Chemotactic Signaling by an *Escherichia coli* CheA Mutant That Lacks the Binding Domain for Phosphoacceptor Partners

Knut Jahreis,† Tom B. Morrison,‡ Andrés Garzón,§ and John S. Parkinson*

Biology Department, University of Utah, Salt Lake City, Utah 84112

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CheA is a multidomain histidine kinase for chemotaxis in Escherichia coli. CheA autophosphorylates through interaction of its N-terminal phosphorylation site domain (P1) with its central dimerization (P3) and ATPbinding (P4) domains. This activity is modulated through the C-terminal P5 domain, which couples CheA to chemoreceptor control. CheA phosphoryl groups are donated to two response regulators, CheB and CheY, to control swimming behavior. The phosphorylated forms of CheB and CheY turn over rapidly, enabling receptor signaling complexes to elicit fast behavioral responses by regulating the production and transmission of phosphoryl groups from CheA. To promote rapid phosphotransfer reactions, CheA contains a phosphoacceptor-binding domain (P2) that serves to increase CheB and CheY concentrations in the vicinity of the adjacent P1 phosphodonor domain. To determine whether the P2 domain is crucial to CheA's signaling specificity, we constructed CheAAP2 deletion mutants and examined their signaling properties in vitro and in vivo. We found that CheAAP2 autophosphorylated and responded to receptor control normally but had reduced rates of phosphotransfer to CheB and CheY. This defect lowered the frequency of tumbling episodes during swimming and impaired chemotactic ability. However, expression of additional P1 domains in the CheA Δ P2 mutant raised tumbling frequency, presumably by buffering the irreversible loss of CheAD2-generated phosphoryl groups from CheB and CheY, and greatly improved its chemotactic ability. These findings suggest that P2 is not crucial for CheA signaling specificity and that the principal determinants that favor appropriate phosphoacceptor partners, or exclude inappropriate ones, most likely reside in the P1 domain.

CheA, a cytoplasmic kinase controlled by transmembrane receptors, plays a central role in the chemotactic signaling pathway of Escherichia coli. CheA autophosphorylates at a histidine residue by using ATP as the phosphoryl donor (11) and then donates its phosphoryl groups to two response regulators, CheB and CheY, thereby activating their signaling functions. Phospho-CheY interacts with the switching apparatus of the flagellar motors to augment clockwise (CW) rotation (5, 42); phospho-CheB regulates a subsequent sensory adaptation process (4, 20). E. coli contains many response regulator proteins whose structures and phosphotransfer chemistries are homologous to those of CheB and CheY (reviewed in references 33, 38, and 43). Although CheA can be made to donate phosphoryl groups to heterologous response regulators in vitro (9, 30, 41), such reactions are relatively inefficient and evidently do not interfere with chemotactic signaling in vivo. This report investigates a structural feature, unique to the CheA subfamily of histidine kinases, that might account for its signaling specificity.

CheA has a modular organization, with different signaling functions allocated to different domains (Fig. 1). The central dimerization (P3) and ATP-binding (P4) domains are flanked by a C-terminal domain (P5) that couples autophosphorylation activity to receptor control (7) and by two N-terminal domains, P1 and P2 (27, 39), that are involved in phosphotransfer signaling. P1 contains the CheA phosphorylation site, His-48, and interacts with the P4 domain during autophosphorylation and with CheB and CheY in subsequent phosphotransfer reactions. The P2 domain binds CheY and CheB with micromolar affinity (18, 39) and increases CheA phosphotransfer rates (36, 37). Binding to P2 does not appear to induce conformational changes in CheB or CheY that appreciably influence their phosphotransfer reactions with the P1 domain, because P2 domains in *trans* do not affect CheA phosphotransfer rates (25, 37).

The P2 domain could conceivably impart specificity to CheA's signaling transactions by increasing the local concentrations of CheB and CheY, but not those of other response regulator proteins, through its binding interactions. To test this "target acquisition" model, we constructed a P2-less mutant of CheA and examined its signaling behavior in vitro and in vivo. We found that the CheA Δ P2 protein had essentially wild-type autophosphorylation activity, which responded normally to chemoreceptor control. However, the mutant protein exhibited defects in phosphotransfer to both CheB and CheY, consistent with an earlier rapid kinetics study that demonstrated a substantial contribution of the P2 domain to the rate of CheA-CheY phosphotransfer (37). CheA Δ P2 mutant cells, presumably owing to reduced phosphotransfer rates, exhibited predominantly counterclockwise (CCW) flagellar rotation and poor chemotactic ability. However, upward adjustment of phospho-CheB and phospho-CheY levels, by buffering the cellular pool of CheA_D2-generated phosphoryl groups with excess P1 domains, dramatically enhanced the chemotaxis proficiency of the CheA Δ P2 mutant. These results demonstrate that the CheA Δ P2 protein, despite lowered phosphotransfer rates,

^{*} Corresponding author. Mailing address: Biology Department, University of Utah, Salt Lake City, UT 84112. Phone: (801) 581-7639. Fax: (801) 581-4668. E-mail: Parkinson@biology.utah.edu.

[†] Present address: Fachbereich Biologie/Chemie, Universität Osnabrück, Osnabrück 49069, Germany.

[‡] Present address: BioTrove, Inc., Woburn, MA 01801-1728.

[§] Present address: Centro Andaluz de Biología del Desarrollo, Universidad Pablo de Olavide/CSIC, E-41013 Seville, Spain.



