Dynamic bimodality of curli expression in planktonic cultures of
*Escherichia coli* is stabilized by cyclic-di-GMP regulation

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Abstract

Curli amyloid fibers are major constituent of the extracellular biofilm matrix formed by bacteria of the Enterobacteriaceae family. Within *Escherichia coli* biofilms, curli gene expression is limited to a subpopulation of bacteria, leading to heterogeneity of extracellular matrix synthesis. Here we show that bimodal activation of curli expression also occurs in well-mixed planktonic cultures of *E. coli*, resulting in stochastic differentiation into distinct subpopulations of curli-positive and curli-negative cells at the entry into the stationary phase of growth. Monitoring curli activation in individual *E. coli* cells growing in a microfluidic device revealed that the curli-positive state is only metastable and it can spontaneously revert during continuous growth in a conditioned medium. The regulation by c-di-GMP is not required for curli gene activation or for differentiation of *E. coli* in subpopulations of curli-producing and curli-negative cells. Instead, we observe that c-di-GMP modulates the probability and dynamics of stochastic curli activation and enhances stability of the curli-positive state.

Keywords: Gene expression, bacteria, biofilm, bistability, differentiation, amyloid fibers
Introduction

Curli amyloid fibers are the key component of the extracellular matrix produced during biofilm formation by *Escherichia coli*, *Salmonella enterica*, and other Enterobacteriaceae [1-9]. In *E. coli* and *S. enterica* serovar Typhimurium, curli genes are organized in two divergently transcribed *csgBAC* and *csgDEFG* operons that share a common intergenic regulatory region [10]. Expression of these operons is under regulation of the stationary phase sigma factor $\sigma^S$ (RpoS) and thus becomes activated during the entry into the stationary phase of growth [4, 11-14]. This activation is achieved by the $\sigma^S$-dependent induction of the transcriptional regulator CsgD, which then controls the expression of the *csgBAC* operon that encodes the major curli subunit CsgA along with the curli nucleator CsgB and the chaperone CsgC [7, 8, 15]. In turn, csgD expression in *E. coli* and *S. Typhimurium* is either directly or indirectly regulated by multiple cellular factors that mediate responses to diverse environmental changes, including both global and specific transcriptional regulators, small regulatory RNAs and second messengers (reviewed in [16-19]).

One of the key regulators of csgD is the transcription factor MlrA [13, 14, 20, 21]. The activity of MlrA depends on cellular levels of bacterial second messenger bis-(3'-5')-cyclic dimeric guanosine monophosphate (c-di-GMP), and in *E. coli* this control is known to be mediated by a pair of the interacting diguanylate cyclase (DGC) and phosphodiesterase (PDE) enzymes, DgcM and PdeR, that form a ternary complex with MlrA [12, 14, 22]. MlrA is kept inactive by binding PdeR, and this interaction is relieved when the latter becomes active as a PDE thus acting as the trigger enzyme [22, 23]. This inhibition is counteracted by DgcM that locally produces c-di-GMP to engage PdeR, as well as by the global pool of c-di-GMP. Besides its enzymatic activity, DgcM might also activate MlrA through direct protein interaction. Another DGC-PDH pair, DgcE and PdeH, provides global regulatory input into the local DgcM-PdeR-MlrA regulation [12, 24].

Previous studies of *E. coli* macrocolony biofilms formed on agar plates showed that curli expression occurs in the upper layer of the colony, but even in this layer its expression remained heterogeneous [25-27], indicating an interplay between global regulation of curli gene expression by microenvironmental gradients within biofilms and its inherent stochasticity. Differentiation of *E. coli* into distinct subpopulations of cells either expressing or not expressing curli was also observed in submerged biofilms formed in liquid cultures, whereby curli expression was associated with cellular aggregation [28]. Furthermore, bi- or multimodality of csgD reporter activity was also observed in the early stationary phase among planktonic cells in *S. Typhimurium* [29, 30] and *E. coli* [27]. Given established c-di-GMP-dependent regulation of CsgD activity, it was proposed that bistable curli expression originates from a toggle switch created by mutual inhibition between DgcM and PdeR, which could act as a bistable switch [27, 31].
In this study we demonstrate that stochastic differentiation of *E. coli* csgBAC operon expression into distinct subpopulations of curli-positive and -negative cells occurs during the entry into the stationary phase in a well-stirred planktonic culture, and thus in absence of any environmental gradients. Similar stochastic and reversible differentiation could be observed among cells growing in conditioned medium in the microfluidic channel. The upstream regulation by c-di-GMP is not required to establish the bimodality of curli expression, but it determines the fraction of curli-positive cells and enhances the stability of curli activation.

**Materials and methods**

**Bacterial strains and plasmids**

All strains and plasmids used in this study are listed in Table S1. Derivative of *E. coli* W3110 [26] that was engineered to encode a chromosomal transcriptional sfGFP reporter downstream of the csgA gene [28] (VS1146) was used here as the wildtype strain. Gene deletions were obtained with the help of P1 phage transduction using strains of the Keio collection [32] as donors, and kanamycin resistance cassette was removed using FLP recombinase [33]. For expression, *dgcE* and *pdeH* genes were cloned into pTrc99A vector [34].

**Growth conditions for planktonic cultures**

Planktonic *E. coli* cultures were grown in tryptone broth (TB) medium (10 g tryptone, 5 g NaCl per liter), supplemented with antibiotics where necessary. Overnight cultures grown at 30°C were diluted 1:100, unless indicated otherwise, in 5-10 ml of fresh TB and grown at 30°C at 200 rpm in 100 ml flasks in a rotary shaker until indicated OD$_{600}$ or overnight (18-25 h; OD$_{600}$ ~ 1.3-1.8). Alternatively, cultures were grown in 96-well plates with linear shaking in a plate reader, with 200 µl culture per well. Where indicated, bacterial cultures were supplemented with either 1 mM L-serine (after 6 h of growth) or 0.1–10 mg/l DL-serine hydroxamate at inoculation.

