The GATC-Binding Protein SeqA Is Required for Bile Resistance and Virulence in *Salmonella enterica* Serovar Typhimurium[⊽]†

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Received 21 July 2007/Accepted 14 September 2007

Disruption of the seqA gene of Salmonella enterica serovar Typhimurium causes defects similar to those described in E. coli: filament formation, aberrant nucleoid segregation, induction of the SOS response, envelope instability, and increased sensitivity to membrane-damaging agents. Differences between SeqA⁻ mutants of E. coli and S. enterica, however, are found. SeqA⁻ mutants of S. enterica form normal colonies and do not exhibit alterations in phage plaquing morphology. Lack of SeqA causes attenuation of S. enterica virulence by the oral route but not by the intraperitoneal route, suggesting a virulence defect in the intestinal stage of infection. However, SeqA⁻ mutants are fully proficient in the invasion of epithelial cells. We hypothesize that attenuation of SeqA⁻ mutants by the oral route may be caused by bile sensitivity, which in turn may be a consequence of envelope instability.

The ability of Salmonella enterica to induce disease in animals relies on the possession of virulence genes, some of which are clustered in pathogenicity islands while others are scattered in the chromosome (16). In Salmonella subspecies I, some virulence genes are plasmid borne (33). In addition to virulence genes, whose products are required only for animal infection, certain housekeeping genes play relevant roles in the interaction of Salmonella with its animal host (15). Examples are two-component systems, such as PhoPQ and RcsBCD, which regulate virulence gene expression in specific stages of the infection process but also play relevant physiological roles outside the animal (34). Another example of a housekeeping function with a role in host-pathogen interactions is DNA adenine (Dam) methylation, which is essential for virulence in Salmonella and other bacterial pathogens (6, 21, 26, 47).

Mutants of S. enterica serovar Typhimurium lacking Dam methylation are severely attenuated in the mouse model, especially by the oral route (12, 18). Virulence attenuation caused by dam mutations is pleiotropic and involves reduced invasion of epithelial cells (12), lowered cytotoxicity after infection of M cells (12), inefficient colonization of Peyer's patches and mesenteric lymph nodes (12, 18), sensitivity to bile and to other DNA-damaging agents produced inside the animal (19, 31), and envelope instability accompanied by leakage of proteins (32). A transcriptomic study of Dam methylation in S. enterica serovar Typhimurium has provided evidence that some of these virulence defects are caused by altered patterns of gene expression (2). However, Dam methylation is not a regulator of gene expression per se but a signaling system for DNA-protein interactions (47), and the proteins that regulate Dam-dependent expression of Salmonella virulence genes remain to be identified.

SeqA was discovered as a protein involved in the methylation/hemimethylation cycle of Escherichia coli DNA (27). SeqA binds to hemimethylated GATC sites formed by DNA replication and regulates activation of the E. coli chromosome replication origin (27). Proper chromosome segregation also requires SeqA (1). Furthermore, SeqA trails the DNA replication fork and may contribute to nucleoid organization in newly replicated DNA (4, 24, 25, 48). Aside from its roles in chromosome replication and nucleoid segregation, SeqA is known to regulate the transcription of certain genes. In bacteriophage lambda, SeqA activates the $p_{\rm R}$ promoter in a GATC methylation-dependent fashion (41). SeqA also acts as a transcriptional coactivator by facilitating binding of the cII transcription factor to the lambda p_{I} and p_{aO} promoters (40). Competition between SeqA and the OxyR repressor for hemimethylated GATC sites has been shown to regulate phase variation in the E. coli agn43 gene (9). These examples raised the possibility that SeqA binding to critical GATC sites might likewise regulate the expression of S. enterica genes.

Here, we examine the consequences of seqA gene disruption in Salmonella enterica serovar Typhimurium and describe analogies and differences between E. coli and S. enterica SeqA⁻ mutants. Of special interest is the observation that disruption of the seqA gene in S. enterica serovar Typhimurium causes attenuation of virulence by the oral route, a trait that may be correlated with sensitivity to bile salts encountered in the host intestine. Transcriptomic analysis does not provide evidence for SeqA-mediated regulation of known virulence genes in Salmonella. In fact, lack of SeqA causes minor changes in the global transcriptional profile of the cell, as previously described in E. coli (25). However, an intriguing observation is that the relatively short list of SeqA-regulated loci in S. enterica includes a significant number of Salmonella-specific genes.

