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Microbial single-cell RNA sequencing by split-pool barcoding

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Single-cell RNA-sequencing (scRNA-seq) has become an essential tool for characterizing gene expression in eukaryotes but current methods are incompatible with bacteria. Here, we introduce microSPLiT, a high-throughput scRNA-seq method for gram-negative and gram-positive bacteria that can resolve heterogeneous transcriptional states. We applied microSPLiT to >25,000 *Bacillus subtilis* cells sampled at different growth stages, creating an atlas of changes in metabolism and lifestyle. We retrieved detailed gene expression profiles associated with known, but rare, states such as competence and prophage induction, and also identified novel and unexpected gene expression states including the heterogeneous activation of a niche metabolic pathway in a subpopulation of cells. MicroSPLiT paves the way to high-throughput analysis of gene expression in bacterial communities otherwise not amenable to single-cell analysis such as natural microbiota.

Gene expression in bacteria is highly heterogeneous even in isogenic populations grown under the same lab conditions. Bacteria can randomly differentiate into subpopulations that assume different roles for the survival of the community; a strategy known as bet hedging (1, 2). For example, gene expression programs governing developmental and stress-response states such as competence or antibiotic resistance may switch on stochastically in a small number of single cells (3–5). Population level gene expression measurements are insufficient to resolve such rare states which, to date, have been characterized only in tractable model systems and through methods such as fluorescence microscopy that can only measure a limited set of reporter genes at a time (6).

Single-cell RNA-seq (scRNA-seq) methods developed for eukaryotic cells can provide comprehensive gene expression profiles for tens of thousands of cells (7–11). Although the need for microbial scRNA-seq has been recognized (12), technical challenges have long prevented adapting scRNA-seq technology to microbes. Specifically, bacteria have low mRNA content, about two orders of magnitude less than human cells (13) and bacterial mRNA is not polyadenylated which makes separation from rRNA challenging. Bacteria have diverse cell walls and membranes which can interfere with the lysis or permeabilization steps required for scRNA-seq. Finally, their small size can hinder microfluidic single-cell isolation. Recent work has begun to address these issues and demonstrated that scRNA-seq methods can be adapted to bacteria.

However, in spite of rapid progress from sequencing just a few cells (13, 14) to performing experiments in a 96-well format (15), these prior approaches remain relatively low-throughput compared to the state-of-the-art in eukaryotic scRNA-seq.

We have managed to overcome the difficulties of performing high-throughput scRNA-seq with bacterial cells with a technique we have named microSPLiT (Microbial Split-Pool Ligation Transcriptomics). We applied microSPLiT to profile gene expression states in >25,000 single *B. subtilis* cells, uncovering both rare and unexpected states present in as little as 0.1% of the population. A technically similar and concurrently developed approach termed PETRI-seq also supports the use of single-cell transcriptomics for gene expression analysis in prokaryotes (16).

Developing microSPLiT

MicroSPLiT builds on SPLiT-seq, a eukaryotic scRNA-seq approach, which labels the cellular origin of RNA through combinatorial barcoding (7). In SPLiT-seq, cells are fixed, permeabilized and mRNA is converted to cDNA through in-cell reverse transcription (RT) with barcoded poly-T and random hexamer primers in a multi-well format. Cells are then pooled, randomly split into a new 96-well plate, and a well-specific barcode is appended to the cDNA through ligation. This split-ligation-pool cycle is repeated and a fourth, optional barcode is added during sequencing library

preparation to ensure that each cell acquires a unique barcode combination with high likelihood (Fig. 1A and fig. S1, A and B).

Because SPLiT-seq does not require cell isolation, it is compatible with a wide range of cell shapes and sizes. Moreover, because SPLiT-seq already uses random hexamer primers, in addition to poly-T primers for RT, we reasoned that it might be suitable for detecting bacterial mRNA. However, a direct application of the mammalian SPLiT-seq protocol to bacteria, not surprisingly, resulted in low total UMI (unique molecular identifier) counts (<100 max UMIs/cell, median 0 mRNA reads/cell) and a bias toward gram-negative over gram-positive bacteria (fig. S1C).

We thus set out to optimize sample processing steps for bacteria. With SPLiT-seq's multiplexing capabilities we tested a range of approaches to cell wall removal and membrane permeabilization. A protocol utilizing a combination of a mild detergent, Tween-20, and lysozyme resulted in the best capture efficiency for both gram-positive *Bacillus subtilis* and gram-negative *Escherichia coli* bacteria. (inset of Fig. 1A and table S1). For the mRNA enrichment, we then tested polyadenylation with *E. coli* Poly(A) Polymerase I (PAP) which preferentially polyadenylates mRNA (17, 18), 5'-phosphate-dependent exonuclease ("Terminator", Epicentre) (19) and reverse transcription with ribosomal RNA-specific probes followed by RNaseH-mediated degradation (fig. S2 and table S1). We found that the treatment of fixed and permeabilized cells with PAP resulted in the highest (about 2.5-fold, or approximately 7% of total RNA) enrichment of mRNA reads (inset of Fig. 1A and fig. S2). We also optimized the fixation protocol as well as the downstream enzymatic reaction conditions [table S1 and (20)]. Notably, we found that RT resulted in cell clumping and that mild sonication after this step was necessary to reliably obtain single cell suspensions (fig. S3).

microSPLiT generates high-quality single-cell RNA-seq data

In order to validate microSPLiT performance on a mixture of gram-positive and gram-negative organisms, we grew *E. coli* MW1255 and *B. subtilis* PY79 cells to OD₆₀₀=0.5 and subjected half of each culture to a brief 47°C heat-shock. We performed a microSPLiT experiment on both samples, using the first barcode as a sample identifier. We prepared and sequenced a cDNA library from 2682 total bacteria from heat-shocked and control treatments and aligned the reads to a combined *B. subtilis*-*E. coli* genome. 99.2% of the putative single cell transcriptomes were unambiguously assigned to a single species (Fig. 1B). The rest is attributed to multi-species cellular aggregates, with true aggregate frequency including same-species aggregates expected to be double the multi-species one (1.6%). We sampled a median of 235 mRNA transcripts per

cell for *E. coli* and 397 for *B. subtilis* (Fig. 1C), or approximately 5-10% of the estimated total mRNA (21), and 3.7% and 9.5% of all detected RNA molecules per cell for each respective species (fig. S4, A and B). We also detected a median count of 3753 and 6033 rRNA per cell for *B. subtilis* and *E. coli* (Fig. 1C) as well as a median number of 18 and 20 tRNA molecules per cell (fig. S4C). We observed 230 median unique genes per cell for *B. subtilis* and 138 for *E. coli* (fig. S4D). The majority of detected genes for both species had on average between 0 and 5 UMIs per gene (fig. S4, E and F). The summed *E. coli* expression data was correlated with independently published bulk transcriptomic data ($r=0.736$, fig. S5A) (16). The most highly expressed genes detected only in the bulk assay but not with microSPLiT encode tRNA species (fig. S5B). When we discarded multiply aligned reads, the proportion of mRNA reads increased to 90.5% for *B. subtilis* and 28.2% for *E. coli* (fig. S6, A and B), while mRNA UMI counts per cell were not strongly affected (fig. S6, C and D).

