Cell shape affects bacterial colony growth under physical confinement

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Evidence from homogeneous liquid or flat-plate cultures indicates that biochemical cues are the primary modes of bacterial interaction with their microenvironment. However, these systems fail to capture the effect of physical confinement on bacteria in their natural habitats. Bacterial niches like the pores of soil, mucus, and infected tissues are disordered microenvironments with material properties defined by their internal pore sizes and shear moduli. Here, we use three-dimensional matrices that match the viscoelastic properties of gut mucus to test how altering the physical properties of their microenvironment influences the growth of bacteria under confinement. We find that low aspect ratio (spherical) bacteria form compact, spherical colonies under confinement while high aspect ratio (rod-shaped) bacteria push their progenies further outwards to create elongated colonies with a higher surface area, enabling increased access to nutrients. As a result, the population growth of high aspect ratio bacteria is, under the tested conditions, more robust to increased physical confinement compared to that of low aspect ratio bacteria. Thus, our experimental evidence supports that environmental physical constraints can play a selective role in bacterial growth based on cell shape.

Most natural habitats host diverse bacterial communities, which actively respond to extrinsic environmental cues that reflect the dynamic properties of their microenvironments $1,2$. Identifying the basis of these feedback-response cycles is critical for describing the evolution of natural microbial populations, characterizing hostsymbiont crosstalk and interdependency, as well as understanding pathogenesis to design effective antibiotics. A major class of such interactions is understood within the framework of biochemical signalling, where small molecules (such as metabolites, growth factors, antibiotics, and chemoattractants) drive cellular perception and consequent responses $3-6$ $3-6$. These processes also directly impact the composition of natural microbial communities, wherein co-existing species exhibit symbiotic, mutualistic, or predatory interactions with each other^{[7,8](#page-10-0)}. However, current experimental approaches designed to explore such interactions rely on bacteria cultured in homogeneous liquid broths or on surfaces of 2D flat plates, which do not

capture the structural complexity of many natural niches. Bacteria often reside in complex and disordered 3D microenvironments like soil, inter-tissue pores, and biological hydrogels such as mucus, which feature a wide range of material properties $9-11$ $9-11$. For example, the porosity and stiffness of soil is affected by its moisture content and granularity 12 . The spatial architecture of tissues also undergoes significant structural alterations as a result of ECM (extracellular matrix) deposition and enzymatic remodelling by fibroblasts and immune cells 13 . Similarly, the viscoelastic properties of mucus vary with diet, inflammatory responses, microbial enzymatic activity, and pathologies like cystic fibrosis $14-16$ $14-16$ (Fig. [1](#page-1-0)a). Furthermore, bacterial colonies physically constrained within these complex 3D matrices experience diffusion-limited growth^{[17](#page-10-0),18}. In contrast, bacteria growing in homogeneously well-mixed liquid cultures have unrestricted access to nutrients, while colonies on a flat agar surface receive oxygen supply from the top, as well as direct nutrient access from the

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red flour beetle is created in BioRender (M. S. (2024) BioRender.com/y37r627). c By packing highly swollen polymeric hydrogel granules beyond jamming concentrations, we design an in vitro 3D growth medium, which provides a granular and internally porous microenvironment for bacterial growth under confinement. d–f The physical properties of the microgel growth medium are highly tunable based on the mass percentage of hydrogel granules relative to liquid LB (wt

approximately match mucosal samples from natural sources 24 . Rheological measurements shown here either apply d a unidirectional shear at different rates to measure the viscosity, or, e a small amplitude (1%) of oscillatory strain at different frequencies to measure the shear moduli of the 3D growth media. The storage modulus (G) is a measure of the elastic solid-like nature, while the loss modulus (G') signifies the viscous properties of the material. f The porosity of the medium, shown here as a complementary cumulative distribution function (1-CDF) of all the inter-particle pore spaces, determines the degree of confinement, and can be tuned by altering the packing fraction of the hydrogel granules.

underlying substratum. Although diffusion-limited zones can be established during growth over time, these usually affect the colony's core once multilayer 3D growth occurs, with cells piling on top of each other¹⁹. However, growth under 3D confinement could introduce such consumption-diffusion limited zones locally around small single cell-derived colonies, despite there being a global abundance of the nutrients in the bulk medium. Hence, growth under 3D confinement becomes fundamentally different from that on standard 2D interfaces. Previous work has investigated colony growth under nutrient-replete 2D conditions, extending growth-generated stresses as a putative mechanism for achieving quasi-3D conformations $20,21$. However, these studies do not fully capture bacterial growth in disordered, confined microenvironments. Growing evidence indicates that bacterial motility is significantly altered by confinement within a granular and porous matrix 22 and the growth of initially smooth, densely-packed bacterial populations under 3D confinement shows generic surface roughening morphodynamics driven by differential nutrient accessibility¹⁸. Despite this, bacterial population level growth and colony organization under complex and disordered confinement remains insufficiently described. Consequently,

whether and how the physical constraints imposed by a 3D growth microenvironment influence microbial populations remains unclear.

In this work, we present experimental evidence for confinementdependent growth dynamics that selectively favour specific bacterial morphologies. We prepare transparent, granular 3D growth media—a special type of porous matrix for direct measurement of bacterial growth, as well as visualization of colony organisation under different biomimetic degrees of confinement—that broadly mimic the structural and viscoelastic properties of mucus, a biological hydrogel. Using several different bacterial strains isolated from the gut mucus of red flour beetles we show how an increase in confinement confers a growth advantage to bacteria with a higher aspect ratio. Our study combines quantitative measurements of bacterial growth and colony morphology, agent-based modelling, numerical calculations, as well as in vitro co-culture experiments, to establish a generalized principle for growth under confinement. By experimentally manipulating cellular morphology, we also demonstrate a remarkable interconvertible behaviour between high and low aspect ratio forms of the same bacterial species, which strongly implicates single-cell morphology as a broad determinant of colony architecture and growth success under physical

confinement. Importantly, we show that population-level variation in growth dynamics can arise without invoking mechanisms such as genetic mutations, behavioural differences, or cellular responses to biochemical cues. Rather, our work suggests that the composition of heterogeneous microbial populations may be strongly dependent on efficient colony organization under increased confinement that selectively favours bacterial species with a high aspect ratio morphology. These principles are valuable for the experimental elucidation and theoretical modelling of microbial dynamics in complex, spatiotemporally varying natural niches.

