Autonomous waves and global motion modes in living active solids

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Elastic active matter—also called an active solid—consists of self-propelled units embedded in an elastic matrix and it resists deformation. This shape-preserving property and the intrinsic non-equilibrium nature make active solids an attractive potential component for self-driven devices, but their mechanical properties and emergent behaviour remain poorly understood. Here, using a biofilm-based living active solid, we observe self-sustained elastic waves with wave properties not seen in passive solids, such as power-law scaling of wave speed with activity. Under isotropic confinement, the active solid develops two topologically distinct global motion modes that can be selectively excited, with a step-like frequency jump at the transition between the two modes. Our findings reveal spatiotemporal order in elastic active matter and may guide the development of solid-state adaptive or living materials.

Active matter consists of living or synthetic units that can convert local free energy input to mechanical work. As an intrinsically non-equilibrium state of matter, active matter displays rich emergent behaviour such as dynamical collective motion and self-organization; it holds great potential for understanding the physics of living matter and for the design of autonomous or self-driven materials with life-like properties. Understanding the mechanical functionality of active matter as a continuum (that is, the ability to generate forces or extract work at length scales much larger than the individual unit) is an essential step to bridge the gap between practical applications and a myriad of fundamental science knowledge in the field learned over the past two decades. However, such understanding has been limited, in part owing to the poor scalability of most experimental systems. Here we use millimetre-sized bacterial biofilms to explore the emergent mechanical behaviour of elastic active matter or active solids. We discover that mass elements in bacterial active solids are self-driven into local oscillatory motion. Under two-dimensional (2D) isotropic confinement, the local oscillation self-organizes into a pair of topologically distinct global motion modes. The mode selection is tunable by varying the activity of the active units; surprisingly, the two modes transit between each other with a sharp, step-like frequency jump at a certain activity threshold. Under anisotropic confinement with a major axis of symmetry, the local oscillation of mass elements is organized in space as self-sustained elastic standing waves. These results are observed in both experiments and numerical modelling with remarkable agreement. Furthermore, our model predicts that the phase of local oscillations is arranged as travelling waves in unconfined space, with the wave speed scaling as the power-law of activity and the wavelength independent of activity. This property of active matter elastic waves is in stark contrast to the counterpart in passive mechanical waves where the wave speed does not depend on the driving amplitude of external stimuli. Our findings reveal unique mechanical properties of active matter and pave the way for investigating the non-equilibrium physics of elastic active matter in continuum. The findings may guide the development of solid-state adaptive or living materials, such as autonomous actuators for soft robotics, programmable tissues and synthetic microbial consortia with mechanical functionalities.

Elastic active matter consists of force-generating units embedded in an elastic matrix and they resist deformation like passive elastic solids; the shape-preserving property makes active solids a superior component for self-driven devices. Owing to the non-equilibrium nature, active solids are predicted to have novel mechanical properties that are not permitted in passive solids. However, active solids have been investigated only in macroscopic analogue models using robotic structures, and an experimental system appropriate for the study of active solids as continuous media is lacking. Motile bacteria are a premier experimental system for active matter studies. We envisaged that motile bacteria embedded in elastic polymer matrices may...
constitute a continuum body of active solid because of the superior scalability, so we turned to bacterial biofilms that consist of densely packed bacterial cells (>10^12 cells per ml) encased by cell-derived extracellular polymers^{42,43}. After testing several commonly used bacterial species, we identified early-stage Proteus mirabilis biofilms^{44} as a prospective active solid. Cells extracted from such early-stage *P. mirabilis* biofilms (~0.5 μm in width and ~2 μm in length) typically retained motility instead of having transitioned into a sessile state, while there was prominent production of extracellular amyloid fibrils matrix (Extended Data Fig. 1 and Methods). In the *P. mirabilis* biofilm, cells are self-propelled by rotating flagellar filaments (~20 nm in diameter and ~5–10 μm in length) appended on cell surfaces, and flagellar rotation is fueled by protonmotive force^{45}. The overall storage modulus (that is, a measure of the elasticity of cell-matrix assembly) of the homogenized early-stage *P. mirabilis* biofilms is higher than the overall loss modulus and ranges from ~10^3 Pa to ~10^9 Pa (Extended Data Fig. 2 and Methods); thus, the early-stage *P. mirabilis* biofilms represent a viscoelastic solid^{46} soft enough to be compliant with the active stress generated by bacteria (~1 Pa for a cell with single flagellum that is ~5–10 μm in length) appended on cell surfaces, and flagellar rotation is fueled by protonmotive force^{45}. Strikingly, we found that mass elements in such bacterial active solids undergo local oscillation of mass elements self-organizes into a pair of topologically distinct, self-driven global motion modes. In one mode, all mass elements being tracked in the biofilm followed a periodic quasi-circular trajectory in a synchronized manner (Fig. 1a, Supplementary Videos 1 and 2, and Methods); hence, the entire system underwent periodic translational motion and we referred to this mode as oscillatory translation. In the other mode, the mass elements followed periodic, synchronized quasi-linear trajectories that can be approximated as concentric circular arcs around the centre of the disk (Fig. 1b), and Supplementary Videos 3 and 4; hence, the entire system underwent global rotation with periodically switching chirality and this mode is referred to as oscillatory rotation. These results confirmed that the mass elements in the biofilms were confined to orbiting about a fixed equilibrium position, as expected in elastic solids; thus, we call the early-stage *P. mirabilis* biofilms a bacterial active solid. Although *P. mirabilis* biofilm is our choice of study, we note that the solid-like global motion can also be found in *Serratia marcescens* (both oscillatory translation and rotation modes) and in *Escherichia coli* (oscillatory rotation mode), suggesting the generality of the findings (see Methods for details). Interestingly, active-solid-like behaviour similar to the oscillatory translation mode reported here has been found in macroscopic robotic systems^{47–50}, which shows a characteristic pattern of a standing wave consisting of periodic, segmented domains with high correlation (Fig. 2d). By contrast, the phase of the collective movement component perpendicular to the major axis (denoted as the transverse component or v_y) does not vary in space (Fig. 2a,b, Supplementary Videos 7 and Methods); meanwhile, the instantaneous magnitude of v_y varies in space as a sinusoidal function (Fig. 2c). These features are also evident in the spatiotemporal autocorrelation of v_y along the major axis, which shows a characteristic pattern of a standing wave consisting of periodic, segmented domains with high correlation (Fig. 2d). In contrast, the phase of the collective movement component parallel to the major axis (denoted as the parallel component or v_x) does not vary in space (Fig. 2a,b), the spatiotemporal autocorrelation of v_x along the major axis shows a pattern of periodic, horizontal lines with high correlation (Fig. 2e). Taken together, the oscillatory motion of mass elements in such bacterial active solids under anisotropic lateral confinement is organized in space as a self-sustained transverse standing wave. We note that the phase of v_x, v_y always differs from that of v_y by π/2 or –π/2; thus, every mass element undergoes oscillatory elliptical motion.