Even with in-cell polyadenylation, the majority of rRNA and mRNA molecules were captured by random hexamer primers, while tRNA were predominantly retrieved by poly-T primers (fig. S7). The mRNA and rRNA UMI counts in each cell were highly correlated ($r=0.97$ and 0.94), as were UMI counts of each detected mRNA captured by poly-T and random hexamer primers ($r=0.87$ and 0.94) (fig. S8). tRNA UMI counts captured by poly-T and random hexamer primers displayed lower correlation (fig. S9) which could be due to the transient native polyadenylation of some tRNA species (22). We found that the 23S and 16S rRNA abundances were highly correlated with each other, while correlation with 5S rRNA was lower (fig. S10). We also quantified the effect of sequencing depth on gene and UMI detection by subsampling analysis, suggesting that additional sequencing would only modestly increase UMI counts (fig. S11).

Next, we tested whether microSPLiT could detect transcriptional responses to heat shock (fig. S12A). Unsupervised clustering identified five distinct clusters which were visualized by t-distributed stochastic neighbor embedding (t-SNE) (Fig. 1D). The first barcode identified two pairs of clusters corresponding to the heat treated and control cultures, and gene expression analysis within each pair further labeled them as corresponding to *B. subtilis* and *E. coli* cells (fig. S12B). Enriched within each heat shock cluster were classical heat shock genes differentially expressed in each of the *E. coli* and *B. subtilis* heat treated clusters as compared to the control clusters (figs. S12, C and D, and S13, A and B) (23, 24).

Unexpectedly, we found an additional small cluster, representing *E. coli* cells from both control and heat treated samples that expressed a different signature of DEAD-box helicase *deaD* induction as well as cold shock genes *cspA-G*, consistent with a transcriptional response to cold (figs. S12C and S13C) (25). This subpopulation of *E. coli* might be

displaying a very rapid response to cold from a brief cold centrifugation step performed as the first step in sample preparation before formaldehyde fixation and is thus likely an artifact of the workflow.

Transcriptional patterns during *B. subtilis* growth in rich medium

Next, we applied microSPLiT to capture transcriptional states across the *B. subtilis* growth curve in a rich medium (LB). In total, we sampled ten optical density (OD) points along the growth curve of the laboratory strain PY79 ranging from OD0.5 (early exponential phase) to OD6.0 (early stationary phase), with one replicate of 4 OD points [(20) and Fig. 2A]. The first barcode was used to record sample identity (i.e., OD) and the replicates are consistent and produced a combined dataset of 25,214 cells (Fig. 2, B and C, and fig. S14). Unsupervised clustering of the combined datasets revealed 14 clusters (Fig. 2B), most of which overlapped with a single OD (Fig. 2C). The most notable exceptions are two smaller clusters that contain cells from multiple ODs: cluster 9 with cells from OD2-3.2 that differentially express myo-inositol metabolism pathway genes, and a very small cluster 13 containing only 36 cells from 5 different OD points uniquely expressing genes associated with the defective PBSX prophage (Fig. 2B and figs. S15 to S18).

We then turned to an analysis of alternative sigma factors which are the primary regulators of prokaryotic RNA polymerase specificity and thus directly shape transcriptional changes in response to environmental conditions (26, 27). To understand whether microSPLiT could capture variation in sigma factor utilization across different growth stages, we averaged expression of genes regulated by each sigma factor, recording for each cluster, both the percentage of cells expressing at least one gene regulated by a given sigma factor and the average intensity of gene expression [Fig. 2D and (20)]. The patterns of sigma factor utilization are largely consistent with the literature, with housekeeping σ^A activity highest at the early growth stage relative to other time points, and the activity of general stress response sigma factor σ^B rising as cells begin to exit from exponential phase (clusters 3-4, OD1.3-1.7) and then declining as cells approach stationary phase (Fig. 2D). Sporulation sigma factors were more active at later ODs, but in a small fraction of cells (clusters 10-12, Fig. 2D) and the extracellular function (ECF) sigma factors were divided into two groups with different patterns of activities [Fig. 2D and (20)]. Additionally, correlations between the sigma factor regulons largely agreed with the concept of molecular time sharing, i.e., the idea that sigma factors compete for RNA polymerases (fig. S19) (28).

To obtain an even finer-grained picture, we inferred the activity profiles of select transcriptional regulators from expression of the genes in their respective regulons (Fig. 2E and

fig. S20) (28). This revealed changes in regulation of carbon utilization, stress responses, metal uptake, developmental decisions and more (Fig. 2E). For example, we observe cellular response to a variety of intrinsic and cell-envelope stresses, as well as temporal activation patterns of a battery of developmental regulators (Fig. 2E). These data indicate that microSPLiT captures known regulatory programs and reveals heterogeneity in a wide range of cellular pathways.

Central carbon metabolism changes and transient activation of alternative carbon utilization pathways

Given the changes in regulation of carbon utilization observed in our analysis of transcriptional regulators, we turned to a more comprehensive examination of carbon metabolism genes enriched in each cluster (Fig. 3A and figs. S15 to S18). When glucose and other preferred sugars are present, they are converted to pyruvate during glycolysis; the primary metabolic route when these sugars are abundant. In these conditions promoting rapid growth, pyruvate is then converted to lactate, acetate, acetoin, and other by-products through fermentation. Upon depletion of preferred sugars, cells redirect the fermentation by-products to be metabolized in the TCA cycle generating additional adenosine triphosphate (ATP) and carbon dioxide (29).

As expected, in the clusters corresponding to early ODs (clusters 0-4) we observe peak expression of genes involved in glycolysis such as glucose permease (*ptsG*), and genes involved in rapid growth and fermentation such as lactate dehydrogenase (*ldh*), pyruvate dehydrogenase (*pdhA*) and acetate kinase (*ackA*) [Fig. 3, A and B, and (20)]. At OD1.7 cells appear to undergo a dramatic transition from glycolysis to gluconeogenesis with multiple genes from the gluconeogenic pathway activated in clusters 7 and 8 [(20) and Fig. 3, A to C, and figs. S16 and S17]. We also find a different pattern of pyruvate production and utilization, together with catabolism of acetoin, another fermentation product, and additional nutrient fluxes into the TCA cycle [(20) and Fig. 3, A to C, and figs. S16 and S17].

