Production and Characterization of Motile and Chemotactic Bacterial Minicells

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ABSTRACT: Minicells are nanosized membrane vesicles produced by bacteria. Minicells are chromosome-free but contain cellular biosynthetic and metabolic machinery, and they are robust due to the protection provided by the bacterial cell envelope, which makes them potentially highly attractive in biomedical applications. However, the applicability of minicells and other nanoparticle-based delivery systems is limited by their inefficient accumulation at the target. Here we engineered the minicell-producing *Escherichia coli* strain to overexpress flagellar genes, which enables the generation of motile minicells. We subsequently performed an experimental and theoretical analysis of the minicell motility and their responses to gradients of chemoeffectors. Despite important differences between the motility of minicells and normal bacterial cells, minicells were able to bias their movement in chemical gradients and to accumulate toward the sources of chemotactants. Such motile and chemotactic minicells may thus be applicable for an active effector delivery and specific targeting of tissues and cells according to their metabolic profiles.

KEYWORDS: chemotaxis, motility, minicell, nanoparticle, drug delivery

INTRODUCTION

Nanoparticles are highly promising as containers for targeted drug delivery in such biomedical applications as tumor therapy, and a large spectrum of different nanoparticle designs has been developed over the recent years. Nevertheless, the efficiency of drug delivery by nanoparticles remained relatively low, with a particular challenge being to enrich nanoparticles within the targeted tissues. One type of nanosized delivery vehicles is bacterial minicells, ~0.5 μm spheres surrounded by a cell envelope, which are spontaneously generated through an aberrant division of bacteria close to cell poles. The production of minicells is particularly frequent in bacterial *min* mutants that have lost control of the cell-division site placement.

Minicells carry no chromosomes and are therefore nonliving, but they can contain plasmid DNA and other cellular components, including metabolic enzymes and cellular machineries that are required for energy generation and for transcription and translation. Hence, minicells are metabolically and biosynthetically active, meaning that—similar to the intact bacteria—they can be utilized as specific biosensors and engineered to express a wide range of toxins, cytokines, tumor antigens, and apoptosis-inducing factors under the control of specific external stimuli. Because of the protection provided by the bacterial cell wall and membranes, minicells are highly robust and do not spontaneously release their content. Their small size enables minicells to penetrate fenestrated blood vessels and to accumulate at tumor sites, where they can be subsequently endocytosed and release their content within the target cells.

Minicells were engineered to target cancer cells via bispecific antibodies and equipped with various payloads including chemotherapeutic drugs or inhibitory RNAs as well as with a secretion system for antigen injection into the host cells.

Intact bacteria can also be utilized as drug delivery vehicles, with the bacterial ability to swim in liquid media providing a particular advantage for efficient delivery. The bacterial swimming motion is typically mediated by the rotation of several flagellar filaments that bundle together to propel the cell, and it can be biased in chemical gradients by the chemotaxis signaling pathway. This pathway perceives temporal changes in chemical stimulation as cells swim in the gradient, and it signals to flagellar motors to modulate the frequency of cell reorientation, thereby increasing the duration of cell runs in a favorable direction. This mechanism of gradient sensing by temporal comparisons of ligand concentration along the swimming path is necessary because of the small size of bacteria, and it is physically limited by a gradual reorientation of the cell body due to Brownian diffusion or—at high cell densities—to the emergent collective motion. Because of this importance of rotational diffusion, the...
processivity of swimming and thus efficiency of chemotaxis can increase with bacterial cell length.\textsuperscript{22,23}

Bacterial chemosensory systems were reported to perceive chemical signals released by the host epithelium\textsuperscript{24−26} and in tumor microenvironments,\textsuperscript{27,28} indicating that chemotaxis could be applicable for specific tumor targeting. Furthermore, flagellar motility can promote an attachment to epithelial cells\textsuperscript{29} and tissue penetration.\textsuperscript{30,31} Motile bacteria can also be specifically loaded with cargo nanoparticles carrying customized therapeutics,\textsuperscript{32} and such bacteriabots are chemotactic as long as the cargo does not strongly reduce their swimming speed.\textsuperscript{33} However, despite these potential advantages for autonomous active delivery, the \textit{in vivo} application of intact bacteria remains severely limited by biosafety concerns.

