Identification of O-antigen polymerase transcription and translation start signals and visualization of the protein in Salmonella enterica serovar Typhimurium

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The wzyrflc gene, encoding the O-antigen polymerase, of Salmonella enterica serovar Typhimurium has been previously cloned and sequenced. In the present work, the wzy transcriptional startpoint was initially identified by primer extension. Next, wzy promoter strength in Escherichia coli K-12 was measured, and was found to be greater than that of the induced lac promoter. To define the Wzy translational startpoint, DNA including the wzy promoter and the putative first five residues of the Wzy protein was fused to the N-terminus of glutathione-S-transferase, and the fusion protein purified by affinity chromatography. N-terminal amino acid sequencing yielded the Wzy translational startpoint. Next, the Wzy protein was C-terminally tagged with the FLAG peptide, and immunoblotting of an S. typhimurium strain expressing a low-copy wzy-FLAG+ gene (five copies per cell) localized the intact Wzy protein in the cytoplasmic membrane of S. typhimurium cells. The Wzy protein was not well-expressed from a multi-copy wzy-FLAG+ plasmid in S. typhimurium, or in E. coli K-12.

Keywords: Wzy, Rfc, lipopolysaccharide, semi-rough

INTRODUCTION

The wzy gene (Reeves et al., 1996), which encodes the 47 472 Da Wzy protein, was previously cloned and sequenced, from Salmonella enterica serovar Typhimurium (henceforth termed S. typhimurium) (Collins & Hackett, 1991). The Wzy protein is the O-antigen polymerase. Thus, wzy mutants of S. typhimurium have a semi-rough phenotype, in which individual lipid A molecules carry at most a single O-antigen unit. If Wzy is synthesized, the typical O-antigen ladder of wild-type smooth S. typhimurium is seen (Collins & Hackett, 1991). The wzy gene of S. typhimurium maps in the 35·6 centisome region on the chromosome (Naide et al., 1965), at a site remote from the rfb gene cluster.

The wzy genes of several other enterobacteria have since been cloned and sequenced. Genes analysed in this manner have been identified as wzy genes either on the basis of homology with other wzy genes, or because the genes complement mutations that result in semi-rough phenotypes. The wzy genes of various S. enterica serovars have received most attention. It is likely that a close homologue of the S. typhimurium wzy gene is active in S. enterica strains of groups A and D1, as the S. typhimurium wzy gene detected homologues in strains of these groups when used as a probe in Southern blotting (Collins & Hackett, 1991), and the same O(1,2) linkage is present between O-antigen units of strains of groups B (typhimurium), A and D1 (Mäkelä, 1965; Nurminen et al., 1971). A group C2 strain of S. enterica also carries a gene with detectable homology to S. typhimurium wzy within a block of group C2-unique genes in the main rfb gene cluster (Brown et al., 1992). A different wzy gene (termed wzyD2H1) (Xiang et al., 1994), located in the main rfb gene cluster, has been identified in S. enterica strains of groups D2 and E1 [which have the same O(1,6) O-antigen linkage], partly on the basis of the fact that this is the only gene common to the rfb clusters of strains of these groups to which no rfb function has been assigned (Xiang et al., 1994). However, a wzy gene of S. enterica group D3 (wzyD3) which

Abbreviation: GST, glutathione-S-transferase.
also gives $\pi(1,6)$ linkage has no detectable sequence similarity to $\text{wzy}^{\text{DE}}$(Curd et al., 1998).

The $\text{wzy}$ genes of some smooth strains of *Escherichia coli* have been identified either by detectable homology to *S. typhimurium* $\text{wzy}$ (Yao & Valvano, 1994) or by the complementation of mutations in $\text{rfb}$ regions (Batchelor et al., 1991; Liu & Reeves, 1994; Lukomski et al., 1996). Complementation analysis has also indicated that the $\text{wzy}$ genes of *Shigella dysenteriae* (Klena & Schnaitman, 1993), *Shigella flexneri* (Morona et al., 1994), *Yersinia enterocolitica* (Zhang et al., 1996) and *Pseudomonas aeruginosa* (Coyne & Goldberg, 1995; De Kievit et al., 1995) are located in their respective $\text{rfb}$ gene clusters.

Putative $\text{Wzy}$ proteins share certain structural features. First, the putative proteins have multiple putative membrane-spanning domains (Collins & Hackett, 1991; De Kievit et al., 1995; Morona et al., 1994; Xiang et al., 1994; Zhang et al., 1996). The GC content of $\text{wzy}$ genes is generally lower than that of the bulk DNA of the bacterium in which the genes are located (Collins & Hackett, 1991; De Kievit et al., 1995). Analysis of codon usage in $\text{wzy}$ genes indicates that ‘rare’ codons (Grosjean & Fiers, 1982) are overrepresented, compared to codon usage by other genes sequenced from bacteria containing $\text{wzy}$ genes (Collins & Hackett, 1991; De Kievit et al., 1995; Morona et al., 1994).

