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DNA topology and adaptation of Salmonella typhimurium to an intracellular environment

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The expression of genes coding for determinants of DNA topology in the facultative intracellular pathogen *Salmonella typhimurium* was studied during adaptation by the bacteria to the intracellular environment of J774A.1 macrophage-like cells. A reporter plasmid was used to monitor changes in DNA supercoiling during intracellular growth. Induction of the dps and spv genes, previously shown to be induced in the macrophage, was detected, as was expression of genes coding for DNA gyrase, integration host factor and the nucleoid-associated protein H-NS. The topA gene, coding for the DNA relaxing enzyme topoisomerase I, was not induced. Reporter plasmid data showed that bacterial DNA became relaxed following uptake of *S. typhimurium* cells by the macrophage. These data indicate that DNA topology in *S. typhimurium* undergoes significant changes during adaptation to the intracellular environment. A model describing how this process may operate is discussed.

Keywords: intracellular growth; DNA topology; gene expression; adaptation; Salmonella typhimurium

1. INTRODUCTION

acterial adaptation to stressful environments is the subject f intense investigation at present and studies performed in itro have helped to identify many of the components used y prokaryotes to survive when under stress. Much of this ork has been carried out with the facultative intracellular athogen, Salmonella typhimurium (see §5). The central nportance of gene regulation to stress responses is obvious nd a great deal of detailed information is available about he processes that control gene expression at the transcriponal and post-transcriptional levels. Almost all of this iformation has been acquired through work with the acteria under in vitro conditions, although attempts have een made to extrapolate from the in vitro work to the in ivo situation. The recent development of techniques that ermit stress responses to be studied directly in the in vivo U tuation has accelerated the pace of this field.

An overview of gene regulatory processes reveals ontrols that operate at a 'local' level (to regulate indiviual promoters, etc.) and others that play a more general r 'global' role. This report deals with regulatory mechanims of the latter class. It is concerned with determinants f the topology of bacterial DNA, each of which has been nown *in vitro* to be capable of modulating the transcriponal profile of the cell. These are DNA gyrase, a type II popisomerase that introduces negative supercoils into NA, an activity that is unique to prokaryotes; DNA opoisomerase I, an enzyme that relaxes DNA and which

acts antagonistically to gyrase; integration host factor (IHF), a sequence-specific DNA-binding protein that places 180° bends into DNA and modulates the function of many bacterial promoters, as well as influencing transposition, site-specific recombination, DNA replication and other DNA transactions; H-NS, a nucleoid-associated protein that binds DNA in a sequence-independent manner (it is thought to recognize and bind at, or close to, regions of intrinsic curvature in DNA) and can regulate transcription from a large number of promoters (almost always negatively) as well as influencing other DNA reactions such as site-specific recombination. In addition to the genes coding for these global regulators, this study also encompasses two well-characterized stress response promoters, those of the plasmid-located spv virulence genes and of the chromosomally linked dps gene that expresses a DNA protection system in starved cells.

2. DNA TOPOISOMERASES

Gyrase is composed of two subunit proteins, GyrA and GyrB, and has an A_2B_2 tetrameric structure. It requires ATP to supercoil DNA negatively (Gellert *et al.* 1976) and this requirement links gyrase activity to the physiological state of the cell. Thus, when bacteria experience certain stresses, such as an upshift in osmolarity or a transition from aerobic to anaerobic growth, gyrase activity is altered and the result is a change in the level of supercoiling in the DNA (Dorman *et al.* 1988; Higgins *et al.* 1988; Hsieh *et al.* 1991*a,b*; Jensen *et al.* 1995; Van Workum *et al.* 1996). Supercoiling imparts free energy to DNA and

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his drives structural transitions in the DNA helix, neluding open complex formation at certain promoters Drlica 1992). In this way, many promoters located in ifferent parts of the genome can respond simultaneously o an alteration in the external environment. The promoers of the gyrase genes are supercoiling responsive. They re inhibited by increases in negative superhelicity but ecome induced when DNA is relaxed (Menzel & Gellert 983, 1987*a*,*b*). Thus, gyrase expression can respond to hanges in supercoiling at the level of transcription and yrase activity can respond to fluctuations in the [ATP]: ADP] ratio at the level of topoisomerase activity.

