c, dashed red).

Analogous to the velocity field, a POD analysis of the vorticity field (Fig 1D),  $\omega = \nabla \times \mathbf{u}$ , provides an enstrophy-ranked set of vorticity modes (Fig. 1e). The vorticity mode ACFs clearly illustrate a similar refinement of the modal vortex size (Fig. 1f) with decreas-

ing enstrophy. Considering their cumulative information content (Fig. 1g), from the full set of 6,300 modes only  $N_{KL} = 162 \text{ modes } (2.5\%) \text{ and } 305 \text{ modes } (4.8\%) \text{ for the}$ energy and enstrophy, respectively, are required to form the Karhunen-Loève (KL) dimension [44], which captures 90% of the total information content. The larger number of modes for the latter suggests that enstrophy is more equally distributed across flow structures compared to the kinetic energy, although this effect may be exacerbated by numerical gradients used to calculate the enstrophy. Importantly, the dynamics of bacterial turbulence in this isotropic, quasi-2D system are accurately captured via a modal representation comprising less than 5% of the size of the original flow field data set. This extremely efficient data compression compares very well to existing high-Reynolds number systems, where 10-60% of the modes are typically used for flow field reconstruction with lower cumulative energy than those demonstrated here [45-49].

Beyond the spatial structure, modal analysis provides insight into the temporal dynamics of the chaotic flow fields. The temporal coefficients,  $a_n(t)$ , describe the instantaneous amplitude (Fig. 1h) for each spatial mode

(Fig. 1b) and fluctuate strongly in time. The ACFs of the temporal coefficients (Fig. 1i) show that the large, high energy flow structures associated with low mode numbers are strongly correlated. While lower energy modes exhibit faster initial decay rates, the envelope of their rapidly fluctuating ACF is comparable to correlation strengths of the high energy modes and the original velocity field (Fig. 1i, red dashed). Thus, not only does POD provide a rigorous framework for concise low-order representations of this model active matter system, but it also establishes new tools to gain deeper insight into their complex structure and dynamics under a variety of physical conditions.

# B. Effect of bacterial activity on mode structure and energy distribution

In addition to cell size, concentration, and other variables [11, 21–24], suspension activity is a critical factor in determining the dynamics of active turbulence [50]. Bacterial activity is quantified by the mean flow speed of the suspension,  $\langle |\mathbf{u}| \rangle$ , and varied here in a controlled manner by exploiting the aerobic properties of the bacteria [11]. Microfluidic chambers were loaded with cells as described previously, and the bacterial suspensions were imaged periodically over  $\approx 30 \, \text{min}$ . As oxygen was slowly depleted, the cell activity decreased from  $> 50 \,\mu\mathrm{m}\,\mathrm{s}^{-1}$ to approximately the single-cell swimming speed. After this point, the cell activity decays rapidly to zero and vortices cease to form, hence these extremely slow fields are not shown. For all activities, the kinetic energy spectra, E(k), as a function of the wave number, k, collapse (Fig. 2a) to previously reported power-law behaviors [4], illustrating the robustness of these results.

Beyond classical measures of collective motion, modal analysis reveals rich changes in the structure and complexity of bacterial turbulence with cell activity, including a diminishing number of dominant modes as the cell swimming speed decreases. The characteristic length scale of the turbulent velocity fluctuations remains approximately constant across all activities (velocity correlation length range of  $\delta \approx 15.4 - 18.5 \,\mu\text{m}$ ) [11, 12]. as shown by the spatial velocity ACF (Fig. 2b). Consequently, the slowdown results in a broadening of the temporal ACF (Fig. 2c; see also Fig. 5). As the cell activity decreases, the mode structures have a qualitatively similar appearance (see Appendix B; Fig. 6). However, the energy distribution shifts to lower order modes (Fig. 2d). For example, the lowest activity examined  $(\langle |\mathbf{u}| \rangle = 14.9 \,\mu\mathrm{m}\,\mathrm{s}^{-1})$  required 65% fewer modes to capture 90% of the kinetic energy relative to the most active suspensions (Fig. 2d, inset). This feature appears consistent with kinetic models [28], which predict that the longest wavelengths are the most unstable, and have a growth rate that scales linearly with activity. Moreover, analysis of the mode correlation lengths,  $\delta$ , reveals that dominant spatial modes emerge (Fig. 2e), which are

characterized by both high energy and long correlation lengths and reflect  $\delta$  for the original flow field (Fig. 2b).