Previously (Collins & Hackett, 1991), it was not possible to view labelled $\text{Wzy}$ protein in minicells of an *E. coli* K-12 strain containing a high-copy-number $\text{wzy}^+$ plasmid, nor was $\text{Wzy}$ detectable when the $\text{wzy}$ gene was under the control of the T7 promoter in a high copy-number plasmid, in an *E. coli* strain expressing the bacteriophage T7 RNA polymerase. Likewise, expression of the *S. flexneri* $\text{Wzy}$ protein could not be detected in *E. coli* K-12 under conditions where the products of adjacent cloned genes were visible after protein labelling (Morona et al., 1994). It has been suggested (Collins & Hackett, 1991) that the high usage of rare codons in the $\text{wzy}$ gene of *S. typhimurium* may greatly limit $\text{Wzy}$ production in *E. coli*, and the concentration of rare codons in the N-terminal region of the *S. flexneri rfc* gene has been proposed to exert an attenuating effect on $\text{Wzy}$ translation (Morona et al., 1994). Indeed, when 2–3 rare codons in the N-terminal region of the *S. flexneri wzy* gene were mutated to synonymous major codons, and the mutated gene then placed under the control of the T7 promoter and the Shine–Dalgarno sequence from gene 10 of phage T7, a low level of $\text{Wzy}$ was detected in whole *E. coli* K-12 cells following induction of the T7 polymerase (provided on a separate plasmid) (Daniels et al., 1998).

Unlike other $\text{wzy}$ genes, that of *S. typhimurium* is not located in the $\text{rfb}$ gene cluster, but at a distant site. The *S. flexneri wzy* gene does not have its own promoter, being part of the $\text{rfb}$ operon. Earlier, no attempt was made to determine (other than by sequence inspection) the transcriptional start point of the *S. typhimurium wzy* gene, and the possibility that $\text{Wzy}$ might be transcribed by readthrough from a plasmid promoter was not formally excluded. Also, because of the difficulties experienced in obtaining $\text{Wzy}$ expression, it was not possible to determine (by N-terminal analysis of purified protein) the translational start point, or to raise antibody against purified $\text{Wzy}$ for use in defining the subcellular localization of the protein. Pulse–chase experiments have shown that polymerization of $\text{O}$-specific lipopolysaccharide chains takes place at the periplasmic face of the inner membrane (McGrath & Osborn, 1991), and it is thus likely that $\text{Wzy}$ might be located in the cytoplasmic-membrane fraction. Also, Daniels et al. (1998) constructed (*S. flexneri*) $\text{Wzy}$–PhoA fusion proteins, and found that fusion proteins with up to 167 aa of $\text{Wzy}$ were located in the ‘membrane fraction’. The fusion points of expressive *lacZ* and *phoA* fusions with *S. flexneri wzy* further indicated that the $\text{Wzy}$ protein was located in the cytoplasmic membrane, with 12 transmembrane segments (Daniels et al., 1998). It remains true, however, that a full-length $\text{Wzy}$ protein has never been visualized, in either whole cells or the cytoplasmic-membrane fraction. The topics of *S. typhimurium wzy* transcriptional and $\text{Wzy}$ translational start signals, the subcellular localization of $\text{Wzy}$ and $\text{Wzy}$ quantification are addressed in this paper.

### METHODS

**Materials.** Enzymes active on DNA were obtained from either Gibco-BRL or Boehringer Mannheim and were used as directed by the suppliers. IPTG was purchased from Amersham Pharmacia. [$\gamma^{32}\text{P}]\text{ATP}$ ($\sim 1 \times 10^{14}$ Bq mmol$^{-1}$) and [$\gamma^{32}\text{P}]\text{dCTP}$ ($\sim 1 \times 10^{14}$ Bq mmol$^{-1}$) were from Amersham Pharmacia. Oligodeoxynucleotide primers were synthesized by Gibco-BRL. Rabbit anti-FLAG peptide antibodies were the product of Santa Cruz Biotechnology. The secondary antibody for ELISA [phosphatase-labelled goat anti-FLAG IgG (heavy and light chains)] was purchased from Pierce Laboratories. The secondary antibody for immuno-blotting (donkey anti-rabbit IgG, conjugated with horseradish peroxidase) was from Amersham Pharmacia. As an ELISA standard, bacterial alkaline phosphatase, C-terminally tagged with the FLAG peptide, was obtained from Eastman Kodak. ELISA plates were purchased from Corning.

