Formation of spatial patterns of cells is a recurring theme in biology and often depends on regulated cell motility. Motility of the rod-shaped cells of the bacterium *Myxococcus xanthus* depends on two motility machineries, type IV pili (giving rise to S-motility) and the gliding motility apparatus (giving rise to A-motility). Cell motility is regulated by occasional reversals. Moving *M. xanthus* cells can organize into spreading colonies or spore-filled fruiting bodies, depending on their nutritional status. To ultimately understand these two pattern-formation processes and the contributions by the two motility machineries, as well as the cell reversal machinery, we analyse spatial self-organization in three *M. xanthus* strains: (i) a mutant that moves unidirectionally without reversing by the A-motility system only, (ii) a unidirectional mutant that is also equipped with the S-motility system, and (iii) the wild-type that, in addition to the two motility systems, occasionally reverses its direction of movement. The mutant moving by means of the A-engine illustrates that collective motion in the form of large moving clusters can arise in gliding bacteria owing to steric interactions of the rod-shaped cells, without the need of invoking any biochemical signal regulation. The two-engine strain mutant reveals that the same phenomenon emerges when both motility systems are present, and as long as cells exhibit unidirectional motion only. From the study of these two strains, we conclude that unidirectional cell motion induces the formation of large moving clusters at low and intermediate densities, while it results in vortex formation at very high densities. These findings are consistent with what is known from self-propelled rod models, which strongly suggests that the combined effect of self-propulsion and volume exclusion interactions is the pattern-formation mechanism leading to the observed phenomena. On the other hand, we learn that when cells occasionally reverse their moving direction, as observed in the wild-type, cells form small but strongly elongated clusters and self-organize into a mesh-like structure at high enough densities. These results have been obtained from a careful analysis of the cluster statistics of ensembles of cells, and analysed in the light of a coagulation Smoluchowski equation with fragmentation.

**Keywords:** collective migration; spatio-temporal pattern formation; self-organization

1. **INTRODUCTION**

Formation of patterns of spatially organized cells is a recurring theme in biology. These processes often depend on regulation of cell motility. For instance, in metazoans, it provides the basis for organ formation during embryogenesis, and in single-celled eukaryotes such as *Dictyostelium discoideum*, it is essential for the formation of fruiting bodies. In bacteria, regulated cell motility is essential for the colonization of diverse habitats as well as for the formation of multicellular structures such as biofilms and fruiting bodies. These pattern-formation processes can be self-organized. For example, *D. discoideum* [1,2] regulates cell aggregation and multicellular organization by secreting and sensing...
the diffusive signal cAMP. In *Myxococcus xanthus*, on the other hand, rippling patterns and the highly complex cellular reorganization leading to fruiting body formation are controlled by a non-diffusing signal, the C-signal \[3\]. Interestingly, collective effects and self-organization can also occur, to a certain extent, in the absence of an explicit signalling mechanism. For instance, hydrodynamic interactions can induce large-scale coherent motion of swimming cells, as recently observed in *Bacillus subtilis* \[4,5\], and a density-dependent diffusivity can lead to aggregation patterns as recently suggested to occur in *Escherichia coli* and *Salmonella typhimurium* \[6\].

*Myxococcus xanthus* is a gliding bacterium that has been used as a model system to study pattern formation \[7\], bacterial social behaviour \[8\] and motility \[9\]. The rod-shaped cells of the bacterium *M. xanthus* move on surfaces in the direction of their long axis using two motility machineries, type IV pili, which requires cell-to-cell contact for its activity because it is stimulated by exopolysaccharides on neighbouring cells \[10\] (giving rise to S-motility), and the gliding motility apparatus that allows cells to move in isolation \[11\] (giving rise to A-motility). Force generation by the A-motility system has been suggested to rely either on slime secretion from the lagging pole \[12\], or on focal adhesion complexes distributed along the cell \[13\]. Cells occasionally reverse their gliding direction with an average frequency of about once per 10 min and the reversal frequency is controlled by the *Frz* chemosensory system \[14\]. In the presence of nutrients, *M. xanthus* cells form coordinate spreading colonies. Upon depletion of nutrients, *M. xanthus* cells initiate a complex developmental programme that culminates in the formation of spore-filled fruiting bodies. Both motility systems as well as reversals are required for the two cellular patterns to form, i.e. spreading colonies and fruiting bodies. It is currently not known how the reversal frequency is regulated except that cell–cell contacts may induce C-signal exchange, which is supposed to stimulate reversals during rippling and to inhibit reversals during aggregation. During fruiting body formation the reversal frequency decreases up to a point where cell movements become nearly unidirectional \[15\] and cells start to display collective motion with the formation of large clusters in which cells are aligned in parallel, making side-to-side as well as head-to-tail contacts and move in the same direction \[16\]. Eventually, cells start to aggregate. Aggregation centres often resemble a cell vortex, at their initial phase.