JOURNAL OF BACTERIOLOGY, Dec. 2007, p. 8496-8502 0021-9193/07/\$08.00+0 doi:10.1128/JB.01156-07 Copyright © 2007, American Society for Microbiology. All Rights Reserved.

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^{*} Supplemental material for this article may be found at http://jb .asm.org/.

⁷ Published ahead of print on 28 September 2007.

TABLE 1. Strain list

^a Omitted for strains described in this study.

MATERIALS AND METHODS

Bacterial strains, plasmids, and phages. The strains of Salmonella enterica used in this study (Table 1) belong to serovar Typhimurium and derive from the mouse-virulent strain ATCC 14028. An exception is DA6522, an LT2 derivative kindly provided by Dan I. Andersson, University of Uppsala, Uppsala, Sweden. E. coli DH5a (49) was the standard host for recombinant plasmids. E. coli CC118 (20) was kindly provided by Victor de Lorenzo, CNB, CSIC, Cantoblanco, Spain. Plasmid pGE108 (Km^r) is a ColE1 derivative carrying a cea::lacZ fusion (35). Plasmid pIZ1574, constructed for this study, is a pBluescript II SK(+) derivative containing the seqA coding sequence from strain ATCC 14028, previously amplified by PCR (see below). Plasmid pIZ1588 is a pBAD18 derivative expressing the S. enterica seqA gene from the arabinose-dependent P_{BAD} promoter (17) and was constructed by cloning an XbaI-HindIII fragment from pIZ1574 onto pBAD18. Transductional crosses using phage P22 HT 105/1 int201 (36; G. Roberts, unpublished data) were used for strain construction operations involving chromosomal markers and for transfer of plasmids among Salmonella strains. The transduction protocol has been described elsewhere (13). To obtain phagefree isolates, transductants were purified by streaking them on green plates (7). Phage sensitivity was tested by cross-streaking them with the clear-plaque mutant P22 H5. Strain SV4861 was constructed by transducing the invJ::Kmr allele from SB302 to ATCC 14028. SB302 is a derivative of strain SL1344 of S. enterica serovar Typhimurium (23), obtained from J. E. Galán, Yale University, New Haven, CT.

Media and chemicals. Luria-Bertani broth (LB) was used as a rich medium. E medium of Vogel and Bonner (45) was used as a minimal medium. Unless otherwise indicated, the carbon source for both LB and E medium was 0.2% glucose. Solid media contained agar at 1.5% final concentration. Deoxycholic acid (sodium salt) and sodium choleate (ox bile extract) were both from Sigma Chemical Co., St. Louis, MO. Antibiotics were used at the final concentrations described previously (44). Green plates were prepared according to the method of Chan et al. (7), except that methyl blue (Sigma) was substituted for aniline blue.

MICs of DOC, ox bile extract, and ethanol. Exponential cultures in LB were prepared as previously described (31). Samples containing around 3×10^2 CFU were transferred to polypropylene microtiter plates (Soria Genlab, Valdemoro, Spain) containing known amounts of sodium deoxycholate (DOC), ox bile extract, or 96% ethanol. After 12 h of incubation at 37°C, growth was visually monitored. Assays were carried out in triplicate.

Construction of an S. enterica SeqA- mutant. The seqA gene from strain ATCC 14028 of S. enterica serovar Typhimurium was amplified by PCR using two 30-nucleotide primers: 5'ATG CGA ATT CAG GGC GGA ATG GTT TCC CAG 3' and 5' ATG CGA ATT CAG CGC ATT GGC GCG ATC TTC 3'. The resulting 1.8-kb fragment, which contained both the seqA coding sequence and the promoter region, was digested with EcoRI and ligated to a dephosphorylated pBluescript II SK(+) vector. Transformation of E. coli DH5 α and a search for Lac- colonies permitted the detection of pBluescript derivatives carrying the seqA insert. One such clone was then used to generate a deletion in the seqA coding sequence. Because seqA is the upstream gene in a two-cistron transcriptional unit that also includes the pgm gene (28), an in-frame deletion 300 bp long was generated to prevent polarity. The deletion was obtained by PCR using primers 5' CGT AAG ATC TCG CAC GCG CGT CTA TTT TG 3' and 5' GCT GAA GAT CTG CTA TAG AGT CTA TCA TCA AC 3'. The resulting fragment was digested with BgIII and ligated to obtain a deleted pBluescript derivative that contained only 243 bp (81 codons) of the seqA coding sequence. The deleted pBluescript derivative was introduced into E. coli DH5a by transformation. Its insert was then excised with KpnI and SacI, cloned onto pDMS197 (11), and propagated in E. coli CC118 \lapla pir (20). One pMDS197 derivative containing the $\Delta seqA$ allele was transformed into E. coli S17-1 λ pir (38). The resulting strain was used as a donor in matings with S. enterica ATCC 14028, selecting Tcr