**Strains and vectors.** *E. coli* K-12 DH5α (Hanahan, 1983) was the host for recombinant plasmids. Promoter strengths were assessed in the background of *E. coli* K-12 JM109 (Yanisch-Perron et al., 1985); *E. coli BL21(DE3) pLysS* (Phillips et al., 1984) was the host for expression of a fusion protein in which part of $\text{Wzy}$ was attached to glutathione-S-transferase (GST) (Phillips et al., 1984). *S. typhimurium* J357 (Cerin & Hackett, 1989) was used to modify plasmid DNA before transformation into other *S. typhimurium* strains. *S. typhimurium* C5 (wild-type) and *S. typhimurium* LV386 (wzy::IS10) have been described previously (Collins & Hackett, 1991). The low-copy plasmids pCL1920 (Lerner & Inouye, 1990), pQF50 (which contains a promoterless *lacZ* gene after a *HindIII* insertion site) (Farinha & Kropinski, 1990), pGEX-2T (Smith & Johnson, 1988) and pUC18 (Norlander et al., 1983), were used. Plasmid pADE206 (wzy$^+$) has been described previously (Collins & Hackett, 1991).

**Growth media.** Minimal M9 medium was prepared as described by Miller (1972) and supplemented prior to use with MgSO$_4$ (0.2 mg ml$^{-1}$), glucose (2 mg ml$^{-1}$) and thiamin-HCl.
251 GATAGCTCTGCTACTGGAGGCGCCGATGATAGGTAGTTAATGAGTGT
-35
301 TAAATGTCATTCAACGCTTGGTTGATACGAGAAGATGATGAGTAAT
-10 Transcription start
351 GATTATAGACTACAGTTACAAATATCTTGGCAGAAGATGTTTCCGTACC
401 ACACCTATTGGCCTAGTGTAATAATTTTTAATACACATTTTTTCTA
451 AAGGCTCTAATGTTAATTTTCACTACATCATGTCATTTAGCTTA........
SD M L I I S Y I A L C L
GTTACTCCCCCTATACTA........
G T S P I L
KpnI gst linker
1671 .....ATAAATAGTTAATACGCCGCAATGCGCGCTTTATTTGCTC
.... N K end
stem-loop
GATTATAAAGATGATGATGAT
D Y K D D D D Insertion to create FLAG tag

Wzy is located in the cytoplasmic membrane

(50 μg ml⁻¹). Luria–Bertani (LB) medium was prepared with Bacto tryptone (10 g l⁻¹), Bacto yeast extract (5 g l⁻¹) and NaCl (10 g l⁻¹). 2XYT medium has been described previously (Collins & Hackett, 1991). Liquid media were solidified, when required, by the addition of 1.5 % Oxoid bacteriological agar. Bacterial growth was at 37°C unless specified otherwise. Ampicillin was added to broth and solid medium at a final concentration of 50 μg ml⁻¹, Spectinomycin was used at 50 μg ml⁻¹ (E. coli) or 500 μg ml⁻¹ (S. typhimurium).

Cell fractionation. Cultures (40 ml) were grown in 2XYT for 3 h. Cells were harvested by centrifugation at 5000 g. The pellet, after washing in 30 mM Tris/HCl, pH 8.1, was resuspended in 0.8 ml 20 % (w/v) sucrose in 30 mM Tris/HCl, pH 8.1. Lysozyme (10 mg ml⁻¹ in 0.1 M EDTA) was added to reach a final concentration of 0.4 mg ml⁻¹. The cells were frozen in an ethanol/dry ice bath for 30 min and then thawed at room temperature. Twelve millilitres of 3 mM EDTA was added and cells were lysed by sonication (Branson sonicator) until the cell lysate became clear (usually 4 × 25 s bursts). Unlysed cells were removed by centrifugation at 5000 g. The lysate was separated into cytoplasmic and total membrane fractions by centrifugation at 200000 g for 1 h. The resulting pellet (total membrane) was dissolved in PBS (0.1 ml), followed by the addition of sodium lauryl sarcosine (Sarkosyl) in 7 mM EDTA, to 1 %. The mixture was incubated at 37°C for 30 min and then centrifuged at 200000 g for 1 h. The supernatant (Sarkosyl-soluble fraction) is the cytoplasmic-membrane fraction, while the pellet is the outer-membrane fraction. The pellet was washed with 10 mM Tris/HCl, pH 7.4 and recentrifuged at 200000 g for 2 h. The final pellet was then dissolved in PBS.

