Chromosome Segregation Proteins of *Vibrio cholerae* as Transcription Regulators

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**ABSTRACT** Bacterial ParA and ParB proteins are best known for their contribution to plasmid and chromosome segregation, but they may also contribute to other cell functions. In segregation, ParA interacts with ParB, which binds to *parS* centromere-analogous sites. In transcription, plasmid Par proteins can serve as repressors by specifically binding to their own promoters and, additionally, in the case of ParB, by spreading from a *parS* site to nearby promoters. Here, we have asked whether chromosomal Par proteins can likewise control transcription. Analysis of genome-wide ParB1 binding in *Vibrio cholerae* revealed preferential binding to the three known *parS1* sites and limited spreading of ParB1 beyond the *parS1* sites. Comparison of wild-type transcriptomes with those of Δ*parA1*, Δ*parB1*, and Δ*parAB1* mutants revealed that two out of 20 genes (VC0067 and VC0069) covered by ParB1 spreading are repressed by both ParB1 and ParA1. A third gene (VC0076) at the outskirts of the spreading area and a few genes further away were also repressed, particularly the gene for an outer membrane protein, *ompU* (VC0633). Since ParA1 or ParB1 binding was not evident near VC0076 and *ompU* genes, the repression may require participation of additional factors. Indeed, both ParA1 and ParB1 proteins were found to interact with several *V. cholerae* proteins in bacterial and yeast two-hybrid screens. These studies demonstrate that chromosomal Par proteins can repress genes unlinked to *parS* and can do so without direct binding to the cognate promoter DNA.

**IMPORTANCE** Directed segregation of chromosomes is essential for their maintenance in dividing cells. Many bacteria have genes (*par*) that are thought to be dedicated to segregation based on analogy to their roles in plasmid maintenance. It is becoming clear that chromosomal *par* genes are pleiotropic and that they contribute to diverse processes such as DNA replication, cell division, cell growth, and motility. One way to explain the pleiotropy is to suggest that Par proteins serve as or control other transcription factors. We tested this model by determining how Par proteins affect genome-wide transcription activity. We found that genes implicated in drug resistance, stress response, and pathogenesis were repressed by Par. Unexpectedly, the repression did not involve direct Par binding to cognate promoter DNA, indicating that the repression may involve Par interactions with other regulators. This pleiotropy highlights the degree of integration of chromosomal Par proteins into cellular control circuits.

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**M**any low-copy-number plasmids and bacterial chromosomes have *parAB* genes for segregating replicated sisters to opposite halves of dividing cells (1, 2). The products of *parAB* are two cis-acting proteins, ParA and ParB. *parS* is a cis-acting site which functions analogously to eukaryotic centromeres. ParA actively moves plasmids/chromosomes that have their *parS* bound by ParB.

In plasmids, the *parAB* genes comprise an operon, which is autorepressed either by ParA or ParB or by a ParA-ParB complex (1). When ParA serves as the repressor, it binds to operator sites unrelated to *parS*. This regulation can be further tightened by the participation of ParB and *parS* (3). In some plasmids, *parS* fulfills both the operator and centromere functions. The regulation of chromosomal *par* genes is known in *Streptomyces coelicolor*, where the *parAB* operon is autorepressed by ParB (4). This seems to be an isolated case, as *parS* sites are not usually found upstream of chromosomal *par* operons, and the domains of plasmid ParA proteins that specifically bind to operator sequences are usually missing from chromosomal ParA (1). The transcriptional regulation of chromosomal *par* genes remains largely unknown.

It is also not known whether Par proteins can regulate transcription of genes other than *parAB*. Such a possibility was suggested in *Bacillus subtilis*, where Soj and Spo0J (ParA and ParB homologues) appeared to control transcription of many sporulation genes (5). Subsequently, the results were reinterpreted to be due to Soj increasing the replication initiation activity of DnaA (6). The increase in DnaA activity turns on a cascade of events that lead to repression of sporulation genes (7). Thus, Soj control of transcription is considered to be indirect, occurring through DnaA.
ParB not only represses promoters that overlap parS but also represses promoters at a distance. The distal promoters are reached by spreading of ParB onto sequences that flank parS (8–11). The spreading can interfere with RNA polymerase interactions with promoter elements, a process termed (gene) silencing. The spreading has also been implicated in the control of DNA replication (8, 12, 13). The spreading can interfere with DNA-protein interactions involved in replication control, which can both promote and interfere with replication initiation, depending upon the situation.