Here, we aim at understanding myxobacterial pattern-formation processes, particularly the contributions by the two motility machineries as well as the cell reversal machinery to the spatial organization of the cells. We study the role of steric interactions, cell adhesion and reversal frequency in the collective dynamics. The question for us is not ‘why’ cells exhibit a given collective behaviour but ‘how’ they do it. In order to identify the role of the two motility machineries and cell reversal machinery, we follow a bottom-up strategy by looking at the collective dynamics of different mutants of increasing complexity. We analyse three *M. xanthus* strains: (i) a mutant that moves unidirectionally without reversing by the A-motility system only—a mutant that has been previously studied by us \[17\], (ii) a unidirectional mutant that is also equipped with the S-motility system, and (iii) the wild-type that, in addition to the two motility systems, occasionally reverses its direction of movement. We characterize the macroscopic patterns mainly through the cluster statistics, in particular in terms of cluster size and shape. We observe that the mutant moving by means of the A-engine displays only collective motion in the form of large moving clusters. The study of its cluster size distribution (CSD) reveals that above a given density, clusters can be arbitrarily large \[17\]. Here, we show in addition that there is a non-trivial scaling of cluster perimeter with cluster size that indicates that the clustering process is neither (fully) random nor as in (equilibrium) liquid–vapour drops \[18\]. We also find that at high densities the collective dynamics changes and cells organize into vortices. The study of the two-engine strain mutant reveals the same phenomenology for these bacteria: collective motion in the form of large moving clusters, a critical density above which clusters can be arbitrarily large, a non-trivial scaling of cluster perimeter with cluster size and vortex formation at high densities. By comparing these two strains, we conclude that unidirectional cell motion induces the formation of large moving clusters at low and intermediate densities, while it results in vortex formation at very high densities (see figure 1a,b). Interestingly, similar collective dynamics has been observed in self-propelled rod models \[19\], a fact that strongly suggests that the combined effect of self-propulsion and volume exclusion interactions is the pattern formation mechanism leading to the observed phenomena.

The study of wild-type cells indicates that cell reversal weakens clustering. Wild-type cells exhibit exponential CSDs at low and intermediate densities, while the scaling of the cluster perimeter with cluster size indicates that clusters are strongly elongated. At high densities, we find that reversing wild-type cells self-organize into a mesh-like structure (see figure 1c).

Wild-type cells, as commented already, exhibit a large variety of self-organized patterns depending on the environmental condition. Our results suggest that only by switching on and off the reversal can cells modify dramatically their collective behaviour, with the suppression of cell reversal leading to collective motion in the form of moving clusters and vortex formation at high densities. This observation is consistent with the observed decrease in reversal frequency in the wild-type upon nutrient depletion, which is followed by the formation of large moving clusters and aggregation of cells. Our findings indicate that these two processes can result from simple steric interactions of the (non-reversing) rod-shaped cells, without the need of invoking any biochemical signal regulation.

The paper is organized as follows. In §2.1, we focus on the spatial self-organization of purely A-motile cells in the absence of cell reversals. The effects induced by the S-motility engine, which include increased cell adhesion, are studied in §2.2, while those due to

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cell reversals in §2.3. In §3, we discuss which collective effects are expected in self-propelled rod models, and interpret the cluster statistics results observed in the experiments in the light of a simple cluster formation theory. We summarize all the results in §4, where we also discuss the implications of the reported findings.

2. CLUSTER STATISTICS

2.1. A-motile non-reversing cells

We start out with the simple mutant A⁺S⁻Frz⁻ that only moves by means of the A-motility system and which is unable to reverse due to an insertion in the frz gene cluster (see §5 for more details about how the strain was generated). This mutant is unable to assemble type IV pili due to deletion of the pilA gene, which encodes the type IV pili subunit, and therefore the S-motility system is non-functional in this mutant. This mutant exhibits relatively weak cell–cell adhesion owing to the lack of type IV pili and the reduced accumulation of exopolysaccharides. This mutant is labelled A⁺S⁻Frz⁻ to indicate that the A-motility engine is on, the S-motility engine is off and the Frz system, i.e. cell reversal, is dramatically reduced.

Control experiments showed that these mutants have a reversal period $\gg 100$ min, whereas the isogenic Frz⁺ strain reversed with a mean reversal period of $\approx 10$ min. In Peruani et al. [17], we showed that this mutant exhibits a transition to a collective motion phase at high enough densities by analysing the dependency of CSD with the packing fraction. Here, we characterize in addition the cluster shape, and show that at densities higher than the one studied in Peruani et al. [17], giant clusters turn into vortices.

Experiments were performed by spotting a drop of cell suspension of the desired density on an agar surface to subsequently monitor the evolution of cell arrangements by taking snapshots of the bacterial colony every 30 min for a total of 8 h. Experiments with cells gliding in isolation indicate an average velocity of $v = 3.10 \pm 0.35 \mu \text{m min}^{-1}$, an average width of about $W = 0.7 \mu \text{m}$ and an average length of $L = 6.3 \mu \text{m}$. This results in a mean aspect ratio of $\kappa = L/W = 8.9 \pm 1.95$ and a cell covering an average area $a = 4.4 \mu \text{m}^2$.

We found that under these conditions, cells organized over time into moving clusters. Time-lapse recordings showed that collisions of cells lead to effective alignment (figure 2d). When the interaction is such that cells end up parallel to each other and move...
in the same direction, they migrate together for a long time (typically >15 min). Eventually, successive collisions allow a small initial cluster to grow in size (figure 2). In the individual clusters, cells are aligned in parallel to each other and arranged in a head-to-tail manner, as previously described [20]. In a cluster, cells move in the same direction. Cluster–cluster collision typically leads to cluster fusion, whereas splitting and break-up of clusters rarely occur. On the other hand, individual cells on the border of a cluster often spontaneously escape from the cluster. These two effects, cluster growth due to cluster–cluster collision and cluster shrinkage, mainly due to cells escaping from the cluster boundary, compete and give rise to a characteristic CSD.