General analytical methods. DNA manipulations followed established methods (Sambrook et al., 1989). Development of Western blots used the Enhanced Chemiluminescence system (Amersham Pharmacia). SDS-PAGE was usually performed on 5 % (stacking gel)–12 % (separating gel) polyacrylamide, and proteins were stained with 0.25 % Coomassie brilliant blue G250. The gel loading buffer contained 2 % SDS and 100 mM DTT (Sambrook et al., 1989). Lipopolysaccharide-specific silver staining of SDS-PAGE gels was performed as described by Hitchcock & Brown (1983). The ELISA method has been published by Ausubel et al. (1993). The GST Gene Fusion System was purchased from Amersham Pharmacia. N-terminal protein sequencing was performed at the Bio-technology Laboratory/NAPS Unit, University of British Columbia, Canada.

Accessing the S. typhimurium wzy sequence. When the wzy sequence is mentioned below, the numbers shown are the nucleotide numbers from GenBank submission M60066, where nt 1 is the first nucleotide in the 1750 bp HindIII fragment carrying the wzy gene of S. typhimurium. Fig. 1 shows the wzy promoter region, the wzy translational start, the C-terminus, and the junctions of fusions of Wzy with GST and the FLAG peptide (see below).

Testing of promoter strength. To assay promoter strengths, PCR fragments, in which various promoters were contained on Spbl–HindIII fragments, were cloned into plasmid pQF50. To amplify the wzy promoter, the primers used were prwzyUP1 (5’-251-GATAGCGTCATGCTATCGGAGGCG-743-3’) and prwzyDOWN1 (5’-344-CATCACAAGCTGTGTTCCG-326-3’). The nucleotides in lower case are substitutions in the sequence (T to G, nt 258; T to A and T to G; nts 336 and 338) to create an Spbl site (prwzyUP1) or a HindIII site (prwzyDOWN1). To insert the lac promoter (with the catabolite-activator-protein binding site and lac operator) into pQF50, the primers used were prlacUP (5’-634-GCA-GCTGGCAcGACCGTTCCCG-812-3’) and plcDOWN (5’-463-TgGTCaACGGCTTTGTTGACG-483-3’), where the numbers refer to the sequence of pUC18 (GenBank accession no. L09136) (Norrander et al., 1983). The nucleotides in lower case are substitutions in the sequence (C to T, nt 258; T to A and T to G; nts 336 and 338) to create an Spbl site (prwzyUP1) or a HindIII site (prwzyDOWN1). To insert the lac promoter into pQF50, the primers used to amplify DNA from plasmid pGEX-2T were prlacUP (5’-65-GCTGTGCAcGTCATGCTATCGGAGGCG-326-3’) and plcDOWN (5’-263-GACATGAAAGCTGTGTTCCG-483-3’), where the numbers refer to the sequence of pUC18 (GenBank accession no. L09136) (Norrander et al., 1983). The nucleotides in lower case are substitutions in the sequence (C to T, nt 624 and A to C, nt 622; T to A, nt 469 and G to T, nt 474) in order to create an Spbl site (prlacUP) or a HindIII site (plcDOWN). To insert the tac promoter into pQF50, the primers used to amplify DNA from plasmid pGEX-2T were ptracUP (5’-65-GCTGTGCAcGTCATGCTATCGGAGGCG-863-3’) and ptracDOWN (5’-623-GACATGAAAGCTGTGTTCCG-243-3’) where the numbers refer to the GenBank sequence U13850 (Smith & Johnson, 1988). The bases in lower case are substitutions in the sequence (G to T, nt 74 and T to C, nt 76; TAC to GCT, nts 254–252) in order to create an Spbl site (ptracUP) or a HindIII site (ptracDOWN). Transformed
strains of *E. coli* K-12, JM109, carrying these plasmids, were grown in M9 minimal medium to ~ 5 x 10^7 cells ml^-1. Cells were pelleted, resuspended in Z-buffer (0.1-0.5 ml) (Miller, 1972) and lysed with 2 drops chloroform and 1 drop 0.1% SDS. The lysates were assayed for β-galactosidase activity as expressed in Miller units (Miller, 1972).

