- 1 Polysaccharide breakdown products drive degradation-dispersal cycles of
- 2 foraging bacteria through changes in metabolism and motility
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#### 41 Abstract

42 Most of Earth's biomass is composed of polysaccharides. During biomass decomposition, 43 polysaccharides are degraded by heterotrophic bacteria as a nutrient and energy source and are thereby 44 partly remineralized into CO<sub>2</sub>. As polysaccharides are heterogeneously distributed in nature, following 45 the colonization and degradation of a polysaccharide hotspot the cells need to reach new polysaccharide 46 hotspots. Even though these degradation-dispersal cycles are an integral part in the global carbon cycle, 47 we know little about how cells alternate between degradation and motility, and which environmental 48 factors trigger this behavioral switch. Here, we studied the growth of the marine bacterium Vibrio 49 cyclitrophicus ZF270 on the abundant marine polysaccharide alginate. We used microfluidics-coupled 50 time-lapse microscopy to analyze motility and growth of individual cells, and RNA sequencing to study 51 associated changes in gene expression. Single cells grow at reduced rate on alginate until they form 52 large groups that cooperatively break down the polymer. Exposing cell groups to digested alginate 53 accelerates cell growth and changes the expression of genes involved in alginate degradation and 54 catabolism, central metabolism, ribosomal biosynthesis, and transport. However, exposure to digested 55 alginate also triggers cells to become motile and disperse from cell groups, proportionally increasing 56 with the group size before the nutrient switch, accompanied by high expression of genes involved in 57 flagellar assembly, chemotaxis, and quorum sensing. The motile cells chemotax toward alginate 58 hotspots, likely enabling cells to find new polysaccharide hotspots. Overall, our findings reveal the 59 cellular mechanisms underlying bacterial degradation-dispersal cycles that drive remineralization in 60 natural environments.

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#### 62 Importance

Polysaccharides, also known as glycans, are the most abundant form of biomass on Earth and understanding how they are degraded by microorganisms is essential for our understanding of the global carbon cycle and the storage and release of CO<sub>2</sub> by natural systems. Although group formation is a common strategy used by bacterial cells to degrade ubiquitous polymeric growth substrates in nature, where nutrient hotspots are heterogeneously distributed, little is known about how cells prepare for dispersal from an exhausted nutrient source and re-initiate degradation of new nutrient patches. By 69 quantifying growth, motility and chemotaxis of individual cells and comparing gene expression changes 70 when populations were exposed to either polysaccharides or their degradation products in the form of 71 digested polysaccharides, we show that bacterial cells alter their behavior when they experience a shift 72 from polymeric to digested polysaccharides: After cells form groups during growth on polymers, the 73 exposure to degradation products made cells motile, enabling dispersal from sessile cell groups and -74 guided by chemotaxis - movement towards new polysaccharide hotspots. Our study sheds light on the 75 cellular processes that drive bacterial growth and behavior during carbon remineralization, an important 76 driver of CO<sub>2</sub> release from biomass in natural systems.

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#### 78 Introduction

Polysaccharides represent the largest fraction of biomass on Earth<sup>1,2</sup> and are constantly degraded and 79 80 remineralized by microorganisms. Polysaccharides, also known as glycans, are long chains of 81 monosaccharide units produced by cells for structural support (e.g., cellulose, chitin, and alginate) or 82 energy storage (e.g., starch or glycogen)<sup>1</sup>. Heterotrophic microbes obtain nutrients and energy from the 83 polysaccharide breakdown products. They often use exoenzymes, either secreted or anchored in the cell membrane<sup>2,3</sup>, to cleave these large polymers into smaller units that can be taken up by cells. The 84 85 formation of dense cell groups is observed during growth on polysaccharides by diverse bacteria, 86 including the soil- and gut-dwelling Bacillus subtilis<sup>3</sup>, the oligotrophic fresh-water Caulobacter crescentus<sup>4</sup>, and several representatives of the copiotrophic<sup>5,6</sup> marine Vibrio spp.<sup>7,8</sup>. It has been 87 88 suggested that "cooperative" growth<sup>3</sup>, where dense cell groups reduce diffusional loss of valuable 89 degradation products and secreted exoenzymes, represents a general principle in polysaccharide degradation<sup>4,8</sup>. Yet, how bacteria subsequently disperse from exhausted polysaccharide sources and 90 91 navigate towards new polysaccharide hotspots remains poorly understood.

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93 To study the cellular mechanisms that govern the switch between polysaccharide degradation and 94 dispersal toward new polysaccharide hotspots, we worked with the ubiquitous marine 95 Gammaproteobacterium *Vibrio cyclitrophicus* ZF270, a degrader of the prevalent marine 96 polysaccharide alginate<sup>6,9</sup>. Alginate is a linear polysaccharide that is produced by brown algae as a cell

97 wall component, as well as by certain bacteria<sup>10</sup>. It can be cleaved into oligomers by endo-acting alginate 98 lyases or into monomers by exo-acting alginate lyases (alginate lyases reviewed here<sup>11</sup>), namely  $\beta$ -Dmannuronic acid and  $\alpha$ -L-guluronic acid<sup>10</sup>. V. cyclitrophicus ZF270 is found predominantly in the large-99 100 particle fractions of coastal water<sup>12</sup>, can attach and form biofilms on the surfaces of multiple 101 polysaccharides including alginate<sup>13</sup>, and was shown to secrete alginate lyases during alginate 102 degradation<sup>8</sup>. In this study, we used microfluidics coupled to automated time-lapse imaging to quantify 103 the growth dynamics, group formation, and motility of V. cyclitrophicus ZF270 at the single-cell level 104 under constant supply of either polymeric alginate or alginate degradation product in the form of 105 digested alginate, as well as upon a transition from alginate to digested alginate. Furthermore, we used 106 RNA-sequencing to compare the gene expression of V. cvclitrophicus ZF270 grown on alginate and 107 digested alginate. We found striking responses to the form of alginate in growth rate, group formation, 108 motility and chemotaxis, as well as in the expression of corresponding genes. Overall, our work 109 provides insights into the metabolic and cellular regulation that allows cells to forage in heterogeneous 110 nutrient-scapes through degradation-dispersal cycles.

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#### 112 Results

#### 113 Extracellular break down of alginate delays population growth

114 To probe the phenotypic and metabolic regulation of bacterial cells during the progressive degradation 115 of a polysaccharide source, we developed an experimental system consisting of the marine bacterium 116 V. cyclitrophicus ZF270 growing on alginate, a highly abundant polysaccharide in the ocean. To mimic 117 the local nutrient environment during the colonization of a new polysaccharide source, we used 0.1% 118 weight per volume (w/v) algae-derived alginate in its soluble form (in the following also simply referred 119 to as "alginate"). Since commercially available alginate breakdown products of specific sizes are limited 120 and expensive, with only one monomeric component available at a considerable cost, we simulate an 121 advanced stage of polysaccharide degradation by supplying cells with digested alginate (0.1% (w/v)). 122 This digested alginate is prepared by treating alginate with a readily available endo-acting alginate lyase 123 (see Materials and Methods). This mimics an environment where degradation products like monomers 124 and oligomers become abundant through the action of extracellular alginate lyases. The commercially available alginate lyase has been shown to produce alginate oligomers of progressively smaller size over extended digestion periods<sup>14,15</sup>. We used Liquid Chromatography-Mass Spectrometry (LC-MS) to analyze the composition of the digested alginate, which had been subject to 48 hours of digestion. We found that digested alginate contained more monosaccharides than untreated alginate, which contained more alginate molecules of higher molecular weight (Fig. S1).

