



Article Evidence for Pentapeptide-Dependent and Independent CheB Methylesterases

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Abstract: Many bacteria possess multiple chemosensory pathways that are composed of homologous signaling proteins. These pathways appear to be functionally insulated from each other, but little information is available on the corresponding molecular basis. We report here a novel mechanism that contributes to pathway insulation. We show that, of the four CheB paralogs of *Pseudomonas aeruginosa* PAO1, only CheB₂ recognizes a pentapeptide at the C-terminal extension of the McpB (Aer2) chemoreceptor ($K_D = 93 \mu$ M). McpB is the sole chemoreceptor that stimulates the Che2 pathway, and CheB₂ is the methylesterase of this pathway. *Pectobacterium atrosepticum* SCRI1043 has a single CheB, CheB_Pec, and 19 of its 36 chemoreceptors contain a C-terminal pentapeptide. The deletion of *cheB_Pec* abolished chemotaxis, but, surprisingly, none of the pentapeptides bound to CheB_Pec. To determine the corresponding structural basis, we solved the 3D structure of CheB_Pec. Its structure aligned well with that of the pentapeptide-dependent enzyme from *Salmonella enterica*. However, no electron density was observed in the CheB_Pec region corresponding to the pentapeptide-binding site in the *Escherichia coli* CheB. We hypothesize that this structural disorder is associated with the failure to bind pentapeptides. Combined data show that CheB methylesterases can be divided into pentapeptide-dependent enzymes.

Keywords: bacterial signal transduction; chemosensory pathways; chemoreceptor; X-ray structure; C-terminal pentapeptide; CheB; methylesterase

1. Introduction

Chemosensory pathways are among the most abundant prokaryotic signal transduction mechanisms [1]. Apart from mediating flagellum based chemotaxis, chemosensory pathways carry out alternative cellular functions like the control of second messenger levels or type IV pili-based motility [1–3]. The key element of a chemosensory pathway is the ternary complex formed by chemoreceptors, the CheA autokinase and the CheW coupling protein. Signaling is typically initiated by signal recognition at the chemoreceptor ligand-binding domain (LBD) that creates a molecular stimulus modulating CheA autophosphorylation and, subsequently, transphosphorylation to the CheY response regulator. The ratio of CheY to phosphorylated CheY (CheY-P) defines the pathway output [4].

The pathway sensitivity is adjusted by the coordinated action of the CheR methyltransferase and the CheB methylesterase that catalyze methylation and demethylation, respectively, of several glutamate residues at the chemoreceptor-signaling domain. It was shown that poorly methylated chemoreceptors have high chemoeffector affinity and high propensity for methylation [4,5]. Genome analyses revealed that genes encoding both enzymes are present in the large majority of chemosensory pathways and are thus among the six core pathway proteins [1].

In *Escherichia coli*, CheR was found to bind to the methylation site of the Tar chemoreceptor with a rather modest affinity, ranging between 100 to 200 μ M, depending on experimental conditions [6]. However, Tar possesses a C-terminal pentapeptide that is tethered to the C-terminal end of the chemoreceptor signaling domain via an unstructured linker [7]. This pentapeptide, NWETF, represents an additional binding site for CheR and CheB [8,9]. CheR from *E. coli* bound NWETF with a *K*_D value affinity of approximately 2 μ M, an affinity that is around 100-fold higher than the affinity for the methylation site [6,10]. It was proposed that CheR binding to the pentapeptide enhanced the local CheR concentration, leading to the optimal adaptation [11,12]. Remarkably, genome analyses indicated that approximately 10% of chemoreceptors possess a C-terminal pentapeptide [9], and experimental studies, as well as sequence analyses of CheR from different species, have shown that this protein family can be subdivided into pentapeptide-dependent and independent enzymes that are either able or unable to bind these pentapeptides [8,13]. The structure of the CheR-pentapeptide complex has been solved [14], and several sequence features at or close to the pentapeptide-binding site have been identified to be specific for each CheR subfamily [8,13].