**Fusion of part of wzy with the gst gene.** To determine the translation startpoint of the Rfc protein, a portion of the *wzy* gene, including the *wzy* promoter and the DNA encoding the first 5 residues of the putative Wzy protein, was fused to the N terminus of the *gst* gene. First, an EcoRI–KpnI fragment, containing the required *wzy* DNA, was cloned into pUC18 and the resulting plasmid then received a KpnI–HindIII fragment containing the required *gst* DNA, to make plasmid pUST013. The primers used to amplify the *wzy* DNA were pr*ozyUP* (5'-39-TACCTTGAGATTCATGCTTG-57-3') and pr*ozyDOWN2 (5'-490-GTCATAAGGCTATGAAATTAT-376-3'). The nucleotides in lower case are substitutions in this sequence (T to G, nt 45; T to G, nt 484; C to T, nt 482 and AT to CC, nts 480–479) to create an EcoRI site (pr*ozyUP2) or a KpnI site (pr*ozyDOWN2). To amplify the *gst*-containing DNA from plasmid pGEX-2T, primers were used were prGSTUP (5'-249-ACAGTAGGATCCCATGAAATTAAATGAAATTATG-376-3') and prGSTDOWN (5'-1041-ACAGACAAAGCATGTCCCTCGGCTCCG-1018-3'), where the numbers refer to the GenBank sequence U13850 (Smith & Johnson, 1988). The bases in lower case are substitutions in this sequence (T–TC to GG–CT, nts 255–260 and G to T, nt 1030) in order to create a KpnI site (prGSTUP) or a HindIII site (prGSTDOWN). As a result of this construction, the first 5 putative residues of Wzy were fused, via 2 residues encoded by the *kpnI* site, to the second residue (after M) of GST (Fig. 1).

**Primer extension.** To determine the transcription startpoint of the *wzy* gene, the primers used in the primer extension experiment were pr*ozyDOWN3 (5'-483-GCAATGATGGAATTTAATAGCAGTCCAG-454-3'), pr*ozyDOWN4 (5'-374-TTCTGATCTGAGTGTGATCATGAAATTATGATC-344-3') and pr*ozyDOWN5 (5'-405-GTTGTTGCATCGGACATCGTCCCTCTG-376-3'). These were end-labelled with [32P]ATP using T4 polynucleotide kinase (Gibco BRL) and purified from a polyacrylamide gel. Total RNA was isolated, from strains grown in LB, using a Stratagene Micro RNA Isolation Kit. RNA (40 µg) was mixed with 0.8 pmol purified labelled primer, and the mixture precipitated with NaCl and ethanol (Sambrook et al., 1989). Reverse transcription was performed using SUPERSCRIPT RNaseH reverse transcriptase (Gibco BRL), and the annealing and subsequent extension reactions were performed as described in the SUPERSCRIPT RNaseH reverse transcriptase protocol, except that actinomycin D and RNasin ribonuclease inhibitor (Promega) were added to the reaction mixture at 50 µg ml^-1 and 2 U µl^-1, respectively. The reaction was stopped by heating at 70 °C for 15 min. RNase treatment, followed by ethanol precipitation, was performed as described by Sambrook et al. (1989). The samples were loaded next to a set of sequencing reactions, with pADE206 as template, which used the same primer. The sequencing reaction was performed using a T7 Sequencing Kit (Amersham Pharmacia).

**Insertion of the FLAG tag at the C-terminus of wzy.** The FLAG tag (DYKDDDDK) (Hopp et al., 1998) was created at the C-terminus of Wzy by insertion of a heterologous sequence between the penultimate and ultimate residue of Wzy (Fig. 1). PCR primers used were pr*ozyUP3 (5’-cattacaaagaatgtgatgatgatgatg-1679-AAATAAGTTATACCGGGCCGAATGCCCGCCTGGTTT-1709-3’) and pr*ozyDOWN6 (5’-ataccatcattgtaacct-1678-ATGTTCTTTAGTAAGACATCCTTATATGC-1648-3’). The heterologous sequence (gat tac aaa gat gat gat gat) in the primer pr*ozyUP3 encodes DYLKDDDDK, while the final K of the FLAG tag is provided by the ultimate codon (AA) of the native *wzy* gene. A sequence complementary to this segment is in the primer pr*ozyDOWN6. Two PCR reactions were performed, using pADE206 as template. The first reaction used Universal primer (5’-GTAAAACGAGCGACGGCTT-3’), from the Amersham Pharmacia T7 sequencing kit and pr*ozyDOWN6, and the other reaction used pr*ozyUP3 and a primer 5’-TCACAGGAAACAGCTTGACTG-3’), which is a modification of the Reverse sequencing primer (Amersham Pharmacia). The two PCR products were mixed and annealed (at the complementing sequences in the primers pr*ozyUP3 and pr*ozyDOWN6). The annealed product was used as a template for a final PCR reaction using the Universal and modified Reverse primers. The final product was digested with HindIII and cloned into pUC18 to give pUST011, or into pCL1920 to give pUST012.

**RESULTS**

**The transcriptional startpoint of the wzy gene**

Cellular RNA from *S. typhimurium* LV386 (*wzy*) and the complemented multi-copy *wzy* strain *S. typhimurium* LV386/pADE206, was subjected to primer extension from pr*ozyDOWN3 to localize the *wzy* transcriptional startpoint (Fig. 2). The *wzy* transcriptional startpoint was shown to be A313 (Figs 1, 2).