transconjugants on E medium plates supplemented with tetracycline (30 mg/ liter). Several Tc^r transconjugants were grown in nutrient broth (without NaCl) containing 5% sucrose. Individual tetracycline-sensitive segregants were then examined for the incorporation of the $\Delta seqA$ allele. Replacement of the chromosomal *seqA* gene by the $\Delta seqA$ construct was detected by PCR amplification, using the same pair of oligonucleotides employed to generate the deletion. One such isolate was propagated as strain SV4752 ($\Delta seqA1$).

RNA isolation, microarray procedures, and data analysis. To prepare cells for RNA extraction, 25 ml of fresh LB was inoculated with a 1:100 dilution from an overnight bacterial culture and grown in a 250-ml flask incubated with shaking at 250 rpm in a New Brunswick Innova 3100 water bath at 37°C. Three biological replicates were performed for each strain, and RNA was extracted at an optical density at 600 nm (OD₆₀₀) of ~0.5 to 0.6 (exponential phase) and ~3 (stationary phase). Specifically, the $\mathrm{OD}_{600} \mathrm{s}$ for RNA extractions from exponential-phase cultures of the wild type were 0.53, 0.52, and 0.53; for exponential-phase cultures of SV4752, 0.60, 0.64, and 0.62; for stationary-phase cultures of the wild type, 3.10, 3.21, and 3.19; and for stationary-phase cultures of SV4752, 3.18, 3.20, and 3.15. RNA extractions were performed as described by Mangan et al. (29), and their quality was assessed on an Agilent 2100 Bioanalyzer. Transcriptomic analyses were performed on a DNA microarray engineered by the Salgenomics consortium of research groups. The Salgenomics microarray contained 6,992 probes (including open reading frames, RNA genes, and intergenic regions) from the genome sequence of S. enterica SL1344 and was developed using genome sequences from the Wellcome Trust Sanger Institute (ftp://ftp.sanger .ac.uk/pub/pathogens/Salmonella). Use of an SL1344 microarray was justified because the ATCC 14028 genome is not completely sequenced; however, pairwise comparisons of sequenced genes indicated that SL1344 and ATCC 14028 are closely related strains (data not shown). Hybridization, microarray scanning, and data analysis were performed as described elsewhere (2; J. Mariscotti and F. Garcia-del Portillo, submitted for publication). Raw transcriptomic data are provided in Tables S1 and S2 in the supplemental material.

Preparation of protein extracts. Bacteria were grown overnight in LB medium at 37°C in nonshaking, static cultures. Under these growth conditions, the final OD₆₀₀ reached ~1.0. Bacteria contained in 10 ml of the culture were centrifuged at 15,000 × g for 10 min at 4°C. The supernatant was filtered with a 0.45-µmpore-size Millipore device, and the extracellular proteins were precipitated with 10% trichloroacetic acid, washed in acetone, and resuspended in cold phosphate-buffered saline, pH 7.4, as previously described (32). Bacteria present in the pellet were directly suspended in cold phosphate-buffered saline, pH 7.4. Finally, an appropriate volume of sodium dodecyl sulfate (SDS)-Laemmli buffer was added to both fractions, extracellular soluble proteins and total bacterium-associated protein, and boiled for 5 min at 95°C. Upon centrifugation (15,000 × g; 5 min; room temperature), the volumes to be load into the gels were adjusted to 1 × 10⁸ bacterial cells (total bacterium-associated proteins) or 5 × 10⁹ bacterial cells (estracellular soluble proteins).