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131 Using this system, we first set out to investigate the growth dynamics of V. cvclitrophicus ZF270 in 132 well-mixed batch cultures containing either alginate or digested alginate as a sole carbon source (Fig. 133 S2A). We found that similar yields were reached in the alginate and digested alginate cultures as 134 indicated by optical density (Fig. S2B and C) and confirmed by colony counts (Fig. S3), suggesting that 135 the same amount of alginate supports a similar yield irrespective of its degree of depolymerization. 136 However, the onset of growth on alginate was delayed by about 7.5 hours compared to growth on 137 digested alginate (Fig. S2D and E). This is consistent with previous observations that in well-mixed 138 environments the lag time of bacteria growing on alginate can be reduced by the external supplementation of alginate lyases<sup>8,14</sup>. Together, these findings indicate that growth on polysaccharides 139 140 such as alginate in well-mixed cultures is initially limited by the extracellular breakdown of the 141 polysaccharide.

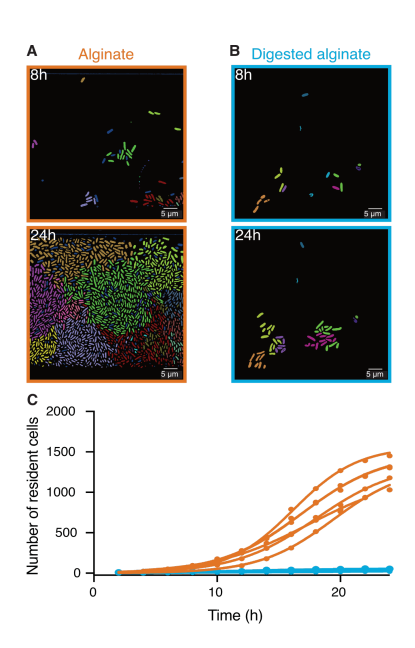
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#### 143 Large cell groups form on alginate but not on digested alginate

144 To better understand how cells react to the changing degree of depolymerization of a polysaccharide 145 source during degradation, we investigated the growth of V. cyclitrophicus ZF270 at the level of single 146 cells on alginate and digested alginate. For this purpose, we grew the cells in microfluidic growth chambers, described in detail by Dal Co et al.<sup>16</sup>, where cells were provided with a continuous flow of 147 148 culture media containing either alginate or digested alginate as carbon source. We found that over 24 149 hours dense groups of more than 1000 cells formed on alginate, filling the entire microfluidic chamber 150 (Fig. 1A). In contrast, cells supplied with digested alginate grew in smaller groups that never exceeded 151 100 cells per chamber (Fig. 1B and C). Reconstructed cell lineages revealed that the large cell groups 152 on alginate formed because cells often did not disperse after division and thereby formed dense cell

153 groups originating from a single cell lineage (Fig. 1A). The lower cell density on digested alginate could 154 be caused by slower growth or by more cells leaving the chambers. As the maximum growth rate both 155 in bulk and on single cell level is similar in alginate and digested alginate (Fig. S2E and F), it is most 156 likely that the lower cell density in digested alginate is caused by cell dispersal. These observations 157 suggest that cells can modulate their propensity to form groups depending on the state of polysaccharide 158 degradation in their local environment. Similar observations were made with a different model system (*Caulobacter crescentus* growing on the polysaccharide xylan)<sup>14</sup>, indicating that group formation on 159 160 polymeric nutrient sources may be a general mechanism of bacteria that degrade polysaccharides 161 extracellularly.





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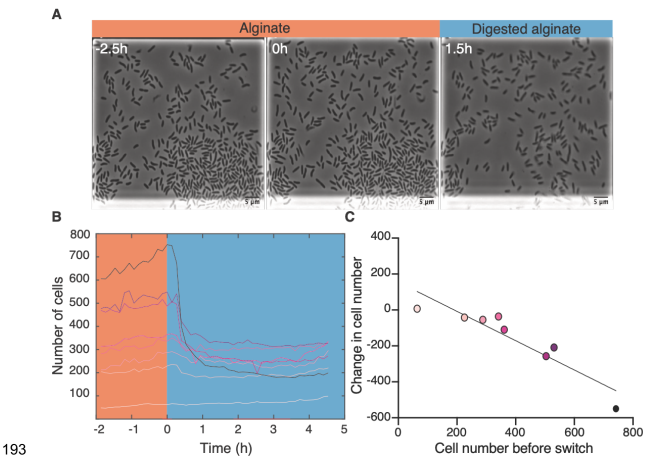
164 Figure 1. Large cell groups form on alginate but not on digested alginate. Representative images at different 165 time points of V. cvclitrophicus ZF270 cells growing in microfluidic chambers, described in detail by Dal Co et 166 al.<sup>16</sup>, with (A) alginate medium or (B) digested alginate medium, both in their soluble form (not visible). Cells are 167 false-colored according to their lineage identities based on cell segmentation and tracking over 24 hours. Cells 168 without identified progenitors are colored in dark blue. See Supplemental Videos S1 (alginate) and S2 (digested 169 alginate) for time-lapse videos. (C) Cell numbers within microfluidic chambers supplied with alginate (orange) 170 are substantially higher than cell numbers within microfluidic chambers supplied with digested alginate (blue) 171 (Logistic growth regression for alginate:  $R^2 = 0.99$ , maximal number of cells = 1217-1564, k = 0.24-0.38 h<sup>-1</sup>; for 172 digested alginate:  $R^2 = 0.86-0.97$ , maximal number of cells = 47-100, k = 0.07-0.4 h<sup>-1</sup>). Circles indicate the number 173 of cells present at a given time point in each chamber ( $n_{chambers} = 7$ ). Data for chambers with alginate originate 174 from D'Souza et al.<sup>8</sup>. Lines are fits of a logistic growth regression line for each condition.

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#### 176 Transition from alginate to digested alginate triggers density-dependent dispersal of cells

177 To investigate the transition from the large cell groups formed on alginate to the small groups formed 178 on digested alginate, we subjected V. cyclitrophicus ZF270 cells grown on alginate to a switch to 179 digested alginate (Fig. 2A). Following the limited cell motility on alginate, this switch led to a rapid 180 decrease in cell density within the growth chambers (Fig. 2B), presumably caused by cell dispersal. As 181 we previously reported for other Vibrionaceae isolates, the growth rate of the cells on alginate was 182 dependent on the local cell density: Initially the growth rate increased with cell density but then 183 decreased at high cell densities, indicating that cell groups can benefit from the sharing of breakdown 184 products generated by each other's exoenzymes, but also increasingly compete for nutrients<sup>8</sup>. This led us to investigate whether cell in larger groups, potentially experiencing stronger nutrient competition, 185 186 might have a higher propensity to disperse after a switch to digested alginate than cells in smaller 187 groups. We indeed found an inverse relationship between the number of cells at the time of the nutrient 188 switch and the following change in cell number, meaning that cells in chambers with many cells were 189 more likely to disperse than cells in chambers with less cells (Fig. 2C). Thus, during the transition from 190 polysaccharides to degradation products, we found that the dispersal rate of cells depends on the size of 191 the cell groups, likely through increased motility of cells in large groups.

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194 Figure 2. Transition from alginate to digested alginate triggers density-dependent dispersal of cells. (A) 195 Representative time-lapse images of V. cyclitrophicus ZF270 cells (phase contrast microscopy) in microfluidic 196 growth chambers that were initially exposed to alginate and then switched to digested alginate. (B) Number of 197 cells in different chambers over time, each chamber indicated by a unique color (n = 8). The carbon source is 198 indicated by the colored background (orange: alginate; blue: digested alginate). See Supplemental Video 3 for a 199 time-lapse video. (C) Negative relationship between the difference in number of cells within the microfluidic 200 growth chambers before and after the nutrient switch. Each circle represents one growth chamber with colors 201 corresponding to (B), and the line depicts a linear regression fit ( $R^2 = 0.85$ , slope = -0.81, p-value < 0.001).