To further understand the findings, we controlled the activity of mass elements in the bacterial active solid by tuning the speed of cells with violet-light illumination^{39} (Extended Data Fig. 4, Supplementary Video 8 and Methods). For bacterial active solids under disk-like isotropic lateral confinement, we found that the oscillatory translation and rotation mode dominates at higher and lower activity, respectively. Surprisingly, the modes transit to each other abruptly at certain activity threshold (Fig. 3a,b) with a sharp, step-like frequency jump (a fold change of 2.04 ± 0.42; mean ± s.d., N = 8) at mode transition (Fig. 3c), which agrees with the approximately twofold frequency difference of the two global motion modes that naturally emerged in disk-shaped bacterial active solids. These results show that activity selectively excites the two global motion modes, revealing a unique emergent mechanical property of active solids. Interestingly, the oscillation frequency of both modes is positively correlated with activity (Fig. 3c). Such activity dependence of the oscillation frequency is also evident in the self-sustained standing waves in bacterial active solids under anisotropic lateral spatial confinement, where the frequency of standing waves increases with activity in a continuous manner (Fig. 3d) rather than taking discrete values as in passive elastic plates. The activity dependence of the oscillation frequency is another unusual mechanical property of active solids.

To rationalize our experimental results, we performed numerical modelling of active solids. We modified the boundary conditions of a particle-based model to describe active solids^{53–55} by considering a collective of overdamped self-propelled particles connected by Hookean springs (with spring constant k_s) and initially arranged in a 2D triangular lattice (Extended Data Fig. 5a). Each particle represents a mass element consisting of 1,000 cells (Methods) and has an intrinsic self-propulsion...
their initial equilibrium positions, and particles initially sitting at the edge experience an additional elastic force pointing radially towards the edge to account for the steric effect of the lateral confinement boundary (Methods and Extended Data Fig. 5a). The position $\mathbf{x}_i$ and the self-propulsion polarity $\mathbf{n}_i$ of the $i$th particle evolve according to the following governing equations (Methods):

$$\mathbf{x}_i = v_0 \mathbf{n}_i + \Xi_i (\mathbf{F}_i + D_i \dot{\mathbf{F}}_i),$$

$$\dot{\mathbf{n}}_i = \beta_i ([\mathbf{F}_i + D_i \dot{\mathbf{F}}_i] \cdot \mathbf{n}_i) \mathbf{n}_i + D_i \dot{\mathbf{F}}_i - \kappa \frac{\mathbf{E}_n}{\mathbf{n}_i},$$

where $v_0$ is the activity of particles and the direction of polarity $\mathbf{n}_i$ coincides with the direction of particles’ self-propelled motion; $\Xi_i$ is a translational mobility tensor; $\mathbf{F}_i$ is the total external elastic force acting on the particle controlled primarily by the local elasticity of the system (corresponding to spring constants as described in the caption of Extended Data Fig. 5a); $\dot{\mathbf{F}}_i$ and $\dot{\mathbf{F}}_i$ are constants. $\mathbf{n}_i$ dynamics is controlled by three terms in equation (2), including a force-induced reorientation, a noise term and a term involving a Landau-type free energy $g = \mathbf{a} - 2 \mathbf{n}_i \cdot \mathbf{n}_i + \mathbf{n}_i \cdot \mathbf{n}_i + \frac{1}{2} \frac{\mathbf{E}_n}{\mathbf{n}_i}$ ($\mathbf{a}$ and $\mathbf{E}_n$ are constants) that penalizes the deviation of $\mathbf{n}_i$ from being a unit vector (Methods). The gradient part $\frac{1}{2} \frac{\mathbf{E}_n}{\mathbf{n}_i}$ in $\mathbf{F}_n$ allows for extending our model to active solids with microscopic geometrical anisotropy and orientational elasticity; nonetheless, here we focused on active solids with isotropic elasticity by setting $\kappa = 0$ in $\mathbf{F}_n$. Details of the model are described in Methods.

We found that all active particles in the modelled active solid are self-driven into local oscillatory motion with homogeneous frequency across space. Under 2D isotropic lateral confinement, our simulations successfully reproduced the two topologically distinct global motion modes observed in circular disk-shaped bacterial active solids (Extended Data Fig. 5 and Supplementary Videos 9 and 10). We scanned over the parameter space and obtained a phase map for the two global motion modes (Fig. 4a). We found that the emergence of the two global motion modes can be selected not only by the particle activity as demonstrated in the experiment but also by the system’s local elasticity; moreover, there was a sharp, step-like frequency jump (approximately twofold change) at the phase boundary (Fig. 4b, c and Extended Data Fig. 6a), in agreement with the experimental result during activity-controlled mode transition shown in Fig. 3c. Informed by the simulation results, we examined the elasticity dependence of the global motion modes in experiment by controlling the temperature of bacterial active solids that scales linearly with the elasticity of polymer networks. Indeed, we observed a transition between the two global motion modes at a certain temperature threshold accompanied by a sharp, approximately twofold frequency change (frequency ratio $2.04 \pm 0.77$, mean $\pm$ s.d., $N = 4$) (Extended Data Fig. 6b–d). A simulation-informed theoretical analysis provides quantitative insights into the frequency relations uncovered in the two emergent modes; see Supplementary Text for details. Essentially, denoting the frequency in the oscillatory translation and rotation mode as $f_t$ and $f_r$, respectively, the theory yields $f_t$ being positively correlated with activity ($f_t \approx \sqrt{v_0}$) and the ratio $f_t/f_r$ independent of activity $v_0$, which are in qualitative agreement with experimental and simulation results.

Under generic anisotropic lateral spatial confinement with a major axis of symmetry (such as in elliptical or rectangular geometry; Extended Data Fig. 7a), our simulations also successfully reproduced the self-sustained transverse standing waves found in experiments with bacterial active solids under anisotropic lateral confinement (Fig. 4d, Extended Data Fig. 7 and Supplementary Video 11). The remarkable agreement between experimental and simulation results prompted us to use the active solid model to explore the wave phenomena of active
Fig. 2 | Self-sustained transverse standing waves in bacterial active solids under anisotropic lateral confinement. a, Time sequence of collective velocity field in an oval-shaped bacterial active solid (Extended Data Fig. 1d) that shows the transverse standing wave (with period \( T = 8.6 \) s) (Methods). The longer side of the rectangular domain shown here is parallel to the major axis of the bacterial active solid. The arrows represent the velocity direction and the colour map indicates the velocity magnitude. Unit of velocity, \( \mu m \cdot s^{-1} \). Also see Supplementary Video 7. b, Spatial distribution of the oscillation phase of orthogonal collective velocity components associated with \( \alpha \). The phase of the parallel (\( \alpha \)) and the transverse (\( \beta \)) collective velocity components are denoted as \( \phi_\alpha \) (top) and \( \phi_\beta \) (bottom), respectively. The positive x axis in the specified coordinate system is parallel to the major axis of the active solid. Scale bars in a and b, 200 \( \mu m \). c, Amplitude distribution of \( \phi_\alpha \), along the major axis of the bacterial active solid. Data are presented as mean values (averaged over a time >60 s) ± s.d. Inset: temporal evolution of \( \phi_\alpha \) profile along the major axis, with colours representing the time associated with \( a \) (blue, 0; red, T/4; yellow, T/2; green, 3T/4). d,e, Spatiotemporal autocorrelation of \( \phi_\alpha \) (d) and \( \phi_\beta \) (e) along the major axis of the bacterial active solid (Methods). The colour map to the right of e indicates the autocorrelation magnitude (a.u.). The data in this figure are from a representative experiment (>10 replicates).

solids in unconfined 2D space, which is currently not attainable in experiments (Methods). We discover that individual particles followed oscillatory quasi-circular motion and the phase of the local oscillation is arranged as travelling waves (Fig. 4c and Supplementary Video 12). While the frequency of this travelling wave scales with particle activity \( \nu_0 \) as \( f = \nu_0^{0.48} \) (Fig. 4f), the wavelength is independent of particle activity (Fig. 4f); thus, the wave speed \( U \) is proportional to frequency and also scales with activity as \( U = \nu_0^{0.48} \). This result is in stark contrast to the counterpart in passive mechanical waves where the wave speed does not depend on the driving amplitude of external stimuli. Notably, the wave speed of such self-generated elastic active matter waves is on the order of a few hundred micrometres per second estimated based on values measured in the standing wave of bacterial active solids (frequency ~0.1 Hz, wavelength ~1–2 mm), which is much smaller than the speed of acoustic waves in ordinary passive solids. Unlike mechanical waves in passive solids, the elastic waves we uncovered in bacterial active solids are self-generated by the activity of mass elements. Our results illustrate that the inherent active nature of mass elements can give rise to unique properties of active matter waves.