![Fig. 2. Identification of the wzy transcription startpoint by primer extension, using the oligonucleotide pr*ozyDOWN3. RNA (40 µg) from S. typhimurium LV386 (wzy) (lane 1), and the complemented wzy strain S. typhimurium LV386/pADE206 (lane 2) was annealed with the wzy-complementary primer pr*ozyDOWN3, and reverse transcription followed (see Methods). A sequencing reaction, using the same primer, was also performed. The letters on the right show DNA sequence, and the bold A at position 313 shows the wzy transcriptional startpoint.](image)
Initial attempts to purify the Wzy protein, from \textit{E. coli} K-12 or \textit{S. typhimurium} cells with the \textit{wzy} gene in low- or multi-copy plasmids, were unsuccessful, possibly due to the fact that the protein is membrane associated (see below). A different approach to determination of the N-terminal amino acid sequence of Wzy was therefore taken. Routinely, all or part of foreign proteins may be fused to the C-terminus of GST without affecting the ability of GST to bind glutathione, so that the fusion proteins may be purified by affinity chromatography. It was thought that it might be possible to fuse a small portion of a foreign protein to the N terminus of GST, also without affecting the glutathione-binding capacity of GST. Accordingly, the entire \textit{wzy} promoter and a sequence containing the first 5 putative residues of Wzy was fused to the second residue (thus, the starting M was omitted) of GST (see Methods) (Fig. 1), and a fusion protein sought by affinity chromatography (Fig. 3). This approach was successful. From an 800 ml culture of \textit{E. coli} BL21(\textit{DE3}) pLysS/pUST013, a total of 3.6 mg Wzy-GST fusion protein was obtained. A portion of this protein was subjected to N-terminal amino acid sequencing and the first 10 residues were LIISGTSPIL (Fig. 1), as previously suggested (Collins & Hackett, 1991). The sequence AAAGG430–454 (Fig. 1) may, as indicated previously (Collins & Hackett, 1991), serve as part of a ribosome-binding site upstream of the \textit{wzy} start codon.

**Visualization of the Wzy protein in cells of \textit{S. typhimurium}**

Initially, it was planned to purify the Wzy protein so that anti-Wzy antibody might be raised. Fusion proteins with all or part of Wzy linked to the C-terminus of GST were uniformly unstable, and were obtained in only small and variable yields. Accordingly, the FLAG tag was inserted at the C-terminus of Wzy (see Methods) (Fig. 1). This insertion did not obviously affect Wzy function (Fig. 4), as the typical O-antigen ladder was obtained when the \textit{wzy} strain, \textit{S. typhimurium} LV386, was complemented with the \textit{wzy–FLAG} gene, resident in either multi- or low-copy plasmids.

Next, FLAG-tagged Wzy protein, expressed in the \textit{S. typhimurium} LV386-based strains containing either the multi- or low-copy \textit{wzy–FLAG} plasmids, was sought by

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**Fig. 3.** Purification of the Wzy–GST fusion protein. The \textit{wzy} promoter and the putative first 5 residues of Wzy were fused to the second residue of GST. The fusion protein was purified by affinity chromatography using agarose-bound glutathione and subsequent glutathione elution (Smith & Johnson, 1988). Lane 1, purified Wzy–GST (50 µg); lane 2, molecular-mass markers (sizes, in kDa, are shown on the right).

**Fig. 4.** Lipopolysaccharides of \textit{S. typhimurium} strains on an SDS-PAGE gel, with lipopolysaccharide-specific silver staining. Each lane received material from 10^7 cells grown in 2XYT. Lane 1, \textit{S. typhimurium} C5 (wild-type); lane 2, \textit{S. typhimurium} LV386 (\textit{wzy}); lane 3, \textit{S. typhimurium} LV386/pUST010 (low-copy \textit{wzy}); the \textit{wzy}’ HindIII fragment was inserted into pC1920; lane 4, \textit{S. typhimurium} LV386/pUST011 (multi-copy \textit{wzy–FLAG}); lane 5, \textit{S. typhimurium} LV386/pUST012 (low-copy \textit{wzy–FLAG}).
immunoblotting (Fig. 5). No genuine FLAG signal was obtained on immunoblots of whole cells of either *S. typhimurium* C5 or *S. typhimurium* LV386 (neither strain expresses a FLAG peptide), although a protein cross-reactive with the anti-FLAG antibodies was present in all strains examined (and is shown, below, to be located in the cytoplasm of *S. typhimurium* LV386/pUST012). Surprisingly, a genuine FLAG signal was also absent upon immunoblots of whole cells of *S. typhimurium* LV386/pUST011 (which carries multi-copy wzy–FLAG), even though such cells clearly expressed active Wzy protein (Fig. 4). The low level of Wzy expression (below the sensitivity level of the immunoblot) in this strain will be discussed later. When whole cells of *S. typhimurium* LV386/pUST012 (which has low-copy wzy–FLAG) were examined, however, FLAG-tagged Wzy protein was visible (Fig. 5c, track 4), and cell fractionation localized the protein to the cytoplasmic membrane (Fig. 5d, track 3).