The CSD—\( p(m,t) \)—indicates the probability of a bacterium to be in a cluster of size \( m \) at time \( t \). Note that throughout the text, the term CSD always refers to this definition. Often, the CSD is alternatively determined as the number \( n_m(t) \) of clusters of size \( m \) at time \( t \). There is a simple relation between these two definitions: \( p_m(t) \propto m n_m(t) \). In experiments, we have observed that the CSD mainly depends on the packing fraction \( \eta \), where \( \eta \equiv \rho a \), with \( \rho \) the (two-dimensional) cell density and \( a \) the average covering area of a bacterium given above. Hence, for all snapshots, first the packing fraction was determined. Then, images with similar packing fraction \( \eta \) were compared and the CSD was reconstructed by determining the CSD for all images within a finite interval of the packing fraction. Very importantly, we find that the CSD \( p(m,t) \) reaches a steady state \( p(m) \) after some transient time, as shown in figure 3. We conclude that the clustering process evolves towards a dynamic equilibrium, where the process of formation of cell clusters of a given size is balanced by events in which clusters of this size disappear either by fusing with other clusters or by losing individual cells from their boundary.

The steady-state CSD \( p(m) \) strongly depends on the packing fraction \( \eta \), with more and more cells moving in larger clusters for increasing packing fraction \( \eta \). This is evident in figure 4, where we observe that at small values of \( \eta \), \( p(m) \) exhibits a monotonic sharp decay with \( m \), while at large \( \eta \) values, \( p(m) \) is non-monotonic, with an additional peak at large cluster sizes. The solid curves in figure 4 are fitted to the raw data by using phenomenological functional forms described in the next section. The CSD here was determined at a fixed time (450 min) after the beginning of each experiment; control experiments at other times (360 and 480 min) revealed practically identical behaviour. We interpret the presence of a peak at large values of \( m \) at bigger values of the packing fractions as the emergence of collective motion resulting in formation of large clusters of bacteria moving in a coordinated fashion. The clustering transition is evident by the functional change displayed by \( p(m) \) monotonically decreasing with \( m \) for small values of \( \eta \), while exhibiting a local maximum at large \( \eta \) values. At a critical value \( \eta_c = 0.17 \pm 0.02 \) that separates different regimes of behaviour, the CSD can be approximated by \( p(m) \propto m^{-\gamma_c} \), with \( \gamma_c = 0.88 \pm 0.07 \). Control experiments with non-motile cells do not exhibit a power-law behaviour in the CSD. For more details, we refer the reader to [17]. Hence, we conclude that without active motion of cells, no comparable transition to clustering occurs. In other words, active motion is required for the dynamical self-assembly of cells.

Now, we turn our attention to the cluster shape, in particular to the scaling of the cluster perimeter \( \Pi(m) \) with the cluster size \( m \). This kind of information can help us to realize how adhesive cells are and which role adhesion plays in the clustering process. If there is surface tension, then clusters should tend to minimize their surface, and they should be round, as observed in liquid–vapour drops [18]. On the other hand, if surface tension is negligible, a cluster can be a very elongated object, with most of the cells on the cluster boundary, and the cluster perimeter is proportional to cluster size. We assume that \( \Pi(m) \propto m^\omega \), where \( m \) denotes the area of the cluster. Thus, it is clear that perimeter exponent \( \omega \) should be 0.5 for round clusters. This would be the case for very adhesive cells exhibiting random movements. If clusters are extremely elongated, then \( \omega = 1 \). We notice that \( \omega = 1 \) would correspond also to a fully random process as observed in percolation theory [18]. In short, the exponent \( \omega \) is then such that \( 0.5 \leq \omega \leq 1 \). Figure 5 shows that for \( \Delta^+S^{-} Frx^{-} \) cells \( \omega = 0.60 \pm 0.03 \), which indicates that the clustering process is non-trivial, that it is neither fully random nor dominated by surface tension (see also figure 1). The scaling of \( \Pi(m) \) with \( m \) plays a central role in the clustering theory we discuss below, where the relation between cluster size statistics and cluster perimeter statistics will be discussed in detail.

As the density increases, typically above \( \eta > 0.26 \), cells do not organize into large moving clusters, and giant clusters evolve into vortices. These vortices are
of cells as suggested in Holmes et al. Notice that these vortices are not disordered aggregates but form by one or several layers of rotating discs whose radii diminish the higher the disc is located in the z-direction. Figure 6 shows a typical example of vortex formation. See the electronic supplementary material for a movie of a vortex of A⁺S⁻Frz⁻ cells at 20× magnification. The real-time duration of the movie is 15 min. Notice that these vortices are not disordered aggregates of cells as suggested in Holmes et al. [21]. Given the fact that vortices are multi-layered structures, phase contrast imaging can provide only limited information regarding the actual cell arrangements inside vortices. A detailed study of vortices requires more sophisticated experimental techniques.

Interestingly, vortex formation has also been observed in other experimental ‘self-propelled rod’ systems as actin–myosin motility assays [22,23] as well as in the actual cell arrangements inside vortices. A detailed study of vortices requires more sophisticated experimental techniques.
We turn our attention to the next simplest mutant:

### 2.2. A- and S-motile, non-reversing cells

Required for the organization of cells into vortices. The S-motility system depends on type IV pili [10]. It is found in the wild-type, while cell reversals are absent.