To conclude, we have discovered an array of mechanical behaviours of elastic active matter that are not permitted in passive solids, including the formation of self-sustained elastic waves with activity-dependent wave properties and the emergence of two topologically distinct global motion modes with activity-dependent mode selection. The activity dependence of these uncovered mechanical behaviours provides simple means to control the emergent mechanical properties of solid-state active matter. In addition, our experimental system, that is, bacterial active solids derived from biofilms, has superior scalability and the convenience of engineering molecule-level interactions using tools from synthetic biology\(^{49-52}\); we expect that bacterial active solids will serve as a valuable experimental platform to explore the non-equilibrium physics of active solids in continuum\(^{49-52}\). Expanding the framework of classical continuum mechanics\(^{53-55}\), lessons learned from bacterial active solids may inspire the design of novel functionalities in adaptive or living materials for soft robotics and biotherapeutics\(^{56-58}\).
Fig. 4 | Global motion modes and self-sustained elastic waves in modelled active solids. a, Phase diagram of global motion modes in modelled active solids under isotropic lateral confinement. The phase diagram is plotted in the plane of particles’ activity \( v \) and system’s local elasticity (with the interparticle spring constant \( k_s \) serving as a proxy; Methods). Triangles and circles represent oscillatory translation and rotation modes, respectively. Each data point in the phase diagram was obtained with 120 simulation runs (Methods). b, Distribution of oscillation frequency (indicated by the colour map) in the phase diagram of a. c, Oscillation frequency of the modelled active solid under isotropic lateral confinement as a function of \( k_s \) (fixing \( k_r = 14 \)). The colour of the data points indicates the mode of global motion (blue, oscillatory translation; red, oscillatory rotation). Data are presented as mean ± s.d. (\( N = 100 \) simulation runs). Empty circles overlaid to the plot are experimental data from Fig. 3c. d, Self-sustained transverse standing waves in a modelled active solid under elliptical lateral confinement. Similar to Fig. 2d, this panel shows the spatiotemporal autocorrelation of the transverse component of particle velocity along the major axis of the elliptical confinement (that is, the abscissa); the pattern of periodic, segmented domains with high correlation is characteristic of a standing wave. The colour map indicates the autocorrelation magnitude (a.u.). Also see Extended Data Fig. 7. e, Self-sustained travelling wave in a modelled active solid in unconfined 2D space. This panel shows the spatiotemporal autocorrelation of the transverse component of particle velocity along the wave propagation direction (that is, the absissa); the pattern of periodic and tilted lanes with high correlation is characteristic of a travelling wave. The component of particle velocity parallel to the wave propagation direction shows the same autocorrelation pattern. The colour map indicates the autocorrelation magnitude (a.u.). f, Oscillation frequency (top) and wavelength (bottom) in unconfined 2D active solid as a function of particle activity \( v \). The black line in the top panel is a linear fit to the log-log plot with a slope of 0.48 ± 0.01 (\( R^2 = 0.997 \)). The data are presented as mean ± s.d. (\( N = 100 \) simulations runs).

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Methods
No statistical methods were used to predetermine sample size.

Bacterial strains
The following strains were used: wild-type P. mirabilis BB2000 and a fluorescent P. mirabilis KAG108 (BB2000 background with constitutive expression of green fluorescent protein (GFP)), from Karine Gibbs, Harvard University, Cambridge, MA; S. marcescens ATCC 274 and its derivative strains from Rasika Harshey, University of Texas at Austin; E. coli HB1 (from Howard Berg, Harvard University, Cambridge, MA) and its derivative strains. Single-colony isolates were grown overnight (~13–14 h) with shaking in LB medium (1% Bacto Tryptone, 0.5% yeast extract, 0.5% NaCl) at 30 °C to stationary phase. For P. mirabilis KAG108, ampicillin (100 µg ml⁻¹) was added to the growth medium to maintain the plasmid.

Agar plates
LB agar (Difco Bacto agar at specified concentrations infused with 1% Bacto Tryptone, 0.5% yeast extract, 0.5% NaCl) was autoclaved and stored at room temperature. Before use, the agar was melted in a microwave oven, cooled to ~60 °C, and pipetted in 10 ml aliquots into 90 mm polystyrene Petri plates. The plates were swirled gently to ensure surface flatness, and then cooled for 10 min without a lid inside a large Plexiglas box before inoculation of bacterial culture. To visualize the amyloid fibrils in P. mirabilis biofilms, the LB agar plates were supplemented with the fluorescence dye thioflavin T (Sigma; catalogue number T3136-5G; at a final concentration of 10 µM) and the plates were placed under the same conditions as specified above. Thioflavin T is commonly used to stain biofilm matrix consisting of amyloid fibrils. The presence of thioflavin T does not affect the experimental phenomena reported in this study.

Fabrication of bacterial active solid
To fabricate the bacterial active solid derived from P. mirabilis biofilms, overnight P. mirabilis cultures were inoculated onto a 0.6% LB agar plate (as described above). To track the motion of mass elements by following the trajectory of single cells embedded in P. mirabilis biofilms, overnight culture of the GFP-tagged P. mirabilis KAG108 was mixed with wild-type P. mirabilis at 0.05% before inoculating the agar plates. The inoculated plates were dried for 10 min without a lid inside the Plexiglas box. The plates were then covered and incubated at 37 °C and 95% relative humidity in an incubator with a water tray for 14 h. Circular P. mirabilis colonies incubated for this duration of time reached a diameter of 1.650 ± 270 µm (mean ± s.d., N = 102) and were at the early stage of biofilm development, with cells therein typically being motile and with prominent production of extracellular amyloid fibrils matrix in the inner region. The height at the centre of the colonies was ~25 µm, and it decreased to ~10 µm over a radial distance of ~700 µm (corresponding to a slope of ~1.2°). The inner region of P. mirabilis colonies with extracellular matrix production (that is, the biofilm region) (Extended Data Fig. 1b) is where we choose to study and refer to as early-stage biofilm or bacterial active solid. As shown in single-cell tracking videos (Supplementary Videos 2 and 4), the orientations of cells embedded in the extracellular matrix appear to lack long-range order (nonetheless, a certain extent of local orientational ordering could not be excluded), and the cell orientations do not rotate during the global motion of the biofilm. The outer rim of P. mirabilis colonies is mostly occupied by immotile cells that have transitioned to the sessile state but expressed little extracellular matrix; the width of this outer immotile rim varies from tens to hundreds of micrometres. This immotile rim serves as the lateral spatial confinement for the bacterial active solid; the lateral confinement is expected to exert steric repulsion that restricts radial but not tangential movement of the mass elements near the edge of the bacterial active solid. Colonies that have developed from an isolated inoculum are circular and disk-shaped. When two inocula happen to be nearby, the colonies arising from these inocula will merge into one oval-shaped colony, with the ratio of the long and short axes controlled by the initial distance between the inocula. The circular and oval-shaped colonies provided the biofilms (or bacterial active solids) with isotropic lateral spatial confinement and anisotropic spatial confinement with a major axis of symmetry, respectively.