While  $\delta$  exhibits relatively minor variations across different activities, the correlation times of the mode amplitudes,  $\tau$ , increase strongly with decreasing cell activity (Fig. 2f). This analysis shows that the most energetic velocity field modes are also responsible for the increase in  $\tau$ . POD analysis enriches observations afforded by traditional metrics (Fig. 2a-c) by identifying the associated flow structures responsible for the observed behaviors and extracts the individual mode dynamics (Fig. 2d-f). Ultimately, insights from the POD analysis are limited due to the isotropic and random nature of bacterial turbulence in this cases. However, as we illustrate below, this isotropic, latterly unbounded system benchmarks the upper bound for the spatial mode complexity, which is dramatically altered by confinement.

## C. Edge modes illustrate the role of confinement in the transition to active turbulence

Boundaries play a fundamental role in regulating active turbulence [50], whereby confinement has been shown to select for vortical flow structures [17] and lead to the formation of self-organized flow patterns [51]. Similar to inertial turbulence, bacterial suspensions [18, 52, 53] and active nematics [29, 52, 54, 55] confined in a channel exhibit a striking transition from stable flow to chaotic dynamics as the separation between lateral boundaries increases. By replicating recent experimental results (Fig. 3a-h) [18], we extend our POD analysis to quantify the effects of confinement on bacterial turbulence. Briefly, a dense suspension of B. subtilis was confined laterally in a series of microfluidic "racetrack" channels (Appendix A4) with varying width, W (Fig. 3a). Strong confinements – where W is approximately less than the quasi-2D vortex size (Fig. 1) - rectify cell activity into a unidirectional flow (Fig. 3c), as shown by the stream-wise averaged flow profile (F(y); Fig. 3f) and normalized net flow rate ( $\psi$ ; Fig. 3b; Appendix A4). As W increases, vortex structures emerge (Fig. 3d) before giving way to chaotic dynamics (Fig. 3e), when W is significantly larger than the quasi-2D vortex size. The formation of vortexes and the subsequent chaotic flow are typified by sinusoidal velocity profiles (Fig. 3g-h) with strong wall velocities and a rapid decrease in  $\psi$  (Fig. 3b), both of which are in very good qualitative agreement with previous work [18]. Reduced order POD representations accurately and compactly capture the emergent active velocity field behaviors across varying channel widths (Fig. 3f-h) throughout this flow transition.

The topologies of the spatial modes,  $\phi_n(\mathbf{x})$ , reveal the fundamental flow structures underpinning the effects of confinement on suspension dynamics (Fig. 3i). For the strongest confinement examined, the highest energy mode encapsulates the nearly-steady, stable flow (Fig. 3i,  $W = 30 \,\mu\text{m}$ ), representing > 70% of the total kinetic en-