CI virulence assays. Eight-week-old female BALB/c mice (Charles River Laboratories, Santa Perpetua de Mogoda, Spain) were used for virulence tests. Groups of three or four animals were inoculated with a 1:1 ratio of two strains. The bacteria were grown overnight at 37°C in LB without shaking. Oral inoculation was performed by feeding the mice with 25 µl of saline (0.9% NaCl) containing 0.1% lactose and 108 bacterial CFU. Bacteria were recovered from the mouse spleens 6 days after inoculation, and the CFU were enumerated on appropriate media. A competitive index (CI) for each mutant was calculated as the ratio between the wild type and the mutant strain in the output divided by their ratio in the input (3). To compare the virulence of a double mutant with that of a single mutant, a "cancelled-out" CI (COI) was calculated. A COI is the ratio between the double mutant and the single mutant in the output divided by their ratio in the input (3). Assays were carried out in triplicate. Student's t test was used to analyze CIs and COIs. The null hypothesis was that CIs were not significantly different from 1. COIs were analyzed with two null hypotheses: (i) the mean COI is not significantly different from 1 and (ii) the mean COI is not significantly different from the CI of the corresponding single mutant. P values of 0.01 or less were considered significant.

 β -Galactosidase assays. Levels of β -galactosidase activity were assayed as described by Miller (30), using the CHCl₃-sodium dodecyl sulfate permeabilization procedure.

Bacterial infection of epithelial cells. Bacteria were grown overnight in LB medium at 37°C under static, nonshaking conditions. HeLa epithelial cells were seeded the day before infection in 24-well plates and grown at 37°C and 5% CO_2 to obtain 80% confluence. Bacteria were added to reach a multiplicity of infection of 10 bacteria/eukaryotic cell. Simultaneous infection with two strains permitted the calculation of a CI for invasion (CII) (37). HeLa cells were infected for 30 min to obtain an infection rate of 70 to 80% of the cell population.



FIG. 1. Photomicrographs of SeqA⁺ and SeqA⁻ S. *enterica* cells during exponential growth, visualized by the fluo-phase combined method of Hiraga et al. (22).

Infected cells were washed three times with phosphate-buffered saline and incubated in fresh tissue culture medium containing 100 mg/liter gentamicin. At 2 h postinfection, the concentration of gentamicin was lowered to 10 mg/liter. The numbers of viable intracellular bacteria were obtained by lysing infected cells with 1% Triton X-100 and subsequent plating. CII calculations and statistical analysis were as described previously (37).

Microscopic observation of cells and nucleoids. Cells were harvested by centrifugation, washed, and resuspended in saline. A small aliquot was spread on a glass slide and dried at room temperature. Nucleoid staining with DAPI (4',6diamino-2-phenyl-indole) and cell visualization followed the "fluo-phase" combined method of Hiraga et al. (22).

RESULTS

Phenotypes of S. enterica SeqA⁻ mutants. In E. coli, SeqA⁻ mutants form very small colonies (27). In contrast, the $\Delta seqA1$ allele did not alter the morphology of S. enterica colonies. The absence of a growth difference associated with a seqA mutation was also observed in liquid cultures: the growth rates of an S. enterica SeqA⁻ strain in both LB and minimal E medium remained largely unaltered (data not shown). Another difference between SeqA⁻ mutants of E. coli and Salmonella was found in phage-plaquing experiments. While phage lambda has been shown to form large plaques on E. coli (46), the lambdoid phage P22 HT and its clear-plaque derivative, H5, formed plaques of similar sizes on both 14028 (seq A^+) and SV4752 $(\Delta seqA1)$. An effect of SeqA on lysogenization, as described for lambda (46), was also ruled out for P22: the plaques formed by P22 HT on strains ATCC 14028 (seq A^+) and SV4752 (Δ seqA1) of S. enterica were identical (small and turbid), and the percentage of clear plaques was below 0.01% in both cases (data not shown). Aside from these differences, SeqA⁻ mutants of S. enterica resemble those of E. coli in the following traits. (i) Nucleoid staining with DAPI and cell visualization following the "fluo-phase" combined method of Hiraga et al. showed filament formation and aberrant nucleoid segregation (Fig. 1). Similar phenotypes have been described in E. coli SeqA⁻ mutants (27). (ii) A pBluescript derivative carrying the wild-type seqA gene (pIZ1574) could not be maintained in a Dam⁻ strain (SV4698). The same construct was easily introduced in a Dam⁺ isogenic strain (ATCC 14028). In contrast, expression of seqA from an arabinose-driven P_{BAD} promoter permitted the introduction of a pBAD-seqA plasmid (pIZ1588) in both Dam⁺ and Dam⁻ hosts. However, a Dam⁻ host carrying pIZ1588 formed normal colonies only in the absence of arabinose (and at low arabinose concentrations). From these observations, we concluded that overproduction of Salmonella SegA is lethal in a Dam⁻ background, as described for *E. coli* (27). (iii) A SeqA⁻ mutant of S. enterica (strain SV4972) showed moderate SOS induction, as previously described for E. coli