Another gene with a supercoiling-sensitive promoter is pA, which codes for topoisomerase I (Tse-Dinh 1985). This monomeric type I topoisomerase relaxes DNA that as become supercoiled past a critical point (Wang 1971). It is thought that the countervailing activities of topoiomerase I and DNA gyrase establish a homeostatic ipercoiling balance in the cell (DiNardo *et al.* 1982; fenzel & Gellert 1983). Topoisomerase I does not have n ATP requirement and uses energy stored in the egatively supercoiled DNA to drive the DNA relaxation eaction (Drlica 1992). The *topA* promoter is activated then DNA supercoiling increases and is inhibited by elaxation (the opposite of the *gyr* gene promoters) (Tse-Dinh 1985).

3. INTEGRATION HOST FACTOR

The IHF is encoded by two unlinked genes, *ihfA* and If B. The protein has a heterodimeric AB structure and inds to the consensus sequence WATCAANNNNTTR where W is a pyrimidine, R is a purine and N is any base). HF binding introduces a bend of up to 180° at the binding te and this is thought to be central to its biological role Ellenberger & Landy 1997; Rice et al. 1996; Travers 1997). t promotes long-range interactions in DNA and between roteins bound at distant sites on the same DNA molecule Goosen & Van de Putte 1995; Nash 1996). IHF is required or the formation of nucleoprotein complexes such as the ambda intasome (Goodman & Nash 1989; Snyder et al. 989) and the type 1 fimbrial invertasome (Blomfield *et al.*) 997; Dorman & Higgins 1987; Eisenstein et al. 1987), both n Escherichia coli. Expression of the *ihf* genes is subject to omplex control; they respond to growth phase, RpoS, uanosine tetraphosphate and are subject to autoregulaon (Aviv et al. 1994). (RpoS is an alternative sigma factor sed by RNA polymerase when bacteria undergo stress Hengge-Aronis 1996); guanosine tetraphosphate is an Ularmone that is synthesized by bacteria during starvation Cashel et al. 1996).) Furthermore, the ihfA gene is part of $\overline{\mathcal{S}}$ n operon with the phenylalanine tRNA synthetase genes, heST. Although *ihfA* has its own promoter, it is also ubject to coregulation with *pheST* (Mechulam *et al.* 1987; 4iller 1984).

4. PROTEIN H-NS

Protein H-NS is a component of the bacterial nucleoid. t binds to DNA and can constrain supercoils. Its ligomeric structure is a matter of disagreement in the terature, although it is probably at least tetrameric Spurio *et al.* 1997). H-NS can form heteromeric complexes with a closely related paralogue, StpA (Cusick & Belfort 1998), and this may be important for the biological roles of both proteins (Dorman et al. 1999; Zhang et al. 1996). H-NS influences the transcription of many genes. Usually it acts as a repressor and the genes under its control usually have other, specific regulators (Atlung & Ingmer 1997; Bertin et al. 1999; Ussery et al. 1994; Williams & Rimsky 1997). Some of these are transcription activators, which directly oppose the negative influence of H-NS (Jordi et al. 1992). H-NS-responsive genes have little in common, apart from a general contribution to the ability of the bacterium to adapt to environmental stress. The mechanism of action of H-NS is a matter of controversy, and it is likely that more than one mechanism of gene regulation is employed. In several cases an association of H-NS binding with a region of DNA curvature has been reported (Yamada et al. 1991), but data from in vitro experiments have caused the importance of curvature in H-NS binding to be questioned (Jordi et al. 1997).

