# Localization of components of the chemotaxis machinery of *Escherichia coli* using fluorescent protein fusions

#### Victor Sourjik and Howard C. Berg\*

Department of Molecular and Cellular Biology, The Biological Laboratories, Harvard University, 16 Divinity Avenue, Cambridge, MA 02138, USA.

#### Summary

We prepared fusions of yellow fluorescent protein [the YFP variant of green fluorescent protein (GFP)] with the cytoplasmic chemotaxis proteins CheY, CheZ and CheA and the flagellar motor protein FliM, and studied their localization in wild-type and mutant cells of Escherichia coli. All but the CheA fusions were functional. The cytoplasmic proteins CheY, CheZ and CheA tended to cluster at the cell poles in a manner similar to that observed earlier for methylaccepting chemotaxis proteins (MCPs), but only if MCPs were present. Co-localization of CheY and CheZ with MCPs was CheA dependent, and colocalization of CheA with MCPs was CheW dependent, as expected. Co-localization with MCPs was confirmed by immunofluorescence using an anti-MCP primary antibody. The motor protein FliM appeared as discrete spots on the sides of the cell. These were seen in wild-type cells and in a fliN mutant, but not in flhC or fliG mutants. Co-localization with flagellar structures was confirmed by immunofluorescence using an antihook primary antibody. Surprisingly, we did not observe co-localization of CheY with motors, even under conditions in which cells tumbled.

## Introduction

Bacterial cells have a signal transduction pathway that allows them to sense and respond to changes in the concentrations of chemical attractants or repellents. In *Escherichia coli*, the addition of attractant or the removal of repellent promotes counterclockwise flagellar rotation or smooth swimming, which carries cells in a favourable direction. The signal transduction pathway includes a number of membrane-bound receptors, including four

Accepted 31 May, 2000. \*For correspondence. E-mail hberg@biosun.harvard.edu; Tel. (+1) 617 495 0924; Fax (+1) 617 496 1114.

methyl-accepting chemotaxis proteins (MCPs: Tsr, Tar, Trg and Tap) and an oxygen receptor (Aer), six cytoplasmic chemotaxis proteins (CheA, CheW, CheR, CheB, CheY and CheZ) and three proteins comprising a switch complex at the cytoplasmic face of the flagellar motor (FliM, FliN and FliG). For reviews and recent structural work, see Bilwes et al. (1999), Djordjevic and Stock (1998), Falke et al. (1997), Kim et al. (1999) and Levit et al. (1998). The chemotactic signal is transmitted by autophosphorylation at His-48 of CheA, phosphoryl group transfer to Asp-57 of CheY (Hess et al., 1988a,b; Borkovich et al., 1989; Sanders et al., 1989) and binding of phosphorylated CheY to FliM (Welch et al., 1993), which stabilizes the clockwise and destabilizes the counterclockwise rotational states (Kuo and Koshland, 1989; Alon et al., 1998; Scharf et al., 1998). Dephosphorylation of CheY is accelerated by CheZ (Hess et al., 1988a).

The assembly of a ternary complex comprising MCP. CheW and CheA was shown to be necessary for CheA activation in vitro (Ninfa et al., 1991; Gegner et al., 1992), where it forms higher order structures (Liu et al., 1997). The formation of a quaternary complex, including CheY, has also been shown in vitro (Schuster et al., 1993). In addition, CheZ was found to interact with the short form of the histidine kinase, CheAs (Wang and Matsumura, 1996; 1997). Immunoelectron microscopy revealed that MCP-CheW-CheA complexes are clustered in vivo, predominantly at the cell poles (Maddock and Shapiro, 1993), but weaker lateral clusters also are observed (Lybarger and Maddock, 1999; Skidmore et al., 2000). CheA and CheW are required for strong clustering, but there is a significant level of CheA-independent clustering (Skidmore et al., 2000). Receptor clustering might be important for the generation of chemotactic signals (Bray et al., 1998; Levit et al., 1998; Duke and Bray, 1999).

The switch complex has been visualized by electron microscopy (Khan *et al.*, 1992; Francis *et al.*, 1994), in which it appears as a cytoplasmic ring (the C-ring) about 45 nm in diameter. The primary components of this ring are FliM and FliN. The remaining component of the switch complex, FliG, appears at the periphery of the MS-ring, where it forms a bridge to the C-ring. FliG is thought to be assembled first on the MS-ring, followed by the cooperative association of FliM and FliN (Zhao *et al.*, 1995;

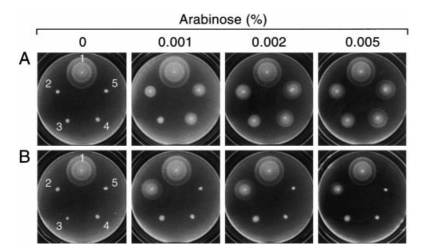
Table 1. Functionality of YFP fusions.

Fusion protein	Functionality <sup>a</sup>	Presence of full-length product <sup>b</sup>
CheY-YFP YFP-CheY CheZ-YFP YFP-CheZ YFP-CheA <sub>L</sub> YFP-CheA <sub>S</sub> FliM-YFP YFP-FliM	+ + + + + + + + + + + + + + - ND + + + + +	+ + + + + +

a. Defined by complementation for spreading in soft agar (Fig. 1) as the ratio of the size of the outer swarm ring of the complementation strain to the size of the outer swarm ring of the wild-type strain, as follows: -, < 0.15;  $\pm$ , 0.15-0.3; + +, 0.4-0.5; + + +, 0.5-0.66; + + + + + . > 0.66.

1996; Kubori et al., 1997; Khan et al., 1998): preparations of flagellar basal bodies that lack FliN also usually lack FliM, and vice versa. However, physical interactions among the three switch proteins, demonstrated both in vitro (Tang et al., 1996) and in vivo (Marykwas and Berg, 1996; Marykwas et al., 1996), suggest that FliM can interact with FliG by itself. Another contradiction is the observation by Oosawa et al. (1994), also made in vitro, that FliM can bind to the MS-ring in the absence of FliG.

Green fluorescent protein (GFP) and its mutants are now widely used to study the localization of proteins in living cells. For general reviews, see Conn (1999); for applications in bacteria, see Margolin (2000). A GFP fusion has already been used to study the localization of one of the chemotaxis proteins, CheZ (M. Manson, personal communication). This construct localizes to the cell poles to what appears to be MCP clusters, in apparent contradiction to the previous result that the interaction of CheZ with CheAs is blocked by the binding of CheAs to CheW (Wang and Matsumura, 1997).



In the present work, we used fusions to YFP (the S65G, V68L, S72A, T203Y GFP mutant) to study the localization in E. coli of the chemotaxis proteins CheY, CheZ and CheA, and the motor protein FliM. The chemotaxis proteins localized in a pattern of clusters in parallel with MCPs, as identified by immunofluorescence. This suggests the existence of a chemotaxis signal complex comprising five proteins: MCPs, CheW, CheA, CheY and CheZ. FliM-YFP localized to the flagellar motors, even in the absence of FliN, but not in the absence of FliG.

