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# A checkpoint control orchestrates the replication of the two chromosomes of *Vibrio cholerae*

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Bacteria with multiple chromosomes represent up to 10% of all bacterial species. Unlike eukaryotes, these bacteria use chromosome-specific initiators for their replication. In all cases investigated, the machineries for secondary chromosome replication initiation are of plasmid origin. One of the important differences between plasmids and chromosomes is that the latter replicate during a defined period of the cell cycle, ensuring a single round of replication per cell. *Vibrio cholerae* carries two circular chromosomes, Chr1 and Chr2, which are replicated in a well-orchestrated manner with the cell cycle and coordinated in such a way that replication termination occurs at the same time. However, the mechanism coordinating this synchrony remains speculative. We investigated this mechanism and revealed that initiation of Chr2 replication is triggered by the replication of a 150-bp locus positioned on Chr1, called *crtS*. This *crtS* replication-mediated Chr2 replication initiation mechanism explains how the two chromosomes communicate to coordinate their replication. Our study reveals a new checkpoint control mechanism in bacteria, and highlights possible functional interactions mediated by contacts between two chromosomes, an unprecedented observation in bacteria.

## INTRODUCTION

Bacteria with secondary chromosomes are frequent and have arisen independently in several taxa (1). This is the case for pathogens such as *Vibrio* or *Burkholderia*, symbionts such as *Rhizobia*, and others. Domestication of large plasmids, after transfer of essential genes, appears to explain the origin of secondary chromosomes (2). Evidence supporting this hypothesis includes the fact that all secondary chromosomes carry plasmid-like replication systems (2). Replication of bacterial chromosomes is regulated at initiation from a single well-conserved origin of replication (*oriC*) under the control of DnaA, the universal initiator of chromosome replication in bacteria (3), whereas plasmids have various types of replication origins. Usually, their replication is controlled by an initiator binding to directly repeated sequences (iterons) or by an antisense RNA (4).

All reported members of the Vibrionaceae family (*Vibrio*, *Listonella*, *Aliivibrio*, and *Photobacterium*) have two chromosomes of uneven size (5). Most of our knowledge on the replication control of secondary chromosomes comes from studies of *Vibrio cholerae*, the causative agent of cholera in humans. *V. cholerae* has two circular chromosomes, a main chromosome (Chr1) of 3 Mbp and a secondary chromosome (Chr2) of 1 Mbp (6). Chr1 replication is initiated at an *oriC*-like origin, *ori1*, by DnaA. Chr2 has a plasmid-like origin, *ori2*, where replication is regulated by a *Vibrio*-specific factor, RctB (7). RctB is a large protein [658 amino acids (AA)] that binds to DNA as a monomer or as a dimer (8). RctB binds and hydrolyzes adenosine 5'-triphosphate (ATP), but unlike DnaA, the ATP-bound form of RctB is inactive (9). Various regulatory mechanisms, such as initiator autoregulation, initiator titration, and origin handcuffing, control the level and activity of RctB [for review, see the study by Val *et al.* (10)]. RctB binds to iterons (12-mer sites) in *ori2*,

which promotes replication initiation. Additionally, RctB binds other regulatory sites. Within *ori2*, RctB binds to 39-mer regulatory sites, which strongly inhibit *ori2* initiation (11). Beyond the origin region of Chr2, chromatin immunoprecipitation with DNA microarray (ChIP-chip) analysis revealed that RctB binds to a locus on Chr2 (coordinates 956828-1030773) containing five iterons and one 39-mer, which inhibits the replication of an *ori2*-driven plasmid (mini-chr2) in *Escherichia coli* (12). RctB was also found to bind a locus on Chr1 sharing no homology with either iterons or 39-mers. This site was shown to act as a replication enhancer of *ori2* by increasing RctB affinity for iterons and decreasing RctB affinity for 39-mers (12). Serial deletions of a DNA fragment containing the Chr1 RctB ChIP-chip binding peak (fragments chr1-2 to chr1-10) showed that the replication-enhancing activity of a mini-chr2 in *E. coli* could be narrowed down to a 70-bp chr1-9 fragment (coordinates 818000-818069). However, the larger (150 bp) chr1-4 fragment (coordinates 817947-818099) was more efficient in enhancing mini-chr2 replication in *E. coli*. The presence of such a site on Chr1 suggested that the two chromosomes communicate with each other during replication. However, the role of this locus in the replication coordination of Chr1 and Chr2 remains a subject of speculation.

Chromosomes replicate during a defined period of the cell cycle, ensuring a single round of replication per cell. Plasmids generally have no such constraint, replicating randomly during the bacterial cell cycle (13). Despite its plasmid origin, Chr2 replication occurs only once per cell cycle (14). Initiation of Chr2 replication is also delayed so that the two chromosomes terminate replication at nearly the same time (15). The mechanism responsible for triggering Chr2 replication at a specific time of the cell cycle remains unknown. Here, we provide new insights into this regulatory process, and we explain how Chr2 monitors the replication status of Chr1 to time its own replication. We show that the Chr1 RctB binding site (12), renamed *crtS* for Chr2 replication triggering site, is crucial for the activation of Chr2 replication. We demonstrate that the replication of *crtS* triggers the replication of Chr2. We also show that the *crtS* locus and *ori2* localize to the same region of the cell during the entire cell cycle and display enhanced physical contacts, suggesting that

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the regulatory mechanisms may involve a structural interplay. This study reveals a new checkpoint control mechanism in bacteria.

## RESULTS

### Marker frequency analysis reveals the relative replication pattern of the two chromosomes of *V. cholerae*

Replication of bacterial chromosomes occurs bidirectionally and terminates in the region opposite to the origin, in the vicinity of the dimer resolution site (*dif*), forming two replicated halves called replichores (Fig. 1A, top) (16). Assuming that Chr1 and Chr2 replicate at the same speed, replication of Chr2 must be delayed so that replication termination of the two chromosomes can be synchronous (15). Control of Chr2 replication initiation might be linked to cell mass or some “timer” on Chr1 that, when replicated, signals initiation on Chr2. We used marker frequency analysis (MFA) to precisely analyze the replication pattern of the two chromosomes of a culture of wild-type *V. cholerae* El Tor N16961 strain (WT) grown under steady-state conditions. MFA provides an unprecedented resolution of the replication timing and the replication fork speed, can pinpoint the origin and the terminus of chromosome replication, and can detect chromosomal rearrangements (17). Indeed, the MFA plot of WT compared to the reference genome sequence (AE003852) registered a discrepancy in Chr1 organization (fig. S1). Rectification of the Chr1 reference sequence corrected the deviation. MFA of WT confirmed that both chromosomes are replicated bidirectionally, with each chromosome having one origin (*ori1* and *ori2*) and one terminus (*ter1* and *ter2*) of replication. The linearity and the slopes of the graphs for the four replichores indicate that replication speed is constant for both chromosomes (Fig. 1A, bottom). This analysis confirmed that Chr1 and Chr2 terminate replication concomitantly and that Chr2 is initiated when about two-thirds of Chr1 is replicated, as hypothesized.

### Chr2 replication initiation depends on the location of a timer region on Chr1

Chr2 replication initiation, after two-thirds of Chr1 has been replicated, could be linked to an unknown mechanism for coordination of replication termination of the two chromosomes, consistent with their relative sizes, 3 and 1 Mbp. To evaluate the impact of the chromosome sizes on their relative timing of replication, we generated two mutants with altered chromosome sizes using a dual site-specific recombination tool to transfer DNA from one chromosome to the other (18). In the CSV2 mutant, Chr1 and Chr2 sizes remained unbalanced at 2.5 and 1.5 Mb, respectively, whereas in the ESC2 strain, the two chromosomes are each at 2 Mb (Fig. 1, B and C). Increasing the size of Chr2 abolished synchronous termination in both ESC2 and CSV2 mutants, with the now larger Chr2 terminating replication after Chr1 (Fig. 1, B and C). MFA also revealed that Chr2 systematically initiates replication when a discrete position along one (or both) Chr1 replicore(s) is being replicated (Fig. 1, B and C). This finding suggested that a region located on Chr1 may trigger Chr2 replication initiation, and should be localized within the first two-thirds of either or both replichores and act as a “checkpoint” in replication of Chr2.