**Growth and quantification of submerged biofilms**

Submerged biofilms were grown and quantified as described previously [28], with minor modifications. Overnight bacterial cultures grown in TB were diluted 1:100 in fresh TB medium and grown at 200 rpm and 30°C in a rotary shaker to OD$_{600}$ of 0.5. These cultures were then diluted in fresh TB medium to a final OD$_{600}$ of 0.05, and 300 µl was loaded onto a 96-well plate (Corning Costar, flat bottom; Sigma-Aldrich, Germany) and incubated without shaking at 30°C for 46 h.

For quantification of biofilm formation, the non-attached cells were removed and the wells were washed once with phosphate-buffered saline (PBS; 8 g NaCl, 0.2 g KCl, 1.44 g Na$_2$HPO$_4$, 0.24 g KH$_2$PO$_4$). Attached cells were fixed for 20 min with 300 µl of 96% ethanol, allowed to dry for
40 min, and stained with 300 µl of 0.1% crystal violet (CV) solution for 15 min at room temperature. The wells were subsequently washed twice with 1x PBS, incubated with 300 µl of 96% ethanol for 35 min and the CV adsorption was measured at OD595 using INFINITE M NANO* plate reader (Tecan Group Ltd., Switzerland). These CV values were normalized to the OD600 values of the respective biofilm cultures.

**Macrocolony biofilm assay**

Macrocolony biofilms were grown as described previously [26]. Briefly, 5 µl of the overnight liquid culture grown at 37°C in lysogeny broth (LB) medium (10 g tryptone, 10 g NaCl, and 5 g yeast extract per liter) was spotted on salt-free LB agar plates supplemented with Congo red (40 µg/ml). Plates were incubated for 8 days at 28°C.

**Fluorescence measurements**

Measurements of GFP expression in an INFINITE M1000 PRO plater reader (Tecan Group Ltd., Switzerland) were done using fluorescence excitation at 483 nm and emission at 535 nm. Relative fluorescence was calculated by normalizing to corresponding OD600 values of the culture. For fluorescence measurements using flow cytometry, aliquots of 40-300 µl of liquid bacterial cultures were mixed with 2 ml of tethering buffer (10 mM KH₂PO₄, 10 mM K₂HPO₄, 0.1 mM EDTA, 1 µM L-methionine, 10 mM lactic acid, pH 7.0). Macrocolonies were collected from the plate, resuspended in 10 ml of tethering buffer and then aliquots of 40 µl were mixed with 2 ml of fresh tethering buffer. Samples were vigorously vortexed and then immediately subjected to flow cytometric analysis using BD LSRFortessa Sorp cell analyzer (BD Biosciences, Germany) using 488-nm laser. In each experimental run, 50,000 individual cells were analyzed. Absence of cell aggregation was confirmed by using forward scatter (FSC) and side scatter (SSC) parameters. Data were analyzed using FlowJo software version v10.7.1 (FlowJo LLC, Ashland, OR, US), applying a software-defined background fluorescence subtraction.

**Microfluidics**

Conditioned medium was prepared by cultivating wildtype *E. coli* in TB in a rotary shaker at 30°C for 20 h, after which the cell suspension was centrifuged at 4000 rpm for 10 min, medium was filter-sterilized and stored at 4°C. Mother machine [35] microfluidics device was designed, fabricated and operated as described in Supporting protocols. *E. coli* cells from the overnight culture in TB were loaded into the mother machine growth sites by manual infusion of the cell suspension through one of the two inlets using a 1-ml syringe. Cells were first allowed to grow at 30 °C for 4 h in fresh TB, then switched to the conditioned TB and cultivated for up to 26 h. Phase contrast and GFP fluorescence images were acquired using a Nikon Eclipse Ti-E
inverted microscope with a time interval of 10 min. Details of image analysis are described in Supporting protocols.

Results

Bimodal curli expression is induced in planktonic culture

In order to characterize curli expression in planktonic culture of *E. coli*, we followed the induction of chromosomal transcriptional reporter of csgBAC operon, where the gene encoding for green fluorescent protein (GFP) was cloned with a strong ribosome binding site as a part of the same polycistronic RNA downstream of csgA [28]. In our previous study of submerged *E. coli* biofilms, this reporter showed bimodal expression both in the surface-attached biofilm and in the pellicle at the liquid-air interface [28]. When *E. coli* culture was grown at 30°C in tryptone broth (TB) liquid medium, this reporter became induced during transition to the stationary phase (Figure 1A), which is consistent with previous reports [12, 14]. The observed induction of curli expression occurred at similar density in the cultures with different initial inoculum size. In both cases the onset of induction apparently coincided with the reduction of the growth rate, which likely occurs due to depletion of amino acids in the medium and induction of the stringent response [36, 37], consistent with proposed role of stringent response in the regulatory cascade leading to curli gene expression [18, 23]. In agreement with that, curli expression was strongly reduced when *E. coli* culture was grown in a concentrated TB medium (Figure S1A) or when TB medium was supplemented with serine (Figure S2A). Moreover, the induction of curli reporter was strongly enhanced by addition of serine hydroxamate (SHX), which is known to mimic amino acid starvation and induce stringent response [38] (Figure S2A).