The onset of the reported active solid motion requires sufficient biofilm matrix expression. Well before the presence of biofilm matrix, the entire colony is liquid-like and cells move freely, displaying bacterial turbulence often seen in dense bacterial suspensions. Once emerged, the active solid motion may last for several hours. Typically, the oscillatory translation mode first appears, and the motion may transit to the oscillatory rotation mode as the expression level of extracellular amyloid fibre matrix further increases and/or the motility of cells decreases. As a colony continues to grow, its thickness increases and cells therein eventually lose motility after ~30 h of growth (most likely due to depletion of nutrients and oxygen within the thick colony); the biofilm then becomes motionless.

Active solid motion can also be found in S. marcescens and E. coli biofilms. In S. marcescens biofilms with an overall storage modulus of ~10–103 Pa (in the experimentally relevant range of strain 0–40%), we found both oscillatory rotation and translation modes (Extended Data Fig. 8 and Supplementary Video 13). Under the same growth conditions as P. mirabilis biofilms, the overall storage modulus of E. coli biofilms is ~10–104 Pa, but it can be lowered by growing E. coli biofilms in a more humid environment (for example, depositing ~5 ml LB broth around the edge of the Petri dish). In E. coli biofilms with an overall storage modulus of ~10–107 Pa, we found that the biofilms can display the oscillatory rotation mode but not the oscillatory translation mode. This result is consistent with the phase map (Fig. 4a), which predicts oscillatory rotation mode at a relatively high elasticity.

Microscopy imaging
All imaging was performed on a motorized inverted microscope (Nikon Ti-E). To examine the expression of extracellular matrix, amyloid fibrils in P. mirabilis biofilms were visualized using a ×10 objective (Nikon CFI Achromat ×10, numerical aperture 0.25, working distance 7.0 mm) via a cyan fluorescence protein filter cube (excitation: 425/26 nm, emission: 479/40 nm, dichroic: 458 nm, FF458-Di02-25×36; Semrock), and images were recorded with a scientific complementary metal–oxide–semiconductor (sCMOS) camera (Andor Zyla 4.2 PLUS USB 3.0). To track the motion of mass elements in bacterial active solids derived from P. mirabilis biofilms, GFP-tagged P. mirabilis KAG108 cells embedded in the biofilm were imaged in epifluorescence using a ×20 objective (Nikon S Plan Fluor ×20, numerical aperture 0.45, working distance 8.2–6.9 mm) and an fluorescein isothiocyanate filter cube (excitation 482/35 nm, emission 536/40 nm, dichroic 506 nm; Semrock), with the excitation light provided by a mercury precentred fibre illuminator (Nikon Intensilight); meanwhile, to verify that the observed single-cell motion coincided with the motion of local mass elements where the cells were embedded, the background motion of P. mirabilis biofilm was imaged in phase contrast through the same optical system, with the illumination light provided by a white-light light-emitting diode (catalogue number MCWHLS; Thorlabs) installed above the microscope stage. Recordings were made with the sCMOS camera at 20 fps and at full frame size (2,048 × 2,048 pixels); fluorescent and phase-contrast images were recorded in alternate frames at 10 fps. The camera was controlled by NISElements (Nikon); the white-light light-emitting diode was switched on only during the acquisition of phase-contrast images and was triggered by 10 Hz Transistor-Transistor Logic signals sent from a custom-programmed Arduino microcontroller that modulated the 20 Hz fire output from the camera. To measure the collective velocity field in bacterial active solids, the sample was imaged in phase contrast with a ×4 (Nikon Plan Fluor ×4, numerical aperture 0.13, working distance 16.5 mm) objective. Recordings were made with the sCMOS
camera at 10 fps and at full frame size (2,048 × 2,048 pixels). To study the dynamics of the global motion modes and elastic waves in bacterial active solids at different activity level of mass elements, the speed of cells in the active solid was tuned by violet-light illumination at 528 mW cm⁻² provided by Nikon Intensilight and passing through the ×4 objective via a 406 nm filter (406/15 nm; FF01-406/15-25, Semrock); in the meantime, phase-contrast images of the active solid were recorded through the same ×4 objective by the sCMOS camera at 10 fps and at full frame size (2,048 × 2,048 pixels). In all experiments, the Petri dishes were covered with a lid to prevent evaporation and air convection. The sample temperature was maintained and controlled via a custom-built temperature-control system installed on the microscope stage.

Rheology of bacterial active solid

Rheological measurements of the bacterial active solid derived from *P. mirabilis* biofilms were performed in a rheometer (Anton Paar Physica MCR 301). The plate used for oscillatory shear measurements was CPS0-1 (diameter 50 mm, angle 0.990° and 99 μm gap; Anton Paar, part number 79040 serial: 20173). The overall storage modulus ($G′$) and loss modulus ($G″$) as a function of strain were measured in shear-strain-amplitude sweep mode (at constant frequency). The oscillation frequency and the temperature of the rheometer were set at 1 Hz and 37 °C, respectively. The temperature of the sample was controlled by the rheometer via a Peltier convection system connected to a water bath. A solvent trap equipped in the rheometer was used to reduce evaporation. Before rheology measurement, the *P. mirabilis* biofilms were gently scraped from Petri plates and collected into an Eppendorf tube, and the tube was heated at 35 °C for ~30 min to sterilize the bacterial cells without causing lysis. We note that this homogenization process might cause a certain disruptive effect to the biofilm structure, and thus the samples prepared for the rheological measurement should be taken as an approximation to the intact biofilms.

Image processing and data analysis for experiments

Images were processed using the open-source Fiji (ImageJ) software (http://fiji.sc/Fiji) and custom-written programs in MATLAB (MathWorks). To track the motion of local mass elements by following the trajectory of embedded single cells, trajectories of cells were obtained based on the recorded fluorescence videos, using a custom-written program in MATLAB published previously⁴⁶. The cells’ trajectories were then used to analyse their velocity and motion pattern. To compute the oscillation phase of individual cells embedded in circular disk-shaped bacterial active solids (insets of Fig. 1a,b), the position of individual cells extracted from their trajectories (lasting longer than 40 s) obtained by single-cell tracking was decomposed as Cartesian components ($x$, $y$) (for bacterial active solids undergoing oscillatory translation) or polar-coordinate components ($r$, $θ$) (for bacterial active solids undergoing oscillatory rotation; $r$ denotes radial component and $θ$ denotes tangential or azimuthal component, and the origin was chosen at the centre of the disk-shaped bacterial active solid). The position components $x(t)$, $y(t)$ and $θ(t)$ were then fitted by sinusoidal functions using the least-squares method in MATLAB in the form of $A \sin(ωt + φ₀)$, where $A$, $ω$ and $φ₀$ are fitting parameters and $(ωt + φ₀)$ was taken as the oscillation phase of individual cells.