#### Results

#### YFP fusions to chemotaxis and motor proteins

In order to investigate the localization of chemotaxis and motor proteins in E. coli cells, we designed several fusions of YFP to the proteins CheY, CheZ, CheA and FliM, as summarized in Table 1. For CheY, CheZ and FliM, both C-terminal and N-terminal fusions were constructed. For CheA, two N-terminal fusions, YFP-CheA<sub>L</sub> and YFP-CheA<sub>S</sub>, were constructed, corresponding to the long and short forms of the CheA protein (Kofoid and Parkinson, 1991). For all fusions, a short amino acid linker (3× or 5× glycine) was used (see Experimental procedures). The fusion genes were expressed under control of the arabinose promoter (pBAD), which allows tight regulation of expression (Guzman et al., 1995). The existence of full-length fusion proteins was verified by immunoblot (see Experimental procedures). YFP fusions to CheY, CheZ, CheA and FliM were tested for their ability to complement corresponding null mutants for spreading in soft agar (Fig. 1; Table 1). CheY, CheZ and FliM fusions complemented the null mutants in an inducerdependent manner, suggesting that all these fusions are functional (although not as efficient as the wild-type proteins). However, the YFP-CheAL fusion did not complement the cheA null mutant. The induction level

Fig. 1. Swarm plates illustrating the ability of different YFP fusion proteins to complement the corresponding null mutants. The concentration of arabinose (as a percentage), the inducer of gene expression for the pBAD constructs, is shown at the top. A. Swarm ring formation by (1) RP437/ pBAD18K wild type; (2) cheY/CheY-YFP; (3) cheY/YFP-CheY; (4) cheZ/CheZ-YFP; and (5) cheZ/YFP-CheZ. B. Swarm ring formation by (1) RP437/ pBAD18K wild type; (2) fliM/FliM-YFP; (3) fliM/YFP-FliM; (4) cheA/YFP-CheAL; and (5) cheA/YFP-CheAs. Swarm plates were supplemented with kanamycin (50 μg ml<sup>-1</sup>).

© 2000 Blackwell Science Ltd, Molecular Microbiology, 37, 740-751

b. Tested by immunoblot.

c. The YFP-FliM fusion complemented the fliM mutant for swimming in liquid media but only poorly for spreading in soft agar.

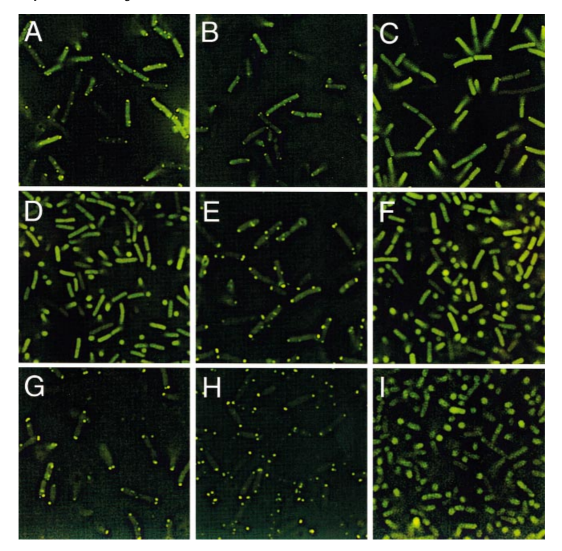


Fig. 2. Localization of YFP fusions to chemotaxis proteins in different mutant backgrounds.

- A. CheY-YFP/cheY.
- B. CheY-YFP/trg.
- C. CheY-YFP/tsr.
- D. CheY-YFP/cheA.
- E. CheZ-YFP/cheZ.
- F. CheZ-YFP/cheA.
- G. YFP-CheA<sub>L</sub>/cheA.
- H. YFP-CheAs/cheA.
- I. YFP-CheAs/tar tsr tap trg.

The arabinose concentration was 0.005%. Results for a larger set of mutant backgrounds are given in Table 2.

that gave an optimal (or nearly optimal) complementation, 0.002% arabinose for FliM-YFP and 0.005% arabinose for all the other proteins, was used for further experiments. Under these conditions, the levels of expression of the fusion proteins were close to the levels of expression of the native proteins ( $\approx$ 70-180%), and the total amount of degraded protein was small, always less than 10% of the amount of native protein (see Experimental procedures). Thus, it seems unlikely that degradation products could account for any of the observed complementation effects.

## Association of YFP fusions with MCP clusters

Both CheY-YFP and YFP-CheY showed a similar pattern of localization. Two types of clusters were observed, intense polar clusters and weaker lateral clusters. Usually, in cells grown to late exponential phase, either one or two clusters were seen at the poles and none or one along the sides of a cell (Fig. 2A). This strongly resembles the clustering of MCPs observed previously by immunoelectron and immunofluorescent microscopy (Maddock and Shapiro, 1993).

Table 2. Localization of YFP fusions

Fusion protein	Background														
	Wild type	cheY	cheZ	tar	tsr	trg	tar tsr tap trg	cheW	cheA	cheA H48Q	fliM flgM	fliM	fliM fliG	fliM fliN	flhC
CheY-YFP	1.40	1.61	1.42	1.19	1.18	1.31	NL	NL	NL	1.45	1.59	NL	ND	ND	NL
YFP-CheY	1.50	1.57	1.49	1.25	1.19	1.36	NL	NL	NL	1.45	1.60	NL	ND	ND	NL
CheZ-YFP	1.45	1.40	1.90	1.23	1.19	1.44	NL	NL	NL	ND	ND	ND	ND	ND	NL
YFP-CheZ	1.57	1.49	1.83	1.30	1.31	1.55	NL	NL	NL	ND	ND	ND	ND	ND	NL
YFP-CheA <sub>I</sub>	1.50	ND	ND	1.32	1.31	1.50	NL	NL	1.88	ND	ND	ND	ND	ND	NL
YFP-CheAs	1.74	ND	ND	1.46	1.52	1.69	NL	NL	2.24	ND	ND	ND	ND	ND	NL
FliM-YFP	1.48	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	1.58	NI	1.30	NI
YFP-FliM	1.42	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	1.55	NI	1.32	NI

The degree of localization was measured as the ratio of the integrated intensity of fluorescence of the localized protein to the integrated intensity of fluorescence of a region of the same size elsewhere in the same cell. More than 40 cells were measured for each fusion construct/strain combination. The standard errors of the measurements were ≤ 0.05. NL, non-localized; NI, non-localized with inclusion bodies in some cells; ND, not determined. See text for further comments.