5. SALMONELLA TYPHIMURIUM

S. typhimurium is a facultative intracellular pathogen and is a useful model for studying gene expression during *in vivo* growth (Finlay & Falkow 1997; Gulig 1996; Jones 1997). A great deal of research has been conducted *in vitro* into the response of S. typhimurium to stress at the level of gene expression (Alpuche-Aranda *et al.* 1992; Foster & Spector 1995; Groisman & Saier 1990). Recently, this work has been extended by a number of *in vivo* studies using reporter gene fusions to stress-regulated promoters. These investigations have identified several genes as being activated during *in vivo* growth or as coding for products that are essential for survival while the bacterium is within the host (Heitoff *et al.* 1997; Hensel *et al.* 1995; Valdivia & Falkow 1997).

Investigations performed with bacteria grown in vitro have illustrated the contributions made to stress responses by genes whose products modulate DNA topology (Dorman 1995). We wished to study the responses of these genes to intracellular growth as a first step in elucidating their involvement in bacterial adaptation to in vivo growth. The work was carried out in the murine macrophage-like J774A.1 cell line (American Type Culture Collection, Manassas, VA, USA), which has been used extensively for the study of intracellular growth (Buchmeier et al. 1993; Francis & Gallagher 1993; Rhen et al. 1993; Uchiya et al. 1999; Wilson et al. 1997). Survival in macrophage requires the bacteria to survive several environmental assaults, including oxidative stress, acid stress and cationic peptides (Foster & Spector 1995; Francis & Gallagher 1993; Groisman 1994; Para-Lopez et al. 1994). The virulent SL1344 strain of S. typhimurium was used because it was fully virulent and was therefore capable of expressing all of the factors required for macrophage survival (Hoiseth & Stocker 1981). The response of S. typhimurium to signals encountered in macrophage includes a role for the pleiotropic regulatory proteins PhoP/PhoQ. This twocomponent signal transduction system controls members of a large regulon of genes negatively or positively in response to intracellular signals (García Véscovi et al. 1996, 1997; Groisman 1994; Groisman et al. 1997; Gunn & Miller

plasmid	promoter insert	insert size (bp)		maximum reporter in vitro TCM	gene induction J774A.1
pGfp2			gfp lacZ		
pGyrB	gyrB	1600		$ imes 0.8 \pm 0.05$	× 5.1 ± 0.8
рТорА	topA	3300 -		$ imes$ 1.6 \pm 0.05	\times 1.3 ± 0.1
pPheST	pheST-ihfA	4700		$ imes$ 1.2 \pm 0.08	× 5.7 ± 1.0
pIhfA	ihfA	860		\times 1.1 ± 0.01	$ imes 6.7 \pm 0.8$
pIhfB	<i>ihfB</i>	4000		$ imes 0.6 \pm 0.04$	$ imes$ 2.8 \pm 0.3
pHns	hns	1600		$ imes$ 1.5 \pm 0.09	× 7.1 ± 1.0
pDps	dps	714		$ imes 0.6 \pm 0.05$	\times 3.0 ± 0.5
pSpv	spv	1574		$ imes 0.5 \pm 0.06$	× 5.1 ± 1.0

open boxes represent coding sequences of genes

) igure 1. A schematic representation of the different promoters assessed for macrophage gene expression in plasmid pGfp2 is nown (not to scale). Plasmid pGfp2 contains contiguous *lac*Z and *gfp* reporter genes downstream of a multiple cloning site Marshall *et al.* 2000). Promoter fragments were amplified with the primer combinations detailed in table 1 (sites engineered into rimers are underlined and named in parentheses). Following cleavage with restriction endonucleases the fragments were cloned nto similarly cleaved pGfp2 in a direction driving reporter gene expression. Open reading frames of genes assessed are indicated y open boxes. Maximal induction levels are shown as fold increase in LacZ expression, similar induction profiles were recovered then assaying for GFP intensity (data not shown). The LacZ levels were assessed using a chemiluminesence assay as previously escribed (Marshall *et al.* 2000). GFP intensity levels were determined using a FACScan (Becton Dickenson, Oxford, UK) with rgon lasers emitting at 488 nm and bacteria were detected by side scatter as previously described (Valdiva *et al.* 1996). The nean fold induction and standard deviations were calculated from a minimum of three independent experiments.