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# 203 Cells growing on digested alginate are more motile and chemotax towards new polysaccharide204 sources

To investigate whether increased cell motility of *V. cyclitrophicus* ZF270 underlies the dispersal of cells observed after a shift to digested alginate, as also previously observed for *C. crescentus* cells on the monosaccharide xylose<sup>4,14</sup>, we quantified the motility of single cells supplied with either alginate or digested alginate. We found that the single-cell swimming speed as well as the swimming distance were

significantly larger for cells supplied with digested alginate compared to cells supplied with alginate,
and that a larger fraction of cells was motile (Fig. 3). This confirmed increased motility as a cellular
response to the exposure to degradation products in the form of digested alginate.

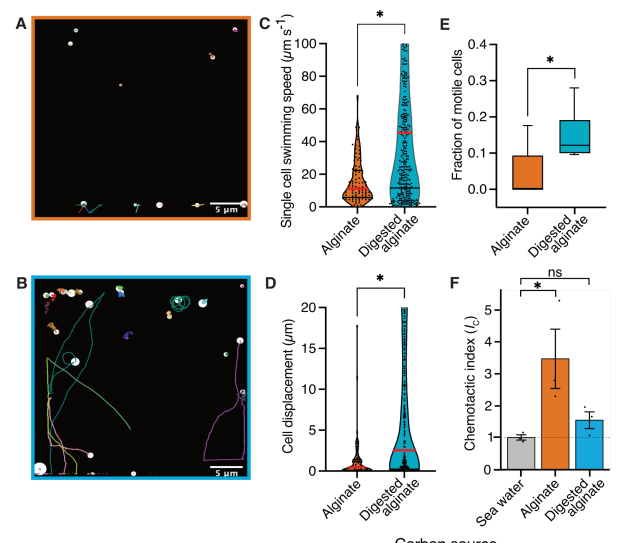
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213 To understand if the motile cells are directed by chemotaxis, we measured the chemotactic strength 214 towards alginate and digested alginate using the In Situ Chemotaxis Assay (ISCA)<sup>17,18</sup>. We found V. *cyclitrophicus* ZF270 to significantly chemotax toward alginate (chemotactic index  $I_c > 1$ ) but not 215 216 toward digested alginate (Fig. 3F). This suggested that the increase in motility is accompanied by 217 chemotaxis toward alginate. Overall, these findings suggest that once the breakdown of a 218 polysaccharide source makes breakdown products available in the local environment, a fraction of cells 219 becomes motile and disperses. These motile cells are guided by chemotaxis towards new polysaccharide 220 sources, allowing them to start a new degradation-dispersal cycle.

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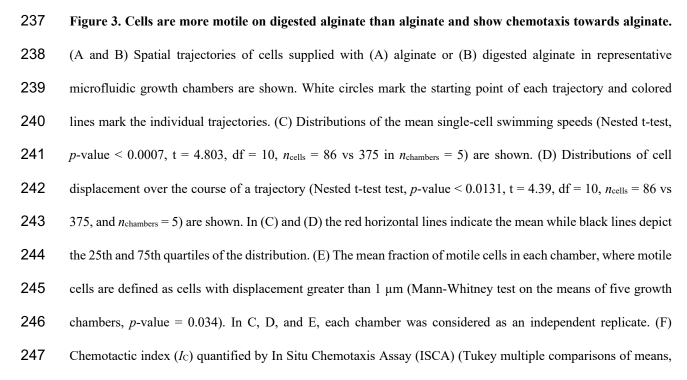
## Altered gene expression in central carbon metabolism, enzyme production, secretion and transporters, motility, and quorum sensing underlies the late-stage alginate degradation and cell dispersal

225 Next, we sought to elucidate the molecular mechanisms underlying the observed phenomenological 226 disparities between cells cultivated on alginate and digested alginate. Due to the challenge of generating 227 knock-out mutants in natural isolates, we used transcriptomics to investigate the differentially expressed 228 genes of V. cyclitrophicus ZF270 under these respective conditions. To obtain a high-quality reference 229 genome of V. cvclitrophicus ZF270, we sequenced and assembled a new reference genome using 230 combined short and long read sequencing (BioProject PRJNA991487). We then grew cultures on either 231 alginate or digested alginate until mid-exponential phase. To understand which cellular functions were 232 affected by the expression changes, we first performed differential gene expression analysis using the 233 software DESeq2. Here, genes exhibiting a log2 fold expression change greater than 0.5 or smaller than 234 -0.5 between the two conditions, with a Benjamini-Hochberg(BH)-adjusted p-value below 0.01, were 235 considered to be differentially expressed (Table S1). Next, we investigated which KEGG categories



236

Carbon source



248 95% family-wise confidence levels as error bars, p-value < 0.05, n = 3). Asterisks indicate statistically significant 249 differences. See Supplemental Videos S4 and S5 for time-lapse videos of swimming cells.

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were enriched in either genes with increased or decreased expression via Gene Set Enrichment Analysis (GSEA)<sup>19</sup> (Table S2) and found nine categories significantly enriched in genes with increased gene expression on digested alginate, and three categories significantly enriched in genes with decreased gene expression (Fig. 4A).

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256 Cells growing on digested alginate showed expression changes in parts of the central metabolism and 257 translation, compared to cells growing on alginate. Specifically, two pathways of the central metabolism were significantly enriched in genes with lower expression on digested alginate ("Valine, leucine and 258 259 isoleucine biosynthesis" and "Propanoate metabolism" with a negative normalized enrichment score 260 (NES), Fig. 4A, S4A and B, Table S3 and S4). We also found the gene set that encodes ribosomal 261 proteins significantly enriched with genes highly expressed on digested alginate (positive NES, Fig. 4A 262 and S4C, Table S5). This implies that cells invest proportionally more of their transcriptome into the production of new proteins, a sign of faster growth<sup>20,21</sup>. Both observations likely relate to the different 263 264 growth dynamics of V. cyclitrophicus ZF270 on digested alginate compared to alginate.

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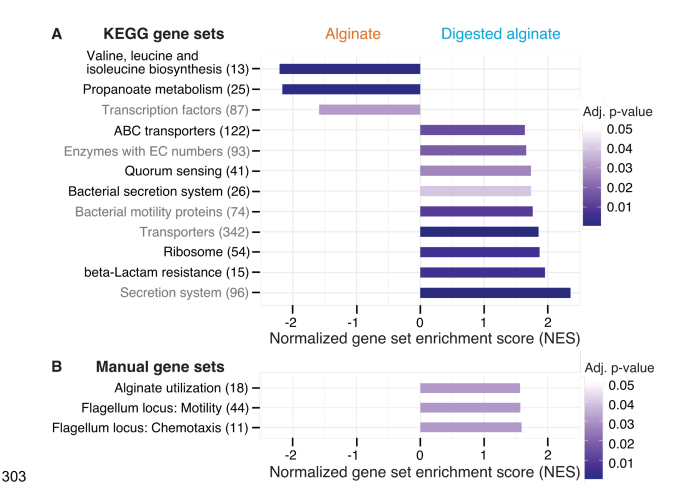
266 The expression of transporters and secretion systems was generally increased in cells growing on 267 digested alginate, compared to cells growing on alginate. This includes genes with a function in 268 "Bacterial secretion systems", namely secretion systems I to VI, which mediate protein export through 269 the inner and outer membranes of Gram-negative bacteria (Fig. 4A and S4D, Table S6). Notably, eleven 270 of the thirteen General Secretion Pathway (GSP) genes (part of the type II secretion system) showed 271 1.1 to 3.4-fold increased expression levels on digested alginate (p-values < 0.01, Fig. 4A, Table S8). 272 The GSP is known to facilitate secretion of various extracellular enzymes like chitinases, proteases, and 273 lipases<sup>22,23</sup>, therefore, the positive enrichment on digested alginate may be linked to the export of 274 extracellular alginate lyases. The KEGG BRITE category "Secretion system" was also enriched on 275 digested alginate (Fig. 4A and S4D, Table S7). It contains additional genes involved in protein export,

276 of which especially genes of type IV pilus assembly, conjugal transfer pilus assembly, and MSHA pilus 277 biogenesis were highly expressed on digested alginate, implicating them as interesting subjects to 278 further research in their role in protein secretion and cell attachment and detachment during 279 degradation-dispersal cycles of polysaccharide-degrading bacteria. Also, the gene set of transporters 280 and in particular ABC transporters were positively enriched (Fig. 4A, Table S10 and S11). The latter 281 showed enrichment especially in genes related to saccharide, iron, zinc, and phosphate transport (Fig. 282 S4E), which suggests that cells growing on digested alginate invest proportionally more of their 283 transcriptome into uptake of not only saccharides but also essential nutrients like iron, zinc, and 284 phosphate, which may become growth-limiting when degradation products are abundantly available.