Compared to CheR, less information is available on CheB function. Much of what we know of this protein family is due to the studies of the enzymes from *E. coli* [15–19] and *Salmonella enterica* sv. Typhimurium [20–23] that share 95% of their amino acid sequence identity (Figure S1). Both species possess a single chemosensory pathway, a single pentapeptide-dependent CheB and two chemoreceptors, Tar and Tsr, that contain a C-terminal pentapeptide. CheB from *E. coli* bound this pentapeptide with much lower affinity ($K_D = 130$ to 160 µM, depending on the method used) [17] as compared to CheR. The affinity of CheB for the pentapeptide is thus too low to increase the local concentration, but the CheB-pentapeptide interaction was found to stimulate methylesterase activity [17]. CheB is composed of a phosphoryl group-accepting receiver domain (REC) and a methylesterase domain [20], and it was shown that REC domain phosphorylation stimulates the catalytic activity of CheB [17].

The mutation or removal of the NWETF pentapeptide from the *E. coli* Tar and Tsr chemoreceptors largely reduced methylation and demethylation *in vivo* and *in vitro* and abolished chemotaxis [11,15,19,24,25]. This results in the paradoxical situation where C-terminal pentapeptides are essential for the functioning of some receptors, like Tar and Tsr, but are absent from many other chemoreceptors that mediate strong chemotactic responses [26–30]. Consequently, the physiological relevance of pentapeptide-dependent chemosignaling remains to be established, but we recently showed that pentapeptide-containing chemoreceptors are more abundant in bacteria that maintain host interactions [9]. In addition, it is only a little clear to which degree there are pentapeptide-independent CheB. To address this latter issue, we report here studies of CheB homologs from the human and plant pathogenic bacterial strains *Pseudomonas aeruginosa* PAO1 [31] and *Pectobacterium atrosepticum* SCRI1043 [32].

P. aeruginosa PAO1 has five gene clusters encoding signaling proteins that assemble into four chemosensory pathways (Figure 1) [33].

These pathways differ in function: whereas the Che pathway mediates chemotaxis [34,35], the Wsp pathway controls the c-di-GMP levels [2], and the Chp pathway was associated with type IV pili-mediated motility and cAMP levels [3,36]. The function of the Che₂ pathway is unknown. As shown in Figure 1, each chemosensory pathway contains a CheR and CheB homolog. Experimental and bioinformatic studies indicate that, of the 26 chemoreceptors, McpB (synonym Aer2) (Figure 1) is the sole chemoreceptor that feeds into the Che₂ pathway [33,37]. Furthermore, it is the only chemoreceptor with a C-terminal pentapeptide [13]. We previously showed that, of the four CheR homologs, the methyltransferase of the Che₂ pathway, CheR₂, was the only homolog that bound

the McpB pentapeptide [13]. Furthermore, binding to this pentapeptide was essential for the CheR₂ interaction and methylation of McpB [13]. We concluded that the specific pentapeptide-CheR₂ interaction is a mechanism that permits the targeting a particular chemoreceptor with a specific CheR [13].

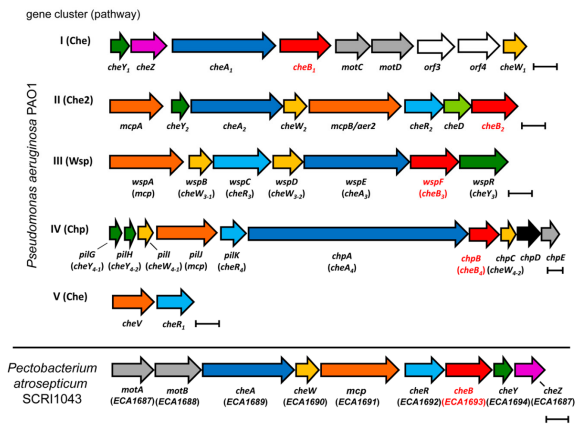


Figure 1. Gene clusters encoding chemosensory signaling proteins in *Pseudomonas aeruginosa* PAO1 and *Pectobacterium atrosepticum* SCRI1043. Genes of the same family are colored in the same color. The genes of the proteins studied in this article are shown in red. Bars, 0.5 kbp.