Results

Mimicking physical properties of the mucosal microenvironment

To understand how the physical properties of their microenvironment regulate bacterial growth, we avoid the use of laboratory strains to minimize potential experimental artefacts arising from laboratory adaptation, such as mutations accumulated during long-term culture in homogeneous liquid cultures or 2D surfaces. Hence, we isolate several bacterial strains from the gut of the red flour beetle (Tribolium castaneum) and identify them using 16S rRNA sequencing. These bacteria naturally reside in a viscoelastic microenvironment, which ensures minimal interference from culturing practices. Together, these strains include a diverse representation of different envelope properties, metabolic modes, motility patterns, and morphologies (Fig. [1](#page-1-0)b). To create a structural mimic of their natural habitat, we design a jammed microgel-based 3D growth medium that mimics the physical and structural properties of mucus (Fig. [1](#page-1-0)c). Previous structural analysis of mucus indicates different regimes of porosities, spanning from nanometre-scale to micron-scale pores $2³$. Further, rheological measurements on mucosal samples from various sources also reveal a broad range of shear moduli (ranging from ~1 Pa to ~100 Pa) and viscosities (ranging from ~10 mPa.s to ~5 \times 10⁵ mPa.s)²⁴. By packing micronscale granules of hydrogels swollen in liquid LB (Luria-Bertani growth medium) beyond their jamming concentration, we manufacture a soft, semisolid 3D matrix with the key physical properties of mucus—especially the porosity (ψ) , shear moduli (G' and G''), and viscosity (η) (Fig. [1](#page-1-0)d–f and Supplementary Figs. 2b, c). The nanometre-scale internal mesh sizes of the polymer chains forming each hydrogel granule allow unimpeded uniform diffusion of nutrients and small molecules throughout the bulk medium^{[25](#page-11-0)-[27](#page-11-0)}. The microparticles themselves pack together to generate micron-scale inter-particle pore spaces (ψ) , which provide a porous and internally disordered matrix for the bacteria. Within this, bacteria can move by exploring the tortuous pore spaces, as well as achieve 3D colony growth by pushing aside the microparticles to locally and reversibly deform the matrix. Importantly, our 3D microgel growth media retain soft solid-like bulk properties under low shear, while being reversibly fluidizable upon application of shear rates higher than the yield stress limit $28,29$ -which is orders of magnitude lower than the internal turgor pressure of typical bacterial membranes. This allows bacterial colonies to internally rearrange the 3D matrix during growth without altering its material properties, allowing us to reproducibly maintain stable mechanical regimes over long-term culture. Using these platforms, we ask how two different degrees of confinement—representing both extremes of the physical regimes commonly observed for naturally occurring mucus—alter bacterial growth.

Bacterial growth varies with confinement

The 3D growth medium comprises a low solid fraction, which lends it high optical transparency— allowing us to directly visualize bacterial growth under physical confinement. Leveraging this physical property, we homogeneously disperse bacteria in both low confinement and high confinement matrices and employ standard optical density-based absorbance measurements for assaying the temporal change in bacterial biomass (Fig. [2](#page-3-0)a). Optical density measurement (A) in these matrices is linearly proportional to the volumetric density of bacterial cells (λ) as obtained by direct counting from micrographs. Our volumetric counting method is quantitively robust as it correlates well with the conventional colony forming units (CFU)-based assay for identical samples (Supplementary Fig. 2e–g). Interestingly, we observe a decrease in the overall biomass production for all bacterial strains with an increase in confinement (Fig. [2b](#page-3-0) and Supplementary Fig. 3). Prior reports suggest that the growth of bacteria is altered under elevated hydrostatic pressure—well above the turgor pressure—which may lead to deformation of the cell shape and induce stress responses $30,31$ $30,31$ $30,31$. However, since the elastic modulus of the microgel growth media is approximately three orders of magnitude lower than the internal turgor pressure of bacterial cells 32 , the observed reduction in overall biomass production is likely not due to compressive deformation of the cell body. To rule out aberrant effects of elevated osmotic stress on bacterial growth, we culture bacteria in liquid LB broth doped with varying amounts of the bioinert polymer PEG-1000 that induces an increase in osmotic pressure up to 100-fold higher than that exerted by our highest degree of 3D confinement. We find minimal effects on bacterial growth (Supplementary Fig. 4a). We further verify that our 3D matrices do not significantly affect cellular morphology. We negate the possibility of body deformation under 3D confinement by morphometrically analysing the cells; in both low and high confinement matrices, cell morphology remains identical, indicating that growth decrease under higher confinement is not due to a decrease in cell sizes (Supplementary Fig. 5a–c). Further, we show that the observed growth patterns cannot be attributed to cell death by quantifying the cell viability in the microgel growth medium. These results show that growth under high confinement does not detrimentally affect the proportion of viable cells in the population (Supplementary Fig. 5d, e). To further ensure that an increase in confinement does not cause detrimental effects on cellular physiology, we also culture a genetically modified Escherichia coli strain—engineered to visually report both the genome's physical organisation using HU (a DNA-binding protein) tagged to GFP, as well as induction of SulA-driven DNA damage responses—in 3D microgel growth media of both low and high degrees of confinement (Supplementary Fig. 5f). We observe that increase in physical confinement neither alters the HU-GFP localization pattern (i.e., no observable change in the genome's spatial arrangement 33) nor induces SulA expression (indicating DNA damage responses are not triggered³⁴). Hence, the decrease in biomass production under higher confinement is likely not due to altered growth patterns such as those triggered by DNA damage-induced stress responses.

To quantify the impact of confinement on growth, we normalize the biomass achieved at each timepoint in the high confinement matrix ($A_{hichl6h}$) to that in the low confinement matrix ($A_{lowl16h}$) for each strain, which gives the relative biomass ratio (ρ) . This parameter initially varies with time, before achieving a plateau over 12–16 h for all strains, indicating the existence of a general, non-species-specific effect of confinement on bacterial growth (Fig. [2](#page-3-0)c). To focus on these effects at a stable time point, we quantify the endpoint biomass ratios at the 16-h mark and observe that strains cluster into two wellseparated groups—those which perform poorly under increased confinement, and those which remain relatively less perturbed (Fig. [2](#page-3-0)c). For additional rigour, we directly benchmark measurements of endpoint absorbance ratios against volumetric density of bacterial cells as obtained by microscopic counting from the same samples. Importantly, we perform these confirmatory tests for cells possessing two different shapes—approximately spherical and rod-like—thereby establishing the relative biomass ratio (ρ) as a generalised quantitative measure of change in bacterial biomass in 3D matrices (Fig. [2d](#page-3-0)).

A comparison of the various strain-specific morphological attributes listed in Fig. [1](#page-1-0)b reveals motility and cell body shape as distinguishing features of the two clusters observed in Fig. [2](#page-3-0)c. Since motility

Fig. 2 | Change in confinement alters bacterial population growth. a We use absorbance-based optical density measurements as an indicator of an increase in biomass over time to quantify bacterial growth in the 3D microgel media. b Representative absorbance-based growth curves over 16 h indicating confinement-dependent growth dynamics for all bacterial strains (mean \pm s.d., $n = 3$ technical replicates); G' - 10 Pa represents a low confinement matrix and G' - 110 Pa represents a high confinement matrix. c Relative biomass ratio values for all bacterial strains across all time points, showing that higher confinement demarcates the different microbial species into two distinct sub-groups, indicated by bar graphs representing the plateau value of the relative biomass ratio at 16 h (mean ± propagated error, which is calculated as described in the statistical analysis, with biological replicates $n = 7$ for S. gallinarum, $n = 5$ for A. radioresistens, $n = 6$ for *E*. faecium, $n = 6$ for *E*. faecalis, $n = 6$ for *P*. mosselli, and $n = 8$ for *E*. hormaechei). d Absorbance measurements from 3D growth media mirror the actual microbial load, determined by measuring volumetric bacterial density via direct counting of cells using 3D confocal microscopy (also see Supplementary Fig. 2e).