Figure 5. (a–c) Cluster perimeter $P(m)$ as a function of the cluster size $m$ for three bacterial strains. The A$^+$S$^-$Frz$^-$ and A$^+$S$^+$Frz$^-$ mutants exhibit roughly the same scaling $P(m) \propto m^\omega$, with $\omega \approx 0.60$ for the A$^+$S$^-$Frz$^-$, $\omega \approx 0.62$ for the A$^+$S$^+$Frz$^-$, suggesting that an increase in adhesion does not have a strong impact on the cluster shape. On the other hand, cell reversals lead to much more elongated clusters, as the scaling of the A$^+$S$^+$Frz$^-$ cells indicates, with $\omega \approx 0.82$.

Figure 6. Vortex formation (circular pattern in the centre) in the non-reversing mutant A$^+$S$^-$Frz$^-$ at high cell density ($\eta > 0.26$). The pattern consists of rotating stacked discs of cells. These structure are observed in both A$^+$S$^-$Frz$^-$ and A$^+$S$^+$Frz$^-$ strains.

Two-dimensional suspensions of sperms [24]. In the latter example, hydrodynamical interactions are supposed to induce the observed pattern, while in the former one the role of hydrodynamic interactions is not well understood; yet in both types of systems, the vortex patterns correspond to vortex arrays. In myxobacteria, on the other hand, hydrodynamical effects can be neglected and vortices do not emerge in a lattice-like arrangement, but rather in a disorganized fashion. At a theoretical level, vortices have been found in active gel theory [25,26]. Whether active gel vortices and those observed in M. xanthus mutants have the same microscopic origin is unclear, but certainly a possibility worth exploring.

In summary, the finding of vortex formation in experiments with A$^+$S$^-$Frz$^-$ indicates that the S-motility system, cell-to-cell signalling and cell reversals are not required for the organization of cells into vortices.

### 2.2. A- and S-motile, non-reversing cells

We turn our attention to the next simplest mutant: A$^+$S$^+$Frz$^-$). These cells contain both motility engines found in the wild-type, while cell reversals are absent. The S-motility system depends on type IV pili [10]. It allows cells to move in a contact-dependent manner, i.e. cells have to be in close proximity for S-motility to become active. As previously reported [10], we find that A$^+$S$^-$Frz$^-$ cells are more adhesive. Our aim is to understand whether the S-motility engine affects the spatial self-organization of cells. We performed the same analysis on A$^+$S$^+$Frz$^-$ cells as described for A$^+$S$^-$Frz$^-$ cells and investigate cell densities close to the obtained critical density. Figure 1 shows that at least at first glance the cluster statistics resembles that obtained with A$^+$S$^-$Frz$^-$ cells. This suggests that the additional motility including its adhesion effect has no significant impact on the organization of cells within a cluster. By looking in more detail at the clustering data, some subtle differences can be revealed. We observe that for all fixed packing fractions $\eta$, the CSD seems to evolve towards a steady state (figure 3). However, the temporal convergence is slower than the one observed for A$^+$S$^-$Frz$^-$ cells. Assuming that CSD after 450 min from the beginning of the experiment is representative of the steady-state CSD, we show in figure 4 the asymptotic behaviour of the CSD with packing fraction $\eta$. The CSDs of the packing fractions $\eta < 0.18$ can be roughly approximated by a power law, $p(m) \propto m^{-\gamma_0}$, with a critical exponent $\gamma_0$ consistent with the one obtained for A$^+$S$^-$Frz$^-$ cells, i.e. $0.81 \leq \gamma_0 \leq 0.95$ (figure 4). On the other hand, the data indicate that a local maximum, as the one described above for A$^+$S$^-$Frz$^-$, emerges for $\eta \approx 0.18$ (figure 4). On the other hand, the cluster shape statistics shows that the scaling of the perimeter $P$ with the cluster mass $m$ is again consistent with the one obtained for A$^+$S$^-$Frz$^-$ cells with $\omega = 0.62 \pm 0.03$ (figure 5). Finally, at sufficiently high densities, these cells also self-organize into vortices.

### 2.3. Wild-type and the effect of cell reversal

We applied the same analysis to the reversing A$^+$S$^+$Frz$^+$ cells that move by means of both motility systems. Figure 1 shows that the spatial organization of wild-type cells is dramatically different from the one observed in the two mutants. Undoubtedly, cell

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**Figure 5. (a–c) Cluster perimeter $P(m)$ as a function of the cluster size $m$ for three bacterial strains. The A$^+$S$^-$Frz$^-$ and A$^+$S$^+$Frz$^-$ mutants exhibit roughly the same scaling $P(m) \propto m^\omega$, with $\omega \approx 0.60$ for the A$^+$S$^-$Frz$^-$, $\omega \approx 0.62$ for the A$^+$S$^+$Frz$^-$, suggesting that an increase in adhesion does not have a strong impact on the cluster shape. On the other hand, cell reversals lead to much more elongated clusters, as the scaling of the A$^+$S$^+$Frz$^-$ cells indicates, with $\omega \approx 0.82$.**

**Figure 6. Vortex formation (circular pattern in the centre) in the non-reversing mutant A$^+$S$^-$Frz$^-$ at high cell density ($\eta > 0.26$). The pattern consists of rotating stacked discs of cells. These structure are observed in both A$^+$S$^-$Frz$^-$ and A$^+$S$^+$Frz$^-$ strains.**
reversal has a strong impact on the macroscopic behaviour of the colony. The CSD after 450 min is exponential for all \( \eta < 0.20 \), as shown in figure 4. The net distance of cell movement is reduced owing to cell reversals, and cells can only form small clusters. On the other hand, clusters exhibit a more elongated shape than those found in experiments with A"S-Frz" and A"S-Frz" cells, as confirmed by the scaling of the perimeter \( \Pi(m) \) which is characterized by a very different exponent \( \omega = 0.82 \pm 0.03 \) (figure 5). The initial monodisperse phase, characterized by an exponential CSD and very elongated clusters, undergoes a transition at packing fractions larger than 0.26. The new arrangement of cells percolates, and the cells organize into a mesh-like structure, as shown in figure 1c.