Segregation, gene silencing, and DNA replication aside, Par proteins contribute to chromosome organization by loading condensin in the vicinity of the replication origin in B. subtilis and in Streptococcus pneumoniae (14–16) and contribute to cell cycle progression and cell division in Caulobacter crescentus (17, 18), cell growth in Mycobacterium smegmatis (19), cell growth and motility in Pseudomonas aeruginosa (20), cell morphology in Pseudomonas putida (21), and cell division in Streptomyces coelicolor (22). It is clear that chromosomal par genes play pleiotropic roles.

Here, we have investigated whether Par proteins control transcription in Vibrio cholerae, a bacterium with two chromosomes (chrI and chrII). Both chromosomes have their own parABS genes (parABS1 for chrI and parABS2 for chrII) (23). Their role in chromosome segregation has been studied in detail (24, 25). The nature of regulation of chromosomal parAB operons, or whether the Par proteins can control transcription of genes other than their own, is largely unknown. We show that ParB binds specifically only to parS sites and can spread to flanking DNA, which results in transcriptional silencing in a minority of cases. Additionally, we found that both ParA1 and ParB1 proteins could silence genes unlinked to parS, apparently without direct binding to promoter DNA. The possibility of involvement of other factors is suggested by the finding that both Par proteins can interact with several V. cholerae proteins.

RESULTS
ParB1 binding to parS1 sites and flanking DNA. We determined genome-wide ParB1 binding in V. cholerae N16961 (CVC796) using chromatin immunoprecipitation with microarray technology (ChIP-chip). When the amount of DNA precipitated by ParB1 antibody was compared with total DNA from the whole-cell extract, DNA in the region containing the three known parS1 sites of chrI (parS1-1, parS1-2, and parS1-3) was selectively precipitated, indicating preferential ParB1 binding to those sites (Fig. 1A; see also Table S1 in the supplemental material). This result is consistent with an earlier finding that identified the sites by bioinformatics analysis and yellow fluorescent protein (YFP)-ParB1 focus formation (23). The modal position of the immunoprecipitated DNA around these sites is shown in Fig. 1B. The data presented in this paper are consistent with the previous results obtained by bioinformatics analysis (23).

FIG 1 (A) ChIP-chip binding profiles of ParB1 in region containing the three parS1 sites (dashed lines) in chrI of V. cholerae. The profiles are shown for wild-type (in black), ΔparA1 (in green), and ΔparB1 (in blue) cells growing exponentially in LB broth. The fold change represents the amount of immunoprecipitated (IP) DNA normalized with respect to input DNA. The profiles represent average signals from three independent experiments. (B) Changes in expression levels of genes around the three parS1 sites (dashed lines) in V. cholerae ΔparA1 (green profile), ΔparB1 (blue profile), and ΔparAB1 (red profile) cells compared with those in WT cells growing exponentially in LB broth. The fold change values are log2 ratios of expression levels in mutants over the WT cells. Data presented are the averages from three independent experiments. The numbers in the boxes indicate the locus tag from VC0060 to VC0079 (abbreviated as 60 to 79). The white and black boxes indicate genes on plus and minus strands, respectively. The boxes outlined in red indicate the genes whose expression was considered changed at least 2-fold in the absence of ParA1 and/or ParB1. The genes 60 to 66 constitute an operon whose promoter is upstream of gene 60. The promoter activity of the operon appears not to be affected by Par proteins, although transcription elongation appears to have been reduced, most likely by opposing ParB1 spreading.
through the two outer peaks (apparently representing binding to DNA peaks roughly corresponded to the three parS1 sites, although the two outer peaks (apparently representing binding to parS1-1 and parS1-3 sites) were shifted slightly toward the central peak, which apparently represents binding to the parS1-2 site. The peaks spread wider (2 to 3 kb at half-maximal height) than the average DNA fragment length (0.5 kb). The peaks were narrower when a similarly fragmented DNA preparation was precipitated with RctB antibody (see Fig. S1) (26). The RctB protein is a site-specific DNA binding protein and is not known to spread along DNA outside its specific binding sites (12). These results are consistent with ParB1 spreading on either side of parS1 sites. The total length covered by ParB1 over three parS1 sites was about 16 kb (chr1 coordinates 59 to 75 kb), although the sites are located within a 7-kb region. This result is also consistent with spreading. The extent of Spo0J (ParB) spreading was about 18 kb in B. subtilis (9). Unlike the situation in B. subtilis, the parS1 sites in V. cholerae are located about 65 kb away from the origin, which is apparently too far to be reached by ParB1 spreading (14, 15).