The four CheB homologs of *P. aeruginosa* were found to play important physiological roles. A *cheB*₁ mutant was nonchemotactic [38], and screening of a 2200-member mutant library for virulence defects in a cystic fibrosis airway *P. aeruginosa* isolate revealed that a *cheB*₂ mutant showed one of the strongest phenotypes in *Caenorhabditis elegans*, a finding that was confirmed by experimentation on mice [39]. A *cheB*₃/*wspF* mutant caused elevated c-di-GMP levels and enhanced biofilm formation [2], due to locking this pathway into an active state. Interestingly, cystic fibrosis airway infections frequently produce rugose small-colony variants (RSCV), and this phenotype could be reverted by the *in trans* expression of *cheB*₃/*wspF*, indicating that *cheB*₃/*wspF* mutations are a very frequent mechanism for generating RSCV morphotypes [40]. As for the *cheB*₄ gene, its mutation caused hyper-piliation [3], prevented swarming, and formed more robust biofilms by stimulating matrix production [41]. Considering the pentapeptide-mediated specific interaction of CheR₂ and McpB, we studied here the interaction of the McpB pentapeptide with the four CheB homologs.

Since *P. aeruginosa* has four CheB paralogs and a single pentapeptide-containing chemoreceptor, we aimed at studying the inverse situation, i.e., a bacterium with a single CheB but multiple chemoreceptors with pentapeptides of different sequences. To this end, we chose *P. atrosepticum* SCRI1043 as a model that has a single CheB and 19 chemoreceptors with a pentapeptide [32]. *P. atrosepticum* is among the top 10 plant pathogens [42] and the causative agent of soft rot diseases in many agriculturally relevant crops [43]. This species belongs, like *E. coli* and *S. enterica*,

to the *Enterobacteriacae* family. Combined data from both strains allows distinguishing between pentapeptide-dependent and independent CheB proteins.

2. Results

2.1. P. aeruginosa CheB₂ Is the Only CheB Homolog That Binds to the McpB Chemoreceptor Pentapeptide

We reported previously that CheR₂ is the only one of the four *P. aeruginosa* CheR homologs that binds the terminal pentapeptide of the McpB chemoreceptor [13]. To verify which CheB homolog interacts with this peptide, the four CheB homologs were overexpressed in *E. coli*, purified, and submitted to microcalorimetric titrations with the pentapeptide of the McpB receptor, GWEEF. The titration of CheB₁, CheB₃, and CheB₄ with the peptide caused small and uniform heat changes that were similar to ligand dilution heats (Figure 2). In contrast, exothermic binding heats were observed for the titration of CheB₂, and a dissociation constant (K_D) of 93 ± 15 µM was derived.

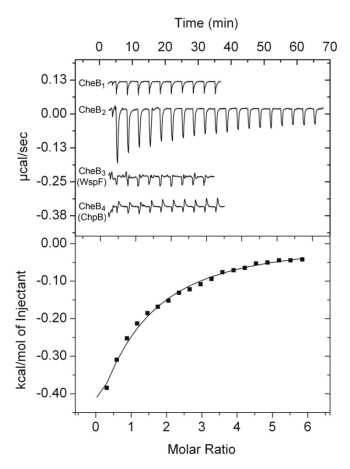


Figure 2. Specificity of the interaction between four CheB homologs of *P. aeruginosa* and the terminal pentapeptide GWEEF of the McpB chemoreceptor. Microcalorimetric titrations of the four CheB homologs (15 to 40 μ M) with 14- μ L aliquots (1 to 7 mM) of the GWEEF pentapeptide. Upper panel: raw titration data; lower panel: integrated, dilution heat-corrected and concentration-normalized peak areas of the titration data for CheB₂. Data were fitted using the "one binding site model" of the MicroCal version of ORIGIN.

This indicated that the CheR and CheB homologs of the Che₂ pathway specifically interact with the only chemoreceptor that feeds into this pathway, McpB (Figure 1). The measured affinity was approximately 180-fold lower than that for the peptide binding to CheR₂ [13] ($K_D = 0.52 \mu$ M), implying that CheR₂ largely outcompetes CheB₂ for binding at the GWEEF pentapeptide. Next, it was investigated whether CheB₂ phosphorylation alters the affinity for the pentapeptide. Since the

phosphorylation half-life of CheB proteins is typically very short [44], we generated stable beryllium fluoride adducts that mimic phosphorylation [45]; however, protein precipitation made any biochemical study impossible. Previous studies showed that the replacement of the phosphoryl group accepting aspartate with glutamate in receiver domains mimics protein phosphorylation [46]. We generated the CheB₂ D55E mutant protein that was titrated with the GWEEF peptide, resulting in a K_D of 56 ± 14 μ M (Figure S2), representing a modest increase in affinity as compared to the native protein.