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286 Both motility and quorum sensing genes increased in expression in cells growing on digested alginate. 287 The positive enrichment of the gene set containing bacterial motility proteins matched the increase in 288 motile cells that we observed in Figure 3E (Fig. 4A, Table S12). The set of quorum sensing genes was 289 also positively enriched in cells growing on digested alginate (Fig. 4A and S4F, Table S13). Quorum sensing is known to control biofilm formation in Vibrio cholerae<sup>24,25</sup> and may orchestrate the density-290 291 dependent dispersal in the presence of degradation products, though the particular signaling cues remain 292 to be uncovered. The strong cellular response to the degree of alginate depolymerization was also 293 emphasized by the finding that transcription factor genes were enriched among genes with decreased 294 expression on digested alginate (Fig.4, Table S14). The set of genes associated to KEGG's beta-lactam 295 resistance category was enriched in genes with increased expression on digested alginate. However, 296 most genes in this gene set are also associated to the KEGG pathways "Transporters", "Quorum 297 sensing", "Chromosome and associated genes", and "Peptidoglycan biosynthesis" and thus the 298 enrichment likely reflects expression changes in these categories and may relate to the difference in 299 growth dynamics (Fig. 4A and S4G, Table S15). Overall, the observed gene expression changes provide 300 insights into the molecular mechanisms underlying the substantial adaptations of V. cyclitrophicus 301 ZF270 cells to the polymeric and digested form of alginate.

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304 Figure 4. Twelve functional gene sets are enriched in genes with increased or decreased expression in cells 305 grown on digested alginate. Gene set enrichment analysis (GSEA) with (A) all KEGG pathways and KEGG 306 BRITE categories as gene sets or with (B) a custom alginate utilization, flagellar assembly, and flagellum-driven 307 chemotaxis gene set was performed comparing the gene counts of the transcriptome of V. cvclitrophicus ZF270 308 cultures grown on digested alginate and alginate (six replicates each). Gene sets with a positive enrichment score 309 were enriched with genes with higher expression in cells grown on digested alginate relative to cells grown on 310 alginate (BH-adjusted p-value < 0.05), whereas gene sets with negative enrichment scores were significantly 311 enriched with genes with decreased expression on digested alginate. The number in brackets indicates the number 312 of genes with unique K number per gene set (A) and the number of genes per gene set (B) within the V. 313 cyclitrophicus ZF270 genome.

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# Growth on digested alginate is associated with increased expression of alginate catabolism, flagellar motility and chemotaxis genes

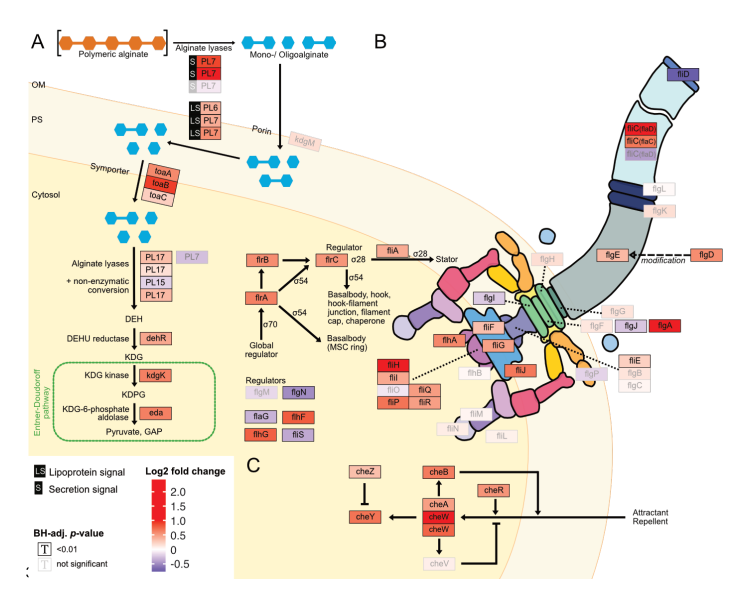
For a more fine-grained understanding of the impact of the form of alginate on the alginate catabolism,we investigated specifically the expression of genes involved in alginate degradation, uptake, and

319 catabolism. In the closed genome of V. cyclitrophicus ZF270, we identified genes that encode CAZymes 320 responsible for alginate degradation, namely alginate lyase genes from the PL6, PL7, PL15, and PL17 321 family. We identified homologs of alginate transporters (porin kdgM, symporter toaA, toaB, and toaC) 322 and metabolic enzymes that shunt into the Entner-Doudoroff pathway (DEHU reductase genes dehR, 323 kdgK, and eda), based on the known genes in the alginate degradation pathway of Vibrionaceae<sup>26,27</sup>. 324 We found that the expression of most of these genes increased significantly on digested alginate relative 325 to alginate (Fig. 4B, Fig. 5A, S8A, Table S16). Surprisingly, also the expression of most alginate lyase 326 genes increased on digested alginate, especially the expression of secreted alginate lyase genes. This 327 indicates the production of "public" exoenzymes despite the abundance of monomeric and oligomeric 328 degradation products in the digested alginate medium.

329

330 The increased motility of cells observed upon exposure to digested alginate (Fig. 3, Fig. 4A) led us to 331 evaluate the expression of motility- and chemotaxis-associated genes across digested alginate and alginate treatments. As flagella are the main mode of motility in the genus Vibrio<sup>28,29</sup>, we focused on 332 333 the expression of flagella-related genes. In the genome of V. cyclitrophicus ZF270 we found a gene 334 cluster that encodes most genes involved in flagellar assembly and that was flanked by chemotaxis 335 genes (hereon called flagellar locus, Fig. S8B, and Table S17). Overall, 21/34 flagellar locus genes were 336 differentially expressed (log2 fold change > 0.5, BH-adjusted *p*-value < 0.01) and the majority (90%) 337 of these differentially expressed genes showed increased expression on digested alginate (Fig. 4B). The 338 flagellar genes *flgA*, *fliC*, and *fliH* and the chemotaxis gene *cheW* showed the strongest overexpression 339 (5.4, 3.8, 2.0, and 2.6-fold, respectively). The expression of flagellar biosynthesis genes in Vibrionacaea occurs by a cascade of gene expression of four classes of genes (Class I - IV)<sup>30</sup>. We found the expression 340 341 of the master regulator of the flagellar biosynthesis regulon, the Class I gene flrA, to be 1.6-fold increased on digested alginate (BH-adj. p-value 3e-19)<sup>31</sup>. The Class II regulatory genes flrBC, 342 343 controlling Class III genes, and *fliA*, controlling Class IV genes, showed 1.5-fold and 1.3-fold increased 344 expression on digested alginate (BH-adj. p-value 1e-12 for flrB, 2e-39 for flrC, 6e-15 for fliA) (Fig. 345 5B)<sup>29,32</sup>. These findings suggest that the increased phenotypic motility observed on digested alginate 346 (Fig. 3) is related to the upregulation of flagellar biosynthesis genes. Additionally, the expression of the

347 flagellum filament, encoded by *fliC* genes of the flagellar locus, was partially increased: *flaD* and *flaC* expression were increased by 3.8 and 1.6-fold, whereas the *flaA* gene was not significantly differentially 348 349 expressed (BH-adj. p-value 7e-37, 2e-19, and 1e-02, respectively) (Fig. 5B, Table S6). This suggests 350 that cells grown on digested alginate have a *flaD*-rich filament composition, which has been shown to alter the swimming and adhesion characteristics of bacterial cells<sup>33,34</sup>. Lastly, we found that most genes 351 involved in chemotaxis and located in the flagellar locus are highly expressed in cells grown on digested 352 353 alginate (Fig. 5C, Fig. 4B) and likely drive the chemotactic activity of V. cyclitrophicus ZF270 during 354 dispersal. Overall, our findings elucidate that cellular responses upon exposure to degradation products 355 manifest in increased expression of genes involved in extracellular alginate breakdown and alginate 356 catabolism as well as flagellar assembly and chemotaxis.