In order to investigate whether curli expression was uniform or heterogeneous across planktonic *E. coli* population, we next measured curli reporter activity in individual cells using flow cytometry. The reporter was induced only in a fraction of cells, and this bimodality of curli expression became increasingly more pronounced at later stages of culture growth, reaching its maximum in the overnight culture (Figure 1B). Thus, the bimodal induction of curli gene expression is observed not only in biofilms but also in a well-mixed planktonic culture. While curli activation was more pronounced in a cell culture growing in an orbital shaker (Figure 1B), bimodality was also observed during culture growth in the plate reader (Figure S1B). Notably, stimulation of curli expression by SHX or its suppression by additional nutrients affected the fraction of positive cells rather than their expression levels (Figure S1B and Figure S2B).

Bimodality of curli expression does not require regulation by c-di-GMP

Subsequently, we investigated dependence of curli expression in planktonic culture on the upstream regulation by c-di-GMP (Figure 2A). As expected, no activation of curli reporter was
observed in the absence of MlrA (Figure 2B). Curli expression was also affected by the lack of enzymes that control MlrA activity at the global (DgcE and PdeH) or local (DgcM and PdeR) level of c-di-GMP regulation (Figure 2C). Consistently with their established regulatory roles, deletions of cyclase genes dgcE and dgcM resulted in nearly all cells being curli-negative, whereas deletions of diesterase genes pdeH and pdeR led to activation of curli expression in the majority of cells within the planktonic population. Notably, in all cases a small fraction of positive (for cyclase knockouts) or negative (for esterase knockouts) cells could be detected, indicating that neither of these knockouts entirely eliminates the bimodality of curli expression. This conclusion could be further confirmed by combined deletions of cyclase and diesterase genes. Removal of the entire global level of c-di-GMP regulation in \( \Delta pdeH \Delta dgcE \) strain led to a bimodal pattern of curli activation (Figure 2D) that was similar to that observed in the wildtype cells. Even more surprisingly, the distribution of curli expression remained bimodal upon removal of the local level of c-di-GMP regulation in \( \Delta pdeR \Delta dgcM \) strain, although the fraction of curli-positive cells was reduced and heterogeneity of their expression levels increased in this background. The bimodality of curli expression was also retained in the quadruple knockout strain lacking all four cyclase and diesterase genes (Figure 2E). Thus, whereas both global and local c-di-GMP-dependent regulation of MlrA activity clearly affect the fraction of curli-positive cells, they are apparently not required to activate curli expression or to establish its bimodality in the planktonic cell population.

*Vibrio cholerae* transcriptional regulator VpsT, a close homologue of CsgD, has been shown to be directly regulated by binding to c-di-GMP [39]. Furthermore, in *S. Typhimurium* c-di-GMP was proposed to regulate csgD expression not only at transcriptional but also at a posttranscriptional level [40]. We thus aimed to verify that *E. coli* curli gene expression was no longer sensitive to the global cellular level of c-di-GMP in the absence of the local PdeR/DgcM regulatory module. Indeed, whereas the overexpression of c-di-GMP cyclase DgcE or phosphodiesterase PdeH had strong impacts on the fraction of curli-positive cells in the wildtype, the quadruple mutant was insensitive to such overexpression (Figure S3), confirming that in this background the expression of the csgBAC reporter is no longer affected by the global pool of c-di-GMP.

**Curli activation shows higher variability in absence of c-di-GMP regulation**

We next explored how the fraction of curli-positive cells in the population depends on the conditions of culture growth, with and without regulation by c-di-GMP. As mentioned above, even in the wildtype strain the fraction of curli-positive cells was smaller in cultures grown in multi-well plates (Figure S1B) compared to the incubation in the flask in an orbital shaker (Figure 1B). However, this reduction in the number of curli-positive cells was much more pronounced for \( \Delta pdeR \Delta dgcM \) or \( \Delta pdeH \Delta dgcE \Delta pdeR \Delta dgcM \) strains, where only a small
fraction of cells became positive under these growth conditions (Figure 3A,B and Figure S4).  
Also individual dgc and pdh gene knockout strains showed reduced activation of curli reporter (Figure S4). Of note, another difference with the flask culture was that the low-fluorescence peak of the wildtype culture was not fully negative but apparently contained a large fraction of cells with incompletely activated curli reporter, which could also be seen in \(\Delta pdeH\) or \(\Delta pdeR\) knockouts but not in the \(\Delta pdeH\ \Delta dgcE\ \Delta pdeR\ \Delta dgcM,\ \Delta pdeR\ \Delta dgcM\) or \(\Delta pdeH\ \Delta dgcE\) strains (Figure 3B and Figure S4). Similar results were obtained even upon prolonged incubation in the plate reader (Figure S5), confirming that the observed difference with the overnight flask culture was not because of the different growth stage. 
We further tested reporter activation under growth conditions that favour biofilm formation. During formation of static submerged biofilms in multi-well plates where cultures are grown without shaking, the overall curli activation in the populations of \(\Delta pdeH\ \Delta dgcE,\ \Delta pdeR\ \Delta dgcM\) or \(\Delta pdeH\ \Delta dgcE\ \Delta pdeR\ \Delta dgcM\) cells (Figure 3C) as well as in the individual knockout strains (Figure S6A) was comparable to that in the culture grown in the orbital shaker (Figure 2B-E). Curli gene activation in individual mutant strains correlated well with the levels of submerged biofilm formation (Figure S6B), although the lack of regulation by c-di-GMP resulted in stronger reduction of the biofilm biomass, consistent with other roles of c-di-GMP in biofilm formation besides curli regulation. 
We also grew all strains in the form of macrocolony biofilms on an agar plate [26]. Interestingly, here the extent of reporter activation in the \(\Delta pdeR\ \Delta dgcM\) and \(\Delta pdeH\ \Delta dgcE\ \Delta pdeR\ \Delta dgcM\) strains was much higher and comparable to that of the wildtype (Figure 3D) and even individual \(\Delta dgcE\) and \(\Delta dgcM\) knockouts showed high fraction of curli-positive cells (Figure S7A), consistent with their stronger Congo red staining compared to the \(\Delta mlrA\) negative control (Figure S7B). Summarily, these results confirm that the regulation by c-di-GMP is not required for (bimodal) curli activation, but also suggest that in absence of this control the fraction of culri-positive cells is more sensitive to growth conditions.  