Additionally, we observe expression of pathways responsible for uptake and utilization of different carbon sources. As the preferred sources of carbon are depleted, the major repressor of alternative carbon utilization pathways CcpA becomes inactive, permitting the cells to catabolize a variety of carbohydrates (30) (Fig. 3, A to C). We find that the activation and suppression of these pathways happen in varying proportions of the cells in each OD sample and appear to follow a temporal order [Fig. 3, A to C; figs. S15 to S18 and S21; and (20)].

Heterogeneous activation of myo-inositol catabolism pathway at intermediate growth stages

Inositol is an abundant resource in soil, and *B. subtilis* is able

to subsist on inositol as its sole carbon source (31). While LB medium is not typically expected to contain myo-inositol (further “inositol”), heterogeneous inositol utilization pathway activation is observed in a small (3-15% across OD1.7-3.2) subpopulation in both of our independent LB growth experiments (cluster 9, see Fig. 3, A and D to E, and fig. S22). The inositol catabolism intermediate, 2-deoxy-5-keto-D-gluconic acid 6-phosphate (DKGP), is responsible for the pathway induction (31). We hypothesize that these results arise from trace amounts of inositol present in the LB medium, potentially from the yeast extract since yeast is capable of inositol production as a precursor to the essential membrane component, phosphatidylinositol.

There are three operons involved in inositol utilization, *iolT* (main transporter), *iolRS* (the first gene is a repressor and the second is a likely dehydrogenase), and *iolA* through *J* (metabolic enzymes, further “*iolAJ*”), with *IolC* producing and *IolJ* cleaving the pathway-activating DKGP intermediate. *iolRS* and *iolAJ* are normally transcribed by σ^A through divergent transcription (32). In the absence of the inducer, *IolR* suppresses transcription of all three operons (33). In addition, *CcpA* represses the *iolAJ* operon in the presence of glucose (34) (fig. S23A). Interestingly, we observe that the pathway suppressor *iolR* gene is more broadly expressed both outside and inside of cluster 9 (Fig. 3, D and E, and fig. S22A).

To validate our findings, we constructed fluorescent reporters of all three operons in the inositol metabolism pathway [transcriptional reporters P_{iolA} -YFP and P_{iolR} -CFP and protein fusion *IolT*-mScarlet-I, (20)]. As expected, we observed widespread expression of all three operons in the presence of inositol as a sole carbon source (fig. S23, B and D). In agreement with our clustering analysis (Fig. 3E), *B. subtilis* cells grown in LB show heterogeneous expression of P_{iolA} -YFP in 22.7% of cells at OD2 and in 44% of cells at OD4, as opposed to cells grown in LB to OD0.7 (threshold set to 1% positive cells in the “off” state OD0.7, Fig. 3, F and G, and fig. S23, C and E). While microSPLiT data shows that the proportion of cells expressing inositol metabolism genes (belonging to cluster 9) drops from 5% at OD3.2 to 0.3% at OD5.3, the accumulation of YFP expressing cells at OD4 is consistent with the delay in fluorescent protein maturation and with the high stability of fluorescent proteins which mainly get cleared from cells by dilution during cell division.

Altogether, our transcriptomics and fluorescent reporter data indicate that the transcription of genes in the inositol metabolism pathway is activated in a heterogeneous fashion in a subpopulation of cells grown in LB between logarithmic and early stationary phases. The source of the activating molecule, as well as the underlying gene regulatory network architecture behind this behavior remain to be determined.

Motility, antimicrobials production, stress response, and metal ion import

Next, we turned to examine a variety of *B. subtilis* behaviors thought to enhance survival in adverse conditions (35). Bacteria universally produce peptide and small molecule antimicrobials that are meant to target both closely and distantly related organisms (Fig. 4A) (36). We observe the expression of at least three broad spectrum antimicrobials – subtilosin, bacillaene, and plipastatin (37–39) – in various fractions of cells across ODs (Fig. 4A and fig. S24). We also see a rise in spore killing factor (SKF) and spore delay protein (SDP) in the last three ODs (fig. S25).

During active growth, *B. subtilis* can morphologically present as filamentous sessile cells or smaller motile cells (40, 41). Similarly, *B. subtilis* populations are expected to be differentiated into surfactin-producing and extracellular matrix-producing bacteria as cell density increases (42). We profiled the fraction of cells expressing motility genes (*fla-che* operon and flagellin encoded by *hag*), which noticeably declines at OD6.0. Meanwhile, surfactin (*urfA-D*) reaches almost 100% detection at OD6.0, consistent with the PY79 strain having defective matrix production genes that cannot negatively regulate *urfA-D* expression [(43), Fig. 4B].

We also found that genes involved in the unfolded protein response such as ClpP associated proteases (*clpP,C,X,E*), McsA and McsB kinases (*mcsA,B*), and chaperonins (*groEL,ES*) peak at OD1.7, the same time as the cells switch from glycolysis to gluconeogenesis (Fig. 4C). Expression of these genes is induced by the transient increase of regulatory sigma factor σ^B activity during normal exponential growth (see Fig. 2D) which is attributed to intrinsic cellular stresses (44). Additionally, we profiled expression of genes associated with metal uptake such as siderophore bacillibactin (*dhbA*) with associated transporter (*feuA*) and manganese transporter (*mntA-D*) [Fig. 4D and (20)].

microSPLiT quantifies a rare stress response

Cluster 13 (36 cells, or 0.142% of total cells, representing ODs between 0.5 and 2.8) contains a rare subpopulation of cells expressing PBSX prophage genes (Fig. 5A). The PBSX element is a defective prophage that is non-infectious but upon induction causes the release of phage-like particles (45) containing 13 kb of random fragmented chromosomal DNA (46, 47). Prophage gene expression is induced by DNA damage (45, 48) and is known to activate in a small fraction of cells during exponential growth (49, 50) (Fig. 5B). The majority of genes (25) differentially expressed in cluster 13 represent known PBSX prophage genes with functions in PBSX prophage-mediated lysis (*xlyA,B*, *xhlA,B*), phage release (*xepA*), and phage replication (*xtmA,B*), and many PBSX-associated genes of unknown function (Fig. 5C). Thus, we not only identify a rare subpopulation of cells in the state of prophage induction, but

also capture the expression of major phage-associated operons. We also identified eleven host genes with known or putative functions expressed in the PBSX prophage cluster (Fig. 5C). Five of these genes have previously been shown to be induced only in PBSX-harboring strains of *B. subtilis* after DNA damage (46). The rest, including a chemoreceptor (*mcpC*), an ATP-binding cassette transporter (*liaL*), a cell wall binding protein (*ykuG*), an ammonium transporter (*amtB*), a sucrose-6-phosphate hydrolase (*sacA*), and a regulatory protein of homologous recombination (*recX*), have not previously been linked to prophage induction (20).