In an effort to quantify the Wzy protein in *S. typhimurium* LV386/pUST012, whole cells of this strain, and of control *S. typhimurium* LV386, were disrupted by boiling in gel-loading buffer, and then used to coat wells of a 96-well tray. An ELISA assay, using a commercial pure FLAG-tagged protein (similarly boiled) as standard, followed (Fig. 6). The Wzy protein level in the (low-copy wzy) strain of *S. typhimurium* LV386/pUST012 was ~25 ng per 10^7 cells (after correction for cross-reaction between the anti-FLAG antibodies and a cytoplasmic protein present in both control and test strains). This calculation assumes that the efficiency of binding to the plastic tray of the FLAG-tagged Wzy protein from solubilized whole cells, was the same as that of the pure FLAG-tagged protein standard, and that the FLAG tag does not affect (inhibit?) the natural degradation rate of the unmodified Wzy protein. If these two conditions are met, it may be calculated that when the wzy gene is present at 5 copies

![Fig. 5](image_url)

**Fig. 5.** Detection of the Wzy protein by immunoblotting. For panels (a) and (c) (10^8 cells per track), whole cells, grown in LB, were solubilized and analysed by SDS-PAGE. Panel (a) was developed with Coomassie blue and panel (c) by immunoblotting, using antibodies against the FLAG tag (1:500 dilution; 0.2 µg protein ml⁻¹) as the primary antibody, and the secondary antibody at a 1:1000 dilution. Lanes 1, *S. typhimurium* C5 (wzy⁺); lanes 2, *S. typhimurium* LV386 (wzy); lanes 3, *S. typhimurium* LV386/pUST011 (multi-copy wzy–FLAG); lanes 4, *S. typhimurium* LV386/pUST012 (low-copy wzy–FLAG). For panels (b) and (d), whole cells of *S. typhimurium* LV386/pUST012 (low-copy wzy–FLAG), grown in LB broth, were fractionated (material from 10^7 cells per track) before SDS-PAGE. Lanes 1, cytoplasmic fraction; lanes 2, whole membrane; lanes 3, cytoplasmic membrane; lanes 4, outer membrane. Panel (b) was developed with Coomassie blue, and panel (d) by immunoblotting, using antibodies against the FLAG tag as the primary antibody. The positions of molecular mass markers, with sizes, are shown on the left. The small arrow (left) marks the position of a cytoplasmic protein, present in all strains, which cross-reacts with the anti-FLAG antibodies. The large arrow (right) shows the FLAG-tagged Wzy protein in whole cells, whole membrane, and the cytoplasmic membrane of *S. typhimurium* LV386/pUST012.
per cell, each cell contains \( \sim 30000 \) copies of the Wzy protein, which would thus comprise \( \sim 1\% \) of total cellular protein (Neidhardt & Umbarger, 1996). As the Wzy protein is a cytoplasmic-membrane protein, it is relevant to mention that this would equate to \( \sim 1 \) molecule of Wzy protein for every 700 molecules of membrane phospholipid (Neidhardt & Umbarger, 1996). If it is assumed that Wzy production from a single wzy gene copy would be one-fifth of the amount seen from the low copy-number wzy plasmid, then Wzy would be present, in wild-type \( S. typhimurium \), at \( \sim 6000 \) copies per cell, comprising \( \sim 0.3\% \) of total protein: one Wzy molecule for each 3500 molecules of phospholipid.

**DISCUSSION**

**Subcellular localization of Wzy**

Pulse–chase experiments have previously shown that polymerization of O-specific lipopolysaccharide chains takes place at the periplasmic face of the inner membrane (McGrath & Osborn, 1991), and it was thus always likely that Wzy might be located in the cytoplasmic-membrane fraction. Daniels et al. (1998) located a (\( S. flexneri \)) Wzy–PhoA fusion with 167 aa of Wzy in a ‘membrane fraction’. It was not possible to view fusions with longer Wzy segments, and it was suggested that this was due to the increasing number of rare codons in the longer wzy segments of such fusions, with consequent poor fusion-protein expression. Also, the locations of expressive lacZ and phoA fusions in the wzy gene were indicative of a cytoplasmic-membrane location for Wzy, with 12 transmembrane segments (Daniels et al., 1998).