2.2. The Signaling Gene Cluster of P. atrosepticum SCRI1043 Encodes a Chemosensory Pathway That Mediates Chemotaxis

Chemosensory pathways can exert a number of different functions like chemotaxis, modulating second messenger levels and type IV pili-based movement [1]. To explore the function of the sole chemosensory gene cluster in *P. atrosepticum* SCRI1043 (Figure 1), we created deletion mutants of the *cheA* and *cheB* genes. Subsequently, we conducted quantitative capillary chemotaxis assays of the wild-type (wt) and mutant strains towards casamino acids. As shown in Figure 3, the wt strain showed strong chemotactic responses, whereas the *cheA* and *cheB* mutants failed to respond, indicating that the chemosensory pathway mediates chemotaxis.

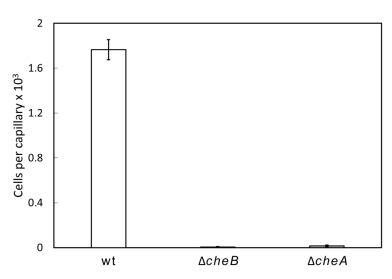


Figure 3. Quantitative capillary chemotaxis assays of wild-type and mutant strains of *P. atrosepticum* SCRI1043 towards 0.1% (*w*/*v*) casamino acids. Data were corrected with the bacteria that migrated into buffer-containing capillaries (225 ± 35). Data are means and standard deviations from three experiments conducted in triplicate. wt: wild-type.

2.3. P. atrosepticum SCRI1043 Contains a Large Number of Chemoreceptors with a C-Terminal Pentapeptide

The chemoreceptor repertoire of *P. atrosepticum* SCRI1043 is illustrated in Figure 4. Previous studies have shown that approximately 14% of bacterial chemoreceptors lack transmembrane regions and are thus involved in the sensing of cytoplasmic signals [47]. No such receptors are present in *P. atrosepticum* SCRI1043, since all 36 chemoreceptors are membrane-bound and possess two transmembrane regions (Figure 4).

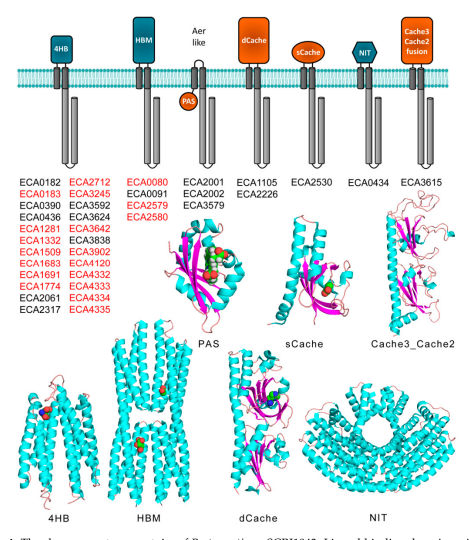


Figure 4. The chemoreceptor repertoire of *P. atrosepticum* SCRI1043. Ligand-binding domains with α/β folds or parallel helices are shown in orange or blue, respectively. Chemoreceptor names in red indicate receptors with C-terminal pentapeptides. Shown below are representative 3D structures of these domains, namely the structure of Tar-LBD (4HB) in a complex with aspartate (PDB ID: 1vlt), McpS-LBD (HBM) in a complex with malate and acetate (PDB ID: 2yfa), Aer2-LBD (PAS) in a complex with heme (PDB ID: 4hi4), TlpQ-LBD (dCache) in a complex with histamine (PDB ID: 6fu4), PscD-LBD (sCache) in a complex with propionate (PDB ID: 5g4z), NasR-LBD (NIT) (PDB ID: 4akk), and a homology model of the ECA3615-LBD (Cache3_Cache2 fusion) generated by SwissModel [48] using PDB ID 4avf as template.

There are three receptors that possess the typical topology and domain arrangement of Aer receptors [49] that mediate aerotaxis (Figure 4). Only two receptors possess a dCache type LBD that are highly abundant sensor domains in chemoreceptors and sensor kinases [50] and respond mainly to different amines [26]. In addition, the repertoire contains one and four receptors with sCache or HBM LBDs, respectively, that are typically organic acid sensors [51]. The alignment of the C-terminal segment of *P. atrosepticum* SCRI1043 chemoreceptors revealed that 19 of them possess a terminal pentapeptide that is tethered to the signaling domain via linker sequences of 29 to 39 amino acids (Figure 5).