We validated spreading of ParB1 in Escherichia coli by silencing of plasmid replication (Fig. 2). When the parS1-1 site was present in the pGB2 vector, transformants carrying such a plasmid could not be selected when ParB1 synthesis was induced. Under identical conditions, cells without induction of ParB1 could grow. In similar experiments, the growth failure was attributed to ParB spreading into the replication initiator gene and silencing its promoter, which is located only 200 bp away from parS (10, 27).

In order to determine the contribution of ParA1 to ParB1 binding and spreading, we performed ChIP-chip using a ΔparA1 strain (CVC797) (Fig. 1A). The ChIP-chip profile of the deletion strain was nearly identical to that of the wild-type (WT) strain, although the peak heights were reduced. This suggests that ParA1 might promote specific binding of ParB1. In contrast, no peaks could be seen in a ΔparB1 strain (CVC1123), as would be expected if the ChIP signals were due to ParB1 binding.

**Gene silencing around parS1 sites.** We performed transcriptome analysis to determine expression levels of genes near the parS1 sites. We focused on 20 genes that were present within the region covered by ParB1 spreading. Only three genes were expressed at levels greater than 2-fold in the ΔparB1 strain compared to their expression in the WT. The genes encoded aminopeptidase P (VC0067), a putative multidrug resistance protein (VC0069), and a universal stress protein A (VC0076) (Fig. 1B). The upstream regions of all three genes (without including the parS1 sites) were tested for ParB1 binding by electrophoretic mobility shift assay (EMSA), but no specific binding was detected (see Fig. S2A in the supplemental material). We also tested the regulation of expression of the three genes by ParB1 by fusing their upstream regions (without parS1) to a promoterless lacZ gene and measuring β-galactosidase activity in WT and ΔparB1 cells (see Fig. S2B). The absence of parB1 did not influence the activity of the first two genes (VC0067 and VC0069) and only marginally increased the expression of the third gene (VC0076). The results of the first two genes are consistent with parS1-dependent spreading and silencing by ParB1. The fact that the third gene is located farthest from the parS1 sites in a region where ParB1 spreading was barely significant yet showed maximal ParB1-mediated silencing suggests that the third gene is silenced by ParB1 without requiring parS1. Why only two out of 20 genes were silenced in the region of significant spreading remains to be determined. The results do indicate that spreading and silencing need not be correlated, as was the case in B. subtilis (9).

We also performed transcriptome analysis in ΔparA1 and ΔparAB1 mutants (Fig. 1B). The expression level of VC0067 and VC0069, but not of VC0076, increased in ΔparA1 cells compared to WT. This again indicates that regulation by Par proteins differs between the first two genes and the third gene. Whereas the silencing of the first two genes appears to require both ParA1 and ParB1, the regulation by ParB1 appears more significant. ParA1 could act by increasing ParB1 binding, as the ChIP-chip data suggest (Fig. 1A). Since ParA1 does not influence ParB1 concentration (28), ParA1 could affect the first two genes by increasing specific DNA binding of ParB1. However, this inference is not supported by the results in ΔparAB1 cells. If ParA1 were acting through ParB1, then the derepression levels in ΔparB1 and ΔparAB1 cells should have been comparable, but they are not. Thus, the regulation may involve factors other than Par proteins, whose activity may depend on ParA1.