TABLE 2. MICs of DOC, ox bile extract, and ethanol

Strain	Genotype	MIC (% [wt/vol])		
		DOC	Ox bile extract	Ethanol
14028 SV4752 SV4698	Wild type ΔseqA1 dam-225::MudJ	6 0.8 0.4	14 6 5	4 0.2 NT ^a

^a NT, not tested.

(27). Induction of the S. enterica SOS system was monitored using a cea::lac fusion carried on plasmid pGE108 (35). This fusion has been extensively used in our laboratory to monitor SOS induction in Salmonella (14, 31). The presence of a seqA mutation in the background increased cea::lac expression three- to fourfold: 40 \pm 16 Miller units in ATCC 14028/ pGE108 versus 188 \pm 23 Miller units in its isogenic derivative, SV4972 (averages of four independent experiments). (iv) In E. coli, the viability of SeqA⁻ mutants requires induction of the SOS system, since the cell division inhibitor SulA is necessary for survival (27). In an analogous fashion, we were unable to transduce the lexA(Ind⁻) allele of DA6522 to SV4972, suggesting that an S. enterica SeqA⁻ LexA(Ind⁻) mutant is nonviable. (v) The antecedent that seqA mutations cause sensitivity to DOC in E. coli (46) suggested the possibility that SeqA⁻ mutants of Salmonella might likewise be sensitive to bile salts. Bile salt sensitivity was tested as previously described (31). The results, summarized in Table 2, indicated that lack of SeqA renders S. enterica sensitive to both DOC and ox bile extract. The MICs of both DOC and bile were similar to those previously described for Dam⁻ strains (31, 32). (vi) Because E. coli SeqA⁻ mutants are sensitive to ethanol (46), we tested whether their Salmonella counterparts had a similar pheno-



FIG. 2. Lack of SeqA causes exacerbated release of proteins into the extracellular medium. TP, bacterium-associated total protein; SP, extracellular soluble protein present in the culture supernatant. The lanes corrrespond to the following strains: 1, wild type (ATCC 14028); 2, Dam⁻ (SV4698); 3, SeqA⁻ (SV4752); 4, Dam⁻ SeqA⁻ (SV4784). Samples loaded in the gel were adjusted to 1×10^8 bacterial cells (TP) or 5×10^9 bacterial cells (SP).



FIG. 3. Graphical representation of CI analyses of SeqA⁻ (SV4752), Dam⁻(SV4698), and SeqA⁻ Dam⁻ (SV4784) strains after oral or intraperitoneal infection of BALB/c mice. The mixed infections performed were as follows: (a) wild type/SeqA⁻, oral infection; (b) wild type/Dam⁻, oral; (c) Dam⁻ SeqA⁻/Dam⁻, oral; (d) SeqA⁻/Dam⁻ SeqA⁻, oral; (e) wild type/SeqA⁻, intraperitoneal; (f) wild type/Dam⁻, intraperitoneal. The indices represented are the means from \geq 3 infections. The error bars represent the standard deviations.

type. The MIC of ethanol was found to be 20-fold reduced in a $SeqA^-$ background (Table 2).

Envelope instability in *S. enterica* **SeqA⁻ mutants.** In previous studies, we showed that DNA adenine methylase mutants of *S. enterica* secrete large amounts of proteins into the culture medium, a trait that can be easily observed in SDS-PAGE gels from culture supernatants (32). The extreme sensitivity to bile and to membrane-damaging agents shown by SeqA⁻ mutants raised the possibility that they could have a similar phenotype. Figure 2 shows an SDS-polyacrylamide gel electrophoresis gel in which the profiles of extracellular proteins present in supernatants of SeqA⁻, Dam⁻, and SeqA⁻ Dam⁻ strains were compared with that of the wild type. Lack of SeqA seems to cause exacerbated release of proteins into the extracellular milieu, to a degree similar to that of Dam⁻ mutants.

