# Evolution of Chemotactic-Signal Transducers in Enteric Bacteria

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The methyl-accepting chemotactic-signal transducers of the enteric bacteria are transmembrane proteins that consist of a periplasmic receptor domain and a cytoplasmic signaling domain. To study their evolution, transducer genes from *Enterobacter aerogenes* and *Klebsiella pneumoniae* were compared with transducer genes from *Escherichia coli* and *Salmonella typhimurium*. There are at least two functional transducer genes in the nonmotile species *K. pneumoniae*, one of which complements the defect in serine taxis of an *E. coli tsr* mutant. The *tse* (taxis to serine) gene of *E. aerogenes* also complements an *E. coli tsr* mutant; the *tas* (taxis to aspartate) gene of *E. aerogenes* complements the defect in aspartate taxis, but not the defect in maltose taxis, of an *E. coli tar* mutant. The sequence was determined for 5 kilobases of *E. aerogenes* DNA containing a 3' fragment of the *cheA* gene, *cheW*, *tse*, *tas*, and a 5' fragment of the *cheR* gene. The *tse* and *tas* genes are in one operon, unlike *tsr* and *tar*. The cytoplasmic domain of Tse is homologous to that of Tsr, but Tas and Tar are much less similar in this region. However, several short sequences are conserved in the periplasmic domains of Tsr, Tar, Tse, and Tas but not of Tap and Trg, transducers that do not bind amino acids. These conserved regions include residues implicated in amino-acid binding.

The methyl-accepting chemotaxis proteins are the bestcharacterized signal transducers of the chemosensory apparatus in *Escherichia coli* and *Salmonella typhimurium*. Homologous proteins are found in a wide range of bacterial species (35).

Four transducers have been characterized in E. coli, and the genes encoding them have been sequenced (5, 6, 24). The Tsr transducer (taxis to serine and away from some repellents) is the receptor for the attractant L-serine and related amino acids and is responsible for chemotaxis away from a wide range of repellents, including leucine, indole, and weak acids (41, 44). Tsr also is involved in thermotaxis (26). The Tar transducer (taxis to aspartate and away from some repellents) is the receptor for L-aspartate and related amino and dicarboxylic acids; Tar also mediates taxis to the attractant maltose via an interaction with the periplasmic maltosebinding protein (14) and taxis away from the repellents  $Co^{2+}$ and Ni<sup>2+</sup> (41). The Trg transducer (taxis to ribose and galactose) interacts with the periplasmic ribose- and galactose-binding proteins to accomplish chemotaxis toward those two sugars (15, 21). The Tap transducer (taxis-associated protein) (7) mediates taxis toward dipeptides via an interaction with the periplasmic dipeptide-binding protein (28; V. Blank, diploma thesis, University of Konstanz, Federal Republic of Germany, 1987).

All transducers characterized to date are composed of the same structural domains (24). The amino terminus resembles the signal peptide of exported proteins, but it is not removed from the mature protein and serves as the first membranespanning region. The next ca. 160 amino acids extend into the periplasm, where they form the receptor domain (23). After a second membrane-spanning region, which seems to act as a stop transfer signal (27), the final 300 or more amino acids are localized in the cytoplasm. This portion of the protein generates the intracellular signals to the flagella (32, 36) and contains the glutamic acid residues that are methylated and demethylated during adaptation to chemotactic stimuli (19, 42). The amino acid sequence in regions containing the methylation sites and signaling domain is highly conserved among the four transducers (5, 24).

The genes encoding the signal transducers are located at different points on the *E. coli* chromosome (1); *tsr* at 99 min, *tar* and *tap* together in the *meche* operon at 41 min, and *trg* at 31 min. The genes are part of the flagellar regulon (20), and their promoters contain the consensus sequence present in flagellar operons that require an alternate  $\sigma$  factor for expression (2, 16, 17).

The tar gene of S. typhimurium also has been sequenced (37). S. typhimurium Tar mediates aspartate but not maltose taxis (9, 31). S. typhimurium also lacks the Tap transducer and therefore shows no taxis toward dipeptides (28). S. typhimurium does respond to serine, ribose, and galactose, so it probably has transducers equivalent to Tsr and Trg. Enterobacter aerogenes and Klebsiella pneumoniae are somewhat more distantly related to E. coli and S. typhimurium than the last two species are to each other (38). E. aerogenes exhibits chemotaxis to aspartate, maltose, and serine (9; M. Dahl, diploma thesis, University of Konstanz, Federal Republic of Germany, 1985), whereas K. pneumoniae is nonmotile.

This work was initiated on the premise that comparison of the transducers from the enteric bacteria will contribute to the study of these proteins. Genes encoding functional transducers were identified in phage  $\lambda$  libraries of chromosomal DNA from *E. aerogenes* and *K. pneumoniae*. We sequenced 5 kilobases (kb) of DNA that encodes serine and aspartate transducers and adjacent chemotaxis-related genes. Analysis of this DNA sequence and of the predicted amino acid sequences derived from it provides information about the function and evolution of proteins involved in chemosensing.

## MATERIALS AND METHODS

**Reagents.** Restriction endonucleases, T4-DNA ligase, Klenow enzyme (DNA polymerase I large fragment), the kit for

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Strain Known markers and properties				
Bacterial strains				
E. coli		39		
LE392	F <sup>-</sup> supF supE hsdR galK trpR metB tonA			
LE392.23	LE392 $\Delta(argF-lac)U169$	39		
RM41	Q358 $F^-$ (r <sup>-</sup> m <sup>+</sup> ) supE $\phi$ 80 <sup>r</sup>	18		
RM42	Q359 P2 lysogen	18		
TG1	pro thi hsd (r <sup>-</sup> ) lac/F' lacI <sup>q</sup> lacZ $\Delta$ M15 ara-14 $\Delta$ (argF-lac)U169 his-4 leuB6 metF159 <sub>am</sub> mtl-1 rpsL136 $\Delta$ (tar-tap)5201 thi-1	8		
VB12	28			
E. aerogenes ATCC 13048	Wild type	13		
K. pneumoniae KAY2026	Wild type	40		
Phage				
M13mp10	M13 sequencing vector	29		
M13mp18	M13 sequencing vector	34		
M13mp19	M13 sequencing vector	34		
λgt4-lac5	cI857(Ts), $lacZ^+$ lacY <sup>+</sup> , helper phage for lysogenizing with $\lambda$ SE6	10		
λŠE6	Kan <sup>r</sup> , low-copy-number phasmid	11		
λSE6-M1	$\lambda$ SE6 carrying <i>E. aerogenes</i> DNA containing 'cheA, cheW, tse, tas, and cheR'	This study		
Plasmids				
pBR322	Ap <sup>r</sup> Tc <sup>r</sup>	4		
pJFG5	pBR322 carrying E. coli tsr	12		
pRK41	pBR322 with 2.4-kb ClaI fragment from S. typhimurium carrying tar,	37		
pMK1	pBR322 with 3.3-kb EcoRI-Aval fragment with E. coli 'cheA, cheW, tar, and tap'	22		
pMD5	pBR322 with 5.0-kb Nrul fragment from $\lambda$ SE6-M1 carrying 'cheA, cheW, tse, tas, and cheR' from E. aerogenes			
pMD6	pBR322 with the 3.5-kb <i>Nde</i> I fragment from pRK41 and the 1.8-kb <i>Nde</i> I fragment from pMK1			
pMD7	pBR322 with the 3.2-kb <i>Ndel</i> fragment from pRK41 and the 4.5-kb <i>Ndel</i> fragment from pMK1			
pMD8	pBR322 with the 2.3-kb <i>Eco</i> RI-StuI fragment from pMD5 carrying 'cheW, tse, and tas'			

TABLE 1. Strains and plasmids

in vitro packaging of phage  $\lambda$  DNA, and reagents for the M13 dideoxy nucleotide sequencing system were purchased from Boehringer Mannheim Biochemicals, Indianapolis, Ind. Isopropyl-1-thio- $\beta$ -D-galactopyranoside was purchased from Serva. The *lacZ* oligonucleotide primer (17-mer) was obtained from Bethesda Research Laboratories, Inc., Gaithersburg, Md., or Boehringer. Deoxy-adenosine-5'- $\alpha$ -[<sup>35</sup>S]thiotriphosphate was purchased from Amersham Corp., Arlington Heights, Illinois. All other chemicals were of reagent grade.