**Curli activation is a stochastic and reversible process stabilized by c-di-GMP**  
In order to investigate the dynamics of curli activation, and the effects of c-di-GMP regulation, at the single-cell level, we utilized “mother machine”, a microfluidic device where growth of individual bacterial cell lineages could be followed in a highly parallelized manner over multiple generations [35] (Figure 4, Figure S8 and Supporting protocols). To activate curli expression in continuously growing single cells by mimicking nutrient depletion, *E. coli* wildtype or \(\Delta pdeH\ \Delta dgcE\ \Delta pdeR\ \Delta dgcM\) cells were first loaded into the mother machine from the overnight culture, allowed to grow in fresh TB medium for several generations and then shifted to the TB medium that was pre-conditioned by growing the batch culture (see Materials and Methods and Supporting protocols). For both strains, a fraction of curli-positive cells was observed at
the beginning of the experiment since cells originated from the overnight culture, but all cells
turned off curli expression after resuming exponential growth in the fresh medium (Figure 4A,B,
Figure S9, Figure S10, Figure S11, Movie S1 and Movie S2). Following shift to the conditioned
medium, cell growth rate was strongly reduced (Figure 4A,C,E and Figure S9A) to
approximately the same low growth rate for both strains. After several generations of slow
growth in the conditioned medium, individual cells of both strains activated curli expression,
while other cells remained in the curli-off state (Figure 4A,C and Figure S11). Importantly, we
observed that after several generations in the curli-on state, individuals cells of both strains
turned curli expression off again during continuous growth in the conditioned medium (Figure
4B,D and Figure S10), and in some cases there was even a second activation event.
Despite these overall similarities, the dynamics of curli reporter activation showed several
differences between the wildtype and the ∆pdeH ∆dgcE ∆pdeR ∆dgcM strain. Most clearly, the
rate of curli activation in individual cells was apparently higher in absence of the c-di-GMP
regulation (Figure 4B,D,F, Figure S10, Figure S12 and Figure S13). Additionally, the induction
showed greater intercellular heterogeneity in the ∆pdeH ∆dgcE ∆pdeR ∆dgcM strain (Figure
4F, Figure S10, Figure S12 and Figure S13). Thus, the control of curli expression by c-di-GMP
reduces the rate but increases the stability of curli induction.

Discussion
Expression of the curli biofilm matrix genes is known to be heterogeneous or even bistable in
communities of E. coli [25-28] and S. Typhimurium [29, 30], which might have important
functional consequences for the biomechanics of bacterial biofilms [27] and for stress
resistance and virulence of bacterial populations [29, 30]. In the well-structured microcolony
biofilms, differentiation into subpopulations with different levels of the curli matrix production is
largely deterministic and driven by gradients of nutrients and oxygen [18]. Bimodality of curli
expression might also emerge stochastically, in a well-mixed population or between cells within
the same layer of the macrocolony [27, 29, 30]. How this bimodality originates within the
extremely complex regulatory network of curli genes [17, 19, 23, 27] remains a matter of
debate. Although earlier studies in S. Typhimurium proposed that bistable expression of curli
might be a consequence of positive transcriptional feedback in csgD regulation [29, 30, 41],
the most recently proposed model attributed bistability to the properties of the c-di-GMP
regulatory switch formed by DgcM, PdeR and MlrA [27, 31]. Furthermore, the dynamics of curli
gene expression in individual cells remains unstudied.
Here, we investigated the bimodal expression of the major curli csgBAC operon at the
population level in the well-stirred planktonic E. coli culture as well as on the single-cell level
in the microfluidic device. Consistent with previous studies [12, 14], the induction of curli
expression in growing E. coli cell population was observed during the entry into the stationary
phase of growth. Curli activation was apparently dependent on depletion of the amino acids from the medium, since it could be suppressed by increasing the levels of nutrients or, specifically, of serine. It might thus be related to induction of the stringent response, and consistently, it was enhanced by the SHX-mediated stimulation of the stringent response. We further observed that activation of the csgBAC operon was strongly bimodal under all tested conditions, even in absence of any nutrient or other gradients. Stochastic nature of this activation was confirmed by incubation of E. coli cells in the microfluidic device, where upon a shift to the conditioned medium only a fraction of the cell population turned on the curli expression. Such differentiation is apparently consistent with previous observations of the bimodal csgD expression in S. Typhimurium [29, 30] and in E. coli [27]. However, whether the bimodality of the csgBAC expression is caused by the bimodal expression of csgD remains clear, since the latter was reported only in the later stationary phase of the culture growth [27, 29, 30], whereas in our experiments the csgBAC reporter showed bimodality already at an earlier stage.