Virulence assays with S. enterica SeqA⁻ mutants. The observation that a SeqA⁻ mutant of S. enterica was sensitive to bile raised the possibility that a seqA mutation might impair Salmonella virulence. BALB/c mice were subjected to mixed infections by inoculating groups of three or four animals with a 1:1 ratio of SeqA⁺ and SeqA⁻ isogenic strains. CI analysis indicated that a seqA mutation caused moderate but significant attenuation of virulence by the oral route but had no effect by the intraperitoneal route (Fig. 3). This observation suggests that the SeqA⁻ mutant is impaired for virulence at the intestinal stage of infection and can be tentatively correlated with the sensitivity to bile salts conferred by a *seqA* mutation. Our finding that lack of SeqA does not attenuate Salmonella virulence by the intraperitoneal route disagrees with a previous report (8). However, it must be noted that the seqA::MudI1734 allele used by Chatti et al. (8) is known to be polar on the downstream gene pgm (5), while we employed a nonpolar, in-frame deletion allele of seqA. Lack of phosphoglucomutase, the product of the pgm gene, in fact has a mild attenuating effect on Salmonella virulence, both orally and intraperitoneally (D. A. Cano and J. Casadesús, unpublished data). This may also explain why the seqA::MudI1734 allele causes a stronger attenuation by the oral route than the $\Delta seqA1$ allele (8).

COI analysis with isogenic SeqA⁻ and SeqA⁻ Dam⁻ strains



FIG. 4. Computer-generated normalized representation of microarray data. The Cy5/Cy3 fluorescence intensity of each locus in the wild type (horizonal axis) is plotted versus the intensity of the same locus in the mutant (vertical axis). Most genes fall near the imaginary bisectrix (slope = 1), indicating identical or similar expression. Genes above the bisectrix show higher expression in the mutant strain, indicating that SeqA represses their expression. Genes below the bisectrix show lower expression in the mutant strain, indicating that SeqA activates their expression.

indicated that a *dam* mutation was epistatic over a *seqA* mutation, both orally and intraperitoneally. The latter observation can be explained by the well-known fact that the SeqA protein does not bind to unmethylated DNA (47). Hence, whatever the function of SeqA in the infection process, it cannot be exerted in the absence of methylated or hemimethylated GATCs. COI analysis confirmed the attenuation of SeqA⁻ mutants by the oral route, since the SeqA⁻/Dam⁻ SeqA⁻ COI was >10-fold lower than the wild-type/Dam⁻ CI (Fig. 3).

The observation that *S. enterica* SeqA⁻ mutants are less virulent upon oral inoculation raised the possibility that lack of SeqA might impair invasion of the intestinal epithelium. This possibility was investigated by comparing the invasion rates of cultured HeLa cells by SeqA⁺ and SeqA⁻ isogenic strains. A noninvading InvJ⁻ mutant (SV4861) was included in these experiments as an internal control. The average invasion rates

 TABLE 3. S. enterica serovar Typhimurium genes with impaired expression in a SeqA⁻ background identified by transcriptomic analysis in exponential-phase cultures^a

Gene	Protein function	SeqA ⁻ mutant/ wild-type expression ratio
glnP	Glutamine ABC transporter	5.53
	permease component	
STM4212	Putative phage tail core protein	2.21
STM1329	Putative inner membrane protein	0.43
STM2911	Putative permease	0.38
STM0284	Putative Shiga-like toxin A subunit	0.38
STM2137	Putative cytoplasmic protein	0.38

^{*a*} SeqA is an activator for genes with relative expression values below 1.0 and a repressor for genes with values above 1.0.

obtained in standard 2-h infection assays of HeLa cells were similar for both SeqA⁺ and SeqA⁻ strains (CII = 0.911) and > 50-fold lower (CII = 0.019) in SV4861 (InvJ⁻), indicating that a *seqA* mutation does not impair invasion of epithelial cells.