996; Soncini & Groisman 1996; Waldburger & Sauer 996). Under *in vitro* growth conditions, specific regulators f this type cooperate with the more global influences of NA topology to modulate the transcriptional profile of he cell (Dorman 1995). Therefore, we wished to ascertain `determinants of DNA topology formed part of the acterium's response to intracellular growth.

6. spv AND dps GENES

In addition to the genes involved in the regulation of NA topology, this study included two S. typhimurium romoters shown previously to be activated in J774A.1 ells. These were from the *spv* and the *dps* genes, each of which has been studied in detail previously in vitro and in \bigcup ivo (Marshall et al. 2000). The spv genes are located on a 0 kb virulence plasmid in the non-typhoid serovars of almonella (Guiney et al. 1995; Gulig 1996; Libby et al. 997). They are required for the establishment of a ystemic infection in the host and have been shown by gnature-tagged mutagenesis and in vivo expression techology to be required for full virulence and to be xpressed during infection (Heitoff et al. 1997; Hensel et al. 0995). The spv locus is composed of a regulatory gene, bvR, coding for a LysR-like transcription activator that ositively regulates both its own gene and the spvABCD peron of structural genes (Fang et al. 1991, 1992; Krause al. 1992; Guiney et al. 1995; Pullinger et al. 1989; heehan & Dorman 1998). Activation of spv transcription occurs during stationary phase *in vitro* and during *in vivo* growth (Chen *et al.* 1995; Kowartz *et al.* 1994). It requires the RpoS stress-response sigma factor, and is modulated by IHF, the leucine-responsive regulatory protein (Lrp), L-leucine, H-NS and cAMP-Crp (the latter probably acting indirectly through its effect on RpoS expression) (Marshall *et al.* 1999; O'Byrne & Dorman 1994*a*,*b*; Robbe-Saule *et al.* 1997). In addition, *spv* requires a negatively supercoiled DNA template for transcription *in vitro* (O'Byrne & Dorman 1994*b*; Marshall *et al.* 1999).

The dps gene is located on the bacterial chromosome, and like spv, it requires the RpoS sigma factor and IHF for full expression; unlike spv, dps is under the control of the OxyR regulator, a redox-sensitive DNA-binding protein (Altuvia *et al.* 1994). Dps protein is produced by starving bacteria, it co-crystallizes with the bacterial DNA and this is thought to protect the nucleic acid from damage while the bacteria are in stationary phase (Wolf *et al.* 1999). The dps promoter has a similar induction profile to that of spv when *S. typhimurium* is growing intracellularly and both promoters have been used successfully to express heterologous antigens in live attenuated vaccine strains (Marshall *et al.* 2000).

7. EXPERIMENTAL RESULTS AND DISCUSSION

Each promoter was cloned from the *S. typhimurium* genome by the polymerase chain reaction using standard methods, it was sequenced to ensure its structural integrity,

