Localization of *phoE*, the Structural Gene for Outer Membrane Protein e in *Escherichia coli* K-12

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To localize the structural gene for outer membrane protein e, mutants resistant to the protein e-specific phage TC45 were isolated and characterized. Three classes of TC45-resistant strains were found: (i) mutants in *phoB*, a regulatory gene for protein e, (ii) mutants with an altered lipopolysaccharide, and (iii) mutants unaltered in the regulation of the *pho* regulon and producing an apparently normal lipopolysaccharide. Mutations in the latter class of mutants are probably altered in the structural gene for protein e and are cotransducible with *proA,B* at min 6 on the chromosomal map. As class (iii) includes mutants with an electrophoretically altered protein e, in both an *nmpA* and an *nmpB* background, we conclude that the structural gene for protein e, designated as *phoE*, is localized at min 6 on the chromosomal map, the gene order being *phoE proA argF*.

Several proteins of the outer membrane of Escherichia coli K-12 are involved in the formation of aqueous pores through which small hydrophilic molecules can pass this membrane (2, 15, 17, 25, 26). Two of them, the products of the genes ompC and ompF, are synthesized constitutively in wild-type strains of E. coli K-12. Another outer membrane protein, protein e (26) (also called Ic [8] or E [5]), was detected in pseudorevertants of mutants lacking both the OmpC and the OmpF protein (26). This protein also has porin properties (14, 21, 26). In wildtype cells of E. coli K-12, the synthesis of protein e can be induced by phosphate limitation (18). Mutations leading to constitutive synthesis of protein e have been localized in either of two loci, nmpA and nmpB at min 83 (6, 21) and min 9 (10, 21), respectively. Recently, we showed that the synthesis of protein e is controlled by the same regulation mechanism as that which controls the synthesis of alkaline phosphatase and that nmpA is identical to one of the genes, phoS, phoT, or pst, which are involved in the uptake of inorganic phosphate, whereas the nmpB gene is identical to the phoR gene, which is a regulatory gene of the pho regulon (23).

Recently, Pugsley et al. (20) reported that probably both the *nmpA* and *nmpB* loci contain structural genes for protein e. Starting with *nmpA* and *nmpB* mutant strains, they isolated TC45-resistant mutants which produce an electrophoretically altered protein e, and these secondary mutations were reported to map in or close to *nmpA* and *nmpB*, respectively. These

data were interpreted as strong evidence for the presence of two structural genes at min 83 and 9, respectively (20). When this information was published, we were analyzing mutants resistant to the protein e-specific phage TC45 to localize the structural gene for this protein. Our results, which are described in this paper, indicate that there is only one structural gene for protein e and that this gene is located at min 6.

MATERIALS AND METHODS

Strains, phages, and growth conditions. All bacterial strains are derivatives of *E. coli* K-12. Their sources and relevant characteristics are listed in Table 1. Cells were grown overnight in yeast broth (13) under vigorous aeration, usually at 37°C. A growth temperature of 30°C was used to check for the presence of protein e in cell envelopes, because protein a, which is another outer membrane protein with the same electrophoretic mobility as protein e in the gel system used (14), is hardly produced at the latter growth temperature (13, 16).

Laboratory stocks of coliphages T3, T4, T7, P1, C21, and TC45 were used.

Genetic techniques. P1 transduction (28) and conjugation (7) were carried out as described previously. Sensitivity to bacteriophages was determined by cross-streaking.

Assays for alkaline phosphatase. Assays for alkaline phosphatase were performed as described previously (23).

Isolation and characterization of cell fractions. After ultrasonic disintegration of the cells, cell envelopes were isolated by differential centrifugation (12). For the isolation of periplasmic proteins, a slightly modified procedure (23) of the EDTA-lysozyme