To verify that CheY-YFP localizes with the MCPs, we analysed the distribution and intensity of CheY-YFP clusters in different mcp, che and fli backgrounds (Table 2; Fig. 2A-D). CheY-YFP clusters were seen in the wild type, cheY, cheZ, tar, tsr and trg backgrounds, but not in a mutant defective in cheA, a mutant defective in cheW or a mutant defective in all four MCPs (tar tsr tap trg). This is consistent with CheY localization on MCP clusters, mediated by CheW and CheA. In tar and tsr strains, which lack either one or other of the major MCP proteins, the intensity of CheY localization at the poles was significantly reduced, and lateral clusters nearly disappeared (Table 2; Fig. 2C). In a trg strain lacking a minor MCP protein, CheY localization was similar to that in the wild type (Table 2; Fig. 2B). Phosphorylation of CheY-YFP by CheA was not required for localization, as shown by the presence of clusters in the cheAH48Q mutant (Table 2). There was no CheY-YFP localization in fliM or cheY fliM mutants, but further investigation showed that this was a result of the negative effect of the fliM deletion on the expression of chemotaxis and receptor genes (Kutsukake and lino, 1994). In the fliM flgM background, where the gene for the anti-sigma factor FlgM responsible for the negative control was deleted, normal localization of CheY-YFP was restored (Table 2).

To determine further whether we might see the localization of CheY-YFP on flagellar motors, we phosphorylated CheY-YFP in vivo under conditions that do not allow clustering of MCPs. First, we used acetate, which is known to cause CheA-independent CheY phosphorylation through the formation of acetyl phosphate (Wolfe et al., 1988; Lukat et al., 1992). The addition of acetate to the cheY cheA strain expressing CheY-YFP yielded no CheY-YFP localization (data not shown), although the cells became tumbly compared with the control, suggesting that the CheY-YFP fusion was phosphorylated and could bind to the motor. We also co-expressed CheY-YFP with a constitutively active cytoplasmic fragment of the Tsr receptor (Ames and Parkinson, 1994) in a cheY tar tsr tap trg strain. Again, the cells tumbled, but no CheY-YFP localization was seen on the motor (data not shown).

Two other chemotaxis proteins, CheZ and CheA, were expected to localize with the MCPs. Co-localization of CheZ-GFP and MCP was shown previously in the laboratory of Mike Manson (personal communication). We observed such localization for both YFP-CheZ and CheZ-YFP (Table 2; Fig. 2E). This localization was dependent on the presence of MCP, CheW and CheA, but not on the presence of CheY (Table 2; Fig. 2F). Both YFP-CheA<sub>L</sub> and YFP-CheA<sub>S</sub> localized in a pattern identical to that of CheY-YFP. This localization was dependent on the presence of MCP and CheW (Table 2; Fig. 2G-H). As the plasmid expressing YFP-CheA<sub>L</sub> also expresses unlabelled CheA<sub>S</sub>, one might expect less fluorescence in the clusters for YFP-CheA<sub>L</sub> because of competition between the two. This appears to be the case, as indicated in Table 2.

To verify the co-localization of the CheY, CheZ and CheA fusion proteins with MCP clusters, we performed additional staining using primary antibody raised against the signalling domain of Tsr, in combination with Texas red-coupled secondary antibody (Fig. 3). Cell fixation with methanol and treatment with lysozyme, necessary to make E. coli cells permeable to antibodies, led to some loss of CheY-YFP, as evidenced by a reduction in the overall level of green fluorescence. At the same time, some cells were not permeabilized enough to allow MCP staining. However, in most cells, presumably cells that were partially lysed, the remaining CheY-YFP spots colocalized precisely with the MCP clusters (Fig. 3A-C). We tried other fixatives, e.g. paraformaldehyde, formaldehyde and glutaraldehyde, but CheY-YFP localization was lost. However, there was no loss of the other fusion proteins,

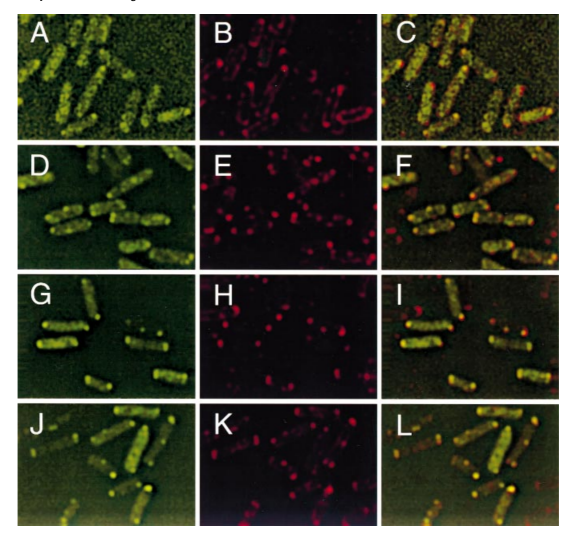


Fig. 3. Association of CheY, CheZ and CheA fusion proteins with MCP clusters, demonstrated by immunofluorescence. Cells expressing YFP fusions were fixed with methanol, stained with anti-Tsr antibody and counterstained with anti-rabbit Texas red-coupled antibody, as described in Experimental procedures.

A-C. cheY cells expressing CheY-YFP.

D-F. cheZ cells expressing CheZ-YFP.

G-I. cheA cells expressing YFP-CheAL.

J-L. cheA cells expressing YFP-CheA<sub>S</sub>.

Left: YFP fluorescence (green); centre: Texas red fluorescence (red); right: superimposition of the images from the left and centre. Orange colour shows co-localization of CheY (C), CheZ (F), CheA<sub>L</sub> (I) and CheA<sub>S</sub> (L) fusion proteins with MCP clusters.

which remained associated with the MCP clusters (Fig. 3D-K).

Results obtained by immunoelectron microscopy suggest that, although CheW and CheA are important for MCP clustering (Maddock and Shapiro, 1993), there is a significant level of clustering in the absence of CheA (Skidmore et al., 2000). To learn whether the mere delocalization of MCPs could account for the absence of localization of the chemotaxis fusion proteins in the *cheW* and cheA backgrounds, we compared MCP localization in wild-type, cheA and cheW strains (Fig. 4A-C). In the absence of CheA or CheW, MCPs were still localized to

the poles (Fig. 4B and C), although not in tight clusters (Fig. 4A). Thus, it is the missing interaction with CheA (for CheY and CheZ) and with CheW (for CheA) that prevents the fusion proteins from being localized to the poles.