To measure the collective velocity field of bacterial active solids, we first performed optical flow analysis based on phase-contrast time lapse videos using the built-in functions of MATLAB with a grid size of 1 pixel × 1 pixel. Before the optical flow analysis, the microscopy images were smoothed to reduce noise by convolution with a Gaussian kernel of standard deviation 1. The results were insensitive to different parameters of smoothing. To examine the uncertainty of the collective velocity measurement, we compared the collective velocity obtained by optical flow analysis and the average velocity of cells in the same field of view obtained by single-cell tracking in fluorescence images. We found that their difference was negligible (~4%). The excellent performance of optical flow velocimetry in our experimental system is due to the highly coherent motion of cells (that is, there is no relative motion between the cell bodies) when the system displays the active solid behaviour. The optical flow analysis yielded space- and time-dependent collective velocity field $v(r, t)$, or in Cartesian coordinates $(v_x(x, y, t), v_y(x, y, t))$. To compute the oscillation frequency, amplitude and phase of local mass elements in bacterial active solids, the obtained collective velocity field was coarse-grained by averaging over windows of a size 26 μm × 26 μm. The Cartesian components of coarse-grained collective velocity $v_x(x, y, t)$ and $v_y(x, y, t)$ at position $(x, y)$ are fitted by sinusoidal functions using the least-squares method in MATLAB in the form of $A \sin(2πf t + φ₀)$, where $A$, $f$, and $φ₀$ are fitting parameters that represent the velocity amplitude, frequency and phase shift of the local oscillation at position $(x, y)$, respectively. The phase of the local oscillation at position $(x, y)$ at time $t$ ($2πf t + φ₀$). As the frequency of the local oscillation is nearly homogeneous in space for a given bacterial active solid (Extended Data Fig. 3), we took the spatial average of the fitted $φ₀$ as the oscillation frequency of the entire system being discussed. To characterize the self-sustained elastic waves in bacterial active solids, the spatiotemporal autocorrelation along certain direction (for example, the major axis of bacterial active solids under anisotropic lateral confinement denoted as $x$-axis) for a collective velocity component (for example, $v_x(x, y, t)$ component perpendicular to the major axis) was computed as $C(Δx, Δt) = \frac{1}{T} \int_{t_0}^{t_0+T} v_x(x, y, t) v_x(x+Δx, y, t+Δt) dt$, where angular brackets (...) indicate averaging over the spatial coordinate on the chosen direction and time $t$.

To compute the spatially averaged collective velocity of bacterial active solids under isotropic lateral confinement, we decomposed the obtained collective velocity field $v(r, t)$ in Cartesian coordinates as $v_x(x, y, t)$ and in polar coordinates as $v_r(r, t)$ and $v_θ(r, t)$ ($v_r$ denotes radial component and $v_θ$ denotes tangential or azimuthal component; the origin of polar-coordinate system is chosen at the centre of the circular disk-shaped bacterial active solid). Each of these collective velocity components were then spatially averaged over the domain within two-thirds of the radius of the bacterial active solid (to exclude the near-boundary region with a larger slope of height variation), yielding the corresponding components of the spatially averaged collective velocity $(V_r(t), V_θ(t))$ and $(V_r(t), V_θ(t))$ of the average collective speed of bacterial active solids (that is, activity in the horizontal axis of Fig. 3c,d). $C(Δx, Δt)$ was computed as the root mean square of the magnitude of collective velocity vectors in the collective velocity field $v(r, t)$ averaged over several oscillation periods. A more accurate definition of cell activity is the self-propulsive force generated by flagellar motility of an average cell. However, we could not make direct measurement of the self-propulsive force and thus the speed of cells becomes an appropriate proxy.

To determine the specific mode of global motion in bacterial active solids under isotropic lateral confinement, we used the relative dominance of the Cartesian or polar-coordinate components of the spatially averaged collective velocity. Specifically, we fitted the spatially averaged collective velocity components $V_r(t)$, $V_r(t)$, $V_θ(t)$, $V_θ(t)$ of the obtained by procedures as described above into sinusoidal functions (using the least-squares method in MATLAB) in the form of $V(r, t) = B \sin(2πf t + φ₀)$, where $B$, $f$, and $φ₀$ are fitting parameters that represent the collective velocity amplitude, oscillation frequency and phase shift, respectively. Note that $f$ is equivalent to the spatial average of the frequency of local oscillations described above. The fitting yielded four sets of parameters ($B$, $f$, $φ₀$) corresponding to the amplitude, oscillation frequency and phase shift, respectively, of each $V(t)$ with the index $i$ being $x$, $y$, $r$, $φ$. In all experiments, we found that $B_0 = 0$ and $f_0 = 0$. If $B_r > B_θ > B_0$, the global oscillatory motion of bacterial active solids was classified as oscillatory translation mode; if $B_r > B_θ > B_0$, the global oscillatory motion was classified as oscillatory rotation mode; for all other cases, the global oscillatory motion of bacterial active solids was
classified as coexistence mode. The coexistence mode was observed both in experiments (Extended Data Fig. 9) and in bead–spring model simulations near the mode transition thresholds at intermediate activity and elasticity (Extended Data Fig. 10). Interestingly, the frequencies of the two global motion modes in a mixed-mode state were always either almost identical \((f - f_v; \text{Extended Data Fig. 9a})\) or doubled \((f_v = 2f_v; \text{Extended Data Fig. 9b})\). In the first scenario \((f - f_v)\), mass elements in the bacterial active solid followed quasi-elliptical trajectories (Extended Data Fig. 9c). By contrast, in the other scenario \((f_v = 2f_v)\), mass elements followed figure-of-eight shaped curves (known as lemniscates) (Extended Data Fig. 9d). Presumably the mode coexistence represents an incomplete mode transition process, reminiscent of a phase-coexistence state during a first-order phase transition.

During the transition between the oscillatory translation mode and the oscillatory rotation mode in bacterial active solids under isotropic lateral confinement while continuously varying bacterial speed or temperature, the oscillation frequency varied with time. Instead of performing sinusoidal fits as described above, we computed the oscillation frequency of global motion at a specific time as the inverse of the time interval between the two adjacent peaks of an appropriate component of spatially averaged collective velocity (choosing one of the Cartesian components for the oscillatory translation mode and the tangential component in polar coordinates for the oscillatory rotation mode, respectively).

**Simulation**

To understand the dynamical phenomena uncovered in the bacterial active solid, we adopted a particle-based simulation framework and considered a collective of overdamped self-propelled particles connected by springs. Each particle in the simulation represents a circular disk-shaped mass element of diameter \(-16\) µm and height \(-20\) µm; thus, a mass element consists of \(-1,000\) cells \((-0.8\) µm in width and \(-2.0\) µm in length), assuming a cell volume fraction of \(-0.2\) in the biofilm. The particles were initially arranged in a 2D triangular lattice truncated by a circular domain with radius \(R\). For the \(i\)th active particle or head, its position \(\mathbf{x}_i\) and self-propulsion polarity \(\mathbf{n}_i\) evolve according to equations (1) and (2). In equation (1), the translational mobility tensor is given by \(M = \alpha_0 \hat{n}_i \cdot \hat{n}_i + \alpha_1 (1 - \hat{n}_i \cdot \hat{n}_i)\) with \(\hat{n}_i\) being the unit vector parallel to \(\mathbf{n}_i\), \(\alpha_0\) (or \(\alpha_1\)) being the parallel (or perpendicular) inverse translational damping coefficients, and \(\mathbf{v}_i\) being a \(2 \times 2\) identity matrix; in equation (2), \(\beta\) is the inverse rotational damping coefficient.

