# archive ouverte UNIGE

http://archive-ouverte.unige.ch

**Article** 

Topological control of the Caulobacter cell cycle circuitry by a polarized single-domain PAS protein

SANSELICIO, Stefano, et al.

#### **Abstract**

Despite the myriad of different sensory domains encoded in bacteria, only a few types are known to control the cell cycle. Here we use a forward genetic screen for Caulobacter crescentus motility mutants to identify a conserved single-domain PAS (Per-Arnt-Sim) protein (MopJ) with pleiotropic regulatory functions. MopJ promotes re-accumulation of the master cell cycle regulator CtrA after its proteolytic destruction is triggered by the DivJ kinase at the G1-S transition. MopJ and CtrA syntheses are coordinately induced in S-phase, followed by the sequestration of MopJ to cell poles in Caulobacter. Polarization requires Caulobacter DivJ and the PopZ polar organizer. MopJ interacts with DivJ and influences the localization and activity of downstream cell cycle effectors. Because MopJ abundance is upregulated in stationary phase and by the alarmone (p)ppGpp, conserved systemic signals acting on the cell cycle and growth phase control are genetically integrated through this conserved single PAS-domain protein.

## Reference

SANSELICIO, Stefano, *et al.* Topological control of the Caulobacter cell cycle circuitry by a polarized single-domain PAS protein. *Nature communications*, 2015, vol. 6, p. 7005

DOI: 10.1038/ncomms8005

PMID: 25952018

Available at: http://archive-ouverte.unige.ch/unige:56084

Disclaimer: layout of this document may differ from the published version.





### **ARTICLE**

Received 20 Oct 2014 | Accepted 24 Mar 2015 | Published 8 May 2015

DOI: 10.1038/ncomms8005

**OPEN** 

# Topological control of the *Caulobacter* cell cycle circuitry by a polarized single-domain PAS protein

Stefano Sanselicio<sup>1</sup>, Matthieu Bergé<sup>1</sup>, Laurence Théraulaz<sup>1</sup>, Sunish Kumar Radhakrishnan<sup>1,†</sup> & Patrick H. Viollier<sup>1</sup>

Despite the myriad of different sensory domains encoded in bacteria, only a few types are known to control the cell cycle. Here we use a forward genetic screen for *Caulobacter crescentus* motility mutants to identify a conserved single-domain PAS (Per-Arnt-Sim) protein (MopJ) with pleiotropic regulatory functions. MopJ promotes re-accumulation of the master cell cycle regulator CtrA after its proteolytic destruction is triggered by the DivJ kinase at the G1-S transition. MopJ and CtrA syntheses are coordinately induced in S-phase, followed by the sequestration of MopJ to cell poles in *Caulobacter*. Polarization requires *Caulobacter* DivJ and the PopZ polar organizer. MopJ interacts with DivJ and influences the localization and activity of downstream cell cycle effectors. Because MopJ abundance is upregulated in stationary phase and by the alarmone (p)ppGpp, conserved systemic signals acting on the cell cycle and growth phase control are genetically integrated through this conserved single PAS-domain protein.

<sup>&</sup>lt;sup>1</sup> Department of Microbiology and Molecular Medicine, Institute of Genetics and Genomics in Geneva (iGE3), Faculty of Medicine/CMU, University of Geneva, Rue Michel Servet 1, Genève 4 1211, Switzerland. <sup>†</sup> Present address: School of Biology, Indian Institute of Science Education and Research, Thiruvananthapuram 695016, India. Correspondence and requests for materials should be addressed to P.V. (email: patrick.viollier@unige.ch)

ellular motility is responsive to external signals such as nutritional changes, but it is also regulated by cues induced systemically during each cell division cycle<sup>1,2</sup>. This latter characteristic has been successfully exploited in motility screens to uncover cell cycle regulators in the model bacterium *Caulobacter crescentus* (herein *Caulobacter*), an aquatic α-Proteobacterium that is easily synchronized by density-gradient centrifugation owing to a cell cycle-regulated capsule<sup>3–6</sup>. Motility is conferred by the polar flagellum, a structure that is required for the dispersal of the swarmer cell type. *Caulobacter* divides asymmetrically into a swarmer cell that resides in a G1-like quiescent state and harbours the flagellum and several pili at the same cell pole, and a replicative (S-phase) cell whose old cell pole is decorated by a cylindrical extension of the cell envelope (the stalk) tipped by an adhesive holdfast (Fig. 1a)<sup>2</sup>. Flagellar motility

along with adhesive properties (conferred by the polar pili and holdfast) and cell division are wired into the *C. crescentus* cell cycle circuitry at the transcriptional level by the master cell cycle regulator CtrA<sup>7</sup>, a DNA-binding response regulator (RR) of the OmpR family<sup>4</sup> whose synthesis is activated in S-phase by the transcriptional regulator GcrA<sup>8</sup>. Following its synthesis, CtrA activates many developmentally regulated promoters, including those of motility, pilus, holdfast and cell division genes<sup>7,9–11</sup>. CtrA also functions as negative regulator of gene expression and, directly and/or indirectly, inhibits firing of the origin of DNA replication (*Cori*). *Cori* fires only once during the *Caulobacter* cell cycle and is bound by CtrA at multiple sites<sup>12,13</sup>.

The DNA-binding activity of RRs such as CtrA is regulated by phosphorylation at a conserved (Asp) residue, a step that is often executed directly by a histidine kinase (HK). Before the

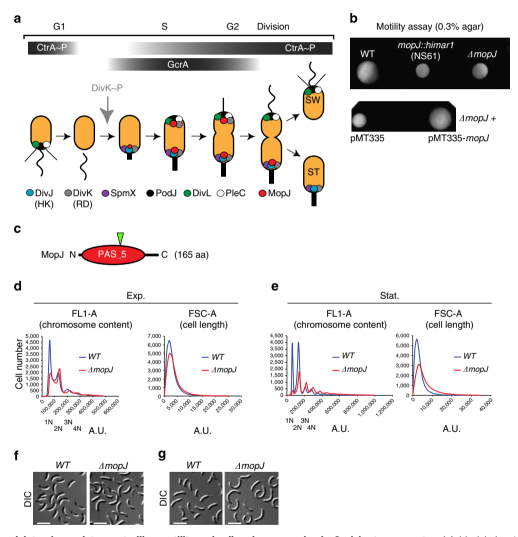


Figure 1 | MopJ, a pleiotropic regulator controlling motility and cell cycle progression in *Caulobacter crescentus*. (a) Model showing the localization of His-Asp signalling proteins and its regulators along with MopJ during *Caulobacter crescentus* cell cycle. The black bars show the abundance of the transcriptional regulators CtrA (and phosphorylated CtrA, CtrA ~ P) and GcrA. The grey arrow indicates when phosphorylation of the DivK receiver domain (RD) by the DivJ histidine kinase (HK) commences. Phosphorylated DivK (DivK ~ P) is present henceforth. The thin vertical line in black represents the flagellum before it rotates (wavy line). The thick vertical line in black represents the stalk. The thin slanted black lines represent the polar pili. (b) (Upper panel) Motility assay on swarm agar plate of *WT, mopJ::himar* and  $\Delta mopJ$  cells. (Lower panel) complementation of  $\Delta mopJ$  cells harbouring the empty vector pMT335 or pMT335-mopJ. (c) Scheme of domain organization of MopJ from N- to C-terminus indicated by the total amino-acid (aa) length. The green arrowhead points to the residue in the *mopJ* coding sequence that was disrupted by the *himar1* transposon insertion. (d,e) FACS analysis of *WT* and  $\Delta mopJ$  cells. Genome content (FL1-A channel) and cell size (FSC-A channel) were analysed by FACS during exponential (d, Exp.) and stationary (e, Stat.) phases in M2G. G1 (1N) and G2 cell (2N) peaks of fluorescence intensity are readily visible (particularly in Stat. phase cells) and are labelled accordingly. S-phase cells are reflected by the broad intermittent signal (absent in Stat. phase cells). (f,g) Differential interference contrast (DIC) images of *WT* and  $\Delta mopJ$  cells during Exp. (f) and Stat. (g) phases in M2G.

phosphate can be accepted by the RRs, the dimeric HK transauto-phosphorylates a conserved histidine (His) in an ATPdependent manner. However, phosphorylation of RRs can also occur indirectly by way of two intermediary components in the His-Asp phosphorvlation pathway, an Asp-containing receiver domain and a His-containing phosphor-transfer domain 14. Complex regulatory schemes emerge when such multicomponent His-Asp pathways are arranged in tandem<sup>15</sup>. The accumulation of phosphorylated CtrA (CtrA~P) underlies such a complex His-Asp pathway topology  $^{15-17}$ . CtrA  $\sim$  P is present in G1-phase, degraded at the G1-S transition by the ClpXP protease<sup>18,19</sup>, to enable the initiation of DNA replication, and then re-synthesized and phosphorylated during S-phase (Fig. 1a). The removal of CtrA~P at the G1-S transition is induced when the DivJ HK phosphorylates its substrate, the DivK receiver domain, which downregulates both the phosphorylation and halflife of CtrA via the DivL tyrosine kinase 15,16,20-24. CtrA abundance and stability are also regulated by the conserved alarmone (p)ppGpp, which tunes cell cycle progression in response to nutritional signals<sup>25-29</sup>.

Superimposed on the temporal events is the spatial regulation of these His-Asp pathway components (Fig. 1a)<sup>30</sup>. At the G1-S transition, DivJ is recruited to the stalked pole by its localization factor SpmX, an event that also is required for optimal DivJ kinase activity *in vivo*<sup>3,31,32</sup>. DivK is enriched at the poles as well, enhancing kinase activity<sup>33</sup>, but a significant fraction of DivK is also dispersed<sup>17,34</sup>. DivK activity is antagonized by the phosphatase PleC that is recruited to the pole opposite the stalk by its localization factor PodJ<sup>32,35-37</sup>. A pleiotropic effector of protein polarization in *C. crescentus* is the polar organizing protein Z that is thought to self-assemble into a polar matrix at both cell poles to sequester proteins like SpmX, DivJ, DivK and others<sup>38-41</sup>. Although the DivJ and SpmX complex is unipolar, DivK localization is bipolar and promoted by SpmX and DivJ activity<sup>17,38</sup>, indicating that DivK localization opposite the stalk is regulated by another factor. The sequestration of DivK to the stalked pole is governed by DivJ, even in the absence of kinase activity<sup>17</sup>.

Here, we report a screen for motility mutants that unearthed a conserved and cell cycle-regulated single-domain PAS (Per-Arnt-Sim)<sup>42</sup> protein (MopJ) that promotes CtrA accumulation during exponential growth and in stationary phase. MopJ is localized to the cell poles where it acts on downstream cell cycle signalling proteins, whereas upstream cell cycle regulators promote MopJ polarization. Our work unveils MopJ as a modulator of the spatio-temporal circuitry controlling bacterial cell cycle progression and as a regulatory node through which stationary phase and (p)ppGpp-dependent nutritional signals are integrated.