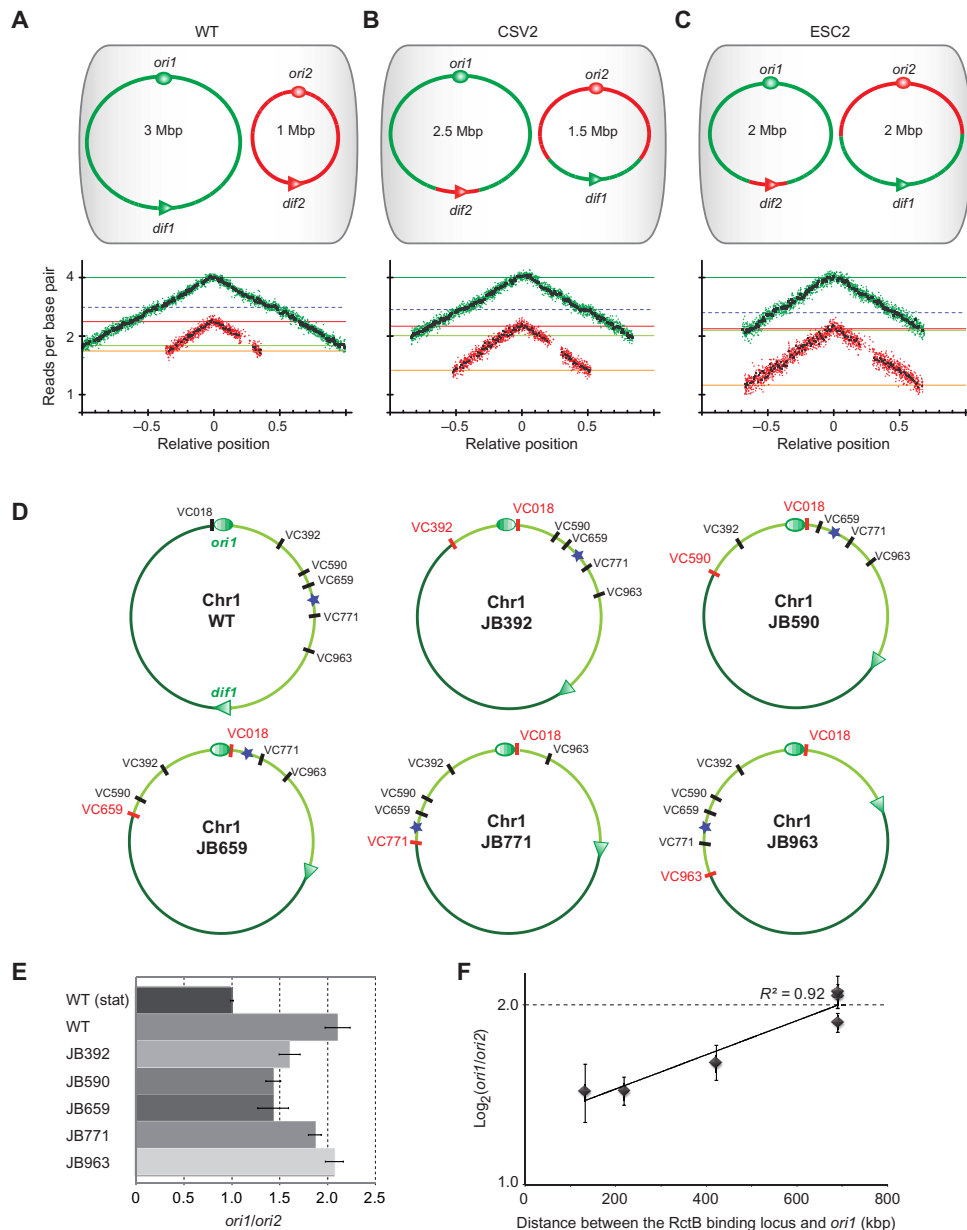
The relative timing of initiation of Chr1 and Chr2 can be followed by qPCR analysis of the *ori1/ori2* ratio. We reasoned that a change in this ratio in isogenic mutants, where Chr1 size was unaltered but replichores were rearranged, would yield a signal for the region of interest.

This was performed by inversion between a fixed intergenic locus (downstream of ORF VC018, that is, near *ori1* on the left replicore) and other intergenic loci located at increasing distances from *ori1* along the right replicore (Fig. 1D). Such inversions either caused no fitness cost (JB392) or were similarly affected (JB590, JB659, JB771, and JB963) compared to WT (fig. S2). We monitored the impact of each inversion on the *ori1/ori2* ratio in exponentially growing cultures (Fig. 1E). In WT, the *ori1/ori2* ratio is around 2, matching the observations of fast-growing *V. cholerae* (15). In mutants JB392, JB590, and JB659, *ori1/ori2* ratios decreased, indicating that the region triggering Chr2 replication may be closer to *ori1*, causing earlier Chr2 replication initiation. In mutant JB771 and JB963, the *ori1/ori2* ratio remains ~2, indicating the wild type-like timing of replication and that the locus triggering Chr2 initiation must be at the same distance from *ori1* as in wild type. These results indicate that a locus located between VC659 and VC771 and its distance from *ori1* play a role in the regulation of Chr2 replication. This region contains the Chr1 RctB binding locus, located in a noncoding region upstream of VC765 (12). Strikingly, the  $\log_2(\text{ori1/ori2})$  ratio increases linearly with the distance between *ori1* and the Chr1 RctB binding locus (Fig. 1F). This observation indicates that the timing of replication of the Chr1 RctB binding locus exerts a control on Chr2 replication initiation.

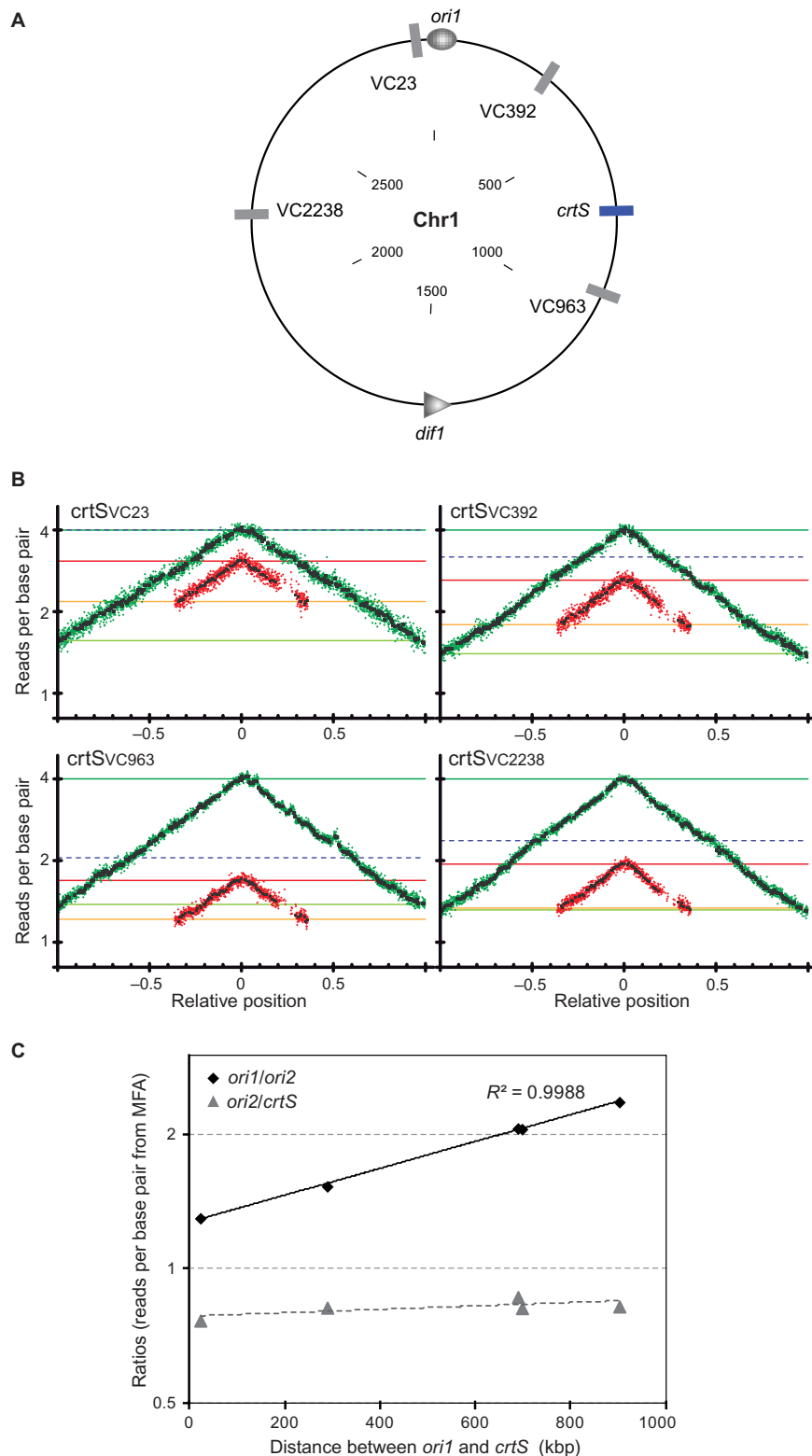
### Replication of the Chr1 RctB binding locus (*crtS*) triggers Chr2 replication initiation

We tested the Chr1 RctB binding locus for its role in the coordination of the timing of replication between Chr1 and Chr2. Hereafter, we will refer to the Chr1 RctB binding locus as *crtS*. *crtS* was relocated to four intergenic loci on Chr1 at varying distances from *ori1* (Fig. 2A). *crtS*-positional mutants display wild type-like fitness and phenotype (fig. S3). Therefore, all the determinants for *crtS* proper function appeared contained within its sequence. MFA showed that in *crtS*<sub>VC23</sub> and *crtS*<sub>VC392</sub>, where *crtS* is closer to *ori1*, Chr2 initiates earlier than in WT (Fig. 2B). In *crtS*<sub>VC2238</sub>, where *crtS* is positioned at the same distance from *ori1* but on the other replicore, Chr2 initiates roughly at the same time as in WT (Fig. 2B). In *crtS*<sub>VC963</sub>, with *crtS* farther from *ori1*, Chr2 initiates later than in WT (Fig. 2B). We calculated the *ori1/ori2* and *ori2/crtS* ratios from the MFA (Fig. 2C). The  $\log_2$  of *ori1/ori2* ratio is linearly correlated with the *ori1-crtS* distance ( $R^2 = 0.9988$ ), suggesting that the timing of replication of *crtS* controls the timing of Chr2 replication initiation. The *crtS/ori2* ratio remains constant (~0.8), indicating that there is a constant delay between *crtS* replication and Chr2 replication initiation.