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able 1.	Primers	(5' - 3')	used for	the PCR	amplification	of p	olasmids
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GICAL	CES	lasmid	primer name	primer sequence	restriction enzyme recognition site ^a		
OLO		Dps	dps up	CGC <u>GGATCC</u> TATATATTCTTACCGG	(BamHI)		
	Š		dps start	CGC <u>GGATCC</u> AATCTCATATCCTCTTGATG	(BamHI)		
		TopA	topA cds	CGG <u>GGTACC</u> GGATCAATCCCCATACG	(KpnI)		
			topA up2	CGC <u>GGATCC</u> TTTGCCGGTATGTACGACGCC	(BamHI)		
		GyrB	gyrBup	ACAT <u>GCATGC</u> TGCTTTCACAACGAAGCC	(SphI)		
			gyrBstart	CGGGGATCCCTTGTCGAAGCGCGCTTTCTCG	(BamHI)		
Γ		Hns	hns up	CGC <u>GGATCC</u> ACTGTCTGAAGATGCCTTCG	(BamHI)		
			hns cds	CGC <u>GGATCC</u> CTTCAACGCTTTCCAGAGTAC	(BamHI)		
\succ	\mathbf{r}	PheST	ihfA cds	CGC <u>GGATCC</u> AACAGATATTCTGACATTTCAG C	(BamHI)		
\mathbf{O}	Η		ihfC	ACAT <u>GCATGC</u> AGGCCGTCAGATGATCATGG	(SphI)		
$\widetilde{\sim}$	Щ	IhfB	ihfB cds	CGC <u>GGATCC</u> GTCTTGGCGGGAATGTGC	(BamHI)		
			ihfB up	CGC <u>GGATCC</u> TGGCACGTACAACGACCACC	(BamHI)		
Ш	\cup	IhfA	ihfA2 val up	CGC <u>GGATCC</u> GTTTACGTTCCAGTTCAG GG	(BamHI)		
I	\bigcirc		ihfA2 val cds	CGC <u>GGATCC</u> AACAGATATTCTGACATTTCAG C	(BamHI)		
H	S	Spv	spv up	CGC <u>GGATCC</u> AACAGGTCAATTAAATCC	(BamHI)		
L		•	spv cds	CCC <u>GGATCC</u> CCTGAAAATAAACAGAATG AAATCC	(BamHI)		
PHILOSOPHICAL RANSACTIONS	0F	The location of the site in the primer sequence is shown by underlining. Ind then fused to promoterless copies of the gfp -M2 gene com Aequorea victoria (Cormack et al. 1996) and the $lacZ$ eporter gene from <i>E. coli</i> , carried in the low copy number lasmid pQF50 (Farinha & Kropinski 1990) derivative, e_{1} determined in the gene form <i>E. coli</i> , carried in the low copy number e_{2} to poisomers that had been more supercoiled in the bacteria migrated fastest in the gel (figure 3). For electro-					

and then fused to promoterless copies of the gfp-M2 gene com Aequorea victoria (Cormack et al. 1996) and the lacZeporter gene from *E. coli*, carried in the low copy number lasmid pQF50 (Farinha & Kropinski 1990) derivative, Gfp2 (figure 1). This plasmid offered two reporters, green uoresence encoded by *gfp* and β -galactosidase encoded by uZ. In the experiments described here, data obtained com monitoring lacZ expression are presented (figure 2), lthough all of the results were validated by gfp expression data not shown). S. typhimurium is naturally deficient in he lacZ gene so it was possible to monitor the levels of its β-galactosidase, from pGfp2 derivatives roduct, arboured in SL1344 strains while these were growing in itro or in vivo. B-galactosidase expression was measured sing the chemiluminescent substrate Galacton Star Clontech, Basingstoke, UK). The cloning strategy and he standard β -galactosidase assay employed have been escribed elsewhere (Marshall et al. 2000). Details of indiidual plasmid constructions are summarized in figure 1.

J774A.1 macrophage-like cells grown in tissue culture nedium (TCM) were infected with SL1344 bacteria at a atio of ten bacteria per macrophage. Following infection, reatment with gentamycin $(20 \,\mu g \,m l^{-1})$ was used to kill acteria that had not been engulfed by the J774A.1 cells. rocedures for the growth of J774A.1 and SL1344 cells nd the monitoring of in vivo gene expression have been Oescribed elsewhere (Marshall et al. 2000). The bacteria rere also inoculated into macrophage-free TCM to deternine background levels of gene expression.

Topoisomer analysis of reporter plasmid supercoiling as used to determine the effects of intracellular growth n DNA topology. The multicopy plasmid pUC18 was ntroduced into strain SL1344 and this strain was used to nfect J774A.1 macrophage-like cells. The pUC18 DNA as recovered from the infected cells and examined elecrophoretically for changes in supercoiling. Due to the w yield of plasmid DNA from the intracellular bacteria, he chloroquine-agarose gels were hybridized with a igoxygenin-labelled pUCl8 probe using the labelling nd detection procedures of Free & Dorman (1997). For

electrophoresis in one dimension, gels contained $2.5 \,\mu g \,m l^{-1}$ chloroquine; under these conditions those topoisomers that had been more supercoiled in the bacteria migrated fastest in the gel (figure 3). For electrophoresis in a second dimension, the topoisomers were passed through a gel containing $20 \,\mu g \,\mathrm{ml}^{-1}$ chloroquine. Here, topoisomers that were more relaxed in the first dimension formed an arc of positively supercoiled topoisomers migrating above the arc of negatively supercoiled topoisomers (Higgins et al. 1988; Wu et al. 1988) (figure 3).