#### Results

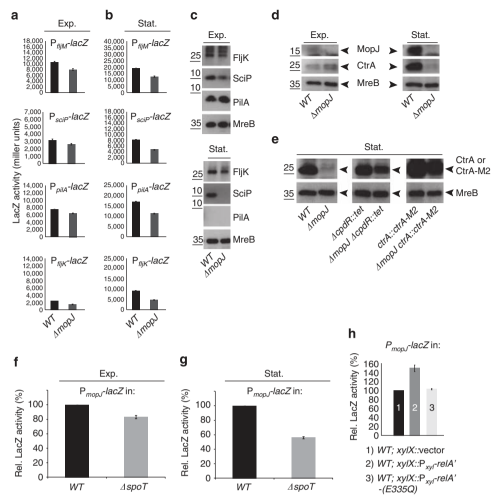
MopJ is a pleiotropic regulator of motility. Our previous comprehensive transposon mutagenesis screen for strains with a motility defect on swarm (0.3%) agar that led to the identification of the spmX gene<sup>3</sup> also yielded one mutant strain (NS61, Fig. 1b) harbouring a *himar1* insertion in the uncharacterized gene  $CCNA\_00999$  (at nucleotide position 1082252 of the *Caulobacter crescentus* wild-type (WT) strain NA1000 (ref. 43). This gene is predicted to encode a single-domain PAS protein (PAS\_5, pfam07310, residues 15–147, Fig. 1c) that is 165 residues in length and is henceforth referred to as MopJ (motility PAS domain associated with DivJ, see below). The PAS domain, a five-stranded antiparallel β-sheet flanked by several α-helices, is generally used to bind small-molecule ligands and is structurally related to the GAF regulatory (GMP-specific and -regulated cyclic nucleotide phosphodiesterase, Adenylyl cyclase and *Escherichia* 

coli transcription factor FhlA) domain often encoded on the same polypeptide<sup>42</sup>. Orthologues of MopJ are also encoded in the genomes of distantly related α-Proteobacteria (Supplementary Fig. 1) such as the animal pathogen Brucella melitensis (BMEI0738), the plant pathogen Agrobacterium tumefaciens (Atu1754) and the plant symbiont Sinorhizobium meliloti (SMc01000). These proteins exhibit a similar domain organization with MopJ, that is, a single PAS 5 domain without any associated regulatory or effector domain (Fig. 1c). The *himar1* transposon in NS61 lies near the middle of the *mop1* gene (that is, after codon 89, Fig. 1c), presumably disrupting its function. In support of this notion, an in-frame deletion of mop!  $(\Delta mop J)$  in WT cells recapitulated the motility defect observed in the mopJ::himar1 mutant on soft agar (Fig. 1b). As the motility defect of  $\Delta mop J$  cells can be corrected by supplying mop J in trans on a plasmid (pMT335-mopl) under the control of the vanillateinducible promoter P<sub>van</sub> (Supplementary Fig. 2a), we conclude that MopJ is a hitherto uncharacterized regulator of motility in

As inactivation of cell cycle regulators often results in pleiotropic defects in C. crescentus including a reduction of motility<sup>5</sup> caused by a change in the fraction of G1-phase cells in the population, we probed for such changes by fluorescenceactivated cell sorting (FACS) analysis of exponentially growing WT and  $\Delta mop I$  cells stained with the nucleic acid dye SYTOX Green (Fig. 1d). A lower G1:G2 cell ratio was observed in  $\Delta mop J$ cultures compared with WT, while the number of S-phase cells seemed unaffected. By contrast, stationary WT or  $\Delta mop J$  cultures contained few S-phase cells, confirming that the majority of cells are no longer replicating (Fig. 1e). Although stationary WT cells have an equal tendency to arrest either in G1- or G2-phase, stationary  $\Delta mop J$  cells are enriched in G2-phase over G1-phase. Consistent with these findings, differential interference contrast (DIC) microscopy and FACS revealed only slight increase in cell length caused by the  $\Delta mopJ$  mutation in exponential phase (Fig. 1d,f), whereas in stationary phase many  $\Delta mopJ$  cells are elongated and appear to divide aberrantly (Fig. 1e,g).

Taken together, our results reveal MopJ as a novel pleiotropic regulator of motility and of normal cell cycle progression in *C. crescentus*. Specifically, MopJ promotes the accumulation of G1-phase cells in exponential phase and of G1- and G2-phase cells in stationary phase.

MopJ acts on the CtrA regulon and is induced by (p)ppGpp. Knowing that MopJ impacts both motility and the abundance of G1 cells, we tested if MopJ affects the expression of the CtrA regulon, as this includes motility, division and G1-phase genes<sup>9,10,12</sup>. In addition, this master transcriptional regulator is a likely target of MopJ because it also regulates the initiation of chromosome replication<sup>4,12</sup>. To this end, we measured if β-galactosidase (LacZ) expression from CrA-regulated promoters is altered in exponential and stationary phase WT and  $\Delta mop J$  cells harbouring  $P_{pilA}$ -,  $P_{sciP}$ - or  $P_{fliM}$ -lacZ promoter probe plasmids (harbouring LacZ under the control of the promoter of the *pilA* pilin gene, the *sciP* repressor gene or the *fljM* minor flagellin gene, Fig. 2a,b) that are directly activated by CtrA<sup>9,10,44</sup> versus a *podJ* promoter reporter (Supplementary Fig. 3A) that is repressed by CtrA. Although only a modest reduction in LacZ activity (14-25%, depending on the CtrAactivated promoter, see Supplementary Table 1) was discernible in exponentially growing  $\Delta mop J$  cells compared with WT cells (Fig. 2a), this difference was accentuated in stationary phase (Fig. 2b and Supplementary Table 1) and even further magnified in a reporter that is indirectly activated by CtrA such as  $P_{fliK}$ -lacZ, in which the promoter of the major flagellin gene *fljK* drives LacZ



**Figure 2 | MopJ promotes CtrA accumulation and is regulated by (p)ppGpp.** (**a,b**) Promoter-probe assays of transcriptional reporters carrying the fljM, sciP, pilA or fljK promoter fused to a promoterless lacZ gene in WT and  $\Delta mopJ$  cells in exponential (**a**, Exp.) and stationary (**b**, Stat.) phase. The graphs show lacZ-encoded β-galactosidase activities measured in Miller Units. (**c**) Immunoblot showing the steady-state levels of the major flagellin FljK, the SciPnegative regulator and the PilA structural subunit of the pilus filament in WT and  $\Delta mopJ$  cells in Exp. and Stat. phase. The steady-state levels of the MreB actin are shown as a loading control (see Supplementary Fig. 4A). (**d**) Immunoblots showing CtrA and MopJ abundance in Exp. and Stat. phase  $\Delta mopJ$  versus WT cells. Steady-state levels of MreB are shown as loading control (see Supplementary Fig. 4B). (**e**) Accumulation of CtrA is increased in  $\Delta mopJ$  cells during Stat. phase either by inactivation of a proteolytic regulator encoded by cpdR ( $\Delta cpdR$ ::tet) or by appending an M2 (FLAG) epitope to the C-terminus of CtrA (ctrA::ctrA-M2). Immunoblot showing the steady-state levels of CtrA in WT,  $\Delta mopJ$ ,  $\Delta cpdR$ ::tet and  $\Delta mopJ$   $\Delta cpdR$ ::tet cells and the levels of CtrA-M2 in ctrA::ctrA-d2 and d3 and d4 and d5 are shown as loading control. (**f**,**g**) Promoter-probe assays of transcriptional reporters carrying the d5 and d6 are shown as loading control. (**f**,**g**) Promoter-probe plasmid in d6 and cells expressing the constitutive active form of d6. d6 RelA (RelA') and the catalytic mutant (RelA'-E335Q) fused to the FLAG tag in the presence of xylose. (**a,b,f,g,h**) Error bar (black bar) is shown as standard deviation (s.d., n = 3).

expression (Fig. 2a,b and Supplementary Table 1).  $P_{fljK}$  is a target of the FlbD activator whose expression in turn is directly and positively regulated by CtrA<sup>9,10</sup>.

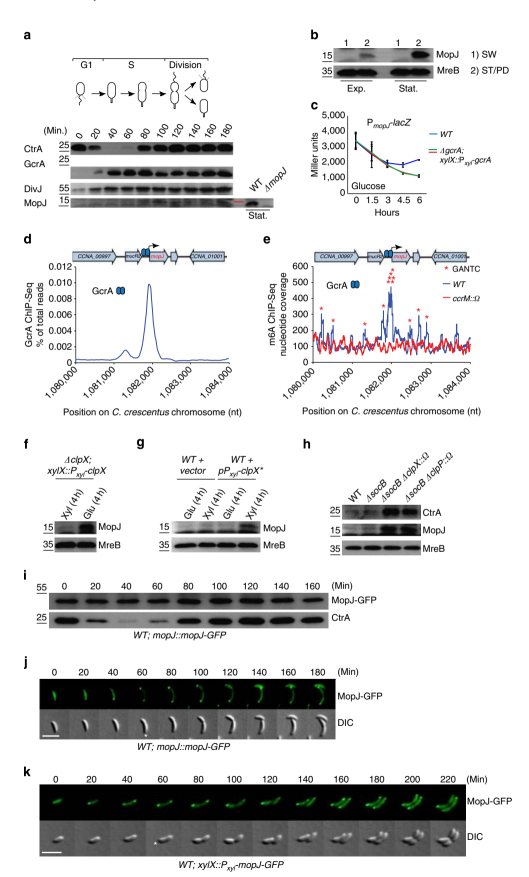
Immunoblotting using polyclonal antibodies to FljK, SciP and PilA confirmed the trends observed with the LacZ transcriptional reporters (Fig. 2c and Supplementary Fig. 4a). (Note that the PilA protein is absent from stationary phase WT cells for reasons that are currently unknown, but is likely operating at the post-transcriptional level (compare Fig. 2b,c and Supplementary Fig. 4a). Immunoblotting using antibodies to CtrA revealed that CtrA abundance is strongly dependent on MopJ in stationary phase (Fig. 2d and Supplementary Fig. 4b), consistent with the exacerbated effects on CtrA and its regulon in stationary  $\Delta mopJ$  cells compared with WT cells. Stabilizing CtrA either by masking the C-terminal recognition motifs for ClpXP in CtrA (using the ctrA:ctrA-M2 allele<sup>45</sup>) or by inactivating the gene encoding the CpdR proteolytic activator of CtrA<sup>16,20</sup> restores CtrA abundance

to near WT levels (Fig. 2e), indicating that CtrA proteolysis by the ClpXP protease contributes to downregulation of CtrA levels in stationary  $\Delta mopJ$  cells.

Next, we raised antibodies to MopJ to probe for commensurate changes in MopJ abundance in stationary phase versus exponential phase. In support of the stationary phase defects of  $\Delta mopJ$  cells described above, we observed that MopJ was barely detectable in lysates from exponential phase cells, but is abundant in lysates from stationary phase cells (Fig. 2d and see Fig. 3a and Supplementary Fig. 4b). As the alarmone (p)ppGpp accumulates in stationary phase when nutrients become exhausted in bacteria 25,46, we asked if the stationary phase induction of MopJ requires the (p)ppGpp-synthase/hydrolase SpoT 27,28,47. To resolve this question, we conducted  $\beta$ -galactosidase (LacZ) measurements in cells harbouring a transcriptional reporter, in which a promoterless lacZ gene is fused to the mopJ promoter ( $P_{mopJ}$ -lacZ). Accordingly, we measured  $P_{mopJ}$ -lacZ activity during

exponential and stationary phase in the presence or absence of SpoT (that is, in WT and  $\Delta spoT$  cells, respectively), observing a strong (50%) reduction of  $P_{mopl}$ -lacZ expression in stationary

cells lacking SpoT (Fig. 2f,g). To demonstrate that (p)ppGpp induction is sufficient for the  $P_{mopJ}$ -lacZ induction even in exponential phase cells, we induced (p)ppGpp synthesis in



exponential phase cells using the constitutively active (p)ppGpp synthase RelA' from  $E.\ coli^{47}$  and detected a commensurate increase in  $P_{mopJ}$ -lacZ activity (Fig. 2h). By contrast, no induction was observed upon induction of the catalytic mutant derivative RelA'-E335Q (Fig. 2h), demonstrating that (p)ppGpp is necessary and sufficient for induction of  $P_{mopJ}$ -lacZ.