**Strains and plasmids.** The bacterial and phage strains and plasmids used in this work are listed in Table 1. *E. coli* VB12 ( $\Delta tsr7021 \ \Delta tar-tap5201$ ) is defective in serine, aspartate, maltose, and dipeptide taxis because of the deletion of the three transducer genes. This strain does not form normal swarms in tryptone soft agar (41) because its steady-state run-tumble behavior is strongly skewed toward running due to the loss of signal input from the missing transducers.

Media and swarm plates. Cells were grown in Luria broth (LB) or minimal medium A supplemented with 0.2% (wt/vol) carbon source, required amino acids at 100  $\mu$ g/ml, and ampicillin or kanamycin at 50  $\mu$ g/ml as needed. Media were prepared by the method of Miller (30). Tryptone swarm plates contained 0.3% Bacto-Agar (Difco Laboratories, Detroit, Mich.), 1% Bacto-Tryptone (Difco), and 0.8% NaCl. Minimal swarm plates contained 0.3% Bacto-Agar, motility salts [10 mM potassium phosphate (pH 7.0), 1 mM

 $(NH_4)_2SO_4$ , 1 mM MgSO<sub>4</sub>, 0.5% (wt/vol) NaCl, and 0.5 µg of FeCl<sub>3</sub> per ml], and 100 µM maltose, L-aspartate, or L-serine. Aspartate and serine swarm plates also contained 1 mM glycerol. Ampicillin or kanamycin was added to swarm plates to a final concentration of 50 µg/ml. The plates were scored after 12 to 20 h of incubation at 32°C.

Cloning of genes coding for chemotactic-signal transducers from E. aerogenes and K. pneumoniae. Chromosomal DNA from E. aerogenes ATCC 13048 and K. pneumoniae KAY2026 was prepared as described previously (39). Genomic libraries were constructed in phage  $\lambda$ SE6 by the method of Elledge and Walker (11), using the E. coli strains RM41 and RM42. These libraries were used to infect strain VB12 ( $\Delta tsr \Delta tar-tap$ ) lysogenized with the helper phage  $\lambda$ gt4-lac5, and a 100-µl sample of washed, infected cells (ca.  $5 \times 10^8$  cells) was applied in a 5-cm-long trough on swarm plates containing kanamycin. With the E. aerogenes library, minimal-aspartate swarm plates were used. With the K. pneumoniae library, tryptone swarm plates were used. Within 24 h of incubation at 32°C in minimal agar or 16 h in tryptone agar, chemotactic swarms spreading away from the trough appeared on some of the plates. Single colonies were isolated from these swarms on LB-kanamycin plates, and phage lysates were prepared from well-aerated cultures of these cells growing in LB medium. Phage production was induced by a 15-min shift from 32°C to 42°C, and lysis of the cells proceeded during continued incubation of the cultures

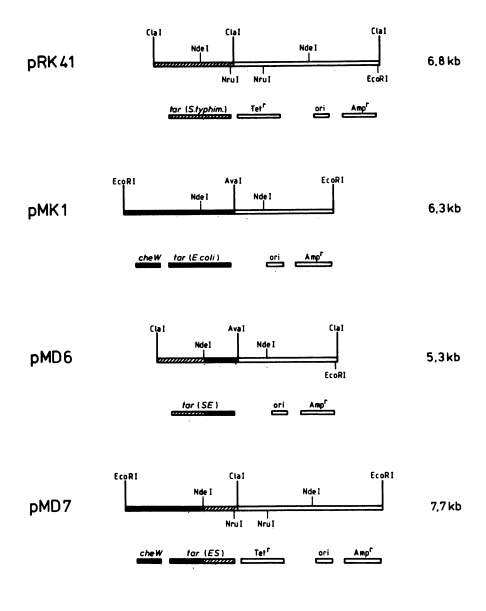
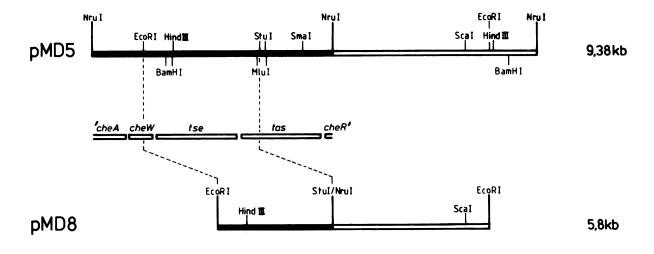


FIG. 1. Construction of *E. coli-S. typhimurium* hybrid *tar* genes. Plasmid pMD6 was made by ligating the 3.5-kb NdeI fragment of pRK41 that contains the 5' end of *S. typhimurium tar* with the 1.8-kb NdeI fragment from plasmid pMK1 that contains the 3' end of *E. coli tar*. Plasmid pMD7 was made by ligating the 4.5-kb NdeI fragment of pMK1 that contains the 5' end of *E. coli tar* with the 3.2-kb NdeI fragment from pRK41 that contains the 3' end of *S. typhimurium tar*. *E. coli* DNA is indicated by black bars, *S. typhimurium* DNA is indicated by hatched bars, and pBR322 DNA is indicated by white bars. The genes carried by each plasmid are indicated below the corresponding restriction map.

at 38°C. The  $\lambda$ SE6 and helper phage were separated by plating dilutions of these lysates with *E. coli* LE392.23 on agar containing 5-bromo-4-chloro-3-indolyl- $\beta$ -D-galactoside;  $\lambda$ SE6 forms white plaques under these conditions, whereas  $\lambda$ gt4-*lac5* forms blue plaques.

From the *E. aerogenes* library, 10 independently isolated, plaque-purified phages were retested to confirm that they conferred an aspartate-chemotaxis-positive phenotype upon infecting strain VB12. One of these phages,  $\lambda$ SE6-M1, was chosen for further analysis. From the *K. pneumoniae* library, 20 plaque-purified isolates were tested for their ability to complement mutations in *E. coli* transducer genes. Phage  $\lambda$  lysates and DNA were prepared by the methods of Silhavy et al. (39). Determination of nucleotide sequence. DNA sequencing was performed by the modified dideoxy-chain termination method of Biggin et al. (3) with deoxyadenosine-5'- $\alpha$ -[<sup>35</sup>S]thiotriphosphate. DNA restriction fragments for sequencing were produced by digesting the purified 5-kb NruI fragment of plasmid pMD5 with the enzymes BamHI, ClaI, EcoRI, EcoRV, HaeIII, HinCII, HindIII, PvuII, Sau3A, SmaI, and StuI. These fragments were ligated into the polycloning sites of DNA from phage M13mp10, M13mp18, and M13mp19 digested with appropriate enzymes. M13 phage were propagated on E. coli TG1.

Computer analysis. The management of sequence information and the subsequent analysis of compiled data were done with the Macintosh DNA Inspector II program (Microsoft



## 1 kb

FIG. 2. Restriction map of the plasmids pMD5 and pMD8. The pBR322 DNA is indicated by the white bar, and *E. aerogenes* DNA is shown as a black bar. The genes present on the plasmids are shown between the two restrictions maps.

Basic Runtime, version 2.11; Microsoft Corp.). The DNA Translate program (Mac 1.25) was kindly provided by J. S. Parkinson (Biology Department, University of Utah).

#### RESULTS

Maltose taxis in cells making hybrid Tar proteins. The difference in the abilities of E. coli and S. typhimurium to sense maltose is logically attributed to differences in the periplasmic domains of Tar from the two species (9, 31), but this suspicion had not been confirmed experimentally. We decided to utilize chimeric transducers, which were previously used to localize functional domains within the proteins (23), to test this inference.

Construction of the hybrid genes (Fig. 1) took advantage of the NdeI site in the tar genes from E. coli and S. typhimurium. Plasmid pMD6 codes for a protein containing the N-terminal 256 residues from S. typhimurium Tar and the C-terminal 297 residues from E. coli Tar, whereas the protein coded by plasmid pMD7 has the N-terminal 256 residues from E. coli Tar and the C-terminal 296 residues from S. typhimurium. These two plasmids were transformed into E. coli VB12 ( $\Delta tsr7021 \Delta tar-tap5201$ ), and the transformants were tested on minimal-aspartate and minimal-maltose swarm plates containing ampicillin.