3. KINETIC MODEL FOR THE CLUSTER STATISTICS

In the following, we outline a generalized kinetic model for the CSD and compare it with the above experimental results. In particular, we want to relate the CSD data and the cluster shape statistics. The model equations are built on the well-established coagulation theory for colloidal particles originally suggested by Smoluchowski [27]—for an early review, see also [28]. A similar phase transition (albeit with different exponents for the CSD at criticality) was recently obtained in a model for reversible polymerization representing a different generalization of the Smoluchowski model [29].

The model studied consists of a system of kinetic equations for the dynamics of the number \( n_j(t) \) of clusters of size \( j \) at time \( t \). It was first proposed in Peruani et al. [19] to describe clustering in simulations of self-propelled rods. The individual cluster dynamics [30], as well as the cluster–cluster interactions [19,30], are strongly simplified in this kinetic theory where the time evolution of the number \( n_j(t) \) of clusters of size \( j \) is simply given by

\[
\begin{align*}
\dot{n}_1 &= 2B_1n_2 + \sum_{k=3}^{N} B_{k} n_k - \sum_{k=1}^{N-1} A_{k,1} n_k n_1, \\
\dot{n}_j &= B_{j+1}n_{j+1} - B_{j} n_j - \sum_{k=1}^{N-j} A_{k,j} n_k n_j \\
&\quad + \frac{1}{2} \sum_{k=1}^{j-1} A_{k,j-k} n_k n_{j-k} \quad \text{for } j = 2, \ldots, N-1 \\
\dot{n}_N &= -B_N n_N + \frac{1}{2} \sum_{k=1}^{N-1} A_{k,N-k} n_k n_{N-k},
\end{align*}
\]

(3.1)

where the dot denotes a time derivative, and \( N \) is the total number of cells in the system. The CSD is then simply obtained from

\[
p(m, t) = \frac{m n_m(t)}{N}.
\]

(3.2)

We have assumed that aggregation of cells occurs only as a result of cluster–cluster collisions. Following earlier work [19,30], the collision rate between clusters of mass \( j \) and \( k \) is defined by

\[
A_{j,k} = \frac{\nu_0 \sigma_0}{\delta} (\sqrt{j} + \sqrt{k}),
\]

(3.3)

where \( \nu_0 \) represents the average speed of individual cells, \( \sigma_0 \) is the average scattering cross section of a single cell, which is assumed to be \( \sigma_0 \approx L + W = \sqrt{a(\sqrt{k} + 1/\sqrt{k})} \), and \( \delta \) is the total area of the system. Equation (3.3) assumes that clusters have a well-defined direction of motion, which means that the equation is not adequate to describe cluster–cluster coagulation in experiments with wild-type cells. This process competes with cluster fragmentation stemming from the escape of individual single cells from the cluster boundary. The fragmentation rate is given by the expression

\[
B_j = \frac{\nu_j^w R_0}{R_j \delta k},
\]

(3.4)

where \( R_0 \) is a proportionality constant that is the only free parameter in the theory that is used to fix the critical value \( \eta \approx R_0^{-1} \) at the same values as in the experiment. The exponent \( \omega \) in the fragmentation rate has an important role: it represents the scaling between the cluster mass \( m \) and the cluster perimeter \( \Pi \), i.e. \( \Pi \approx m^\omega \). If one assumes large clusters of approximately circular shape, then \( \omega = \frac{1}{2} \); this special case has been previously studied in Peruani et al. [19]. Instead one considers that cells form elongated narrow clusters, where practically all cells are near the boundary, then a choice of \( \omega = 1 \) is appropriate. In practice, the value of \( \omega \) will depend on the number \( j \) of particles in the respective cluster. For simplicity, we study only the limiting cases \( \omega = \frac{1}{2} \) and \( \omega = 1 \) and compare the resulting CSD to the experimental findings. According to the model, the exponent \( \gamma_0 \) only depends on the scaling of \( \Pi(m) \), i.e. the exponent \( \omega \), while the critical packing fraction \( \eta_c \) is a non-universal quantity. The analysis of equations (3.1), performed by direct numerical integration using a fourth-order Runge–Kutta method, reveals that for \( \eta \leq \eta_c \), the scaling of \( p(m) \) takes the form

\[
p(m) \propto m^{-\gamma_0} \exp \left( -\frac{m}{m_0} \right),
\]

(3.5)

while above it, i.e. for \( \eta > \eta_c \), the scaling is

\[
p(m) \propto m^{-\gamma_0} \exp \left( -\frac{m}{m_1} \right) + Cm^{\gamma_1} \exp \left( -\frac{m}{m_2} \right),
\]