ParB1 appears to be the main regulator of the third gene, VC0076, since the regulation did not differ much between WT and ΔparA1 cells (Fig. 1B). We conclude that the requirements of spreading and ParA1 are considerably less for VC0076 than for the other two genes.

**Gene silencing outside the ParB1 spreading region.** Transcriptome analysis also revealed that the expression levels of several genes outside the ParB1 spreading region can change in parA1, parB1, and parAB1 deletion strains (see Table S2 in the supplemental material). We selected six genes whose expression changed (up or down) most significantly in at least one of three mutants tested (Table 1). We validated expression level changes for these genes by fusing the gene regulatory regions to a promoterless lacZ gene. The change was most significant in the case of ompU (VC0633) in both transcriptome and lacZ fusion assays (Table 1). In the transcriptome analysis, the expression of the ompU gene increased in both ΔparA1 and ΔparB1 cells and syn-
ergistically in ΔparA1, ΔparB1, and ΔparAB1 cells. These results indicate that both ParA1 and ParB1 can independently repress the expression of ompU (as in VC0067 and VC0069) and that the two proteins might cooperate in their repressive activities on ompU (unlike the situation in VC0067 and VC0069). Likewise, the lacZ fusion assays in ΔparA1, ΔparB1, and ΔparAB1 cells revealed that the ompU promoter activity increased in all three deletion strains, confirming that both ParA1 and ParB1 are involved in the repression of the ompU promoter (Fig. 3A). The synergistic effect seen in the transcriptome analysis was not evident in the gene fusion assay, the reason for which remains to be understood. Both assays, however, suggest that ParA1 and ParB1 can control ompU expression.

In order to determine the region required for repression by ParA1 and ParB1, we fused various lengths of the ompU promoter region to lacZ and compared promoter activities between WT and ΔparAB1 cells (Fig. 3B). We found that the repression is most efficient when the ompU promoter fragment contains 118 bp upstream of the transcription start site.

Notably, ompU is one of the genes whose promoter is significantly activated by ToxRS proteins (29). The ompU promoter contains three ToxR binding sites in the upstream regulatory region. To determine if the regulation of the promoter by Par proteins depends on ToxRS, we determined the promoter activity with and without ToxRS in E. coli (Fig. 3C). As reported, the

| Gene          | Function                        | Expression levela | lacZ fusion |
|---------------|---------------------------------|-------------------|-------------|
| VC0321 (tufB) | Elongation factor Tu            | 4.23              | 2.85        | 1.42        | 1.56        |
| VC0470 (dns)  | Extracellular DNase             | 1.16              | 0.20        | 0.14        | 4.72        |
| VC0633 (ompU) | Outer membrane protein          | 4.86              | 3.89        | 120.26      | 2.13        |
| VC0706        | Sigma-54 modulation protein, putative | 0.52 | 45.89 | 93.70        | 1.95        |
| VCA0519 (fruR) | Fructose repressor             | 0.83              | 0.92        | 0.21        | 1.84        |
| VCA0676 (napF) | Iron-sulfur cluster-binding protein | 11.39             | 1.48        | 0.96        | 1.84        |

a Expression levels in mutant cells were in comparison to those in WT cells.
activity of the *ompU* promoter increased dramatically in the presence of ToxRS, but the repression by ParAB1 could be seen in both the absence and the presence of ToxRS. These results indicate that *ompU* promoter repression by ParAB1 does not require ToxRS.