Thus, MopJ acts on CtrA and its regulon, especially in stationary phase, and (p)ppGpp signalling by SpoT induces MopJ at the transcriptional level in stationary phase cells.

Coordinated synthesis of MopI and CtrA in S-phase. The functional relationship between MopI and CtrA is further reinforced by their concurrent accumulation during the cell cycle. coordinated by the S-phase regulator GcrA. Immunoblotting revealed that MopJ appears during the cell cycle coincident with or slightly ahead of CtrA and follows the accumulation of GcrA and DivJ (Fig. 3a and Supplementary Fig. 5a,b). Separation of exponential and stationary phase WT cells into swarmer (G1-phase) and stalked/pre-divisional (S-phase) cell fractions showed MopJ to be absent from the former and present in the latter (Fig. 3b and Supplementary Fig. 5c). The MopJ expression pattern matches that of the TipF flagellar regulator, an unstable protein that is expressed from a GcrA-activated promoter in early S-phase and is subsequently proteolysed in a manner that depends on the ClpXP protease<sup>48</sup>. Measuring LacZ activity in GcrA-depleted cells harbouring the  $P_{mopJ}$ -lacZ reporter, we observed a strong reduction after 6h of depletion relative to WT (Fig. 3c) or to GcrA-replete cells (Supplementary Fig. 3b), indicating that GcrA also acts positively on this promoter. In support of this, previous chromatin immunoprecipitation coupled to deep-sequencing experiments revealed that GcrA binds P<sub>mopJ</sub> (Fig. 3d) efficiently in vivo and that it harbours N6-methyladenosine (m6A, Fig. 3e) marks, a hallmark of a class of promoters that fire in S-phase and that require methylation for efficient activation and binding of GcrA in vitro and in vivo<sup>49,50</sup>.

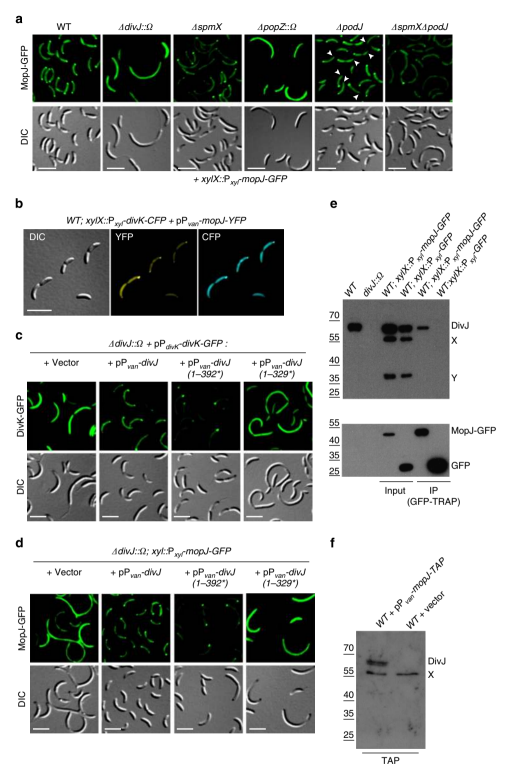
Interestingly, MopJ features two terminal alanine residues that resemble (class I) C-terminal recognition motifs of the ClpXP protease<sup>51</sup> and MopJ was previously found associated with the ClpP protease in pull-down assays<sup>52</sup>. We therefore investigated if ClpXP regulates MopJ abundance *in vivo*. To this end, we used immunoblotting to monitor the abundance of MopJ in cells from which the ClpX ATPase component of the ClpXP proteolytic machine had been depleted (Fig. 3f) or in cells intoxicated with a dominant negative version of ClpX (Fig. 3g)<sup>53</sup>. We found steady-state levels of MopJ to be elevated under these conditions. Moreover, the abundance of MopJ and CtrA, a known ClpXP substrate<sup>18</sup>, is elevated upon disruption of the *clpX* or the *clpP* gene in Δ*socB* mutant cells (Fig. 3h). (Note that inactivation of

clpX or clpP is lethal in WT cells, but no longer in  $\Delta socB$  cells, in which the gene encoding the SocB toxin has been deleted<sup>54</sup>). Finally, the levels of a MopJ variant harbouring GFP fused to the C-terminus (MopJ-GFP) expressed from  $P_{mopJ}$  no longer fluctuate during the cell cycle (Fig. 3I and Supplementary Fig. 5d) and confers motility to a comparable level as the untagged (WT) version (Supplementary Fig. 2b).

In sum, a combination of GcrA-controlled synthesis and ClpXP-dependent proteolysis restricts MopJ accumulation to S-phase and ensures that re-synthesis of CtrA and its reinforcement factor MopJ (see above) is linked, thus optimally reinforcing CtrA accumulation and function. Moreover, as MopJ is also induced in stationary phase by (p)ppGpp, an adaptive (nutritional) feed exists into the cell cycle.

Bipolar localization of MopJ requires the DivJ kinase. Superimposed on the temporal regulation of MopJ is its cell cyclecontrolled polar sequestration revealed in time-lapse fluorescence microscopic imaging of live mopJ::mopJ-GFP cells (Fig. 3j). Although diffuse fluorescence was observed in G1-phase (swarmer) cells, MopJ-GFP localizes first to the stalked pole during the transition into S-phase (stalked) cells and adopted a bipolar localization until MopJ-GFP dispersed from the opposite pole at the time of cell division (Fig. 3j). MopJ-GFP localized in a similar way when expressed constitutively from the xylose-inducible  $P_{xyl}$ promoter (Fig. 3k), compared with expression from its native promoter in S-phase, providing further evidence that the cell cycle-regulated localization occurs even when MopJ is expressed ectopically. Imaging of MopJ-GFP (Fig. 4a) or GFP-MopJ (Supplementary Fig. 6) expressed from the xylX locus in unsynchronized C. crescentus cultures also revealed monopolar and bipolar localization patterns. Consistent with the time-lapse imaging, MopJ-GFP is bipolar in pre-divisional cells, but is dispersed from the pole opposite the stalk in deeply constricted cells (Figs 3j,k and 4a), a characteristic also observed for the DivK RR that is phosphorylated by the DivJ kinase (Fig. 1a). Further reinforcing the parallels between MopJ and DivK (bi)polarity, we observed that MopJ-YFP and DivK-CFP co-localize in C. crescentus WT cells (Fig. 4b). This prompted us to explore if MopJ and DivK rely on a common localization mechanism. Indeed, in the absence of DivJ, DivK-GFP delocalized<sup>34</sup> (Fig. 4c) and MopJ-GFP as well (Fig. 4a,d and Supplementary Fig. 7a,b). Moreover, a truncated version (residues 1-392) of DivJ containing the HisKA domain, but lacking the ATPase (HATPase\_c) domain, is sufficient to direct MopJ-GFP and DivK-GFP to the stalked pole, but no longer to the opposite pole (Fig. 4c,d and Supplementary Fig. 7a,b). Thus, as was reported for DivK-GFP previously, MopJ-GFP requires DivJ kinase activity to be

**Figure 3 | Multi-layered spatiotemporal regulation of MopJ.** (a) Immunoblots showing steady-state levels of MopJ, GcrA, CtrA and DivJ in synchronized WT cells and MopJ in stationary (Stat.) phase (**a**, right, see Supplementary Fig. 5a,b). The red line marks the position of MopJ. (**b**) MopJ levels in exponential (Exp.) or stationary (Stat.) phase swarmer (SW) and stalked (ST)/predivisional (PD) cells detected using antibodies to MopJ (see Supplementary Fig. 5c). (**c**) LacZ measurements (Miller units) of  $P_{mopJ}$ -lacZ in WT (blue) and two independent  $\Delta gcrA xylX::P_{xyl}$ -gcrA (red and green) mutants, during 6 h (hours) of inhibition (glucose) of gcrA expression. Note the decline in activity upon dilution of overnight cells is due to the prior induction in stationary phase by (p)ppGpp (see Fig. 2f-h and Supplementary Fig. 3b). Error bar (black bar) is shown as s.d. (n = 3). (**d**) Occupancy of GcrA (blue ovals) at the mopJ locus in WT cells as determined by chromatin immunoprecipitation coupled to deep-sequencing (ChIP-Seq) using polyclonal antibodies to  $GcrA^{50}$ . Occupancy is shown as percentage of total reads in the precipitated sample as a function of the chromosome coordinates. (**e**) Trace of N6-methyladenosine (m6A) marked DNA at mopJ locus in WT cells (blue trace) and  $\Delta ccrM$ :Ω (red trace) from ChIP-Seq experiment performed with antibodies to mopJ occupancy is shown as percentage of total nucleotide coverage in sequencing as a function of the chromosome coordinates at the mopJ locus. Stars show the positions of 5'-GANTC-3' sequences that are methylated by the CcrM methyltransferase at the N6 position of adenine. (**f-h**) Immunoblots showing the steady-state levels of MopJ in  $\Delta clpX \times ylX::Pxyl-clpX$  cells with (Xyl, xylose) or without (Glu, glucose) induction of  $P_{xyl}$  for 4 h (**f**), upon induction of the dominant negative form ClpX\* from  $P_{xyl}$  for 4 h (**g**), and in WT,  $\Delta socB$  and  $\Delta socB\Delta clpP::\Omega$  double mutants (**h**) along with CtrA and MreB (control). (**i**) Immunob



**Figure 4 | Polar localization of MopJ requires the DivJ kinase and the PopZ polar scaffold.** (a) Fluorescence and DIC images of live *C. crescentus WT*,  $\Delta divJ:\Omega$ ,  $\Delta spmX$ ,  $\Delta popZ:\Omega$ ,  $\Delta popJ$  or  $\Delta podJ\Delta spmX$  cells expressing MopJ-GFP under the control of xylose promoter ( $P_{xyl}$ ) at the xylX locus. White arrowheads point to the flagellated pole. (b) Co-localization of DivK-CFP expressed from  $P_{xyl}$  at the xylX locus in WT cells and MopJ-YFP expressed from  $P_{van}$  on pMT374. The corresponding DIC image is also shown. (c) Fluorescence and DIC images of  $\Delta divJ:\Omega$  cells expressing DivK-GFP fusion from pMR10-divK-GFP and various versions of DivJ, including full-length DivJ, DivJ-392 (short version of divJ encoding a truncated protein without kinase and dimerization domains) under the control of the vanillate promoter ( $P_{van}$ ) from pMT335. The control harbouring the empty vector is also shown. (d) Fluorescence and DIC images of  $\Delta divJ:\Omega$  cells expressing MopJ-GFP under the control of  $P_{xyl}$  and the different forms of DivJ from pMT335 as described in c. (e) Co-immunoprecipitation (IP) of DivJ with MopJ-GFP from a GFP-TRAP affinity matrix (ChromoTek GmbH). Precipitated samples were probed for the presence of DivJ and GFP by immunoblotting using antibodies to DivJ (upper) and GFP (lower). Cell lysates used as input are also shown. (f) Tandem-affinity-purification (TAP) of MopJ-TAP from  $P_{van}$  on pMT335 followed by immunoblotting of the samples for the presence of DivJ using polyclonal antibodies to DivJ. In e and f, X and Y denote nonspecific proteins that react with the antiserum or degradation products.

sequestered to the pole opposite the stalk, but its sequestration to the stalked pole occurs independently of kinase activity<sup>17</sup> (Fig. 4c,d). A further truncated version of DivJ (residues 1–329) no longer supported polar localization of MopJ-GFP and DivK-GFP (Fig. 4c,d and Supplementary Fig. 7a,b), indicating that the dimerization/phosphor-acceptor (HisKA) domain and/or a determinant encoded from residues 329 to 392 is critical to recruit MopJ and DivK to the stalked pole. By contrast, expression of full-length DivJ from a comparable plasmid reinstated bipolarity upon MopJ-GFP and DivK-GFP (Fig. 4c,d and Supplementary Fig. 7a,b). As both derivatives of DivJ can still localize to the stalked pole (albeit 50% less efficient for the shorter version, Supplementary Fig. 8), this experiment confirms that residues 329-392 of DivJ encode for the localization determinants of MopJ and DivK. We conclude that DivJ integrates the bipolar localization of two dissimilar effectors, MopJ and DivK, to the stalked pole and, in early pre-divisional cells, also to the opposite pole.