Cells containing pMD6 formed normal swarm rings only with aspartate, whereas strains containing pMD7 formed swarm rings on both types of plates. Thus, the inability of *S*. *typhimurium* to respond to maltose is a property of the first 256 amino acids of Tar, which encompass the periplasmic domain.

**Transducer genes from K.** *pneumoniae.* The differences in transducer identity and function between E. coli and S. *typhimurium* raise a question about the transducer complement of other enteric bacteria. K. *pneumoniae* presents an especially intriguing case, since cells of this species are nonmotile. Twenty phage isolated from a K. *pneumoniae* genomic library restored the ability of E. coli strain VB12 to form swarm rings in tryptone soft agar. Lysogens of strain VB12 containing these phage were then screened on serine, aspartate, and maltose swarm plates (data not shown). Five

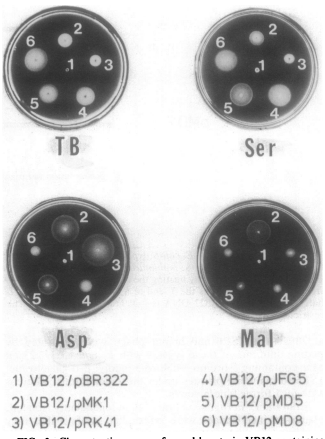


FIG. 3. Chemotactic swarms formed by strain VB12 containing different plasmids. The plasmid in each strain is indicated by the number next to the swarm as follows: 1, pBR322; 2, pMK1; 3, pRK41; 4, pJFG5; 5, pMD5; 6, pMD8. The swarm plates used were tryptone (TB), minimal-serine (Ser), minimal-aspartate (Asp), and minimal-maltose (Mal). All plates contained 50 µg of ampicillin per ml.

phage presumably carry a serine transducer gene, since they confer the ability to form swarm rings in serine soft agar but not in aspartate or maltose soft agar. Fifteen phage did not restore swarm rings with these three attractants. These phage must carry genes for transducers mediating responses to other compounds present in the tryptone extract.

**Transducer genes from** *E. aerogenes.* Unlike *K. pneumoniae*, *E. aerogenes* is motile. It also responds to many of the same attractants as *E. coli*, suggesting that these two species should have at least some similar transducers. To compare the structure and function of a related transducer from the two bacteria, we set out to clone the *tar* gene equivalent from *E. aerogenes*.

Phage  $\lambda$ SE6-M1, isolated from the *E. aerogenes* genomic library, is able to complement the defect in aspartate taxis of strain VB12. DNA prepared from this phage was digested with *Nru*I, and a 5-kb fragment was ligated into the *Nru*I site of plasmid pBR322 to yield pMD5 (Fig. 2). This plasmid and the reference plasmids pBR322, pJFG5 (*E. coli tsr*<sup>+</sup>), pMK1 (*E. coli tar*<sup>+</sup>), and pRK41 (*S. typhimurium tar*<sup>+</sup>) were transformed into *E. coli* VB12, and the chemotactic phenotypes of the transformants were tested on swarm plates (Fig. 3). Plasmid pMD5 restored the ability to form swarm rings with both aspartate and serine, but not with maltose. None of the plasmids restored the ability to form swarm rings with L-prolyl-L-leucine, a good attractant for dipeptide chemotaxis (data not shown).

DNA sequence analysis (described below) revealed that the 5-kb NruI fragment in pMD5 contained two genes that encode transducers. One of these genes, including its promoter, was located on a 2.3-kb EcoRI-StuI restriction fragment that was inserted into the EcoRI and NruI sites of pBR322 to generate pMD8 (Fig. 2). This plasmid was transformed into strain VB12, where it restored the ability to form swarm rings with serine but not with aspartate (Fig. 3). The transducer gene on plasmid pMD8 was named *tse* (taxis to serine), because it complemented the defect in serine taxis caused by a *tsr* mutation. The transducer gene present on pMD5 but not on pMD8 was named *tas* (taxis to aspartate) because it was required to complement the defect in aspartate taxis caused by a *tar* mutation.

Nucleotide sequence of chemotaxis-related genes from E. aerogenes. The nucleotide sequence of the 5-kb NruI restriction fragment from plasmid pMD5 was determined from both strands between two and five times, and the amino acid sequences of the five longest open reading frames were predicted (Fig. 4). Based on the similarity of these sequences to their homologs in E. coli (33), three of the open reading frames correspond to cheW and fragments of the 'cheA and cheR' genes. The open reading frames corresponding to tse and tas were identified by their resemblance to transducer genes from E. coli and S. typhimurium.

Comparison of Che proteins from E. aerogenes and E. coli. The open reading frame for the 3'-terminal portion of cheA from E. aerogenes (Fig. 4) codes for 226 amino acids, of which 176 (78%) are identical to residues within the last 226 amino acids of CheA from E. coli (33). One stretch of 19 amino acids has only one match, but we do not believe our sequence is in error because neither alternate reading frame in this region gives a better match. Without these 19 residues the percent identity rises to 85% (175 of 207 amino acids).

The 5' region of *cheR* in our sequence contains an open reading frame coding for 100 amino acids starting with the ATG codon at nucleotides 4715 through 4717 (Fig. 4). This codon is preceded by the sequence GAGCT at nucleotides 4707 through 4711, which may serve as a ribosome-binding

site. (The C does not match the Shine-Dalgarno consensus sequence.) Alternatively, translation could start at the ATG codon at nucleotides 4769 through 4771. Significant homology between the predicted CheR amino acid sequences of E. *aerogenes* and E. *coli* begins at the glutamine preceding the second methionine residue. Of the next 82 residues, 68 (83%) are identical in the two species. The sequences preceding this glutamine bear little similarity in the two species.

The predicted amino acid sequence of the CheW protein of *E. aerogenes* is 165 residues long and begins with the methionine coded by the ATG triplet at nucleotides 715 through 717 (Fig. 4). This protein is 77% identical to CheW from *E. coli* and 79% identical to CheW from *S. typhimurium* (Fig. 5). The CheW proteins of the latter two species are 92% identical.

Comparison of Tse and Tas with signal transducers from E. coli and S. typhimurium. The predicted amino acid sequences of Tse and Tas can be aligned with the sequences of four other signal transducers (Fig. 6). The amino acid sequences of the periplasmic domains of the six transducers are compared in Table 2. Tse and Tsr are more like each other than any of the other transducers, being almost as similar as the E. coli and S. typhimurium Tar proteins. Surprisingly, the periplasmic portion of Tas has a low percent identity with all of the other transducers, exhibiting similarities only in a few, limited regions. The periplasmic portion of Tap diverges even more widely.

# DISCUSSION

The properties of *E. coli-S. typhimurium* Tar hybrids suggest that differences in the N-terminal halves of the two proteins (presumably in their periplasmic receptor domains) determine whether they function in maltose taxis. Since *E. aerogenes* carries out both aspartate and maltose taxis (19), the periplasmic domain of its Tar transducer might share structural features with *E. coli* Tar that contribute to its interaction with maltose-binding protein. These features could be recognizable as amino acid sequences present in *E. coli* and *E. aerogenes* Tar but not in *S. typhimurium* Tar. Our interest in making this comparison led us to isolate transducer genes from *E. aerogenes*. As the work progressed, the broader goal emerged of using a comparative approach to study the evolution of the signal transducers.

The nonmotile bacterium K. pneumoniae contains at least two transducer genes that can function in E. coli. One of them codes for a serine transducer. No defined substrates have been identified for the other K. pneumoniae transducers, which overcome the general chemotactic defect of strain VB12.

Why have transducer genes been retained in a nonmotile species? In *E. coli*, the transducer genes are expressed late in the regulatory cascade of the flagellar regulon (2, 20), and their expression is blocked by *fla* mutations that inhibit flagellar synthesis. If comparable mutations have occurred in *K. pneumoniae*, then the transducer genes would not be expressed in that species, and the selective pressure to delete them would be relatively low.

Two transducer genes were found on a 5-kb DNA fragment from *E. aerogenes*. The *tse* gene encodes a serine transducer, since it complements an *E. coli tsr* mutation. The *tas* gene is required to restore aspartate taxis in an *E. coli tar* mutant. Since we do not have a construct in which *tas* is expressed in the absence of *tse*, we have not demonstrated that Tas alone functions as an aspartate transducer, although that is the simplest assumption.