## FliM-YFP localization on flagellar motors

In addition to the localization of chemotaxis proteins on MCP clusters, we also observed the localization of FliM on flagellar motors. Although FliM-YFP was much more efficient than YFP-FliM in complementing a fliM mutant for spreading in soft agar (Fig. 1B, swarms 2 and 3), both

© 2000 Blackwell Science Ltd, Molecular Microbiology, 37, 740-751

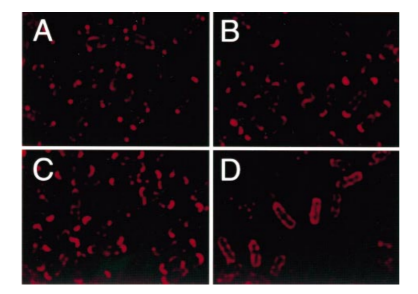


Fig. 4. Localization of MCPs in different mutant backgrounds, demonstrated by immunofluorescence. Wild type (A), cheA (B), cheW (C) and tar tsr tap trg (D) cells were fixed and stained with anti-Tsr antibody, as described in the legend to Fig. 3. In wild-type cells, MCPs are tightly clustered (A). In the cheA or cheW background, they remain localized at the cell poles, but more diffusely (B and C). This localization is not observed in the mcp strain.

constructs complemented the cells for flagellation and motility and showed the same pattern of localization (Table 2; Fig. 5A). This pattern showed several spots (up to 10), more or less evenly distributed along the cell body. As FliM is known to be a part of the cytoplasmic C-ring of the flagellar motor, these spots probably correspond to motors distributed at random on the surface of the cell. In some cells, a bright fluorescent spot was also observed at the end of the cell, presumably corresponding to an inclusion body. The localization of FliM-YFP was dependent on another constituent of the cytoplasmic C-ring, FliG (Fig. 5C), which is known to be assembled on the C-ring before FliM (Zhao et al., 1995). The absence of yet another C-ring component, FliN, did not abolish FliM-YFP localization (Fig. 5B), in agreement with the observation that FliM binds to FliG in the absence of FliN (Marykwas et al., 1996; Tang et al., 1996). The formation

of a single protein aggregate (inclusion body) by FliM-YFP in some fliM cells, in many fliM fliG cells or in a flhC strain suggests that proper targeting to the motor reduces the free concentration of FliM and, hence, the likelihood that it forms inclusion bodies. This is consistent with the finding that, under normal growth conditions, wild-type FliM tends to form aggregates of high molecular weight (Zhao et al., 1996).

The localization of FliM-YFP to flagellar motors was confirmed by immunofluorescence. We used a primary antibody against flagellar hooks and a secondary antibody coupled with Texas red. These experiments were carried out in a fliC strain that lacks flagellar filaments but has hooks (Fig. 6). An additional mutation, clpP, which inactivates a component of one of the major E. coli proteases, was introduced in the fliM fliC background strain (GP90). This reduces the level of FliM-YFP

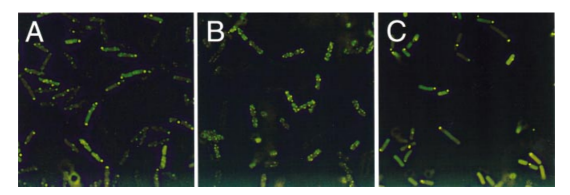


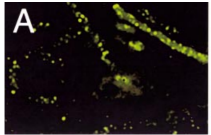
Fig. 5. Localization of FliM-YFP in different mutant backgrounds. A fliM

B. fliM fliN.

C. fliM fliG.

The arabinose concentration was 0.002%.

© 2000 Blackwell Science Ltd, Molecular Microbiology, 37, 740-751





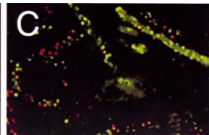


Fig. 6. Association of FliM-YFP with flagellar motors, demonstrated by immunofluorescence. fliM fliC cells expressing FliM-YFP were stained with antihook antibody and counterstained with anti-rabbit Texas red-coupled antibody, as described in Experimental procedures.

A. FliM-YFP (green).

B. Flagellar hooks (red).

C. Superimposition of the images in (A) and (B). All the red spots (hooks) are close to or overlap with the green spots (FliM-YFP). Additional green spots represent immature motors (motors with C-rings but without hooks).

degradation to <2% without interfering with FliM-YFP function. As shown in Fig. 6C, green FliM-YFP spots (Fig. 6A) co-localized with red hook spots (Fig. 6B). However, not all of the green FliM-YFP spots had a corresponding red hook spot. This is expected, as many motors that have yet to synthesize hooks contain FliM (Aizawa, 1996). However, the brightest FliM-YFP foci were not associated with the hooks. Currently, we do not have an explanation for this fact. We do not know why incomplete motors should be brighter. And it seems unlikely that the brighter spots are inclusion bodies, because there is generally only one inclusion body formed per cell in the fliG background (Fig. 5C), even when the cells are made filamentous by growth in the presence of cephalexin (data not shown).

## **Discussion**

Two protein complexes are of particular importance in bacterial chemotaxis: the receptor complex, consisting of MCPs and associated proteins; and the switch complex, assembled on the cytoplasmic face of the flagellar motor. The clustering of receptor complexes adds an additional level of complexity and could be important for signal amplification (Bray et al., 1998; Levit et al., 1998; Duke and Bray, 1999). We used YFP fusions to study the localization of a number of the components of these complexes. Except for CheA, the fusions were functional and restored chemotaxis in the corresponding null mutant background. The present results (together with the work of M. Manson, in preparation) provide direct visualization of receptor and switch complexes in living cells.

Our data imply the existence of a stable complex in which CheW, CheA, CheY and CheZ co-localize with MCP. We did not make a CheW fusion, but CheW was required for association of the other components, in agreement with previous studies (Ninfa et al., 1991; Gegner et al., 1992; Maddock and Shapiro, 1993). The deletion of either of the genes that encodes a major MCP,

tar or tsr, decreases the degree of localization of CheA, CheY or CheZ, but it does not abolish it. This is consistent with the suggestion that different MCP proteins are intermixed in the same clusters (Stock and Levit, 2000). None of the fusion proteins formed clusters in strains that were deleted for all of the MCPs or for CheW; therefore, their localization is not an artifact resulting from the formation of inclusion bodies. CheW is required for localization of CheA, and CheA is required for localization of both CheY and CheZ; however, CheY and CheZ localize independently. As MCPs localize at the poles in the absence of CheW or CheA, the localization of CheY and CheZ must result from their binding to the MCP-CheW-CheA ternary complex, not to MCP directly.

We were also able to observe assembly of FliM-YFP on the switch complex. If a motor had a hook, as indicated by immunofluorescence, it also had a switch complex, as indicated by fluorescence of FliM-YFP. However, not all motors had hooks. FliM localization was not observed in a flhC strain, where flagellar synthesis does not occur. Nor was it observed in the absence of FliG, in support of a mediatory role for FliG in FliM attachment. FliN was not required for the FliM localization as long as FliG was present, consistent with the biochemical evidence for FliM, FliG binding (Marykwas and Berg, 1996; Marykwas et al., 1996; Tang et al., 1996). A possible explanation for the absence of FliM in basal body preparations from fliN mutants (Zhao et al., 1995) is that FliN stabilizes the attachment of FliM to FliG, which is otherwise lost in the purification procedure. We did observe somewhat lower intensity of FliM localization in a fliN background.