growth (mean ± propagated error, which is calculated as described in the statistical analysis; for A. radioresistens $n = 3$ technical replicates for absorbance measurements; while for the manual counting 317 cells were pooled from three unique micrographs for the low confinement gel, and 198 cells were pooled from four unique micrographs for the high confinement gel—all sourced from the same biological replicate; and for the non-motile E . coli, $n = 4$ technical replicates for absorbance measurements; while for the manual counting 1163 cells and 828 cells were pooled from five unique micrographs each for the low confinement gel and high confinement gel, respectively—all sourced from the same biological replicate). e–g Motility does not confer significantly greater fitness benefits under confinement, as tested using a motile and non-motile strain of E. coli (the latter carries a deletion in the flagellar protein-encoding gene flhDC), which show no significant differences between the growth curves (mean \pm s.d., $n = 3$ technical replicates) or the relative biomass ratio (mean \pm propagated error, $n = 5$ biological replicates).

can conceivably allow better dispersion of bacteria in the 3D matrix 2^2 thereby improving overall population-level growth through better nutrient accessibility and lesser local crowding—we first test the impact of motility on enhancing growth by employing both a motile (wild-type) and non-motile strain of E. coli (harbouring a deletion of a flagellaencoding gene). Surprisingly, both strains perform similarly under different degrees of confinement as captured by their relative biomass ratio (ρ) (Figs. 2e–g and Supplementary Fig. 5g, h), invalidating motility as the dominant feature mediating the growth patterns observed under increased confinement. Consequently, we revisit the set of speciesspecific morphological attributes and note that while all poorlyperforming strains comprise low aspect ratio (approximately

spherical) cells, the better-performing strains have high aspect ratio cells (rod-like). These effects are not due to different biological species possessing intrinsically different growth rates. Although we observe that the absolute growth of certain coccoidal species is higher than some bacilli in the low confinement matrices, the relative biomass ratio metric specifically captures the performance of a given strain across two different degrees of confinement and is hence agnostic to how any other species fares. Using this, we observe a specific trend favouring a particular class of bacteria over others only when analysed along the shape axis. This observation forces a significant realignment of our perspective on growth under confinement, implicating cellular morphology as the attribute critical to growth success under spatial constraints.

Single-cell morphology dictates colony shape

Unlike growth on homogeneous 2D interfaces or in well-mixed liquid media, growth under physical confinement imposes spatial constraints and consequent nutrient limitations on bacterial colonies. To determine how different single-cell morphologies affect colony architecture, we employ agent-based simulations using the CellModeller platform³⁵, maintaining all growth parameters constant and only altering the cellular aspect ratio. We simulate growth undertaken by low aspect ratio (approximately spherical) and high aspect ratio (rodshaped) cells (Fig. [3](#page-5-0)a and Supplementary movie 1). While most high aspect ratio bacilli (rod-shaped) cells such as E. coli exhibit cell division patterns that produce progeny cells oriented along the parental cell's longitudinal axis, several low aspect ratio coccoid (spherical) species show planes of division that are either randomly or perpendicularly positioned to previous division planes, which allow a higher degree of freedom for the orientation of progeny cells following division³⁶. We incorporate these patterns in our simulations by implementing that high aspect ratio bacteria divide and produce progeny oriented along the length of the parent cell, whereas low aspect ratio bacteria produce progeny oriented along a randomly chosen direction. Under these conditions, striking differences emerge, wherein the high aspect ratio bacteria generate elongated colonies, while the low aspect ratio bacteria give rise to rounded 3D colonies. We further characterise these using a parameter termed the shape factor (ϕ) , defined as the ratio between the perimeter and square root of area for the projection of a given colony. A low value for the shape factor (ϕ) indicates that the colony has a relatively lower exposed surface area as opposed to the enclosed volume. We find that the low aspect ratio bacteria predominantly organise into colonies with a low shape factor, and the high aspect ratio bacteria form elongated, high shape factor colonies

Fig. 3 | Bacterial morphology determines colony architecture and competitive outcomes. a We perform agent-based simulations of 3D colony growth for both high and low aspect ratio cells. In these simulations, the progeny of high aspect ratio cells exhibit a directional bias along the parental axis, whereas those of low aspect ratio cells are more flexible to grow along any random axis. **b** Rod-shaped high aspect ratio E. coli cells give rise to elongated colonies under confinement as they proliferate by pushing their progeny outwards (similar phenomena have been observed for at least three biological replicates; representative micrograph has been shown from a replicate with $n \geq 40$ distinct individual colonies imaged). c Treatment with 5 μg/ml of the MreB-perturbing compound A22 produces approximately spherical E. coli cells, which result in a rounded colony architecture (similar phenomena have been observed for at least three biological replicates; representative micrograph has been shown from a replicate with $n \geq 10$ distinct individual colonies imaged). d Naturally spherical E . faecalis cells, stained here with Calcein-AM green, show division and colony formation patterns similar to that predicted by the simulations for low aspect ratio cells, as well as the drug-treated spherical E. coli cells (similar phenomena have been observed for at least three biological replicates; representative micrograph has been shown from a replicate with $n \geq 50$ distinct individual colonies imaged). Micrographs in **b**-d are generated by capturing the front view of a 3D volumetric projection. e, f Quantitative analysis of colony morphology using the shape factor (ratio between perimeter and square root of the area obtained from 2D projections of colonies) shows a significant difference between high and low aspect ratio cells, for both computational simulations ($n \ge 200$ snapshots, from ten simulations) and experiments ($n \ge 200$

(Fig. 3e and Supplementary Fig. 6d, e). Hence, single-cell morphology appears to strongly dictate overall colony architecture.

We experimentally test this by live imaging the 3D colony growth of non-motile rod-shaped E. coli under high confinement matrices and observe patterns consistent with our computational model. Rodshaped E. coli cells undertake volumetric growth along the parental axis, and push their progeny cells outward, leading to elongated colony architecture (Fig. 3b, Supplementary Fig. 6a, and Supplementary movie 2). Next, to test these patterns using a spherical bacterial species, we fluorescently label E. faecalis cells using Calcein-AM Green. Time-lapse imaging shows that these coccoid cells also follow a colony growth mode similar to that predicted for low aspect ratio cells by the simulations, by organizing into well-rounded colonies (Fig. 3d and Supplementary Figs. 6c and 8d). Further, we directly alter the morphology of non-motile E. coli using sub-MIC dosages of the MreBperturbing compound A22³⁷. Sub-lethal concentrations (5 μ g/ml) of the drug induce aberrant division, rendering the rod-shaped E. coli into spherical cells. Again, as predicted by our computational model, the A22-treated low aspect ratio cells assume a rounded colony morphology (Fig. 3c, Supplementary Fig. 6b, and Supplementary movie 2). By quantitatively analysing the shape factor (ϕ) of the max projected fluorescent micrographs of colonies formed by rod-shaped and spherical cells, we find both the experimental and computational data to be in good agreement, with the single cell morphology being strongly implicated as the key determinant of colony-level organisation under 3D growth conditions (Fig. 3f).