**ParA1 and ParB1 interactions with other V. cholerae proteins.** Since direct binding of ParA1 and ParB1 without requiring parS1 in *cis* could not be demonstrated, we considered the alternative possibility that Par proteins function by interaction with other proteins. A candidate approach using the bacterial two-hybrid (B2H) assay confirmed that ParA1 and ParB1 do not interact with ToxR or *OmpU* (see Fig. S3 and Text S1 in the supplementary material). We showed above that Par regulation of *ompU* does not require ToxR and that the *ompU* gene is not autorepressed (data not shown). To identify interacting proteins, we took a global approach using both bacterial and yeast two-hybrid (Y2H) screening systems using ParA1 and ParB1 proteins as baits (see Table S3). We identified known interactions of ParA1 with ParB1, ParB1 with itself (28, 30), and ParA1 with HubP (VC0998) (30), indicating the effectiveness of our screening approach. The *cyaA* gene (VC0122) was also identified several times in our B2H screening system, which is based on reconstitution of CyaA activity. A number of other proteins were also found to interact with ParA1 and/or ParB1 (see Table S3). The relevance of these interactions in the regulation of *ompU* or other genes remains to be determined.

We note that although we used three different two-hybrid systems to achieve maximum coverage and reproduced several known interactions, overlapping interactions were not found among the three systems. Previous studies have shown that different versions of yeast two-hybrid (Y2H) vectors produce complementary interactions rather than overlapping interactions (31, 32), which may explain our findings. However, we cannot exclude the possibility of missing open reading frames (ORFs) among three different prey libraries.

**parAB1 operon is not autorepressed.** In plasmids such as P1 and F, the N-terminal region of ParA is important for transcriptional autorepression, but many chromosomal ParA proteins, including *V. cholerae* ParA1, lack this domain (1). How the chromosomal par genes are regulated remains unknown. To determine the regulation of *parAB1* genes in *V. cholerae*, we located the promoter for parA1 within the upstream sequence of the *gidB* gene, using the 5′ RACE (rapid amplification of cDNA ends) method (Fig. 4A). To determine whether the promoter is the only one that directs transcription of both parAB genes, we created three separate fusions to a gfp reporter gene. The region fused consisted of either the parAB1 genes with the promoter identified by 5′ RACE (parAB1p) or parAB1 genes without that promoter or only the parB1 gene. Measurement of fluorescence intensities revealed that only the parAB1 fragment with the promoter showed increased fluorescence compared to that of the vector without any insert (Fig. 4B). When the inserts were parAB1 and parB1, the fluorescence intensities were comparable to that of the vector-alone control. These results suggest that the parB1 gene is transcribed from the parA1 promoter and that parAB1 genes constitute an operon.

We then asked whether the *parAB1* operon is autorepressed, as is usually the case in plasmids (1). We fused the *parAB1* promoter to a promoterless *lacZ* gene present in pMLB1109 (resulting in pBJH17) and transferred it to the *E. coli* chromosome using phage λ (resulting in CVC1881). We determined the activity of the promoter both before and after induction of ParA1 only, ParB1 only, or both ParA1 and ParB1 (Fig. 4C). We previously showed that the induction increases the protein levels at least 16-fold (28). The promoter activity did not change upon induction of any of the proteins, indicating that ParA1 and ParB1 do not control their
own promoter. ChIP-chip analysis indicated that ParA1 and ParB1 proteins also did not bind in the vicinity of the parAB1 promoter (see Table S1 in the supplemental material). The transcriptome analysis also revealed that the levels of parA1 transcript in ∆parB1 cells and parB1 transcript in ∆parA1 cells remain comparable to those in the WT, indicating that expression of parAB1 genes is not controlled by ParA1 and ParB1 proteins (see Table S2). The expression levels of the parA1 gene in ∆parB1 cells and the parB1 gene in ∆parA1 cells were very low, as would be expected. We previously showed that ParA1 and ParB1 protein levels do not change in ∆parB1 and ∆parA1 strains, respectively, compared to their levels in the WT (28). From these results, we conclude that the parAB1 operon is not autorepressed.