In contrast to those previous interpretations of the csgD expression pattern as bistability, our data suggest that curli activation is only transient, and therefore bimodal but not bistable. Under conditions of continuous cell growth in the microfluidic device, the activation of curli expression is followed by its inactivation, indicating a pulsatile activation of the curli-positive state. Pulsing in expression was proposed to be common to many gene regulatory circuits [42], although only few well-studied examples such as stress response and differentiation in Bacillus subtilis [43, 44] are available. Pulsatile expression has also been recently described in E. coli for the upstream regulator of curli, RpoS [45], as well as for the flagellar regulon [46, 47] that is anti-regulated with curli [28]. However, in neither of these cases did pulsing lead to apparent bimodality of expression, and their relation to the observed pulses in curli expression thus remains to be seen.

How does such stochastic pulsing of curli expression observed at the single-cell level lead to the differentiation into two very distinct subpopulations in the batch planktonic culture? This could be likely explained by the timing of curli activation, and by the duration of these observed expression pulses: Since the curli expression is only turned-on during transition to the stationary phase, individual cells can stochastically activate curli genes just before the culture growth ceases. In contrast to the continuous culture, subsequent reversion to the curli-negative state by inactivation and dilution by cell division is no longer possible in the stationary batch culture, and these initially curli-expressing cells will thus remain positive.

Importantly, the regulation by c-di-GMP is not required for the bimodal curli expression in E. coli, since the differentiation into distinct subpopulations still occurred when the control of CsgD expression by c-di-GMP was abolished, as observed both in the planktonic cultures and in the microfluidic device. The level of csgBAC expression in curli-positive cells was also little
affected by the c-di-GMP control. Nevertheless, this control is important during the
establishment of bimodality: Firstly, global levels of c-di-GMP determine the fraction of curli-positive cells, via regulation of MlrA activity, which is respectively inhibited by PdeR when c-di-GMP is low and activated by DgcM at high c-di-GMP [22, 23]. Secondly, the c-di-GMP-dependent control affects the dynamics of curli gene pulsing, with the faster but more heterogeneous activation of curli expression in the absence of the c-di-GMP control. Thirdly, and possibly related to the previous observation, this control might also ensure that the fraction of curli-positive cells is less variable dependent on the growth conditions of *E. coli* population. Thus, our study reveals a novel regulatory function exhibited by the second messenger c-di-GMP, in stabilizing the bimodal gene expression.