Transcriptomic analysis of SeqA-dependent gene expression. Wild-type ATCC 14028 and its isogenic SeqA⁻ derivative, SV4752, were grown in LB medium, and RNA extraction for microarray analysis was carried out in exponential- and stationary-phase cultures. Use of these two conditions was justified by the fact that SeqA binds to methylated DNA, found in both exponential- and stationary-phase cells, and hemimethylated DNA, present in dividing cells only (48). The experiments were based on the rationale that higher RNA content in a SeqA⁻ background would indicate that SeqA represses gene expression in the wild type. In turn, lower RNA content in a SeqA⁻ background would indicate that SeqA activates gene expression in the wild type. As shown in the computer image diagrams in Fig. 4, the RNA contents for most S. enterica loci were similar in SeqA⁺ and SeqA⁻ hosts, in both exponentialand stationary-phase cultures, suggesting that SeqA-regulated genes are rare in Salmonella. A similar conclusion was drawn from a SeqA transcriptomic study of E. coli (25). A number of SOS genes showed moderately increased expression in a SeqA⁻ background, as previously described for *E. coli* (25). Aside from SOS genes, a small number of genes showed differences higher than the twofold cutoff arbitrarily chosen as a threshold. In exponential-phase cultures (Table 3), two SeqArepressed genes and four SeqA-activated genes were found. In stationary-phase cultures (Table 4), 4 SeqA-repressed genes and 32 SeqA-activated genes were found. A noteworthy observation is that a significant proportion (16/42) of such genes are Salmonella-specific and in most cases have unknown functions. The existence of six SeqA-regulated genes with unkown functions that are not Salmonella specific is likewise intriguing, since it raises the putative number of SeqA-regulated genes with unknown functions to about half of the total detected.

DISCUSSION

The cell division abnormalities observed in SeqA⁻ mutants of *S. enterica* closely resemble those previously described in their *E. coli* counterparts, which have been the subject of more

 TABLE 4. S. enterica serovar Typhimurium genes with impaired expression in a SeqA⁻ background identified by transcriptomic analysis in stationary-phase cultures^a

Gene	Protein function	SeqA ⁻ mutant/ wild-type expression ratio
rpsH	30S ribosomal protein S8	2.94
cbiT	Precorrin-8w decarboxylase	2.84
STM3098	Putative transcriptional regulator	2.70
STM2133	Putative cytoplasmic protein	2.69
misL	Putative autotransporter	0.40
kdpA	Potassium-transporting ATPase subunit A	0.40
ydhL	Putative oxidoreductase	0.39
glgP	Glycogen phosphorylase	0.39
STM2590	Tail assembly protein I like	0.39
argE	Acetylornithine deacetylase	0.39
gudT	Putative D-glucarate permease	0.39
csgG	Putative curli operon transcriptional regulator	0.38
STM0053	Putative transcriptional regulator	0.38
<i>yrfF</i>	Putative inner membrane protein	0.38
rpsA	30S ribosomal protein S1	0.38
STM2609	DNA packaging-like protein	0.38
yeaR	Putative cytoplasmic protein	0.38
ygiN	Putative cytoplasmic protein	0.37
menF	Menaquinone-specific isochorismate synthase	0.37
STM2743	Putative cytoplasmic protein	0.37
STM1671	Putative regulatory protein	0.37
STM4493	Putative cytoplasmic protein	0.37
chaB	Cation transport regulator	0.36
yijP	Putative integral membrane protein	0.36
STM4192	Putative cytoplasmic protein	0.36
deoA	Thymidine phosphorylase	0.36
yidE	Hypothetical protein	0.35
avrA	Secreted effector protein	0.35
sbcC	ATP-dependent double-stranded DNA exonuclease	0.35
ybeU	Putative cytoplasmic protein	0.34
STM4493	Putative cytoplasmic protein	0.34
glgP	Glycogen phosphorylase	0.34
STM4015	Putative cytoplasmic protein	0.33
adi	Catabolic arginine decarboxylase	0.32
yeeY	Putative transcriptional regulator	0.30
narJ	Nitrate reductase 1 delta subunit	0.24

^{*a*} SeqA is an activator for genes with relative expression values below 1.0 and a repressor for genes with values above 1.0.

numerous and deeper studies (1, 4, 10, 25, 27, 39, 46, 48). Other analogies between *E. coli* and *Salmonella* SeqA⁻ mutants include SOS induction, requirement for the SOS regulon for viability, envelope instability, and sensitivity to bile salts and other envelope-damaging agents. Given these analogies, the observation that a *seqA* mutation does not alter the morphology of P22 plaques is anecdotal and should not blur the tentative conclusion that the well-known roles of SeqA in *E. coli* chromosome replication and nucleoid organization (1, 4, 25, 27, 39, 48) can be tentatively extended to a relatively close bacterial genus like *Salmonella*. This view is consistent with the 87% amino acid identity found between the *E. coli* and *Salmonella* SeqA proteins (data not shown).