CTC GCC ANA GCG GCG CAG GGG CTG GCG GTC ACG GAC ACG ATC ACT GAT GAN GAG GTC GGA ATG CTT ATT TTT GCG CCG GGC TTT TCA ACC GCG GAA Leu Ala Lys Ala Ala Ala Gln Gly Leu Ala Yal Thr Asp Thr <u>Het Ser Asp</u> Glu <u>Glu Yal</u> Gly <u>Het Leu Ile Phe Ala Pro Gly Phe Ser Thr Ala G</u>U ACC GAC GTC TCT GGC GGC GTC GGC ATG GAC GTC GTC AAA CG ATA TT CAG GAG ATG GGC GGT CAC GTA GAA ATC CAT TCC CGT GCG GGC AAA Thr Amp Val Ser Gly Arg Gly Val Gly Met Amp Val Val Lys Arg Amm The Gln Glu Met Gly Gly Him Val Glu The Him Ser Arg Ala Gly Lys GGG ACC <u>Gly Thr</u> 243 270 297 324 TTG TTG CCG CTA ACG CTC GCC ATC CTC GAC GGC ATG TCG GTC AAG GTC AAT GAA GAG GTC TTT ATT CTG CCG CTC AAC GCG GTG ATG GAA TCG CTG Leu Leu Pro Leu Thr Leu Ala Ile Leu Asp Gly Met Ser Val Lys Val Asn Glu Glu Val Phe Ile Leu Pro Leu Asn Ala Val Met Glu Ser Leu ATT CGT ATT CAG CCG CAG CGC GAA GAC CTG CAT CCA ATG GCC GGC GGC GAG CGG ATG CTG CAG GTT CGC GGC GAG TAT CTA CCG CTG GTG GAG CTC TAC CGG GTG TTT GAA GIn Pro Gin Arg Glu Asp Leu His Pro Met Ala Gly Gly Glu Arg Met Leu Gin Val Arg Gly Glu Tyr Leu Pro Leu Val Glu Leu Tyr Arg Val Phe Glu TGT GCC Cys <u>Ala</u> GGG GCG ANA ACC GAG GCC ACT CAG GGC ATC GTG GTG ATT CTG CAA AGC GCC GGC CGC CGT ANT GCG CTG GTG GTG GAT CAA CTG ATC GGC CAG CAC CAG<sup>C/</sup>TGT GTG Gly Ala Lye Thr Glu Ala Thr Gln Gly lle Val Val Ile Leu Gln Ser Ala Gly Arg Arg Asn Ala Leu Leu Val Asp Gln Leu Ile Gly Gln His Gln Cys Val Lys GAA ACG AAT TAC CGC AAA GTG CCG GGA ATT TCC GCG GCG ACG ATC CTC GGC GAC GGC AGC GTG GCG CTG ATC GTC GAC GTG TCG GCG CTG CAA ATG CTC AAT Glu Thr <u>Asn Tyr Arg Lys Val Pro Gly Ile Ser Ala Ala Thr Ile Leu Gly Asp Gly Ser Val Ala Leu Ile Val Asp Val Ser Ala Leu Gln</u> Met Leu <u>Asn</u> 675 702 <u>chem</u> 729 756 CGG GAA AAG CTG CTG AGC GCA GCG GCC GCA TAA CGA CTC ATC TCA TCA AAT TAA CTG GTG CAG ACC ATG GCA GGA TTA GCA ACC GTC AGC AAA TTG GCT GGC GAA ACG <u>Arg Glu</u> Lys Leu Leu Ser Ala Ala <u>Ala Ala</u> \* MET Ala Gly Leu Ala Thr Val Ser Lys Leu Ala Gly Glu Thr GTA GGT CAG GGG TTT TTA ATC TTT ACC CTC GGC AAT GAA GAA TAC GGC ATC GAT ATC CTG AAA GTG CAG AAG ATC CGC GGC TAT GAC CAG GTG ACG CGG ATC GCC AAC Val Gly Gln Ala Phe Leu Ile Phe Thr Leu Gly Asn Glu Glu Tyr Gly Ile Asp Ile Leu Lys Val Gln Lys Ile Arg Gly Tyr Asp Gln Val Thr Arg Ile Ala Asn 912 ACC CCG GAT TTC ATC AAA GGC GTC ACC AAT CTG CGC GGG GTG ATC GTG CGG ATT ATC GAC CTG CGG GTA AAA TAT GCC CAG GGC GTC TCT TAT GAT GAA AAC ACG Thr Pro Asp Phe Ile Lys Gly Val Thr Asn Leu Arg Gly Val Ile Val Pro Ile Ile Asp Leu Arg Val Lys Tyr Ala Gln Gln Gly Val Ser Tyr Asp Glu Asn Thr 999 1026 1053 1080 GTG GTT ATC GTG CTT AAC TTC GGC CAG CGG GTG GTG GGG ATT GTG GTC GAC GGG GTC TCC GAC GTG TTG TCT CTT ACC GCC GAA CAG ATC CGC CCG GCG GCA TTC Val Val Ile Val Leu Asn Phe Gly Gln Arg Val Val Gly Ile Val Val Asp Gly Val Ser Asp Val Leu Ser Leu Thr Ala Glu Gln Ile Arg Pro Ala Pro Glu Phe 1184 CCA GTG ACG ATG GCG ACC GAA TAT CTC ACC GGT CTT GGC GCG CTC GGA GCG CTG TTG ATC CTT GTG GAT ATC GAA AAG CTC ACC GAA GAG ATG GCG CTC GTC Ala Val Thr Met Ala Thr Glu Tyr Leu Thr Gly Leu Gly Ala Leu Gly Ala Leu Leu 11e Leu Val Asp 11e Glu Lys Leu Leu Ser Thr Glu Glu Met Ala Leu Val GAT AAC GTC GCC AAA AGC CAC TAA GCA ATC GGG CGC GCC GGT AAA AAT AGT CCC CGC CTG GC<u>T AAA</u> GTT CCC CTC CCG TAC <u>GCC GAT AAC C</u>CT TTC AGT CAC ATA CGT Asp Asn Val Ala Lys Ser His \* 1324 LEE 1352 1379 1 ANA GCC TGG CCG TTC AGG TTC <u>CAGGAAGG</u>GGA AAC ATG TTT AAT CGT ATT AAG GTC GTC ACC AGT CTC TTA TTA GTG CTG GTG CTA TTT GGC GCA TTG CAG CTG ATT SD NET Phe Asn Arg Ile Lys Val Val Thr Ser Leu Leu Val Leu Val Leu Phe Gly Ala Leu Gin Leu Ile GGC GGT CTG TTT TTT TCG TCG TCG AAA GGC GAT AAA GAG AAC TTT ACC GTC CTG CAA ACC ATC CGT CAG CAG CAG TTG CTG CTG AGT GAA AGT CGG GTC GAT Gly Gly Leu Phe Phe Ser Ser Leu Lys Gly Asp Lys Glu Asp Phe Thr Val Leu Gin Thr 11e Arg Gin Gin Gin Leu Leu Ser Giu Ser Arg Val Asp CAG GCG CGT AAC TCC CTG AAC CGC GCA GGG ATC CGC TAC ATG ATG GAT ACC AAA ATC GGC AGC GGC GCG ACT ATC GAC GAG CTG CTG GCG AAA GCG GAA AAA Gin Ala Arg Asn Ser Leu Asn Arg Ala Gly Ile Arg Tyr Met Met Amp Thr Asn Lys Ile Gly Ser Gly Ala Thr Ile Asp Glu Leu Leu Ala Lym Ala Glu Lym AAG CTG GCG CGC GAG CGC AAC TAC ACC GCC TAT GAA AAA ATC CCG CAG GAC CCG CAG GAT CCT CAG GCG ACG GAA AAG CTT AAG CAG CAG TAT GGC ATC CTG Lys Leu Als Arg Als Glu Arg asn Tyr Thr Als Tyr Glu Lys Ile Pro Gin Asp Pro Gin Asp Pro Gin Als Thr Glu Lys Leu Lys Gin Gin Tyr Gly Ile Leu

FIG. 4. Nucleotide sequence of the 'cheA-cheW-tse-tas-cheR' region from E. aerogenes. The predicted amino-acid sequences for the longest open reading frames are given below. The amino acids underlined in the 'cheA and cheR' gene fragments are identical with residues in the corresponding genes of E. coli. The inverted triangles in 'cheA indicate single amino acid deletions relative to E. coli, and the bracket denotes an inserted amino acid. The underlined sequence designated "flagellar operon consensus" corresponds to the consensus sequence found in the promoters of flagellar operons in E. coli (2, 16). The underlined sequences designated SD refer to regions surrounding the predicted Shine-Dalgarno sequences for the tse and tas genes.