Here we report a system that combines advantages provided by small and chromosome-less minicells with the chemotactic capability of motile bacteria. We engineered an \textit{Escherichia coli} strain that produces minicells with an inducible expression of the flagellar system and investigated their motility and chemotaxis. We demonstrate that, at higher levels of flagellar gene expression, these minicells are well-motile, despite their small size and hence small number of flagella and faster rotational diffusion. Moreover, although minicells were previously shown to contain functional chemosensory complexes,\textsuperscript{33,34} it was unclear whether their swimming could be fast and processive enough to enable a proper functioning of the bacterial chemotaxis strategy. We show that, despite these potential limitations, the chemotactic efficiency of minicells is comparable to that of regular bacteria, and we develop an analytical model of the minicell motility that can largely account for our experimental observations. This proof-of-concept implementation of motility and chemotaxis in minicells makes it possible to further increase the efficiency of minicell-based drug delivery as well as its specific targeting relying on chemical gradients emanating from particular microenvironments such as tumors.

\section*{RESULTS AND DISCUSSION}

\textbf{Engineering \textit{E. coli} for an Inducible Production of Flagellated Minicells.} To generate minicells, we used a derivative of the \textit{E. coli} strain MG1655 that carries a deletion of the \textit{minCDE} operon encoding the division-site positioning system. This deletion results in frequent cell divisions at cell poles, pinching off multiple minicells.\textsuperscript{3} We further introduced an A115V amino acid replacement in the actin-like protein \textit{mreB}, which decreases the \textit{E. coli} width and therefore leads to the production of minicells with even smaller diameter.\textsuperscript{34} Whereas cells of the wild-type strain MG1655 are \(\sim 2.3\, \mu\text{m}\) long and \(\sim 1\, \mu\text{m}\) wide when grown in tryptone broth (TB) (Figure 1A–E), the \textit{minCDE mreB\textsuperscript{A115V}} strain also acquired a spontaneous (apparently adaptative) deletion that inactivated the operon encoding \textit{flhDC}, the upstream master regulator of the flagellar regulatory network (Figure 2A), thus effectively shutting down the expression of all flagellar genes. Consequently, both minicell-producing mother cells (Figure 1A) and minicells (Figure 1B) were not flagellated. In order to tune the levels of flagellar and chemotaxis proteins, we engineered this strain to express the \textit{flhDC} operon from a plasmid under an arabinose-inducible promoter. The induction of \textit{flhDC} expression in minicell-producing mother cells indeed led to the increased activity of the flagellin (\textit{flIC}) promoter (Figure 2B,C) that is representative for the expression of flagellar and chemotaxis genes (Figure 2A).\textsuperscript{33,36} Consistently, the activation of flagellar gene expression led to the appearance of flagellar filaments in both mother cells (Figure 1C and Figure 3A,B) and in minicells (Figure 1D,F,G and Figure 3C,D). On average, flagellar filaments in minicells were \(\sim 7.5\, \mu\text{m}\) in length (Figure 1G and Figure 4A), similar to the length of flagellar filaments in the parental MG1655 cells (Figure 1C). The filament length remained constant over the whole range of the \textit{flhDC} expression levels, above the initial activation threshold (Figure 1G and Figure 4A), whereas the number of flagellar motors increased with induction up to a maximum of approximately two motors per minicell (Figure 1F and Figure 4B).

\textbf{Motility and Chemotaxis of Flagellated Minicells.} To investigate whether flagellated minicells generate enough energy to power the rotation of flagellar motors, and whether a rotation of one to two flagellar filaments produces a sufficient force to processively propel the minicell, we next compared the motion of nonflagellated and flagellated minicells. Consistent with them lacking an active propulsion system, trajectories of
nonflagellated minicells were clearly Brownian (Figure 2A). In contrast, trajectories of flagellated minicells showed significantly persistent swimming (Figure 2B and Figure S5), comparably to the trajectories of the parental MG1655 cells (Figure S5D). Cell tracking confirmed that, as expected for diffusive behavior, the mean squared displacement (MSD) increases linearly as a function of time for nonflagellated minicells (Figure 2D) and that the distribution of their displacements is Gaussian (Figure S5A). For flagellated minicells and MG1655 cells, the MSD grows quadratically with time (Figure 2D), which is characteristic for a ballistic motion. Distributions of displacements were also similar for flagellated minicells and MG1655 cells (Figure S5A), with minor differences being likely explained by a slightly higher fraction of minicells that were nonmotile, ∼10–30% (independent of flhDC induction) compared to 5–10% for MG1655, as determined by microscopy analysis (see Materials and Methods). This subpopulation of nonmotile minicells might arise from cell or flagella damage during the culture preparation or from the residual heterogeneity of the flhDC induction (Figure S2C). The average velocity of swimming minicells saturated below 15 μm/s (Figure 2C), which is significantly lower than the velocity of MG1655 cells (24 μm/s). In contrast, the duration of runs was longer for the swimming minicells (Figure 2E). Finally, consistent with their small size, swimming minicells were reoriented more rapidly due to the rotational diffusion, being thus less able to maintain their swimming direction than normal E. coli cells (Figure 2F).