Confinement selectively favours rod-shaped cells

A comparison of the relative biomass ratio (ρ) for rod-shaped and spherical E. coli reveals that alteration in shape severely compromises growth success under increased confinement, mirroring the trends exhibited by high and low aspect ratio gut-derived bacteria. Under similar concentrations of drug treatment (5 μg/ml A22), no significant growth differences are observed in liquid media. (Figs. 3g, h and Supplementary Fig. 6f, g). In fact, the relative biomass ratio of the spherical *E. coli* is quite similar to that of the naturally low aspect ratio cells (approximately $\rho \le 0.5$). Together, these observations highlight how cellular morphology enforces differences in the colony architecture of rod-shaped and spherical cells during growth under confinement. We further find that these trends depend only on the cellular individual colonies). g Shaken liquid cultures in LB media for both rod-shaped E. coli and A22-treated spherical E. coli cells do not show any significant differences in their overall growth (mean \pm s.d., for $n = 3$ technical replicates). **h** Relative biomass ratios for rod-shaped and spherical E. coli indicate that rod-shaped cells are less affected than spherical ones under increased confinement, i.e., have ρ values closer to 1 (mean ± propagated error, which is calculated as described in the statistical analysis, with biological replicates $n = 5$ for rod-shaped non-motile *E*, *coli* and $n = 4$ for A22-treated spherical non-motile E , coli). **i**, **j** Direct visualization and endpoint counting of cells from 3D co-culture assays including both low aspect ratio (A. radioresistens or E. faecalis) and high aspect ratio cells (motile E. hormaechei or nonmotile E. coli) show that the latter enjoy a growth advantage under increased confinement, at the expense of the former. However, competitive cultures between two high aspect ratio strains do not show significant enrichment of either strain (mean \pm s.d., for *E. hormaechei* vs *E. faecalis, n* = 2 biological replicates, each with ≥400 individual cells pooled from ≥10 unique micrographs for each gel type; for non-motile E. coli vs E. faecalis, n = 2 biological replicates, each with ≥300 individual cells pooled from five unique micrographs for each gel type; for non-motile E . coli vs A. radioresistens, $n = 3$ biological replicates, each with ≥180 individual cells pooled from ten unique micrographs for each gel type; for non-motile E. coli vs E. hormaechei, $n = 2$ biological replicates, each with ≥130 individual cells pooled from ten unique micrographs for each gel type). k Analysis of genomic DNA extracts from similar 3D co-culture setups confirms that the growth of high aspect ratio bacteria is more robust to increase in physical confinement compared to that of low aspect ratio cells ($n = 3$ biological replicates).

shape rather than other environmental variables. To test this hypothesis, we first alter the nutrient composition for the 3D matrices from rich LB broth to minimal media (M9) supplemented with glucose and amino acids (Supplementary Fig. 7a, b). When cultured in the minimal media matrices, high aspect ratio E. coli and low aspect ratio A. radioresistens mirror the relative biomass ratio trends observed in LBbased matrices, indicating that the relative biomass ratio patterns are independent of nutrient composition (Supplementary Fig. 7c–e). The above results lead to the prediction that all else being equal, rodshaped bacteria should have a growth advantage over spherical bacteria under higher confinement. We additionally verify this using numerical calculations of bacterial colony growth, wherein we consider two different colony architectures—ellipsoidal and spherical formed by high aspect ratio rod-shaped bacteria and low aspect ratio spherical bacteria, respectively. The effective local concentration of a nutrient inside a 3D bacterial colony is a result of two processes: replenishment of small nutrient molecules via unimpeded diffusion and depletion of nutrients via consumption. The competition between consumption and diffusion sets a depth of 10 μ m from the surface of a 3D bacterial colony beyond which the nutrient is completely depleted^{[18](#page-10-0)}. Factoring in this previously established limit on effective nutrient availability inside a 3D bacterial colony—which sets a limit on the actively proliferating fraction of a given 3D bacterial colony—we estimate the proportion of actively growing bacterial cells in colonies of either geometry with different sizes (Supplementary Fig. 6h, i). Ellipsoidal colonies comprising rod-shaped cells retain a higher proportion of cells in the actively growing zone as opposed to spherical colonies of comparable sizes. Together, the above results indicate that high aspect ratio bacteria produce elongated colonies, which allow better nutrient access in a 3D matrix, thereby enabling superior growth under increased confinement.

To test whether this effect provides a competitive advantage to high aspect ratio cells in a mixed population, we carry out co-culture assays that combine both high aspect ratio and low aspect ratio cells across two different degrees of confinement (Fig. 3i–k and Supplementary Fig. 8a–c). For this, we prepare an inoculum with equal parts of two different bacterial species that we homogeneously disperse in both low and high-confinement 3D growth matrices. More specifically, we perform competitive co-culture assays between the following five combinations: (i) high aspect ratio motile E. hormaechei

and low aspect ratio non-motile E. faecalis, (ii) high aspect ratio nonmotile *E. coli* and low aspect ratio non-motile *E. faecalis.* (iii) high aspect ratio non-motile E. coli and low aspect ratio non-motile A. radioresistens, (iv) high aspect ratio motile *E. hormaechei* and high aspect ratio non-motile E , coli, and (v) high aspect ratio motile P . mosselii and high aspect ratio non-motile E. coli. Following 16 h of growth, we mechanically disrupt the colonies in each matrix into homogeneously dispersed single-cell suspensions. Subsequently, we tally the number density of each type of bacteria using microscopy to obtain the relative strain ratios for each competing pair. As expected, the relative proportion of high aspect ratio cells in the population significantly increases with an increase in confinement, at the cost of a large decrease in the abundance of low aspect ratio cells (Fig. [3i](#page-5-0), j and Supplementary Fig. 8a). Interestingly, an increase in confinement does not alter the relative abundance of two high aspect ratio species in direct competition (Fig. [3](#page-5-0)j and Supplementary Fig. 8b). Parallelly, we also perform bulk genomic DNA extraction and quantify the relative abundance of each strain using 16s rRNA amplicon sequencing. Our bulk genomic analysis qualitatively mirrors the trends observed with microscopic tallying of the number density: despite identical nutrient media, rod-shaped bacteria gain a substantial advantage over spherical bacteria under high confinement (Fig. [3](#page-5-0)k and Supplementary Fig. 8c). These results thus validate high aspect ratio morphology as a beneficial attribute under spatially confined growth. Of course, in many ecological settings, inter-species interactions are also heavily influenced by cross-feeding, cooperative and antagonistic behaviours, fluctuations in the environment, resistance to invasion, predation, and chemical cross-signalling^{[3](#page-10-0)-[8](#page-10-0)}. Our platform presents the capability to conserve the heterogeneous chemical and behavioural landscapes introduced by such variables, and study how physical confinement acts on top of these processes. This is because the porous nature of our matrix allows for unimpeded small mole-cule diffusion^{25–[27](#page-11-0)} regardless of the degree of confinement, thus ensuring that modulation of the mechanical regime can occur independent of all other factors.