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Figure 1: Bimodal activation of curli gene expression in *E. coli* planktonic cultures. *E. coli* cells carrying genomic transcriptional reporter of *csgBAC* operon were grown in liquid tryptone broth (TB) medium at 30 °C under constant shaking. (A) Optical density (OD$_{600}$) and relative fluorescence (fluorescence/OD$_{600}$; AU, arbitrary units) of the culture during growth in a plate reader, starting from two different dilutions of the overnight culture. Error bars indicate standard error of the mean (SEM) of 10 technical replicates. (B) Distribution of single-cell fluorescence levels in cultures grown to indicated OD$_{600}$ or overnight (ON; 25 h) in an orbital shaker, measured by flow cytometry.
Figure 2: Regulation of curli expression by c-di-GMP. (A) Current model of regulation of curli gene expression by c-di-GMP in *E. coli*, adapted from [27]. The regulation is mediated by two pairs of diguanylate cyclases (DGCs; blue) and phosphodiesterases (PDEs; orange). PdeH and DgcE control global level of c-di-GMP, whereas PdeR and DgcM mediate local c-di-GMP-dependent regulation of curli gene expression by controlling activity of transcription factor MlrA, which activates another curli-specific transcription factor CsgD. (B-E) Flow cytometry measurements of curli gene expression in *E. coli* planktonic cultures grown overnight in flasks in an orbital shaker, shown for the wildtype (WT) and ΔmlrA knockout (B), and individual (C), double (D) and quadruple (E) knockouts of DGC or PDE enzymes, as indicated. Fraction of positive cells in the population (mean of three biological replicates ± SEM) is indicated for each strain. Note that the scale in the y axes is different for individual strains to improve readability.
Figure 3: Dependence of curli expression on c-di-GMP regulation under different growth conditions. (A) Relative fluorescence of transcriptional curli reporter in planktonic culture grown in a plate reader. Error bars indicate SEM of 5 technical replicates. (B-D) Flow cytometry measurements of curli expression in the wildtype (WT) and indicated knockout strains after 24 h of growth either in a plate reader (B), as submerged biofilms (C) or as macrocolony biofilms (D). Measurements for other knockouts are shown in Figures S4, S6A, S7A. Fraction of positive cells in the population (mean of three biological replicates ± SEM) is indicated for each strain. Note that the scale in the y axes is different for individual strains to improve readability.
Figure 4: Impact of c-di-GMP regulation on dynamics of curli induction in individual cells. *E. coli* cells in a microfluidic device (mother machine) were shifted from a fresh to conditioned TB medium after 4 h of growth to induce curli expression. (A-D) Examples of image time series and single-cell fluorescence traces for the wildtype (WT) (A,B) and for the ΔpdeH ΔdgcE ΔpdeR ΔdgcM strain disabled in the c-di-GMP regulation (C,D) growing in a mother machine in one experiment. Expression of the curli reporter is indicated by the green overlay on the phase contrast image. (E) Median instantaneous growth rate — fold rate of change in length — of cells grown in microfluidics experiments as described in (A). The switch to conditioned medium is at time zero, as indicated. Shaded area is the interquartile range. The number of cells in the device varies with time, but is on average *n* = 296 for WT and *n* = 522 for the quadruple knockout. (F) Median curli expression profile for single-cell traces aligned by the time at which they exceed a threshold of 10^3 fluorescence units. Shaded area is interquartile range; *n* = 128 for WT and 230 for the quadruple knockout.
Figure S1: Dependence of curli gene expression on nutrient levels. Wildtype *E. coli* cultures were grown as in Figure 1A but with different indicated concentrations of TB. (A) Bacterial growth and activity of transcriptional curli reporter. Error bars indicate SEM of 6 technical replicates. (B) Distribution of single-cell fluorescence levels after 24 h of growth in a plate reader measured by flow cytometry. Note that the scale in the y axes is different for individual conditions to improve readability.
Figure S2: Stimulation of curli gene expression by stringent response. Wildtype E. coli cultures were grown as in Figure 1A but with addition of either indicated concentrations of serine hydroxamate (SHX) at inoculation point or 1 mM serine after 6 h of growth. (A) Bacterial growth and activity of transcriptional curli reporter. Error bars indicate SEM of 6 technical replicates. (B) Distribution of single-cell fluorescence levels after 24 h of growth in a plate reader measured by flow cytometry. Note that the scale in the y axes is different for individual conditions to improve readability.
Figure S3: Decoupling of curli gene expression from c-di-GMP regulation in the absence of PdeR/DgcM regulatory module. *E. coli* wildtype (WT) cells or cells lacking c-di-GMP regulatory enzymes (ΔpdeH ΔdgcE ΔpdeR ΔdgcM) were transformed with either empty pTrc99a plasmid (control) or with pTrc99a plasmid carrying dgcE (pTrc99a::dgcE) or pdeH (pTrc99a::pdeH) genes. Expression from the vector was induced with 1 μM IPTG. Bacteria were grown overnight in flasks with shaking and cultures were subjected to the flow cytometry analysis. Note that the scale in the y axes is different for individual strains to improve readability.
Figure S4: Curli gene expression in cultures grown in a plate reader. *E. coli* cells were grown as in Figure 3A,B. Flow cytometry measurements for indicated knockouts are shown. Measurements in other strains are shown in Figure 3B. Fraction of positive cells in the population (mean of three biological replicates ± SEM) is indicated for each strain. Note that the scale in the y axes is different for individual strains to improve readability.
Figure S5: Curli gene expression upon prolonged cultivation in a plate reader. Wildtype (WT) and (ΔpdeH ΔdgcE ΔpdeR ΔdgcM) knockout strains were grown in a plate reader as in Figure 2 A,B, but for 36 h. (A) Induction of transcriptional curli reporter. Error bars indicate SEM of 10 technical replicates. (B) Distribution of single-cell fluorescence levels in populations of both strains after 36 h of growth measured by flow cytometry.
Figure S6: Curli gene expression in submerged biofilm cultures. (A) Distribution of single-cell fluorescence levels in populations of indicated knockout strains after 46 h of submerged biofilm culture growth, measured by flow cytometry. Measurements in other strains are shown in Figure 3C. Fraction of positive cells in the population (mean of three biological replicates ± SEM) is indicated for each strain. Note that the scale in the y axes is different for individual strains to improve readability. (B) Biofilm formation by indicated strains, quantified using crystal violet (CV) staining. Error bars indicate SEM of 3 independent replicates.
Figure S7: Curli expression in macrocolony biofilms. (A,B) Flow cytometry measurements of curli expression (A) and images of microcolonies of indicated strains (B) after 8 days of growth. Flow cytometry measurements for other strains are shown in Figure 3D. Fraction of positive cells in the population (mean of three biological replicates ± SEM) is indicated for each strain. Note that the scale in the y axes is different for individual strains to improve readability.
Figure S8: Design of the microfluidic mother machine chip. (A) Schematic overview on the channel layout, featuring four supply channels (green) for cell inoculation and media supply. (B) Detailed view of the area marked by a rectangle in (A), showing the switching junction and a part of the mother machine cultivation sites (blue). The junction is formed by two inlets, leading to one central supply channel. The control of the pressure at each inlet allows to choose the medium flowing through the supply channel, and, ultimately, to the mother machine cultivation sites. Residual medium flows out through the waste channels located on both sides from the central channel. Medium flowing through the supply channel exits the chip through one outlet. (C) Detailed view of the area marked in (B) by a rectangle, showing the mother machine cultivation sites. Each of the four channels contains 57 mother machine cultivation sites, which contain 30 mother machine traps with widths of 0.9, 1, or 1.1 µm. The mother machine traps feature a 0.3 µm wide constriction on the bottom, preventing the mother cell from exiting the trap while allowing medium perfusion. The supply channels (green) are 8 µm in depth, the mother machine traps (blue) are 0.8 µm in depth. (D) On-chip medium switching visualized by merged phase contrast and mCherry images of the channel junction. Media are supplied through separate inlets (top and bottom), which are separated in the center of the channel by a PDMS barrier. The direction of flow is indicated by white arrows. Water was supplied through the top inlet, while a 0.2 µM sulforhodamine B solution was supplied through the bottom inlet, visualizing the flow pattern in the junction. The pressure at the top inlet was kept constant at 200 mbar. Depending on the pressure set at the bottom inlet, it is possible to select which one of the two media flows into the central supply channel to the mother machine growth sites.
Figure S9: Growth rates and fraction of curli expressing cells over time. Stationary phase cells were introduced into mother machine devices, supplied with fresh medium and then switched to conditioned medium after four hours of growth, as in Figure 4. (A) Median instantaneous growth rates for the wildtype and for the \( \Delta pdeH \Delta dgcE \Delta pdeR \Delta dgcM \) strain disabled in the c-di-GMP regulation. Growth rate drops rapidly and cells switch on curli expression after a switch to conditioned medium. Shaded area is interquartile range. (B) Fraction of cells with fluorescence exceeding 1000 units. (C) Number of detected cells. Two biological replicates (r1 and r2) were performed for each strain; data for the r1 replicate are also shown in Figure 4. Note that in the experiment r2 cells were only imaged after medium switching.
Figure S10: Single-cell traces of cell fluorescence for all cells for the wildtype and for the c-di-GMP-regulation disabled strain. Data are from the same biological replicates (r1 and r2) as in Figure S9.
Figure S11: Distributions of curli expression at different time points in the microfluidics experiment. Shown are kernel density estimates of curli expression in the wildtype WT (A) and in the c-di-GMP-regulation disabled strain (B) at selected time points, for the replicate presented in Figure 4. In time order, for WT, n = 18, 158, 332, 317, 335, 342 and 285, and for the ΔpdeH ΔdgcE ΔpdeR ΔdgcM strain, n = 25, 247, 525, 546, 581, 546 and 544.
Figure S12: The rate and variability of curli induction for the wildtype and for the c-di-GMP-regulation disabled strain. (A) Median curli expression, (B) mean curli expression, (C) number of cells for traces from both microfluidics experiments (r1 and r2) aligned by the time at which they exceeded a threshold of $10^3$ fluorescence units. Shaded area is interquartile range in (A) or standard error in (B). Compared to the WT, the rate of curli induction is faster but traces show more variability in a mutant without the global or local c-di-GMP regulatory modules.
Figure S13: Distributions of curli induction parameters for the wildtype and for the c-di-GMP-regulation disabled strain. Shown are histograms of the times at which a threshold of $10^3$ fluorescence units were crossed (A), the maximum rates of increase in fluorescence (B), or the fluorescence amplitudes at the first peak (C) for cell traces from both microfluidics experiments. Though switch timing is similar, rates of curli induction are faster and peak amplitudes more variable in a mutant without the global or local c-di-GMP regulatory modules.
### Table S1. *E. coli* strains and plasmids used in this study.