Since *E. aerogenes* responds chemotactically to maltose and synthesizes maltose-binding protein (9), it should have a maltose transducer. Our attempt to isolate this transducer was based on the assumption that it should also be an aspartate receptor. In *E. aerogenes*, however, maltose taxis could be mediated by a minor transducer (like Trg or Tap in *E. coli*) that is unable to restore normal chemotactic behavior to the smooth-swimming VB12 strain. Thus, a search for the gene encoding the maltose transducer of *E. aerogenes* should be repeated with a  $tsr^+$  tar *E. coli* strain, which shows normal swimming behavior but no aspartate or maltose taxis.

The genes on the 5-kb DNA fragment from *E. aerogenes* appear in the order 5' *cheA-cheW*-promoter-*tse-tas-cheR* 3' (Fig. 4). This segment spans the 3' and 5' ends of the *E. aerogenes* equivalents of the *E. coli mocha* and *meche* operons (33). In *E. coli* the first two genes in the *meche* operon are *tar* and *tap*, whereas in *S. typhimurium tar* is the sole transducer gene in the *meche* operon. The *che* genes are in the same relative location in all three species.

The 36 nucleotides between cheA and cheW in *E. aero*genes show no obvious similarity to the corresponding 23-nucleotide regions in *E. coli* or *S. typhimurium*. The untranslated region between cheW and the start codon of the first gene in the *meche* operon also varies considerably among the three species. This region contains 122 nucleotides in *E. aerogenes*, 147 nucleotides in *E. coli*, and 243 nucleotides in *S. typhimurium*. The proposed rho-independent site for transcription termination distal to *cheW* in *E. coli* is missing in *E. aerogenes*, as is the perfect 13-base inverted repeat found 3' to the tandem TAA stop codons of *cheW* in *S. typhimurium*.

The region (underlined in Fig. 4) around the Shine-Dalgarno sequence of the first gene in the *meche* operon is conserved in all three species. So is the consensus sequence (also underlined in Fig. 4) identified in promoters of flagellar operons of *E. coli* (2, 16). Conservation of these regulatory sequences is consistent with the observation that the *tse* and *tas* genes are expressed in *E. coli*. No significant homology was detected between the *tse-tas* (85 base pairs) and *tar-tap* (48 base pairs) or the *tas-cheR* (91 base pairs) and *tap-cheR* (16 base pairs) intergenic regions.

Some regions of the predicted amino acid sequences of the 'CheA and CheR' polypeptide fragments of *E. aerogenes* are more similar to their counterparts in *E. coli* than others (Fig. 4). The highly conserved sequences presumably represent regions of special structural and functional importance. The CheW proteins from *E. aerogenes*, *E. coli*, and *S. typhimu*-