The particle’s self-propulsion velocity (that is, \(\mathbf{v}_i \mathbf{n}_i\) in equation (1)) in our model represents the velocity of emergent self-propelled motion of a mass element, which presumably results from alignment of self-propulsive forces of motile cells within the mass element. Here the direction of polarity vector \(\mathbf{n}_i\) of a particle or mass element can be regarded as the direction of statistically averaged self-propulsive forces of all cells in the mass element; the magnitude of \(\mathbf{n}_i\) is variable and reflects the degree of alignment of cells’ self-propulsive forces. Meanwhile, similar to the average collective speed in the horizontal axis of Fig. 3c,d, the particle activity \(v_0\) should be interpreted as a proxy of the magnitude of self-propulsive force generated by flagellar motility of a typical cell in the mass element. The spontaneous alignment of self-propulsive forces of motile cells in a mass element is probably due to cell–matrix interactions within the mass element; at steady state, we expect that the self-propulsive forces tend to be well aligned to a common direction and thus the polarity \(\mathbf{n}_i\) approaches a unit vector (or the particle’s self-propulsion speed \(v_\parallel, \mathbf{n}_i\) approaching \(v_\parallel\); note that this is different from the particle’s apparent speed). In the model, this scenario is realized by the Landau-type free energy term \(F_{\text{e}}\), which controls the magnitude of \(\mathbf{n}_i\) and penalizes the deviation of \(\mathbf{n}_i\) from being a unit vector.

As described in the main text and Extended Data Fig. 5a, three types of Hookean spring are involved in the model, namely an interparticle spring, a restoring spring and a boundary spring with spring constants \(k_b, k_r\), and \(k_p\), respectively. The elastic interaction between mass elements mediated through the biofilm matrix is characterized by the springs with stiffness \(k_b\); the springs with stiffness \(k_r\) account for the adhesion between the bacterial active solid and the substrate, which ensures that the centre of mass of the system is confined to orbiting about a fixed equilibrium position; the springs with stiffness \(k_p\) account for an effective steric effect of the lateral confinement boundary and maintain the in-plane geometrical shape of the bacterial active solid. Therefore, the external elastic force experienced by the \(i\)th particle \(\mathbf{F}_i\) is a sum of three terms, \(\mathbf{F}_i = \mathbf{F}_{\text{e}} + \mathbf{F}_\text{adh} + \mathbf{F}_{\text{conf}}\), with each term given below.

The first term \(\mathbf{F}_{\text{e}}\) is a sum of elastic forces between the \(i\)th particle and its spring-connected neighbours (denoted as set \(S_i\)):

\[
\mathbf{F}_{\text{e}}(i) = \sum_{j \in S_i} -k_b(\mathbf{r}_{ij} - l_0)\mathbf{r}_{ij}/|\mathbf{r}_{ij}|.
\]

where \(l_0\) is the equilibrium distance of interparticle springs with spring constant \(k_b\), and \(\mathbf{r}_{ij} = \mathbf{x}_i - \mathbf{x}_j\). The second term \(\mathbf{F}_{\text{adh}} = -k_r(\mathbf{x}_i - \mathbf{x}_{\text{bound}})\) is a linear elastic restoring force that pulls the \(i\)th particle towards its equilibrium position \(\mathbf{x}_{\text{bound}}\). The third term \(\mathbf{F}_{\text{conf}}\) accounts for the steric effect of the lateral confinement boundary and thus it applies to particles initially sitting at the edge only; for instance, under circular lateral confinement, such particles are those with the position satisfying \(|\mathbf{x}_i - \mathbf{x}_{\text{bound}}| < l_0\), where \(R\) is the radius of the circular lateral confinement and \(\mathbf{x}_{\text{bound}}\) is the equilibrium position of the centre of the modelled active solid. \(\mathbf{F}_{\text{conf}}\) is a linear elastic restoring force pointing towards the edge in parallel to the normal direction \(\mathbf{n}_{\text{bound}}\) (that is, the unit vector perpendicular to the lateral confinement boundary) given by

\[
\mathbf{F}_{\text{conf}}(i) = -k_p(\mathbf{x}_i - \mathbf{x}_{\text{bound}}) \cdot \mathbf{n}_{\text{bound}}.
\]

As active units in the bacterial active solid (each active unit consisting of an associated group of motile cells) are densely packed, an active unit would find it more difficult to move sideways than to move along its polarity direction due to steric hindrance. This indicates that \(a_0 \ll a_1\) for active particles in the model; for convenience we chose \(a_0 = 1\). All simulations were carried out with \(a_0 = 10, \beta = 1, D = 0.0001, D_0 = 0.01, \tau = 1, \alpha = 0.1\), initial or equilibrium interparticle distance \(l_0 = 65\) and time step \(\Delta t = 0.0005\). The emergence of the two global oscillation modes does not require the presence of noise (that is, setting both \(D_0\) and \(D_{\text{ch}}\) to zero does not affect the results); also the simulation results are robust as long as \(D_{\text{ch}} \leq 10^3\) (with the ratio between \(D_0\) and \(D_{\text{ch}}\) fixed).

To match with experimental results, we set one time unit as \(1\)) and one length unit as \(0.25\) µm. For simulations of active solids under isotropic lateral confinement, the radius of the circular simulation domain was chosen as \(R = 15d\) with the particle number \(N = 511\); for simulations with elliptical lateral confinement, the semi-major and semi-minor axis of the simulation domain were \(a = 240d\) and \(b = 9d\), respectively, with the particle number \(N = 2,419\) for simulations in unconfined 2D space, the size of the simulation domain was \(480d \times 480d\) with the particle number \(N = 231,121\). Also, for simulations with either isotropic or anisotropic lateral spatial confinement, we kept the ratios between the interaction spring constants \(k_s, k_r, k_p\) and \(k_s = k_r/20\) and \(k_p = k_s/270\) unless otherwise specified; the values of \(k_s\) and \(k_r\) relative to \(k_p\) were selected to best fit the experimental results, including mode selection and approximately twofold frequency change during the transition. For simulations in unconfined 2D space, the boundary spring constant \(k_b\) (for the interaction between cells and the lateral confinement boundary) was set to 0, while the ratio between \(k_b, k_r, k_p\) was kept as \(k_b = k_r = k_p/270\). All simulations began with a random orientational distribution of particle polarity. Despite the similarity between our model and the models developed in refs. 10,46, we did not observe the persistent translation and rotation predicted in these earlier studies. The difference is due to the different boundary conditions we used (including lateral spatial confinement and substrate adhesion).

We note that the spring constant \(k_b\) can be mapped to the shear modulus (storage modulus) \(G\) obtained via rheological measurement.
in the experiment. Modelling the bulk part of the biofilm as a triangular elastic spring network, the relationship between Young’s modulus $E$ and bulk spring constant $k_b$ can be written as $E = \frac{2k_b}{\sqrt{3}H_e}$, where $H_e$ is the height of the biofilm ($H_e = 20 \mu m$). Noting that $G = \frac{k}{211+\nu}$ and assuming the Poisson’s ratio of the biofilm $\nu = 0.5$, we have $k_b = \sqrt{3H_e}/2 \approx 3\sqrt{3G}H_e$ As $G$ is in the range of $10^2$–$10^3$ Pa (Extended Data Fig. 2), we have $k_b = 10^{-10^3}$ pN $\mu m^{-1}$. To convert the parameters in physical units to dimensionless ones for simulations, we choose one length unit as $\xi = 0.25 \mu m$, one velocity unit as $\bar{v} = 0.25 \mu m \ s^{-1}$, and one unit of spring stiffness as $\bar{k} = 1,000$ pN $\mu m^{-1}$. With these basic units, the typical value of $k_b$ estimated above ($5.0 \times 10^3$ pN $\mu m^{-1}$) is expressed as $-5\xi$; therefore we chose $k_b$ ranging from 1 to 20 in the simulations. Other simulation parameters are expressed in units of a combination of $\xi$, $\bar{v}$ and $\bar{k}$. For instance, the full expression of the inverse translational damping coefficient at its chosen value should be $a_\parallel = 10 e^{-k^1 \xi^1}$.