(3.6)

with \( \gamma_1 \), \( \gamma_2 \), \( m_1 \), \( m_2 \) and \( C \) being constants that depend on \( \eta \) and system size. Equation (3.5) is the result of a system size study of equation (3.1) at the critical point (not shown), while equation (3.6) is just an educated guess. Equations (3.5) and (3.6) have been used to fit the experimental data for the CSDs in the different strains of myxobacteria shown in figure 4. For \( \eta < \eta_c \), for either A"S-Frz" or A"S-Frz" cells, we find using equation (3.5) that \( \gamma_0 \in [0.80, 0.95] \) and \( m_0 \in [20,1300] \) (\( m_0 \approx 20 \) for \( \eta = 0.04 \) and \( m_0 \approx 1300 \) for \( \eta = 0.16 \)). Nevertheless, the critical exponent \( \gamma_0 \) has been estimated by the method explained in §5, where \( \gamma_0 \) is found to be \( \gamma_0 = 0.88 \pm 0.07 \). For wild-type cells, the distribution is strongly dominated by an exponential tail. Using equation (3.5), we find \( \gamma_0 \in [0, 0.63] \) and \( m_0 \in [10, 120] \).
Through equation (3.1), it can be shown that \( n_0 \) is a function of \( \eta \) that increases as \( \eta \) is approached from below as observed in figure 4. According to the kinetic model, the critical packing fraction \( \eta_c \) is defined by \( p(m) \propto m^{-\eta_c} \) at \( \eta = \eta_c \) as long as \( m \) is below the total number of cells \( N \) in the system. In contrast, for \( \eta < \eta_c \), the function \( p(m) \) clearly exhibits an exponentially decaying tail at larger \( m \), as observed in the experiments with A^S Frz^+ and A^S Frz^- cells (figure 7). The theoretical CSD \( p(m) \) was obtained by numerical integration from an initial condition with \( n_i = N \) and \( n_i = 0 \) for \( i \geq 2 \). The values of the variables \( n_i \) of equation (3.1) reached constant steady values after sufficiently large integration times. The steady-state \( p(m) \) was found to depend only on the packing fraction \( \eta \) for a given perimeter scaling characterized by \( \omega \). For both values of \( \omega \) studied, we find a transition from an exponentially decaying CSD, described by equation (3.5) for low densities, to a non-monotonic CSD, described by equation (3.6), consisting of a power-law behaviour for small cluster sizes and a peak, local maximum, at large cluster sizes (figure 7). Upon closer inspection of the model results, one recovers distinctly different exponents for the different model assumptions regarding \( \omega \). \( \gamma_0 = 1.3 \) for \( \omega = \frac{1}{2} \) and \( \gamma_0 = 0.85 \) for \( \omega = 1 \). Both choices of \( \omega \) give reasonable qualitative agreement with the experimental data shown in figure 4 (figure 7). Moreover, we find that the exponent of the CSD is non-universal and depends sensitively on the choice of the fragmentation rate in equation (3.4). We expect that changes in the collision rate for the cluster will have a similarly strong effect, as discussed below.

The clustering model given by equation (3.1) allows us to study the relationship between the perimeter scaling (characterized by an exponent \( \omega \)) and the CSD exponent \( \gamma_0 \). Equation (3.1) also predicts the existence of two CSDs, depending on the packing fraction \( \eta \), i.e. equations (3.5) and (3.6). These two predicted distributions are found in experiments with A^S Frz^- and A^S Frz+ cells. For the wild-type cells, only the CSD given by equation (3.5) is found. In this context, it is interesting to note that the results shown in figure 7 imply that for \( \omega = 1 \), one needs to assume a much lower fragmentation rate—indicated by a much larger value of the parameter \( R_0 \)—than for \( \omega = 0.5 \) to obtain the same critical \( \eta \). Beyond the apparent agreement between the CSD exhibited by equation (3.1) and the experimentally obtained CSDs for A^S Frz^- and A^S Frz+ cells, there are important differences. To obtain a critical exponent \( \gamma_0 \) close to 0.88, \( \omega \) has to be large—specifically, close to 1, while the experimental measurements on \( \langle R^2 \rangle \) revealed \( \omega \sim 0.6 \). There are several possibilities that could explain this discrepancy. For instance, the assumption that the cluster–cluster coagulation is proportional to the square root of the cluster mass has to be revised. An estimation of the scaling of the effective scattering cross section of a cluster with its mass, as well as an accurate measurement of the functional dependency of cluster speed with cluster mass would allow us to determine the correctness of equation (3.1). Unfortunately, such measurements are extremely difficult to obtain. Nevertheless, the apparent discrepancy suggests that a possible generalization of the presented clustering theory would include a modification of equation (3.3).

4. DISCUSSION

In order to identify the role of the two motility machineries as well as cell reversal machinery in the special collective dynamics of M. xanthus, we have analysed three bacterial strains: (i) a mutant that moves unidirectionally without reversing by the A-motility system only, (ii) a unidirectional mutant that is also equipped with the S-motility system, and (iii) the wild-type that is equipped with the two motility systems and occasionally reverses its direction of movement. The study of the two non-reversing mutants revealed the same phenomenology. At low and intermediate densities, non-reversing cells display collective motion in the form of large moving clusters, with a critical density above which clusters can be arbitrarily large. At the critical density, the two non-reversing strains exhibit a CSD characterized by roughly the same critical

\[
\eta_0 = 0.17, \quad \eta_0 = 0.27
\]