In support of the notion that a physical interaction between MopJ and DivJ underlies the localization of MopJ to the stalked pole, co-immunoprecipitation experiments revealed that DivJ is pulled-down with MopJ-GFP from cellular lysates of a WT strain expressing MopJ-GFP from  $P_{xyl}$  (Fig. 4e). By contrast, DivJ was not detectable in comparable pull-downs from lysates of the WT strain expressing GFP from P<sub>xyl</sub> (Fig. 4e). Furthermore, DivJ was also recovered with MopJ in tandem-affinity purification experiments<sup>55</sup> conducted with lysates of WT cells expressing MopJ-TAP from P<sub>van</sub> (Fig. 4f). Having shown that MopJ resides in a complex with DivJ, we then tested whether overexpression of MopJ can affect the DivJ-regulated subcellular distribution of the other DivJ-interacting effector DivK<sup>17,34</sup> (Fig. 5a,b). Live-cell imaging of DivK-GFP in cells overexpressing MopJ from Pvan revealed an enhancement of DivK-GFP polar localization and a concomitant reduction of the cytoplasmic fluorescence that correlated with the strong induction of MopJ upon the addition of vanillate (Fig. 5a,b), without affecting DivK~P levels noticeably by in vivo phosphorylation analysis (Fig. 5c). However, we found that  $\Delta mop J$  cells are sensitized towards elevated DivK levels from a plasmid expressing DivK under the control of  $P_{xyl}$ , suggesting that extra DivK<sup>17</sup> further curbs the CtrA pathway in  $\Delta mop J$  cells (in which activation of the CtrA regulon is already disturbed, see above) to inhibit growth (Fig. 5d) and division (Fig. 5e) more than in WT cells. Taken together, these experiments show that MopJ binds DivJ and depends on it for its own polar localization, while promoting the polar sequestration of DivK and attenuating its activity.

Examining possible effects on downstream components, we found that MopJ overexpression not only alters the subcellular distribution of DivK, but also that of its interaction partner, the tyrosine kinase DivL<sup>24</sup> that localizes primarily to the pole opposite the stalk<sup>15,22,56</sup> (Fig. 1a). Overexpression of MopJ caused a near fivefold increase in the fraction of bipolar DivL-GFP (expressed from the native promoter at the *divL* locus in lieu of endogenous DivL; Fig. 5f and Supplementary Fig. 9). Conversely, in cells lacking MopJ, DivL-GFP was largely delocalized (Fig. 5g) and a divL::Tn5 mutation (in which Tn5 is inserted upstream of the region encoding the HisKA domain) did not impact MopJ-GFP localization (Supplementary Fig. 10). The previous observation that DivL is delocalized in the absence of DivJ<sup>56</sup> is consistent with our findings, as we showed above that DivJ is also required for the polarization of MopJ-GFP. Intriguingly, DivL localization is thought to be dependent on the onset of DNA replication<sup>57</sup>. The result that MopJ is required for DivL localization and MopJ expression requires GcrA is noteworthy, as it could provide an explanation for the finding that DivL polarization (and the concomitant re-accumulation of

CtrA that this event regulates) requires S-phase entry<sup>57</sup>. GcrA expression and S-phase entry are known to be dependent on the replication initiator DnaA<sup>58</sup>.

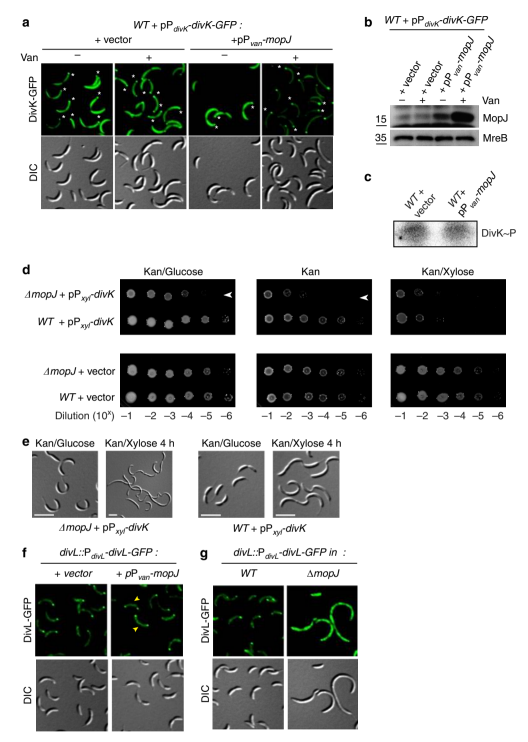
In sum, MopJ is both target (via DivJ) and effector (via DivK and DivL) of the *Caulobacter* spatiotemporal cell cycle network and is thus perfectly positioned to reinforce CtrA with the synthesis of both proteins in S-phase and is genetically linked as expression of both proteins is directly induced by GcrA.

A second localization pathway controls polarization of MopJ. Although DivJ is necessary for the bipolar localization of MopJ, two findings indicate that it is not sufficient for bipolarity. First, MopJ is localized to the pole opposite the stalk (Fig. 4a), where no focus of DivJ is observed<sup>32</sup>. Second, MopJ is still polarized when DivJ is delocalized by inactivation of its localization factor SpmX (that is, in  $\Delta spmX$  mutant cells, Fig. 4a and Supplementary Fig. 11)<sup>3</sup>. Taken together, this indicates that other factors must promote polarization of MopJ. As it is also conceivable that DivJ simply targets MopJ to the membrane or fastens it there, we also explored whether the bipolar scaffolding protein PopZ that forms adhesive patches at both cell poles<sup>26,36,38–40</sup> might keep MopJ-GFP polarized. Live-cell imaging revealed that MopJ-GFP is dispersed in  $\Delta popZ::\Omega$  mutant cells (Fig. 4a). As PopZ has also been likened to a molecular plug that prevents diffusion away from the cell pole, we predicted that an additional factor promotes MopJ-GFP localization to the pole opposite the stalk. The PodJ polarity protein is known to attract regulatory proteins of CtrA to the pole opposite the stalk<sup>36,37,59,60</sup>. We observed MopJ-GFP to be monopolar in  $\Delta podJ$  mutant cells, unable to assemble into foci at the pole opposite the stalk regardless of whether SpmX was present or not (Fig. 4a and Supplementary Fig. 11). This dependence of MopJ-GFP on PodJ for localization to the flagellated pole not only supports the existence of distinct recruitment mechanisms of MopJ-GFP for each cell pole in WT cells, but also provides an explanation of why DivL is no longer localized to the flagellated pole in  $\Delta podJ$  cells<sup>59</sup>. It also points to a possible explanation of how DivJ might influence MopJ localization to the pole opposite the stalk. As the expression of the gene encoding the PerP periplasmic protease that cleaves PodJ is strongly and directly upregulated by CtrA in ΔdivJ::Ω mutant cells<sup>9,10,61</sup>, we hypothesize that DivJ promotes the localization of MopJ at the pole opposite the stalk indirectly through PodJ (or another DivJ-dependent protein).

#### Discussion

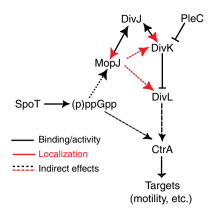
A single-domain PAS protein, MopJ, is not only regulated by cell cycle cues, but also integrates nutritional and growth phase regulatory signals to exert topological control over conserved components of a bacterial cell cycle network (Fig. 6). The identification of MopJ enabled us to illuminate two important and hitherto enigmatic cell cycle events. The first conundrum was how the master regulator CtrA can re-accumulate in S-phase<sup>4</sup>, a time when DivK is still phosphorylated and signals the removal of CtrA<sup>30,34</sup>. Unless the proteolytic cascade is attenuated and/or the rate of synthesis is potentiated such that it exceeds proteolysis, CtrA accumulation should be suppressed or curbed. We found that the PAS-domain protein MopJ indeed enhances CtrA accumulation and acts on DivK and its target DivL15. MopJ re-accumulation is temporally linked with that of CtrA through the S-phase-specific transcriptional activator GcrA that binds the *ctrAP1* and the *mopJ* promoter<sup>8,49,50</sup>, thus reinforcing CtrA re-synthesis with the coordinated synthesis of its enhancing factor MopJ in S-phase.

By discovering that MopJ acts at the same subcellular site as the CtrA His-Asp phosphorelay components DivJ, DivK and DivL,



**Figure 5 | MopJ affects DivK localization and function.** (a) Fluorescence and DIC imaging of WT cells harbouring pMR10-divK-GFP and pMT335- $P_{van-mopJ}$  with (+) or without (-) induction (50 μM of vanillate) of MopJ from  $P_{van}$  on pMT335. The controls harbouring pMR10-divK-GFP and the empty vector (pMT335) are also shown. The position of stalks are indicated by a white asterisk. (b) Immunoblot showing the steady-state levels of MopJ in strains used in **a**. Steady-state levels of MreB are shown as a loading control. (c) In vivo phosphorylation of cultures with  $^{32}P$  followed by immunoprecipitation reveals DivK  $\sim$  P levels in WT cells harbouring pMT335 or pMT335-mopJ ( $P_{van}$ -mopJ) after with vanillate (50 μM). (d) Effect of  $\Delta mopJ$  mutation on viability in the presence of extra DivK. Efficiency of plating assays of WT and  $\Delta mopJ$  cells harbouring the empty vector (pMR10) or the  $P_{xyl}$ -divK expression plasmid (pMR10- $P_{xyl}$ -divK). Serial tenfold dilutions of cells plated on PYE agar with kanamycin and glucose (kan + gluc, glucose represses  $P_{xyl}$ ), kanamycin (kan, background expression from  $P_{xyl}$ ) and kanamycin and xylose (kan + xyl, xylose induces  $P_{xyl}$ ) agar. (e) DIC (differential interference contrast) imaging of cells described in d in glucose or 4 h after induction with xylose (0.3%). The medium was supplemented with kanamycin (5 μg ml  $^{-1}$ ). (f) Effect of MopJ overexpression on DivL-GFP localization. The empty vector (pMT335) or the  $P_{van}$ -mopJ plasmid (pMT335-mopJ) was transformed into  $divL::P_{divL}$ -divL-GFP cells and the resulting cells imaged by DIC and fluorescence microscopy 4 h after induction with vanillate (50 μM). Yellow arrowheads point to cells with bipolar DivL-GFP. (g) Effect of mopJ deletion on DivL-GFP localization. The  $divL::P_{divL}$ -divL-GFP allele was transduced in WT and  $\Delta mopJ$  cells and the resulting cells imaged by DIC and fluorescence microscopy.