TABLE 1. Characteristics of bacterial strains^a

Strain	Characteristics	Source ^b or reference
PC0479	F ⁻ thr leu thi pyrF thy ilvA his lacY argG tonA rpsL cod dra vtr glpR, protein e ⁻	PC
CE1108	ompB471 phoS200, protein e ⁺ derivative of PC0479	14, 23
CE1174	phoB201, protein e derivative of CE1108	23
CE1182	TC45-resistant protein e^- derivative of CE1108, resistant to T3, T7, and P1	This study
CE1183	TC45-resistant protein e ⁻ derivative of CE1108, sensitive to C21	This study
CE1184	TC45-resistant protein e ⁺ derivative of CE1108, sensitive to C21	This study
CE1185	TC45-resistant protein e ⁻ phoE203 pro derivative of CE1108	This study
CE1186	TC45-resistant protein e^- phoE204 pro derivative of CE1108	This study
CE1187	TC45-resistant protein e ⁻ phoE205 pro derivative of CE1108	This study
CE1202	TC45-resistant protein e* phoE202, derivative of CE1108	N. Overbeeke
CS809	F ⁻ metB65 his-53 proC24 cyc-1 xyl-14 lacY29 rpsL77 tsx-63 ompF ompC171 nmpA104 nmpA114, protein e*	20
CS812*°	F ⁻ metB65 his-53 proC ilvY227 cyc-1 xyl-14 lacY29 rpsL77 tsx-63 ompF ompC171 nmpB105 nmpB116, protein e*	20
JF694	ilv his purE proC metB rpsL cyc xyl lacY tsx ompF ompC nmpA, protein e^+	5
CE1188	proA, protein e ⁺ derivative of CE1108	This study
PC0031	Hfr R4 argF gal tonA phx rel	PC
C9 ^d	Hfr Cav relA1 tonA22 pit-10 spoT1 phoR18 T2	A. Garen via CGSC
CE1196	Hfr R4 thr leu proA argF argI phx rel bgl phoS21	This study

^a Genotype descriptions follow the recommendations of Bachmann and Low (1).

method of Willsky and Malamy (29) was used. The protein patterns of the cell fractions were analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis as described previously (12). For the isolation of ³²P-labeled lipopolysaccharide (LPS) and its characterization by means of paper chromatography, we used the method described by Boman and Monner (3). Isobutyric acid-1 N NH₄OH (5:3, by volume) was used as the solvent in paper chromatography.

RESULTS

Isolation and characterization of TC45-

resistant mutants. Starting with strain CE1108, a phoS strain which produces protein e constitutively, spontaneous mutants resistant to the protein e-specific phage TC45 were isolated. Three classes of TC45-resistant mutants were found. The characteristics of these mutants are summarized in Table 2. Class 1 mutants lack, in addition to protein e, alkaline phosphatase (Table 2) and the other periplasmic proteins of the pho regulon (data not shown). The mutation in one of these mutants, strain CE1174, has previ-

^b PC, Phabagen Collection, Department of Molecular Cell Biology, Section Microbiology, State University of Utrecht, Utrecht, The Netherlands; CGSC, E. coli Genetic Stock Center, Department of Human Genetics, Yale University School of Medicine, New Haven, Conn. (B. J. Bachmann, curator).

^c This strain is a derivative of CS812, obtained from A. Pugsley. In contrast to strain CS812, this strain carries an additional *proC* mutation.

^d As the mutation which causes resistance to phage T2 and disappearance of the *ompF* protein is 40% cotransducible with *pyrD*, it probably is localized in the *ompF* gene (R. van Boxtel, J. Tommassen, and B. Lugtenberg, unpublished data).

ously been located in the phoB gene, which codes for a positive regulatory protein of the pho regulon (23). Class 2 mutants have an altered LPS, as indicated by their changed sensitivity towards the LPS-specific phages T3, T7, and P1 or C21 (11). One subclass of these mutants, of which strain CE1182 is an example, has largely decreased quantities of protein e (<10%) and has become simultaneously resistant to the phages T3, T7, and P1. Like the parental strain, it is still resistant to phage C21. This phage pattern indicates that these strains have a heptose-deficient LPS (7, 24). The isolation of such mutants could be expected since it has been reported that mutants with a heptose-deficient LPS contain strongly decreased amounts of protein e (14). A second subclass of TC45-resistant LPS mutants has become sensitive to phage C21 and is still sensitive to phages T3, T7, and P1, suggesting that their LPS is altered such that it is galactose deficient but contains heptose (27). This subclass is heterogeneous with respect to the residual amounts of protein e, as some strains, e.g., strain CE1183, have very low levels of this protein (<10%), whereas in others, e.g., strain CE1184, the amount of protein e is slightly or not at all affected (>80%). It should be noted that also in the latter case, phage TC45 resistance is due to a decreased rate of phage adsorption.