We note that the activity of the parAB1 promoter was high in both E. coli and V. cholerae. A single-copy promoter fusion showed about 4,000 Miller units of activity in E. coli (Fig. 4C), and the same fusion in multicopy showed about 2,500 Miller units of activity in V. cholerae (data not shown). The parAB1 genes could also be expressed from the upstream gidAB promoter as revealed by a reverse transcription-PCR (RT-PCR) assay (see Fig. S4 in the supplemental material). However, the activity of the gidAB promoter was about 10-fold lower than that of the parAB1 promoter, and the gidAB promoter was also not regulated by ParA1 and/or ParB1 (data not shown).

DISCUSSION

Par proteins as transcription factors. Here, we show that in V. cholerae, parAB1 genes of chromosome I constitute an operon, which is usually the case in plasmids, but unlike the situation in plasmids, the operon is not autorepressed. Par proteins, however, can influence transcription of other genes by acting either separately or together. ParB1 silences VC0067 and VC0069 genes that are near parS1 sites, apparently by spreading. ParA1 could also influence the expression of these two genes either by reducing ParB1 binding to parS1 (Fig. 1A) or by unknown mechanisms. ParB1 can also repress genes not linked to parS1 (VC0076 and ompU). VC0076 was regulated mainly by ParB1, whereas ompU was regulated by both ParA1 and ParB1. To the extent studied, it is clear that the Par proteins can regulate transcription without necessarily requiring parS sites or other and, remarkably, without direct binding to the promoter region.

Direct binding was considered unlikely for the following reasons. ParA1 ChIP-chip data did not reveal any significant binding above background (data not shown). ParA1 also does not contain the specific DNA binding domains that are often present in plasmid ParA proteins (1). ParB1 binding that could be detected unambiguously by ChIP-chip (Fig. 1A; see also Table S1 in the supplemental material) and EMSA (see Fig. S2A) was absent from the upstream regions of genes that it represses.

The upstream regions of VC0067, VC0069, VC0076, and ompU genes do not reveal any sequence similarity that would suggest the presence of a binding site for a common regulator. We considered whether the indirect regulation is through VC0067 and VC0069, the only two genes that ParA1 and ParB1 control in a parS-dependent manner, and that they in turn regulate other genes. VC0067 and VC0069 function as aminopeptidase P and multidrug resistance proteins, respectively, which are unlikely to be transcription regulators. Orthologous genes (ypdF and mdtL, respectively) which do not have a parABS system are present in E. coli, and yet the V. cholerae ompU gene was regulated by ParA1 and ParB1 when tested in E. coli. These reasons lead us to think that the regulation is indirect, the possible mechanisms of which are discussed below.

Par proteins are known to interact with two global regulators. In B. subtilis and S. pneumoniae, Spo0J/ParB loads SMC protein near the replication origin, which has been suggested to organize the origin region of the chromosome (14, 15). SMC proteins are known to condense chromosomes in general and therefore can have a global effect on DNA transactions by changing DNA topology. The effect can be more pronounced in local regions, as for example the origin and terminus regions, because of interactions with proteins such as ParB and TopI/V, respectively (14, 15, 33). ParA (Soj) of B. subtilis also interacts with DnaA, which can turn on a checkpoint response that ultimately controls the expression of many sporulation genes (7). Both ParA and ParB have significant nonspecific binding and in principle can have a role in transcriptional control, but this role remains to be demonstrated.

Pleiotropy of chromosomal Par proteins. Plasmid and chromosomal Par proteins, although largely related, appear to be functionally distinct. The proteins are major players in plasmid segregation, whereas their contribution to chromosome segregation is often modest (2, 34, 35). The main distinguishing feature of chromosomal Par proteins is their ability to affect functions besides segregation, as discussed in the introduction. Here we show that ParA1 and ParB1 proteins control expression of genes that have been implicated in the control of drug resistance (VC0069), the stress response (VC0076), and pathogenesis (VC0633) of V. cholerae. In addition to interaction with SMC and DnaA, ParB binding to MipZ has been demonstrated to regulate cell division in C. crescentus (17, 18). Thus, DNA binding aside, protein-protein interactions contribute to the pleiotropy of Par proteins. Several proteins are found to interact with Par proteins in B2H and Y2H screening systems. Although these interactions need to be validated by independent methods, they suggest that there is a wider spectrum of cell functions regulated by Par proteins, involving both global and specific regulators.