To further test whether, despite these differences in their swimming behavior, motile minicells are capable of performing chemotaxis, we probed the motility of minicells in gradients of α-methyl-D,L-aspartate (MeAsp), a nonmetabolizable analogue of aspartate and potent chemoattractant for E. coli. We first used a previously described microfluidic device (Figure 3A inset) that allows measurements of the chemotactic drift of a bacterial population in a steady linear chemical gradient.35,37 A significant population drift up the MeAsp gradient could be observed, with the drift velocity of the minicell population growing with flhDC gene expression (Figure 3A). This drift velocity apparently increased linearly as a function of the average number of flagellar motors of the minicells (Figure 3B). Thus, although a single flagellum might already be...
sufficient to propel a minicell at nearly maximum speed, the efficiency of the chemotaxis is apparently higher for minicells that are propelled by two flagella. This higher efficiency might stem from physical effects of the number of flagella on the tumbling rate and processivity of swimming and/or from an increased expression of chemotaxis proteins at higher levels of flhDC induction (Figure S2A and Supporting Information). Notably, even at the highest induction of flagellar genes the chemotactic drift of the minicells ($v_{\Delta m} = 0.30 \pm 0.07 \mu m/s$) remained lower than the one of the parent MG1655 cells ($v_{\Delta m} = 2.1 \pm 0.1 \mu m/s$).

We further confirmed that chemotactic minicells can efficiently accumulate toward the sources of chemoeffectors, by using another microfluidic device where MeAsp is continuously released at the end of the channel (Figure 3C,D and Figure S6). This device mimics natural situations where chemoeffectors are released by a source, such as tumor tissue. Consistent with their ability to perform chemotaxis, minicells showed an increased migration into the observation channel, thus accumulating toward the source of attractant (Figure 3E,F). No accumulation was observed in the absence of MeAsp in the microfluidic chamber (Figure 3E,G) or for nonmotile minicells (Figure 3E,H).