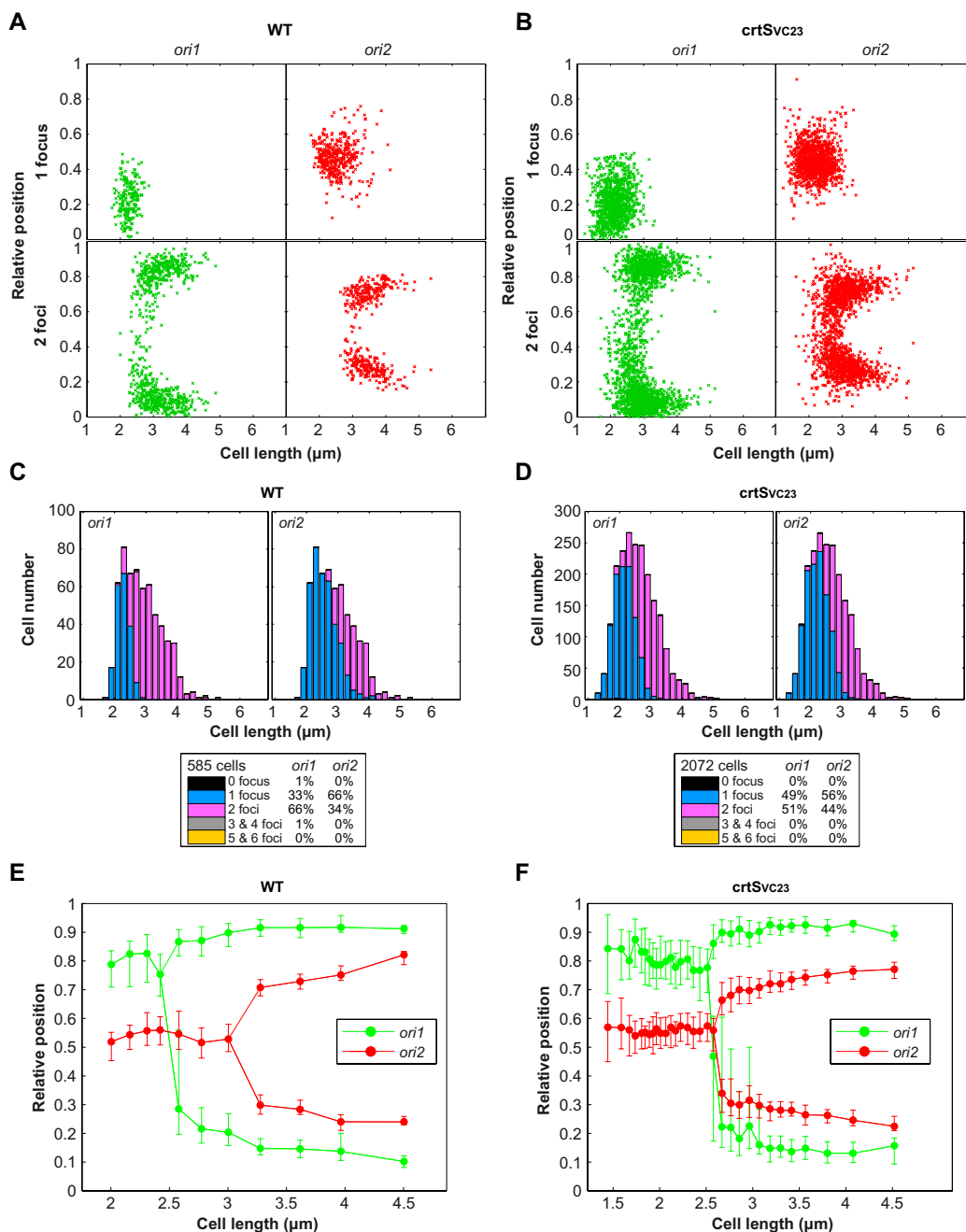
To confirm these observations, we tracked pairwise combinations of fluorescently labeled chromosomal positions using epifluorescence microscopy (19). We compared the distribution of *ori1* with *ori2* foci, VC783 (near *crtS*) with *ori2* foci, and *ter1* with *ter2* foci in both WT and *crtS*<sub>VC23</sub> strains (Fig. 3 and figs. S4 and S5). Images of exponentially growing cells were acquired, and cell length, along with the position of each tagged loci, was followed. WT cells ranged in size from 2  $\mu\text{m}$  (newborn cells) to 4.5  $\mu\text{m}$  (dividing cells), with two *ori1* foci appearing in cells from 2.5 to 3  $\mu\text{m}$  and two *ori2* foci from 3 to 3.5  $\mu\text{m}$  (Fig. 3, A, C, and E, and fig. S6, left). This observation indicates that *ori1* is replicated and segregated before *ori2*, as previously observed (19). In *crtS*<sub>VC23</sub>, cells with two *ori1* and two *ori2* foci appeared in the same cell size range from 2.5 to 3  $\mu\text{m}$  (Fig. 3, B, D, and F, and fig. S6, right), consistent with the MFA results showing that *ori2* is replicated shortly after *ori1* (Fig. 2B). In WT, duplication of VC783 foci



**Fig. 1. Chr1 and Chr2 replication coordination is promoted by the presence of a timer on Chr1 and not by the requirement to terminate their replication synchronously.** (A) Top: Genome structure of wild-type (WT) *V. cholerae*. Ovals indicate the origins of replication (*ori1* and *ori2*) and triangles show *dif* sites (*dif1* and *dif2*) on Chr1 (green) and Chr2 (red). Bottom: MFA of exponentially growing WT cultures using a corrected reference sequence of Chr1 (fig. S1).  $\text{Log}_2$  of number of reads starting at each base (normalized against reads from a stationary phase WT control) is plotted against their relative position on Chr1 and Chr2. Positions of *ori1* and *ori2* are set to 0 for a better visualization of the bidirectional replication. Any window containing repeated sequences is omitted; thus, the large gap observed in the right arm of Chr2 consists of filtered repeated sequences within the superintegron (28). Green (Chr1) and red (Chr2) dots indicate the average of 1000-bp windows; black dots indicate the average of 10,000-bp windows. Dark green, light green, red, and orange lines indicate *ori1*, *ter1*, *ori2*, and *ter2* number of reads, respectively; dashed blue lines indicate the Chr1 RctB binding locus (12). The same color code is used for all MFA figures. (B and C) Genomic variants CSV2 (Chr1 = 2.5 Mbp and Chr2 = 1.5 Mbp) (B) and ESC2 (Chr1 = 2 Mbp, Chr2 = 2 Mbp) (C) are the same as in (A). The genetic exchanges made between Chr1 and Chr2 are shown in green and red. (D) Chr1 map of WT and genomic variants (JB392, JB590, JB659, JB771, and JB963) with large chromosomal inversions around a fixed locus (VC018) and other loci located at increasing distances from *ori1* (VC392, VC590, VC659, VC771, and VC963), respectively. For each genomic mutant, the loci flanking the DNA inversion are shown in red. The left (dark green) and right (light green) replichores are separated by *ori1* (oval) and *dif1* (triangle). The position of the Chr1 RctB binding locus is indicated by a blue star. (E) Histogram representing quantitative PCR-measured *ori1/ori2* ratios from relative gDNA quantification of exponentially fast-growing strains (gDNA from WT stationary culture was used for normalization). Bars display means ( $\pm$ SD) of at least three experiments. (F)  $\text{Log}_2(\text{ori1/ori2})$  plotted as a function of the distance between *ori1* and the RctB binding locus displays a linear relationship ( $R^2 = 0.92$ ). Dots show means ( $\pm$ SD) of three experiments.



**Fig. 2. Timing of replication of the Chr1 RctB binding locus (*crtS*) controls the timing of initiation of Chr2.** (A) Circular map of WT Chr1 showing the native location of *crtS* (blue bar) and the various loci where *crtS* was relocated (gray bars) with respect to *ori1* (oval) and *dif1* (triangle). The inside scale designates DNA size in kilobase pair. (B) MFA of relocated *crtS* mutants (*crtS*<sub>VC23</sub>, *crtS*<sub>VC392</sub>, *crtS*<sub>VC963</sub>, and *crtS*<sub>VC2238</sub>). The dashed blue lines indicate the number of reads of the loci where *crtS* has been relocated. (C)  $\log_2(\text{ori1}/\text{ori2})$  and  $\log_2(\text{ori2}/\text{crtS})$  plotted as a function of the distance between *crtS* and *ori1* in kilobase pair. The ratios were calculated as the ratios of the number of reads per base pair (from MFA) for each designated loci.



**Fig. 3. Segregation of *ori2* (but not *ter2*) occurs earlier when *crtS* is transposed near *ori1*.** (A and B) Plot showing the position of *ori1* (left panel) and *ori2* (right panel) foci inside WT (A) and mutant *crtS<sub>VC23</sub>* (B) cells. Foci are oriented longitudinally relative to the old pole of the cell as a function of cell length. The old pole of the cells was defined as the closest pole to an *ori1* focus. The x axis represents the cell length (in micrometers). The y axis represents the relative position of the focus in bacterial cells, 0 being the old pole and 1 the new pole. Snapshot images of 585 WT and 2072 *crtS<sub>VC23</sub>* mutant cells were analyzed. (C and D) Histograms displaying the amount of cells that exhibit zero, one, two, three, four, or five and six fluorescent foci according to cell size (in micrometers) in WT (C) and mutant *crtS<sub>VC23</sub>* (D) cells. (E and F) By correlating the longitudinal position of *ori1* and *ori2* foci as a function of cell length, the segregation choreographies of the *ori1* and *ori2* were reconstituted throughout the cell cycle of WT (E) and mutant *crtS<sub>VC23</sub>* (F) bacterial cells. Cells were classified according to their size and grouped by 30 to define each size interval. For most loci and cell length intervals, there were cells with either a single focus or two separated foci, the relative proportions of each type varying as a function of cell length. Only the position of the foci corresponding to the dominant cell type in each cell length interval was plotted. The median positions of the observed foci (filled circles), along with the 25th to 75th percentiles (error bars), were plotted for each cell size bin. The x axis represents the cell length (in micrometers). The y axis represents the relative position of the focus in bacterial cells (0, new pole and 1, old pole).