However, we were not able to observe localization of CheY fusion proteins to the switch complex, even under conditions in which cells tumbled. This might be a signalto-background problem, as there are many more CheY molecules in the cell than there are FliM binding sites. On the other hand, there are a relatively large number of components with which to build CheW-CheA-MCP

complexes and a relatively small number of molecules of FliM (see Bray et al., 1993).

An obvious advantage in the use of YFP (or other GFP) fusions is that the architecture of receptor and motor complexes can be probed in living cells. Our results complement those obtained by immunofluorescence and protein purification.

#### **Experimental procedures**

## Strains and plasmids

E. coli strains used in this work were derived from strain RP437, a K-12 derivative that is wild type for chemotaxis (Parkinson and Houts, 1982). All strains and plasmids are listed in Table 3. LB (Luria-Bertani) medium or tryptone broth (TB; 1% tryptone, 0.5% NaCl) was used for E. coli cultures. Ampicillin was used at 100 µg ml<sup>-1</sup> and kanamycin

at 50 µg ml<sup>-1</sup>. Chemotaxis tests were performed on TB soft agar plates (1% tryptone, 0.5% NaCl, 0.3% Difco agar).

#### DNA methods

E. coli plasmid DNA was purified using the QIAprep Spin Miniprep Kit (Qiagen). Polymerase chain reaction (PCR) was performed in a MiniCycler (MJResearch) using Pwo DNA polymerase (Boehringer Mannheim). PCR primers used in this study are listed in Table 4; they were from Integrated DNA Technologies. Sequencing was carried out at the Harvard MCB sequencing facility.

## Strain construction

An in frame deletion in *cheY* (codons 8–122) was generated in vitro using two PCR steps, as described by Higuchi (1989), with pRL22 as template and the primers listed in Table 4.

Table 3. Bacterial strains and plasmids used in this study.

	Reference or source		
Wild type for chemotaxis	Parkinson and Houts (1982)		
$\Delta$ che $A$	J. S. Parkinson		
$\Delta$ che $Z$	J. S. Parkinson		
cheW	J. S. Parkinson		
$\Delta$ tar	J. S. Parkinson		
$\Delta \mathit{tsr}$	J. S. Parkinson		
<i>trg::</i> Tn10	J. S. Parkinson		
$\Delta$ (flhC $-$ flhA)	J. S. Parkinson		
	J. S. Parkinson		
	Tang and Blair (1995)		
	D. F. Blair		
	D. F. Blair		
	D. F. Blair		
	P. Danese		
•	This work		
$\Delta$ cheY fliM::cam $\Delta$ flgM	This work		
pVS11, pVS30, pVS50, pVS53, pVS56, pVS59	Guzman <i>et al</i> . (1995)		
Expression vector, P <sub>BAD</sub> promoter, Ap <sup>R</sup> ; parent of pVS13	Guzman <i>et al.</i> (1995)		
Expression vector, P <sub>BAD</sub> promoter, Cm <sup>R</sup> , pACYC ori; parent of pVS15, pVS17	Guzman et al. (1995)		
YFP expression plasmid, Ap <sup>R</sup>	Clontech		
	Matsumura et al. (1984)		
	Tang and Blair (1995)		
	Hess <i>et al.</i> (1987)		
	Oosawa <i>et al.</i> (1988)		
	This work		
' ' B	This work		
	This work		
the state of the s	This work		
	Ames and Parkinson (1994		
	G. J. Phillips, personal gift		
	This work		
	This work This work		
	ΔcheA ΔcheZ cheW Δtar Δtsr trg::Tn10 Δ(flhC-flhA) Δ(tar-tap) Δtsr trg::Tn10 fliM null strain, fliM::cam fliM null strain, in frame fliM deletion fliM fliN null strain fliM fliG null strain fliM::cam fliC::Tn10 clpP::cam recA::kan ΔcheY, in frame cheY deletion ΔcheY fliM::cam ΔflgM  Expression vector, P <sub>BAD</sub> promoter, Km <sup>R</sup> ; parent of pVS1, pVS5,		

Table 4. Primers used in this study.

Primer	Sequence <sup>a</sup>	Priming site <sup>b</sup>			
Fusions					
CheY-EYFP					
PCHEY	5-'CCGGACAGGAGCTCCGTATTTAAATC-3'	-37→-11 (cheY)			
VIC1	5'-GCTCACCACTCCCCCCCCCCCCAGTTTCTCAAAGAT-3'	384→366 ( <i>cheY</i> compl) <sup>c</sup>			
VIC2	5'-GGCGGAGGAGTGGTGAGCAAGGGCGAGGAG-3'	2→21 ( <i>eyfp</i> )			
VIC3	5'-TCAGTTGGAATTCTAGAGTC-3'	749→729 ( <i>eyfp</i> compl)			
EYFP-CheY		(3)			
VIC20	5'-TCGCCACCGAGCTCAGGAGTGTGAAATGGTGAGCAAGGGCGAGGAG-3'	1→20 ( <i>eyfp</i> )			
VIC11	5'-TCCGCCTCCCCTTGTACAGCTCGTCCATG-3'	716→698 ( <i>eyfp</i> compl)			
VIC10	5'-GGAGGCGGAGCGGAGTGGCGGATAAAGAAC-3'	2→17 (cheY)			
VIC18	5'-GTCAGCAGGTCTAGATTGATGGTTGC-3'	429→405 ( <i>cheY</i> compl)			
FliM-EYFP		` ',			
VIC12	5'-GCTGTAGAGCTCTTTTATTCTGCGATAACGAC-3'	-20→0 ( <i>fliM</i> )			
VIC9	5'-TCCGCCTCCGCCTCCTTTGGGCTGTTCCTCGTT-3'	1001→948 ( <i>fliM</i> compl)			
VIC21	5'-GGAGGCGGAGCGGAGTGGTGAGCAAGGGCGAGGAG-3'	2→21 ( <i>eyfp</i> )			
VIC3	(See above)	,			
EYFP-FliM					
VIC20	(See above)				
VIC11	(See above)				
VIC54	5'-GGAGGCGGAGCGGAGTGGGCGATAGTATTCTTTCTCAAGCTG-3'	2→27 (fliM)			
VIC55	5'-CCGGATTCTAGATGTCACTCATTTGGGCTG-3'	1022→993 (fliMcompl)			
CheZ-EYFP		, , , ,			
VIC61	5'-ATGTTTGAGCTCCAGGGCATGTGAGG-3'	-33→-8 ( <i>cheZ</i> )			
VIC60	5'-TCCGCCTCCGCCTCCAAATCCAAGACTATCCAAC-3'	641→623 ( <i>cheZ</i> compl)			
VIC21	(See above)				
VIC3	(See above)				
EYFP-CheZ					
VIC20	(See above)				
VIC11	(See above)				
VIC58	5'-GGAGGCGGAGGCGGACAACCATCAATCAAACCTGC-3'	7→25 ( <i>cheZ</i> )			
VIC59	5'-TCGCCTTCTAGACCGCCTGATATG-3'	699→675 (cheZcompl)			
EYFP-CheA <sub>L</sub>					
VIC20	(See above)				
VIC11	(See above)				
VIC62	5'-GGAGGCGGAGGCGGAGATATAAGCGATTTTTATCAG-3'	10→30 ( <i>cheA</i> )			
VIC63	5'-GTTACATTCTAGATACCGGTCATATTG-3'	2007→1981 (cheAcompl)			
EYFP-CheA <sub>S</sub>					
VIC20	(See above)				
VIC11	(See above)				
VIC64	5'-GGAGGCGGAGCGGAGTGCAAGAACAGCTCGACGC-3'	292-310 (cheA)			
VIC63	(See above)				
cheY deletion					
VIC14	5'-ATCGGCCT <u>TCTAGA</u> TGTGTTGTTCCATTC-3'	-298→-268 ( <i>cheY</i> )			
VIC15	5'-TCCTCACATGCCCAGTTTAAGTTCTTTATCCGCC-3'	20→2 (cheYcompl)			
VIC16	5'-CTGGGCATGTGAGGATGCG-3'	378→396 (cheY)			
VIC19	5'-ATCTGGCAGAATTCTCGTGTATCTG-3'	759→734 ( <i>cheY</i> compl)			