Figure 7. Theoretical predictions for the CSD. The CSD from the kinetic model depends on the scaling of the fragmentation \( B_\omega \propto m^{-\omega} \) (equation (3.4)). The figure shows the results for the two limiting cases: (a) \( \omega = 0.5 \) (and \( R_0 = 0.58 \)) corresponding to round clusters; (b) \( \omega = 1 \) (and \( R_0 = 11.9 \)), implying elongated clusters. Notice that in theory, as well as in experiments, the CSD can be well approximated by a power law at the critical packing fraction \( \eta_c \), while for \( \eta > \eta_c \), a peak at large cluster sizes emerges, a signature of dynamic self-assembly into larger moving clusters. Other model parameters: \( k = 9 \), \( a = 4.4 \mu m^2 \) and \( \delta = 699 \times 522 \mu m^2 \) (see equations (3.3) and (3.4)).
exponent $g_0 \sim 0.88$. Even though the two-engine strain is supposed to be more adhesive than the single A-engine strain, we found a similar non-trivial scaling of cluster perimeter with cluster size characterized by an exponent $\omega \sim 0.6$. This finding indirectly shows that the clustering process is, for both strains, neither fully random nor an equilibrium one as in liquid–vapour drops [18]. In order to connect the statistics on cluster size and cluster shape, we derived a Smoluchowski coagulation theory with fragmentation, where we related the scaling of cluster perimeter with cluster size with the fragmentation kernel. The proposed theory allows us to understand the cluster formation process as a dynamic self-assembly process in the absence of adhesion. It predicts the existence of a steady-state CSD which is a function of the cell density and perimeter exponent $\omega$, and a functional change of the CSD above a critical density. In addition, the proposed theory predicts that the critical exponent $g_0$ depends on the perimeter exponent $\omega$ only. In summary, the theoretical clustering model provides a qualitative description consistent with the experimental measurements, and explains why if the value of $\omega$ is similar for both strains, the value of $g_0$ has to be also similar. We observe that similar spatial organization has been observed in self-propelled rod simulations using either rigid [19] or elastic [31] elongated particles. We found that at high densities the collective dynamics changes and cells organize into vortices. This finding cannot be accounted for by the proposed clustering theory, but it is reminiscent of what is observed in self-propelled rod simulations at high densities, though in experiments vortices seem to be stable structures while in simulations, vortices are unstable. By comparing these two non-reversing strains, we conclude that unidirectional cell motion induces the formation of large moving clusters at low and intermediate densities, while it results in vortex formation at very high densities. In the light of the clustering theory and given the remarkable similarity with self-propelled rod simulations, we suggest that the spatial self-organization in these two strains occurs in the absence of biochemical signal regulations and as a result of the combined effect of self-propulsion and volume exclusion interactions. All these results strongly suggest that the combination of self-propulsion and steric interaction is a valid pattern-formation mechanism that could be also at play in recent experiments with *Escherichia coli* [32] and driven actin filaments [22], which makes us wonder about the connection between this mechanism and the large body of work on simple models of self-propelled particles where spontaneous segregation and long-range orientational has been reported [33–41].

The study of wild-type cells has revealed that cell reversal affects dramatically the collective dynamics. We found that wild-type cells exhibit CSDs exponentially distributed at low and intermediate densities. On the other hand, we measured the scaling of the cluster perimeter with cluster size characterized by a large exponent $\omega \sim 0.8$, which indicates that clusters are strongly elongated in comparison with those found in experiments with the two non-reversing mutants. Finally, we observed that at high densities, cells self-organize into a mesh-like structure. A qualitative understanding of this macroscopic behaviour is still missing. A comparison of the two non-reversing strains and wild-type cells suggests that only by switching on and off the reversal can cells modify dramatically their collective behaviour, with the suppression of reversal leading to collective motion in the form of moving clusters and vortex formation at high densities. We note that this observation is consistent with the observed decrease in reversal frequency in the wild-type upon nutrient depletion, which is followed by the formation of large moving clusters and aggregation of cells.

At a more speculative level, our results suggest that the cell density and the rod shape of the cells may play an essential role for bacteria to achieve collective motion [42,43]. According to self-propelled rod simulations, an elongated cell shape strongly facilitates collective motion by promoting the formation of larger clusters. Another hint that the rod shape of the moving bacteria is important for collective motion is provided by the empirical observation that many bacteria undergo a dramatic elongation of their cell shape before assembling into larger groups, e.g. in *Vibrio parahaemolyticus* [44] or *B. subtilis* [45]. Finally, the reported results increase the plausibility of earlier biological hypotheses [42], that multicellular organization may be achieved by regulating the cell density via proliferation and cell length by direct developmental control.

5. MATERIAL AND METHODS

5.1. Bacterial strains

The fully motile strain DK1622 (A+S+Frz+) was used as a wild-type [46] and all other strains used are derivatives of DK1622. The non-reversing strain DK8505 [47] is referred to as A+S+Frz−. To generate SA2407 cells, here referred to as A+S+Frz−, the frz loss-of-function allele frzCD::Tn5 lac Ω536 from DK8505 [47] was introduced into the ΔpilA strain DK10410 [48], which is unable to assemble type IV pilus, using standard procedures [49]. To generate SA2082 (ΔpilA, romR::nptII), the non-motile *M. xanthus* mutant referred to as A+S+Frz−, the romR::nptII loss-of-function allele from SA1128 [50] was introduced into DK10410. All strains used had a doubling time of approximately 5 h in CTT liquid medium at 32°C. Notice that the relaxation time of spatial patterns is below 120 min, which implies that the doubling time has a weak effect on the spatial patterns.