microSPLiT captures a rare stochastically induced developmental state

Under stress or nutrient limitation, a small fraction (2-5%) of *B. subtilis* cells undergoes stochastic transient differentiation into a state of natural competence, characterized by the ability to uptake extracellular DNA and integrate it into the chromosome (Fig. 5D) (49). The master transcriptional regulator of competence ComK is activated via a positive feedback loop, inducing expression of a suite of >165 genes involved in a variety of cellular processes in addition to DNA uptake (51). Competence is expected to naturally occur under nutrient limitation. We thus separately subclustered the last two OD points (OD5.3 and 6.0). UMAP embedding revealed a small cluster (62 cells, or 4.6% of cells at OD 5.3 and 6) expressing a distinct transcriptional signature of the competent state, or K-state (Fig. 5, E and F). The observed frequency of competence is comparable to previous reports [3-10%, (51, 52)]. The most enriched gene was *comGA*, as expected from prior transcriptomic data (53, 54), followed by the succinyl-CoA synthetase (*sucCD*) operon which is induced in competent cells (51, 53, 55). We also see enrichment of genes encoding the DNA uptake machinery: *comF* and *comE* operons, the response regulator (*rapH*) which represses sporulation development in competent cells (54), genes necessary for processing of internalized ssDNA such as *recA* along with genes for single-strand DNA binding proteins SsbA and SsbB (51), and other genes related to DNA processing [Fig. 5F and (20)]. Overall, we capture the majority of genes associated with the state of competence as defined in two previous microarray studies (53, 54) as well as other approaches (56). In addition, we found four genes not previously linked to the competent state [Fig. 5F and (20)].

Discussion

We applied microSPLiT to *B. subtilis* cells growing in liquid rich medium, which is not associated with cellular heterogeneity. Nevertheless, we found a variety of subpopulations displaying differential gene expression of select metabolic, stress response or developmental pathways. In particular, we identified a myo-inositol catabolism pathway, which was

activated only in a fraction of cells at later OD points in a distinct temporal fashion. We anticipate microSPLiT to be broadly useful in identifying heterogeneous cell states in more varied environments, such as in multi-species biofilms and natural microbiota.

We were able to detect subpopulations of cells as rare as 0.142% (Fig. 5A), pointing to microSPLiT's potential to uncover physiologically relevant rare cell states, such as persistence, that are hard to study by bulk or low-throughput methods. The regulators for many such states are not well known and currently reporters or mutants producing the desired state at a higher frequency cannot be engineered. Even for better understood and inducible states, such as prophage induction by UV irradiation, microSPLiT can produce state-specific transcriptional signatures free of artifacts introduced by the perturbation.

In order to use microSPLiT on complex natural communities, the protocol will likely need to be further optimized, particularly the permeabilization and mRNA enrichment steps as cell wall and membrane composition vary among bacteria. However, alternate treatment for different subsamples may still provide optimal results. In addition, we experienced lower mRNA counts from bacteria in stationary phase as opposed to logarithmic growth phase (figs. S26 and S27), consistent with their slower growth rate and smaller cell size at this stage. Although the resulting data still reliably identified rare cell states, such as the K-state, further improvement of the protocol should increase sensitivity for applications to slower dividing bacteria or challenging environmental conditions. Finally, it may become desirable to increase cell retention, currently about 25% (20) between the RT step and sub-library preparation, as the method is applied to sparse natural communities rather than lab-grown cultures. Still, we expect microSPLiT to provide an exciting new dimension to studies of bacterial gene expression heterogeneity and community behavior facilitated by the potential scalability to millions of bacterial cells and single-cell resolution without the need for constructing reporters.

Materials and methods summary

Experimental methods

Bacterial culture

Escherichia coli MW1255, a derivative of MG1655, and *Bacillus subtilis* PY79 overnight cultures were inoculated into fresh 1.25x LB Lennox medium (Sigma-Aldrich) at 1:1000 or 1:250, respectively, and grown at 37°C with shaking (200rpm). For the heat shock experiment, upon reaching the OD₆₀₀ = 0.5, half of each culture was transferred to the separate 37°C incubator where the temperature was increased to 47°C, and kept there with shaking (200rpm) for 8 min from the time the temperature stabilized. Both control and heat-treated samples then were immediately centrifuged at 4°C,

5000 rcf for 5 min, and resuspended in cold formaldehyde. Since we found a cluster in these data that may represent an artifact of the cold centrifugation conditions, the *B. subtilis* growth curve samples were instead centrifuged at room temperature, 10000 rpm for 1 min before fixation.

Fixation and permeabilization

For the steps below, centrifugation was performed at 4°C, 5000 rcf for 5 min. Following centrifugation, the bacterial pellet was resuspended in 1mL of fresh, cold, 4% formaldehyde solution (in 1x PBS) and incubated at 4°C overnight. The next morning, cells were centrifuged and resuspended in 1ml cold 100mM Tris-HCL+RI ('RI' indicates that SUPERase-In RNase Inhibitor, ThermoFisher, was added to a final concentration of 0.1 U/μL). Cells were centrifuged and resuspended in 250ul of 0.04% Tween-20 in 1x PBS, then permeabilized for 3 min on ice. We then added 1ml of cold PBS+RI, centrifuged the cells and resuspended in 200ul lysozyme mix per sample on ice as follows: 0.1M Tris-HCL pH7, 0.05M EDTA, 2.5mg/ml lysozyme, 0.25U/ml SUPERase-In. We incubated the samples at 37°C in the thermocycler for 15 min as we found that precise timing of lysozyme incubation is critical to maintaining cell integrity at later stages of the protocol. Following the cell wall digestion step, we immediately added 1ml of cold PBS+RI, centrifuged the cells and counted the cells stained with SYTO9 using Accuri C6 flow cytometer. For the species-mixing experiment, we mixed the *E. coli* and *B. subtilis* cells at equal proportions and took 0.6M cells for each of heat shocked and control samples. For the *B. subtilis* growth curve experiments, we took 0.25M cells for each OD sample.