FIG. 1. CheA domain organization and signaling activities. The relative lengths of the CheA domains (P1 to P5) and linkers (L1 and L2) are proportional to the sizes of their primary structures. The N-terminal P1 domain accepts a phosphoryl group at His-48 via interaction with the ATP-binding P4 domain. The C-terminal P5 domain enables chemoreceptors, assisted by the CheW coupling protein, to control CheA autophosphorylation activity in response to attractant or repellent stimuli. The ensuing motor response and subsequent sensory adaptation are controlled by the levels of phosphorylated CheY (Y~P), which enhances CW flagellar rotation, and phosphorylated CheB (B~P), which hydrolyzes glutamyl-methyl esters in the signaling domains of the chemoreceptors. The phosphorylated forms of these response regulators turn over rapidly through self-catalyzed hydrolysis. The CheZ protein further enhances dephosphorylation of $Y \sim P$ (but not $B \sim P$). CheY and CheB acquire their phosphoryl groups from phospho-CheA. The P2 domain of CheA is not essential for these phosphotransfer reactions but rather enhances their rate by reversibly binding the unphosphorylated forms of CheY and CheB to increase phosphoacceptor concentrations in the vicinity of the P1 phosphodonor.

retains sufficient signaling specificity for chemotactic responses. Thus, although the P2 domain augments the phosphotransfer process through target acquisition, it does not play a critical role in CheA signaling specificity. We conclude that the main specificity determinants that prevent excessive cross talk with heterologous response regulators reside in the P1 domain of CheA.

MATERIALS AND METHODS

Bacterial strains. The bacterial strains used in this work (Table 1) were all derivatives of the *E. coli* K-12 strain RP437, our reference wild-type strain for chemotaxis studies (32).

Plasmids. Plasmids central to this work are listed in Table 1. All are derivatives of pBR322 (6) and confer resistance to ampicillin. All except pEK46 and pTM46 carry *lac1* and an IPTG (isopropyl- β -D-thiogalactopyranoside)-inducible P_{tac} promoter driving transcription of a multiple cloning region containing the ribosomebinding site and translation start from the *cheY* gene of *E. coli* (29).

Media and culture conditions. T broth (10 g of tryptone and 5 g of NaCl per liter) was routinely used for growth of bacterial strains, generally at 35°C. IPTG was purchased from Promega Corp. (Madison, Wis.). Ampicillin was purchased from Sigma Chemical Co. (St. Louis, Mo.) and used at a final concentration of 100 μ g/ml in solid and liquid media.

Behavioral assays. Chemotactic ability was measured on semisolid tryptone agar (T broth plus 2.6 g of agar per liter) (31). For strains harboring plasmids, ampicillin was added to a final concentration of 50 μ g/ml. Flagellar rotation patterns were determined by cell tethering with antiflagellar serum (31).

Construction of pKJ9. Pertinent features of the *cheA* plasmid pKJ9 have been described previously (10). It was constructed by PCR amplification of the *cheA* coding region in RP437 with primers kj81 and kj82 (Table 1). The PCR fragment was blunt ended by treatment with the Klenow fragment of DNA polymerase and inserted in the forward orientation at the Smal site of pBluescript SK(+) (2). The *cheA* coding region was subsequently excised at the flanking PstI (upstream) and BamHI (downstream) sites in the pBluescript vector and inserted at the corresponding sites into the pTM30 vector (Table 1).

Strain, plasmid, or oligonucleotide	Relevant properties	Reference
Strains		
RP437	<i>thr-1 leuB6 his-4 metF159 eda-50</i> (chemotaxis wild type)	32
RP3098	$srlC::Tn10-300 \Delta(flhD-flhB)4$	35
RP9005	$recD$::mini-Tn10-1903 Δ (motB-cheA)m1111	34
RP9535	RP437 $eda^+ \Delta(cheA) 1643$	19
UU1120	RP437 eda^+	This work
UU1121	RP437 $eda^+ \Delta(cheA)150-247\Omega PA1$ (cheA $\Delta P2$)	This work
Plasmids		
pTM30	IPTG-inducible P_{tac} expression vector	27
pEK46	pUC118 motA-motB-cheA-cheW	14
pTM46	pEK46 Δ (cheA)150–247 Ω PA1 (CheA Δ P2)	This work
pKJ9	pTM30 (<i>cheA</i>)	This work
pKJ9-1	pKJ9 (cheA) SacII-150	This work
pKJ9-2	pKJ9 (cheA) SacII-160	This work
pKJ9-1.1	pKJ9-1 Δ (<i>cheA</i>)150–247 Ω PA1 (CheA Δ P2)	This work
pKJ9-1.2	pKJ9-1 Δ (cheA)150–247 Ω PA2	This work
pKJ9-2.1	pKJ9-2 Δ (cheA)160–247 Ω PA1	This work
pKJ9-2.2	pKJ9-2 Δ (cheA)160–247 Ω PA2	This work
pAG3	pKJ9 <i>cheA</i> (1–149)	10
Oligonucleotides		
kj-1	5'-CAGCACCAGCTGCTCCACCGCGGGCTAGCGCC-3' (PA linker [top strand])	This work
kj-2	5'-GGCCGGCGCTAGCCCGCGGTGGAGCAGCTGGTGCTGGC-3' (PA linker [bottom strand])	This work
kj-3	5'-AGTGAACCGCGGGATGAGCAG-3' (SacII site at <i>cheA</i> codon 150)	This work
kj81	5'-GTGAGCATGGATATAAGC-3' (upstream <i>cheA</i> primer)	This work
kj82	5'-TCAGGCGGCGGTGTTCGC-3' (downstream <i>cheA</i> primer)	This work
kj63-3	5'-CGCCGCGGCGAATTATCC-3' (SacII site at <i>cheA</i> codon 160)	This work