(near *crtS*) occurs shortly before *ori2* foci duplication (fig. S4, C and E), which is in accordance with MFA, showing a delay between *crtS* replication and *ori2* replication (Fig. 1A). In *crtS*<sub>VC23</sub>, when *crtS* is no longer near VC783, *ori2* foci duplicate before VC783 foci (fig. S4, D and F), consistent with MFA results (Fig. 2B). Duplications of the *ter1* and *ter2* foci in WT were only visible at the end of the cell cycle in dividing cells (fig. S5, A, C, and E, and fig. S6, left). Surprisingly, most *crtS*<sub>VC23</sub> cells with two *ter1* and two *ter2* foci appeared in the same cell size range as WT (fig. S5, B, D, and F, and fig. S6, right). Because MFA studies showed that *ter2* is replicated long before *ter1* in *crtS*<sub>VC23</sub> mutant (Fig. 2B), this result indicates that duplicated *ter1* and *ter2* remain colocalized at mid-cell until cell division occurs.

To further probe the influence of *crtS* doubling on the initiation of Chr2 replication, we analyzed two mutant strains, *crtS*<sub>WT/VC23</sub> and *crtS*<sub>WT/VC2238</sub>, carrying two chromosomal copies of *crtS*. The *crtS*<sub>WT/VC23</sub> mutant carries the native *crtS* site, as well as an extra copy near *ori1* (VC23), so that one *crtS* is replicated before the other. The two *crtS* copies of the *crtS*<sub>WT/VC2238</sub> mutant are located at equal distances from *ori1* on each replichore; thus, the two *crtS* are replicated at the same time. Both mutants displayed a significant loss of fitness compared to WT, with *crtS*<sub>WT/VC2238</sub> being the least affected (fig. S7). Both *crtS*<sub>WT/VC23</sub> and *crtS*<sub>WT/VC2238</sub> mutants displayed a wider cell size range, from 2  $\mu$ m (newborn cells) to 5.5  $\mu$ m (dividing cells) (Fig. 4 and fig. S8), indicating a flaw in the cell cycle control. Tracking of fluorescently labeled *ori1* and *ori2* loci reveals that most *crtS*<sub>WT/VC23</sub> newborn cells exhibit two *ori2* foci but only one *ori1* focus (Fig. 4, A and B) compared to WT (Fig. 3A). Most *crtS*<sub>WT/VC23</sub> dividing cells present two *ori1* foci and four *ori2* foci (Fig. 4, A to C), suggesting that whereas Chr1 goes from one to two copies, Chr2 goes from two to four copies per cell cycle. Similar results were obtained with the *crtS*<sub>WT/VC2238</sub> mutant (fig. S8). In both mutants, daughter cells usually receive one copy of *ori1* and two copies of *ori2*, whereas WT daughter cells normally receive a single copy of both *ori1* and *ori2* (Fig. 4, C to E). Sporadically, in both mutant strains, we observed that cell division occurs asymmetrically, leading to uneven chromosome partitioning. The proportion of cells with three *ori2* foci in *crtS*<sub>WT/VC2238</sub> (15%) is equivalent to the proportion of cells with one *ori2* foci (14%) (Fig. 4E), suggesting that these cells could have arisen from the asymmetric cell division of four *ori2* foci cells. However, *crtS*<sub>WT/VC23</sub> cells display a larger fraction of cells containing three *ori2* foci (30%) and a smaller fraction of cells with only one *ori2* foci (7%) (Fig. 4D), meaning that uneven chromosome partitioning alone does not explain the existence of three *ori2* foci cells. We speculate that in *crtS*<sub>WT/VC23</sub>, one of the two *ori2* duplicates before the other. These differences between *crtS*<sub>WT/VC23</sub> and *crtS*<sub>WT/VC2238</sub> suggest that the difference in location and/or timing of replication of the extra *crtS* site alters the synchrony of initiation of multiple origins. These observations are puzzling because they do not fit with the *E. coli* paradigm where multiple origins initiate in synchrony (20). To explain that one *ori2* is duplicated before the other in *crtS*<sub>WT/VC23</sub>, we hypothesized that the duplication of one *crtS* 3 triggers the firing of only one *ori2*. This may require a direct contact between the replicated *crtS* and *ori2*.

### Chromosome conformation capture reveals a preferential contact between Chr1 and Chr2

To test for the possibility that Chr1-located *crtS* site regulation of Chr2 replication initiation involves physical contacts between the two chromosomes, we applied chromosome conformation capture [3C; (21)] to exponentially growing cultures in rapid (LB) and slow [minimal medium

(MM)] growth conditions (Fig. 5A and fig. S9). As expected, the two chromosomes in the resulting contact map turned out as two well-individualized entities, each exhibiting a strong diagonal signal reflecting frequent contacts between adjacent loci (Fig. 5A) (22). These chromosomes presented local domains of increased contact frequencies, separated by barriers (fig. S10), similar to the chromatin interaction domains (CIDs) previously described in *Bacillus subtilis* (23, 24) and *Caulobacter crescentus* (25). The intrachromosomal contact maps of Chr1 and Chr2 differ with respect to the presence of a secondary perpendicular diagonal that reflects the bridging of left and right replichores by cohesin complexes in other bacteria (23–25). Whereas Chr1 did not present such a signal, Chr2 displayed a secondary diagonal revealing radically different folding of the two chromosomes (Fig. 5A). The conversion of contact maps into three-dimensional (3D) structures (fig. S9C and movies S1 and S2) (26) shows that whereas Chr1 adopts a largely open structure, Chr2 folds into a helicoidally shape with its two replichores tightly interlaced. It appears from this matrix that Chr2 folds into a structure similar to that of *B. subtilis* (23, 24) or *C. crescentus* (27), whereas Chr1 adopts a structure closer to the one observed for *E. coli* chromosome (22). The superintegron, a large gene capture and excision system localized on Chr2 (28), defines a clear CID surrounded by two highly transcribed genes (fig. S10). This observation suggests a specific topological structure, which can reflect the generally low transcription in this DNA element.

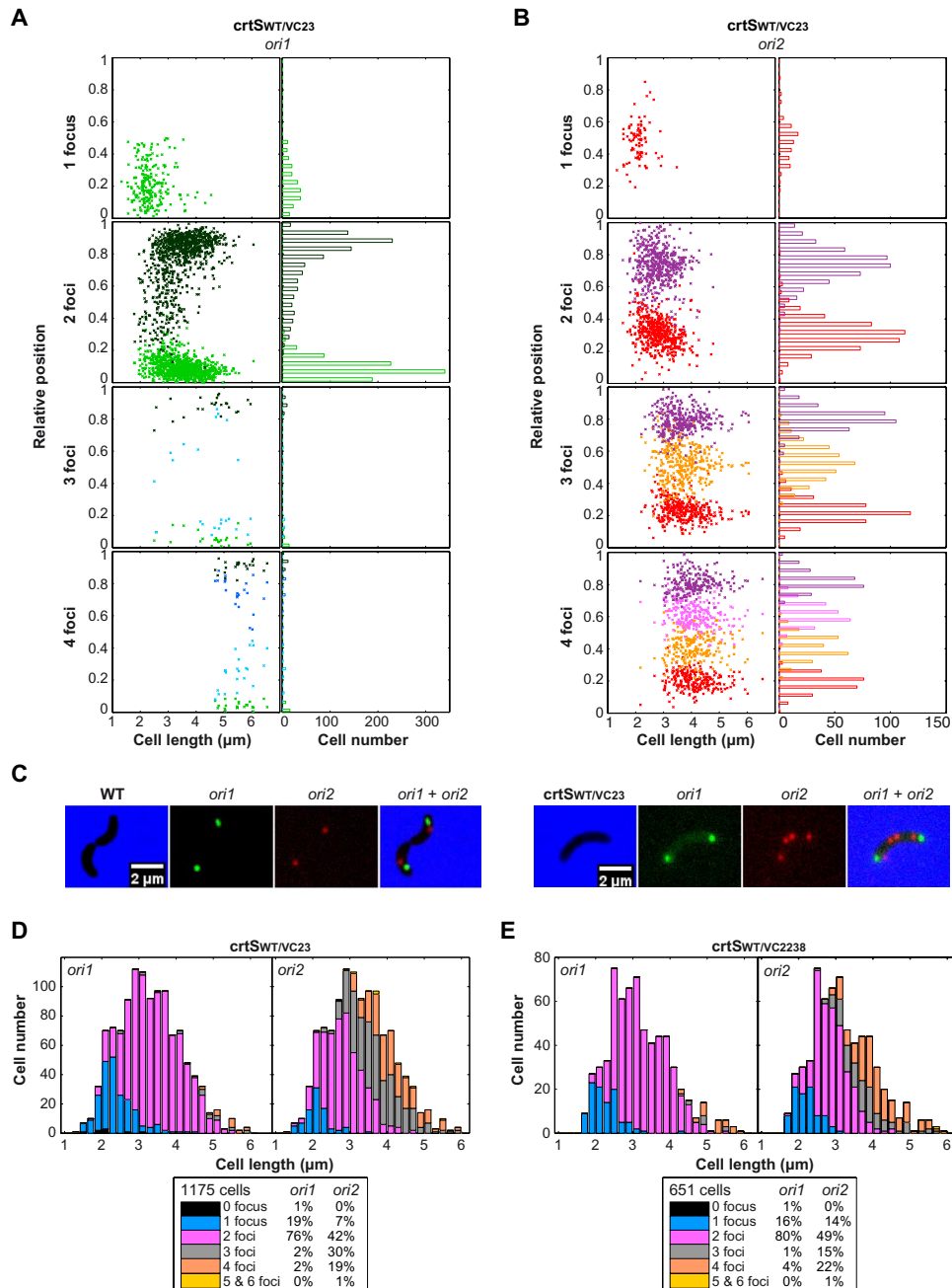
Overall, the chromosomes of *V. cholerae* present known features of chromosome organization in bacteria, as well as new ones. Indeed, the contact maps unveiled trans contacts between the two chromosomes at an unprecedented resolution. The two replichores of Chr2 present enrichment in contact with the bottom third of Chr1 replichore, as indicated by the cross-shaped trans contacts (Fig. 5B). These contacts initiate at *dif* sites and extend along the length of the chromosome up to *ori2* for Chr2 and midway to Chr1. The spatial proximity of the terminus regions (*ter*), surrounding *dif1* and *dif2*, is a striking driver of organization, with strong contacts between the *ter* regions, visible in both growth conditions (fig. S9A). The circos representation of the strongest interactions of 100 kbp surrounding *dif2* (Fig. 5C) illustrates the enrichment of contacts between the *ter* regions. This is also visible in the 3D contact maps (fig. S9C and movies S1 and S2). These results are consistent with the imaging of the *ter1* and *ter2* showing a colocalization of the regions to the mid-cell at the end of the cell cycle (fig. S5E). The circos representation of the contacts made by a 50-kbp window centered on *ori2* reveals preferential contacts with the right replichore of Chr1 (Fig. 5D). These contacts initiate at *crtS* and become stronger immediately after. More precisely, the contacts involve a region on the Chr2 left replichore adjacent to *ori2*, pointing toward a mechanical interplay that would drive this interaction.