In-cell polyadenylation

In order to enrich for mRNA capture, we performed in situ polyadenylation with *E. coli* Poly(A) Polymerase I (PAP) from NEB. For 0.25M cell samples, the reaction proceeded in 50μl volume, and in 100μl volume for the 0.6M cell samples. For each 0.6M cells bacterial pellet, we added 66μl of water, 4μl of SUPERase-In, 10μl of 10x PAP Buffer, 10μl of 10mM ATP, and 10μl of PAP. The reaction mixture was incubated at 37°C for 30min, then centrifuged upon addition of 1ml cold PBS+RI. We also added 1μL of 10% Tween-20 in order to make the cells easier to pellet. The cells then were resuspended in 0.5ml of cold PBS+RI.

Reducing aggregate formation

Two steps were crucial to break down cell aggregates and reduce the doublet rate: first, we vortexed and double-filtered the bacterial cells prior to reverse transcription; and second, we performed sonication with double filtration right after the reverse transcription.

Following the polyadenylation step, cells in 500ul of PBS-RI were vortexed for 1 min on the highest setting, filtered

through 10μm pluriStrainer (pluriSelect) by pipetting through the membrane, then filtered again through 1μm pluriStrainer with gentle suction, and finally, right before adding to the reverse transcription wells, vortexed again on the highest setting for 1 min.

Following the reverse transcription step and after resuspension in 2 mL of cold PBS+RI, cells were vortexed for 1 min on the highest setting, filtered through 10μm and 1μm pluriStrainer as above, and briefly sonicated at 10% power for 5s on ice for 1 pulse (Sharpertek Ultrasonic Cell Crusher) followed by immediate distribution to the ligation plate. We found that the sonication step can be replaced by a second vigorous vortexing step, with the roughly 2-fold increase in resulting detected doublet rate (0.7% to 1.3%).

In-cell reverse transcription

Like in SPLiT-seq, the first round of barcoding occurs through an in situ reverse transcription (RT) reaction. Cells are split into up to 48 wells, each containing barcoded well-specific reverse transcription primers. We used both random hexamer and anchored poly(dT)₁₅ barcoded RT primers in each well at the ratio of 1:2 (2.5μM random hexamer + 5μM poly(dT)₁₅). In addition to primers, each RT well had a mix of 1x RT Buffer, 0.25U/μL RNase Inhibitor (Enzymatics), 0.25U/μL SUPERase-In RNase Inhibitor, 500μM dNTPs each (ThermoFisher), 7.5% of PEG8000, 20U/μL of Maxima H Minus Reverse Transcriptase (ThermoFisher). We pipetted 4 μL of cells at about 1M cells per mL in PBS-RI into every well, in the total resulting RT reaction volume of 20ul. The plate was incubated in a thermocycler for 10 min at 23°C followed by 50°C for 50 min. RT reactions were pooled back together and after adding 9.6 μL of 10% Triton X-100, cells were centrifuged for 5 min at 3000g at 4°C in a swinging bucket rotor centrifuge. The supernatant was removed and cells were resuspended in 2 mL of cold 1X PBS-RI. The cells then underwent two rounds of filtration and sonication as described above.

In-cell ligations

The oligonucleotide plates for the second and third barcoding round were prepared as previously described (7). We prepared a 2.04 mL ligation mix containing 727.5 μL of RNase-free water, 500 μL 10X T4 Ligase buffer (NEB), 20 μL T4 DNA Ligase (2000 U/μL, NEB), 30 μL RNase inhibitor (40 U/μL, Enzymatics), 12.5 μL SuperaseIn RNase Inhibitor (20 U/μL), and 750 μL of 50% PEG8000. This ligation mix and the 2 mL of sonicated and filtered cells in 1X PBS were added to a basin and mixed thoroughly to make a total of 4.04 mL. The ligation steps were performed as in the SPLiT-seq protocol (7), except we did vigorous vortexing combined with the double filtration technique as described above where the protocol called for a filtration step.

Lysis and sublibrary generation

After the third round of barcoding, we performed a final vigorous vortexing and double filtration step as described above. Then 70 μ L of 10% Triton-X100 was added to the cell solution before spinning it down for 5 min at 3000G and 4°C. We carefully aspirated the supernatant, leaving about 30 μ L to avoid removing the pellet. We then resuspended the cells in 4 mL of wash buffer (4 mL of 1X PBS, 40 μ L of 10% Triton X-100 and 10 μ L of SUPERase-In RNase Inhibitor) and spun down for 5 min at 3000G and 4°C. We then aspirated the supernatant and resuspended in 50 μ L of PBS+RI. After counting cells, we aliquoted them into sublibraries (in 1.7 mL tubes). After adding the desired number of cells to each sublibrary, we brought the volume of each to 50 μ L by adding 1x PBS and froze the cells at -80°C overnight. Next morning, we flash-thawed the cells and added 50 μ L of 2X lysis buffer (20 mM Tris (pH 8.0), 400 mM NaCl, 100 mM EDTA (pH 8.0), and 4.4% SDS) and 10 μ L of proteinase K solution (20mg/mL). We incubated cells at 55°C for 2 hours with shaking at 200 rpm to lyse the cells and reverse the formaldehyde crosslinks.

Computational methods

Alignment and generation of cell-gene matrices

The data preprocessing and alignment was performed using a modified SPLiT-seq pipeline (7), where the cDNA reads were mapped to either a combined *B. subtilis* – *E. coli* genome [ASM904v1.45 and ASM80076v1.37 from EnsemblBacteria (57)] or only the *B. subtilis* 168 genome (ASM904v1.45 from EnsemblBacteria) using STAR with the splicing isoform detection switched off (58). In addition, we kept the highest-scored multimapping reads, assigning a fractional count based on the number of equally-good alignments, since bacterial genomes are known to contain overlapping CDSs. We then generated a matrix of gene counts for each cell (N x K matrix, with N cells and K genes).

Processing of data from the heat shock experiment

Clustering and data analysis for the species-mixing experiment with heat shock treatment was performed using Scanpy (59). We only kept transcriptomes with the number of total reads higher than 200. Then, we removed the ribosomal and tRNA reads from the data, retaining only reads representing the mRNA counts for both species. We further filtered cells based on the mRNA counts, retaining cells expressing >100 reads and >100 genes, and additionally filtered the genes retaining the genes expressed in >5 cells. We then applied standard Scanpy normalization and scaling, dimensionality reduction, and clustering as described in the Scanpy tutorial (59, 60). The clusters were produced by Louvain graph-clustering method and manually inspected for the top differentially expressed genes. After inspection, three pairs of transcriptionally similar clusters with fewer differentially

expressed genes were merged, resulting in clusters 1, 2 and 3 in Fig. 1D.