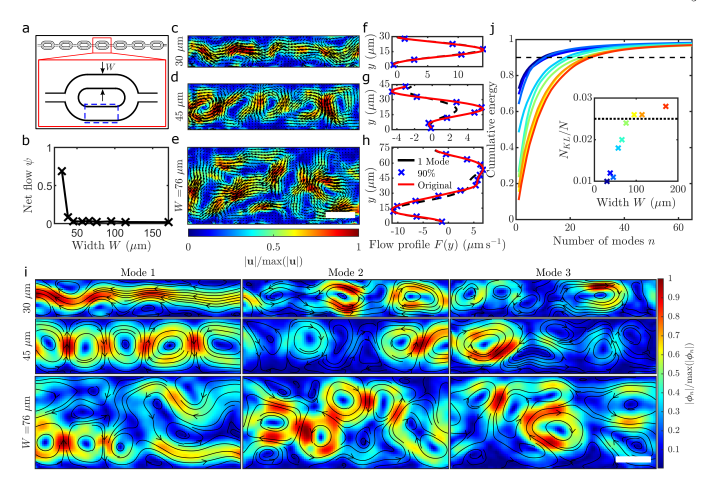


FIG. 3. POD unveils the role of edge modes in the transition to active turbulence under confinement. (a) Schematic of microfluidic racetrack channels of varying widths  $(30 \, \mu \text{m} \leq W \leq 171 \, \mu \text{m})$  to study effects of confinement on bacterial turbulence [18]. Straight observation regions (dashed blue) were fixed at 150  $\mu$ m long. (b) Normalized net flow rate,  $\psi$ , illustrates the transition from unidirectional flow for small W to turbulent flow at large W, which agrees well with previous studies [18]. (c-e) Normalized instantaneous velocity fields for various channel widths. Scale bar, 25  $\mu$ m. (f-h) Stream-wise averaged flow profiles, F(y), for original flow fields (red; corresponding to (c-e)) compared to a 90% POD reconstruction (blue crosses) and single mode reconstruction (black dashed) using modes from (i). (i) The three highest energy (normalized) POD modes for various W. Scale bar, 25  $\mu$ m. (j) Cumulative POD mode energy increases with decreasing channel width. Black dashed line indicates the 90% energy KL dimension. (Inset)  $N_{KL}$  as a fraction of the total number of modes, N, plateaus as the channel approaches quasi-2D conditions (black dotted).

ergy (Fig. 3j). At intermediate channel widths, mode 1 highlights the emergent vortex chain, where higher-order modes have similar topologies that describe stream-wise fluctuations of the vortexes (Fig. 3i,  $W = 45 \,\mu\text{m}$ ). Importantly, in weak confinement prior to transitioning to isotropic, quasi-2D turbulence, the most energetic mode reveals coherent, near-wall activity (mode 1; Fig. 3i,  $W = 76 \,\mu\text{m}$ , see also Appendix B Fig. 7). Such edge modes are consistent with both previous experimental observations of confined bacterial suspensions [18] and the pinning and stabilization of defects to walls in active nematics simulations [50], which are thought to regulate the transition to active turbulence. In contrast, lower energy modes (modes 2-3) describe chaotic fluctuations in the center of the channel away from boundaries, akin to quasi-2D turbulence modes (Fig. 1b). These highly energetic modes (Fig. 3i) also bear striking resemblance to the unstable eigenmodes predicted by kinetic models [56], suggesting that POD and related techniques [57] may provide direct insights into the stability of experimentally measured active suspensions. The kinetic energy distribution across modes is a strong indicator of the onset of bacterial turbulence under confinement (Fig. 3j), which is exemplified by the rapid drop of the kinetic energy in mode 1 from 70% to 10% as W increases from  $30 \,\mu\mathrm{m}$  to  $171 \,\mu\mathrm{m}$ . Moreover, the number of modes,  $N_{KL}$ , to capture 90% of the system energy exhibits a striking increase with W (Fig. 3j, inset) that plateaus at  $W = 95 \,\mu\mathrm{m}$  (or  $\approx 2.5$  vortex lengths in the quasi-2D). The fractional  $N_{KL}$  for large W is consistent with the benchmark established in the previous quasi-2D turbulence results (i.e. 2.5% of the total modes; Fig. 3j, inset, dashed; see also Fig. 1) and indicates that the system has reached the fully chaotic regime.