### *crtS* is crucial for Chr2 replication initiation at *ori2*

To assess the importance of *crtS* on Chr2 replication, we deleted a 150-bp sequence (coordinates 817950-818100), which closely corresponds to the chrI-4 sequence (coordinates 817947-818099) deleted by Baek and Chatteraj (12). Baek and Chatteraj (12) reported that chrI-4-deleted mutants showed minimal phenotypic changes and no growth defects, indicating that its action was probably modest. In contrast, our  $\Delta$ *crtS* mutants exhibited strong fitness defects and suffered marked physiological changes, with a large proportion of filamentous cells (Fig. 6A, top, and fig. S11). To corroborate the marked phenotype of  $\Delta$ *crtS* mutants with a problem in *ori2* replication initiation, we deleted

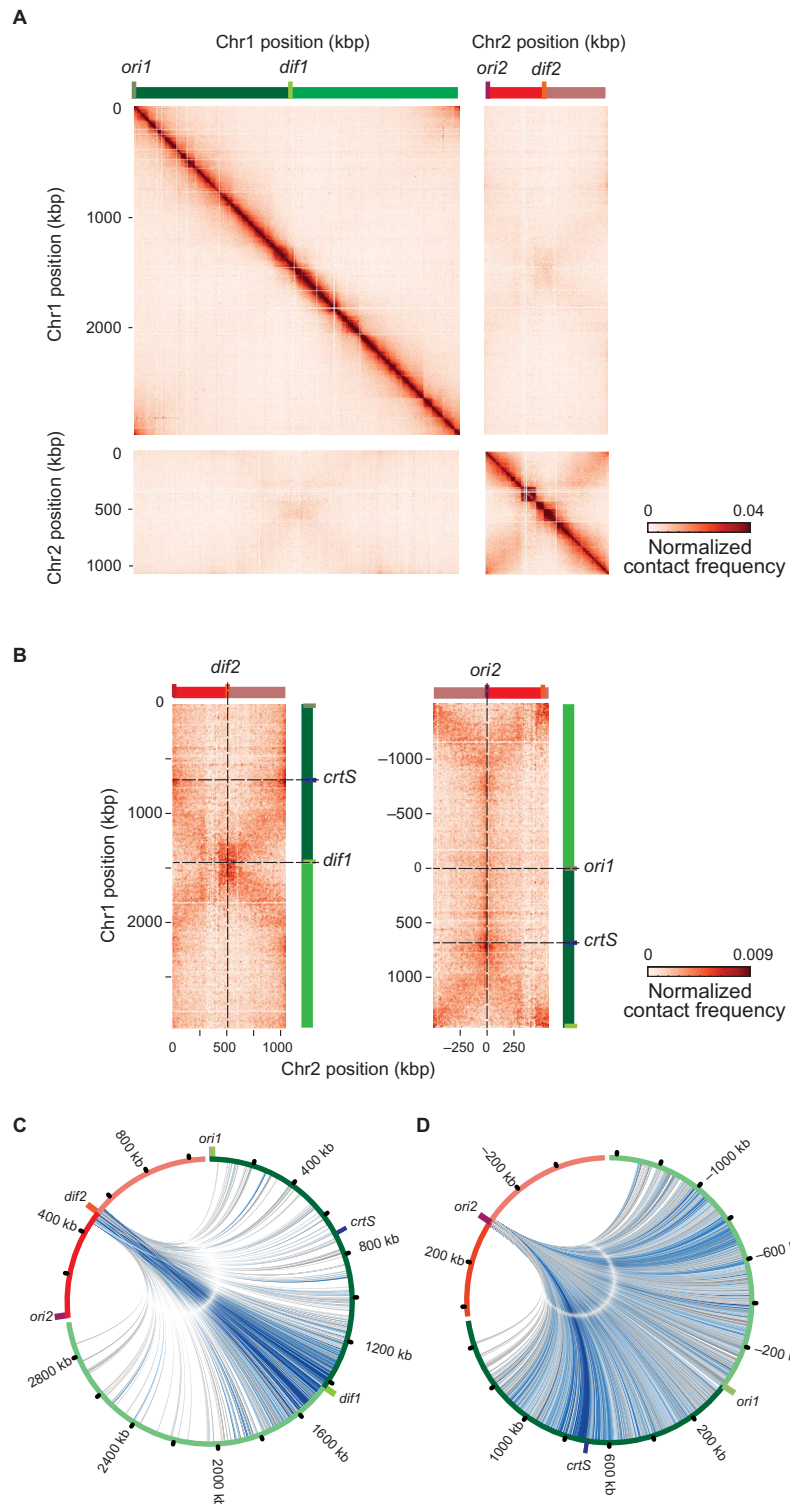
*crtS* in a *V. cholerae* strain where *ori2* was replaced by a second *ori1* locus and is thus no longer dependent on RctB (ICO1) (18). ICO1 $\Delta$ *crtS* mutants displayed no marked filamentous phenotypes and no additional growth defects compared to ICO1 (Fig. 6A, bottom, and fig. S11), confirming that the  $\Delta$ *crtS* phenotype is linked to a defi-

ciency in replication initiation of Chr2 at *ori2*. By performing MFA on mutant  $\Delta$ *crtS* #7 (with two separate chromosomes), we could detect the location of *ori2* by the overrepresentation of sequences (sharp peak), indicating that replication initiation still occurs at *ori2*. However, the MFA also highlighted an imbalance in the copy number of Chr1

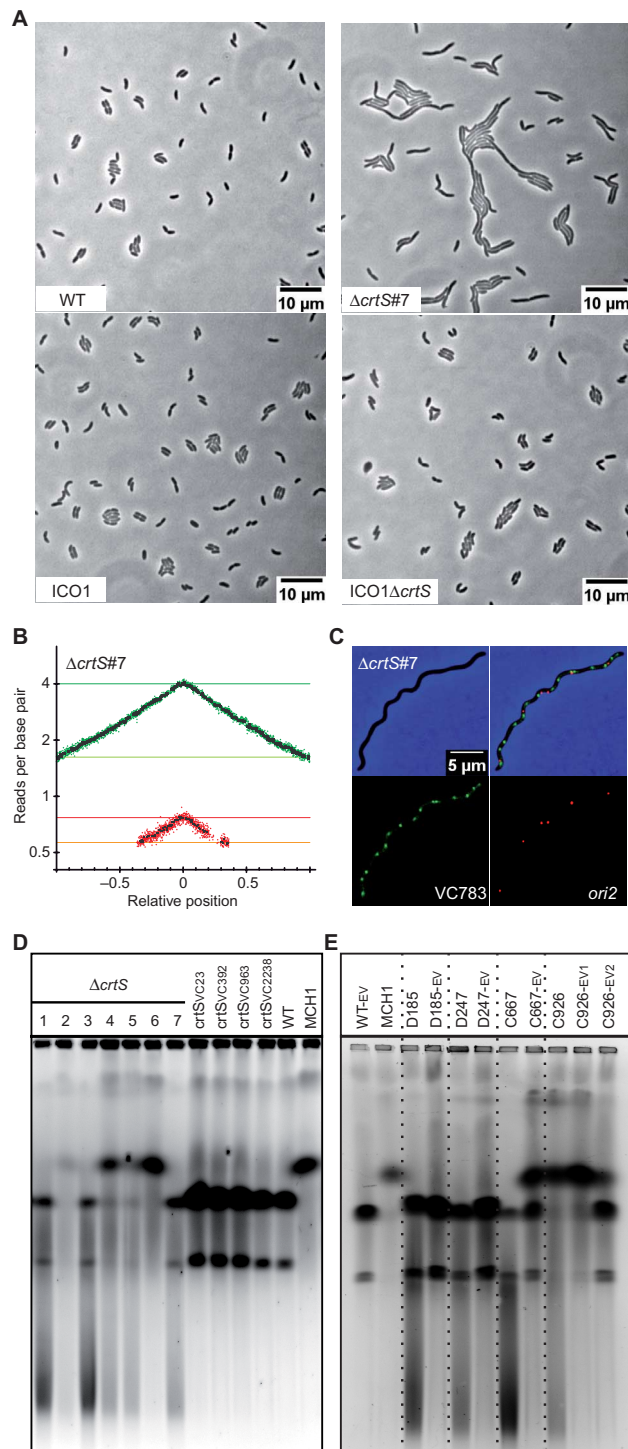


**Fig. 4. Two chromosomal copies of *crtS* doubles Chr2 copy number.** (A and B) Position of *ori1* (A) and *ori2* (B) foci inside *crtS<sup>WT/VC23</sup>* cells. Foci are oriented longitudinally relative to the old pole of the cell as a function of cell length. The old pole was defined as the closest pole to an *ori1* focus. On the left panel, the x axis represents the cell length (in micrometers). On the right panel, the x axis indicates cell number. The y axis represents the relative position of the focus in bacterial cells, 0 being the old pole and 1 the new pole. (C) Representative pictures of dividing cells (WT and *crtS<sup>WT/VC23</sup>*) observed by fluorescence microscopy. Cells were fluorescently labeled near *ori1* and *ori2*. Merged pictures of *ori1* (green) and *ori2* (red) and phase-contrast (blue) micrographs show 2x more red spots than green spots in mutant *crtS<sup>WT/VC23</sup>*. (D and E) Amount of *crtS<sup>WT/VC23</sup>* (D) and *crtS<sup>WT/VC2338</sup>* (E) cells exhibiting zero, one, two, three, four, or five and six *ori1* foci (left panel) and *ori2* foci (right panel) according to cell size (in micrometers).