All 15 isolated class 3 mutants, e.g. CE1185, CE1186, and CE1187, lack protein e. They have an apparently wild-type LPS, as judged by their sensitivity pattern towards the described phages and by the chromatographic mobilities of the isolated ³²P-labeled LPS. Since all mutants still produce alkaline phosphatase constitutively, the

lack of protein e cannot be due to reversion to $phoS^+$ or to mutation of the phoB gene. Thus, the most likely explanation is that these strains have an altered structural gene for protein e.

Genetic localization of the structural gene for protein e. In contrast to their parental strain CE1108, 14 out of 15 class 3 TC45-resistant mutants appeared to be dependent on exogenous proline for growth. In transduction experiments, this proline auxotrophy was always strongly coupled with TC45 resistance, indicating that the mutations are caused by deletions of part of the genome including the structural gene for protein e and at least one of the pro genes. On the E. coli K-12 chromosome, three genes are known to be involved in proline synthesis, namely, proA and proB at min 6 and proC at min 8.8 (1). proA and proB are proximal markers in crosses with Hfr strain R4, whereas proC is a late marker (Fig. 1). To localize the proline auxotrophy and TC45 resistance markers in class 3 mutants, we crossed three mutant strains, namely, CE1185, CE1186 and CE1187, with Hfr R4. Proline prototrophy appeared to be an early marker in these conjugation experiments, and for each cross, all 50 tested pro+ transconjugants were TC45 sensitive. So we conclude that the TC45 resistance marker in the class 3 mutants is localized close to proA and

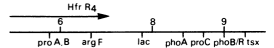


Fig. 1. Marker positions in the region of min 6 to 9 of the E. coli chromosome (1). The origin and direction of transfer of Hfr strain R4 are indicated.

Table 2. Characteristics of three classes of TC45-resistant mutants of strain CE1108^a

	Strain		Sensitivity to:				1100000
Class		TC45	T3, T7, P1	C21	AP	Protein e	Defective in:
Parent	CE1108	S	S	R	c	+	phoS
1	CE1174	R	s	R	unind	-	phoB
2	CE1182	R	R	R	c	_	LPS
	CE1183	\mathbf{R}	S	\mathbf{s}	c	_	LPS
	CE1184	R	S	s	c	+	LPS
3	CE1185	R	s	R	c	_	$phoE^b$
	CE1186	\mathbf{R}	S	\mathbf{R}	c	_	$phoE^b$
	CE1187	R	s	\mathbf{R}	c	_	$phoE^b$
	CE1202	R	s	\mathbf{R}	c	*	$phoE^b$

^a R and S, Resistant and sensitive, respectively; AP, synthesis of alkaline phosphatase; c and unind., constitutive and uninducible, respectively; + and -, presence or absence, respectively, of protein e in cell envelope preparations; *, a protein e with altered mobility in sodium dodecyl sulfate-polyacrylamide gels.

^b phoE is the proposed abbreviation for the structural gene for protein e.

proB at min 6.

Recently, Pugsley et al. (20) isolated TC45resistant mutants from nmpA and nmpB strains, and they localized these mutations in or close to nmpA at min 83 and nmpB at min 9, respectively. Some of their mutants produced an electrophoretically altered protein e. Since mutations that cause an altered protein are very likely to be in the structural gene for this protein, they suggested that there are two structural genes for protein e, one at min 83 and one at min 9. Assuming that the existence of three structural genes for protein e is unlikely, we therefore reinvestigated the localization of the two mutations which cause an electrophoretically altered protein e (designated as protein e* in this paper) in two mutant strains obtained from Pugsley. A third strain with an e* protein, strain CE1202, which was isolated in our lab, was also included in these experiments. The major outer membrane proteins of the e⁺ strain CE1108 (Fig. 2b, d, and h) and of three TC45-resistant mutant strains which produce an altered protein e (Fig. 2a, c, and e) are shown in Fig. 2. Strain CS809 (Fig. 2a) is a TC45-resistant mutant isolated from an nmpA strain. This strain was isolated by Pugsley et al., and the TC45 resistance mutation $(nmpA^*)$ was mapped at min 83 in or close to nmpA (20). Strain CS812* (Fig. 2c) is a TC45-resistant mutant isolated from an nmpB (phoR) strain. The TC45 resistance mutation (nmpB*) in this strain was localized at min 9 in or close to the nmpB gene (20). The third protein e* strain, strain CE1202 (Fig. 2e), was isolated in our laboratory as a TC45-resistant strain.