358 Figure 5. Digested alginate increases expression of genes involved in alginate degradation, uptake and 359 catabolism, as well as flagellar assembly and chemotaxis. Genome-wide differential expression analysis where 360 the log2 fold changes of gene expression on digested alginate compared to alginate is shown for (A) alginate 361 lyases (PL6, PL7, PL15, PL17), transporters (porin kdgM, symporter toaB, symporter toaC), and metabolic 362 enzymes shunting into the Entner-Doudoroff pathway (DEHU reductase DehR, kdgK, eda), (B) genes of the 363 flagellar locus associated with flagellar assembly and (C) adjacent chemotaxis genes. Genes displayed in (B) and 364 (C) are part of the KEGG pathways "Bacterial motility proteins" and "Bacterial chemotaxis". Differential 365 expression analysis was performed to compute the Benjamini-Hochberg-adjusted Wald test p-value ("BH-adj. p-366 value", text color and box outline color) and log2 fold change (box fill color) for each gene (box). For better 367 visibility, genes that exhibited a log2 fold gene expression change greater than 1 (i.e., doubling of expression) or 368 less than -1 (i.e., halving of expression) are designated maximum intensity of red or blue, respectively. Genes with 369 BH-adj. p-value smaller than 0.01 were considered significantly differentially expressed. In (A), the location of 370 the gene products was based on Figure 1 of Wargacki et al.<sup>26</sup> with the exception of the alginate lyases (PL6, PL7, 371 PL15, PL17) which were placed based on their signal peptides (S: extracellular, LS: membrane-embedded, none: 372 cytosolic). In (B) and (C) the gene location and depiction were based on the KEGG pathway "Flagellar assembly" (map02040), "Bacterial chemotaxis" (map02030), and Figure 3 of Rajagopala et al.<sup>35</sup>. Genes without known 373 374 cellular location were omitted here but displayed in the genomic architecture in Fig. S5. Arrow: activation; dashed 375 arrow: modification; "flat" arrow: inhibition; OM: outer membrane; PM: periplasm; IM: inner membrane; PL: 376 polysaccharide lyase family; kdgM: oligogalacturonate-specific outer membrane porin; toaABC: oligoalginate 377 symporter; DEH: 4-deoxy-L-erythro-5-hexoseulose uronic acid; dehR: DEH reductase; KDG: 2-keto-3-deoxy-378 gluconate; kdgK: KDG kinase; KDPG: 2-keto-3-deoxy-6-phosphogluconate; eda: KDG-6-phosphate aldolase; 379 GAP: glyceraldehyde 3-phosphate; ED: Entner-Doudoroff; ns: not significant, i.e. BH-adj. p-value > 0.01.

### 380

#### 381 Discussion

382 On Earth organic carbon is mostly present in the form of polysaccharides<sup>1,2</sup>, which are often in a 383 particulate state and form a heterogeneous resource landscape. Over the last years, the study of 384 extracellular bacterial degradation of polysaccharides has revealed that bacterial growth on 385 polysaccharides increases with increased cell density, enabling cells to benefit from the exoenzymes 386 and extracellular degradation products of surrounding cells otherwise lost to diffusion ("cooperative 387 growth")<sup>36–38</sup>. However, cells have been observed not only to aggregate on polysaccharide sources, but 388 also to leave them before the source is depleted<sup>13,38</sup>. This has thus far been interpreted with optimal 389 foraging theory, which weighs the diminishing returns of harvesting resources from a nutrient hotspot 390 against the predation and opportunity costs which ensue<sup>39,40</sup>, but the cellular mechanisms that drive the 391 cell dispersal remain unclear.

392

393 Our work links these observations and connects them to the cellular mechanisms that underlie the 394 degradation-dispersal cycles of bacterial degraders, which we see as basal drivers of the biogeochemical 395 processing of polysaccharides in heterogeneous nutrient-scapes. When bacteria encounter a new source 396 of biomass, their local environment likely contains few mono- or oligosaccharides but is rich in 397 polysaccharides which usually require extracellular breakdown (Fig. 6, "Finding a new nutrient 398 source"). General concepts of how bacteria recognize the presence of large biopolymers remain elusive, 399 but it was proposed that "sentry" enzymes are constitutively expressed at a basal level to cleave mono-400 or oligosaccharides from polysaccharides, which cells can take up and which prime their metabolism for the degradation of the respective polysaccharide<sup>41,42</sup>. It is not known yet how widespread the concept 401 402 of sentry enzymes may be, but the observation of a constitutively expressed PL7 and PL15 family alginate lyase gene in Z. galactanivorans<sup>41</sup> is mirrored in our work by the constant expression of an 403 404 extracellular PL7 family alginate lyase gene, which may act as sentry enzyme that helps to initiate 405 alginate degradation when cells encounter alginate.

406

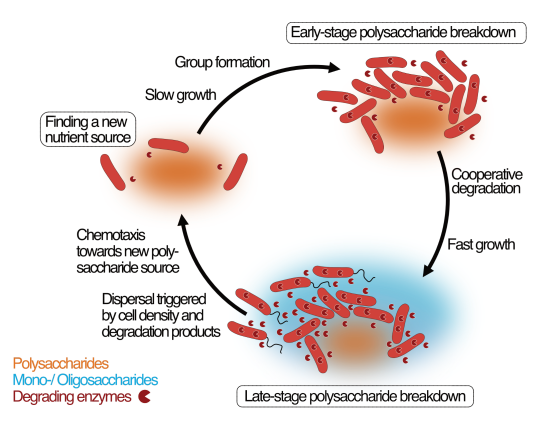
407 It has been previously found that cells grow slowly until a critical cell density for efficient cooperative polymer breakdown is reached<sup>36</sup>, observed here as the formation of large cell groups. In this stage, cells 408 409 start to benefit from the diffusing degradation products and exoenzymes of cells in their local 410 environment and reach their maximal growth rate (Fig. 6, "Early-stage polysaccharide breakdown"). 411 We found that cells grown on degradation products reached their maximal growth earlier and showed 412 increased expression of ribosomal biosynthesis, enzyme secretion, especially of secreted alginate 413 lyases, transporters, quorum sensing and expression changes in the central carbon metabolism. The 414 secretion of secreted alginate lyases might seem surprising and wasteful in a monomer-rich environment. We speculate that enhanced enzyme secretion supports the continued growth of detached bacteria's offspring that remain on the particle, as observed in the case of *V. splendus* 1AO1 during chitin particle degradation<sup>43</sup>. Another reason may be that cells primarily rely on intracellular monosaccharide levels to trigger the upregulation of genes associated with polysaccharide degradation and catabolism, as seen in *E. coli* across various carbon sources<sup>44,45</sup>, unable to modulate the expression of degrading enzymes based on the extracellular presence of monomers.