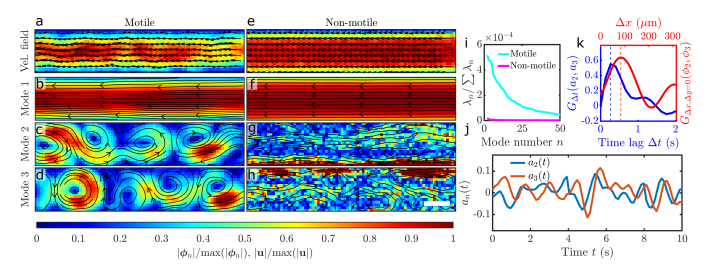


FIG. 4. POD modes capture intermittent bacterial turbulence in externally applied flows. (a) Instantaneous normalized velocity field of a dense, active bacterial suspension flowed through a rectangular microchannel exhibits strong flow intermittency. Velocity field data obtained from previously published work [43]. (b-d) Three most energetic (normalized) POD modes for the motile suspension in (a). Mode 1 resembles a Poiseuille-like background flow, whereas modes 2 and 3 are anti-phase vortexes that capture travelling active flow structures. (e) Instantaneous normalized velocity field of a non-motile cell suspension under the same conditions as (a), which shows a near-parabolic flow profile. (f-h) Three most energetic (normalized) POD modes for the non-motile suspension in (e). Mode 1 resembles a Poiseuille-like background flow, while modes 2 and 3 capture only minor noise fluctuations. Scale bar,  $100 \,\mu\text{m}$ . (i) Fractional POD mode energy for the motile (cyan) and non-motile (magenta) suspensions corresponding to (a-d) and (e-h), respectively. (j) Temporal amplitudes  $a_2$  (blue) and  $a_3$  (orange) of the motile suspension, corresponding to spatial modes 2 and 3 in (c) and (d), respectively. (k) Phase shifts of the temporal mode amplitudes (blue) and spatial modes (red) for modes 2 and 3 of the motile suspension are indicative of the traveling turbulent structures shown in (a).  $G_{\Delta t}$  is the temporal cross-correlation of mode amplitudes shown in (j), and  $G_{\Delta x,\Delta y=0}$  is the spatial cross-correlation in the stream-wise direction of POD modes shown in (c-d).

# D. Evolution of active turbulence structures in externally driven flow

Driven ambient flows are known to strongly modify the dynamics of active suspensions [43, 58, 59], leading to novel rheological [19, 20] and transport [43] properties. Pressure-driven flows of dense, swimming bacterial suspensions through microfluidic channels were recently observed to exhibit significant flow intermittency [43]. Active vortex structures interact with the imposed external flow to produce sporadic periods of near-parabolic and plug-like flow in motile suspensions of B. subtilis (Fig. 4a), compared to non-motile cells with steady flow (Fig. 4e). Here, we investigate this phenomenon through a POD analysis of previously published data [43]. The most energetic velocity field mode for the motile (Fig. 4b) and non-motile (Fig. 4f) suspensions are both representative of the parabolic background flow. For the active suspension, vortex chain structures (Fig. 4c-d) characterize higher-order modes  $(n \geq 2)$ , responsible for the observed flow intermittency. In contrast, the featureless higher-order modes for the non-motile cells (Fig. 4g-h) comprise a comparatively minute fraction of the total kinetic energy (Fig. 4i), commensurate with their steady, homogeneous flow. The non-trivial higher-order modes of the swimming bacterial suspension also give direct insights into the dynamics of the active flow structures.

The vortex chain modes,  $\phi_2(\mathbf{x})$  and  $\phi_3(\mathbf{x})$ , both appear periodic in space with a wavelength,  $\ell = 320 \,\mu\text{m}$ , and a cross-correlation analysis reveals a phase difference of  $\Delta x = 84.5 \,\mu\text{m}$  (Fig. 4k). Similarly, the temporal coefficients (Fig. 4j-k),  $a_2(t)$  and  $a_3(t)$ , exhibit a period and phase difference  $T = 1.27 \,\mathrm{s}$  and  $\Delta t = 0.333 \,\mathrm{s}$ , respectively. Similar to Fourier mode analysis, a linear combination of (minimum) two, orthogonal POD modes (Fig. 4c-d) can be used to describe the propagation of turbulent flow structures. Hence, we estimate the propagation velocity of the observed intermittent flow features (Fig. 4a) as a phase velocity  $v_P = \ell/T \approx 252 \,\mu\mathrm{m}\,\mathrm{s}^{-1}$ . This result is comparable to the flow structure velocity  $(\approx 235 \,\mu\mathrm{m\,s^{-1}})$  measured by manual tracking (see Appendix A 5) [43]. A principle application of POD is the identification of coherent flow structures [60–63], which is effectively demonstrated here in the case of active turbulence.