5.2. Cluster formation experiments

Cultures of *M. xanthus* were grown in CTT liquid medium [51] at 32°C with shaking to an estimated density of $7 \times 10^9$ cells ml$^{-1}$. Subsequently, cells were diluted to densities of $0.5 \times 10^8$ ml$^{-1}$, $1.0 \times 10^8$ ml$^{-1}$, $1.5 \times 10^8$ ml$^{-1}$, $1.75 \times 10^8$ ml$^{-1}$ and $2 \times 10^8$ ml$^{-1}$, respectively. Cell densities were confirmed by counting the colonies manually on CTT agar plates and by counting the number of cells using a counting chamber. Thirty microlitres of aliquots of cells were transferred to a microscope slide covered with a 1.0 per cent agar pad.

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in 0.5 per cent CTT medium. The time point at which the cell drop was completely absorbed in the agar was set as \( t = 0 \). For each cell density, 16 slides were prepared and every 30 min (starting at 30 min) up to 480 min, a sample was analysed by microscopy using a Leica DFC 350FX camera. Twenty phase-contrast images were taken at 20× magnification across a spot. After 480 min, a short time-lapse movie was taken to verify that cells and clusters were migrating.

### 5.3. Image analysis

Clustering images were taken at 20× magnification. Images contain cell clusters as dark regions, often surrounded by a light halo. Cluster boundaries were detected in a multi-step processing queue. After initial image normalization, edge detection via the Canny–Deriche algorithm was applied for two different levels of spatial detail. Both edge images were superimposed subsequently. Next, edges were filtered out that surround halos and other non-cluster objects. Finally, all incomplete detections were revised manually in a post-processing step. The areas of the clusters in pixels were extracted using an implementation of the processing queue in the image processing tool IMAGEJ (http://rsbweb.nih.gov/ij/). The number of cells inside a cluster, i.e. the cluster size, was estimated as the area of a cluster divided by the mean area covered by a single cell, which was found to be 150 pixels at 20-fold magnification. According to this definition, a cluster is a connected group of cells, regardless of their orientation. Packing fraction estimates per image were obtained as the ratio of area covered by cells and the whole area of the image (1392 × 1040 pixels corresponding to 699 × 522 μm).

### 5.4. Statistical analysis

After applying the image analysis procedure described above to a given image \( I \), corresponding to a given packing fraction, a large array of various cluster sizes is obtained, and \( n_I(m, t) \) can be computed. We represent by \( n_I(m, t) \) the number of clusters of size \( m \) in the image \( I \). To build the CSD, we make use of all the available images corresponding to the given packing fraction \( \eta \). Let the auxiliary function \( g(m, t) \) be \( g(m, t) = m n_I(m, t) \). The average of this function reads

\[
g(m, t) = \frac{1}{M} \sum_i g_I(m, t),
\]

where \( M \) is the number of available images. To cope with the sparseness of the data for large cluster sizes, we implemented several binning procedures, in particular, linear and exponential binning. In the following, we explain the exponential binning procedure. The cluster size range is divided into bins, the first bin contains all clusters of size \( s \), \( 0 < s \leq 1 \), the second bin all clusters of size \( 1 < s \leq 2 \), the third bin, \( 2 < s \leq 4 \), etc. The \( n \)th bin contains cluster sizes \( 2^{n-1} < s \leq 2^n \). It is useful to define the function

\[
g_{\text{bin}}(n, t) = \sum_{i(n)} \sum_l g_I(m, t),
\]

where \( c(n) = 2^n \). The binned CSD is defined as

\[
p_{\text{bin}}(c(n), t) = \frac{g_{\text{bin}}(n, t)}{C(c(n) - c(n - 1))}.
\]

Thus, \( \sum_m p_{\text{bin}}(m = c(n), t)(c(n) - c(n - 1)) = 1 \). It is worth noticing that if the underlying CSD \( p(m) \) is a power law characterized by an exponent \( \gamma \), i.e. \( p(m) \sim m^{-\gamma} \), the exponential binning procedure given by equation (5.3) results in \( p_{\text{bin}}(m) \sim m^{-\gamma} \). On the other

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hand, if the underlying CSD $p(m)$ is an exponential, i.e. $p(m) \sim \exp(m/m_0)$, the exponential binning leads to $p_{\text{bin}}(m, t) \sim m^{-g} \exp(m/m_0)$. In the text, for simplicity we referred to $p_{\text{bin}}(m, t)$ just as $p(m, t)$.

In what follows, we explain how the critical exponent has been measured. At the critical packing fraction $\eta_c$, the CSD is a power law (with an exponential cut-off due to the finite number of cells). The problem consists of identifying the critical packing fraction $\eta_c$ and the critical exponent $y_0$. Assuming that we know $y_0$ at $\eta_c$, the following transformation yields a constant:

$$y(m) = p(m; \eta_c) m^{y_0} = W,$$

where $W$ is a constant and the equality holds true for $1 < x < x_{\text{cut-off}}$, where $x_{\text{cut-off}}$ denotes the beginning of the cut-off. The value of $W$ is the average value of $y(m)$ in the interval $1 < x < x_{\text{cut-off}}$. This means that if we plot $y(m)$ versus $m$, we observe a horizontal line at the critical packing fraction $\eta_c$. We can measure how close we are to the horizontal line by computing

$$
\sigma^2(\eta, \gamma) = \sum (y(m) - W)^2 .
$$

By minimizing equation (5.5) with respect to $\eta$ and $\gamma$, the critical packing fraction and critical exponent can be obtained. Figure 8 illustrates the procedure. In the critical packing fraction lies between 0.16 and 0.18 for either $A^+$ or $A^-$.

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