(see Discussion).



**Figure 6 | Proposed action of MopJ on the** *Caulobacter* **cell circuitry.** Summary scheme depicting the interactions of MopJ with the cell circuitry as determined in this work. Effects on protein localization are shown in red, whereas interactions shown in black designate direct interactions that generally result in activation (arrow, DivJ-DivK) or inhibition (T-bar, PleC-DivK, DivK-DivL). The scheme also shows the effects of the alarmone (p)ppGpp in inducing MopJ and the maintenance of CtrA<sup>25,26,28,47</sup>

we were able to identify MopJ as an important missing link connecting DivL polarization to DivJ (Fig. 6) and likely to the onset of DNA replication. MopJ polarization requires the polar organizer PopZ and the DivJ kinase. With the previous findings that polar localization of DivJ is itself dependent on PopZ assembly at polar sites<sup>38,40</sup>, our work extends this spatiotemporal regulatory network by showing that MopJ associates with DivJ (Fig. 4), enhances the sequestration of DivK to the cell poles (attenuating DivK function, Fig. 5), governs subcellular localization of DivL and facilitates CtrA accumulation and expression of its target promoters (Fig. 2).

The stationary phase and (p)ppGpp-based induction of MopJ unveils a concerted feed-forward mechanism underlying signal integration during bacterial cell cycle progression, acting on an essential master transcriptional regulator in different phases of growth (Fig. 6). Why MopJ abundance increases in stationary phase remains to be determined, but our experiments suggest that this inductions occurs at the transcriptional level as the activity of the mopJ promoter is elevated in stationary phase and is inducible in nutrient replete conditions when (p)ppGpp levels are raised artificially in exponential phase (Fig. 2). As (p)ppGpp signals nutrient starvation by globally reprogramming RNA polymerase and is induced in stationary phase in many bacteria<sup>46</sup>, the induction of MopJ is likely mediated at the level of promoter activity. It is also possible that MopJ synthesis is further reinforced at the level of protein stability by inhibiting ClpXPmediated proteolysis (Fig. 3). It is not obvious why MopJ should be induced in stationary phase when nutrients are exhausted to enhance the cell cycle regulator CtrA. However, it is known that CtrA levels are maintained in the presence of (p)ppGpp<sup>26,28,47</sup>, and our results raise the possibility that this effect is (at least partially) mediated by (induced) MopJ. In light of the fact that the number of cells undergoing DNA replication is strongly diminished in stationary phase, with cells accumulating predominantly in G1 and G2 phase (Fig. 1), along with the fact that CtrA acts on replication control<sup>4,12</sup>, we suspect that CtrA is responsible or at least contributes to this replication arrest. In support of this, cells no longer exhibit a normal stationary phase arrest in the absence of MopJ (Fig. 1), and the abundance of CtrA is strongly reduced under these conditions (Fig. 2). Such stationary-phase-induced replication arrest may be further accentuated by other (p)ppGpp-dependent mechanisms, such as

the reduction of the replication initiator DnaA, an unstable protein<sup>62,63</sup>, when (p)ppGpp levels are high<sup>26,28,47</sup>. It is also noteworthy that in contrast to this specific cell cycle control mechanism on the level of a master regulator, (p)ppGpp induces a general growth arrest and a state of antibiotic tolerance (known as persistence) in *E. coli* and likely other bacteria through a shutdown of macromolecular synthesis by type II toxin–antitoxin systems that are activated indirectly upon accumulation of (p)ppGpp<sup>64</sup>.

(p)ppGpp-induced replication arrest mechanism(s) are also operational in other  $\alpha$ -Proteobacteria<sup>25</sup>, as a nutritional downshift induces a cell cycle arrest in *S. meliloti*<sup>65</sup>. Although CtrA is not known to bind the origin of replication in *Sinorhizobium fredii* directly<sup>10</sup>, CtrA may inhibit replication initiation indirectly as was recently also suggested to be the case for *Caulobacter*<sup>13</sup> and reinforcement of CtrA by induced MopJ presents an appealing and conceivable possibility. This is mechanistically distinct from the (p)ppGpp-based replication arrest at the level of DNA primase described for the Gram-positive soil bacterium *Bacillus subtilis*<sup>66</sup>, a member of the Firmicutes, clearly showing that evolution has found elegant solutions to the same problem.

#### Methods

**Growth conditions.** Caulobacter crescentus NA1000 (ref. 43) and derivatives were cultivated at 30 °C in peptone yeast extract (PYE)-rich medium or in M2 minimal salts plus 0.2% glucose (M2G) supplemented by 0.4% liquid PYE<sup>5</sup>. E. coli S17-1 and EC100D (Epicentre Technologies) were cultivated at 37 °C in Luria Broth (LB)-rich medium. 1.5% agar was added into M2G or PYE plates and motility was assayed on PYE plates containing 0.3% agar. Antibiotic concentrations used for C. crescentus include kanamycin (solid: 20  $\mu$ g ml $^{-1}$ ; liquid: 5  $\mu$ g ml $^{-1}$ ), tetracycline (1  $\mu$ g ml $^{-1}$ ), spectinomycin (liquid: 25  $\mu$ g ml $^{-1}$ ), spectinomycin/streptomycin (solid: 30 and 5  $\mu$ g ml $^{-1}$ , respectively), gentamycin (1  $\mu$ g ml $^{-1}$ ) and nalidixic acid (20  $\mu$ g ml $^{-1}$ ). When needed, D-xylose or sucrose was added at 0.3% final concentration, glucose at 0.2% final concentration and vanillate 500 or 50  $\mu$ M final concentration. For the experiments in stationary phase in PYE, cultures with an OD<sub>660nm</sub> > 1.4 were used.

Swarmer cell isolation, electroporation, biparental mating and bacteriophage  $\varphi$ Cr30-mediated generalized transduction were performed as described in ref. 5. Briefly, swarmer cells were isolated by Ludox or Percoll density-gradient centrifugation at 4 °C, followed by three washes and final re-suspension in prewarmed (30 °C) M2G. Electroporation was done using 1 mm gap electroporation cuvettes at 1.5 kV in an Eppendorf 2510 electroporator and 6 ml exponential phase cells that had been washed three times in sterile water. Biparental matings were done using exponential phase *E. coli* S17-1 donor cells and *Caulobacter* recipient cells washed in PYE and mixed at 1:10 ratio on a PYE plate. After 4–5 h of incubation at 30 °C, the mixture of cells was plated on PYE harbouring nalidixic acid (to counter select *E. coli*) and the antibiotic that the conjugated plasmid confers resistance to. Generalized transductions were done by mixing 50 µl ultraviolet-inactivated transducing lysate with 500 µl exponential phase recipient cells, incubation for 2 h, followed by plating on PYE containing antibiotic to elect for the transduced DNA.

**Bacterial strains, plasmids and oligonucleotides.** Bacterial strains, plasmids and oligonucleotides used in this study are listed and described in Supplementary Tables 3–5, respectively.

**Co-immunoprecipitation.** When the culture (50 ml) reached  $OD_{660nm} = 0.4$ –0.6 in the presence of xylose, cells were harvested by centrifugation at 6,000g for 10 min. The pellet was then washed in 10 ml of dilution/wash buffer (10 mM Tris-HCl (pH 7.5); 150 mM NaCl; 0.5 mM EDTA) and lysed for 15 min at room temperature in 1 ml of lysis buffer (dilution/wash buffer + 0.5% NP40, 10 mM MgCl<sub>2</sub>, two protease inhibitor tablets (for 50 ml of buffer; Complete EDTA-free, Roche), 1 × Ready-Lyse lysozyme (Epicentre), 50 U of DNase I )Roche)). Cellular debris was removed by centrifugation at 7,000g for 15 min at 4 °C. The supernatant was incubated for 1 h (or overnight) at 4 °C with GFP-Trap\_A beads (ChromoTek GmbH) previously washed three times with 500  $\mu$ l of dilution/wash buffer. The sample was then centrifuged at 2,500g for 2 min at 4 °C and the supernatant was removed. The beads were washed three times with 500  $\mu$ l of dilution/wash buffer and finally resuspended in 2 × SDS sample buffer (50 mM Tris-HCl (pH 6.8), 2% SDS, 10% glycerol, 1%  $\beta$ -mercaptoethanol, 12.5 mM EDTA, 0.02% Bromophenol Blue), heated to 95 °C for 10 min and stored at -20 °C.

**Tandem affinity purification (TAP).** The TAP procedure was based on that described by ref. 55. Briefly, when the culture (11) reached  ${\rm OD_{660nm}}\,{=}\,0.4{-}0.6$  in the presence of  $50\,\mu{\rm M}$  vanillate, cells were harvested by centrifugation at 6,000g for  $10\,{\rm min}$ . The pellet was then washed in 50 ml of buffer I (50 mM sodium phosphate at

pH 7.4, 50 mM NaCl, 1 mM EDTA) and lysed for 15 min at room temperature in 10 ml of buffer II (buffer I + 0.5% n-dodecyl-β-D-maltoside, 10 mM MgCl<sub>2</sub>, two protease inhibitor tablets (for 50 ml of buffer II; Complete EDTA-free, Roche), 1 × Ready-Lyse lysozyme (Epicentre), 500 U of DNase I (Roche)). Cellular debris was removed by centrifugation at 7,000g for 20 min at 4 °C. The supernatant was incubated for 2 h at 4 °C with IgG Sepharose beads (GE Healthcare Biosciences) that had been washed once with IPP150 buffer (10 mM Tris-HCl at pH 8, 150 mM NaCl, 0.1% NP40). After incubation, the beads were washed at 4 °C three times with 10 ml of IPP150 buffer and once with 10 ml of TEV cleavage buffer (10 mM Tris-HCl at pH 8, 150 mM NaCl, 0.1% NP40, 0.5 mM EDTA, 1 mM DTT). The beads were then incubated overnight at 4 °C with 1 ml of TEV solution (TEV cleavage buffer with 100 U of TEV protease per millilitre (Promega)) to release the tagged complex. CaCl<sub>2</sub> (3 μM) was then added to the solution. The sample with 3 ml of calmodulin-binding buffer (10 mM β-mercaptoethanol, 10 mM Tris-HCl at pH 8, 150 mM NaCl, 1 mM magnesium acetate, 1 mM imidazole, 2 mM CaCl2, 0.1% NP40) was incubated for 1 h at 4°C with calmodulin beads (GE Healthcare Biosciences) that previously had been washed once with calmodulin-binding buffer. After incubation, the beads were washed three times with 10 ml of calmodulin-binding buffer and eluted five times with 200 µl IPP150 calmodulin elution buffer (calmodulin-binding buffer substituted with 2 mM EGTA instead of CaCl2). The eluates were then concentrated using Amicon Ultra-4 spin columns (Ambion).