## III. CONCLUSION

In this work, we established the application of proper orthogonal decomposition to gain new physical insights into the collective motion of dense suspensions of swimming bacteria. Far from boundaries, POD analysis of quasi-2D bacterial turbulence revealed that the spatial and temporal structure of the underlying velocity field modes strongly correlated with kinetic energy, which naturally ranks the mode structures in decreasing size. Decreasing cell activity shifts energy to lower-order modes that are responsible for increasing the correlation time [4, 12]. For confined cell suspensions, our results accurately capture the transition to chaotic flow [18], and the resulting velocity field modes illustrate the importance of boundary interactions in regulating the dynamics, as predicted by active nematics simulations [29]. In external flows, higher-order velocity field modes describe the propagation of coherent active flow structures that are responsible for the observed flow intermittency [43]. Despite their prevalence in inertial turbulence and macroscale flow applications [37], the potential insights gained from POD and related techniques in microscopic flows have largely been overlooked, due to the typically linear behavior of low-Reynolds number flows. However, as illustrated here, the nonlinear, chaotic nature of active matter flows presents copious opportunities for mode analysis techniques to yield new insights into the dynamics and potentially the stability [57] of these complex systems. For example, beyond the effects of cell activity, POD analysis could provide additional insights into the effects of cell size, suspension density, nutrient availability, media/substrate, and viscosity on the structure [11, 21–24] and movement ecology [31] of collective motion. Furthermore, the compact descriptions of flow field data afforded by POD have recently been used in applications for machine learning of hydrodynamic equations [30], classification of wakes [64], and identification of ice flow drivers in glaciological processes [65]. As demonstrated by our present work, POD also provided an efficient, highly-compact description of experimental data for collective bacterial motility, where we anticipate that such low-dimensional representations will help facilitate future data-driven models of active turbulence [66, 67].

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# Appendix A: Methods

# 1. Cell culturing

Wild-type Bacillus subtilis bacteria (strain OI1084) were taken from  $-80^{\circ}$ C frozen stock and streaked onto 1.5% agar plates prepared with Terrific Broth (TB,

Sigma). Plates were incubated at 25°C for 24 hours, after which time a single colony from the plate was used to inoculate an overnight liquid TB culture at 30°C with shaking (200 rpm). The bacterial suspension was then subcultured (1.5 ml of cell culture into 60 ml of pre-warmed TB) and grown at 35°C and 200 rpm for 6 hours to mid-log phase (OD<sub>600</sub>  $\approx$  0.2). Immediately prior to experiments, dense cell suspensions ( $\sim 10^{10}$  cells.ml<sup>-1</sup>) were prepared by centrifugation at 5000 g for 5 minutes, and the pellet was resuspended with 2  $\mu$ l of fresh TB media.

### 2. Microfluidics and image analysis

Polydimethlysiloxane (PDMS) microfluidic channels were fabricated through soft lithography [68] and plasma bonded to No. 1 thickness glass coverslips. The PDMS channels were thinly cast to ensure ample diffusion of oxygen to the bacterial suspension and prolonged cell activity. Dense cell suspensions were gently loaded into the microfluidic devices via pipette, and the channel inlet and outlet were sealed with wax to prevent residual flows. For all experiments, bacterial suspensions were imaged with brightfield illumination on an inverted microscope (Nikon Ti-E) using a sCMOS camera (Zyla 5.5, Andor Technology). Time-resolved velocity fields,  $\mathbf{u}(\mathbf{x},t)$ , of the bacterial suspensions were measured by performing Particle Image Velocimetry (PIV) using PIVLab [69] implemented in MATLAB. The subsequent velocity fields were then lightly smoothed using a Guassian kernel with a standard deviation of one PIV pixel  $(1.73 \,\mu\text{m})$  in space and one frame (9.5 ms) in time. Vorticity fields,  $\omega(\mathbf{x},t)$ , were computed from measured velocity fields using central differencing (MATLAB). The POD was performed on the resulting velocity and vorticity fields using builtin singular value decomposition functions in MATLAB.

#### 3. Quasi-2D cell suspensions with varying activity

To capture the dynamics of dense quasi-2D bacterial suspensions in the absence of lateral wall effects (Fig 1-2), large microfluidic chambers  $(2 \text{ mm} \times 2 \text{ mm} \times 26 \mu\text{m})$ were prepared, where the side length corresponds to  $\approx 80$ correlation lengths of the turbulent bacterial suspension. Imaging was performed with a  $20\times$  objective (0.45 NA) at 105.5 fps. We imaged a 442  $\mu$ m square region in the center of the 2 mm square channel, corresponding to  $\sim 30$ correlation lengths from all boundaries. Bacterial activity (Fig. 2) was varied using a previously established approach [11, 12]. Briefly, the cell suspensions were imaged periodically over the course of 30 minutes, during which time the cell swimming speed naturally decayed due to oxygen depletion, where the decay rate of cell activity was controlled via the thickness of the PDMS device. Without the need to manipulate the cell suspension, this approach ensured that the bacterial concentration was constant across varying activity levels. Seven data sets were captured in total, which consisted of 6,300 frames each ( $\approx 1 \,\text{min}$  per video) and a  $2-3 \,\text{min}$  delay between acquisitions.