**Modeling of Motility and Chemotaxis of Minicells.** In order to better understand physical limitations on the swimming and chemotaxis of minicells, we used a common model for the chemotactic drift of *E. coli* (Supporting Information). The chemotactic drift is described in this model as a function of biochemical properties of the signaling pathway as well as of the physical parameters that characterize cell swimming. We assumed that the functioning of the chemotaxis pathway in minicells is similar to that of the normal cells. Two important physical parameters, which are affected by the cell dimensions, are the rotational diffusion coefficient for swimming cells $D_r$ and the tumble persistence time $\tau_T$ (Figure 4). According to our data (Figure 2F), minicells are less able to keep swimming in a given direction, with the rotational diffusion coefficient of the minicells being $D_{ri} = 1.3 \pm 0.1 s^{-1}$ and therefore much larger compared with $D_{ri} = 0.1 s^{-1}$ for normal cells. This increase is well-accounted for by a simple model of rotational diffusion that considers the rotation of the cell body and the flagellum (Supporting Information).
Information). The tumble persistence time measures the time it takes for a cell to randomize its direction of motion via tumbling. It is expected to be \( \tau_T = \tau_0/(1 - \langle \cos(\Delta \theta_T) \rangle) \), where \( \tau_0 \) is the mean run duration, and \( \Delta \theta_T \) is the angular change in direction during a tumble. For minicells, the angular change \( \Delta \theta_T \) is expected to be larger compared to normal cells because of their smaller body, whereas the run duration \( \tau_0 \) is expected to increase because of their smaller number of flagella. Indeed, the run duration of minicells was \( \tau_0^\text{min} = 1.9 \pm 0.1 \text{ s} \), compared with \( \tau_0^\text{norm} \approx 1.6 \pm 0.1 \text{ s} \) for normal cells (Figure 2E). This moderate increase could be well-accounted for by an effective veto model for bacterial tumbling (Supporting Information). We also assumed a complete randomization of the swimming direction of minicells during a tumble (\( \langle \cos(\Delta \theta_T) \rangle = 0 \)), in contrast to only a partial reorientation for longer normal cells (\( \langle \cos(\Delta \theta_T) \rangle \approx 1/2 \)) as reported previously. Assuming the biochemical properties of the chemotaxis pathway are unchanged in the minicells, we predict a chemotactic velocity for the minicells \( v_{ch} = 0.18 \pm 0.04 \mu \text{m/s} \). This estimate is in very good agreement with the experimentally observed value (Figure 3B), confirming that the difference in chemotactic ability between normal and minicells is primarily due to the difference in their physical properties, especially their increased rotational diffusion coefficient. The model also highlights that, besides cell propulsion, flagella play another essential role in ensuring that minicells are capable of chemotaxis, namely, by reducing rotational diffusion and therefore stabilizing the direction of the minicell motion. Nevertheless, the model seems to underestimate their chemotaxis efficiency, which might be either because of the oversimplified model assumptions or due to the slightly different signaling parameters of minicells, such as higher concentrations of chemotaxis proteins or faster signaling due to shorter distances between the chemosensory complexes and flagellar motors.

**Concluding Remarks.** Concluding, we observed that *E. coli* minicells that were engineered to have high levels of a flagellar gene expression are motile. The swimming pattern of these minicells was different from that of the parental *E. coli* cells, with minicells exhibiting a lower swimming velocity and directional persistence but increased run duration. These differences were consistent with the mathematical model describing minicell motility, and they could be accounted for by their small size and thus faster rotational diffusion as well as by the smaller number of flagella per minicell. Despite the potential major impact of these factors on the bacterial chemotaxis strategy, minicells were capable of following chemical gradients and accumulating toward sources of chemoattractants. Given increasing evidence that chemotactic bacteria can follow local chemical gradients to accommodate specific sites within their animal hosts, 24–26 motility and chemotaxis could therefore be used to largely enhance the efficiency and specificity of the delivery of various drugs and protein and nucleic acid effectors that can be carried by minicells.

**Analysis of Swimming Velocity and Chemotaxis.** The average swimming and chemotactic drift velocity of minicells were measured as described previously. 35,36,53 Swimming velocity and chemotactic drift velocity of purified minicells were measured by recording the cell motion in a poly-(dimethylsiloxane) (PDMS) microchannel using phase-contrast microscopy (Nikon TI Eclipse, 10x objective with numerical aperture (NA) = 0.3, CMOS camera EoSens 4CXP). The cell motion was analyzed as described previously both via Fourier-based algorithms, 34,35 for measuring the swimming velocity and chemotactic drift, and cell tracking 53 for measuring mean squared displacements, run durations, and swimming persistence. A suspension of 100 µM MeAsp in a tethyng buffer (6.15 mM K2HPO4, 3.85 mM KH2PO4, 100 µM ethylenediaminetetraacetic acid (EDTA), 1 µM L-methionine, 10 mM lactic acid, pH 7.0) was used to generate a chemical gradient in PDMS chambers. All data were analyzed using ImageJ (https://imagej.nih.gov/ij/) with custom-written plugins.

The chemotactic accumulation of minicells in response to releasing gradients of MeAsp was measured with a microfluidic device described previously, 38–40, with a slight modification. For the microfluidic devices preparation, 0.3% agarose was added to fill the whole microfluidic chamber. Ten microliters of tethering buffer each was then added into both source and sink sides. Afterward, purified minicells were added into the sink pore and allowed to diffuse into the observation channel for 2 h. A solution of 2 mM MeAsp was then added to the source pore and allowed to gradually diffuse through the agarose gel into the observation channel. The minicell density in the observation channel was monitored over time, starting immediately after compound addition, using Nikon Ti-E inverted fluorescence microscope with a 20x objective lens and Lumencor SOLA-SEII equipped with Andor Zyla SCiMOS camera.