**Fig. 5. Intra- and interchromosomal interactions in *V. cholerae*.** (A) Normalized and filtered genomic contact map obtained from an asynchronous population of WT cells growing exponentially in LB. x and y axes represent genomic coordinates of each chromosome centered on *dif* sites (light green bar, *dif1*; orange bar, *dif2*). Origins of replication are shown as a dark green bar (*ori1*) and a dark red bar (*ori2*). Chr1 and Chr2 are represented by dark green (right arm) or light green (left arm) and dark red (right arm) or light red (left arm). The color scale reflects the frequency of contacts between two regions of the genome (arbitrary units), from white (rare contacts) to dark red (frequent contacts), and is conserved across all panels of all figures. (B) Interchromosomal contact map centered on *dif* (left panel) or *ori* sites (right panel). The *crtS* site is indicated as a blue bar. (C and D) Circos representation of interactions of 100 kbp (20 bin) around *dif2* with Chr1 (C) and circos representation of interactions of 50 kbp (10 bin) around *ori2* with Chr1 (D).



**Fig. 6. *crtS* is crucial for Chr2 replication initiation at *ori2*.** (A) Phenotype of *crtS*-deleted mutants in WT and ICO1. Representative pictures of phase-contrast microscopy of live cells growing on LB agar pads: WT,  $\Delta crtS\#7$  (with two separate chromosomes), ICO1, and  $ICO1\Delta crtS$ . (B) MFA of  $WT\Delta crtS\#7$ . (C) Representative picture of a filamentous  $\Delta crtS$  cell observed with fluorescence microscopy. Loci near *crtS* (VC783) and *ori2* were fluorescently labeled. Merged pictures of VC783 (green) and *ori2* (red) and phase-contrast (blue) micrographs show a higher number of green spots than red spots. (D) Ethidium bromide-stained pulsed-field gel electrophoresis (PFGE) of native gDNA. From left to right: Independent clones of *crtS*-deleted mutants ( $WT\Delta crtS\#1$  to  $\#7$ ), mutants with relocated *crtS* ( $crtS_{VC23}$ ,  $crtS_{VC392}$ ,  $crtS_{VC963}$ , and  $crtS_{VC2238}$ ), and WT (two chromosomes) and MCH1 (one synthetic fused chromosome), which are used as size reference (18). (E) Same as (D). From left to right:  $WT_{-EV}$  (control), MCH1 (one synthetic fused chromosome), and independent clones of  $\Delta crtS$  mutants before (D185, D247, C667, and C926) and after a 200-generation evolution ( $_{-EV}$ ). For C926, samples were harvested after 100 ( $_{EV1}$ ) and 200 generations ( $_{EV2}$ ) to track the tendency of the fused chromosome to revert to two separate chromosomes.

and Chr2 with an *ori1/ori2* ratio of 5.2 and a *ter1/ter2* ratio of 2.9 (Fig. 6B). This imbalance was also observed by imaging fluorescently labeled Chr1 and Chr2 loci in a  $\Delta crtS$  mutant strain. Figure 6C shows a typical  $\Delta crtS$  filamentous cell with more VC783 foci than *ori2* foci, whereas in WT, the two loci duplicate around the same time (fig. S4E), and most cells display the same number of VC783 and *ori2* foci (fig. S4C). Time-lapse fluorescence microscopy of  $\Delta crtS$  cells shows that filaments grow and accumulate many Chr1 foci (movie S3). These filaments divide irregularly and occasionally give rise to a Chr2-less cell, which does not grow (movie S4).

PFGE revealed a high instability in the genome structure of  $\Delta crtS$  mutants. In many instances, the two chromosomes had spontaneously fused (Fig. 6D;  $\Delta crtS$  #2 and #4 to #6), and in all  $\Delta crtS$  mutants, a smear was clearly visible in the PFGE (Fig. 6D;  $\Delta crtS$ ), indicating either genomic DNA (gDNA) degradation or a high level of genomic instability. *crtS*-positional mutants, however, did not display gDNA degradation/instability, thus evidencing the proper activity of *crtS* when transposed to ectopic chromosomal positions (Fig. 6D;  $crtS_{VC23}$ ,  $crtS_{VC392}$ ,  $crtS_{VC963}$ , and  $crtS_{VC2238}$ ). The high instability observed in the genome of  $\Delta crtS$  mutants suggested a positive selection toward chromosome fusion. To test for this constraint, we grew four independent cultures of  $\Delta crtS$  mutants for 200 generations to investigate spontaneous fusion and other potential suppressors. PFGE analysis was performed on samples collected at the beginning and at the end of the experiment. At generation 0, three mutants (D185, D247, and C667) carried two chromosomes, whereas the fourth (C926) had already undergone a fusion. After 200 generations, both C926 and C667 displayed a mixed population of cells with fused and separate chromosomes, whereas D185 and D247 showed no sign of chromosome fusion (Fig. 6E). Evolved mutants significantly improved their fitness over 200 generations (fig. S12). Moreover, the DNA smear trademark of a  $\Delta crtS$  deletion was no longer detectable in any of the evolved mutants (D185<sup>-EV</sup>, D247<sup>-EV</sup>, C667<sup>-EV</sup>, and C926<sup>-EV2</sup>) (Fig. 6E). Microscopy and flow cytometry showed that D185<sup>-EV</sup> recovered a wild-type morphology with less than 1% of aberrantly sized cells (fig. S13). However, D247<sup>-EV</sup>, C667<sup>-EV</sup>, and C926<sup>-EV2</sup> populations still displayed some filamentation. MFA of D247, before experimental evolution, is very similar to the MFA of  $\Delta crtS$  #7 and shows that no genome rearrangement had yet occurred (fig. S14). However, MFA of D185 shows that gene amplification (triplication) had occurred before experimental evolution and reverted in the evolved mutant D185<sup>-EV</sup> (fig. S14). Stress-induced gene amplification has been observed in *E. coli* and may confer a selective advantage to cells under stress (29). After 200 generations, MFA of D185<sup>-EV</sup> and of D247<sup>-EV</sup> reveals an up-regulation of *ori2* initiation, presumably compensating for the lack of *crtS* activation, as shown by the increase in the Chr2/Chr1 copy number ratio compared to their parental strains (fig. S14). Such a remarkable rescue of phenotype of all  $\Delta crtS$  mutants over 200 generations suggested that compensatory mutations were required to restore, even partially, the growth defects. D185<sup>-EV</sup> whole-genome sequencing revealed a single mutation, leading to a F230S substitution in RctB, not present in D185. D247<sup>-EV</sup> genome sequence also revealed a single mutation (C to A transversion), 25 nucleotides upstream of the *rctB* start codon within the 29-mer RctB binding site, which is important for autorepression of RctB expression and *ori2* initiation negative regulation via handcuffing with iterons (30). Further analysis of the *ori2* regions of C667<sup>-EV</sup> and C926<sup>-EV2</sup> revealed two new single mutations in RctB, which were absent from their parental strain (table S1). In all four mutants,  $\Delta crtS$  phenotype was partially rescued by

*rctB*-related mutations, further supporting the direct connection between *crtS* and RctB.

## DISCUSSION

Approximately 10% of bacterial species have genomes split over multiple chromosomes. The machineries controlling replication initiation of secondary chromosomes are always of plasmid origin (2). An important difference between plasmids and chromosomes is that the latter replicate once, and only once, per cell cycle. In *V. cholerae*, the two chromosomes satisfy this rule; furthermore, they are known to have a synchronous termination of replication. However, it remains unclear how this is coordinated. Here, we show that the timing of Chr2 replication initiation directly depends on the position of a short intergenic sequence (*crtS*) on Chr1. Our results provide strong evidence that the replication of *crtS* triggers initiation of Chr2 replication. Additionally, we reveal preferential contacts between Chr1 and Chr2, suggesting the existence of mechanistic functional interactions between these two chromosomes. This study unravels a novel checkpoint control mechanism allowing the replication coordination of multiple chromosomes in bacteria.

### *crtS* is crucial for Chr2 replication

Here, deletion of *crtS* severely impairs growth and is associated with filamentation and DNA damage (Fig. 6A, top, and fig. S11). The large cell size heterogeneity in  $\Delta crtS$  mutants prevented statistically robust analysis based on fluorescence microscopy data. Nevertheless, filaments containing fewer Chr2 than Chr1 were consistent with MFA analysis showing Chr2 underrepresentation in  $\Delta crtS$  mutants (Fig. 6, B and C). In  $\Delta crtS$  filaments, we observed more VC783 foci than *ori2* foci, suggesting that failure of proper regulation/activation of Chr2 replication is responsible for the observed filamentation. It was reported that deletion of the *chrI-4* sequence ( $\Delta chrI-4$ ) led only to mild phenotypic changes (12).  $\Delta chrI-4$  mutants were also characterized by a delay in *ori2* foci duplication (12). Why  $\Delta crtS$  differs from  $\Delta chrI-4$  phenotype previously characterized could be explained by some differences in the way these sequences were deleted (see the Supplementary Materials). We further show that *crtS* is directly involved in the regulation of replication initiation at *ori2* through an RctB-associated mechanism because its deletion does not affect the physiology of the ICO1 mutant, in which Chr2 replication is no longer dependent on RctB at *ori2* (Fig. 6A, bottom). Strains with *crtS* relocated to different positions along Chr1 did not show any of the  $\Delta crtS$  phenotypes, indicating that the 150-bp *crtS* DNA sequence is both necessary and sufficient to trigger Chr2 replication initiation, independently of its genetic context (Fig. 6D and fig. S3).