421

422 We show that cells respond to the exposure to degradation products with dispersal from dense cell 423 groups by means of increased flagellum-driven swimming (Fig. 6, "Late-stage polysaccharide 424 breakdown"), decreasing their local cell number. This finding matches with previous observations of cells leaving biopolymer particles before they are depleted<sup>13,38</sup>. A plausible explanation for this density-425 426 dependent dispersal is that cells in larger groups compete with each other for nutrients and space, while 427 not profiting from cooperative degradation anymore due to the abundance of degradation products. Motility has also been shown to increase the encounter rate of cells with sources of nutrients<sup>46,47</sup>, 428 429 suggesting motility as a strategy that allows cells to escape from the ensuing competition. Previous 430 work in Caulobacter crescentus demonstrated that a flagellum knock-out mutant formed larger cell groups, resulting in reduced growth rates due to intercellular competition<sup>4,14</sup>. While it would be 431 432 interesting to study non-motile mutants of V. cyclitrophicus ZF270, the non-tractability of natural 433 isolates makes direct tests of molecular mechanisms difficult. Additionally, subjecting cells separately 434 to the two monomeric units of alginate or oligomers of defined size could improve our understanding 435 of the specific molecules that trigger motility, but this was experimentally not feasible. We also found 436 that the presence of degradation products increased the expression of chemotaxis genes and detected 437 that cells chemotaxis toward polymeric alginate, but not digested alginate, analogous to previous 438 findings on chitin<sup>46-48</sup>. Even though this may first seem counterintuitive, it makes sense in the light of 439 the dispersal process: chemotaxis towards degradation products would prevent cells from leaving a 440 nutrient source, whereas chemotaxis towards polymers may increase encounters with a fresh nutrient source<sup>49</sup> (Fig. 6, "Finding a new nutrient source"), on which cells can re-initiate degradation. This 441

- 442 strategy allows cells to alternate between degradation and dispersal to acquire carbon and energy in a
- 443 heterogeneous world with nutrient hotspots 50-53.

#### 444



445

446 Figure 6. Bacterial growth and regulation on patches of polysaccharides. Based on our findings we propose 447 a conceptual model where degradation of polysaccharides proceeds after the encounter of a polymer source by 448 bacterial cells that have a basal exoenzyme production of "sentry" enzymes ("Encounter of a new carbon source"). 449 This phase is succeeded by group formation, which enables cells to benefit from exoenzymes of neighboring cells 450 and diffusing degradation products ("Early-stage polysaccharide breakdown"). The following phase includes 451 cooperative extracellular degradation of the polysaccharide source, further increasing the concentration of 452 available degradation products. These degradation products trigger the overexpression of alginate degrading, 453 importing, and catabolizing enzymes, ensuring swift polysaccharide degradation ("Late-stage polysaccharide 454 breakdown"). The increased pool of breakdown products also cues flagellar swimming and chemotaxis in a 455 subpopulation of cells towards new polysaccharide sources, thus restarting the cycle of degradation of polymeric 456 carbohydrates by heterotrophic bacteria. Cells and molecules are not drawn to scale. Dark red pie symbols: 457 intracellular and extracellular polysaccharide-degrading enzymes; orange shading: a polymeric carbon source; 458 blue shading: monomeric or oligomeric degradation products.

#### 459 Conclusion

460 The heterogeneous landscape of polysaccharide hotspots in natural systems requires bacteria to 461 effectively break down polymeric carbohydrates as well as readily ensure dispersal to new nutrient 462 hotspots. Our findings show that the degree of depolymerization of the polysaccharide influences this 463 decision, altering the growth dynamics, metabolic activity, and motility of cells. Our study also 464 contextualizes the surprising finding that foraging bacteria majorly leave polysaccharide particles before the last third of the particle is consumed<sup>38</sup>. Dispersal from a partially degraded carbon source 465 466 may serve several purposes: (i) escaping competition that ensues within large cell groups, (ii) ensuring the spread of a part of the clonal population to new environments as bet hedging strategy $^{40,54}$ , here 467 468 guided by chemotaxis towards new nutrient hotspots, (iii) preventing whole populations degrading a 469 sinking marine particle or a deposited sediment particle to be buried in depth where nutrient hotspots 470 become sparse<sup>38,55</sup>, and/or (iv) increase the genetic variation in bacterial populations<sup>40</sup>. However, 471 dispersal may also occur when a nutrient source offers a surplus of carbon while other essential nutrients 472 become limiting, as the increased expression of iron, zinc, and phosphate transporters in cells grown on 473 digested alginate suggested. These findings emphasize that metabolic molecules can also act as triggers 474 of dispersal, expanding upon the current perspective of dispersal in biofilms as a reaction to dispersal cues like NO, signalling molecules, nutrient starvation, and oxygen starvation<sup>56</sup>. The study of bacterial 475 476 motility on increasingly complex biomass particles will reveal the role of the nutrient composition of 477 the present nutrient hotspot on the bacterial decision-making. Overall, these new insights into the 478 cellular mechanisms and regulation that drive degradation-dispersal cycles contribute to our 479 understanding of the microbially driven remineralization of biomass, and factors that modulate this 480 process. The open questions of how bacteria sense polysaccharides in their environment, which cell 481 signaling pathways integrate the presence of degradation products in the cellular decision-making of 482 degradation and dispersal, and to what extent cell populations coordinate this decision, present an 483 exciting avenue of further research.

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- 485
- 486

#### 487 Materials and Methods

#### 488 Bacterial strains, media and growth assays

Vibrio cyclitrophicus ZF270 (available through Culture Collection Of Switzerland; Accession number: 489 490 2043) cells were cultured in Marine Broth (DIFCO) and grown for 18 hours at 25 °C. Cells from these 491 cultures were used for growth experiments in Tibbles Rawling (TR) salts minimal medium<sup>57,58</sup> 492 containing either 0.1% (weight/volume) algae-derived alginate (referred to as "alginate") (Sigma 493 Aldrich, CAS-number 9005-38-3) or 0.1% (weight/volume) digested alginate. The digested alginate 494 was produced by enzymatically digesting 2% alginate with 1 unit ml<sup>-1</sup> of alginate lyase (Sigma Aldrich, 495 CAS-number 9024-15-1) at 37 °C for 48 hours. Carbon sources were prepared in nanopure water and 496 filter sterilized using 0.40 µm Surfactant-Free Cellulose Acetate filters (Corning, USA). Well-mixed 497 batch experiments in alginate or digested alginate medium were performed in 96-well plates (Greiner 498 Bio) and growth dynamics were measured using a microwell plate reader (Biotek, USA). Assays were initiated as described previously<sup>59</sup>. Briefly, 1 ml from a culture grown for 18 h on Marine broth was 499 500 centrifuged at 5000 g in 1.5 ml microfuge tubes for 5 minutes. The supernatant was discarded and the 501 cell pellet was subjected to two rounds of washing with the basal TR salts medium. The cell-pellet was 502 then resuspended in 1 ml of TR salts medium and 5  $\mu$ l of this suspension inoculated into 195  $\mu$ l TR medium with either carbon source ( $\sim 10^5$  colony forming units (CFUs ml<sup>-1</sup>) in a 96-well plate (Greiner 503 504 Bio). The optical density (600 nm) was then measured every 15 minutes for 40 hours. All measurements 505 had six biological replicates.

506

#### 507 *Alginate oligosaccharide measurements*

508 Oligosaccharide measurements were performed using liquid chromatography time of flight mass 509 spectrometry (LC-QTOF-MS). Samples were prepared by diluting 1:20 in milliQ water and 5  $\mu$ L of 510 sample was injected per measurement. Chromatographic separation was performed using an Agilent 511 1290 stack, using an Agilent HILIC-Z column (2.7  $\mu$ m particles, 2.1 x 50 mm). Mobile phase A 512 contained 10% acetonitrile (Fisher Scientific) and 0.1% medronic acid (Agilent), and Mobile phase B 513 contained 90% acetonitrile and 0.1% medronic acid. The separation was performed as follows: Mobile 514 phase B 100% for 1 minute, gradient to 30% phase B over 3 minutes, 30% phase B for 30 seconds, and

515	equilibration of 100% phase B for 5 minutes. The flow rate was 400 $\mu$ L min <sup>-1</sup> at 30 °C. Samples were
516	measured using an Agilent 6520 mass spectrometer in negative mode, in 4 GHz high-resolution mode.
517	Data analysis was performed in Agilent Quantitative Analysis software.