In nature, cells exhibit morphological alterations under specific environmental conditions. For instance, uropathogenic E. coli transition between coccoid and rod-like morphologies during infection progression³⁸. These transitions are also subject to chemical signalling and nutrient availability. These are best illustrated by examples such as those of V. cholerae, which shows a curved rod-to-sphere transition in response to both low temperatures or L-arabinose treatment, as well as L. monocytogenes that undergoes a similar transition during biofilm formation on surfaces $39-42$ $39-42$ $39-42$. Such shape changes likely confer survival advantages under suboptimal conditions such as extreme environmental changes or nutrient starvation. However, it is also conceivable that a sphere-to-rod transition might prove particularly beneficial purely in terms of nutrient availability in porous 3D environments such as tissues and soil by promoting more efficient growth and proliferation. We test this hypothesis using P. sanguinis, a species wherein the spherical form can grow in an elongated fashion as a rod—similar to previous reports on P. denitrificans^{[43,44](#page-11-0)} (Fig. 4a, Supplementary Fig. 8e, f, and Supplementary movie 3). We assay the growth of P. sanguinis across two different degrees of confinement and find that it mimics the relative biomass ratio trend observed for high aspect ratio bacteria. This further validates that the elongated colony structure of high aspect ratio strains confers a relative advantage under increased degrees of confinement.

Thus, we conclude that the aspect ratio of individual cells is a key determinant of the relative biomass ratio under 3D confined growth. To test this, we measure the cell body morphology of all the different bacterial strains used in our experiments and plot their individual aspect ratio by measuring the major axis and minor axis of the cell body. Distribution of these aspect ratios captures two different populations, separated by their morphology (Fig. 4b). Superimposing

Fig. 4 | Single-cell morphology enables physical confinement to selectively favour elongated colony architecture. a Growth assays indicate that P. sanguinis performs at par with rod-shaped bacteria across two different degrees of confinement (mean \pm s.d., $n = 3$ technical replicates). **b** Measurements of the aspect ratios of bacterial cell bodies for all ten strains used in this study, measured from samples in the late exponential phase ($n \geq 80$ individual cells represented as lognormal distribution, with boxes indicating the means, and bounds of the boxes indicating \pm s.d.). c Microbial growth under confinement, as described using a phase space defined by the relative change in biomass production under increased confinement (the relative biomass ratio) vs single cell morphology (mean ± propagated error for the relative biomass ratio, and mean ± s.d. for the single cell aspect ratio).

this with our previous results on the relative growth ratios across two different degrees of confinement, we observe a remarkable compartmentalization of the different bacterial strains into two nonoverlapping domains. This phase space—defined by single-cell morphology and consequent growth performance under elevated confinement—underscores the idea that with an increase in confinement, single-cell morphology is the primary determinant of growth success in complex environments (Fig. 4c). Effectively, we demonstrate that the selective pressure exerted by the physical microenvironment in terms of confinement-induced spatial constraints strongly favours high aspect ratio cellular morphology due to its efficient colony organisation.

Discussion

Our work provides experimental evidence implying that physical confinement can play a selective role in deciding bacterial growth

fitness within their natural niches. High aspect ratio bacteria leverage growth anisotropy to form elongated colonies with higher surface area, allowing better nutrient accessibility as opposed to the spherical colonies formed by low aspect ratio bacteria. Increased confinement strongly selects high aspect ratio cells for more efficient growth—a trend that holds even for high and low aspect ratio forms of the same bacterial strain, indicating a remarkable interconvertible behaviour governed solely by cellular shape. Importantly, we demonstrate the potential to use such constraints to alter bacterial community composition under biomimetic mechanical regimes. Thus, our results represent an important conceptual advancement towards experimentally modelling bacterial growth within complex natural habitats.

In microbial ecology, the present mechanistic understanding of predator-prey interactions, niche partitioning, and community organization largely relies on a framework comprising genetic mutations, behavioural patterns, and biochemical signalling⁴⁵⁻⁴⁹. Notably, our platform does not preclude such chemical interactions. Instead, the porous and permeable nature of our 3D matrix presents an opportunity to capture the dynamics of a chemical signalling landscape coupled with different mechanical regimes, enabling a more comprehensive experimental recapitulation of natural niches. While the permeable nature of our 3D matrix allows unimpeded small molecule diffusion, several biological niches such as soil and aquifers resemble semi-permeable porous media with non-deformable, rigid physical barriers. We expect that the reduced nutrient availability and severe spatial restrictions in such contexts significantly restrict colony growth and organization. Recreating such regimes in vitro will require the development of semi-permeable 3D scaffolds. Further, within the temporal window of our assays, we do not observe significant contributions from either aberrant phenotypes or perturbed physiologies. However, environmental pressures in the form of nutrient deprivation, high mechanical stiffness, and antagonistic interspecies interactions are known to induce the expression of stress genes, alter metabolism, trigger extreme phenotypes (such as filamentous growth), and favour increased mutational rate^{[50](#page-11-0)–[55](#page-11-0)}. Such factors are likely to affect long-term bacterial growth— hence, future work that investigates bacteria under similar conditions over extended durations could provide interesting insights into the long-term effects of confinement on bacterial communities. Presumably, our system can be engineered to recreate such effects by altering the nutrient media composition and tuning the mechanical stiffness of the matrix. We speculate that such environmental constraints will selectively favour a subset of adaptive mutations with the potential to alter evolutionary trajectories.

Furthermore, it has been reported that spatial patterning within mixed microbial communities can be strongly driven by single-cell morphology⁵⁶. This suggests an interesting direction for future exploration. Given the selective pressure enforced by elevated physical confinement favouring rod-shaped bacteria over spherical ones, how do existing spatial patterns get altered when the microenvironmental mechanics change? Conversely, does physical confinement alter the evolution of spatial architecture within mixed microbial communities such as biofilms? Finally, our work highlights the need to explore the role of physical confinement in diverse niches that may be selected for varied life history strategies. For instance, short-term rapid growth and expansion are not universally preferred traits. In adverse conditions such as in the presence of toxins or predators when longterm cell survival is prioritized, a transient shift towards quiescence or dense collective organisations with minimally exposed surface areas may be more advantageous $57-61$ $57-61$ $57-61$. Such conditions and strategies may override the selection of cell shape imposed by physical confinement and should be explored further.

Current descriptions of microbial growth invoke either genetic mutations or biochemical signalling as the primary mechanisms via which bacteria respond to their microenvironment. By contrast, we argue that the spatial constraints imposed by their microenvironment exert a more general effect, irrespective of specific organismal biology. Our conceptual framework can be employed to generate coarsegrained predictions of population dynamics under different biophysical regimes. Understanding how bacteria grow in complex, disordered 3D environments such as tissues, mucus, and soil, is also central for combating antibiotic resistance, improving agricultural practices, and bioremediation—underscoring the high-value practical applications of our work.