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TAC GGC GGC GTG TCG GAG GTG TAC CAG GTG CTG GGC GGA GGG TGT AAA ATC AAC GCC TTC TTC GAC CAG CCG ACG AAAA TAC CAG GAC GAT TTC GAG CAG ACC TAT Tyr Gly Ala Leu Ser Glu Leu lle Gin Leu Leu Gly Gly Gly Cys Lys Ile Asm Ala Phe Phe Asp Gin Pro Thr Gin Lys Tyr Gin Asp Asp Phe Glu Gin Thr Tyr AAC GCC TAT CTG CAG CAA AAC GGC AAG CTG TAT CAA ATC GCC GTT GAC GCT AGC AAC AGC TCG TAT AGC TCG GCG ATC TGG ACC TTA ATT GTG GTC ATC ATC ATC GTG GTG Asm Ala Tyr Leu Gin Gin Asm Giy Lys Leu Tyr Gin Ile Ala Val Asp Ala Ser Asm Ser Ser Tyr Ser Ser Ala Ile Trp Thr Leu Ile Val Val Ile Ile VI Val Val CTG GCG GCG ATC GTC GGC GTG GTG GTT GGC TATC CAC CAT ATC CTG GTG CGC CTG CTG AAC CGC ATG ATA GAA CAC ATC AAA CGG ATC GCG CTG GGC GTC GGCG 2081 2162 CCT ATT CCG GTG ACC AGC CCT ATG GAA ATC GGC GTG CTG GCG GCC AGC CTC AAG GAC ATG GAA ACG GTA AGC GGA GGC GGG GGG GGG GAT PTO 11e PTO Val Thr Ser Arg Aan Glu 11e Gly Val Leu Ala Ala Ser Leu Lys His Met Gln Aan Glu Leu 11e Glu Thr Val Ser Gly Val Arg Gln Gly Ala Asp GTC GAA ACC ATG CAC GAC ATC GCC AGC AGC TCG CAG AAA ATT GCC GAT ATC ACC GGC GTT ATC GAC GTA ATC GCC TTC CAG ACC AAC ATC CTG GCG CTT AAC GCC GCG Val Glu Thr Met His Asp Ile Ala Ser Ser Ser Gln Lys Ile Ala Asp Ile Thr Gly Val Ile Asp Val Ile Ala Phe Gln Thr Asn Ile Leu Ala Leu Asn Ala Ala 2513 2594 GTA GAA GOG GOG GOT GOG GGT GAA CAA GGT CGC GOG TTC GOG GTC GTG GOG GGC GAA GTC CGC AAC CTG GOG CAG GCC CAG GOG GCC AAA GAG ATC AAA GOG Val Glu Ala Ala Arg Ala Gly Glu Glu Glu Gly Arg Ala Phe Ala Val Val Ala Gly Glu Val Arg Asn Leu Ala Gln Arg Ser Ala Gln Ala Ala Lys Glu 11e Lys Ala 2621 2675 CTG ATC GAA GAC TCG GTC AAC CGT GTG GAT ATG GGC TCG GTA CTG GTG GAA AGC GCC GGC GAC ACC ATG GGC GAT ATC GTC AAT GCG GTC ACC CGC GTG ACC GAC ATC Leu lie Glu Asp Ser Val Asn Atg Val Asp Met Gly Ser Val Leu Val Glu Ser Ala Gly Asp Thr Met Gly Asp Tie Val Asn Ala Val Thr Arg Val Thr Asp Tie 2729 2756 2783 2810 ATG GGT GAA ATC GCC TCT GCT TCC GAT GAA CAG AGC CGC GGT ATC GAC CAG GTC GGC CAG GCG GTA CGG GAA ATG GCT TCC GAT GAA CAG AAC GCC TCG CTG GTC Met Gly Glu Ile Ala Ser Als Ser Asp Glu Gln Ser Arg Gly Ile Asp Gln Val Gly Gln Ala Val Arg Glu Met Asp Arg Val Thr Gln Gln Asm Ala Ser Leu Val GAG GAG TCG GCC TCG GCC GCC GTG GAA GAG CAG GCC AGG CTG CTG ACG CAG TCG GTT GCC GTC TTC CGC CTG AAG TCG GAA GGG CAG GAA GAG TAT AAA GCG CCG Glu Glu Ser Ala Ser Ala Ala Leu Glu Glu Glu Gln Ala Ser Leu Leu Thr Gln Ser Val Ala Val Phe Arg Leu Lys Ser Glu Gly Gln Glu Glu Tyr Lys Ala Pro 2945 2972 2999 3026 GTC AGC AAT AAA ACC GCG CCT GCC GCC ATC GCT ACG CAT AAA AAA ACC AGC GCC AGC GAC TAC CAG GAT AAC TGG GAG ACG TTC TAA CGG GTA CCG GCC GCG ATA CCC Val Ser Asn Lys Thr Ala Pro Ala Ala Ile Ala Thr His Lys Lys Thr Ser Ala Ser Asp Tyr Gln Asp Asn Trp Glu Thr Phe \* 3053 3080 tas 3105 3132 TAC ACG CGG CGG CCT CTC TTG CGT TGT AAG CCG GCC GAG ACC GGC TAC <u>CAA GGA GA</u>T TGC T SD HET Phe Asn Arg Ile Arg Ile Ser Thr Ser Leu Phe Leu Leu Leu 3159 3186 3213 3240 ATC TCC TTT TGC ATC ATG CAA CTG ATC AGC ACC GGG CTC TCT TAC GTC CGG TAC GCG CCG ACA ACC ACA ATC TTG AAC GTA TTA CCG TCA GTA GCC AGC AGG GAT GCG Ile Ser Phe Cys Ile Met Gln Leu Ile Ser Thr Gly Leu Ser Tyr Val Arg Tyr Ala Pro Thr Thr Ile Leu Asn Val Leu Pro Ser Val Ala Ser Thr Asp Ala CTT AGC CTG AGC TGG GTA TCG CTG TTA CAG GCG AGA AAT ACC CTT AAC CGC GCC GGT ACC CGC GGC GCT GAA GGT GCA GGA GGA GGT GGA AGT GCG CTG ATG GCG Leu Ser Leu Ser Trp Val Ser Leu Leu Gin Ala Arg Asn Thr Leu Asn Arg Ala Gly Thr Arg Gly Ala Glu Gly Ala Ala Gly Ala Gly Gly Ser Ala Leu Met Ala GGG CCC GCA GCT CGC TGC AAA AAG CCG ATC TCT ATT TTA ACC AGT TCC TCG ACA CGC CTC GCG CGG ATG AGC AGG AGT TGG CCG ATG CCA CGC GTG ATA GCT Ala Pro Ala Ala Arg Cys Lys Lys Pro lle Ser lle Leu Thr Ser Ser Thr Arg Leu Ala Arg Met Ser Arg Asn Ser Ser Trp Pro Met Pro Arg Val Ile Ala ACG AGA ATC TAC GCG CTG CCC GAT GTC GTC GTC GAA AAT CGC AAC CTG CAG GCC TTT ATG GAC CAG CCG ACG AAA ATA CAG GAC CGC TTT GAG GCC GAC Thr Arg Ile Tyr Ala Leu Arg Glu Leu Ile Val Phe Leu Glu Asn Arg Asn Leu Gln Ala Phe Met Asp Gln Pro Thr Gln Lys Ile Gln Asp Arg Phe Glu Ala Asp CAN TAT CTG CAG CTG GCG AAG GCG ACG ACC GAT GAA GCC AGC GCC TCC AGC CAG GAG GCC TAC GCG TTG GTT CCA ATC TGG TTG GTT GCC GGG GCG GTG CTG Gin Tyr Leu Gin Leu Ala Lys Ala Thr Thr Asp Glu Ala Ser Ala Ser Ser Gin Gin Ala Tyr Ala Leu Val Pro Ile Trp Leu Val Ala Gly Ala Val Leu CTC TTG GTA GTC ACC CTC AGC GGG ATG TGG TGG CTG CGA ACC ATG CTG GTG CAG CCG CTG AAT ATT ATT CGC GGC CGAT CTT Leu Leu Val Val Thr Leu Ser Ala Met Trp Trp Leu Arg Thr Het Leu Val Gln Pro Leu Asn Ile Ile Arg Gly His Phe Glu Arg Ile Ala Ser Gly Asp Leu TCC GCA CCG ATT GAG GTC TAT GGG GCT AAC GAA ATC AGC CAG CTA TAT GCC AGC CTG CAG CGC ATG CAA CAG TCG CTG ATT GAT ACC GTC GGC GGC GGC GGC GGC GGC GGC GAC GGC Ser Ala Pro 11e Glu Val Tyr Gly Arg Aan Glu 11e Ser Gln Leu Phe Ala Ser Leu Gln Arg Met Gln Gln Ser Leu Ile Gly Thr Val Gly Ala Val Arg Amp Gly GCG GAG TCG ATT CTT ATC GGC CTG GAG GAG ATT GCC GAA GGC AAC AAC GAT CTC TCC TCG CGC ACC GAA CAA CAG GAC GGT GGG GAA GAG ACC GCC GCC GGC AGT ATG Ala Glu Ser lie Leu ile Giy Leu Gin Glu ile Ala Glu Giy Asn Asn Asp Leu Ser Ser Arg Thr Glu Gin Gin Asp Arg Ser Leu Glu Glu Thr Ala Ala Sar Met GAG CAG CTA ACG GCG ACG GTA AAA CAG AAT GCC GAT AAC GCC CGC CAG GCA TCG CAG CTG GCG CGT GAT GCC TCC TCC TCC ACC GCC GCC AAA CGG ACG CTG GCG GAT GAC Glu Gln Leu Thr Ala Thr Val Lys Gln Asm Ala Asp Asm Ala Arg Gln Ala Ser Gln Leu Ala Arg Asp Ala Ser Ser Thr Ala Ala Lys Arg Thr Leu Ala Asp Asp 4131 4158 4212 GTG GTC ACC ACG ATG CAC GAT ATC GCC AAC AGT TCG CAA AAA ATC GGC GCG ATT ACC AGC GTC ATC GAC GGC ATC GTC TAC GAG ACC AAT ATT CTG GCG CTT AAC GGC Val Val Thr Thr Met His Asp lie Ala Asm Ser Ser Gin Lys lie Gly Ala lie Thr Ser Val lie Asp Gly lie Ala Phe Gin Thr Asm lie Leu Ala Leu Asm Ala 4239 4266 4293 4320 GCG GTG GAG GCG GCG GCC GGC GAA CAG GGG CGC GGC TTT GCG GTG GCC GGT GAA GTG CGC AAC CTG GCC AGC GCC AGG GCG GCA AAA GAA ATC AAG Ala Val Glu Ala Ala Arg Ala Gly Glu Gln Gly Arg Gly Phe Ala Val Val Ala Gly Glu Val Arg Asn Leu Ala Ser Arg Ser Ala Gln Ala Ala Lys Glu Ile Lys CTG CTG ATC GAT GAA TCC GTC AGC CGC GTC AAA CAC GGC TCG GTG GTG GTG GTG GAA AAT TCC GGC GCC ACC ATG CAG GAC ATC GTG CGC TCG GTC ACC CGG GTC ACC GAC Leu Leu lle Amp Glu Ser Val Ser Arg Val Lys His Gly Ser Val Leu Val Glu Asn Ser Gly Ala Thr Met Gln Amp Ile Val Arg Ser Val Thr Arg Val Thr Amp 4455 4482 4509 4536 ATC ATG GGC GAA ATC GCC TCC GCC TCC GAT GAG CAG AGC CGC GGC ATC GAG CAG GTG ACG CAG GCG TTA CCC AGA TAT CAG GTC ACG CAA CAG AAC GCC GCC GTG GTG Ile Met Gly Glu Ile Ala Ser Ala Ser Asp Glu Gln Ser Arg Gly Ile Glu Gln Val Thr Gln Ala Leu Pro Arg Tyr Gln Val Thr Gln Gln Asn Ala Ala Leu Val GTG GAN TOG GCT TCC GCC GCC GCG GTG GAA GAG CAG GCG GTA ACC CTC GCC GAT GCA GTG GCG GTA TTC CGT CGC GAC GAC GAC TAA TTG TCG CGC CGC GGG ATA TAC Val glu ser Ala Ser Ala Ala La Leu glu glu glu glu gli thr Leu Ala Thr Leu Ala Asp Ala Val Phe Arg Leu Ala Asp Asp \* GA CCC GGT GGA TTG TCA AAC CGC ATT AAA CGC TCG TAT TTG CCG GTG AGC TGG C ATG AAG CAA GAC GAC ATC AAC GCG GCG CGT GAA HET Lys Gin Asp Asp Iie Asn Aia Aig Giu GGA TCG GCC GTG GCG CAG ATG GCC GTC GTC GCG GTC TCC GAC GCG CAT TTT CGC CGC ATC AGC CAG CTT ATC CAG CGC GCG GGG ATC GTG GCG CGC GCA Gly <u>Ser</u> Ala Val Ala <u>Gin Met</u> Ala Gin <u>Arg Leu</u> Pro <u>Leu Ser Aep Ala His Phe Arg Arg Ile Ser Gin Leu Ile Tyr Gin Arg Ala Gly Ile Val Leu Ala</u> Pro <u>His</u> CGC GAG ATG GTG TAC AAC CGG CTG GTG CGC CGT TTA CGT CTG GGC GAC TAT CTG GGC GAC TAC CTG GCG CTG GAA AGC GAC CGC GAC AGC GCC GAC Arg Glu <u>Met yal Tyr Asn Arg Lau yal Arg Arg Lau Arg Lau Lau Gly</u> 11e His <u>Asp Phe Gly</u> Asp <u>Tyr Lau</u> Ala <u>Lau Lau Glu Ser</u> Asp Pto <u>His Ser</u> Ala <u>G</u>lu TGG CAG GCG TTT ATC AAT GCG CTG ACC ACC AAC CTG ACC GCC TTT TT C TTD GIn Als Phe Ile Asn Als Leu Thr Thr Asn Leu Thr Als Phe Phe