TABLE 1. Strains, plasmids, and oligonucleotides

Construction of CheA\DeltaP2 derivatives of pKJ9. The *cheA* coding region was excised from pKJ9 as a PstI-BamHI fragment and inserted into the pALTER-1 phagemid (Promega Corp.). Primers kj-3 and kj63-3 (Table 1) were used to introduce a SacII site at *cheA* codon 150 or 160, according to the supplier's instructions for the Promega in vitro mutagenesis system. The desired *cheA* mutations were confirmed by DNA sequencing and then transferred back into pTM30 as a PstI-BamHI restriction fragment, yielding pKJ9-1 and pKJ9-2 (Table 1). To anneal oligonucleotides kj-1 and kj-2 (Table 1), 6 µg of each were mixed in an Eppendorf tube with annealing buffer (125 mM Tris [pH 7.6], 15 mM MgCl₂, 10 mM dithiothreitol) in a total volume of 20 µl, incubated for 3 min at 95°C, and then allowed to cool to room temperature over the course of an hour. Plasmids pKJ9-1 and pKJ9-2 were digested with SacII and EagI and then ligated with 10 µl of the annealed linker mixture to produce plasmids pKJ9-1.1 and pKJ9-2.1 (Table 1). In the same manner, a second copy of the linker was inserted into these plasmids to create pKJ9-1.2 and pKJ9-2.2 (Table 1).

Construction of UU1121. The *cheA* Δ P2 deletion was transferred from pKJ9-1.1 to pEK46 by restriction fragment replacement, yielding plasmid pTM46. Linearized pTM46 was used to transform RP9005, and the transformation mixture was streaked onto semisolid tryptone plates. Motile transformants migrating away from the streak of nonmotile recipient cells were picked and shown to carry the *cheA* Δ P2 deletion by PCR tests. The *cheA* Δ P2 allele in one of these recombinants was subsequently transferred to RP437 by cotransduction with the *eda* locus (31). Transductants were screened by PCR to identify ones that had inherited the donor *cheA* Δ P2 allele. One strain carrying *cheA* Δ P2 (UU1121) and an isogenic *cheA*⁺ strain (UU1120) were kept for further analysis.

Protein purification. CheA and CheA Δ P2 (13). CheW (3), CheB (12), CheY (21), and CheZ (13) were expressed from plasmids in strain RP3098 and purified by published procedures.

Phosphorylation assays. All reactions were carried out in phosphorylation buffer (50 mM Tris-HCl [pH 7.5], 50 mM KCl, and 5 mM MgCl₂) at room temperature. CheA autophosphorylation assays were performed as described previously (3). ³²P-CheA and ³²P-CheAΔP2 were purified and used in CheB and CheY phosphotransfer assays as described previously (28). Receptor coupling assays were done with membranes containing the serine receptor Tsr as described previously (26). In all assays, 2 μ I-reaction samples were removed at various times and added to 10 μ I of sodium dodecyl sulfate protein sample buffer (15) to stop the reaction. Reaction products were separated by electrophoresis on sodium dodecyl sulfate-containing 16.5% polyacrylamide gels and quantified with a Molecular Dynamics PhosphorImager (28).

RESULTS

Construction of CheA Δ P2. We based our design of a P2-less version of CheA on the structure of FrzE, a CheA homologue of Myxococcus xanthus that lacks a P2 domain (1, 23, 24). In FrzE a proline- and alanine-rich linker, approximately 130 residues long, connects the P1 and P3/P4 domains. In the CheA proteins of E. coli and Salmonella enterica serovar Typhimurium, the P2 domain is flanked by flexible linkers, 20 to 30 residues long, whose sequences are not highly conserved (27) (Fig. 1). Accordingly, we replaced the P2 coding region, and joined the flanking L1 and L2 segments, with a prolineand alanine-rich (PA) linker that was similar to, but shorter than, the linker in FrzE. To vary the length of the connection between the P1 and ATP-binding domains, in case it proved to be critical for CheA autophosphorylation activity, the construction strategy allowed for insertion of multiple, tandem copies of the PA linker.

CheA constructs with deletions of P2 were made in plasmid pKJ9, which carries a wild-type *cheA* gene expressed from an inducible P_{tac} promoter (10). The construction details are summarized in Fig. 2. We first introduced a unique SacII restriction site at either codon 150 or 160 in the L1 coding region (Fig. 2A). The sequence change at codon 160 did not alter the amino acid it specified (arginine), whereas the change at codon 150 did (glutamine to arginine). Both mutant plasmids, pKJ9-1 and pKJ9-2, complemented a $\Delta(cheA)$ host strain (RP9535) as



FIG. 2. Strategy for constructing *cheA* genes with deletions of P2. (A) Features of *cheA* flanking the P2 coding segment. A SacII restriction site was introduced at codon 150 (pKJ9-1) or codon 160 (pKJ9-2) to permit excision of the P2 coding segment by double digestion with SacII and EagI enzymes. (B) The PA linker. The boxed sequence depicts the double-stranded linker made by annealing the complementary single-stranded oligonucleotides kj-1 and kj-2 (Table 1). The PA linker carried a SacII-compatible overhang at one end and an EagI-compatible overhang at the other for joining the P2-excised coding segments. An internal SacII site in the PA linker permitted additional excision-replacement cycles for creating tandem PA linker inserts.

proficiently as did the parent plasmid (data not shown), demonstrating that neither change had any discernible effect on CheA function. The *cheA* coding segment between the introduced SacII sites in L1 and a unique EagI restriction site in L2 was then replaced with the double-stranded oligonucleotide shown in Fig. 2B to introduce the PA linker. The linker oligonucleotide contained an internal SacII site to permit introduction of additional copies of the PA linker in the same manner. In all, we constructed four P2-less CheA plasmids (pKJ9-1.1, pKJ9-1.2, pKJ9-2.1, and pKJ9-2.2) which had either one or two tandem copies of the PA linker beginning at either codon 150 or 160 in the L1 segment (Table 1).