The results in this paper are in full agreement with these data. In this work, the full-length \( S. typhimurium \) Wzy protein is shown to be located, by immunoblotting of an attached FLAG tag, in the cytoplasmic membrane.

**Transcription of wzy**

When an extract of (wild-type) \( S. typhimurium \) C5, which contains a single copy of the wzy gene, was used for primer extension, no labelled product was detectable on a gel such as that shown in Fig. 2 (data not shown). The sensitivity of this method thus requires that the wzy gene be present in a multi-copy state. Transcription of the wzy gene from a single startpoint in the multi-copy wzy plasmid, pADE206, was noted in \( S. typhimurium \) (Fig. 2), and this transcript was clearly translated into functional Wzy protein (Collins & Hackett, 1991). It is reasonable to assume that the transcription startpoint of wzy in pADE206 is the same as that used in wild-type
S. typhimurium. It was further shown that the \textit{wzy} gene has its own strong promoter.

**Level of translation of Wzy**

Previous attempts (Collins & Hackett, 1991) to detect Wzy translation used systems (maxicells, minicells, T7-polymerase-mediated transcription from a T7 promoter) requiring that Wzy be translated in polymerase-mediated transcription from a T7 promoter) requiring that Wzy be translated in E. coli K-12; as suggested (Collins & Hackett, 1991), it seemed possible that such expression might be suboptimal, as the Wzy protein has no obvious function in \textit{E. coli} K-12. This suggestion may have merit. Thus, when pUST012 (which allows good levels of Wzy protein expression in a \textit{S. typhimurium} background; Fig. 6) was transformed into various \textit{E. coli} K-12 strains and Wzy protein sought, after SDS-PAGE of lysed whole cells, by immunoblotting with anti-FLAG antibodies, no Wzy protein was detected (data not shown). The absence of obvious Wzy in \textit{E. coli} K-12 is not due to poor expression of the \textit{wzy} promoter in \textit{E. coli} K-12. It may be that the lack of function for Wzy in \textit{E. coli} K-12 causes the protein to be targeted by the protein-degradation systems of the cell, resulting in a steady-state level of Wzy which is very low or zero.

It was also suggested previously (Collins & Hackett, 1991) that the presence of many attenuating codons in \textit{wzy} might cause Wzy to be translated at a low level. With the multi-copy \textit{wzy} plasmid, pUST011, in \textit{S. typhimurium}, no Wzy protein was seen, and it is possible that the simultaneous initiation of many Wzy protein molecules, with consequent high demand for attenuating tRNA species, results in completion of only a few Wzy protein molecules, so that Wzy protein levels are below the detection level of the immunoblot (Fig. 5) but still adequate to complement the chromosomal \textit{wzy} mutation (Fig. 4). Daniels et al. (1998) placed the \textit{S. flexneri} \textit{wzy} gene downstream of the T7 promoter and the Shine–Dalgarno sequence from gene \textit{10} of phage T7, induced the T7 polymerase, and found that expression of labelled Wzy increased (from undetectable to weak) when 2–3 rare codons in the N-terminal region of \textit{wzy} were mutated to major synonyms. This is the most direct evidence that translation attenuation by rare codons is a controlling factor in Wzy production.

If only a few Wzy molecules are being synthesized at any time (from the low-copy \textit{wzy} plasmid pUST012), the protein completion rate may be much higher than seen when \textit{wzy} is present in a high copy-number plasmid (Fig. 5). This greater Wzy protein completion rate, coupled with possible good stability of the completed protein (embedded, and active, in the cytoplasmic membrane), appears to result in a steady-state level of Wzy equivalent to \( \approx 0.3\% \) of bacterial protein/\textit{wzy} copy, in \textit{S. typhimurium}.

The main conclusions of this study are that the \textit{wzy} transcriptional startpoint has been identified at residue A313 and confirms that the \textit{S. typhimurium} \textit{wzy} gene has its own promoter, with its \(-10\) and \(-35\) sequences, as predicted previously (Collins & Hackett, 1991). The \textit{wzy} promoter strength was assessed and lies between that of the \textit{lac} and \textit{tac} promoters, N-terminal amino acid sequencing of a Wzy–GST fusion protein confirmed the translation start point which agreed with sequence prediction. Using FLAG as an antibody-reactive tag, a Wzy–FLAG protein was visualized by SDS-PAGE and was localized to the inner membrane. The level of Wzy in wild-type \textit{S. typhimurium} was estimated to be \( \approx 6000 \) copies/cell by ELISA analysis.

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**REFERENCES**


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