Data analysis for bead–spring model simulations
All calculations and data analysis in active solid simulations followed the same procedures as described above for bacterial active solids, except that the local mass elements and the associated collective velocity field $\mathbf{v}(r, t)$ obtained by optical flow analysis were now replaced by the active particles and their velocities. For modelled active solids under isotropic lateral confinement (that is, in circular geometry), we found that the oscillation amplitude of the radial component of spatially averaged velocity of active particles was also nearly zero (Extended Data Fig. 5e), which is similar to that in experiments (Fig. 1f). To obtain the phase diagram of global motion modes in modelled active solids under isotropic lateral confinement (Fig. 4a), 120 independent simulation runs were performed for each parameter set ($\bar{v}_o, k_b$), and the mode of emergent global motion was determined for each simulation run at steady state. The mode that emerged in $>50\%$ of the 120 simulation runs was taken as the dominant mode of global motion at the parameter set ($\bar{v}_o, k_b$), and the mean oscillation frequency at ($\bar{v}_o, k_b$) was computed by averaging the frequencies of those simulation runs displaying the dominant mode.

Reporting summary
Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

Data availability
The data supporting the findings of this study are included within the paper and its Supplementary Information. All other data that support the plots within this paper and other findings of this study are available from the corresponding author upon reasonable request.

Code availability
The custom codes used in this study are available from the corresponding author upon request.

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Author contributions
H.X. discovered the phenomena, designed the study, performed experiments, performed simulations, and analysed and interpreted the data. Y.H. and R.Z. developed the theory and improvised the simulations. Y.W. conceived the project, designed the study, and analysed and interpreted the data. R.Z. and Y.W. supervised the study. Y.W. wrote the first draft and all authors contributed to the revision of the manuscript.

Competing interests
The authors declare no competing interests.