All four mutant plasmids behaved similarly in preliminary tests. Their CheA expression levels were identical to that of pKJ9 over a range of IPTG concentrations (data not shown), indicating that the proteins with deletions of P2 were similar in synthesis and stability to the wild type. Moreover, at IPTG concentrations above 10 µM, both the parental and mutant plasmids caused episodes of CW flagellar rotation in tethered cells of RP9535 [Δ (*cheA*)] (data not shown), indicating that the mutant proteins were capable of autophosphorylation and subsequent phosphotransfer to CheY. However, unlike pKJ9, the mutant plasmids failed to complement RP9535 for chemotaxis at any level of induction (data not shown), suggesting that one or more of the CheA signaling reactions was impaired. To examine the activities of CheA with a deletion of P2 in more detail, we purified and characterized the mutant protein made by plasmid pKJ9-1.1, which carries the $\Delta(cheA)150-247\Omega$ PA1 alteration, in which one copy of the PA linker replaces cheA codons 150 to 247. Hereafter, we refer to this cheA allele as *cheA* Δ P2 and to its gene product as CheA Δ P2.

In vitro assays of CheA function. The CheA reactions involved in phosphorelay signaling are summarized in Fig. 3. CheA catalyzes its own phosphorylation, using ATP, and then serves as a phosphodonor in transfer reactions catalyzed by CheB or CheY. Finally, the phosphoryl groups on CheB and CheY are released as inorganic phosphate. In the case of phospho-CheY, the dephosphorylation step can be accelerated



FIG. 3. The phosphorelay signaling reactions of chemotaxis. Only the net forward rates are depicted: CheA autophosphorylation (which is subject to modulation in signaling complexes with chemoreceptors and CheW), phosphotransfer from phospho-CheA (P~CheA) to CheB and CheY, and the subsequent dephosphorylation of phospho-CheB (P~CheB) and phospho-CheY (P~CheY). Note that dephosphorylation of P~CheY (but not that of P~CheB) is accelerated by CheZ.

by CheZ. In chemoreceptor signaling complexes, CheA autokinase activity responds to the signaling state of the receptor. CW-signaling (e.g., unliganded) receptors stimulate the CheA autophosphorylation rate more than 100-fold, whereas CCWsignaling (e.g., attractant-bound) receptors inhibit CheA to approximately its uncoupled level of activity.

Based on the reaction scheme depicted in Fig. 3, we devised assays to compare the autophosphorylation, receptor control, and phosphotransfer abilities of CheA and CheA Δ P2. Two general considerations influenced our experimental designs. (i) The autophosphorylation and phosphotransfer reactions are readily reversible (10, 40). Accordingly, we chose reactant concentrations and stoichiometries that minimized the reverse reactions to facilitate comparison of their net forward rates, which are the ones depicted in Fig. 3. (ii) At physiological component concentrations, the phosphotransfer and receptorstimulated autophosphorylation reactions are exceedingly fast and cannot be directly measured by manual methods (17, 36, 37). To evaluate those activities, we chose conditions that either slowed the reactions to directly measurable rates or enabled us to determine the steady-state levels of reaction intermediates and thereby infer their relative turnover rates. The rationale, design, and analysis of those experiments are detailed in the relevant sections below.

Autophosphorylation of CheA Δ P2. When CheA is not coupled to receptor control, its autophosphorylation rate (Fig. 3, k_1) depends mainly on ATP concentration ($K_m \approx 200 \ \mu$ M) and is low enough to measure by manual methods (half-life of ~15 s). Thus, CheA and CheA Δ P2 were mixed with a saturating level of [γ -³²P]ATP (1 μ M), and the formation of radiolabeled protein was monitored over time (Fig. 4). Both reactions followed pseudo-first-order kinetics, but the autophosphorylation activity of CheA Δ P2 differed from that of CheA in two respects. First, the rate constant was about twofold greater for the mutant protein (0.024 ± 0.005 s⁻¹ for CheA and 0.046 ± 0.011 s⁻¹ for CheA Δ P2 [averages and standard deviations of

five determinations]). The higher specific activity of CheA Δ P2 is most likely a consequence of changing the length and nature of the segment connecting the substrate histidine to the ATPbinding site in the catalytic center, which could allow more frequent productive encounters between them. Second, the steady-state phosphorylation level of CheAAP2 was consistently severalfold higher than that of wild-type CheA (Fig. 4). We have not investigated the basis for this difference, but it may be that the forward (ATP-mediated) and backward (ADPmediated) reactions are differentially affected by the removal of the P2 domain or that a greater proportion of the His-48 substrate sites are accessible to catalysis in the mutant protein. Alternatively, CheA Δ P2 might be phosphorylated at one or more sites in addition to His-48, but this seems unlikely in view of the fact that all of its phosphoryl groups are available for transfer to CheB and CheY (see below).