Processing of B. subtilis data from the growth in rich media experiment

Clustering and data analysis for the combined 10 samples of *B.S.* grown in rich medium was performed using Scanpy (59) and verified with Seurat v3 (61) and UNCURL (62). Experiment 1 sampled OD points 0.5, 1.0, 1.7, 2.0, 2.8, and 3.2, while in experiment 2 we collected OD points 0.5, 1.0, 1.3, 1.7, 2.8, 3.5, 5.3, and 6.0. For the data from both experiments separately, we discarded any transcriptomes with the number of total reads fewer than 200. Then, we filtered the data and retained only reads representing the mRNA counts (excluding the ribosomal and tRNA reads). Finally, we combined the data matrices together. Since the read depth decreased for the higher OD samples (fig. S26), for selecting the highest quality data, we implemented differential thresholds for each OD in the combined matrix, retaining top 75% of the cells by read counts for each OD sample. This resulted in retention of 25,214 transcriptomes from both experiments. Finally, we performed batch correction through Scanpy, using a python implementation of ComBat (63, 64). Cells that passed the QC were clustered using a pipeline described in previous studies (7, 60).

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SUPPLEMENTARY MATERIALS

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Materials and Methods

Figs. S1 to S27

Tables S1 to S4

References (65–89)

MDAR Reproducibility Checklist

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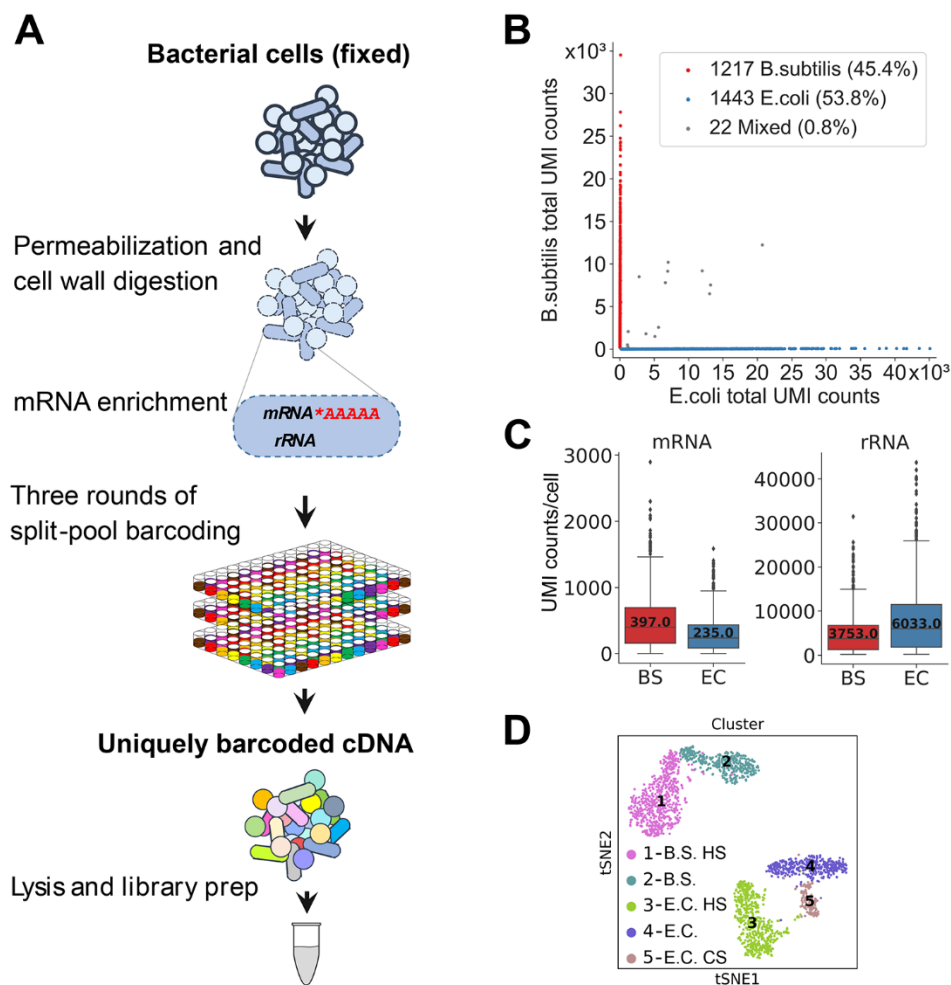


Fig. 1. MicroSPLiT development and validation. (A) MicroSPLiT method summary. Fixed bacterial cells are permeabilized with Tween-20 and lysozyme. The mRNA is then polyadenylated in-cell with *E. coli* Poly(A) Polymerase I (PAP). The cellular RNA then undergoes three rounds of combinatorial barcoding including in-cell reverse transcription (RT) and two in-cell ligation reactions, followed by lysis and library preparation. (B) Barnyard plot for the *E. coli* and *B. subtilis* species-mixing experiment. Each dot corresponds to a putative single-cell transcriptome. Total UMI (unique molecular identifier) counts for all types of RNA are plotted. (C) mRNA and rRNA UMI counts per cell for both species. Error bars represent 95% confidence intervals. (D) t-stochastic neighbor embedding (t-SNE) of the data from heat shock experiment showing distinct clusters. “HS” – heat shock, “CS” – cold shock [see (20)].