**β-Galactosidase assays.** β-Galactosidase assays were performed at 30 °C as described previously³. 50 μl of cells grown in PYE at OD<sub>660nm</sub> = 0.1–0.6 were lysed with chloroform and mixed with 750 μl of Z buffer (60 mM Na<sub>2</sub>HPO<sub>4</sub>, 40 mM NaH<sub>2</sub>PO<sub>4</sub>, 10 mM KCl and 1 mM MgSO<sub>4</sub> heptahydrate). 200 μl of ONPG (4 mg ml $^{-1}$  o-nitrophenyl-β-D-galactopyranoside in 0.1 M KPO<sub>4</sub> pH 7.0) was added and the reaction timed. When a medium-yellow colour developed the reaction was stopped with 400 μl of 1 M Na<sub>2</sub>CO<sub>3</sub>. The OD<sub>420nm</sub> of the supernatant was determined and the units were calculated with the equation:  $U = (\mathrm{OD}_{420nm} \times 1000)/(\mathrm{OD}_{660nm} \times \text{time}$  (in min) × volume of culture (in ml)). For the stationary phase experiments, stationary phase cultures were diluted in PYE to obtain an OD<sub>660nm</sub> = 0.1–0.6 and quickly lysed with chloroform. Following steps were performed as above. For GcrA depletion experiments in PYE using strain NA1000 Δ*gcrA*::Ω *xylX*::P*xyl-gcrA*, PYE supplemented with 0.3% xylose overnight cultures were harvested and washed three times with PYE, and then restarted in appropriate PYE supplemented with 0.3% xylose or PYE supplemented with 0.2% glucose medium for 0, 1.5, 3, 4.5 and 6 h at 30 °C. Experimental values represent the averages of three independent experiments.

MopJ purification and production of antibodies. MopJ-short protein, lacking the first N-terminal 45 residues, was expressed from pET28a in *E. coli* Rosetta (DE3)/pLysS (Novagen) and purified under native conditions using Ni<sup>2+</sup> chelate chromatography. A 5-ml overnight culture was diluted into 11 of pre-warmed LB. When cells reached OD<sub>660nm</sub> = 0.3–0.4, 1 mM isopropyl-β-D-thiogalactoside was added to the culture and growth continued. After 3 h, cells were pelleted and resuspended in 25 ml of lysis buffer (10 mM Tris HCl (pH 8), 0.1 M NaCl, 1.0 mM b-mercaptoethanol, 5% glycerol, 0.5 mM imidazole Triton X-100 0.02%). Cells were sonicated (Sonifier Cell Disruptor *B-30*; Branson Sonic Power. Co.) on ice using 12 bursts of 20 s at output level 5.5. After centrifugation at 4,300g for 20 min, the supernatant was loaded onto a column containing 5 ml of Ni-NTA agarose resin pre-equilibrated with lysis buffer. Column was rinsed with lysis buffer, 400 mM NaCl and 10 mM imidazole, both prepared in lysis buffer. Fractions were collected (in 300 mM Imidazole buffer, prepared in lysis buffer) and used to immunize New Zealand white rabbits (Josman LLC).

**Immunoblot analysis.** Pelleted cells were re-suspended in  $1 \times SDS$  sample buffer (50 mM Tris-HCl (pH 6.8), 2% SDS, 10% glycerol, 1% β-mercaptoethanol, 12.5 mM EDTA, 0.02% Bromophenol Blue), heated to 95 °C for 10 min and stored at - 20 °C. Protein samples were separated by SDS-polyacrylamide gel electrophoresis and blotted on polyvinylidenfluoride membranes (Merck Millipore). Membranes were blocked for 1 h with Tris-buffered saline (TBS), 0.05% Tween-20 and 5% dry milk and then incubated for an additional 1 h with the primary antibodies diluted in TBS, 0.05% Tween-20, 5% dry milk. The different polyclonal antisera to PilA (1:5,000 dilution), FljK (1:10,000), CtrA (1:10,000) and SciP (1:2,000) were used as described before<sup>10</sup>. The other antibodies were used at following dilutions: anti-GcrA (1:10,000)<sup>8</sup>, anti-MreB (1:20,000)<sup>67</sup>, anti-DivJ (1:10,000)<sup>32</sup> and anti-MopJ (1:4,000). Commercial monoclonal (Roche, CH) and polyclonal antibodies to GFP were used at 1:5,000 and 1:10,000 dilutions, respectively. The membranes were washed four times for 5 min in PBS and incubated for 1 h with the secondary antibody (HRP-conjugated donkey anti-rabbit antibody, Jackson ImmunoResearch) diluted 1:20,000 in TBS, 0.05% Tween-20 and 5% dry milk. The membranes were finally washed again four times for 5 min in PBS and revealed with Immobilon Western Blotting Chemoluminescence HRP substrate (Merck Millipore) and Super RX-film (Fujifilm).

**Microscopy**. PYE or M2G cultivated cells in exponential growth phase were immobilized using a thin layer of 1% agarose. For time-lapse experiments, synchronized cells were immobilized using a thin layer of 1% agarose in M2G

supplemented with 0.4% PYE. Fluorescence and contrast microscopy images were taken with an Alpha Plan-Apochromatic  $\times$  100/1.46 DIC(UV) VIS-IR oil objective on an Axio Imager M2 microscope (Zeiss) with acquisition at 535 nm (enhanced green fluorescent protein), 580 nm (yellow fluorescent protein (YFP)) and 480 nm (cyan fluorescent protein (CFP); Visitron Systems GmbH) and a Photometrics Evolve camera (Photometrics) controlled through Metamorph V7.5 (Universal Imaging). Images were processed using Metamorph V7.5.

Fluorescence-activated cell sorting. FACS experiments were performed as described previously<sup>50</sup>. Cells in exponential growth phase ( $OD_{660nm} = 0.3-0.6$ ) or in stationary phase (diluted to obtain an OD<sub>660nm</sub> = 0.3-0.6), cultivated in M2G, were fixed in ice-cold 70% ethanol solution. Fixed cells were re-suspended in FACS staining buffer, pH 7.2 (10 mM Tris-HCl, 1 mM EDTA, 50 mM NaCitrate, 0.01% Triton X-100) and then treated with RNase A (Roche) at 0.1 mg ml<sup>-1</sup> for 30 min at room temperature. Cells were stained in FACS staining buffer containing 0.5 μM of SYTOX Green nucleic acid stain solution (Invitrogen) and then analysed using a BD Accuri C6 flow cytometer instrument (BD Biosciences). Flow cytometry data were acquired and analysed using the CFlow Plus V1.0.264.15 software (Accuri Cytometers Inc.). 20,000 cells were analysed from each biological sample. The forward scattering (FSC-A) and Green fluorescence (FL1-A) parameters were used to estimate cell sizes and cell chromosome contents, respectively. Experimental values represent the averages of three independent experiments. Relative chromosome number was directly estimated from the FL1-A value of NA1000 cells treated with 20 µg ml<sup>-1</sup> Rifampicin for 3 h at 30 °C, done in ref. 50. Rifampicin treatment of cells blocks the initiation of chromosomal replication, but allows ongoing rounds of replication to finish.

In vivo phosphorylation-immunoprecipitation experiments. In vivo phosphorylation measurements by  $^{32}\mathrm{P}$  labelling of WT cultures harbouring pMT335 or pMT335-mopJ after induction with vanillate (50 µM), followed by immunoprecipitation with antibodies to DivK as described in the study by Radhakrishnan et al.^3. Briefly, a freshly grown colony was picked from a PYE plate, washed with M5G medium lacking phosphate and was cultivated overnight in M5G with 0.05 mM phosphate to an optical density of 0.3 at 660 nm. One millilitre of culture was labelled for 4 min at 28 °C using 30 µCi of  $\gamma$ -[32P]ATP. Upon lysis, proteins were immunoprecipitated with 3 µl of anti-DivK antiserum and Protein A agarose (Roche, CH) and the precipitates were resolved by SDS-polyacrylamide gel electrophoresis and radiolabelled DivK was quantified.

#### Strain constructions

 $NA1000~\Delta mopJ$ . Deletions were introduced using SacB-based counterselection using 3% sucrose. Briefly, pNPTS138- $\Delta mopJ$ -KO was first introduced into NA1000 (WT) by intergeneric conjugation and then plated on PYE harbouring kanamycin (to select for recombinants) and nalidixic acid to counter select E.~coli donor cells<sup>5</sup>. A single homologous recombination event at the  $CCNA\_00999$  locus of kanamycin-resistant colonies was verified by PCR. The resulting strain was grown to stationary phase in PYE medium lacking kanamycin. Cells were then plated on PYE supplemented with 3% sucrose and incubated at 30 °C. Single colonies were picked and transferred in parallel onto plain PYE plates and PYE plates containing kanamycin. Kanamycin-sensitive cells, which had lost the integrated plasmid due to a second recombination event, leaving a deleted version of mopJ behind ( $\Delta mopJ$ ), were then identified for disruption of the mopJ locus by PCR.

NA1000  $\Delta spm X \Delta pod J$ . pNPTS138- $\Delta spm X$ -KO<sup>3</sup> was first introduced into NA1000  $\Delta pod J^{36}$  by intergeneric conjugation and then plated on PYE harbouring kanamycin (to select for recombinants) and nalidixic acid to counter select *E. coli* donor cells. A single homologous recombination event at the  $CCNA\_02255$  locus of kanamycin-resistant colonies was verified by PCR. The resulting strain was treated as above.

NA1000 mopJ::himar1. mopJ::himar1 allele (himar1 insertion at nt 1082252,  ${\rm Kan}^{\rm R}$ ) was transduced into NA1000 from NS61 using  $\varphi$ Cr30-mediated generalized transduction.

Strains harbouring pMT335 or pMT335-derived plasmids. pMT335 or pMT335-mopJ was introduced by electroporation and then plated on PYE plates harbouring gentamycin.

 $NA1000~\Delta socB~\Delta clpP::\Omega~$  and  $\Delta socB~\Delta clpX::\Omega$ . The  $\Delta clpP::\Omega~$  (Spc<sup>R</sup>) or  $\Delta clpX::\Omega~$  (Spc<sup>R</sup>)<sup>19</sup> allele was introduced into NA1000  $\Delta socB$  as described in ref. 54 by  $\phi$ Cr30-mediated transduction followed by selection on plates harbouring spectinomycin.

 $NA1000 \ xylX::P_{xyl}$ - $clpX^*$ . pMO88 (ref. 68) was introduced into NA1000 by electroporation and then plated on PYE harbouring tetracycline. The resulting strain was grown to stationary phase in PYE medium containing tetracycline.

 $NA1000~ctrA::ctrA-M2~and~\Delta mopJ~ctrA::ctrA-M2$ . The ctrA::ctrA-M2 allele<sup>45</sup> was introduced into NA1000 or  $\Delta mopJ$  by  $\varphi$ Cr30-mediated transduction on PYE plates harbouring kanamycin.

 $NA1000~\Delta cpdR$ :: $tetR~and~\Delta mopJ~\Delta cpdR$ ::tetR. The  $\Delta cpdR$ ::tet (Tet<sup>R</sup>) allele was transduced from  $\Delta cpdR$ :: tet<sup>16</sup> into NA1000  $\Delta mopJ$  followed by selection on PYE plates harbouring tetracycline.

Strains harbouring pMR10 or derivatives. Plasmids were introduced by electroporation, followed by selection of transformants on PYE plates harbouring tetracycline (pMR20) or kanamycin (pMR10).

Strains harbouring xylX::Pxyl-mopJ-GFP. pCWR282 (pXGFP4-mopJ) was introduced into *C. crescentus* by electroporation and then plated on PYE harbouring kanamycin (to select for recombinants). A single homologous recombination event at the xyl locus of kanamycin-resistant colonies was verified by PCR.

Strains harbouring xylX:: $P_{xyl}$ -divJ-GFP, xylX:: $P_{xyl}$ -divJ392-GFP or xylX:: $P_{xyl}$ -divJ329-GFP. pXGFP4-divJ, pXGFP4-divJ392 or pXGFP4-divJ329 were introduced into *C. crescentus* by electroporation and then plated on PYE harbouring kanamycin (to select for recombinants). A single homologous recombination event at the xyl locus of kanamycin-resistant colonies was verified by PCR.