Receptor control of CheAAP2 autophosphorylation. CheA and CheA Δ P2 were mixed with the coupling protein CheW and membranes containing the serine chemoreceptor, Tsr, to form ternary signaling complexes. The autophosphorylation rate of CheA in such complexes is typically too high to measure directly without rapid-reaction instruments (40). To slow the receptor-stimulated autophosphorylation reaction, we used a limiting ATP concentration (45 nM) and determined the initial rate of the reactions (before ADP product levels had risen sufficiently to cause a significant reverse reaction). We fitted the CheA and CheA Δ P2 data points to single exponentials to obtain a pseudo-first-order rate constant for each reaction. Under these conditions, the autophosphorylation activities of both CheA and CheAAP2 were stimulated about 150-fold (data not shown). The formation of phospho-CheA Δ P2 was about twofold faster and its extrapolated steady-state level was about twofold higher than those for phospho-CheA (data not shown), reminiscent of the activity differences seen in the absence of receptor coupling. No stimulation was seen for either CheA protein if CheW was omitted from the coupling reactions, implying that CheA Δ P2 forms ternary signaling complexes in the same manner as CheA (data not shown). Moreover, saturating levels of the attractant serine reduced



FIG. 4. Receptor-uncoupled autophosphorylation activity of CheA Δ P2. Reaction mixtures contained purified CheA and CheA Δ P2 (5 μ M) and [γ -³²P]ATP (1 mM). The data points were fitted to an exponential function to determine the pseudo-first-order rate constant and steady-state phosphorylation level of each reaction (10).



FIG. 5. Receptor-uncoupled phosphotransfer from phospho-CheA Δ P2 to CheB (A) and CheY (B). Reaction mixtures contained CheB or CheY at the indicated concentrations and, in the case of CheY, 0.5 μ M CheZ to augment turnover of phospho-CheY. Phosphotransfer was initiated by addition of 0.5 μ M ³²P-CheA or ³²P-CheA Δ P2, and the phosphodonor levels were monitored for 600 s (14 time points) for CheB and for 120 s (seven time points) for CheY to determine their half-lives ($t_{1/2}$) at each phosphoacceptor concentration tested.

autophosphorylation activity to approximately the same extent in both CheA and CheA Δ P2 signaling complexes (2.7-fold versus 4.5-fold [averages of seven determinations]). In summary, the receptor coupling experiments demonstrated that CheA Δ P2 can form receptor signaling complexes that regulate its autophosphorylation ability in the same manner as wild-type CheA.

Phosphotransfer by CheA Δ **P2.** Phosphotransfer from phospho-CheA to CheY occurs on a millisecond time scale at physiological reactant concentrations (k_{cat} of ~750 s⁻¹) (22, 36). In a study that used rapid-reaction instruments, Stewart et al. found that phosphotransfer from phospho-CheA Δ P2 to CheY was about 25-fold slower than that from wild-type CheA but still quite fast under physiological conditions (37). To extend these studies to CheB, we devised two experimental paradigms for comparing the phosphotransfer abilities of CheA and CheA Δ P2 without the use of rapid-reaction instruments. In describing those experiments, we refer to the reactions and rate constants summarized in Fig. 3.

In the first experiment, CheA and CheA Δ P2 were prelabeled by autophosphorylation with $[\gamma^{-32}P]ATP$. We then monitored the dephosphorylation of purified phospho-CheA and phospho-CheA Δ P2 upon addition of CheB (k_2) or CheY (k_4). The reactant stoichiometries were adjusted so that only a small fraction (<20%) of the phosphoacceptor molecules would be phosphorylated at steady state (data not shown), thereby ensuring that hydrolysis of phospho-CheB (k_3) or phospho-CheY (k_5 , assisted by CheZ) was not rate limiting in the overall dephosphorylation reaction. To slow the phosphotransfer reaction, these experiments were done at protein concentrations well below reported physiological levels. Under these conditions, we determined the half-times for the phospho-CheA and phospho-CheA Δ P2 dephosphorylation reactions at several concentrations of CheB and CheY (Fig. 5). At the lowest phosphoacceptor concentrations, dephosphorylation of phospho-CheAAP2 by both CheB (Fig. 5A) and CheY (Fig. 5B) was approximately fourfold slower than that for phospho-CheA $(4.0 \pm 0.9 \text{ for CheB} \text{ and } 3.5 \pm 1.3 \text{ for CheY})$. At higher phosphoacceptor concentrations, dephosphorylation was faster and the difference in phosphotransfer ability was less apparent (Fig. 5). At nominally physiological phosphoacceptor concentrations ($\sim 1 \mu M$ and above), CheA and CheA Δ P2 were indistinguishable as phosphodonors in this assay (data not shown).

To compare the phosphodonor abilities of phospho-CheA and phospho-CheADP2 at more physiologically relevant concentrations, we set up a coupled reaction sequence (Fig. 3) beginning with CheA or CheA Δ P2 autophosphorylation (k_1), followed by phosphotransfer to CheB (k_2) or CheY (k_4) , and ending with dephosphorylation of phospho-CheB (k_3) or phospho-CheY (k_5) . We devised conditions in which phosphotransfer was the rate-limiting step in the overall flow of phosphate groups from ATP to P_i and examined the effect of phosphoacceptor concentration on the steady-state levels of phospho-CheB and phospho-CheY. To ensure that the phosphodonor supply was not rate limiting in the overall reaction scheme, we assembled stimulatory chemoreceptor signaling complexes and used a saturating concentration of ATP to accelerate the autophosphorylation step. Because CheA and CheA Δ P2 have similar receptor-stimulated autophosphorylation rates at subsaturating ATP levels (see above), we assumed that their rates would also be comparable at ATP excess. Moreover, we assumed that those rates would not be influenced by changes in the phosphoacceptor concentration. At an initial concentration of 1 mM ATP, the level of phosphodonors (phospho-CheA or phospho-CheA Δ P2) reached steady state by 10 s and remained at steady state for at least 30 s (data not shown). Over that interval, less than 5% of the ATP was consumed in the course of the reactions (data not shown). Under these conditions, we found that the proportion of CheA or CheA Δ P2 molecules in the phosphorylated form at steady state dwindled with increasing amounts of phosphoacceptor in the reaction mixtures (data not shown), indicating that until the concentrationdependent phosphotransfer reaction outpaced the concentration-independent autophosphorylation reaction, phosphodonor production was not rate limiting. The phosphoacceptor concentrations that produced half-maximal steady-state levels of phospho-CheA were similar to their in vivo levels ($\sim 2 \mu M$ for CheB and $\sim 1 \mu M$ for CheY [data not shown]). In this concentration range, the steady-state level of phospho-CheB or phospho-CheY in the reactions should reflect the ratios of their phosphotransfer to dephosphorylation rates $(k_2/k_3 \text{ or }$ k_4/k_5). We assumed that the rates of the dephosphorylation reactions $(k_3 \text{ and } k_5)$ were independent of phosphoacceptor concentration and the nature of the phosphodonor (phospho-CheA or phospho-CheA Δ P2). Thus, any phosphodonor-dependent differences in the steady-state levels of phospho-CheB or phospho-CheY should reflect differences in their respective phosphotransfer rates $(k_2 \text{ or } k_4)$.