Plasmid DNA isolated from S. typhimurium SL1344 cells that had been incubated in TCM alone showed a broad distribution of topoisomers similar to that seen in extracts from bacteria grown in standard laboratory media (Dorman et al. 1988; Higgins et al. 1988) and this distribution varied by just one or two topoisomers over time (figure 3b). In contrast, plasmid DNA removed from SL1344 that had infected J774A.1 macrophage-like cells showed a distribution that became progressively relaxed with time and included a novel fast-migrating species that increased in intensity with time spent in the macrophage (figure 3a). Electrophoresis with different concentrations of the chloroquine intercalator and twodimensional electrophoresis (figure 3c) showed that this species was composed of positively supercoiled DNA, i.e. plasmid DNA that had been highly relaxed in the bacteria.

The discovery that in vivo growth led to the production of a population of highly relaxed plasmids raised the possibility that bacterial DNA lost negative supercoils as a result of life within the macrophage. This finding is analogous to the effect of starvation on bacteria grown in vitro (Balke & Gralla 1987) and normally elicits a response at the level of gene transcription. In particular, genes required to compensate for the change in DNA topology might have been expected to have been induced, whereas genes whose products are unhelpful would not. We studied the response to intracellular growth of a panel of S. typhimurium promoters likely to be involved in adaptation to DNA relaxation.

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igure 2. Induction kinetics for *S. typhimurium* promoters studied *in vivo* following uptake by J774A.1 macrophages. Macrophage ifection and LacZ assays were performed as described previously (Marshall *et al.* 2000). β -galactosidse activity was expressed as ght units released per viable bacterium (light units per colony forming unit) following infection relative to the activity in the ioculum sample, and is the mean of at least three independent experiments for studies with J774A.1 cells. Squares, data points or intracellular bacteria; diamonds, data points for bacteria in TCM alone.

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igure 3. DNA supercoiling in intracellularly grown *S. typhimurium*. (*a*, *b*) One-dimensional agarose gel electrophoresis through % (w/v) agarose gel in 1 × tris borate (TBE) containing 2.5 mg chloroquine ml⁻¹. Topoisomer distribution in inoculum sample anes 1); after 40 min (lanes 2); after 80 min (lanes 3); after 120 min (lanes 4); after 180 min (lanes 5). (*a*) Topoisomer distribution of pUC18 following incubation of SL1344(pUC18) in TCM in 5% CO₂ at 37 °C. (*b*) Topoisomer distribution of plasmid UC18 before and following infection of macrophages with SL1344(pUC18). (*c*) Two-dimensional agarose gel electrophoresis rough 1% (w/v) agarose gel in 1 × TBE, containing 2.5 µg chloroquine ml⁻¹ in the first dimension and 20 µg chloroquine ml⁻¹ in the second dimension. Sample 1, pUC18 topoisomer distribution in *S. typhimurium* inoculum sample (corresponds to lane 1 f b); Sample 4, pUC18 topoisomer distribution of more negatively supercoiled topoisomers present in the inoculum but absent the intracellular bacteria is highlighted (X) in b and c. The novel fast migrating species unique to the intracellularly ecovered bacteria is arrowed in b and c.