To determine whether the $nmpA^*$ mutation in strain CS809 is indeed localized at min 83 in or close to the nmpA gene, we transduced strain PC0479 to ilv^+ , using P1 grown on CS809. A total of 30 ilv^+ transductants were tested, and 16 of them produced alkaline phosphatase constitutively, indicating that they received the nmpA (pho) gene from donor strain CS809.

These 16 transductants were all TC45 sensitive. The protein pattern of the cell envelopes of these transductants was analyzed on gels, and, as is shown for one of the transductants in Fig. 2 (slot f), they appeared to have a normal (wild-type) protein e. In addition, by selecting for ilv^+ transductants, it was not possible to transduce the e⁺ strains CE1108 and JF694 to TC45 resistance with P1 grown on $nmpA^*$ strain CS809. For these reasons, we conclude that the $nmpA^*$ mutation in strain CS809 is not localized in or close to the nmpA gene at min 83.

To determine whether the $nmpB^*$ mutation in strain CS812* is indeed localized at min 9 in or close to the nmpB (phoR) gene, as reported in reference 20, we transduced strain CS812* to $proC^*$ with P1 grown on strain C9. Two hundred pro^* transductants were tested. No contransduction of TC45 sensitivity with $proC^*$ was found, whereas normal cotransduction percentages for $lacY^*$ (14%) and tsx^* (43%) were obtained. Therefore, we conclude that the $nmpB^*$ mutation of strain CS812* is not localized between lacY and tsx in or close to nmpB (phoR) at min 9.

To determine whether the mutations which cause an e* protein are localized in the same region of the chromosome where the TC45-resistant pro mutations are mapped (class 3 mutants), we transduced e+ strain CE1188 to proA+ with P1 grown on strain CE1202. A total of 200 pro⁺ transductants were tested, and 194 (97%) appeared to be TC45 resistant. Also, the TC45 resistance markers in the strains CS809 and CS812* were cotransducible with proA (97%). The cell envelope protein patterns of two $proA^+$ TC45-resistant transductants of strain CE1188 with P1 grown on strains CS809 and CS812* are shown in Fig. 2g and i, respectively. The results show that an altered protein e was transduced into strain CE1188 together with proA⁺. Therefore, the mutations resulting in an altered electrophoretic mobility of protein e in the three

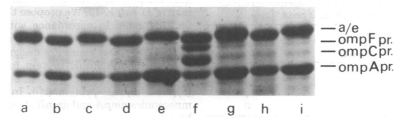


FIG. 2. Sodium dodecyl sulfate-polyacrylamide gel electrophoresis patterns of the cell envelope proteins of strain CE1108 (b, d, and h), nmpA* strain CS809 (a), nmpB* strain CS812* (c), e* strain CE1202 (e), an ilv* pho transductant of PC0479 with P1 grown on CS809 (f), a proA* TC45-resistant transductant of CE1188 with P1 grown on CS809 (g), and a proA* TC45-resistant transductant of CE1188 with P1 grown on CS812* (i). Only the relevant part of the gel showing the proteins with apparent molecular weights between 40,000 (40K) (protein e) and 35K (OmpA protein) is presented.

strains CS809, CS812*, and CE1202 are all localized at min 6, very close to *proA*. The mutations are therefore probably all in the same gene, for which we propose the name *phoE*.