In the CheB titration experiment, the steady-state level of phospho-CheB rose substantially more slowly with phospho-CheA (Fig. 6A). The difference in the CheY titration was equally dramatic (Fig. 6B). (Note, as detailed in the legend to Fig. 6, that CheZ was added to the CheY titration to ensure that dephosphorylation of phospho-CheY was not rate limiting.) These results imply that phosphotransfer to both CheB and CheY was substantially slower from phospho-CheA $\Delta P2$ than from phospho-CheA. The reduction in the phosphotransfer rate of CheA $\Delta P2$ appeared to be similar for both phosphoacceptors. At physiological reactant concentrations (e.g., 1 μ M phosphodonor and 1 to 10 μ M phosphoacceptor), phospho-CheA $\Delta P2$ supported



FIG. 6. Steady-state levels of phospho-CheB (A) and phospho-CheY (B) in chemoreceptor-activated reaction systems. Reaction mixtures contained 0.5 μ M CheA and CheA Δ P2, 4 μ M membrane-embedded Tsr (serine chemoreceptor), 4 μ M CheW, and variable amounts of CheB or CheY. The CheY reaction mixtures also contained 0.5 μ M CheZ to augment phospho-CheY turnover. The phosphorylation cascade was initiated by addition of 1 mM [γ^{-32} P]ATP, and the levels of phospho-CheB and phospho-CheY were measured after 10 and 20 s, when the reactions had reached steady-state.

approximately fivefold-lower steady-state levels of phospho-CheB or phospho-CheY than did phospho-CheA. We conclude that the P2 domain of CheA enhances the CheB and CheY phosphotransfer reactions to similar extents.

Behavior of a *cheA* Δ P2 mutant. The *cheA* Δ P2 allele was transferred from pKJ9-1.1 into the E. coli chromosome by homologous recombination, as detailed in Materials and Methods. The resulting *cheA* Δ P2 mutant (UU1121) showed reduced chemotactic ability on semisolid tryptone agar, spreading at about 25% the rate of UU1120, an isogenic cheA⁺ control strain (Fig. 7A). However, the mutant colonies still exhibited cell bands characteristic of chemotactic movements, indicating that the behavioral defect of UU1121 was not complete. Conceivably, the normal receptor-mediated regulation of CheA Δ P2 autophosphorylation activity permits the mutant cells some stimulus control over flagellar rotation despite a reduced rate of phosphotransfer to CheB and CheY. We compared the unstimulated flagellar rotation patterns of *cheA* Δ P2 and *cheA*⁺ strains by cell tethering (Fig. 7B). Whereas most of the wildtype cells exhibited frequent reversals of motor rotation, more than two-thirds of the mutant cells rotated exclusively in the CCW direction during the 15-s observation period. The low frequency of CW rotation in UU1121 is consistent with a reduced rate of phosphotransfer from CheA Δ P2 to CheY and most likely accounts for its chemotactic disability.

Phenotypic rescue of a *cheA* Δ P2 mutant by free P1 domains. Upon prolonged incubation on semisolid agar, UU1121 gave rise to pseudorevertants with enhanced chemotactic ability (data not shown). If its CCW-biased flagellar rotation and poor chemotactic ability are due to the reduced phosphotransfer activity of CheA Δ P2, mutations that either augment the steady-state pool of phospho-CheY or otherwise increase the CW bias of the flagellar motors could conceivably alleviate those behavioral defects. To test this idea, we examined the behavior of UU1121 cells containing high levels of CheA[1-149], the P1 domain of CheA. At high stoichiometries, P1 fragments can efficiently exchange phosphoryl groups with CheB and CheY, in effect creating a reservoir of CheA-generated phosphoryl groups (10). Because phospho-P1 has a much longer half-life than phospho-CheB and phospho-CheY, CheA[1-149] fragments should serve to buffer the loss through dephosphorylation of signaling phosphates from CheB and CheY and thereby raise the steady-state levels of phospho-CheY and phospho-CheB.