А	
ECA0182	LADAVSAFKIPSYSHGNAGSYESVPMSTPTLSLALARKE
ECA0183	LAEAVSTFKLLSYGNGKTASYASAPTRTPTLSLAPAAAKNQSNND NWTTF
ECA1774	LARTVSVFNLGASYKSAALNRKTETPALAAPKNNRAEKTSAKGELA DWTTF
ECA2061	LSHAVAAFRI
ECA3642	LAETVSQFRLGNGHQIARTPAAAASLTLRPALAAPGKSGISAGEG DWTSF
ECA4332	LVELMKVFIVEGGSSQRIAPPLKRPSSAKFSLANPKGSAGSNNQ NWEQF
ECA4333	LVELMKVFIV <u>E</u> SGSSQRTTPELKRPSSAKLSLASPKGRTKSDSQ NWETF
ECA4120	LVTLMNHFHLRGTPAARPAPMAKKAQTARLALAPVGNTQD NWEKF
ECA4334	LLELMGVFKLNGIQTKAPRLTSQVKQPAAPRLALASKSGHTSSD NWETF
ECA4335	LVELMGVFKIDGTQSQRAVPQVTTLSRPKLALAGNSSNT NWETF
ECA2530	LNEQAQELSRTVEQFRVDESAGSYLALGAR
ECA0436	LRDAVRFFKVNQDHLLRIH
ECA1281	LQNAVEVFKI <u>NQAVAQEHRAASASSLAALPKSLLPKPTSAGSSNANWETF</u>
ECA1691	LNQAVAVFRL <u>SEDTGSFRRTTQATAGQKPVLLAPSVNGGKKAKEGSSTD</u> NWETF
ECA1683	LTRAVATFKL <u>SSHLSSGHSAPARPNALAAKGRSSLALPRQANTENG</u> NWETF
ECA3902	LTQAVAVFKL <u>SGIVQQVRSSLPKSAPQPRLAPAMAIAGSSKGNSNQ</u> NWETF
ECA3838	LNLGVSRFHLM
ECA1332	LNQTVSLFQLSDTQSALQVAAKPVRKAQAIAPRAGKALPTSSD NWEKF
ECA1509	LSAVVDVFNLDSDSDQQTAFSRPAIAAPVHRAVAQSTTPLLSVHGRHGE GWEKF
ECA2712	LTEAVSVFQLSAAEAPRRPQQRLAEKAPAAQKPMLLAAAGGKKGNAND NWETF
ECA2579	LESMVANFRLSENEGRKPKANISGLPPQQKYLPPAAKQTQDS GWTTF
ECA2580	LEKLLEHFRVSQSDNRVASRASSSIPRHTLPKSVSAKAASSESDWTSF
ECA0080	LATLMSVFRISDKDVARLQGSNTGNPNSGNKATARLPTLASRDNGND NWTTF
ECA0091	LKQAVSVFRLANAQHDDTPAGIAFNNQPHRLHAPR
ECA2226	LAQTIEHFRLEQQHALPHALLR
ECA3592	LAESMVQFKVQSQEFAAIGRF
ECA3624	LMRSMALFQVEPRLS
ECA0390	LQQSVSRFQIARENREMDNVLPGLRQNITLADAR
ECA3245	LSQLVGQFIVGQIASSSLIPALASVPSGLSAPRLASAKNKNALAQDEAGWQRF
ECA1105	LNSVVGAFRV
ECA0434 ECA2001	MESLVSHFKVDDSAPQQPLQHALLSR
ECA2001 ECA2002	MSEAVSVFSIPR MVOAASVFSLSR
ECA2002 ECA3579	INVQAASVESLSR INSAINVYGS
ECA3615	LNSAINVIGS LSELVSVFRI
ECA3015 ECA2317	LNTSVSLFILPSTEADVNPMIDQREMTQRIPMMG
ECAZ31/	"THI2A2PT TEADANEWIDGKEWIGKIEWNG
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Figure 5. C-terminal pentapeptides at *P. atrosepticum* chemoreceptors. (**A**) C-terminal section of the sequence alignment of *P. atrosepticum* SCRI1043 chemoreceptors. Pentapeptides are in boldface, and the linker sequences are underlined. Residues in green are highly similar. (**B**) Sequence logo of the 19 pentapeptides. The figure was generated using Weblogo (https://weblogo.berkeley.edu/logo.cgi).