DNA gyrase would be expected to play a prominent ole in the restoration of lost negative supercoiling. To do his it would require an upward shift in the [ATP]: ADP] ratio. Even in the absence of sufficient ATP (as in arving cells) to allow gyrase to function in negative upercoiling, the gyr gene promoters would be expected to e induced directly by DNA relaxation. In TCM alone, he gyrB promoter showed no induction during a 180 min cubation. In contrast, it was induced by approximately vefold when the bacteria were inside macrophage for the ame period (figure 1). This was in agreement with the nown behaviour of the equivalent promoter from *E. coli* then studied *in vitro* (Menzel & Gellert 1983, 1987*a*,*b*). In ontrast, the promoter for the *topA* gene (encoding the DNA relaxing enzyme topoisomerase I) showed no induction in TCM or in macrophage (figure 1). Since the *topA* promoter from *E. coli* is known to be inhibited by DNA relaxation, this finding was in keeping with the plasmid topoisomer data described above.

Other studies have suggested that the IHF may play a role in maintaining DNA topology in a form favourable to certain promoters under conditions where a general DNA relaxation takes place (Porter & Dorman 1997). In vitro, this relaxation occurs in stationary phase, and IHF levels have been shown to increase during this period of the growth curve, at least in *E. coli* (Ditto *et al.* 1994). An earlier study had shown that one of the genes coding for IHF, the *ihfA* gene (designated *mig-23* for 'macrophage-inducible

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ene 23'), was possibly induced when S. typhimurium grows tracellularly (Valdivia & Falkow 1987). In that study, the ecombinant plasmid tested contained just a segment of the *heT* gene where the *ihfA* promoter was located; it had not cluded the complete *pheST* operon and its promoter as rell as that of *ihfA*. We constructed a plasmid that included he complete *pheST-ihfA* operon and obtained an approxinately fivefold induction in the [774A.1 cells with no inducon in TCM alone (figure 1). To resolve the issue of the ossible contribution of the *pheST* promoter, a derivative lasmid was made in which the expression of the reporter enes was under the control of the *ihfA* promoter alone. \succ he data obtained with this plasmid showed that *ihfA* was - iduced approximately sixfold during intracellular growth \square ut not in TCM alone (figure 1). This shows that the *ihfA* romoter responds to growth in macrophage and this \mathcal{L} sponse is not modulated by a contribution from the *pheST* romoter. The promoter from the ihfB gene was also tested nd found to be inducible intracellularly but not in growth redium alone (figure 1). (The response of the ihfBromoter to intracellular growth had not been assessed reviously.) These data for *ihf* gene expression are in eeping with results obtained from *in vitro* studies showing hat these genes are induced when bacteria are starved or ndergo environmental stress (Aviv et al. 1994; Ditto et al. 994).

Successful establishment of an infection requires the acterium to multiply in the intracellular state (Finlay & alkow 1997). Previous work performed in vitro with E. coli as shown that expression of the nucleoid-associated rotein H-NS is coupled to DNA synthesis such that in apidly growing bacteria the hns gene is induced while in uiescent cells it is repressed (Free & Dorman 1995). The ns promoter was induced strongly in S. typhimurium cells rowing in macrophage and not induced in TCM alone figure 1). This suggests that while the bacteria may have een stressed when associated with the J774A.l cells, they vere still able to operate an efficient cell cycle, thus reating a demand for hns gene expression. A previous udy has indicated that intracellular S. typhimurium cells row rapidly and require protein synthesis for survival. s the bacteria adapt to the intracellular environment of he macrophage, they switch from this rapid growth node to a survival state that does not require protein ynthesis (Abshire & Neidhardt 1993). The kinetics of hns iduction revealed here, with expression declining 2 h ost-infection (figure 2), concurs with the findings of this arlier study.

As positive controls for the experiments described here, ne dps and spv promoters were also studied. Each had een shown previously to be induced during intracellular rowth (Marshall *et al.* 2000) and induction was also etected in this study (figures 1 and 2).