- a. Introduced restriction sites (Sacl, Xbal or EcoRI) are underlined; sequences encoding glycine linkers are marked in italics.
- **b.** Relative to the transcriptional start site (+1) of the corresponding gene.
- c. Complementary strand.

Outer primers were designed to contain *EcoRI* (VIC19) and *Xbal* (VIC14) restriction sites, and these sites were used to clone the fragment into the temperature-sensitive pAMPTS vector. The resulting construct, pVS20, was transformed in *E. coli* and grown on LB plates with ampicillin at 30°C. Transformants were streaked on LB–ampicillin plates and grown overnight at 42°C, allowing only the growth of cells that integrated the pVS20 construct into the chromosome. From these plates, single colonies were picked, grown for 24 h in LB at 30°C without selection, plated at serial dilutions on LB–ampicillin plates and grown overnight at 30°C. The colonies were then tested for ampicillin resistance and chemotaxis on soft agar plates.

## Construction of YFP fusion proteins

YFP fusions to chemotaxis proteins and FliM were constructed using PCR. The target gene and the  $\it eyfp$  gene (Clontech) were amplified using primers with complementary overhangs, encoding either a  $3\times$  or a  $5\times$  glycine linker (Table 4). The resulting DNA fragments were annealed and amplified in a second round of PCR to form a fragment encoding a fusion protein with either a  $3\times$  Gly or a  $5\times$  Gly linker between the target protein and YFP. Outer primers were designed to contain  $\it Sacl$  and  $\it Xbal$  restriction sites, and these were used to clone fragments into the arabinose-inducible pBAD18K expression vector. The final constructs

were sequenced to ensure no PCR mistakes. Expression of full-length fusion proteins was verified by immunoblot using antibodies against FliM, CheY, CheZ and/or YFP (see below). Immunoblots were quantified using the program NIH IMAGE. The levels of expression of fusion proteins (compared with native proteins) estimated from immunoblots were as follows: CheY-YFP,  $\approx 90\%$ ; YFP-CheY,  $\approx 70\%$ ; CheZ-YFP,  $\approx 110\%$ ; YFP-CheZ,  $\approx 70\%$ ; YFP-CheA<sub>L</sub>,  $\approx 120\%$ ;  $YFP-CheA_S, ~\approx 180\%; ~FliM-YFP, ~\approx 150\%; ~YFP-FliM,$  $\approx$  100%. The total levels of degradation products were: CheY-YFP and YFP-CheY, <3%; FliM-YFP and YFP-FliM, < 10%; CheZ-YFP, < 10%; YFP-CheZ, not detected; YFP-CheA<sub>L</sub> and YFP-CheA<sub>S</sub>, not detected. FliM-YFP degradation was further reduced to <2% by the introduction of a clpP mutation (ClpP is a component of the ClpAP protease) in the background strain (GP90) without loss of motor function. Functionality of fusion proteins was tested by the restoration of chemotaxis on soft agar plates supplemented with kanamycin (50 µg ml<sup>-1</sup>) and arabinose (0.001-0.005%).

#### **Immunoblots**

Immunoblots were performed as described previously with minor modifications (Scharf et al., 1998). Motile cells expressing fusion proteins were grown as described below. Whole-cell extracts were prepared from 10 ml samples. Cells were washed once with PBS, resuspended in 300 µl of PBS and lysed by sonication. SDS-PAGE loading buffer ( $3\times$ ) was added, samples were boiled at 95°C for 5 min, and 5 µl of each sample was loaded onto an SDS gradient (8-15%) polyacrylamide gel. After separation by electrophoresis, the proteins were electroblotted to a Hybond ECL nitrocellulose membrane using a tank blot device (Bio Labs, Harvard University) for 3 h at 60 V in transfer buffer (25 mM Tris, 192 mM glycine, 0.05% SDS, 20% methanol, pH 8.3). Blots were blocked overnight at room temperature in TBS-T [20 mM Tris-HCl, 140 mM NaCl, 0.1% (v/v) Tween 20, pH 7.6] with 5% blocking reagent (instant non-fat dry milk) on a rocking platform. Blots were then incubated with primary antibodies in TBS-T for 2 h at room temperature. Monoclonal anti-GFP antibodies (Clontech) were used at a 1:1500 dilution, monoclonal anti-CheY and anti-CheZ antibodies (Scharf et al., 1998) at a 1:1000 dilution and polyclonal anti-FliM antibodies (a gift from David Blair) at a 1:1000 dilution. Blots were washed three times (15 min each) with TBS-T, incubated for 2 h with sheep anti-mouse (or anti-rabbit) horseradish peroxidase-linked secondary antibodies (Amersham) diluted 1:2500 in TBS-T, washed again and detected using an ECL kit (Amersham), as described by the manufacturer.

#### Growth conditions and fluorescence microscopy

Motile cell cultures were grown in tryptone broth (TB) with kanamycin (50 μg ml<sup>-1</sup>) at 33°C. To obtain motile cells expressing YFP fusions for fluorescence measurements, overnight cultures were diluted 1:100 in TB containing arabinose (0.002-0.005%, as indicated in the text) and allowed to grow for 4 h in a rotary shaker. A cell suspension

(100 µl) was applied to a polylysine-coated coverslip, incubated for 5 min and washed three times with tethering buffer (Block et al., 1983) before microscopy.