In total, there were nine different pentapeptide sequences—among which, NWETF, the pentapeptide of the *E. coli* and *S. enterica* sv. Typhimurium receptors, was the most abundant and present in eight chemoreceptors (Table S1). Interestingly, the 19 pentapeptide-containing chemoreceptors possess either 4HB or HBM-type LBDs that correspond to either single- or double-module four-helical bundle domains, respectively (Figure 4). The linkers showed no apparent sequence similarities (Figure S3) and were predicted to be mainly unstructured (Figure S4).

2.4. P. atrosepticum SCRI1043 CheB Fails to Recognize Pentapeptides

We subsequently overexpressed and purified *P. atrosepticum* CheB (CheB_Pec) to study its interaction with the pentapeptides present in SCRI1043 chemoreceptors. However, microcalorimetric titrations conducted with all nine pentapeptides (Table S1) and at different analysis temperatures

did not show any sign of binding. This was an unexpected finding, since CheB_Pec shares 86% sequence identity with the *E. coli* CheB that was shown to bind in its unphosphorylated form the free or receptor-born NWETF pentapeptide [15–17]. To verify whether the N-terminal His-tag at CheB_Pec may potentially prevent binding, the His-tag was enzymatically removed from CheB, but no binding was observed in isothermal titration calorimetry (ITC). Since the phosphorylation of *E. coli* CheB greatly enhanced its methylesterase activity [17], we hypothesized that phosphorylation may be a necessary prerequisite for pentapeptide binding to CheB_Pec. To verify this hypothesis, we generated purified CheB_Pec containing a beryllium fluoride adduct that mimics phosphorylation [45,52]. However, microcalorimetric titrations did not evidence binding.

2.5. Three-Dimensional Structure of a Pentapeptide-Independent CheB Methylesterase

The above results suggest that the single CheB in a strain that harbors 19 chemoreceptors with a pentapeptide is unable to bind any of these pentapeptides. One possible explanation may be that the protein is unfolded or present in an inactive conformation. To address this issue, we crystallized CheB_Pec and solved its three-dimensional structure at a resolution of 2.3 Å (Figure 6).

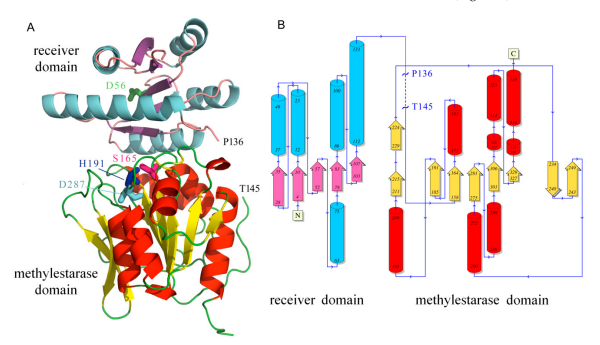


Figure 6. The three dimensional structure of *P. atrosepticum* CheB. (**A**) Ribbon diagram of the structure. The gap observed due to lacking electron density is indicated (P136 to T145). The phosphoryl group accepting aspartate (D56), as well as the residues that form the methylesterase catalytic triad (S165-H191-D287), are shown in stick mode. (**B**) Secondary structure elements: tubes: α -helix; arrows: β -strand. The figure was produced using PDBsum [53].

The enzyme is composed of an N-terminal REC domain and a C-terminal methylesterase domain that are connected by a linker of approximately 25 amino acids (Figure 6). The asymmetric unit contains five CheB_Pec chains that can be closely superimposed onto each other, resulting in C α root mean square deviation (RMSD) values of 0.23 to 0.61 Å (Figure S5 and Table S2). These five chains can also be closely aligned with the structure of *S. enterica* sv. Typhimurium CheB [20,23], as evidenced by RMSD values between 1.21 to 1.47 Å (Table S2), as well as onto the receiver and catalytic domains of *Thermotoga maritima* CheB (PDB ID: 3t8y and 3sft). In each CheB_Pec chain, there was a gap due to a lacking electron density that, depending on the chain, extended from amino acids 137 to 144–150. Lacking electron density is generally attributed to the corresponding protein segment being disordered. The gap was flanked by a segment with high B-factor values, indicative of significant protein flexibility in this region [54] (Figure S6).