### Replication of *crtS* orchestrates the replication of Chr1 and Chr2

The replication pattern of WT and mutant strains was investigated by MFA. WT MFA results show that the copy number of *ori1* is higher than that of *ori2*, whereas the copy numbers of *ter1* and *ter2* are similar (Fig. 1A), in agreement with former hypotheses and results, suggesting a synchronized termination (15). Displacing the *crtS* locus to different positions on Chr1 modified Chr1 and Chr2 replication synchronization (Fig. 2, A and B). The *ori1/ori2* ratio is perfectly correlated with the distance between *ori1* and *crtS* (Fig. 2C), demonstrating that Chr2 replication initiation timing is dependent on this parameter, with *crtS* replication triggering Chr2 replication. According to MFA, there is a

constant delay between *crtS* replication and the firing at *ori2* (Fig. 2C), which roughly corresponds to the replication of 200 kbp, suggesting that the transfer of information to *ori2* is not immediate. This delay may correspond to the time needed to activate RctB and the *ori2* initiation system. Therefore, both the position of *crtS* and this delay account for the Chr1 and Chr2 termination synchrony. The *crtS* position may have been selected throughout evolution by the constraint imposed by this activation delay.

### Transient genome remodeling suppresses *crtS* deletion

Spontaneous chromosome fusions are common suppressor events of *crtS* deletion, but these marked rearrangements are transient and can revert back to a two-chromosomal genomic configuration over time (Fig. 6, D and E). We previously showed that recombination-induced chromosome fusions occur spontaneously within wild-type populations of *V. cholerae* (31). These rearrangements are quickly reversed or outcompeted, probably as a result of slower growth linked to an increase in replication duration due to the larger chromosome size and/or metabolic cost due to gene dosage imbalance. Fused chromosomes can be stabilized upon depletion of Dam methylase, which is essential for Chr2 replication initiation (31). Here, we hypothesize that chromosome fusions are a transient and conveniently accessible suppressor step when confronted with *crtS* deletion, ensuring replication of Chr2 through piggybacking of the Chr1 replication machinery. The higher cost of this genome state would subsequently favor the acquisition and maintenance of compensatory mutations, permitting the restoration of the wild type-like two-chromosome structure. This was observed in mutant C926-*EV2*, which acquired a compensatory mutation in RctB (R195C) and resolved the chromosome fusion (Fig. 6E and table S1).

### RctB-associated mutations compensate $\Delta$ *crtS* by altering Chr2 initiation regulation

Our 200-generation evolution experiment results showed that the phenotype of  $\Delta$ *crtS* mutants was largely rescued by the selection of spontaneous mutations in the origin region of Chr2. Although the phenotypes of the evolved  $\Delta$ *crtS* mutants vary, they all involve an increase in fitness, a decrease in cell filamentation, and an absence of gDNA degradation compared to their parental strain (figs. S12 and S13). MFA of the evolved *crtS* mutants showed a higher Chr2/Chr1 copy number ratio, suggesting an up-regulation of *ori2* initiation (fig. S14). This rapid acquisition of suppressor mutations can explain why our observations on  $\Delta$ *crtS* mutants differ from those of Baek and Chattoraj (12). Considering the rapid genome remodeling and acquisition of compensatory mutations in *ori2* of  $\Delta$ *crtS* mutants, we speculate that the  $\Delta$ chrI-4 mutant previously characterized (12) has acquired compensatory mutations to overcome the marked effect of *crtS* deletion. All  $\Delta$ *crtS* compensatory mutations were directly related to RctB (table S1), including three in the coding sequence (R195C, F230S, and L357I). RctB is a large, relatively poorly characterized protein that presents a 70-AA C-terminal region (451 to 521) involved in both iterons and 39-mer binding, as well as in dimerization (8). Previous *rctB* mutants were selected in the *E. coli* heterologous host for causing replication overinitiation of mini-Chr2 (9, 32–34). These “copy-up” mutants of RctB show reduced dimer binding to iterons (32), reduced dimerization (33), or reduced 39-mer binding (32) or fail to bind ATP (9), all consistently preventing RctB to repress replication. Most of these mutations were found outside the 70-AA C-terminal region (34), implying that important functions for *ori2* replication control remains to be identified in RctB. One of our  $\Delta$ *crtS* suppressor mu-

tants (R195C) was already documented as a copy-up mutant (34). An additional mutation was found in the promoter region of RctB within the 29-mer RctB binding site, possibly impairing binding. Up-regulation of Chr2 replication initiation could result from the suppression of RctB autorepression leading to an increase in RctB expression, or from the abolition of the 29-mer negative regulatory function by handcuffing with iterons (30). The mutations obtained in our study were selected directly in *V. cholerae* on Chr2, therefore taking into account the native physiological levels of all partners interacting to regulate Chr2 replication initiation. The experimental evolution screening of *crtS* mutants provides a promising approach to unravel new regulatory factors involved in *V. cholerae* Chr2 replication.

### Chr1 and Chr2 termini cohesion before cell division

MFA shows that *V. cholerae* Chr1 and Chr2 finish replication at *ter1* (around *dif1*) and *ter2* (around *dif2*), respectively, in a nearly synchronous manner (Fig. 1A). In WT, both *ter1* and *ter2* are recruited early to mid-cell and remain together until the end of the cell cycle (fig. S5E), and 3C analysis shows strong interactions between the two *ter* regions (Fig. 5C). When Chr2 replication completes before Chr1 in *crtS*<sub>VC23</sub> mutant (Fig. 2B), *ter2* foci relocate earlier to mid-cell but remain at mid-cell until cell division, and segregate roughly at the same time as *ter1* foci, like in WT (fig. S5, E and F, and fig. S6). These results suggest that *ter1* and *ter2* localization at mid-cell, where septum formation occurs at the end of the cell cycle, is important to coordinate their proper segregation before cell division.

In bacteria, chromosome replication, segregation, and cell division are precisely orchestrated mechanisms. In *E. coli*, the Ter domain relocates to mid-cell as replication completes and becomes accessible to the divisome machinery for the final steps of cell division, including chromosome compaction, dimer resolution, and decatenation (35–38). MatP, a DNA binding protein, bridges distant *matS* sites within the Ter domain to organize it into a compact structure (39, 40) that interacts with the divisome to coordinate segregation and cell division (38). Chromosome dimer resolution at *dif* is mediated by the XerCD site-specific recombinases and the DNA translocase FtsK, which is anchored at the septum (36). In *V. cholerae*, MatP is known to play a role in Ter confinement of the two chromosomes at mid-cell after replication (41) and both Chr1 and Chr2 require the septal protein FtsK to resolve their dimers (42). Therefore, *ter1* and *ter2* cohesion at mid-cell may be important for their proper segregation, which may be coordinated by MatP or another mechanism that remains to be found.

### Proposed mechanisms that mediate the signal between *crtS* replication and *ori2* initiation

Our study shows that Chr2 initiation requires *crtS* replication (Fig. 2C), meaning that either DNA conformational changes caused by the passage of the replication fork across *crtS*, or the doubling in copy number of *crtS* after duplication triggers Chr2 replication. The binding activity of RctB to chrI-4 (*crtS*) was observed by ChIP-chip (12). It was reported that RctB binding could only be observed in vitro by DNase I footprinting when chrI-4 was carried by a supercoiled plasmid (12). Mutations of the protected bases abolished the enhancer activity of chrI-4, suggesting a direct interaction between chrI-4 and RctB (12). Baek and Chattoraj (12) suggested that chrI-4 could act as a DNA chaperone to remodel RctB to an active form with altered DNA binding activities. Indeed, by using a mini-Chr2 in *E. coli*, they show that addition of chrI-4 in trans abolishes the requirement for DnaK/J chaperones, which

are normally important to promote RctB-dependent initiation at *ori2* (12, 32). Initiator remodeling upon interaction with their cognate DNA binding sites has been observed for iteron-bearing plasmids (43), as well as for *E. coli* DnaA (44). RctB could form a bidentate protein simultaneously using two DNA binding domains to contact two DNA loci (for example, *crtS* and *ori2*). In mutants with two chromosomal copies of *crtS*, we observed that most of newborn cells have two *ori2* and only one *ori1* (Fig. 4 and fig. S8), suggesting that it is the duplication of the *crtS* sequence itself that triggers Chr2 replication initiation, and that the duplication of the single-copy *crtS* sequence in wild type is limiting initiation at *ori2*. Once *crtS* is duplicated, dimers of RctB could simultaneously contact two *crtS* sites, allowing a higher DNA binding affinity to iterons. In mutants with two chromosomal copies of *crtS*, a large proportion of cells divide with two *ori1* and four *ori2* foci, giving rise to newborn cells already having one *ori1* and two *ori2* foci (Fig. 4, C to E). Hence, passage of the replication fork across *crtS* might still be an option for triggering Chr2 replication if the signal had been sent in the mother cell. *crtS* contains two GATC sites that are substrates for Dam methylase. RctB binding is sensitive to the methylation state of iterons sites, which need to be fully methylated (45). The passage of the replication fork across *crtS* would generate transiently hemimethylated GATC sites that may affect RctB binding. However, mutation of the two GATC sites had no impact on *crtS* function. Passage of the replication fork also generates single-stranded DNA on the template for lagging-strand synthesis. RctB could recognize a DNA hairpin structure formed by the single-stranded state of *crtS*, thereby modifying its binding affinities for iterons and/or 39-mers. A better understanding of the genetic determinants enclosed within the *crtS* sequence will provide new clues to decipher the *crtS*-RctB-mediated mechanism that triggers Chr2 replication, and perhaps a better understanding of the mechanisms controlling the activation/deactivation of RctB.