518

#### 519 *Microfluidics and time-lapse microscopy*

Microfluidic experiments and microscopy were performed as described previously<sup>14,16,60</sup>. Cells were 520 521 imaged within chambers of a PDMS (Sylgard-Dow) microfluidic chip that ranged in size from 60-120  $\times$  60  $\times$  0.56 µm ( $l \times b \times h$ ). Within these chambers, cells can attach to the glass surface and experience 522 523 the medium that diffuses through lateral flow channels. Imaging was performed using IX83 inverted 524 microscope systems (Olympus, Japan) with automated stage controller (Marzhauser Wetzlar, 525 Germany), shutter, and laser-based autofocus system (Olympus ZDC 2). Chambers were imaged in 526 parallel on the same PDMS chip, and phase-contrast images of each position were taken every 8 or 10 527 minutes. The microscopy unit and PDMS chip were maintained at 25 °C using a cellVivo microscope 528 incubation system (Pecon GmbH).

529

#### 530 Motility assays

531 Cells were grown for 10 hours in Marine Broth (DIFCO) after which 10  $\mu$ l of culture was used to 532 inoculate culture tubes (Greiner) containing 5 ml of TR medium with either 0.1% alginate or 0.1% 533 digested alginate. After 6 hours of growth at 25 °C, 2  $\mu$ l of cell suspension was inoculated into 534 microfluidic growth chambers. Cells within six replicate chambers were then imaged with the phase-535 contrast channel at a high frame rate (125 Hz, i.e., frames s<sup>-1</sup>) using the same microscopy setup described 536 above.

537

#### 538 *Chemotaxis assays*

539 To assess whether polymeric alginate and digested alginate attract *Vibrio cyclitrophicus* ZF270, we 540 used the *In Situ* Chemotaxis Assay<sup>17,18</sup> (ISCA), a microfluidic device consisting of a  $5 \times 5$  array of 541 microwells that can be individually loaded with solutions of different chemicals (110 µl each). Once 542 the ISCA is deployed in an aqueous environment, the chemicals diffuse out of the wells through a small 543 port, creating chemical gradients which will guide chemotactic bacteria inside the wells of the device<sup>17,18</sup>. Vibrio cvclitrophicus ZF210 was plated on Marine Agar (BD Difco) from a glycerol stock 544 545 and grown for 16 h at 27 °C. A single colony was then incubated in 10% Marine Broth (BD Difco) in 546 0.22-µm filtered artificial seawater (Instant Ocean, Spectrum Brands) and grown overnight at 27 °C and 547 180 rpm. The culture was diluted down to 1 x 10<sup>6</sup> cells ml<sup>-1</sup> in 0.22 µm filtered artificial seawater (Instant 548 Ocean, Spectrum Brands) to perform the chemotaxis experiment. Both chemoattractants (alginate and digested alginate) were diluted in sterile seawater (35 g l<sup>-1</sup>; Instant Ocean, Spectrum Brands) at a final 549 550 concentration of 0.1% and then filtered with a 0.2 µm filter (Millipore) to remove particles and potential 551 contaminants. Within the ISCA, one full row of five wells was used per chemoattractant as technical 552 replicates. The chemoattractants were injected in triplicate ISCA with a sterile 1 ml syringe (Codau) 553 and needle (27 G, Henke Sass Wolf). A last row containing 0.2 µm-filtered seawater acted as negative 554 control accounting for cells swimming in the device by random motility only. Experiments were 555 conducted by incubating the ISCAs for 1 h in the diluted Vibrio cyclitrophicus ZF270 culture. Upon 556 time completion, a sterile syringe and needle were used to retrieve the content of the wells and 557 transferred to 1 ml microfuge tubes resulting in a pooling of a row of 5 wells containing the same 558 sample. Sample staining was performed with SYBR Green I (Thermo Fisher) and the chemotactic 559 response was quantified by counting cells using flow cytometry. The strength of the chemotactic 560 response was determined by the mean chemotactic index  $(I_{\rm C})$ , defined as the ratio of the number of cells 561 found in each chemoattractant to the number of cells in control wells containing filtered seawater (so 562 that attraction corresponds to  $I_{\rm C} > 1$ ).

563

#### 564 *Culturing and harvesting cells for transcriptomics*

Cells were grown for 18 hours in Marine Broth (DIFCO) after which 1 ml of culture was centrifuged at 5000 g in 1.5 ml microfuge tubes for 5 minutes. The supernatant was discarded and the cell pellet was subjected to two rounds of washing with the basal TR salts medium. The cell pellet was then resuspended in 1 ml of TR salts medium and 250  $\mu$ l of this suspension were used to inoculate 100 ml flasks (Schott-Duran) containing 10 ml of TR medium with either 0.1% alginate or 0.1% digested alginate. This was done in parallel for six flasks. Once cultures in the flasks reached mid-exponential

571 phase (10 hours and 15 hours after inoculation for digested alginate and alginate, respectively) and had 572 approximately the same OD (0.39 for digested alginate and 0.41 for alginate), 2 ml of cultures were 573 harvested for RNA extraction. Samples were stabilized with the RNprotect reagent (Qiagen) and RNA 574 was extracted using the RNeasy mini kit (Qiagen).

575

576 Sequencing and gene annotation of Vibrio cyclitrophicus ZF270

577 Long read sequencing using the Oxford Nanopore Platform (Long read DNA sequencing kit) and short 578 read sequencing using the Illumina platform (Illumina DNA Prep kit and IDT 10bp UDI indices, and 579 sequenced on an Illumina NextSeq 2000 producing 2x151bp reads) was performed by the Microbial 580 Genome Sequencing Center, Pittsburgh, USA (MiGS), to create a new closed reference genome of V. 581 cyclitrophicus ZF270 (BioProject PRJNA991487). Annotation of this genome was done with RASTtk 582 (v2.0, Rapid Annotation using Subsystem Technology tool kit<sup>61</sup>). Additionally, KEGG Ontology identifiers ("K numbers") were annotated with BlastKOALA (v2.2)<sup>62</sup>. Dedicated annotation of alginate 583 584 lyase genes was performed by homology search for proteins belonging to the PL5, PL6, PL7, PL14, PL15, PL17, PL18, PL31, PL36, or PL39 family<sup>63</sup> by dbCAN2 (v9.0)<sup>64</sup>. Enzymes for alginate transport 585 586 and metabolism were identified by BLASTn-search<sup>65,66</sup> of gene sequences of Vibrio splendidus 12B01, 587 which were previously identified as minimum genetic prerequisites for alginate utilization and enabled alginate degradation when cloned into  $E. \ coli^{26}$ . 588

589

590 *Location prediction of alginate lyases* 

591 Signal peptides were annotated using SignalP (v.5.0)<sup>67</sup>, and LipoP (v.1.0)<sup>68</sup>. SignalP discriminated 592 between 1) Sec/SPI: "standard" secretory signal peptides transported by the Sec translocon and cleaved 593 by Signal Peptidase I (SPI), 2) Sec/SPII: lipoprotein signal peptides transported by the Sec translocon 594 and cleaved by Signal Peptidase II (SPII), and 3) Tat/SPI: Tat signal peptides transported by the Tat 595 translocon and cleaved by SPI. LipoP discriminates between 1) SPI: signal peptide, 2) SpII: lipoprotein 596 signal peptide, and 3) TMH: n-terminal transmembrane helix. All predictions were in agreement, apart 597 from one PL7 (gene 1136176.5.peg.4375) which was predicted by LipoP as cytoplasmic and by SignalP 598 as equally likely cytoplasmic as containing a lipoprotein signal peptide.