This study addresses a specific aspect of bacterial hysiology during infection, namely what are the conseuences of the infection process for the topology of acterial DNA and for the expression of genes coding for roteins that modulate DNA topology? It appears that ifection of the J774A.1 cell line causes bacterial DNA to ecome relaxed. As one might predict, the relaxationhibited promoter of the topA gene remains repressed nder these conditions. In contrast, the gyrB gene, coding or one of the subunits of DNA gyrase, is induced strongly. This is consistent with the production of extra copies of gyrase to restore supercoiling levels to values favourable to most DNA-based molecular transactions. The level of gyrB expression declines after 3 h, suggesting that by this stage post-infection supercoiling is restored at the gyrB promoter (figure 2).

The spv promoter used in this study illustrates some of the complexities arising from our data. It is supercoiling responsive and has been shown previously to be inhibited when DNA becomes relaxed (Marshall et al. 1999). However, in vitro data show that this inhibition requires the IHF protein; in the absence of this accessory factor, spv expression remains high even when DNA is relaxed (Marshall et al. 1999). This points to a subtle interplay between IHF and gyrase in the regulation of spv transcription. The *in vivo* data presented here show that spv is induced when the global level of DNA supercoiling is low. This may point to a shortage of IHF in vivo (consistent with the strong induction of the genes that code for this protein) or simply point to additional complexities inherent in the in vivo situation. For example, the leucineresponsive regulatory protein (Lrp) also regulates spv expression in vitro (Marshall et al. 1999) and this may have a positive modulatory influence when the bacterium is growing in association with macrophage. Alternatively, the negative influence of H-NS on spv expression described previously in vitro (O'Byrne & Dorman 1994b; Robbe-Saule et al. 1997) may not operate under the in vivo conditions studied here, and this is consistent with our data on *hns* gene induction in macrophage (figures 1 and 2) suggesting that not all H-NS-binding sites in the bacterial genome are occupied.

The data presented here allow a tentative model to be advanced to describe early events in intracellular adaptation from the perspective of DNA topology. Three lines of evidence suggest that DNA becomes relaxed early in the process. These are a direct observation of reporter plasmid topology by electrophoresis in the presence of a DNA intercalator, the positive response of the DNA relaxation-activated gyrB promoter, and the lack of a response from the DNA relaxation-inhibited topA promoter. Why does the DNA become relaxed? The most likely explanation is that the negative supercoiling activity of bacterial gyrase is inhibited as a result of interactions with the mammalian cells. This could arise from an unfavourable shift in the [ATP]:[ADP] ratio. As ATP levels fall and ADP levels rise, the negative supercoiling activity of gyrase is inhibited, although it can continue to contribute to ATP-independent relaxation of DNA (Drlica 1992). DNA topoisomerase I, and to some extent the other two topoisomerases in the cell, might also be expected to contribute to the general relaxation of genomic DNA. The induction of the genes encoding IHF may be regarded as a mechanism for dealing selectively with unfavourable alterations in DNA topology. Those promoters with appropriately located IHF-binding sites (such as spv) may be protected from inhibition as the DNA relaxes through the creation of a nucleoprotein complex that preserves promoter function. Other IHFdependent processes, such as DNA replication, transposition and site-specific recombination, may also continue to function. Other evidence suggesting that DNA replication continues under the conditions of our experiments comes

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om the observation of hns gene expression. A demand or hns transcription has been shown in vitro to be created y DNA synthesis (Free & Dorman 1995) and the in vivo ata presented here suggest that this process operates uring the macrophage infection. This is consistent with he data of Abshire & Neidhardt (1993) showing that igorous protein synthesis occurs for approximately 2 h blowing macrophage engulfment of S. typhimurium. The verall picture that emerges is one of stressed bacteria ith abnormally relaxed levels of DNA supercoiling that ontinue to grow and express genes that are critical for daptation to this environment. This situation is in > harked contrast to that described for invasion of epithelia - y S. typhimurium. Their increased negative supercoiling is 🗳 n essential prerequisite for expression of invasion funcons (Galán & Curtiss 1990). Thus, DNA topology Ontributes to different types of host-pathogen inter-Oction, and it does so in a dynamic way. Understanding • ow transitions between the different phases of the infecon affect DNA topology and gene expression will form he next phase of the work.

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