Methods

Preparation and characterization of 3D growth media

We prepare the 3D matrices by dispersing dry hydrogel granules of Carbopol C-980 (Lubrizol) at either a 0.50% or 0.85% $(w/v\%)$ concentration in 2% ($w/v\%$) liquid LB (Sigma, LB Lennox) media. While preparing minimal media matrices, we replace LB media with M9 media (BD Difco, M9 Minimal Salts) supplemented with 0.2% glucose and 0.2% milk protein hydrolysate as a source of amino acids. The hydrogel suspension is vigorously stirred for at least 2 h to ensure complete hydration of the dry granules. Since the Carbopol C-980 is a negatively charged hydrogel, we neutralize the pH of this suspension to 7.4 using 10 N NaOH. At pH 7.4 the hydrogel granules or microgels swell maximally to an extent when individual granules jam against each other forming a transparent and disordered solid matrix. We determine the viscoelastic properties of jammed microgels using oscillatory shear rheology on an Anton Parr (MCR302e) rheometer with a roughened cone-plate measuring tool. We adjust the solid fraction of 3D growth media to achieve an elastic storage modulus range between 10 Pa and 110 Pa, that spans the measured range of biological mucosal samples. To measure the yield stresses and viscosity of the hydrogels, we apply unidirectional shear at varying rates while recording the shear stress response. As shown in Supplementary Fig. 2b, our microgels behave as yield stress materials, which are reversibly fluidizable upon application of the threshold shear rate but retain a soft solid-like nature below this regime. The porosity of the microgels is determined using 200 nm fluorescent tracer particles. We track the thermal diffusion of these beads through the inter-particle pore spaces. Since the internal mesh sizes of the hydrogel granules are smaller than 100 nm, the beads are occluded from these regions. Hence, the beads only explore volumes that are also accessible to bacteria (which are up to 10-fold larger in dimension than the bead size). We disperse a dilute solution of the tracer particles in both low and high-confinement matrices. Subsequently, we acquire images within a time interval of 17 ms using the 40× objective of an inverted laser-scanning confocal microscope (Nikon A1R HD25) to track their thermal diffusion-guided trajectory. Individual beads are tracked with sub-pixel precision using a custom-written MATLAB script based on the Crocker-Grier algorithm⁶², and the mean-square displacement (MSD) is quantified as a function of lag time. Over short time scales, the particles freely explore the pore spaces. However, over longer time scales, their motion becomes constricted by the porous medium and its dead-end channels. This implies that the particle MSDs will increase linearly with time for unimpeded motions, while over longer time scales the pore boundaries will make the MSDs plateau. The characteristic smallest pore space dimensions explored by the particles are then given by the square root of these plateau values added to the particle diameter. We calculate the 1-CDF (complementary cumulative distribution function) of the measured pore spaces, which represents the overall pore size distribution within our 3D microgel growth media. Herein, the value of the 1-CDF at any given pore size represents the fraction of the pores which are larger than the given pore size. For instance, a 1-CDF value of 0.5 represents the 50th percentile of pore sizes. Similarly, a 1-CDF value of 0.05 denotes that 5% of the pores are larger than the corresponding pore dimension. (Supplementary Fig. 2c). We also show that under the culturing conditions followed in

this work, our microgel samples do not undergo any significant alterations to the rheological properties or bulk volumes. For this, we take the low confinement matrix, and maintain it at 30 °C for up to 24 h under humid conditions (Supplementary Fig. 2a). This is similar to the assay set-up for all our biological experiments. Further, we recover and pool together the microgel samples from these plates and carry out rheological measurement, as well as direct weighing to establish that both the rheological properties and the overall sample volume remain conserved (Supplementary Fig. 2h–j).

Isolation, in vitro culture, and characterization of bacterial strains

We isolate seven different microbial species from red flour beetles (T. castaneum; the red flour beetles were collected by DA from a flour warehouse in India and have been maintained by DA for ~10 years at NCBS) and identify these by Sanger sequencing of the 16S rRNA gene using the universal 16S primers 27 F (5'-AGAGTTTGATCCTGGCTCAG-3'), and 1492 R (5'-GGTTACCTTGTTACGACTT-3'). Following the alignment of both forward and reverse sequences, a homology search using NCBI BLAST is carried out to identify the closest bacterial species for each such amplicon (16S sequences provided for each isolate in Supplementary Table 1). Primary isolates from these are cultured in 2% LB broth and cryopreserved as glycerol stocks $(20\% v/v)$. For each experiment, a shaken culture of 1% stock dilution in LB broth is grown at 30 °C for ~20 h to achieve a stationary phase culture. From this, a 1% inoculum is added to fresh 3D growth media and homogeneously dispersed by gentle pipette mixing. To promote better growth in the case of E. faecalis and E. faecium, overnight cultures are directly set using 3D hydrogel media. We also employ two strains of *E. coli*-one, the wild-type motile strain, and the second carrying a deletion in the flagellar gene flhDC, that renders it non-motile. These E. coli strains were a kind gift from Prof. Robert Austin, of Princeton University, which constitutively express cytoplasmic green fluorescent protein (GFP) and have been previously used to study bacterial motility in 3D porous media^{22,25}. For antibiotic treatment of E , coli with A22, we maintain a sub-MIC concentration of 1 μg/ml in LB and 5 μg/ml in 3D media. To characterize the bacterial morphology, we dilute samples using either LB or 0.5% 3D growth media and spread these on a flat agar pad $(1\% (w/v)$ agar in water) placed inverted onto a glass bottom dish. Images are acquired using a point-scanning laser confocal microscope. To measure the cellular morphology, we segment micrographs in ImageJ and use in-built functions to quantify the cell body area, length, and aspect ratio.

We ascertain the suitability of LB broth as a nutrient medium for all bacterial strains by assaying their growth in shaken liquid cultures (Supplementary Fig. 1). We separately assess the growth performance of these strains under non-shaken culturing conditions in liquid LB, results from which show that static culturing conditions in liquid media lead to less efficient growth for most microbial strains used in our study (Supplementary Fig. 4b). While shaken culture ensures well-mixed condition for uniform nutrient and oxygen availability throughout the bulk culture volume, in non-shaken culture progressive densification may occur over time due to cells settling under gravity. In contrast, physical confinement manages to maintain an initially homogeneous distribution of bacterial density throughout the bulk 3D volume of the material, while nutrient availability is also similar throughout due to the freely permeable and porous nature of the microgel matrix. Hence, both shaken and nonshaken culturing conditions are fundamentally different from the conditions encountered by bacteria embedded in our 3D matrix and hence growth performance in these conditions should not be compared.

We also quantify the viability of cells grown in the high confinement matrix. In all samples, we use propidium iodide (fluorescence in red) as a marker for dead cells, since this compound stains cells with compromised membrane integrity. We also use Calcein-AM Green as a marker for viable cells in the case of *E. faecalis* which shows good uptake of this compound. For the E. coli strains, we use the constitutively expressed green fluorescent protein signal as a marker for viable cells. For viability testing, we culture different bacterial strains in the high confinement matrix for 24 h, following which we dilute the samples in liquid LB, vigorously mix using vortexing to break apart colonies into single cells under mechanical shear, and incubate with the appropriate staining solutions, before spreading them on flat agar pads for image acquisition. To test for genomic DNA damage-triggered responses, we use E. coli strains carrying a constitutively-expressing GFP-tagged histone protein (HU-GFP) that marks the DNA, along with mCherry-tagged DNA damage response-triggered protein (SulAmCherry), received as a kind gift from Dr. Anjana Badrinarayan of NCBS. These are cultured for 24 h in microgels of different stiffnesses and imaged using a confocal microscope. As an additional control, we induce DNA damage in an exponential-phase culture of bacterial cells cultured in liquid LB by treating with 5 μg/ml of the DNA damageinducing drug MMC, which exhibits strong SulA-mCherry expression.