599

600 Transcriptomic analysis: Sequencing, pre-processing, differential expression analysis, and functional
601 analysis

Sequencing (12 M reads, 2 x 50 bp) of the isolated RNA was performed by MiGS after rRNA depletion using RiboZero Plus (Ilumina). cDNA libraries were prepared using an Illumina DNA Prep kit and IDT 10 bp UDI indices, and sequenced on an Illumina NextSeq 2000. Preprocessing of the raw reads was carried out as follows: Quality control was performed with FastQC (v0.11.9)<sup>69</sup> and reads were trimmed with Trimmomatic (v0.38)<sup>70</sup>; the high-quality reads were mapped to the reference genome (described above) with Bowtie2 (v2.3.5.1)<sup>71</sup>; binarization, sorting, and indexing were done with Samtools (v1.10)<sup>72</sup>; gene counts were computed with the featureCount function of Subread (v2.0.1)<sup>73</sup>.

Differential expression analysis was performed with DESeq2  $(v1.30.1)^{74}$ . In brief, DESeq2 normalizes 609 610 the raw read counts with normalization factors ("size factors") to account for differences in sequencing 611 depth between samples. Subsequently, gene-wise dispersion estimates are computed for each gene 612 separately using maximum likelihood, and then shrunk toward the values predicted by the dispersion-613 mean dependence curve to obtain final dispersion values. Finally, DESeq2 fits a negative binomial 614 model to the read counts and performs significance testing using the Wald test. Here reported *p*-values 615 result from the Wald test of read counts from the digested alginate condition compared to the alginate condition and were adjusted for multiple testing by Benjamini-Hochberg correction<sup>75</sup> as implemented 616 617 in the p.adjust function of base R (v4.1.2). The reported log2 fold changes indicate the log2(DESeq2-618 normalised reads in digested alginate condition / DESeq2-normalised reads in alginate condition) for 619 each gene.

620 Visualization of gene maps was performed in R with the ggplot2 package (v3.4.0) and the extension621 gggenes (v0.4.1) by David Wilkins.

For systematic functional analysis we performed gene set enrichment analysis  $(GSEA)^{19}$  using the fgsea function of the fgsea package (v1.20.0) with minimal number of unique genes per gene set "minSize" = 5 and number of permutations "nPermSimple" = 1000000. In brief, GSEA takes the full gene list ranked by log2 fold change and annotated with K numbers as input and determines whether the member genes of any KEGG pathway are randomly distributed throughout the ranked gene list or whether they

are primarily found at the top or bottom<sup>19</sup>. This is quantified by the enrichment score (ES), which 627 628 corresponds to a weighted Kolmogorov-Smirnov-like statistic. The ES of each gene set is normalized 629 to the mean enrichment of random samples of the same size to account for the size of the set, yielding 630 the normalized enrichment score (NES). To estimate the significance level of the enrichment score, the 631 *p*-value of the observed enrichment score is calculated relative to a null distribution that was computed 632 from permuted data. The estimated significance level was adjusted to account for multiple hypothesis 633 testing. As gene sets we chose all KEGG pathways and KEGG BRITE categories (as noted in Table S1 634 in column "KEGG pathway") within all genes of V. cyclitrophicus ZF270 annotated with a KEGG 635 Ontology identifier ("K number"). Visualization of differential expression levels in KEGG pathways was performed with the R package pathview  $(v1.35.0)^{76}$ . We also formed gene sets of the genes 636 637 associated with alginate utilization (see "Sequencing and gene annotation of Vibrio cyclitrophicus 638 ZF270", Table S16), of the genes of the flagellar locus that map to the KEGG pathways "Bacterial 639 motility proteins", and of the genes of the flagellar locus that map to the KEGG pathways "Bacterial 640 chemotaxis" (as noted in Table S17) within all genes of V. cyclitrophicus ZF270.

641

#### 642 Image analysis

643 Cells within microscopy images were segmented and tracked using ilastik (v1.3) ("pixel classification 644 workflow" and "tracking with probabilities workflow"). Phase contrast images were used for alignment, 645 segmentation, tracking and linking. Images were cropped at the boundaries of each microfluidic 646 chamber. The lineage identity of each single cell was assigned by ilastik's tracking plugin and visualized 647 by coloring the segmented cells, respectively. The growth rate of each cell was computed as the change 648 of cell area over time, i.e., via a linear regression of the single-cell area over the time between 649 consecutive cell divisions, based on ilastik's segmentation and tracking output. Cells that were tracked 650 over less than three frames were excluded. Measurement of swimming speeds and displacement of cells 651 was performed using ilastik (v1.4), ImageJ (v2.3) and Trackmate (v7.5.2). Briefly, cell-segmentation 652 ("pixel classification workflow" in ilastik) and tracking ("animal tracking workflow" in ilastik) were 653 performed using the high frame rate phase contrast images in ilastik (v1.4). Cell trajectories and 654 properties were then computed using the output of the ilastik workflow in Trackmate.

655

#### 656 Dispersal analysis

For the analysis on the dispersal of cells (Fig. 2), we computed the cell number as the total number of cells within a microfluidic chamber. The change in the number of cells was computed by subtracting the number of cells before the medium switch (i.e., average number of cells between t = 1.9 to 2.1 h) from the number of cells after the medium switch (i.e., average number of cells between t = 3.9 to 5.5 h).

662

#### 663 Datasets and statistical analysis

664 All batch experiments were replicated 3 to 6 times. Growth curves were analyzed in Python (v3.7) using the Amiga package (v1.1.0)<sup>77</sup> and GraphPad Prism (v8, GraphPad Software). The microscopy dataset 665 666 consisted of eight chambers each, corresponding to the eight replicates shown in Figure 1 and Figure 2. 667 These were grouped into two biological replicates wherein each biological replicate was fed by media 668 through a unique channel in a microfluidic chip. Cells with negative growth rates were excluded from 669 the analysis after visual curation, as they represented artefacts, mistakes in segmentation or linking 670 during the tracking process, or non-growing deformed cells. Each chamber was treated as an 671 independent replicate. Comparisons were considered statistically significant when p < 0.05 or when the 672 False Discovery Rate (FDR)-corrected q was smaller than 0.05. FDR corrections were applied when 673 multiple t tests were performed for the same dataset. Measures of effect size are represented by the  $R^2$ 674 or eta<sup>2</sup> value. All statistical analyses were performed in GraphPad Prism v9.0 (GraphPad Software, 675 USA), R v4.1.2, RStudio v1.1.463 (Posit, USA).

676

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682

683 Data Availability: Sequencing data is available on NCBI, BioProject PRJNA991487, upon publication. 684 further All data deposited ERIC Open (https://opendata.eawag.ch/, is on 685 https://doi.org/10.25678/0008MH), available upon publication. All code will be made publicly 686 accessible on https://github.com/Microbial-Systems-Ecology/publications upon publication.

687

688 **Competing interests:** The authors declare no competing interests.

689

690 Author Contributions: GD, AS, JS, and JK conceived the research along with MA. GD designed and

691 performed the experiments. AS developed and implemented the algorithms to compute single-cell

692 growth rates and visualize lineage identities from segmented and tracked cells. AS analyzed and

693 visualized the RNA-seq data. JK conceived and implemented the cell dispersal analysis. EC conducted

- the ISCA experiment. SP carried out the mass spectrometry measurements. AS and GD analyzed the
- data with inputs from JK, JS, CM, MA, and RS. AS wrote the manuscript with inputs from GD, OS,

696 MA, JK, JS, EC, SP, CM, and RS.

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698

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