We found that pAG3, an IPTG-inducible P1 expression plasmid, produced a dramatic improvement in the chemotactic ability of UU1121 (Fig. 8). In the absence of IPTG inducer, pAG3-containing cells spread on soft agar plates at essentially the same rate as vector-containing control cells (Fig. 8A). However, in the presence of IPTG, pAG3-containing cells spread much faster than the control cells. At maximal induction (320 μ M IPTG), UU1121/pAG3 spread at ~75% of the wild-type rate, about threefold faster than control cells (Fig. 8B). This improvement in chemotactic ability was accompanied by an increase in the proportion of cells with frequent episodes of CW flagellar rotation (Fig. 8C), suggesting that the aberrant steady-state swimming pattern of UU1121, rather than a defect in stimulus detection per se, is primarily responsible for its poor chemotactic ability.

DISCUSSION

Signaling defects of CheA Δ P2. To investigate the contribution of the P2 domain to chemotactic signal processing by CheA, we constructed several mutant proteins in which the P2 domain had been replaced by a flexible linker and characterized one of them in detail. We found that extirpation of P2 had no deleterious effect on the CheA autophosphorylation reaction. In fact, the mutant protein autophosphorylated about twice as fast as wild-type CheA and reached roughly twofoldhigher steady-state levels. CheA autophosphorylation is a *trans* reaction in which the phosphorylation site from one subunit of the dimer interacts with the dimerization and ATP-binding domains from the other subunit. At saturating ATP concentrations, the rate of this reaction is determined mainly by the frequency of collisional encounters between the P1 domain and the P3/P4 domain of the other subunit. However, the P1



FIG. 7. Chemotaxis phenotypes of UU1121 (*cheA* Δ P2). (A) Colony morphology on semisolid agar. Colonies of UU1120 (*cheA*⁺) and UU1121 (*cheA* Δ P2) were transferred with a toothpick to a semisolid tryptone agar plate and incubated at 35°C for 8 h. (B) Flagellar rotation pattern. Strains UU1120 (*cheA*⁺) and UU1121 (*cheA* Δ P2) were analyzed by cell tethering as described in Materials and Methods. Rotating cells were assigned to five categories: exclusively CCW or CW, predominantly CCW or CW with some reversals (CCW-R and CW-R, respectively), or with frequent reversals and no significant directional bias (CCW/CW).



FIG. 8. Suppression of CheA Δ P2 chemotaxis defects by free P1 domains. The chemotaxis phenotypes of strain UU1121 (*cheA* Δ P2) carrying pAG3 (expressing CheA[1-149] under IPTG-inducible control) are compared to those of UU1121 carrying pTM30 (vector control). (A) Rates of colony expansion on semisolid tryptone agar containing various IPTG concentrations. The diameters of colonies incubated at 35°C were measured at three or more time points to calculate an expansion rate. (B) Colony sizes on semisolid tryptone agar containing 320 μ M IPTG. The plate was incubated for 8 h at 35°C. (C) Flagellar rotation patterns. The rotation patterns of tethered UU1121/pAG3 cells, grown without inducer or with 320 μ M IPTG, were classified as described in the legend to Fig. 7B.

phosphorylation sites in a wild-type dimer are probably in steric conflict (16). The absence of the P2 domain in CheA Δ P2 might be expected to enhance the flexibility of the P1-P3/P4 connection, which could facilitate the autophosphorylation reaction by alleviating steric clashes between the two subunits.

In ternary complexes with CheW and the serine receptor (Tsr), CheA Δ P2 responded like wild-type CheA to chemoreceptor control: the rate of autophosphorylation activity was approximately 100-fold greater than the receptor-uncoupled rate and was subject to down-regulation by serine stimuli. Evidently, the P2 domain of CheA plays no important role in the assembly or subsequent autophosphorylation behavior of receptor signaling complexes.

The principal biochemical difference between CheA Δ P2 and wild-type CheA was the speed of their phosphotransfer reactions with CheB and CheY. We showed with several qualitative reaction schemes that CheA Δ P2 had reduced rates of phosphotransfer to both CheB and CheY, consistent with the findings of a more precise kinetic analysis of the CheY reaction (37). Most importantly, CheA Δ P2 phosphotransfer was at least 10-fold slower than that of wild-type CheA in ternary signaling complexes, which would most likely reduce the steady-state phosphorylation levels of the CheB and CheY response regulators in vivo.

Chemotaxis without P2. CheA Δ P2 cells exhibited predominantly CCW flagellar rotation, consistent with a reduced intracellular level of phospho-CheY. If phospho-CheB levels are also reduced in the CheA Δ P2 mutant, we would expect the cells to have elevated receptor methylation states owing to feedback control of receptor signals by the sensory adaptation system. Although we did not examine this prediction in the present work, the low tumbling frequency of the mutant cells demonstrates that their adaptation system was not able to fully compensate for the decreased flux of phosphates to CheB and CheY. Nevertheless, CheA Δ P2 cells exhibited slow chemotactic spreading on semisolid medium, indicative of functional chemoreceptor control of CheA Δ P2 activity.

If the aberrant tumbling pattern of the CheA Δ P2 mutant is

responsible for its low rate of chemotactic spreading, mutations that raise the tumbling frequency should improve its chemotactic ability. Indeed, the CheA Δ P2 mutant readily acquires mutations that enhance its rate of chemotactic spreading, suggesting that it may be amenable to a number of bypasstype suppression mechanisms. Preliminary genetic analyses of such pseudorevertants has revealed a variety of suppressor mutations located in or near the main cluster of chemotaxis and flagellar genes at min 42 on the *E. coli* chromosome (A. Garzón and C. Jensen, unpublished results). These suppressors might include flagellar switch mutations, partial *cheZ* lesions, *cheY* overproducers, signal-biased receptor mutations, and other alterations that promote CW flagellar rotation without compromising phosphorelay signaling.