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Extended Data Fig. 1 | *Proteus mirabilis* colonies at the early stage of biofilm development with prominent production of extracellular polymer matrix. (a) Phase-contrast image of a circular disk-shaped *P. mirabilis* colony grown for 14 hr at 37 °C after inoculation with overnight culture (Methods). Scale bar, 500 μm. (b) Fluorescent image of extracellular amyloid fibrils matrix labelled by Thioflavin T (Methods) in the *P. mirabilis* biofilm shown in panel a. The outer rim of *P. mirabilis* colonies at this development stage is mostly occupied by immotile cells that have transitioned to the sessile state but expressed little extracellular matrix. The width of this outer immotile rim varies from tens to hundreds of μm across different colonies; in the case of panel a, the immotile rim ranges from radius $R = 860 \mu m$ (measured from the colony centre) to $R = 1154 \mu m$ (i.e., the colony edge), spanning a width of ~300 μm. The inner region of the colonies with Thioflavin T fluorescence (that is, the biofilm region; enclosed by the dashed circle in panels a,b) is where we choose to study and refer to as early-stage biofilm or bacterial active solid, such as the fields in main text Figs. 1c,d and 3a. The immotile outer rim of the colony serves as the lateral spatial confinement for the bacterial active solid. Scale bar, 500 μm. (c) Enlarged view of the centre of panel b. Scale bar, 100 μm. (d) Phase-contrast images of oval-shaped *P. mirabilis* colonies with various values of the eccentricity. Scale bars, 500 μm. The field in main text Fig. 2a corresponds to the stripe region enclosed by the dashed rectangle in panel d.
Extended Data Fig. 2 | Dynamic moduli of the bacterial active solid derived from early-stage *P. mirabilis* biofilms. Storage ($G'$; red) and loss ($G''$; blue) shear modulus of the bacterial active solid as a function of strain were measured by bulk rheometry with a cone-plate rheometer in shear-strain-amplitude sweep mode (at constant frequency) (Methods). The oscillation frequency and the temperature of the rheometer were set at 1 Hz and 37 °C, respectively. Data are presented as mean ± S.D. (N = 3). The solid and dashed lines are guides to the eye.
Extended Data Fig. 3 | Spatial distribution of local oscillation frequency in bacterial active solids. Panels a and b shows the frequency distribution in circular disk-shaped bacterial active solids undergoing global oscillatory translation and oscillatory rotation, respectively, with the colour bars indicating the magnitude of local oscillation frequency (unit: Hz). Scale bars, 500 μm. The frequency at the centre of panel b is absent because the velocity there is vanishing. Panel c shows the frequency distribution in a rectangular region of an oval-shaped bacterial active solid that displays the transverse standing wave, with the colour bar indicating the magnitude of local oscillation frequency (unit: Hz). The longer side of the selected region is parallel to the major axis of symmetry of the bacterial active solid. Scale bar, 200 μm.
Extended Data Fig. 4 | Tuning single-cell speed of \textit{P. mirabilis} via violet-light illumination. To obtain this plot, cells were extracted from the \textit{P. mirabilis}-based bacterial active solids that were undergoing either global oscillatory translation or global oscillatory rotation; the extracted cells were mixed with 0.02\% Tween 20 and deposited on 0.6\% LB agar surface, forming a quasi-2D dilute bacterial suspension drop. \textit{P. mirabilis} cells in the prepared quasi-2D dilute bacterial suspension drop were continuously illuminated by 406 nm violet light starting from $T = 0$ s while being tracked in phase-contrast microscopy (Methods); note that for single-cell tracking, a 20x objective lens was used. The speed of an individual cell at a specific time T was computed based on its trajectory tracked from $(T-0.5)$ s to $(T + 0.5)$ s; the single-cell speeds computed from $(T-25)$ s to $(T + 25)$ s were then averaged and taken to be the mean cell speed at T. Data shown in the plot was normalized by the mean speed at $T = 0$ s (that is, the free-swimming speed of cells without blue light illumination; $V_0 = 26.7 \pm 6.9$ $\mu$m/s; mean$\pm$S.D., N = 2500). Data are presented as mean $\pm$ S.D. (N = 2500).
Extended Data Fig. 5 | Emergent global motion modes in modelled active solid under isotropic lateral confinement. (a) Schematic diagram of the bead–spring model for active solid under two-dimensional isotropic lateral confinement (Methods). The model consists of $N = 511$ self-propelled particles (black solid circles). Every nearest-neighbour pair of particles is connected by an interparticle spring with spring constant $k_b$ (red). The particles also experience elastic forces due to substrate adhesion and lateral spatial confinement (see main text) via a restoring spring (green) and a boundary spring (blue) with spring constant $k_s$ and $k_r$, respectively. The three spring constants together determine the system’s local elasticity. In simulations the interparticle spring constant $k_b$ was used as a proxy for the system’s local elasticity, with the ratios between $k_b$, $k_s$ and $k_r$ fixed. (b, c) Representative trajectories of particles in the modelled active solid that underwent global oscillatory translation (panel b) and oscillatory rotation (panel c). Most particles (except those very near the centre or the boundary) followed periodically oscillating quasi-circular trajectories (at relatively high activity; panel b; Supplementary Video 9) or quasi-linear concentric trajectories (at relatively low activity; panel c; Supplementary Video 10) with highly synchronized phases (insets of panel b,c), in the same manner as the motion of matrix-embedded cells in the experiments undergoing global oscillatory translation (main text Fig. 1a) or rotation (main text Fig. 1b), respectively. Black dot in each panel indicates the centre of the simulation domain. Scale bars represent 1/3 of the interparticle distance at equilibrium and colour map indicates time. Insets: Oscillation phases of individual particle’s velocity components plotted in the same way as in insets of Fig. 1a,b. Simulation parameters: $v_0 = 15$ (panel b) or $v_0 = 3$ (panel c), $k_b = 12$, $k_r = 0.6$, and $k_s = 0.044$. (d, e) Temporal dynamics of spatially averaged particle velocity in the modelled active solid in global oscillatory translation mode (panel d) or oscillatory rotation mode (panel e). The velocity was averaged over all particles in the simulation and then decomposed as Cartesian (yellow and blue traces) and polar-coordinate components (red: tangential or azimuthal component; green: radial component). The spatially averaged particle velocity in the two emergent modes was characterized by distinct temporal dynamics in Cartesian or polar coordinates similar to that found in the experiments (main text Fig. 1e,f). Simulation parameters are identical to those used in panels b,c.
Extended Data Fig. 6 | Elasticity dependence of the global motion modes in active solids under isotropic lateral confinement. (a) Oscillation frequency of global motion modes in modelled active solids as a function of $\lambda_c$ (fixing $v_0 = 4$). Colour of data points indicates the mode of global motion (blue: oscillatory translation; red: oscillatory rotation). Data are presented as mean +/- S.D. (N=100 simulation runs). (b) Temperature dependence of $P.\ mirabilis$ single-cell speed. $P.\ mirabilis$ cells in quasi-2D dilute bacterial suspension drops (prepared in the same manner as described in the caption of Extended Data Fig. 4) were tracked in fluorescent microscopy while the environmental temperature was varied from 24 °C to 50 °C with a custom-built temperature-control system (Methods). As shown in the plot the speed of cells only changed slightly in this temperature range (up to ~15%). The mean speed of cells at a specific temperature was computed based on 1-s segments of cell trajectories tracked in a 200-s time window. Data shown in the plot was normalized by the mean speed at temperature 24 °C. Data are presented as mean +/- S.D. (N = 2500). (c,d) Transition of global motion modes in bacterial active solids controlled by temperature. Panel c shows the temperature dependence of oscillation frequency in the bacterial active solid during mode transition. Colour of data points indicates the mode of global motion (blue: oscillatory translation; red: oscillatory rotation). Panel d shows the temporal dynamics of spatially averaged collective velocity during transition from the oscillatory rotation mode to the oscillatory translation mode following the decrease of temperature. The spatially averaged collective velocity was decomposed as Cartesian components (yellow and blue traces; upper part of panel d) and polar-coordinate components (red: tangential or azimuthal component, green: radial component; lower part of panel d). Data in panels c,d were from a representative experiment (>5 replicates).
Extended Data Fig. 7 | Self-sustained transverse standing waves in modelled active solids under anisotropic lateral confinement. This figure is associated with main text Fig. 4d. Simulation parameters: \(v_0 = 10, \ k_b = 4, \ k_r = 0.2, \) and \(k_s = 0.015.\) (a) Schematic diagrams of the bead–spring model for active solid under two-dimensional anisotropic lateral confinement (Methods). The elliptical confinement (left) mimics the anisotropic lateral confinement geometry of oval-shaped bacterial active solids used in main text Fig. 2. Rectangular lateral confinement (right) produces similar simulation results. (b) Time sequence of particle velocity field in a modelled active solid under elliptical lateral confinement that displays the transverse standing wave. The longer side of the rectangular domain shown here is in parallel to the major axis of the elliptical confinement. T denotes the period of oscillation. Arrows represent velocity direction and colour map indicates velocity magnitude. (c) Spatial distributions of local oscillation frequency and phase associated with panel b. The local oscillation frequency is homogeneous in space (upper part of the panel, with the colour bar indicating the magnitude of frequency). The phase of parallel (\(v_x\); parallel to the major axis) and transverse (\(v_y\); perpendicular to the major axis) component of particle velocity is denoted as \(\phi_x\) (lower) and \(\phi_y\) (upper), respectively. Scale bar under panel c is shared by panel b and represents 7.7 times of the interparticle distance at equilibrium. (d) Averaged amplitude distribution of \(v_y\) along major axis of the modelled active solid. Data are presented as mean ± S.D. (N=100 simulation runs). Inset: temporal evolution of \(v_y\) profile in panel b along the major axis of the modelled active solid, with colours representing time (blue: 0; red: T/4; yellow: T/2; green: 3T/4). (e) Spatiotemporal autocorrelation of \(v_x\) along the major axis of the modelled active solid (that is, the abscissa of the figure); the pattern of periodic, horizontal lanes with high correlation is similar to that seen in experiment (main text Fig. 2e). Colour map at right side indicates the autocorrelation magnitude (a.u.).
Extended Data Fig. 8 | Emergent global motion modes in bacterial active solids derived from *S. marcescens* biofilms. Temporal dynamics of spatially averaged collective velocity in the global oscillatory translation mode (panel a) and oscillatory rotation mode (panel b). The spatially averaged collective velocity was decomposed as Cartesian (yellow and blue traces) and polar-coordinate components (red: tangential or azimuthal component; green: radial component). In the oscillatory translation mode, the polar-coordinate components are negligible; in the oscillatory rotation mode, both the radial and the Cartesian components are negligible.
Extended Data Fig. 9 | Coexistence of two global motion modes in bacterial active solids under isotropic lateral confinement. (a, b) Temporal dynamics of spatially averaged collective velocity in circular disk-shaped bacterial active solids where the oscillatory translation and oscillatory rotation modes co-existed with identical frequencies ($f_t = f_r = 0.15\text{Hz}$; panel a) or with the frequency of the oscillatory translation mode doubled ($f_t = 0.14\text{Hz}, f_r = 0.07\text{Hz}$; panel b). The velocity was decomposed as Cartesian (yellow and blue traces) and polar-coordinate components (red: tangential component; green: radial component).

(c, d) Panels c and d display representative trajectories of mass elements in bacterial active solids analysed in panels a and b, respectively. In each panel the trajectories were obtained by integrating the spatially averaged collective velocity over a $50\text{\mu m} \times 50\text{\mu m}$ domain located ~500$\mu$m from the centre of the disk-shaped bacterial active solid (black dot) at different polar angles. The trajectories were brought close to the centre for better visualization, and thus the scale bars (panel c, 10$\mu$m; panel d, 20$\mu$m) apply to the trajectories only. Colour map indicates time (unit: s).
Extended Data Fig. 10 | Coexistence of two global motion modes in modelled active solid under isotropic lateral confinement. (a,b) Temporal dynamics of collective velocity of the system where oscillatory translation and oscillatory rotation modes co-existed with identical frequencies ($f_t = f_r = 0.10$; panel a) or with the frequency of the oscillatory translation mode doubled ($f_t = 0.13$, $f_r = 0.07$; panel b). The collective velocity was averaged over all particles in the simulation and then decomposed as Cartesian (yellow and blue traces; upper part of each panel) and polar-coordinate components (red: tangential or azimuthal component, green: radial component; lower part of each panel).

Simulation parameters: panel a, $v_0 = 5, k_b = 12$; panel b, $v_0 = 9, k_b = 16$; the ratios between $k_s, k_t$ and $k_r$ are fixed (Methods). (c,d) Panels c and d display the trajectory of representative particles in the simulations analysed in panels a and b, respectively. In each panel the particle was chosen at ~5 times of the equilibrium interparticle distance from the centre of the circular simulation domain (black dot) at different polar angles. The trajectories were brought close to the centre for better visualization, and thus the scale bars (indicating 1/3 interparticle distance at equilibrium) apply to the trajectories only. Colour map indicates time.
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