Growth assays

We assay microbial growth in 3D growth media using absorbance-based optical density measurements at 600 nm. Samples prepared as described above are dispensed as 200 μl replicates into 96-well plates, and readings are acquired at 10-min intervals for 16 h using a multi-mode plate reader (Varioskan Lux) with temperature control (set to 30 °C). All data are normalized by subtracting the initial timepoint value (except for antibiotic-treated samples, where we normalize using the lowest absorbance value to account for initial cell death). This gives a measure of the net increase in biomass over time. For 3D co-culture assays, all parameters are maintained identical to the above, except the initial inoculum, wherein both competing strains are added in similar proportions by normalizing the optical density of overnight-grown cultures. To maintain identical culturing conditions as followed for the individual growth assays, we dispense these samples in a 96-well plate and grow them for 16 h at 30 °C. Following this, we pool together all the technical replicates for each gel type and homogeneously mix the sample. From this, a 20 μl aliquot is used for direct bacterial counting, with the rest being processed for genomic DNA extraction (described below). For growth experiments where we vary the increase in osmotic pressure, we use varying amounts of the inert polymer PEG1000 doped in liquid LB. Here, we maintain the elevation in osmotic pressures to levels similar to the high confinement matrix (~110 Pa), as well as 10-fold and 100-fold higher. The osmotic pressures of these solutions are approximated using the equation $\pi = i cRT$, where π represents the osmotic pressure, i denotes the van't Hoff index, c is the molar concentration of solute, R is the ideal gas constant, and T is the temperature.

Direct microscopic counting of single cells

Briefly, we set up growth curve assays for the non-motile E. coli ΔflhDC, as described above, in multiple triplicate sets, for both low confinement and high confinement matrices. At specific time points separated by 3-h intervals, we obtain an optical density measurement for each set. Each individual replicate belonging to this set is harvested, mechanically sheared to break up colony-like clumps into single cells and diluted using LB (Supplementary Fig. 2d). We scale the dilution factor for each timepoint following the first one to achieve approximately the same final density of bacterial cells. These samples are then mixed with microgel (to prevent cells from settling during the counting procedure) in a 1:1 ratio, following which 200 μl of this sample is dispensed into a glass-bottomed 35 mm dish and fixed 3D volumes from each are imaged using a confocal microscope. The total number of bacteria is tallied using automated thresholding and segmentation functions available in ImageJ, as well as the 3D Cell Counter plugin. Using this, we calculate the number of bacteria per ml from the original

sample and plot this against the measured absorbance value. We observe a roughly linear relation between the microscopically counted number of cells and the optical density of the sample, confirming that absorbance values provide an accurate estimate of the total bacterial biomass in 3D growth media (Supplementary Fig. 2e). We also directly benchmark the volumetric counting method against the conventional colony forming units (CFU)-based quantification method. Herein, we prepare samples of bacteria inoculated in the microgel growth media. These are diluted in liquid LB and imaged using confocal microscopy to obtain the volumetric cell density as described above. Following this, the same samples are serially diluted using liquid LB and plated on LB agar. Colonies resulting from these are counted the next day and tallied against the statistics obtained from direct volumetric counting. We find a good agreement between both these metrics, which together with the optical density vs volumetric counting approach, establish that direct counting using microscopy is a quantitatively rigorous method which compares well against both optical density and CFUbased quantification methods (Supplementary Fig. 2f, g).

To verify this for the relative biomass ratio metric, we set up growth curve assays for both A. radioresistens and the non-motile E. coli ΔflhDC, as described above, in both low and high confinement matrices. Immediately after the growth curve assay concludes, we pool together all replicate samples of each bacterial strain and each specific confinement type in separate microcentrifuge tubes. This is well-mixed to ensure homogeneous dispersion of the bacteria. Twenty microlitres of this is added to 1 ml of 0.50% microgel (to prevent cells from settling during the counting procedure) and mixed thoroughly to break up the colonies into single cells. Two hundred microliters of this sample are dispensed into a glass-bottomed 35 mm dish and imaged using a confocal microscope. From the images, individual bacteria are microscopically identified and tallied. We then compare the ratio of endpoint absorbances from 0.85% microgel to 0.50% microgel for both bacterial strains with the ratio of numbers obtained by direct counting for identical samples.

For the co-culture experiments, we prepare an inoculum with equal parts of both high aspect ratio cells (motile E. hormaechei or nonmotile E. coli) and low aspect ratio cells (either A. radioresistens or E. faecalis), that we homogeneously disperse in both low and high confinement 3D growth matrices. Following 16 h of growth, we mechanically disrupt the colonies in each matrix into homogeneously dispersed single-cell suspensions. Subsequently, we spread these on flat agar pads (in the case of the E. coli vs A. radioresistens combination, we also include one replicate with volumetric 3D cell counting) and tally the number density of each type of bacteria using microscopy to obtain the relative number of cells belonging to each strain type per sample.

Colony morphology experiments

For live-cell imaging of bacterial colonies, we use 35 mm plastic dishes with a central circular cavity (15 mm in diameter) sealed off with a glass coverslip for optimal quality imaging. Samples for assessing the colony morphology are prepared exactly as described above for the growth curve experiments. While the E . coli strain used in this study shows constitutive expression of the green fluorescent protein, we use the cell viability dye Calcein-AM Green to stain E . faecalis cells (for live imaging, the dye is directly added to the microgel growth media). Two hundred microlitres of well-mixed samples are dispensed into the glass-bottomed cavity of these dishes. Using a sterile coverslip, we scrape the top of this sample to ensure a smooth, flat layer. To prevent evaporative losses, we gently layer the top of the sample with 1.5 ml of mineral oil (Sigma). This oil is permeable to gases; hence the microgelsuspended bacteria are not cut off from oxygen supply. Plates prepared in this manner are placed in an incubator maintained at 30 °C for 16 h. Following this, we acquire z-stack images of the bacterial colonies using confocal microscopy. Colony morphology is analysed by segmenting 2D projections of the individual bacterial colonies, followed by calculation of the colony area and perimeter using in-built functions in Fiji-Imagel. Using this, we calculate a parameter termed the shape factor, defined herein as the ratio between the perimeter and the square root of the area (for a circle, the shape factor equals 3.54). This quantity gives an estimate of how much colony shapes differ across various species and treatment conditions.

DNA extraction and 16S rRNA amplicon sequencing

For genetic analysis of 3D media-grown and liquid (t_0) , initial inoculum) co-culture samples, we carry out genomic DNA extraction. Three hundred microliters of each sample is incubated at 65 °C for 2 h with 200 μl of Promega nuclei lysis buffer, followed by the addition of 200 μl Promega protein precipitation solution and 100 μl of chloroform to promote phase separation. The DNA-containing fraction is separated out by centrifugation, to which 300 μl isopropyl alcohol is added to precipitate the nucleic acids fraction. The resultant pellet is washed twice with 80% ethanol, following which we elute the genomic DNA in 40μ l nuclease-free water. To determine the change in the relative proportion of the two co-cultured strains, we amplify the V3–V4 hypervariable region of the bacterial 16S rRNA gene using the modified Illumina 10N primer 63 and sequence amplicons on the Illumina Miseq platform (300 × 2 paired-end, obtaining ~75,000–140,000 reads per sample). We analyse sequence data using the DADA2 pipeline 64 in R, which gives us the relative abundance of each strain in each sample.