Strains harbouring pMR10-divK-GFP and pMT335 or pMT335-mopJ. pMT335-mopJ was introduced into strains harbouring pMR10-divK-GFP ( $P_{divK}$ -divK-GFP) by electroporation and then plated on PYE harbouring kanamycin and gentamycin.

 $NA1000 \ mopJ::P_{mopJ}-mopJ-GFP$ . pGFP4-mopJ was introduced into NA1000  $\Delta mopJ$  by electroporation and then plated on PYE harbouring kanamycin (to select for recombinants). A single homologous recombination event at the mopJ locus of kanamycin-resistant colonies was verified by PCR.

 $NA1000~xylX::P_{xyl}$ -GFP-mopJ. pXGFP4-C1-GFP-mopJ was introduced into NA1000 by electroporation and then plated on PYE harbouring kanamycin. A single homologous recombination event at the xyl locus of kanamycin-resistant colonies was verified by PCR.

NA1000  $xylX::P_{xyl}$ -divK-CFP  $P_{van}$ -mopJ-YFP. pMT374-mopJ-YFP was introduced into NA1000  $xylX::P_{xyl}$ -divK-CFP<sup>17</sup> by electroporation and then selected on PYE plates harbouring tetracycline and kanamycin.

Strains harbouring placZ290 or derivatives. placZ290 or derivatives<sup>10</sup> were introduced into *C. crescentus* strains by electroporation and then plated on PYE harbouring tetracycline.

NA1000~xylX::pXTCYC4~placZ290-PmopJ.~placZ290-PmopJ~was~introduced~into~NA1000~xylX::pXTCYC4~(ref.~47)~by~electroporation~and~then~plated~on~PYE~harbouring~gentamycin~and~tetracycline.

NA1000 harbouring xylX::pXTCYC4<sup>47</sup>, xylX::pXTCYC4-relA'-FLAG or xylX::pXTCYC4-relA'(E335Q)-FLAG and placZ290-PmopJ. placZ290-PmopJ was introduced into NA1000 xylX::pXTCYC4 (ref. 47), xylX::pXTCYC4-relA'-FLAG or xylX::pXTCYC4-relA'(E335Q)-FLAG<sup>47</sup> by electroporation and then plated on PYE harbouring gentamycin and tetracycline.

NA1000 divL::divL-GFP and  $\Delta$ mopJ divL::divL-GFP. divL-GFP::kan allele<sup>69</sup> (Kan<sup>R</sup>) was transduced into NA1000  $\Delta$ mopJ using  $\phi$ Cr30-mediated generalized transduction.

divL::divL-GFP strains harbouring pMT335 or derivatives. Plasmids were introduced into divL::divL-GFP strains by electroporation and then plated on PYE harbouring kanamycin and gentamycin.

EC100D pET28a-mopJshort. pET28a-mopJshort was introduced into EC100D by electroporation and then plated on LB harbouring kanamycin.

#### **Plasmid constructions**

pCWR296 (pNPTS138-\(\Delta\)mopJ-KO). The plasmid construct used for mopJ (CCNA\_00999) deletion was made by PCR amplification of two fragments. The first, a 568-bp fragment (nt 1081464–1082022, NA 1000 genome coordinates, flanked by an EcoRI site at the 5'end and a BamHI at the 3'end) was amplified using primers delmopJ\_1-BamHI and delmopJ\_1-EcoRI (sequences for all the primers used in this work can be found in Supplementary Table 5). The second, a

447-bp fragment (nt 1082446–1082880, flanked by a *Bam*HI site at the 5'end and a *Hind*III site at the 3'end) was amplified using primers delmopJ\_2-HindIII and delmopJ\_2-BamHI. These two fragments were first digested with appropriate restriction enzymes and then triple ligated into pNTPS138 (M.R.K. Alley, Imperial College London, unpublished) that had been previously restricted with *Hind*III and *EcoR*I. This construct deletes 423 nt of the *mopJ*-coding sequence (nt 1082023–1082445, or codons 14–154 of the annotated CCNA\_00999 coding sequence).

pMT335-mopJ (pUG16). The mopJ-coding sequence (nt 1081990–1082481, NA1000 genome) was PCR amplified from the NA1000 strain using the mopJ-NdeI and mopJ-EcoRI primers. This fragment was digested with NdeI/EcoRI and cloned into NdeI/EcoRI-digested pMT335 (ref. 70).

pMT335-divJ. The divJ-coding sequence (nt 1220499–1222292) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pMT335.

pMT335-divJ392. The divJ392-coding sequence (short divJ form without sequence encoding for kinase domain, nt 1220499–1221674) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ392-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pMT335.

pMT335-divJ329. The divJ329-coding sequence (short divJ form without sequence encoding for dimerization and kinase domains, nt 1220499–1221485) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ329-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pMT335.

pXGFP4-divJ. The divJ-coding sequence without stop codon (nt 1220499–1222289) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ\_fusion-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pXGFP4 (M.R.K. Alley, Imperial College London, unpublished).

pXGFP4-divJ392. The divJ392-coding sequence (short divJ form without sequence encoding for kinase domain, nt 1220499–1221674) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ392fusion-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pXGFP4.

pXGFP4-divJ329. The divJ329-coding sequence (short divJ form without sequence encoding for dimerization and kinase domains, nt 1220499–1221485) was PCR amplified from the NA1000 strain using the divJ-NdeI and divJ329fusion-MunI primers. This fragment was digested with NdeI/MunI and cloned into NdeI/EcoRI-digested pXGFP4.

pET28a-mopJshort. The mopJ-coding sequence lacking the first 132 bp from the start codon (nt 1082122–1082481) was PCR amplified from the NA1000 strain using the mopJshort-NdeI and mopJ-EcoRI primers. This fragment was digested with NdeI/EcoRI and cloned into NdeI/EcoRI-digested pET28a (Novagen).

pCWR282 (pXGFP4-mopJ). The mopJ-coding sequence without stop codon (nt 1081990–1082478) was PCR amplified from the NA1000 strain using the mopJ-NdeI and mopJfusion-BamHI primers. This fragment was digested with NdeI/BamHI and cloned into NdeI/BamHI-digested pXGFP4.

pGFP4-mopJ. The mopJ-coding sequence without stop codon and the upstream region of 640 bp (nt 1081990–1082478) were PCR amplified from the NA1000 strain using the mopJ-NheI and mopJfusion-BamHI primers. This fragment was digested with NheI/BamHI and cloned into pXGFP4, previously digested with NheI/BamHI to remove the PxyIX locus.

pXGFP4-C1-GFP-mopJ. The mopJ-coding sequence (nt 1081993–1082481), in which start codon ATG was replaced with GTC, was PCR amplified from the NA1000 strain using the mopJfusion-BglII and mopJ-EcoRI primers. This fragment was digested with BglII/EcoRI and cloned into BglII/EcoRI-digested pXGFP4-C1 (M.R.K. Alley, Imperial College London, unpublished).

<code>placZ290-PmopJ</code>. The <code>mopJ</code> promoter region ( - 228 to +78 relative to the ATG, nt 1081990–1081992) was PCR amplified using PmopJ-EcoRI and PmopJ-XbaI primers using NA1000 chromosomal DNA as a template. This fragment was digested by <code>EcoRI-XbaI</code> and cloned into <code>EcoRI-XbaI-digested placZ290</code> promoter probe vector.

#### References

- Guttenplan, S. B. & Kearns, D. B. Regulation of flagellar motility during biofilm formation. FEMS Microbiol. Rev. 37, 849–871 (2013).
- Skerker, J. M. & Laub, M. T. Cell-cycle progression and the generation of asymmetry in Caulobacter crescentus. Nat. Rev. Microbiol. 2, 325–337 (2004).
- Radhakrishnan, S. K., Thanbichler, M. & Viollier, P. H. The dynamic interplay between a cell fate determinant and a lysozyme homolog drives the asymmetric division cycle of *Caulobacter crescentus*. *Genes Dev.* 22, 212–225 (2008).
- Quon, K. C., Marczynski, G. T. & Shapiro, L. Cell cycle control by an essential bacterial two-component signal transduction protein. Cell 84, 83–93 (1996).
- Ely, B. Genetics of Caulobacter crescentus. Methods Enzymol. 204, 372–384 (1991).
- Ardissone, S. et al. Cell cycle constraints on capsulation and bacteriophage susceptibility. eLife 3 doi: 10.7554/eLife.03587 (2014).
- Laub, M. T., Shapiro, L. & McAdams, H. H. Systems biology of Caulobacter. Annu. Rev. Genet. 41, 429–441 (2007).
- Holtzendorff, J. et al. Oscillating global regulators control the genetic circuit driving a bacterial cell cycle. Science 304, 983–987 (2004).
- Fiebig, A. et al. A cell cycle and nutritional checkpoint controlling bacterial surface adhesion. PLoS Genet. 10, e1004101 (2014).
- Fumeaux, C. et al. Cell cycle transition from S-phase to G1 in Caulobacter is mediated by ancestral virulence regulators. Nat. Commun. 5, 4081 (2014).
- Laub, M. T., McAdams, H. H., Feldblyum, T., Fraser, C. M. & Shapiro, L. Global analysis of the genetic network controlling a bacterial cell cycle. *Science* 290, 2144–2148 (2000).
- Quon, K. C., Yang, B., Domian, I. J., Shapiro, L. & Marczynski, G. T. Negative control of bacterial DNA replication by a cell cycle regulatory protein that binds at the chromosome origin. *Proc. Natl Acad. Sci. USA* 95, 120–125 (1998).
- Bastedo, D. P. & Marczynski, G. T. CtrA response regulator binding to the Caulobacter chromosome replication origin is required during nutrient and antibiotic stress as well as during cell cycle progression. Mol. Microbiol. 72, 139–154 (2009).
- West, A. H. & Stock, A. M. Histidine kinases and response regulator proteins in two-component signaling systems. Trends Biochem. Sci. 26, 369–376 (2001).
- Tsokos, C. G., Perchuk, B. S. & Laub, M. T. A dynamic complex of signaling proteins uses polar localization to regulate cell-fate asymmetry in *Caulobacter crescentus*. Dev. Cell 20, 329–341 (2011).
- Biondi, E. G. et al. Regulation of the bacterial cell cycle by an integrated genetic circuit. Nature 444, 899–904 (2006).
- Lam, H., Matroule, J. Y. & Jacobs-Wagner, C. The asymmetric spatial distribution of bacterial signal transduction proteins coordinates cell cycle events. *Dev. Cell* 5, 149–159 (2003).
- Chien, P., Perchuk, B. S., Laub, M. T., Sauer, R. T. & Baker, T. A. Direct and adaptor-mediated substrate recognition by an essential AAA + protease. *Proc. Natl Acad. Sci. USA* 104, 6590–6595 (2007).
- Jenal, U. & Fuchs, T. An essential protease involved in bacterial cell-cycle control. EMBO J. 17, 5658–5669 (1998).
- Iniesta, A. A., McGrath, P. T., Reisenauer, A., McAdams, H. H. & Shapiro, L. A phospho-signaling pathway controls the localization and activity of a protease complex critical for bacterial cell cycle progression. *Proc. Natl Acad. Sci. USA* 103, 10935–10940 (2006).
- Hung, D. Y. & Shapiro, L. A signal transduction protein cues proteolytic events critical to *Caulobacter* cell cycle progression. *Proc. Natl Acad. Sci. USA* 99, 13160–13165 (2002).
- Iniesta, A. A., Hillson, N. J. & Shapiro, L. Cell pole-specific activation of a critical bacterial cell cycle kinase. *Proc. Natl Acad. Sci. USA* 107, 7012–7017 (2010).
- Wu, J., Ohta, N. & Newton, A. An essential, multicomponent signal transduction pathway required for cell cycle regulation in *Caulobacter. Proc.* Natl Acad. Sci. USA 95, 1443–1448 (1998).
- Wu, J., Ohta, N., Zhao, J. L. & Newton, A. A novel bacterial tyrosine kinase essential for cell division and differentiation. *Proc. Natl Acad. Sci. USA* 96, 13068–13073 (1999).
- Boutte, C. C. & Crosson, S. Bacterial lifestyle shapes stringent response activation. Trends Microbiol. 21, 174–180 (2013).
- Lesley, J. A. & Shapiro, L. SpoT regulates DnaA stability and initiation of DNA replication in carbon-starved *Caulobacter crescentus*. J. Bacteriol. 190, 6867–6880 (2008).
- Stott, K. V. et al. (p)ppGpp modulates cell size and the initiation of DNA replication in *Caulobacter crescentus* in response to a block in lipid biosynthesis. *Microbiology* 161, 553–564 (2015).
- Boutte, C. C., Henry, J. T. & Crosson, S. ppGpp and polyphosphate modulate cell cycle progression in Caulobacter crescentus. J. Bacteriol. 194, 28–35 (2012).
- Boutte, C. C. & Crosson, S. The complex logic of stringent response regulation in *Caulobacter crescentus*: starvation signalling in an oligotrophic environment. *Mol. Microbiol.* 80, 695–714 (2011).
- Tsokos, C. G. & Laub, M. T. Polarity and cell fate asymmetry in Caulobacter crescentus. Curr. Opin. Microbiol. 15, 744–750 (2012).