The energy spectrum, E(k), in 2D (Fig. 2a) was calculated through the Wiener-Khinchine theorem [4] as the Fourier transformation of the equal-time two-point velocity correlation function:

$$E(k) = \frac{k}{2\pi} \int d^2R e^{-i\mathbf{k}\cdot\mathbf{R}} \langle \mathbf{v}(\mathbf{r}, t) \cdot \mathbf{v}(\mathbf{r} + \mathbf{R}, \mathbf{t}) \rangle.$$

Autocorrelation functions (ACFs) for a given field  $\chi(\mathbf{r},t)$  were calculated as the mean time-averaged two-point correlation:

$$C_r(\Delta r) = \frac{\langle \chi(\mathbf{r}, t) \cdot \chi(\mathbf{r} + \Delta r, t) \rangle}{\langle \chi^2(\mathbf{r}, t) \rangle},$$

where  $\chi$  is  $\mathbf{v}$  or  $\phi$ , and the correlation length  $\delta$  is defined as  $C_r(\Delta r = \delta) = 1/e$ .

## 4. Cell suspensions under varying confinement

To quantify POD modes for bacterial suspensions under varying degrees of confinement (Fig. 3), experiments from [18] were replicated with minor modifications. Microfluidic devices (19  $\mu$ m deep) were fabricated, which comprised a series of racetrack geometries connected by narrow inlets (30  $\mu$ m wide). Nine racetrack widths were chosen between  $30 \,\mu\mathrm{m} < W < 171 \,\mu\mathrm{m}$  in fractional increments of our measured characteristic vortex size (38  $\mu$ m) taken as the minimum point in the spatial ACF in quasi-2D conditions (Fig. 1C, dotted red line). Cell suspensions were imaged at  $40 \times$  (0.6 NA) magnification and  $100\,\mathrm{fps}$ for 10s per channel width. All confined microfluidic geometries were simultaneously loaded with the same cell suspension and imaged within the first 10 min of loading to ensure consistent bacterial activity. Experiments were repeated across multiple days with freshly cultured bacteria to verify repeatability. The net flow  $\psi$  was calculated following the definition in [18]:  $\psi = |(\Sigma \mathbf{u} \cdot \hat{\mathbf{e}}_x) / (\Sigma ||\mathbf{u}||)|$ , where  $\hat{\mathbf{e}}_x$  is the unit vector along the principal channel direction and the sum is over all PIV sub-windows over 5 s (500 frames) of video.  $\psi = 1$  indicates unidirectional circulation flow around the racetrack, and  $\psi = 0$  corresponds to a globally stationary suspension. The stream-wise flow profiles, F(y), were generated from the velocity field by averaging over time and space as  $F(y) = \langle \mathbf{u}(x,y,t) \cdot \hat{\mathbf{e}}_x \rangle_{(x,t)}$ , where  $\hat{\mathbf{e}}_x$  was chosen as defined in [18], such that F(y) is on average positive.

## 5. Cell suspensions under externally imposed flow

The data corresponding to Fig. 4 was obtained from a previously published work [43], where the relevant experimental details are repeated here for convenience. Dense

suspensions ( $\approx 1.3-2\times 10^{10}$  cells/ml) of wild-type B. subtilis (strain OI1085) were flowed through a PDMS microfluidic chamber ( $200\,\mu\mathrm{m}$  wide  $\times~320\,\mu\mathrm{m}$  deep) via a syringe pump at a flow rate of  $Q=25\,\mu\mathrm{l/hr}$ . Imaging was performed under brightfield conditions at  $30\times$  magnification. For motile cell suspensions, 19 individual videos (4 s duration) were acquired at 370 fps. Velocity fields were quantified through ghost particle velocimetry [70] and averaged such that each pixel is the average of  $32\times 32$  pixel square over 50 frames. Image data and velocity fields for non-motile cell suspensions were acquired under the same conditions, but with a dead suspension of B. subtilis. Vortical regions in the flow were identified using the Okubo-Weiss parameter [71, 72] for manual tracking of travelling flow structures.