A large fraction of *crtS*<sub>WT/VC23</sub> cells display three *ori2* foci, suggesting that one of the two *ori2* duplicates before the other (Fig. 4D). These observations are different from *E. coli*, where multiple origins initiate in synchrony (20). To accommodate for a generation time shorter than the replication time in fast-growing cells, overlapping rounds of replication occur with multiple origins of replication. In *E. coli*, multiple origins fire synchronously, such that either 2, 4, or 8 (that is,  $2^n$ ) origins of replication are present at the same time (20). In *E. coli*, several mechanisms are responsible for the coordinated initiation of multiple origins (DnaA titration, regulatory inactivation of DnaA, origin sequestration, and DnaA reactivation sequences) (46). All these mechanisms control the availability of the active form of DnaA for initiating replication from *oriC*. If the control of *ori2* initiation by *crtS* was done only by controlling the availability of the RctB active form, we would expect a similar synchrony in the firing of multiple *ori2*, and this would be observed by cells containing only  $2^n$  *ori2* foci (for example, 2 or 4). Because a large portion of *crtS*<sub>WT/VC23</sub> cells have three *ori2* foci, we hypothesize that the duplication of one *crtS* triggers the firing of only one *ori2*. This reinforces the hypothesis that firing could necessitate a contact between *crtS* and *ori2*. Fluorescence microscopy of WT fluorescently labeled near *crtS* (VC783) and *ori2* loci shows that the two loci localize to the same region throughout the cell cycle (fig. S4E). The contacts between *ori2* and Chr1 revealed by 3C may be caused by the simultaneous binding of RctB to *ori2* and *crtS*. The most frequent contacts between *ori2* and Chr1 occur downstream of *crtS* (Fig. 5D). A possible explanation is that, following the duplication of the *crtS* locus, the replication machineries of Chr1 and Chr2 are in the vicinity of each other

until the end of replication of the two chromosomes. Nonreplicating cells (that is, stationary phase) lose the cross-shape contacts observed between Chr1 and Chr2 replichoes during exponential growth (Fig. 5B, right), suggesting that replication is indeed responsible for the contacts of the two chromosomes along their chromosomal arms. Overall, the 3C analysis of the *V. cholerae* chromosomes points to a direct interplay between 3D organization and replication regulation. How trans topological contacts would drive a functional interaction between the two chromosomes remains unknown.

### Concluding remarks

Bacteria can be subjected to sudden environmental changes and must adapt their replication rate to their growth rate. Fast-growing bacteria often initiate multiple overlapping rounds of DNA replication. Inter-related mechanisms control the availability of the active form of DnaA to regulate chromosome initiation, some of which are modulated by growth rate (46). *V. cholerae* is a fast-growing bacterium that encounters a broad spectrum of habitats (for example, aquatic environments and human beings). Here, we describe a system for the replication control of multiple chromosomes. We demonstrate that Chr2 surveys the replication of Chr1 via the *crtS* locus and only initiates replication when this site has been replicated. Repositioning *crtS* closer to or further from the *ori1* causes Chr2 initiation to be initiated earlier or later, respectively, in the cell cycle. If the signal is not received (for example,  $\Delta$ *crtS* mutant), initiation of Chr2 will be impaired and the cell will filament, accumulating many copies of Chr1. We suggest this mechanism to be an example of a bacterial cell cycle checkpoint; coordination of passage from early cell cycle, with only Chr1 replicating, into the late part of the cell cycle, with both chromosomes replicating. The location of *crtS* on Chr1 dictates the timing of replication initiation at *ori2* so that replication of the two chromosomes terminate simultaneously. This, in turn, ensures coordinated and faithful segregation of the chromosomes in line with cell division. The *crtS*-RctB-mediated checkpoint control of Chr2 initiation is a simple and flexible mechanism to ensure a cell cycle-specific time setting of Chr2 replication relative to Chr1 replication. This novel mechanism is an elegant and cost-effective way for secondary chromosomes to benefit from the already well-adapted replication regulatory system of the host main chromosome.

Replication of eukaryotic genomes is initiated from multiple origins located on each chromosome, enabling the complete genomic replication within the S phase of the cell cycle (47). Thus, bacteria with multiple replicons may share similarities with eukaryotes in the control of their genomic replication. However, it is observed in eukaryotes that (i) not all origins are activated within a single replication round and (ii) activated origins do not start replication simultaneously (48). The observed tight control of initiation at *ori2* is different from the stochastic nature of initiation at eukaryotic origins (49). Nevertheless, we envisage that replication-dependent controls, such as the checkpoint described here, may contribute to the orchestration of the complex eukaryotic DNA replication.

## MATERIALS AND METHODS

### Experimental design

Our objective was to determine the regulatory pathway responsible for triggering Chr2 replication at a specific point of the cell cycle, so that the two chromosomes terminate replication at nearly the same time. To

achieve this, we created several *V. cholerae* derivatives with altered chromosome organization and monitored their replication pattern to determine whether this synchronization was directly linked to the replication of a specific region in Chr1. We used genome engineering, MFA, microscopy, and chromosome conformation capture to identify the elements involved in this regulatory pathway.

### Bacterial strains and plasmids

The bacterial strains and plasmids used in this study are listed in table S2. Large genome rearrangements were performed following the procedures described by Val *et al.* (18). Details are given in the Supplementary Materials.

### Quantitative PCR

qPCR was performed on the gDNA of cells growing exponentially in LB at 37°C [OD<sub>450</sub> (optical density at 450 nm), ~0.15]. Primers used to determine *ori1/ori2* ratios are listed in table S3. Details are given in the Supplementary Materials.

### Marker frequency analysis

MFA was performed on the gDNA of cells growing exponentially in LB (glucose) at 30°C (OD<sub>450</sub>, ~0.15). Libraries were sequenced using an Ion Proton sequencer (Life Technologies). MFA was performed essentially as described by Skovgaard *et al.* (17). Details are given in the Supplementary Materials.

### Pulsed-field gel electrophoresis

PFGE was done following the procedure described by Val *et al.* (10). Details are given in the Supplementary Materials.

### Fluorescence microscopy

All Chr1 loci were labeled with a *parS*<sub>PMT1</sub> site, and Chr2 loci were labeled with a *lacO* array. The genes encoding for yGFP-Δ30ParB<sub>PMT1</sub> and LacI-mCherry protein fusions were inserted in the *lacZ* gene using plasmid pAD19 (table S2) (19). Cultures for microscopy were grown in minimal fructose medium to limit replication rounds to once per cell cycle. Microscopy observations and data analysis were performed following procedures and using MATLAB scripts already described by David *et al.* (19). Details are given in the Supplementary Materials.

### Chromosome conformation capture

3C libraries were built, sequenced, and analyzed as previously described (23). For all the data, we generated matrices with bins of 5 kbp using a four-base cutter (Hpa II). Both matrices (LB and MM) showed a strong correlation together using Spearman's rank correlation coefficient ( $\rho = 0.488$ ,  $P < 10$  to 64). Experimental details are given in the Supplementary Materials.

## SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at <http://advances.sciencemag.org/cgi/content/full/2/4/e1501914/DC1>

#### Materials and Methods

fig. S1. Chromosomal inversion detected around *ori1* when WT sequences are mapped against the National Center for Biotechnology Information (NCBI) reference genome AE003852.

fig. S2. Large DNA inversions either caused no fitness cost (JB392) or were similarly affected (JB590, JB659, JB771, and JB963).

fig. S3. Complementation of Δ*crtS* filamentous phenotype by addition of an ectopic chromosomal copy of *crtS*.

fig. S4. Duplication and segregation of VC783 and *ori2* foci in WT and mutant *crtS*VC23 throughout the cell cycle.

fig. S5. Duplication and segregation of *ter1* and *ter2* foci in WT and mutant *crtS*VC23 throughout the cell cycle.

fig. S6. *ori2* foci duplicate earlier when *crtS* is located near *ori1*.

fig. S7. The addition of an extra copy of *crtS* affects the growth of *V. cholerae*.

fig. S8. Doubling in *ori2* copy number in mutant with two chromosomal copies of *crtS*.

fig. S9. Comparison of global chromosome organization of *V. cholerae* in different growth conditions.

fig. S10. Comparison of directional index analysis at a 100-kbp scale with transcription and GC content for fast-growing cells.

fig. S11. Δ*crtS* mutants have a fitness defect, whereas ICO1Δ*crtS* shows no additional growth defect.

fig. S12. Fitness improvement of Δ*crtS* mutants by the acquisition of compensatory mutations.

fig. S13. Loss of filamentation phenotype of Δ*crtS* mutants by the acquisition of compensatory mutations.

fig. S14. MFA of *crtS* mutants before and after acquisition of compensatory mutations.

fig. S15. The effect of MFA normalizations.

table S1. Compensatory mutations obtained after evolution of Δ*crtS* mutants.

table S2. List of plasmids and bacterial strains.

table S3. Primers used in qPCR.

movie S1. 3D representations of the contact map from fig. S9 (exponential growth of the WT in LB).

movie S2. 3D representations of the contact map from fig. S9 (exponential growth of the WT in MM).

movie S3. Time-lapse fluorescence microscopy of Δ*crtS* filamentous cells, tagged at VC783 (Chr1) and near *ori2* (Chr2), growing on an M9 MM agar pad supplemented with fructose and thiamine.

movie S4. Time-lapse fluorescence microscopy of Δ*crtS* filamentous cells, tagged at VC783 (Chr1) and near *ori2* (Chr2), growing on an M9 MM agar pad supplemented with fructose and thiamine.

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