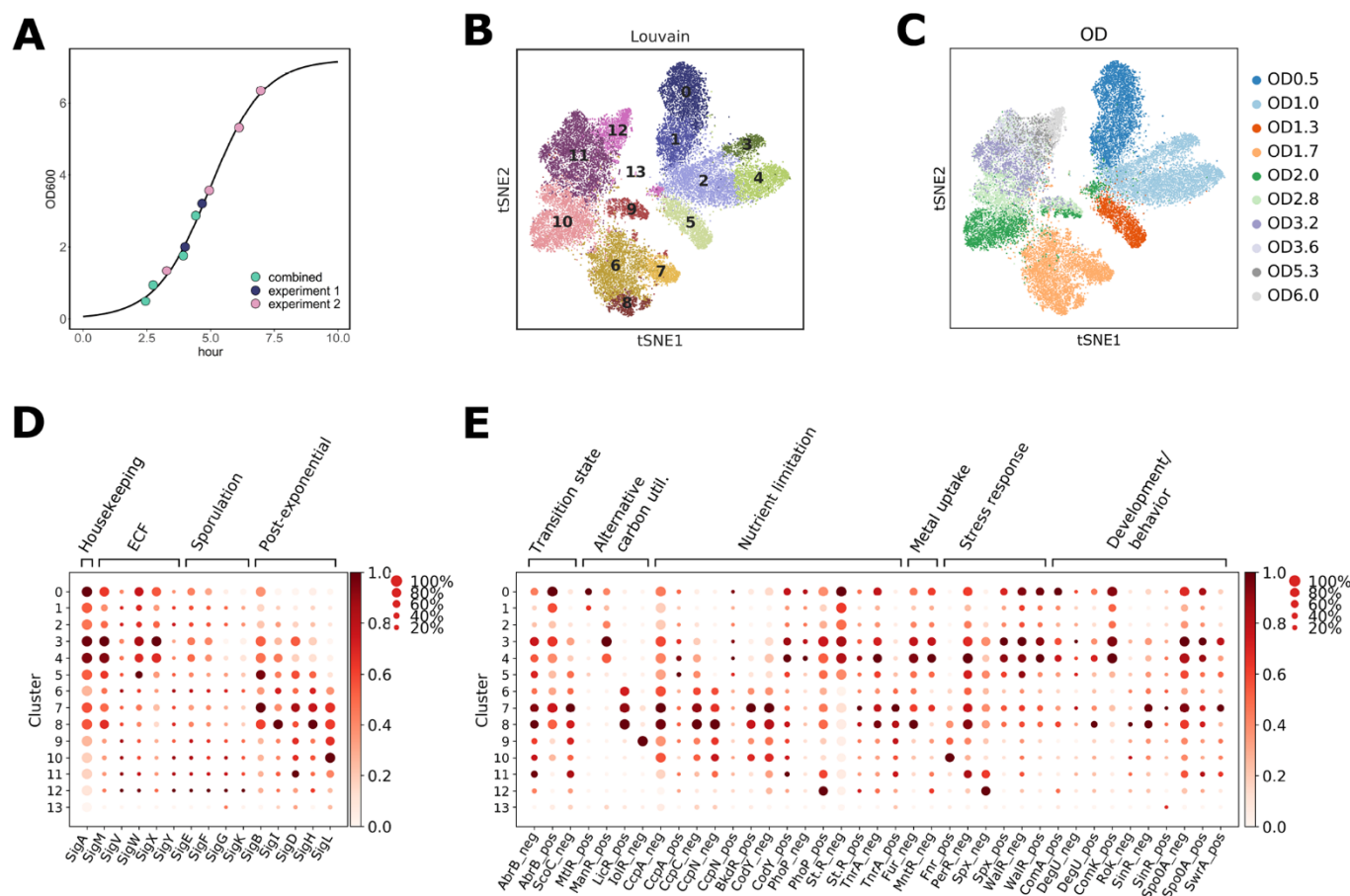


Fig. 2. MicroSPLIT detects global transcriptional states during *B. subtilis* growth. (A) Optical density (OD) points sampled in two experiments overlaid on the growth curve from experiment 2 (20). (B and C) t-SNE embedding of the combined growth curve data colored by (B) cluster or (C) OD. (D) Inferred normalized sigma factor activity for each cluster, calculated from averaged expression of genes regulated by each sigma factor. The size of each dot indicates the proportion of cells in the cluster in which the sigma factor is active, while the color indicates the average activity normalized from 0 to 1 across all clusters for each sigma factor. (E) Inferred activity of select transcriptional regulators per cluster, calculated and normalized for all clusters as above, plotted as in (D). “Neg” indicates that activity was calculated for the genes known to be negatively regulated by this TR, and “pos” indicates the activity was calculated for the genes positively regulated by the given TR.