| Strains | Relevant genotype | Reference |
|---------|-------------------|-----------|
| W3110   | W3110 derivative with functional RpoS | [1] |
| VS1146  | W3110 csgA::csgA_RBS_sgfp | [2] |
| VS1857  | VS1146 ΔmlrA | This work |
| VS1732  | VS1146 ΔpdeH | This work |
| VS1720  | VS1146 ΔdgcE | This work |
| VS1258  | VS1146 ΔpdeR | This work |
| VS1257  | VS1146 ΔdgcM | This work |
| VS1717  | VS1146 ΔpdeH ΔdgcE | This work |
| VS1713  | VS1146 ΔpdeR ΔdgcM | This work |
| VS1729  | VS1146 ΔpdeH ΔdgcE ΔpdeR ΔdgcM | This work |

### Plasmids

| | Expression vector; *Ptrc* promoter inducible by isopropyl-β-D-thiogalactopyranoside (IPTG); pBR ori; Ap<sup>R</sup> | Reference |
|---|----------------------------------------------------------|-----------|
| pTrc99a | | [3] |
| pVS2689 | pTrc99a::pdeH | This work |
| pVS1644 | pTrc99a::dgcE | This work |

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Supporting protocols

Design and fabrication of the mother machine. A microfluidic device was used for time-resolved studies of *E. coli* growth and curli expression, which features mother machines traps for the observation of one-dimensional cell growth. The design of the microfluidic device is shown in Figure S8. The channel structure of the microfluidic device features four independent channels with a depth of 8 µm, allowing the realization of four conditions, e.g. different media compositions, in one experiment. Each channel contains two inlets enabling on-chip switching between two different media, which are supplied at defined pressures. By controlling the pressure ratio between the two inlets, the medium supplied at the higher pressure flows through the central channel following the junction, while the other medium is pushed away from the junction and flows out through the waste channel (Figure S8D). The medium which flows through the central channel reaches the mother machine cultivation sites before exiting the chip through a single outlet.

The device was produced using a two-layer soft lithography method as described previously[1]. Based on our in-house made design of the channel layout, a 100 mm silicon wafer was produced by e-beam lithography (ConScience, Sweden). The wafer contains the channel layout as a positive relief. The mother machine traps, which are shown in blue in Figure S8A-C, were structured by etching the wafer by 0.8 µm, giving the mother machine the appropriate vertical dimension for cell trapping. The supply channels, which are shown in green in Figure S8A-C, are implemented as photoresist structures with a height of 8 µm on the wafer. The wafer served as a master mold for liquid polydimethylsiloxane (Sylgard 184 PDMS, VWR International GmbH, Germany), which was mixed at volumetric ratio of 7:1 with a cross-linking agent, degassed in a desiccator for 30 minutes and poured over the wafer to a height of approximately 4 mm and thermally cured at 80 °C overnight. The cured PDMS was peeled off from the wafer and manually cut into separate chips. Inlet and outlet holes were punched with a 0.75 mm punching tool (Robbins True-Cut Disposable Biopsy Punch 0.75mm with Plunger, Robbins Instruments, USA). The surface of the chip was cleaned by a rinse with isopropanol and the application of adhesive tape (tesafilm, Germany) prior to bonding. The chip was irreversibly bonded to a glass substrate by applying oxygen plasma to both chip and glass surfaces (Diener Femto, Diener GmbH, Germany) and bringing the treated surfaces together. The bond was strengthened by storing the bonded device in the oven at 80 °C for two minutes. The device was mounted on an inverted fluorescence microscope (Nikon Eclipse Ti, Nikon Corporation, Japan) equipped with an incubator. The microscope setup included an Andor Zyla 4.2 sCMOS camera (Oxford Instruments, UK), an objective with 100x magnification (Plan Apochromat λ Oil, NA=1.45, WD=170 µm; Nikon, Japan) and a perfect focus system (Nikon Corporation, Japan) for focus drift compensation.
Growth experiments. *E. coli* cells were allowed to grow in TB medium until the stationary phase prior to inoculation into the device. Conditioned medium was prepared by cultivating the wildtype *E. coli* cells in TB medium for 20 hours, after which the cell suspension was centrifuged at 4000 rpm for ten minutes, the medium was filter sterilized and stored at 4°C. The mother machine growth sites were loaded with the undiluted cell suspension by manual infusion of the cell suspension through one of the two inlets using a 1-ml syringe. The connection between the syringe and the chip was realized by tygon tubing (Tygon S-54-HL, inner diameter = 0.51 mm, outer diameter = 1.52 mm, VwR International GmbH, Germany) in combination with blunt dispensing needles (General purpose tips, inner diameter = 0.41 mm, outer diameter = 0.72 mm, Nordson EFD, USA). Medium flow was controlled by programmable pressure regulators (LineUP FlowEZ, FLUIGENT, France), which generated flow by applying pressure on 50-ml medium reservoirs (P-CAP series, FLUIGENT, France). Fresh and conditioned TB medium were respectively filled in separate reservoirs, and each one was pressurized by one module of the pressure regulator. After the cell inoculation both media were connected to the inlets of the channel via tygon tubing and blunt dispensing needles. The pressure at the inlet of the fresh TB medium was set to 200 mbar and remained constant throughout the experiment. During the selection of the positions for imaging the pressure at the inlet of the conditioned medium was set to 250 mbar, allowing the conditioned medium to flow through the junction to the mother machine growth sites and thereby maintaining the stationary state of the cells. At the beginning of imaging, the pressure at the inlet of the conditioned medium was reduced to 150 mbar and programmed to increase back to 250 mbar after 4 hours of on-chip cultivation, thereby activating a medium switch from fresh to conditioned TB medium. Phase contrast and GFP fluorescence images were acquired with a time interval of 10 min.