## Appendix B: Supporting Figures

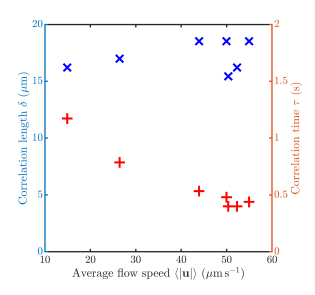


FIG. 5. Variations in the velocity correlation length and time scales with cell activity. As the suspension activity (proxied by average flow speed  $\langle |\mathbf{u}| \rangle$ ) decreases, the correlation length  $\delta$  (blue crosses) remains approximately constant, which is consistent with previous reports [11], but the correlation time  $\tau$  (red pluses) increases.

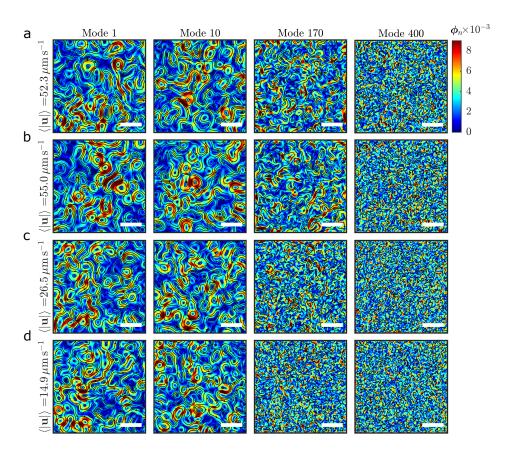


FIG. 6. Demonstration of mode structure changes with decreasing activity (row a-d) and increasing mode numbers (columns; see also Fig. 2). (a,b) Two different experimental realizations with similar cell activity levels and similar correlation lengths/times exhibit variations in mode structure for the same mode number due to the random nature of bacterial tubulence. As the mode number increases, typical feature size decreases towards noise (i.e. Mode 400, see also Fig. 1bc). (c,d) As the activity decreases, the dominant spatial scales are captured in a lower range of mode numbers (see also Fig. 2e), with noisy modes also occurring at lower mode numbers in the decomposition (see also Fig. 1g). Scale bars are  $100 \, \mu \text{m}$ .

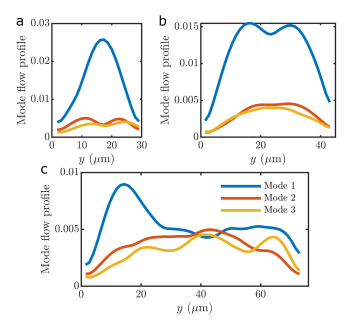


FIG. 7. Effect of confinement on individual mode flow profiles for (a)  $W = 30 \,\mu\text{m}$ , (b)  $W = 45 \,\mu\text{m}$ , (c)  $W = 76 \,\mu\text{m}$  (see Fig.3i). Here, the mode flow profiles are calculated by averaging individual spatial modes along the channel, which are scaled by their respective energies (i.e.  $\langle \sqrt{\lambda_n} \phi_n(x,y) \cdot \hat{\mathbf{e}}_x \rangle_{(x)}$ ), demonstrating the energy distribution across the y-direction of the channel. At the narrowest width (a;  $W = 30 \,\mu\text{m}$ ), mode 1 contains the majority of the energy of the system (see also Fig. 3j) and reflects the pumping state of the system under this confinement, with the subsequent modes providing small corrections to the flow profile. In the intermediate case (b;  $W = 45 \,\mu\text{m}$ ), the peak of mode 1 broadens due to the pinning vortical flow at this confinement. As the system transitions towards bulk turbulence (c;  $W = 76 \,\mu\text{m}$ ), the flow energy shifts to the walls - demonstrating the previously reported coherent, near-wall activity [18] – and the higher order modes now contribute significantly to the central portion of the channel

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