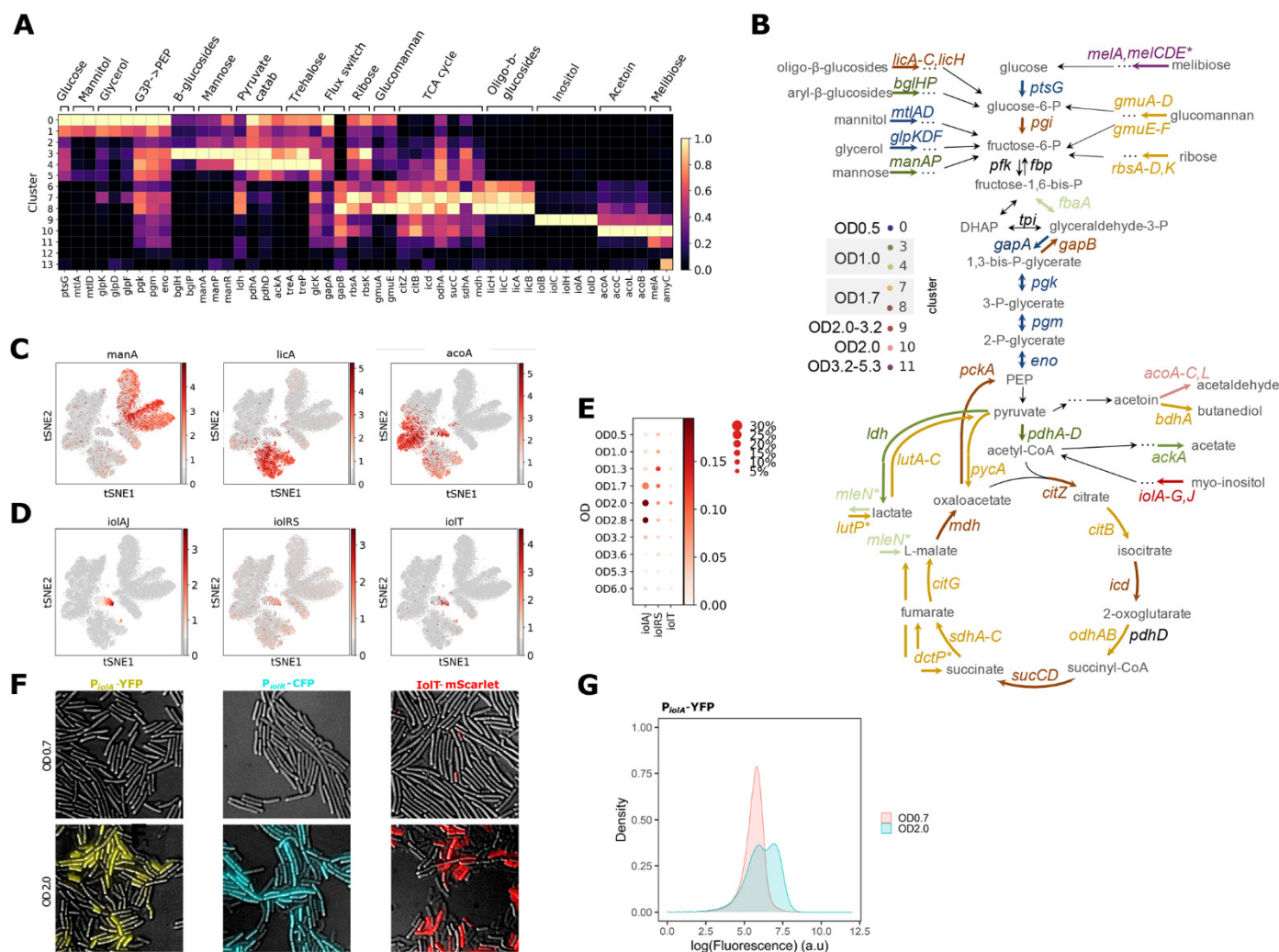


Fig. 3. Central carbon metabolism changes and alternative carbon sources utilization during *B. subtilis* growth. (A) Normalized expression of genes from select metabolic pathways and central carbon metabolism shown per cluster. Gene expression shows distinct carbon utilization programs associated with different clusters and growth states. (B) Schematic of the central carbon metabolism pathway showing alternative carbon sources, metabolic products and genes in the pathway. The genes are color-coded according to the cluster they are highest expressed in. (C) Expression of select genes from (A) overlaid on the t-SNE plot to illustrate the differential patterns of activation. (D) Expression of each of the three inositol utilization operons, averaged across all genes in a given operon, and overlaid on the t-SNE plot. (E) Activities of the three inositol utilization operons across ODs. The size of each dot indicates the proportion of cells in each OD sample expressing any of the genes in the selected operon, while the color shows the average expression of the genes in a given operon. (F) Fluorescence and DIC microscopy overlays of *B. subtilis* expressing P_{iolA} -YFP (left), P_{iolR} -CFP (middle) or $IolT$ -mScarlet-I (right) grown in LB to OD0.7 (top row) or OD2.0 (bottom row). (G) Flow cytometry of P_{iolA} -YFP strain grown to OD0.7 or 2.0.

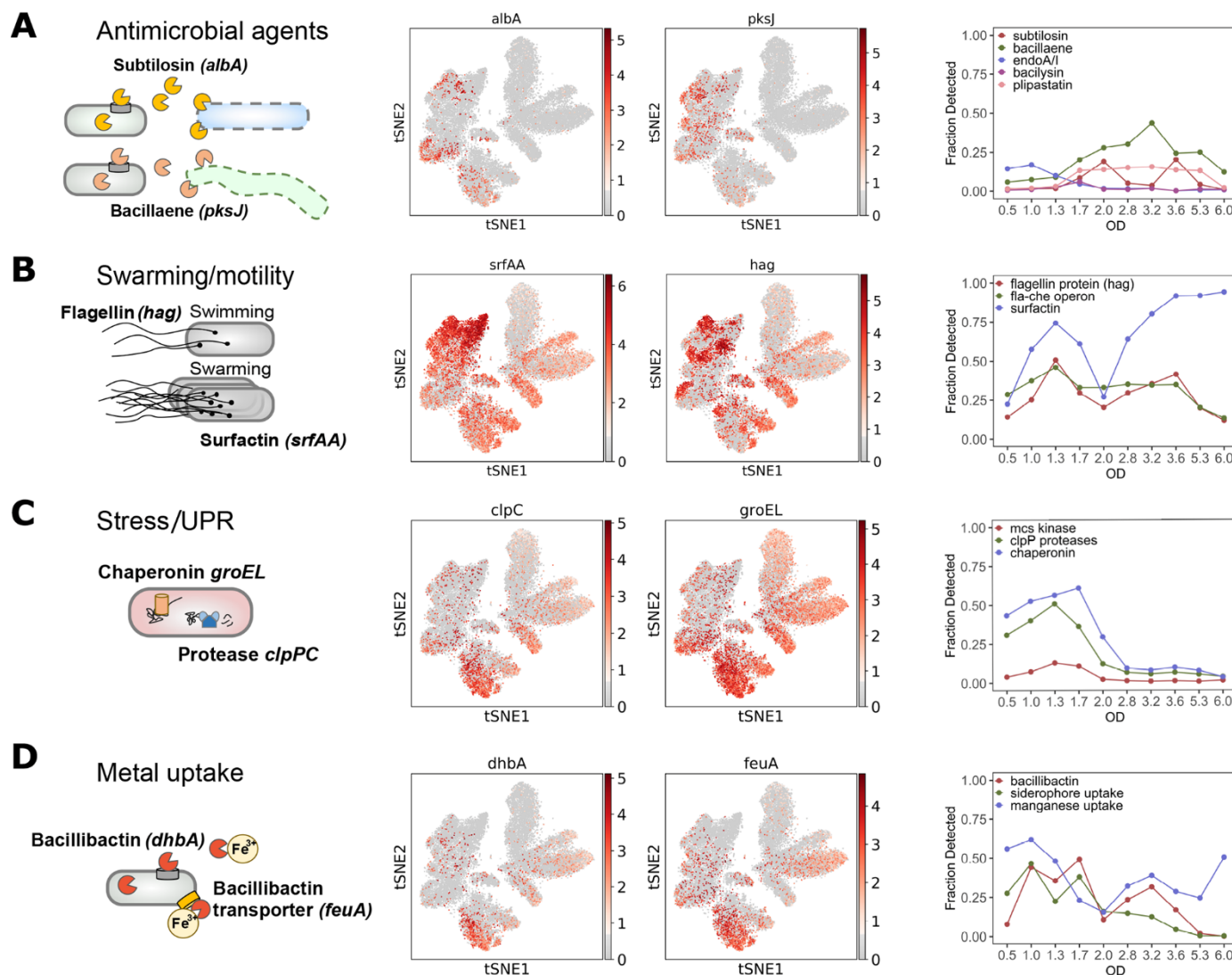


Fig. 4. Intrinsic stress responses and developmental gene expression. Left to right: Pathway diagram. Overlays of expression of genes representative of each pathway on the t-SNE. Fraction of cells expressing at least one of the genes in the indicated operon as a function of OD. (A) Antimicrobial agents [subtilisin (*albA*) and bacillaene (*pksJ*)] and endoA toxin-antitoxin system. (B) Swarming and motility [surfactin (*srfAA*) and flagellin (*hag*)]. (C) Intrinsic stress and unfolded protein response (UPR) [GroEL chaperonin (*groEL*) and ClpCP protease (*clpC*)]. (D) Iron [bacillibactin (*dhbA*) and siderophore transporter (*feuA*)] and manganese uptake. The genes used for fractional plots are listed in table S2.

Microbial single-cell RNA sequencing by split-pool barcoding

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