We also note that the trends observed using 16S rRNA amplicon sequencing-based methods are largely qualitative in nature. This is primarily due to experimental limitations with designing efficient cell lysis protocols for both competing strains, as well as difficulties with optimal DNA extraction from cells embedded in 3D microgel matrices (wherein, cross-reactivity and charge interactions of the extraction reagents with the scaffold material cannot be entirely discounted). In this regard, the flexibility afforded by the matrix in terms of cell extraction and visualisation allows for parallel strategies such as direct counting of individual cell types, enabling a precise estimation of relative bacterial loads even from co-culture setups. However, direct visualisation and counting also come with certain limitations. For instance, this strategy is only amenable for bacteria whose colonies can be sufficiently disrupted using mechanical shear without lysing the cells. Second, unless the competing strains are unambiguously distinguishable based on morphology (rod vs sphere) or visual contrast (e.g., fluorescently tagging/dye-based labelling or differential colourimetric staining), counting is not a reliable method. This limits its applicability for a select few combinations of bacterial strains. We believe that further work towards optimizing both 16S rRNA amplicon sequencing and visualisation-based approaches, as well as developing hybrid workflows on a case-by-case basis will significantly improve the applicability of our 3D microgel matrix towards enabling a rigorous quantitative estimation of species-specific bacterial loads from mixed microbial communities cultured in vitro in 3D.

Simulating bacterial growth

We simulate 3D colony growth using a previously published agentbased platform (CellModeller³⁵) which models bacterial cells as spherocylinders. We modify the original code by altering the aspect ratio of these objects by explicitly defining the radius and overall volume, which together adjust the cell's length. Using this, we simulate 3D growth of either high aspect ratio "rods", or low aspect ratio "spheres". We trigger the cells to divide once they achieve a target volume greater than twice the defined volume for a single cell, with both resultant daughter cells starting off with the initial volume defined for a single cell. Further, we constrain the progeny cells produced by rods to maintain orientation along the parental axis, whereas the progeny cells produced by spherical cells are allowed to reorient in random directions. We also simulate different degrees of physical confinement by varying the environmental drag (Γ) on the growing bacterial cells.

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Statistical analysis

For each strain of bacteria, we record the $OD₆₀₀$ for a minimum of four sets of biological replicates in both low-confinement and highconfinement matrices. Each set of biological replicates for each type of matrix contains OD_{600} data from 2 to 4 technical replicates. For simplicity, consider the data from high confinement to have a mean A_{hich} and standard deviation dA_{hich} , while for low confinement this would be A_{low} and dA_{low} , respectively. This can be considered as two random variables A_{high} and A_{low} , with errors of dA_{high} and dA_{low} , respectively. Now, the relative biomass ratio (ρ) can be written as:

$$
\rho = A_{\text{high}} / A_{\text{low}} \tag{1}
$$

where $d\rho$ is the propagated error from the technical replicates of both A_{high} and A_{low} . This can be formulated as:

$$
d\rho = \sqrt{\left(\left(\frac{\partial \rho}{\partial A_{\text{high}}}\right) dA_{\text{high}}\right)^2 + \left(\left(\frac{\partial \rho}{\partial A_{\text{low}}}\right) dA_{\text{low}}\right)^2}
$$
(2)

ffi

Now, $\frac{\partial \rho}{\partial A_{\text{high}}} = \frac{1}{A_{\text{low}}}$ and $\frac{\partial \rho}{\partial A_{\text{low}}} = -\frac{A_{\text{high}}}{A_{\text{low}}}$, hence, we get: ffi

$$
d\rho = \sqrt{\left(\left(\frac{1}{A_{\text{low}}}\right) dA_{\text{high}}\right)^2 + \left(\left(-\frac{A_{\text{high}}}{A_{\text{low}}^2}\right) dA_{\text{low}}\right)^2}
$$
(3)

Multiplying and dividing R.H.S. by ρ , we get:

$$
d\rho = \rho^* \sqrt{\left(\frac{dA_{\text{high}}}{A_{\text{high}}}\right)^2 + \left(\frac{dA_{\text{low}}}{A_{\text{low}}}\right)^2}
$$
(4)

Hence,
$$
\rho = A_{\text{high}} / A_{\text{low}} \pm \rho^* \sqrt{\left(\frac{dA_{\text{high}}}{A_{\text{high}}}\right)^2 + \left(\frac{dA_{\text{low}}}{A_{\text{low}}}\right)^2}
$$
 (5)

Now, consider the relative biomass ratios of any n such biological replicates, i.e., some ρ_1 , ρ_2 , ... ρ_3 , from *n* independent experiments. The average of these will be given by:

$$
\rho_{\text{avg}} = (\rho_1 + \rho_2 + ... + \rho_n)/n
$$
 (6)

where, $d\rho_{avg}$ should represent the propagated error from all the individual relative biomass ratio values. This can be represented as:

$$
d\rho_{\text{avg}} = \sqrt{\left(\left(\frac{\partial \rho_{\text{avg}}}{\partial \rho_1}\right) d\rho_1\right)^2 + \left(\left(\frac{\partial \rho_{\text{avg}}}{\partial \rho_2}\right) d\rho_2\right)^2 + ... + \left(\left(\frac{\partial \rho_{\text{avg}}}{\partial \rho_n}\right) d\rho_n\right)^2}
$$
(7)

Herein, we know that $\frac{\partial \rho_{avg}}{\partial \rho_1} = \frac{\partial \rho_{avg}}{\partial \rho_2} = \frac{\partial \rho_{avg}}{\partial \rho_n} = \frac{1}{n}$, which reduces the above to:

$$
d\rho_{\text{avg}} = \sqrt{(d\rho_1)^2 + (d\rho_2)^2 + ... + (d\rho_n)^2}/n
$$
 (8)

Hence, the net relative biomass ratio for n biological replicates, each with its own technical replicates, will be given by:

$$
\rho_{\text{avg}} = (\rho_1 + \rho_2 + \rho_3 + ... + \rho_n) / n \pm \sqrt{(d\rho_1)^2 + (d\rho_2)^2 + (d\rho_3)^2 + ... + (d\rho_n)^2} / n
$$
\n(9)

Reporting summary

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

Data availability

The experimental data and the simulation results that support the findings of this study are available in Figshare [https://doi.org/10.6084/](https://doi.org/10.6084/m9.figshare.25757580) m9.fi[gshare.25757580.](https://doi.org/10.6084/m9.figshare.25757580) The 16S rRNA sequencing data of all beetlederived primary isolates have been deposited in GenBank, accession codes for each of which are listed in Supplementary Table 1. The coculture data used in this study are available in NCBI SRA Bioproject accession no. PRJNA1169755. Source data are provided with this paper.

Code availability

The Cellmodeller codes that were used in this study are already publishe[d35](#page-11-0). All modifications are available in Figshare [https://doi.org/](https://doi.org/10.6084/m9.figshare.25757580) 10.6084/m9.fi[gshare.25757580.](https://doi.org/10.6084/m9.figshare.25757580)

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Author contributions

T.B. conceptualized and supervised the study. T.B., M.S., D.A., and P.S. contributed towards the study design. M.S. performed the experiments, simulations, and data analysis, with help from M.G. P.S. characterized the gut-derived microbial strains and performed the genetic analysis. C.K. helped with the computational simulation setup. M.S. and T.B. wrote the manuscript with help from D.A. and C.K. T.B. acquired primary funding support. All authors provided critical feedback on the manuscript.

Competing interests

The authors declare no competing interests.

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