There was a very satisfactory overall structural alignment of the CheB structures from *S. enterica* sv. Typhimurium and *P. atrosepticum* SCRI1043 (Figure 7A). Major deviations in this alignment showed a small region in the receiver domain, as well as in both flanking regions of the gap (Figure S7). The segment in the *S. enterica* sv. Typhimurium structure corresponding to the gap in CheB_Pec was characterized by low mean B-factors, namely 20 ± 7 for chain A and 30 ± 9 for chain B, indicative of a well-ordered structure.

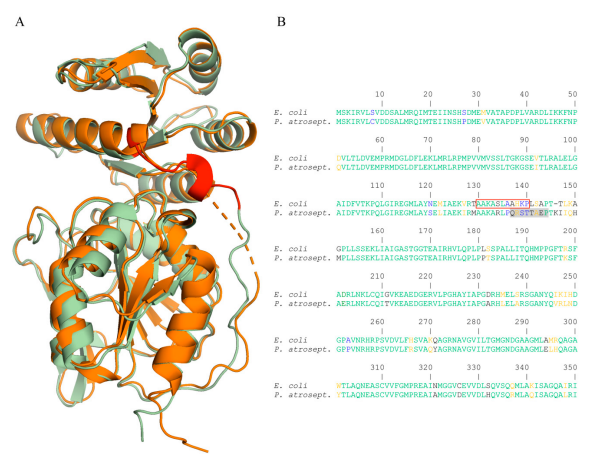


Figure 7. Structural and sequence features related to the capacity of CheB to recognize C-terminal pentapeptides. (**A**) Structural alignment of CheB from *P. atrosepticum* (orange) and *S. enterica* sv. Typhimurium (green, PDB ID: 1A2O). The amino acid segment identified as being the pentapeptide-binding site [16] is shown in red. (**B**) Sequence alignment of *Escherichia coli* K-12 and *P. atrosepticum* SCRI1043 CheB. The amino acids that form the pentapeptide-binding site in the *E. coli* enzyme are boxed in red. The gap in the CheB_Pec structure is shaded in grey. The alignment was done using the CLUSTALW algorithm of the NPS@ software [55]. The Gonnet protein weight matrix was used; gap opening and gap extension penalties were 10.0 and 0.1, respectively. Residues in green are identical, orange highly similar, blue weakly similar, and black dissimilar.

2.6. The Region Corresponding to the Pentapeptide-Binding Site in E. coli CheB Is Disordered in *P. atrosepticum CheB*

Having provided evidence that CheB_Pec is a correctly folded protein that resembles closely the *S. enterica* sv. Typhimurium structure, the question as to why it does not bind pentapeptides remained. The answer to this question may be related to studies that have identified the pentapeptide-binding site at *E. coli* CheB. This binding site was found to comprise amino acids 130 to 140 (colored in red in Figure 7A,B) and is located on the C-terminal extension of the REC domain and N-terminal part of the linker [16]. The inspection of the sequence alignment of CheB from *E. coli* and *P. atrosepticum* SCRI1043 revealed a high degree of sequence divergence in the pentapeptide-binding area (Figure 7B).

Importantly, a large part of the *E. coli* CheB pentapeptide-binding site overlaps with the gap observed in the CheB_Pec structure (shaded in grey in Figure 7B). We therefore hypothesize that the structural disorder of CheB_Pec in the region homologous to the pentapeptide-binding site in *E. coli* may be related to the failure to bind pentapeptides. In the case of CheR, distinct sequence features were identified for the pentapeptide-dependent and independent forms [8,13]. In contrast, the sequence alignment of pentapeptide-dependent and independent CheB did not reveal any obvious conserved sequence features.