For double staining with FliM-YFP and polyclonal antihook antibody (Ishihara et al., 1983), cells from a 1 ml culture were resuspended in 100 µl of tethering buffer and incubated with antibody at a 1:500 dilution for 20 min. Cells were washed three times with tethering buffer and incubated in 100 µl of tethering buffer with secondary goat anti-rabbit Texas redconjugated antibodies (1:300 dilution; Molecular Probes) for another 20 min. After incubation, cells were washed with tethering buffer, applied to a polylysine-coated coverslip, washed with tethering buffer once more and imaged.

For double staining with CheY-YFP and anti-Tsr antibody, cells from a 1 ml culture were fixed using methanol as described previously (Teleman et al., 1998). Fixed cells were placed on a polylysine-coated coverslip, allowed to dry fully and treated with lysozyme (2 mg ml<sup>-1</sup>) in GTE buffer (50 mM glucose, 25 mM Tris, 1 mM EDTA, pH 7.5) for 10 min. Coverslips were incubated with a blocking solution (2% BSA in PBS, pH 7.5) overnight at 4°C, incubated in 2% BSA-PBS with anti-Tsr antibodies (1:500 dilution; a gift from Sandy Parkinson) for 2 h at room temperature, washed 10 times with PBS, incubated with secondary goat anti-rabbit Texas red antibody (1:300 dilution in 2% BSA-PBS) for 2 h at room temperature, washed with PBS again and imaged.

Fluorescent microscopy was performed using a Delta Vision deconvolution microscope and program package (Applied Precision). YFP images were taken using a bandpass excitation filter (480-500 nm) and either a longpass emission filter (510 nm) or, if cells were also stained with Texas red, a bandpass emission filter (509-547 nm). Texas red images were taken using a bandpass excitation filter (541-569 nm) and a bandpass emission filter (581-654 nm). When necessary, images were quantified using the program NIH IMAGE. Deconvoluted images were prepared for final publication using Adobe Photoshop 5.5 and a Tektronix Phaser 450 printer.

# Acknowledgements

We thank Paul Danese for inspiration and help during the early phases of this study. This work was supported by grant Al16478 from the National Institute of Allergy and Infectious Diseases and by the Rowland Institute for Science.

### References

Aizawa, S.-I. (1996) Flagellar assembly in Salmonella typhimurium. Mol Microbiol 19: 1-5.

Alon, U., Camarena, L., Surette, M.G., Aguera y Arcas, B., Liu, Y., Leibler, S., et al. (1998) Response regulator output in bacterial chemotaxis. EMBO J 17: 4238-4248.

Ames, P., and Parkinson, J.S. (1994) Constitutively signalling fragments of Tsr, the Escherichia coli serine chemoreceptor. J Bacteriol 176: 6340-6348.

Bilwes, A.M., Alex, L.A., Crane, B.R., and Simon, M.I. (1999) Structure of CheA, a signal-transducing histidine kinase. Cell **96**: 131–141.

© 2000 Blackwell Science Ltd, Molecular Microbiology, 37, 740-751

- Block, S.M., Segall, J.E., and Berg, H.C. (1983) Adaptation kinetics in bacterial chemotaxis. J Bacteriol 154: 312-323.
- Borkovich, K.A., Kaplan, N., Hess, J.F., and Simon, M.I. (1989) Transmembrane signal transduction in bacterial chemotaxis involves ligand-dependent activation of phosphate group transfer. Proc Natl Acad Sci USA 86: 1208-
- Bray, D., Bourret, R.B., and Simon, M.I. (1993) Computer simulation of the phosphorylation cascade controlling bacterial chemotaxis. Mol Biol Cell 4: 469-482.
- Bray, D., Levin, M.D., and Morton-Firth, C.J. (1998) Receptor clustering as a cellular mechanism to control sensitivity. Nature 393: 85-88.
- Conn, P.M. (ed.) (1999) Green fluorescent protein. Methods Enzymol 302.
- Djordjevic, S., and Stock, A.M. (1998) Structural analysis of bacterial chemotaxis proteins: components of a dynamic signaling system. J Struct Biol 124: 189-200.
- Duke, T.A.J., and Bray, D. (1999) Heightened sensitivity of a lattice of membrane receptors. Proc Natl Acad Sci USA 96: 10104-10108.
- Falke, J.J., Bass, R.B., Butler, S.L., Chervitz, S.A., and Danielson, M.A. (1997) The two-component signaling pathway of bacterial chemotaxis: a molecular view of signal transduction by receptors, kinases, and adaptation enzymes. Annu Rev Cell Dev Biol 13: 457-512.
- Francis, N.R., Sosinsky, G.E., Thomas, D., and DeRosier, D.J. (1994) Isolation, characterization and structure of bacterial flagellar motors containing the switch complex. J Mol Biol 235: 1261-1270.
- Gegner, J.A., Graham, D.R., Roth, A.F., and Dahlquist, F.W. (1992) Assembly of an MCP receptor, CheW, and kinase CheA complex in the bacterial chemotaxis signal transduction pathway. Cell 18: 975-982.
- Guzman, L.-M., Belin, D., Carson, M.J., and Beckwith, J. (1995) Tight regulation, modulation, and high-level expression by vectors containing the arabinose P<sub>BAD</sub> promoter. J Bacteriol 177: 4121-4130.
- Hess, F., Oosawa, K., Matsumura, P., and Simon, M.I. (1987) Protein phosphorylation is involved in bacterial chemotaxis. Proc Natl Acad Sci USA 84: 7609-7613.
- Hess, F., Oosawa, K., Kaplan, N., and Simon, M.I. (1988a) Phosphorylation of three proteins in the signaling pathway of bacterial chemotaxis. Cell 53: 79-87.
- Hess, J.F., Bourret, R.B., and Simon, M.I. (1988b) Histidine phosphorylation and phosphoryl group transfer in bacterial chemotaxis. Nature 336: 139-143.
- Higuchi, R. (1989) Using PCR to engineer DNA. In PCR Technology: Principles and Applications for DNA Amplification. Erlich, H.A. (ed.). New York: Stockton, pp. 61-70.
- Ishihara, A., Segall, J.E., Block, S.M., and Berg, H.C. (1983) Coordination of flagella on filamentous cells of Escherichia coli. J Bacteriol 155: 228-237.
- Khan, I.M., Reese, T.S., and Khan, S. (1992) The cytoplasmic component of the bacterial flagellar motor. Proc Natl Acad Sci USA 89: 5956-5960.
- Khan, S., Zhao, R., and Reese, T.S. (1998) Architectural features of the Salmonella typhimurium flagellar motor switch revealed by disrupted C-rings. J Struct Biol 122: 311-319.