To see whether the phoE gene is localized clockwise or counterclockwise from proA. threefactor transductions were carried out with strain CE1196 as the recipient and strain CS809 as the donor strain. A total of 100 pro⁺ transductants were tested, and 98 appeared to be TC45 resistant, whereas 29 were $argF^+$. This shows that argF is cotransducible with proA, although it is located more distant than phoE from proA. Therefore, the gene order is phoE proA argF or proA phoE argF. To discriminate between these two possibilities, we also selected for arg⁺ transductants in the same transduction experiment. By selecting for arg^+ transductants, $argI^+$ as well as $argF^+$ transductants can be expected. Since argI is located in a clearly distinct region of the chromosome, at min 96 (1), this influences only the absolute but not the relative cotransduction percentage of proA and phoE with arg^+ . The results of this transduction experiment are summarized in Table 3. Since proA+ is linked more closely than phoE to arg+, the gene order is phoE proA argF. Similar results were obtained when strains CS812* or CE1202 were used as donor strains.

DISCUSSION

To localize the structural gene for outer membrane protein e, mutants resistant to the protein e-specific phage TC45 were isolated. Three classes of TC45-resistant mutants were found: (i) *phoB* mutants, (ii) LPS mutants, and (iii) mutants probably altered in the structural gene for protein e. The results with TC45-resistant LPS mutants (Table 2) show that a considerable part of the LPS molecule is somehow required for the presence of normal amounts of protein e in the outer membrane. Heptose-proficient LPS

TABLE 3. Three-factor transduction involving phoE, argF, and proA^a

Unselecte	ed marker	No. of transductants	
phoE	proA	No. of transductants	
+	_	150	
_	+	46	
+	+	4	
		0	

^a P1 transduction was carried out with strain CS809 ($phoE\ proA^+\ argF^+$) as donor, with strain CE1196 ($phoS\ phoE^+\ proA\ argF\ argI$) as the recipient, and with selection of Arg⁺ transductants. The presence of the wild-type or mutant phoE allele was determined by testing for sensitivity to phage TC45.

obviously does not suffice for this purpose. An even larger part of the LPS, probably including the galactose residue, is required for phage TC45 receptor activity of protein e in vivo, consistent with and extending the observation that LPS stimulates the phage TC45 receptor activity of protein e in vitro (4).

Out of 15 strains expected to have an altered structural gene for protein e, 14 had been simultaneously mutated in proA or proB, probably because the mutations arose by deletions extending over the structural gene for protein e and one or both of these pro genes. The observation that no pro⁺ revertants of the mutant strains could be obtained supports this notion. A possible explanation for the many deletion mutations may be the presence of insertion elements in this region of the chromosome (9), since insertion elements can generate deletions (22).

Recently, Pugsley et al. (20) isolated TC45resistant mutants from nmpA and nmpB strains, and they localized the mutations in or close to the nmpA and nmpB genes, respectively. Some of their mutant strains produced a protein e with an altered mobility on sodium dodecyl sulfatepolyacrylamide gels (e* proteins). Since mutations that cause an altered protein are most likely localized in the structural gene for this protein, the authors suggested that there are two structural genes for protein e, one at min 83 and one at min 9. Since our data on TC45resistant mutants suggested a position at min 6 for the structural gene for protein e, we reinvestigated the localization of two mutations which cause an altered protein e. In addition, we localized the mutation of strain CE1202 which results in a similar phenotype. Our results showed that all of these mutations, no matter whether they had been isolated from an nmpA or an nmpB strain, were localized very close to and counterclockwise from proA. We therefore conclude that there is only one locus for the structural gene for protein e at min 6 of the chromosomal map of E. coli K-12. We propose the name phoE for this gene. In accordance with the way in which other outer membrane proteins with known structural genes are designated, we propose the name PhoE protein for the protein formerly designated as protein e (26), Ic (8), E (5), or NmpA,B protein (20). In addition, the mnemonics *nmpA* and *nmpB* should be replaced by pho.

Protein e has in common with the constitutively formed porins OmpC protein and OmpF protein that it has general pore properties (14, 21, 26). Recent experiments have shown that it is even immunologically related to the other two

porins (19). It therefore is clear that experiments with a cloned porin gene will provide information on porins in general. A potential problem with cloning of porin genes is that overproduction of porins could be lethal to the cell. However, cloning of the phoE gene may be easier than cloning of the structural genes for the other two porins as protein e is an inducible protein which usually is not expressed in a pho^+ background.

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