To show more directly that the CheA Δ P2 mutant was proficient at chemotactic signaling, we contrived to elevate its steady-state intracellular levels of phospho-CheB and phospho-CheY by expressing additional P1 domains in the cells, a tactic previously shown to restore chemotactic ability to a CheA Δ P1P2 mutant lacking both the P1 and P2 domains (10). Indeed, elevated P1 levels improved the chemotactic ability of the CheA Δ P2 mutant to about 75% of that of the wild type. The probable signaling mechanism for the P1 helping effect is shown in Fig. 9. In the CheA Δ P1P2 system described previously (10), the free P1 domains have to acquire phosphoryl groups through trans interactions with the ATP-binding domains of the CheA Δ P1P2 molecules before relaying them to CheB and CheY. In the CheAAP2 system, the P1 domain resident in the CheA Δ P2 molecule most likely excludes free P1 domains from interacting with the ATP-binding and catalytic center, as demonstrated in several analogous cases (10, 16). Thus, essentially all phosphoryl groups transmitted to CheB and CheY probably pass through the P1 domain tethered to the CheA Δ P2 molecules (Fig. 9). The free P1 domains most likely influence the flagellar bias of CheA Δ P2 by participating in reversible exchange reactions with phospho-CheB and phospho-CheY (10), as demonstrated previously (10). Because phospho-P1 is much less susceptible to hydrolysis, the ex-



FIG. 9. Model of chemotactic signaling in a CheA Δ P2 mutant containing free P1 domains. CheA Δ P2 autophosphorylates at essentially normal rates and responds normally to chemoreceptor coupling control. However, without the target-acquiring P2 domain, the reduced rate of phosphotransfer to CheB and CheY cannot support efficient chemotaxis. A stoichiometric excess of free P1 domains may enhance chemotactic ability by buffering the CheA Δ P2-generated pool of signaling phosphates through reversible exchange reactions with B \sim P and Y \sim P. See text for additional details of the model.

change reactions should slow the irreversible loss of phosphoryl groups from CheB and CheY, thereby serving to buffer the phosphoryl pool. Using an empirical relationship between cellular phospho-CheY levels and CW rotational bias derived by Cluzel et al. (8), we estimate that P1 buffering would need to increase intracellular phospho-CheY levels only about 15% (from ~2.4 to ~2.8 μ M) to produce the observed CW shift in CheA Δ P2 flagellar rotation pattern. This modest increase is consistent with the extent of P1 buffering seen in our previous in vitro study (10). In summary, our model proposes that the free P1 domains create a phosphoryl group reservoir that serves to augment the steady-state phosphorylation levels of CheB and CheY.

This signaling model predicts that P1-assisted CheA Δ P2 cells might show delayed motor responses to stimuli, such as attractant increases, that suppress CheA autophosphorylation because the ensuing drop in the phospho-CheY level would be slowed by replenishment from the phospho-P1 buffer. This delay could be largely offset by an accompanying increase in CheZ activity, but the case for stimulus control of CheZ remains controversial. Thus, an increased response latency may be one of the factors that limits the chemotactic efficiency of P1-assisted CheA Δ P2. However, on semisolid media, where metabolism-generated attractant gradients are steep, response latency is probably a less critical factor than tumbling rate. Under such conditions, the cells might show more profound defects in their response to CheA-activating stimuli, such as repellent increases, because the model provides no obvious way to accelerate the flux of phosphoryl groups to CheY. This analysis assumes that the relatively slow P2-independent phosphotransfer reaction is the rate-limiting step, as indicted by our steady-state measurements with receptor-coupled CheA Δ P2. Again, this predicted signaling defect could be offset by a concomitant reduction in CheZ activity, allowing phospho-CheY levels to rise through slowed dephosphorylation. These considerations suggest that the P1-assisted CheA Δ P2 signaling

system might prove valuable in exploring the possible stimulus control of CheZ activity.

The signaling role of P2. This study and a previous study (37) demonstrate that the principal signaling role of the P2 domain is to facilitate the transfer of CheA phosphoryl groups to CheB and CheY through reversible binding reactions. Despite the specificity of P2 binding behavior, two observations indicate that it is not the primary determinant of signaling specificity in the chemotaxis phosphorelay. First, the unassisted P1 domain is capable of rapid phosphotransfer to CheB and CheY, suggesting that it can discriminate among potential phosphoacceptor partners. If a discrete binding interaction is involved, it must have very low affinity, because the P1 phosphotransfer reactions to CheB and CheY are not readily saturable (37). Alternatively, P1 residues might promote phosphotransfer specificity by controlling access to the donor phosphohistidine or by participating with target residues in the phosphotransfer chemistry. Second, although the reactivity of P1 with heterologous response regulators has not been examined in detail, the proficient chemotaxis of the P1-assisted CheA Δ P2 mutant might mean that P1 is not a highly promiscuous phosphodonor.

The P2 domain seems to be a relatively recent evolutionary embellishment to CheA. It was probably acquired not because it enhanced signaling specificity, but rather because it accelerated signaling reactions that were already quite specific. For example, a P2-promoted increase in the cell's phospho-CheY level might have increased turning episodes while swimming, which was possibly an improved foraging pattern. Whatever the evolutionary scenario, in the right genetic background the P2 domain is largely dispensable, at least for chemotactic movements on semisolid media. Moreover, some CheA homologues, such as FrzE of *M. xanthus*, lack a P2 domain and may have never had one. Instead, their signaling partner is covalently tethered to the phosphodonor molecule, ensuring a rapid phosphotransfer reaction. The P1 domain, a more ancient and essential component of the CheA kinase, probably carries the critical (and still poorly understood) determinants of signaling specificity in the chemotaxis phosphorelay.

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