An interesting trait of *Salmonella* SeqA⁻ mutants is their virulence attenuation by the oral route. In vitro assays have indicated that SeqA⁻ mutants are not impaired for invasion of

epithelial cells, suggesting that attenuation is not caused by deficient interaction with the intestinal epithelium. Furthermore, global gene expression (microarray) analysis has not unveiled altered expression of any known locus involved in *Salmonella* infection (either at the intestinal stage or during systemic infection). Hence, a tentative explanation for oral attenuation in SeqA⁻ mutants may be their sensitivity to bile salts. During passage through the small intestine, *Salmonella* faces periodic release of bile, and the extreme sensitivity of SeqA⁻ mutants to bile salts may cause massive cell killing in the *Salmonella* population with concomitant reduction of virulence. In fact, SeqA⁻ mutants are not attenuated upon intraperitoneal infection, which spares the bacterial population from exposure to bile in the host intestine.

The cause of bile sensitivity associated with *seqA* disruption remains unknown. In *E. coli*, SeqA⁻ mutants show altered membrane permeability (46) and abnormal phospholipid composition (10), which may explain their increased sensitivity to a number of dyes. Our observation that the envelope of *S. enterica* SeqA⁻ mutants is likewise unstable can be tentatively correlated with bile sensitivity, because unconjugated bile salts can enter the cell by diffusion (43). How a *seqA* mutation can impair membrane stability is an interesting enigma. In *E. coli*, SeqA has been shown to associate with the *E. coli* outer membrane in a replication cycle-dependent manner (39). Thus, a structural role of SeqA in envelope stability cannot be discounted (46). An alternative explanation is that SeqA might regulate the expression of genes involved in the stability of the cell membranes, a possibility also considered in *E. coli* (42).

Our conclusion that SeqA does not play a major role in gene regulation in S. enterica is in agreement with a previous study carried out with E. coli (25). This view does not exclude the possibility that SeqA could play subtle roles, hardly visible by transcriptome analysis, in the expression of certain genes. For instance, the SeqA/OxyR competition described in the E. coli agn43 gene (9) would likely be overlooked in a microarray study. Another possibility, not investigated here, is that SeqA might control bacteriophage P22 promoters, as shown for the $p_{\rm R}$, $p_{\rm I}$, and $p_{\rm aQ}$ promoters of bacteriophage lambda (40, 41). Despite these caveats, our study has detected a small number of Salmonella genes whose expression is impaired in a SeqA⁻ background. The list includes both loci activated by SeqA and loci repressed by SeqA, and one-third of such genes (16/42) are Salmonella specific. However, impaired gene expression in a SeqA⁻ background does not imply a role for SeqA in the control of transcription. A well-documented example of this kind involves DNA adenine methylase mutants of E. coli and Salmonella, which show increased expression of the SOS regulon without any direct effect of Dam methylation on LexAcontrolled transcription (25, 47). In a similar fashion, the existence of SeqA-regulated genes might reflect physiological changes occurring in response to a seqA mutation rather than a direct role of SeqA in transcriptional control.

ACKNOWLEDGMENTS

This work was supported by grants BIO2004-3455-CO2-02 and GEN2003-20234-CO6-03 (to J.C.) and BIO2004-3455-CO2-01 and GEN2003-20234-CO6-01 (to F.G.D.P.), all from the Spanish Ministry of Education and Science and the European Regional Fund. A.I.P. was supported by a predoctoral fellowship from the Fundación Ramón Areces. M.J. and I.S. were supported by FPI predoctoral fellowships

from the Spanish Ministry of Education and Science. M.G.P. is an investigator supported by a contract of the Ramón y Cajal program of the Spanish Ministry of Education and Science.

We thank Dan I. Andersson, Victor de Lorenzo, and Jorge Galan for providing bacterial strains and Roberto Balbontín for advice on microarray experiments.

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