- Kim, K.K., Yokota, H., and Kim, S.-H. (1999) Four-helicalbundle structure of the cytoplasmic domain of a serine chemotaxis receptor. Nature 400: 787-792.
- Kofoid, E.C., and Parkinson, J.S. (1991) Tandem translation starts in the cheA locus of Escherichia coli. J Bacteriol 173: 2116-2119.
- Kubori, T., Yamaguchi, S., and Aizawa, S.-I. (1997) Assembly of the switch complex onto the MS ring complex of Salmonella typhimurium does not require any other flagellar proteins. J Bacteriol 179: 813-817.
- Kuo, S.C., and Koshland, D.E. (1989) Multiple kinetic states for the flagellar motor switch. J Bacteriol 171: 6279-6287.
- Kutsukake, K., and lino, T. (1994) Role of the FliA-FlgM regulatory system on the transcriptional control of the flagellar regulon and flagellar formation in Salmonella typhimurium. J Bacteriol 176: 3598-3603.
- Levit, M.N., Liu, Y., and Stock, J.B. (1998) Stimulus response coupling in bacterial chemotaxis: receptor dimers in signalling arrays. Mol Microbiol 30: 459-466.
- Liu, Y., Levit, M., Lurz, R., Surette, M.G., and Stock, J.B. (1997) Receptor-mediated protein kinase activation and the mechanism of transmembrane signaling in bacterial chemotaxis. EMBO J 16: 7231-7240.
- Lukat, G.S., McCleary, W.R., Stock, A.M., and Stock, J.B. (1992) Phosphorylation of bacterial response regulator proteins by low molecular weight phospho-donors. Proc Natl Acad Sci USA 89: 718-722.
- Lybarger, S.R., and Maddock, J.R. (1999) Clustering of the chemoreceptor complex in Escherichia coli is independent of the methyltransferase CheR and the methylesterase CheB. J Bacteriol 181: 5527-5529.
- Maddock, J.R., and Shapiro, L. (1993) Polar location of the chemoreceptor complex in the Escherichia coli cell. Science 259: 1717-1723.
- Margolin, W. (2000) Green fluorescent protein as a reporter for macromolecular localization in bacterial cells. Methods **20**: 62-72.
- Marykwas, D.L., and Berg, H.C. (1996) A mutational analysis of the interaction between FliG and FliM, two components of the flagellar motor of Escherichia coli. J Bacteriol 178: 1289-1294.
- Marykwas, D.L., Schmidt, S.A., and Berg, H.C. (1996) Interacting components of the flagellar motor of Escherichia coli revealed by the two-hybrid system in yeast. J Mol Biol 256: 564-576.
- Matsumura, P., Rydel, J.J., Linzmeier, R., and Vacante, D. (1984) Overexpression and sequence of the Escherichia coli cheY gene and the biochemical activities of the CheY protein. J Bacteriol 160: 36-41.
- Ninfa, E.G., Stock, A., Mowbray, S., and Stock, J. (1991) Reconstitution of the bacterial chemotaxis signal transduction system from purified components. J Biol Chem 266: 9764-9770.
- Oosawa, K., Hess, J.F., and Simon, M.I. (1988) Mutants defective in bacterial chemotaxis show modified protein phosphorylation. Cell 53: 89-96.
- Oosawa, K., Ueno, T., and Aizawa, S.-I. (1994) Overproduction of the bacterial flagellar switch proteins and their interactions with the MS ring complex in vitro. J Bacteriol 155: 265-274.
- Parkinson, J.S., and Houts, S.E. (1982) Isolation and
- © 2000 Blackwell Science Ltd, Molecular Microbiology, 37, 740-751

- behavior of Escherichia coli deletion mutants lacking chemotaxis functions. J Bacteriol 151: 106-113.
- Sanders, D.A., Gillece-Castro, B.L., Stock, A.M., Burlingame, A.L., and Koshland, D.E., Jr (1989) Identification of the site of phosphorylation of the chemotaxis response regulator protein, CheY. J Biol Chem 264: 21770-21778.
- Scharf, B.E., Fahrner, K.A., Turner, L., and Berg, H.C. (1998) Control of direction of flagellar rotation in bacterial chemotaxis. Proc Natl Acad Sci USA 95: 201-206.
- Schuster, S.C., Swanson, R.V., Alex, L.A., Bourret, R.B., and Simon, M.I. (1993) Assembly and function of a quaternary signal transduction complex monitored by surface plasmon resonance. Nature 365: 343-347.
- Skidmore, J.M., Ellefson, D.D., McNamara, B.P., Couto, M.M.P., Wolfe, A.J., and Maddock, J.R. (2000) Polar clustering of the chemoreceptor complex in Escherichia coli occurs in the absence of complete CheA function. J Bacteriol 182: 967-973.
- Stock, J., and Levit, M. (2000) Signal transduction: hair brains in bacterial chemotaxis. Curr Biol 10: R11-R14.
- Tang, H., and Blair, D.F. (1995) Regulated underexpression of the FliM protein of Escherichia coli and evidence for a location in the flagellar motor distinct from the MotA/MotB torque generators. J Bacteriol 177: 3485-3495.
- Tang, H., Braun, T.F., and Blair, D.F. (1996) Motility protein

- complexes in the bacterial flagellar motor. J Mol Biol 261: 209-221.
- Teleman, A.A., Graumann, P.L., Lin, D.C.-H., Grossman, A.D., and Losick, R. (1998) Chromosome arrangement within a bacterium. Curr Biol 8: 1102-1109.
- Wang, H., and Matsumura, P. (1996) Characterization of the CheAs/CheZ complex: a specific interaction resulting in enhanced dephosphorylating activity on CheY-phosphate. Mol Microbiol 19: 695-703.
- Wang, H., and Matsumura, P. (1997) Phosphorylating and dephosphorylating protein complexes in bacterial chemotaxis. J Bacteriol 179: 287-289.
- Welch, M., Oosawa, K., Aizawa, S.-I., and Eisenbach, M. (1993) Phosphorylation-dependent binding of a signal molecule to the flagellar switch of bacteria. Proc Natl Acad Sci USA 90: 8787-8791.
- Wolfe, A.J., Conley, M.P., and Berg, H.C. (1988) Acetyladenylate plays a role in controlling the direction of flagellar rotation. Proc Natl Acad Sci USA 85: 6711-6715.
- Zhao, R., Schuster, S.C., and Khan, S. (1995) Structural effects of mutations in Salmonella typhimurium flagellar switch complex. J Mol Biol 251: 400-412.
- Zhao, R., Amsler, C.D., Matsumura, P., and Khan, S. (1996) FliG and FliM distribution in the Salmonella typhimurium cell and flagellar basal bodies. J Bacteriol 178: 258-265.