FIG. 4—Continued

J. BACTERIOL.

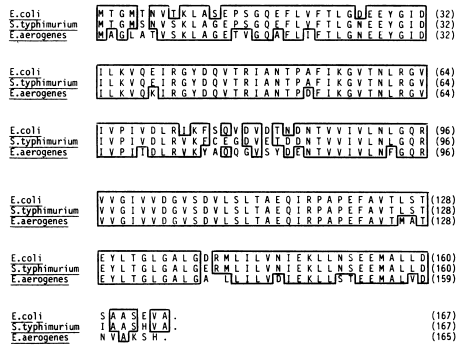


FIG. 5. Comparison of the predicted amino acid sequences of CheW from *E. coli* (33), *S. typhimurium* (43), and *E. aerogenes*. Identical residues are boxed. The  $\beta$ - $\alpha$ - $\beta$  structural motif (45) extends over residues 128 through 148 and 155 through 160.

rium also have a pattern of conserved and variable regions (Fig. 5). CheW has been proposed to contain a nucleotidebinding site (43). An 11-residue consensus sequence that determines the  $\beta$ - $\alpha$ - $\beta$  motif of the binding site has been described (45). All three species have the identical amino acid at each of the 11 positions, and 10 of the 11 residues conform to the consensus. Other highly conserved segments of CheW may correspond to sites of interaction between CheW and the signaling domains of the transducers (P. Ames, J. Chen, C. Wolff, and J. S. Parkinson, Cold Spring Harbor Symp. Quant. Biol, in press).

Tse and Tas have the typical structural features of signal transducers (Fig. 6): a positively charged N terminus (residues 1 through 6) followed by a hydrophobic membrane-spanning region (residues 7 through 30), a hydrophilic periplasmic domain (residues 31 through 190), a second hydrophobic membrane-spanning region (residues 191 through 210), and a hydrophilic cytoplasmic domain (residue 210 through the C terminus). Of 187 residues that are identical in the six transducers in Fig. 6, 168 are within the cytoplasmic domain and 145 are within a stretch of 236 amino acids (residues 280 through 516 in Tsr).

The K1 and R1 tryptic peptides of Tsr (residues 295 through 317 and 483 through 507) contain the sites of covalent methylation (19). Within the region corresponding to K1, Tse and Tsr are identical at 22 of 23 residues and Tas and Tsr are identical at 21 of 23 residues. The same three methylation sites, which are Glu-Glx sequences, are present in Tse and Tas as well as in Tsr and Tar. Within R1, Tse and Tsr are identical at 21 of 25 residues and the two methylation sites are present. Tas and Tsr are identical at 20 of 25 residues, but one of the methylation sites may be absent in Tas due to the replacement of the Glu-Glu sequence at residues 492 and 493 of Tsr with Val-Glu in Tas.

Another highly conserved stretch runs from residues 360 through 407 of Tsr. The same 48 residues are present in Tar, and there is only one difference in Tap. Tse and Tas share 44

of the 48 residues with Tsr. In Trg, only 37 of these 48 residues are identical to Tsr (5), underscoring the proposed early divergence of Trg from the other transducers. The strong conservation of this region and of the K1 and R1 peptides emphasizes their importance in signaling and adaptation, respectively (Ames et al., in press).

The Tse and Tsr proteins are 60% identical in their periplasmic domains (Fig. 6 and Table 2). Even their cytoplasmic domains resemble one another more closely than they do those of other transducers. The periplasmic domain of Tas bears little resemblance to that of any other known transducer, including Tap, which like Tas is the product of the second gene in a *meche* operon, or Tar, which is a functional analog of Tas. Given the dissimilarity of Tar and Tas, it is not surprising that Tas does not participate in maltose taxis.

The amino acid transducers Tsr, Tse, Tas, and Tar share three sites that are partially or wholly absent in Tap and Trg: (i) Arg at residues 64, 69, and 73 (Arg-64 is present in Trg); (ii) Glu-Leu-Ile at residues 138 through 140; (iii) Gln-Pro-Thr-Gln at residues 154 through 157 (Pro-155 and Gln-157 are present in Tap). Genetic studies indicate that some of these residues are important in chemoreception. E. coli mutants in which Arg-64, Arg-69, or Arg-73 of Tar is replaced by other residues have a drastically reduced ability to carry out aspartate taxis, and the apparent affinity for aspartate is reduced by a factor of  $10^2$  or more (46). Furthermore, a mutation affecting the Arg-64 residue of E. coli Tsr greatly reduces the ability of cells to sense serine (25). The three Arg residues may form a positively charged pocket that interacts with a common element of amino acids, such as their  $\alpha$ -carboxyl group (25, 46; Ames et al., in press). Other data suggest that Thr-156 of Tsr and Tar is important for serine (25) and aspartate (C. Wolff, personal communication) sensing.

Tse (with 557 residues) and Tas (with 510 residues) are the largest and smallest transducers, respectively, known in the

Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	O * * O * * O * * * * * * * * * * * * *	* * * 0 LNRAGI (72) LNRAGI (72) LNRAGI (71) LSRSAV (72) LSRSAV (72) LNKAGT (72)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	O*** O O**O O ** * O RYMMDQNNIGSGSTYAELNESASISLKQAEKNWADYEAL PRDPRQSTAAAAEIKRNYDDIYHNALA RYMMDDNNKIGSGATIDELLAKAEKEKLARAERNYTAYEKIPQDPRQDPQATBKLKQQYGILYGALS RGAEGAAGAGGSALMAAPAARCKKPISILTSSSTRLARNSRNSSWPMP RVIATRI Y ALR RMMMDSSNQQSNAKY ELLDSABKTLAQAATHYKKKSHAPLPEMVATSRNIDEK YKNYTALT RMMMDSNQQSSAKTD LLQNAKTLAQAATHYKKKSHAPLPEMVATSRNIDEK YQYAALA LTALSYPADD IKT LNTTABASLT9STTLFKSFMANTAGNEHVRGLQKETEKSF ARWHNDLE	E L I Q L L (144) $E L I V F L (136)$ $E L I D Y L (141)$
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	D O 000 *** O * ** O * 00 * **0 O O O * *** **	A         I         YG         (211)           Y         T         L         S         (203)           I         L         YA         (208)         (208)           I         L         YA         (208)         (208)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	VW PGIKASLVAPMARLIDSIBHIAGGDLVKPIBVDGSAENGALASIRHMQCALRGTVGDVRAGA VWNGIHHILVRPLARMIEHIKBIAGDLTQPIPVTSRAEIGVLAASLKHMQAELIETVSGVRQGA AWWW LRTHLVQPLAIIRGHPEBIAGGDLSAPIBVYGBNEISQLFASLQRMQQSLIGTVGAVRDGA WYGIBRALLTPLAKJIAHIBBIAGGDLSAPIBVYGBNEISQLFASLQRMQQSLIGTVGAVRDGA WYGIBRALLTPLAKJIAHIBBIAGGNLAATLTIDGBSENGDLAQSYSHMQRSLLDTVTVTVRBGS SLWW TBRKMIVQPLAIJGSHPDSIAAGNLARTLTVSGRNGIGELAGSYSHMQRSLIDTVTVVRBGS SLWW TBRKMIVQPLAIJGSHPDSIAAGNLARTIAVYGBNEJTATFASLKTMQQALRGTVSDVRKGS	NAIYSG (280) ADRSIDG (282) ESILIG (274) DAIYAG (278) DAIYAG (278) SOAIYSG (278) GQEMHIG (276)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	DOO A SE JATG NND LSS RTE QQAA SLEETAAS MEQLTATVKQ NAENARQASHLALSASETAQ RGG KVV DN ASESAAG NND LSS RTE QQAA ALEETAAS MEQLTATVKQ NAENARQAS QLALSASETAQ KGG KVV DN LQEJAEG NND LSS RTE QQAA ALEETAAS MEQLTATVKQ NADNARQAS QLAR DASSTAA KRTLAD TREJAAG NTD LSS RTE QQAS ALEETAAS MEQLTATVKQ NADNARQAS QLAQ SASDTAA KRTLAD TREJAAG NTD LSS RTE QQAS ALEETAAS MEQLTATVKQ NADNARQAS QLAQ SASDTAQ KG KVV DG TREJAAG NTD LSS RTE QQAS ALEETAAS MEQLTATVKQ NADNARQAS QLAQ SASDTAQ KG KVV DG TREJAAG NTD LSS RTE QQAS ALEETAAS MEQLTATVKQ NADNARQAS QLAQ SASDTAQ KG VV DG I AEJVAG NND LSS RTE QQA A SLA QTAAS MEQLTATVGQ NADNARQAS E TAK NAA TTAQ GG G VQ VS T	VVETMH (354) VVTTMH (345) VVKTMH (350)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	A DIS TS SQRIA DIISVIDGIA FQTNILALNA A VEAARAGEQGRGFA VVAGEVRNLAQRSAQAAREIK DIASS SQRIA DIITGVIDVIA FQTNILALNA A VEAARAGEQGRAFA VVAGEVRNLAQRSAQAAREIK DIASS SQRIA DIITGVIDVIA FQTNILALNA A VEAARAGEQGRGFA VVAGEVRNLASRSAQAAREIK EIADS SKRIA DIISVIDGIA FQTNILALNA A VEAARAGEQGRGFA VVAGEVRNLASRSAQAAREIK EIIADS SKRIA DIISVIDGIA FQTNILALNA A VEAARAGEQGRGFA VVAGEVRNLASRSAQAAREIK EIIADS SKRIA DIISVIDGIA FQTNILALNA A VEAARAGEQGRGFA VVAGEVRNLASRSAQAAREIK EIIADS SKRIA DIISVIDGIA FQTNILALNA VEAARAGEQGRGFA VVAGEVRNLASRSAQAAREIK	L I E D S         (426)           L I D E S         (417)           A L I E D S         (422)           A L I E D S         (422)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap	VG K VD VG STL VESAGET MA EI VSAVTR VT DI MGEIASAS DE QSRG IDQ VG LAVAEM DR VT QQ NA AL VN RVD MG SYL VESAGDT MG DI VNAVTR VT DI MGEIASAS DE QSRG IDQ VG LAVAEM DR VT QQ NA AL VSRVD MG SYL VENSGAT MQ DI VNAVTR VT DI MGEIASAS DE QSRG IDQ VG QAVREM DR VT QQ NA AL VSRVD TG SYL VENSGAT MQ DI VRSVTR VT DI MGEIASAS DE QSRG IDQ VALAVSEM DR VT QQ NA AL VSRVD TG SYL VESAGET MNNI VNAVTR VT DI MGEIASAS DE QSRG IDQ VALAVSEM DR VT QQ NA AL VSRVD TG SYL VESAGET MNNI VNAVTR VT DI MGEIASAS DE QSRG IDQ VALAVSEM DR VT QQ NA AL VSRVD TG SYL VESAGET MNNI VNAVTR VADI MGEIASAS DE QSRG IDQ VALAVSEM DR VT QQ NA SL VNRVQ QG SKL VNNAATMI DI VNSVTR VT DI MGEIASAS DE QSRG IDQ VALAVSEM DR VT QQ NA SL	* (496) LVEESAS (498) LVEESAS (498) LVVESAS (498) LVQESAS (494) LVQESAS (494) LVEE AS (491)
Tsr Tse Tas Tar <sub>e</sub> Tar <sub>s</sub> Tap Fi	* * * O * * AAAA LEEQAIS BLITE AVA VFRI QQQQ RETSAVVKTVT PAAP. AAA LEEQAIS BLITOS VAVFRIKSEGQEEYKAPVS NKTAPAAIAT HKKTSAS DYQD NWETF. AAAA LEEQAITILADAVAVFRIADD. AAAA LEEQAIS BLITOAVSAFRIJAS PLT NKPQTPSRPA SEQPPAOPRIRIAE ODPNWETF. AAA LEEQAIS BLITOAVSAFRIJAS PII V NKPMRISVNAQSGNTP OS L AAB DDANWETF. VATEQLANQAD BLISS RVAVFTIE EHEVARHESVQITN CASGILK.	(536) (557) (510) (553) (551) (535)

FIG. 6. Comparison of the predicted amino acid sequences of signal transducer proteins Tsr (*E. coli*), Tse (*E. aerogenes*), Tas (*E. aerogenes*);  $Tar_E$  (*E. coli*),  $Tar_S$  (*S. typhimurium*), and Tap (*E. coli*). The sequences are aligned to give the maximal number of residue identities among the six transducers, using the corrected sequence of Tsr (12, 25, 27). This alignment introduces some gaps, most of which can reasonably be explained by the deletion or insertion of codons in particular lineages. The alignment for a sequence (covering residues 194 through 212 of Tsr) that includes most of the second membrane-spanning region contains the most gaps and has the lowest qualitative confidence level. Residues identical in all six transducers, or in all except Tap, are boxed. Symbols: \*, positions at which residues are identical in Tse and Tse but different in the other transducers. For all other positions, residues that are the same in two or more transducers are printed boldface and underlined. The residue count for each transducer is indicated within parentheses at the end of each line.

enteric bacteria. Most of this variation in length occurs at the extreme C termini of the proteins, which are among the most highly variable segments (Fig. 6). It is striking that the C termini of Tse and Tar are of about the same length and that the last five residues are identical. These features may be correlated with the location of *tse* and *tar* at the 5' end of the *meche* operon. Tsr, encoded by a gene located elsewhere on the chromosome, has a very different C terminus, although the Tsr protein is otherwise homologous with Tse.

Within the enteric bacteria there appear to be at least five evolutionary lines among the transducers. Trg, which looks least like the others, may represent the fusion of the N terminus of a membrane transport protein with a C-terminal domain common to all transducers (5). The remaining four groups (Tsr-Tse, Tar, Tas, and Tap) are similiar enough that they probably arose by mutational divergence of intact genes. Their progenitor may have been an amino acid receptor of rather low specificity. Crucial features of the binding site may be conserved in the three Arg residues and the sequences at positions 138 through 140 and 154 through 157. Subsequent gene duplication would have allowed individual transducers to evolve a higher affinity for certain substrates. The interaction with substrate-binding proteins (maltose-binding protein for Tar, dipeptide-binding protein for Tap) could have elaborated later.

*E. coli*, *S. typhimurium*, and *E. aerogenes* all have one or two transducer genes in their *meche* operons, but the genes occur in three different combinations. It may be that all transducers other than Trg evolved at this locus and that some of the genes were then deleted or moved to new sites by chromosome rearrangements. Alternatively, duplicated genes may have been redistributed on the chromosome

 
 TABLE 2. Sequence identities in the periplasmic domain of chemotactic signal transducers<sup>a</sup>

Amino acid	No. (%) of identical residues in:					
	Tse	Tas	Tar <sub>E</sub>	Tars	Тар	
Tsr	100 (60)	43 (26)	53 (32)	51 (31)	32 (19)	
Tse		50 (30)	52 (31)	50 (30)	27 (16)	
Tas			35 (21)	33 (20)	29 (17)	
Tar <sub>E</sub>			. ,	113 (68)	35 (21)	
Tars					35 (21)	

" The amino acid sequences for the six transducers, aligned as in Fig. 6, were compared for the region encompassing residues 31 to 193 of Tsr. The region contains 167 residue positions, although no single transducer has an amino acid at each position. The Table presents the number of residue identities over these 167 positions in pairwise combinations of transducers. Since the deletion or insertion of a codon can occur during gene evolution, the absence of an amino acid at the same position in two or more transducers was also scored as an identity. The percent identity is given in parentheses; it would be altered only slightly if shared gaps were not counted as identities.

before their evolutionary divergence. Genes evolved elsewhere could then have reinvaded the *meche* operon by recombining with the resident transducer genes. The study of transducer evolution will be enhanced by information from a wide taxonomic spectrum of bacteria.

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