**Analysis of Cell-Tracking Data.** The mean squared displacement \( \text{MSD}(t) = \langle (r_i(t) - r_i(t_0))^2 \rangle \) and \( r_i(t_0) = (x_i(t_0), y_i(t_0)) \) the two-dimensional (2D) position of particle \( i \) at time \( t_0 \) was computed as a function of the lag time \( t \), averaging over particles \( i \) and initial times \( t_0 \). The MSD was displayed until a lag time of \( t = 6 \text{ s} \), corresponding to one-tenth of the duration of the movies (61 s, 2500 frames) and above which statistics gets poor (less than 10 independent time steps per average). For the quantification of the tumbling rate and swimming persistence, trajectories were sorted into swimmer and nonswimmer based on their radius of gyration \( R_g = m r B^{115V} \) point mutation. 50 Green fluorescent protein (GFP) promoter reporter for flhC (pAM109) 35,36 was constructed based on pUA66, 51 pBAD18 or pBAD24 vectors carrying flhDC genes were used to express FlhDC. 52 sfGFP was expressed using a pTrc99a-backbone-based vector.

**Minicell Production and Purification.** For the minicell production, overnight cultures were inoculated into TB supplemented with kanamycin and grown at 30 °C with shaking (180 rpm) for 8 h. When necessary, different concentrations of arabinose (0%, 0.001%, 0.01%, 0.1%) were added in the culture to induce FlhDC expression, and 50 μM isopropyl β-D-1-thiogalactopyranoside (IPTG) was used to induce the sfGFP expression. For minicell purification, parental cells were first removed by the cell culture by centrifugation at 10 000g for 20 min, and minicells were subsequently harvested by centrifugation at 40 000g for 20 min.

**Materials and Methods**

**Strains and Plasmid Construction.** The *E. coli* strain MG1655 was used as the wild-type for all experiments. A *minCDE* deletion was conducted using λ red recombination relying on pKD46. 48 The kanamycin resistance cassette was removed using pCP20. 49 pKOV was used to generate

\[ \text{MSD}(t) = \frac{\langle (r_i(t) - r_i(t_0))^2 \rangle}{\langle (r_i(0))^2 \rangle} \]

\[ v_{ch} = \frac{\langle \Delta v \rangle}{\langle \Delta t \rangle} \]

\[ \text{Nicolle}^T \]
viaveraging over initial times and swimming cells, only. Quantification of tumbling rate was also performed as described previously. For measuring the swimming persistence, the instantaneous swimming direction was defined as the 2D instantaneous velocity measured on a 10 frames (0.25 s) wide sliding window normalized to its norm (\(u_0(t) = v(t)/v(t)\)). The time autocorrelation of \(u(t)\) was computed by averaging over initial times and swimming cells, \(\langle u(t)u(0)\rangle = \langle u(t + \tau_0)u(\tau_0)\rangle_{\tau_0}\).

Quantification of Flagellar Length and Number of Minicells. Purified minicells were suspended in a tethering buffer with 10% glycerol and frozen at −80 °C before analysis by electron microscopy. For the sample preparation, 5 μL of the minicell suspension was applied onto hydrophilized carbon-coated copper grids (400 mesh). After a brief wash with filtered water, bacteria were stained with 2% uranyl acetate. All samples were analyzed using a JEOL JEM-2100 transmission electron microscope with an acceleration voltage of 120 kV. For the image acquisition, an F214 FastScan CCD camera (TVIPS; Gauting) was used. The flagellar number and length of minicells were quantified manually with ImageJ.

Promoter Activity Analysis. The activity of the gfp reporter of the fliC promoter was assayed using a BD LSRFortessa SORP cell analyzer (BD Biosciences) as described previously.^

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acssynbio.1c00012.

Mathematical models for minicell motility, supporting figures (PDF)

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Author Contributions

(B.N. and R.C.) These authors contributed equally. B.N., R.C., and V.S. designed the research. B.N. performed the experiments. B.N. and R.C. analyzed the data. R.C. performed the mathematical modeling. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Funding

This work was supported by the Max Planck Society. R.C. acknowledges support from the Deutsche Forschungsgemeinschaft, Grant No. CO 1813/2–1.

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

We thank T. Heimerl, E. Kaganovitch, and N. Krink for their help with experiments.

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