- Sciochetti, S. A., Lane, T., Ohta, N. & Newton, A. Protein sequences and cellular factors required for polar localization of a histidine kinase in *Caulobacter* crescentus. J. Bacteriol. 184, 6037–6049 (2002).
- Wheeler, R. T. & Shapiro, L. Differential localization of two histidine kinases controlling bacterial cell differentiation. Mol. Cell 4, 683–694 (1999).
- Paul, R. et al. Allosteric regulation of histidine kinases by their cognate response regulator determines cell fate. Cell 133, 452–461 (2008).
- Jacobs, C., Hung, D. & Shapiro, L. Dynamic localization of a cytoplasmic signal transduction response regulator controls morphogenesis during the Caulobacter cell cycle. Proc. Natl Acad. Sci. USA 98, 4095–4100 (2001).
- Matroule, J. Y., Lam, H., Burnette, D. T. & Jacobs-Wagner, C. Cytokinesis monitoring during development; rapid pole-to-pole shuttling of a signaling protein by localized kinase and phosphatase in *Caulobacter. Cell* 118, 579–590 (2004).
- Viollier, P. H., Sternheim, N. & Shapiro, L. Identification of a localization factor for the polar positioning of bacterial structural and regulatory proteins. *Proc. Natl Acad. Sci. USA* 99, 13831–13836 (2002).
- Hinz, A. J., Larson, D. E., Smith, C. S. & Brun, Y. V. The Caulobacter crescentus
  polar organelle development protein PodJ is differentially localized and is
  required for polar targeting of the PleC development regulator. Mol. Microbiol.
  47, 929–941 (2003).
- Bowman, G. R. et al. Caulobacter PopZ forms a polar subdomain dictating sequential changes in pole composition and function. Mol. Microbiol. 76, 173–189 (2010).
- Bowman, G. R. et al. A polymeric protein anchors the chromosomal origin/ ParB complex at a bacterial cell pole. Cell 134, 945–955 (2008).
- Ebersbach, G., Briegel, A., Jensen, G. J. & Jacobs-Wagner, C. A self-associating protein critical for chromosome attachment, division, and polar organization in *Caulobacter. Cell* 134, 956–968 (2008).
- Laloux, G. & Jacobs-Wagner, C. Spatiotemporal control of PopZ localization through cell cycle-coupled multimerization. J. Cell Biol. 201, 827–841 (2013).
- Henry, J. T., Crosson, S. & Ligand-binding, P. A. S. domains in a genomic, cellular, and structural context. *Annu. Rev. Microbiol.* 65, 261–286 (2011).
- Marks, M. E. et al. The genetic basis of laboratory adaptation in Caulobacter crescentus. J. Bacteriol. 192, 3678–3688 (2010).
- Skerker, J. M. & Shapiro, L. Identification and cell cycle control of a novel pilus system in Caulobacter crescentus. EMBO J. 19, 3223–3234 (2000).
- Domian, I. J., Quon, K. C. & Shapiro, L. Cell type-specific phosphorylation and proteolysis of a transcriptional regulator controls the G1-to-S transition in a bacterial cell cycle. *Cell* 90, 415-424 (1997).
- Potrykus, K. & Cashel, M. p)ppGpp: still magical? Annu. Rev. Microbiol. 62, 35–51 (2008).
- Gonzalez, D. & Collier, J. Effects of (p)ppGpp on the progression of the cell cycle of Caulobacter crescentus. J. Bacteriol. 196, 2514–2525 (2014).
- 48. Davis, N. J. et al. De- and repolarization mechanism of flagellar morphogenesis during a bacterial cell cycle. *Genes Dev.* 27, 2049–2062 (2013).
- Fioravanti, A. et al. DNA binding of the cell cycle transcriptional regulator GcrA depends on N6-adenosine methylation in Caulobacter crescentus and other Alphaproteobacteria. PLoS Genet. 9, e1003541 (2013).
- Murray, S. M., Panis, G., Fumeaux, C., Viollier, P. H. & Howard, M. Computational and genetic reduction of a cell cycle to its simplest, primordial components. *PLoS Biol.* 11, e1001749 (2013).
- Flynn, J. M., Neher, S. B., Kim, Y. I., Sauer, R. T. & Baker, T. A. Proteomic discovery of cellular substrates of the ClpXP protease reveals five classes of ClpX-recognition signals. *Mol. Cell* 11, 671–683 (2003).
- Bhat, N. H., Vass, R. H., Stoddard, P. R., Shin, D. K. & Chien, P. Identification of ClpP substrates in *Caulobacter crescentus* reveals a role for regulated proteolysis in bacterial development. *Mol. Microbiol.* 88, 1083–1092 (2013).
- Osteras, M., Stotz, A., Schmid Nuoffer, S. & Jenal, U. Identification and transcriptional control of the genes encoding the *Caulobacter crescentus* ClpXP protease. *J. Bacteriol.* 181, 3039–3050 (1999).
- Aakre, C. D., Phung, T. N., Huang, D. & Laub, M. T. A bacterial toxin inhibits DNA replication elongation through a direct interaction with the beta sliding clamp. *Mol. Cell* 52, 617–628 (2013).
- Puig, O. et al. The tandem affinity purification (TAP) method: a general procedure of protein complex purification. Methods 24, 218–229 (2001).
- Sciochetti, S. A., Ohta, N. & Newton, A. The role of polar localization in the function of an essential *Caulobacter crescentus* tyrosine kinase. *Mol. Microbiol.* 56, 1467–1480 (2005).
- Iniesta, A. A., Hillson, N. J. & Shapiro, L. Polar remodeling and histidine kinase activation, which is essential for *Caulobacter* cell cycle progression, are dependent on DNA replication initiation. *J. Bacteriol.* 192, 3893–3902 (2010).
- 58. Collier, J. Regulation of chromosomal replication in *Caulobacter crescentus*. *Plasmid* **67**, 76–87 (2012).
- 59. Curtis, P. D. *et al.* The scaffolding and signalling functions of a localization factor impact polar development. *Mol. Microbiol.* **84**, 712–735 (2012).
- Duerig, A. et al. Second messenger-mediated spatiotemporal control of protein degradation regulates bacterial cell cycle progression. Genes Dev. 23, 93–104 (2009).
- Chen, J. C. et al. Cytokinesis signals truncation of the PodJ polarity factor by a cell cycle-regulated protease. EMBO J 25, 377–386 (2006).

- Jonas, K., Liu, J., Chien, P. & Laub, M. T. Proteotoxic stress induces a cell-cycle arrest by stimulating Lon to degrade the replication initiator DnaA. *Cell* 154, 623–636 (2013).
- Gorbatyuk, B. & Marczynski, G. T. Regulated degradation of chromosome replication proteins DnaA and CtrA in *Caulobacter crescentus*. *Mol. Microbiol*. 55, 1233–1245 (2005).
- Maisonneuve, E. & Gerdes, K. Molecular mechanisms underlying bacterial persisters. Cell 157, 539–548 (2014).
- De Nisco, N. J., Abo, R. P., Wu, C. M., Penterman, J. & Walker, G. C. Global analysis of cell cycle gene expression of the legume symbiont Sinorhizobium meliloti. Proc. Natl Acad. Sci. USA 111, 3217–3224 (2014).
- Wang, J. D., Sanders, G. M. & Grossman, A. D. Nutritional control of elongation of DNA replication by (p)ppGpp. Cell 128, 865–875 (2007).
- Figge, R. M., Divakaruni, A. V. & Gober, J. W. MreB, the cell shapedetermining bacterial actin homologue, co-ordinates cell wall morphogenesis in Caulobacter crescentus. Mol. Microbiol. 51, 1321–1332 (2004).
- Potocka, I., Thein, M., ØSterås, M., Jenal, U. & Alley, M. R. Degradation of a Caulobacter soluble cytoplasmic chemoreceptor is ClpX dependent. J. Bacteriol. 184, 6635–6641 (2002).
- Hughes, H. V. et al. Protein localization and dynamics within a bacterial organelle. Proc. Natl Acad. Sci. USA 107, 5599–5604 (2010).
- Thanbichler, M., Iniesta, A. A. & Shapiro, L. A comprehensive set of plasmids for vanillate- and xylose-inducible gene expression in *Caulobacter crescentus*. *Nucleic Acids Res.* 35, e137 (2007).

#### **Acknowledgements**

Funding support is from SNF grant #31003A\_143660 (to P.H.V.) and Fondation pour des bourses d'études Italo-Suisses (to S.S.). We thank Sean Crosson (University of

Chicago), Jim Gober (UCLA), Kathleen Ryan (UC Berkeley), Lucy Shapiro (Stanford University), Urs Jenal (University of Basel), Mike Laub (MIT), Christine Jacobs-Wagner (Yale University) and Justine Collier (University of Lausanne) for materials, Gaël Panis for help with illustrations and FACS.

#### **Author contributions**

S.S., M.B., S.K.R. and P.H.V conceived and designed the experiments. S.S., M.B., L.T., S.K.R. and P.H.V. performed the experiments. S.S., M.B., S.K.R. and P.H.V analysed the data. S.S. and P.H.V. wrote the paper.

#### **Additional information**

Supplementary Information accompanies this paper at http://www.nature.com/ naturecommunications

Competing financial interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

**How to cite this article:** Sanselicio, S. *et al.* Topological control of the *Caulobacter* cell cycle circuitry by a polarized single-domain PAS protein. *Nat. Commun.* 6:7005 doi: 10.1038/ncomms8005 (2015).



This work is licensed under a Creative Commons Attribution 4.0 International License. The images or other third party material in this

article are included in the article's Creative Commons license, unless indicated otherwise in the credit line; if the material is not included under the Creative Commons license, users will need to obtain permission from the license holder to reproduce the material. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/