Analysis of microfluidics data. Cells were segmented from phase contrast images by making use of a fully convolutional neural network based on the U-net architecture [2]. A set of manually curated cell outlines was prepared for training (1105 outlines) and validation (346 outlines) of the network. The training set was augmented by scaling, rotation, flipping and addition of white noise. A U-net of depth three, with 8, 16 and 32 filters along the contracting path, was trained to predict cell interiors from phase contrast images. Phase contrast images were normalized by subtracting the median and scaling to intensities expected between the 2nd and 98th percentiles. Cell interiors were defined from the curated outlines by filling each outline and then subjecting it to two rounds of morphological erosion. The erosion step ensured that neighbouring cells predicted by the network were well separated, such that distinct cell instances could be clearly identified simply by thresholding the prediction and labelling.
connected regions. After instance identification, two rounds of morphological dilation restored each mask to its original size. Finally, a smooth outline for each cell was obtained as a two-dimensional spline defined by equidistant knots placed on the mask edge.

To track cells between time points, we applied a length conservation strategy for the cells along each trench. At each time point, we ordered cell outlines by their depth in the trench, with deepest cells first. We then attempted to match, in order, a cell outline in time point $t$ with one or more cell outlines in time point $t+1$, chosen such that the sum of their cell lengths would be conserved within some threshold tolerance. In the trivial case, the length of the first outline at time point $t$ would match that of the first outline at time point $t+1$. In the event of cell division, the cell length at time point $t$ would match the sum of the first two cell lengths at time point $t+1$.

Since cells may grow in length between time points, we also initialised a growth rate parameter for each cell that biased the expected cell lengths for time point $t+1$ as a fold-increase in length. To enable adaptation to the true growth rate, the growth rate parameter was updated by a lagging average over 20 time points. To increase robustness to errors in segmentation, we additionally allowed state transitions from one to many and many to one, and built a proposal tree, which branched for all valid assignments lying within the length thresholds. We searched for the proposal with the lowest average fold-change in matched lengths, but limited branching by retaining only the 10 best proposals for subsequent nodes (cell outlines) in the tree. The length thresholds were deliberately set loosely such that the (sum of) cell length(s) at $t+1$ could decrease at most five-fold or increase at most two-fold relative to the (sum of) cell length(s) at $t$. This increased the number of valid proposals, but was important in cases where the growth rate estimate was poor. For transitions where one cell outline split into more than two, or transitions where multiple outlines merged into one cell, a new label was generated for the corresponding cells at $t+1$. We made one exception to this labelling strategy to account for occasional ambiguity in segmentation near division events, where a cell segmented as two sister cells could later be segmented as a single mother cell. Specifically, when two sister cells — i.e., cells that were previously involved in a division event — merged into one, the label was set back to that of the mother; at the next division event, the labels of the sister cells were also retained. Finally, note that any outlines below a minimum size threshold of 50 pixels were ignored. All errors in tracking were manually curated.

Cell length was estimated from cell regions as the ‘major axis length’ of the Matlab regionprops function — the major axis of the ellipse with same normalised second central moment as the region. Instantaneous growth rates were estimated from the derivative of a smoothing spline fitted to the logarithm of cell length over each cell division cycle. Knots for the spline were placed at intervals of at least 15 time points. Single-cell fluorescence traces were quantified from the median fluorescence within each outline. Background fluorescence varied as a function of time due to the accumulation of cells at some trench exits, so we corrected for
background fluorescence in each trench at each time point using the median value of all non-cell pixels. Fluorescence traces were characterised along branching lineages and were smoothed with a Savitzky-Golay filter of order 3 and window length 21. The derivative of the filter was used to obtain the maximum rate of fluorescence increase. In cases where multiple descendants shared a common peak event before branching, we counted that event only once.

References

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