Conservation of Par functions. Here, we show that V. cholerae Par proteins share many of the properties of Par proteins seen in other bacteria. For example, ParB1 binds in a site-specific manner to its cognate centromeric sites and can spread to neighboring sequences, as was initially shown in B. subtilis (36). In both bacteria, ParA1 has little contribution to the spreading process. Spreading can silence some, but not all, genes encountered by spreading. As in B. subtilis, ParA1 and ParB1 do not regulate their own genes. The similarities, particularly with B. subtilis, suggest that the characteristics of chromosomal Par proteins have been conserved since the two bacteria diverged more than a billion years ago. Recently, Par proteins were shown to be global transcription regulators in P. aeruginosa (37). In this study, both ParA and ParB were shown to regulate many genes, apparently indirectly.

MATERIALS AND METHODS

Strains and plasmids. V. cholerae and E. coli strains and plasmids used in this study are listed in Table S4 in the supplemental material.

ChIP-chip assay. ChIP-chip assays were performed using a custom Agilent 8-by-60K V. cholerae oligonucleotide microarray, as described previously (12). Vibrio cholerae cells were cultured in LB broth at 37°C to exponential phase and cross-linked with 1% formaldehyde for 30 min at room temperature. ParA1-DNA or ParB1-DNA complexes were immunoprecipitated using ParA1 or ParB1 antibody (Biosource International), respectively. Fold changes were calculated by dividing precipitated DNA with...
(Cy5) signals by the total input (Cy3) signals from three independent experiments.

Transcriptome analysis. Vibrio cholerae cells were cultivated in LB broth at 37°C to exponential phase. Total RNA from wild-type, ΔparA1, ΔparB1, or ΔparAB1 cells was purified using RNAProtect bacterial reagent and RNeasy mini kits (Qiagen) according to the manufacturer’s protocol. Using the purified RNA, CDNA was synthesized, labeled, and hybridized to the Agilent V. cholerae microarray as described elsewhere (38). Fold changes were calculated by dividing the signals from deletion mutants (Cy5) by the signal from the WT (Cy3) from three independent experiments.

Plasmid replication assay. Transformants containing an inducible source of parB1 (pBJH15) and either pB2 vector or its parS1–1–carrying derivative (pB2H105) were grown on LB agar plates at 37°C for 1 day under drug selection for both plasmids. Isopropyl-β-D-thiogalactopyranoside (IPTG) (0.5 mM) was added to the plates when induction of ParB1 was desired (12).

Transcription assay by reporter gene fusion. Promoter activities were measured using promoter-cloning vectors with a reporter gene, which was either lacZ (pMLB1109) or gfp (pFBH65), in exponentially growing cells in LB broth as described elsewhere (12). Fluorescence intensities were measured using a Victor2 (Wallac) microplate reader, and β-galactosidase activities were determined as described previously (39).

5’ RACE. The 5’3’ RACE kit was used according to the manufacturer’s protocol (Roche). CDNA was synthesized and amplified using primers BJH114 and BJH115 (see Table S5 in the supplemental material), respectively. RNP-protect bacterial reagent and RNeasy mini kits (Qiagen) were used to purify RNA. The pGEM-T Easy vector system (Promega) was used to clone amplified cDNA.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at http://mbio.asm.orglookup/suppl?doi=10.1128/mBio.01061-14/-/DCSupplemental.

Text S1, DOCX file, 0.2 MB.
Figure S1, TIF file, 3 MB.
Figure S2, DOCX file, 4 MB.
Figure S3, TIF file, 14 MB.
Figure S4, TIF file, 6.3 MB.
Table S1, PDF file, 7.1 MB.
Table S2, PDF file, 0.6 MB.
Table S3, DOCX file, 0.1 MB.
Table S4, DOCX file, 0.1 MB.
Table S5, DOCX file, 0.1 MB.

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