3. Discussion

Many bacteria contain multiple paralogs of signaling proteins that form part of different chemosensory pathways [1]. A central question is whether or to what degree there is a specificity of interactions between the different homologs of signaling proteins and chemoreceptors. Furthermore, bacteria contain frequently a significant number of chemoreceptors, of which some possess C-terminal pentapeptides that are generally considered additional binding sites for CheR and CheB [9]. P. aeruginosa has four chemosensory pathways and a single chemoreceptor that contains a C-terminal pentapeptide. We showed previously that exclusively CheR₂ but not any of the remaining three CheR homologs of *P. aeruginosa* binds to the McpB pentapeptide [13]. Here, we show that the same holds for the four CheB homologs of *P. aeruginosa*, since CheB₂ was the only homolog that interacted with the McpB-derived pentapeptide. The data thus show that, exclusively, the CheR and CheB homologs encoded by the Che2 gene cluster (Figure 1) bind to McpB, the only receptor predicted to stimulate the Che₂ pathway [33]. This pathway is essential for the full virulence of *P. aeruginosa* [39,56], but its precise function still needs to be determined [37]. McpB and Che₂ pathway homologs are widespread in pathogenic and nonpathogenic γ -Proteobacteria [57], suggesting a function that is not exclusively associated with virulence. Interestingly, this pentapeptide is present in most McpB homologs, and it was concluded that this motif represents a fundamental feature of the McpB-like family [57]. We propose that a major reason for the pentapeptide conservation is its capacity to bind the pathway-specific CheB₂ and CheR₂ homologs [57], corresponding to a mechanism permitting pathway isolation. The pathway isolation of two-component systems has been extensively studied, particularly in the Laub laboratory [58], but the corresponding knowledge for chemosensory pathways is scarce. The data available on P. aeruginosa indicate that these four pathways are isolated [33], and the findings of our study may represent one of the corresponding mechanisms. Genome analyses of bacteria with pentapeptide-containing chemoreceptors showed that strains containing a single pentapeptide-containing chemoreceptor are the most abundant (approx. 2500 genomes) [9]. Future research will show whether the corresponding pentapeptides exert a similar function in these species.

The dissociation constant for the binding of CheB₂ to the McpB pentapeptide (93 μ M) is well below the corresponding value determined for the CheR₂-pentapeptide interaction of 0.52 μ M [13]. However, the values for the CheR and CheB binding to the *P. aeruginosa* pentapeptide GWETF are in the same range as those reported for the *E. coli* CheB and CheR binding to the NWETF pentapeptide, namely K_D values of 2 μ M [10] and 130 to 160 μ M for the CheR [17] and CheB, respectively. This may suggest that the much lower pentapeptide affinity of CheB as compared to CheR may be a more general feature. Studies have so far shown that there are pentapeptide-dependent and pentapeptide-independent CheR methyltransferases [8,13,14,19]. Sequence and structural features in the CheR β -subdomain, responsible for pentapeptide binding, were identified that account for the capacity or incapacity to bind pentapeptides [8,13]. We show here that, in the analogy to CheR, CheB methylesterases also form pentapeptide-dependent and independent subfamilies. The reason for the failure of CheB_Pec to bind pentapeptides may be related to the structural disorder in the segment homologous to the pentapeptide-binding site in E. coli CheB. Bioinformatic studies are required to establish the evolutionary history of CheB proteins in Enterobacteria to assess which subfamily evolved first. However, in contrast to CheR, sequence analyses of this region in pentapeptide-dependent and independent CheBs did not permit to identify a feature that can be associated with the capacity to bind pentapeptides.

Chemotaxis is particularly relevant for the virulence of plant pathogens [59]. This is also reflected at the genome level, since more than 90% of plant pathogens, compared to 50% for the bacterial average, possess chemosensory pathways [59]. In addition, the average number of chemoreceptors in plant pathogens, 33, is well superior to the bacterial average of 14 [59,60]. In contrast to the relevance of chemotaxis in plant pathogens, there is little information on the corresponding molecular mechanisms. *P. atrosepticum* is among the top 10 plant pathogens [42] and a suitable model to study chemosensory signaling in a plant pathogen. The chemoreceptor repertoire of the strain SCRI1043 shows a number of unusual features: (1) Remarkably, 67% of its chemoreceptors possess a 4HB LBD, which is well above the bacterial average of approx. 31% [61]. This chemoreceptor family is characterized by its versatility, as reflected in the broad range of ligands recognized (i.e., amino acids, boric acid, inorganic phosphate, aromatic acids, citrate, etc.); its capacity to recognize ligands with high [62,63] and low specificity [29,64]; and its ability to bind ligands directly [51] or via ligand-binding proteins [65]. (2) Apart from the three Are-like receptors that possess cytosolic LBDs (Figure 4), there are no cytosolic chemoreceptors, suggesting that, primarily, extracellular signals are sensed. (3) The abundance of chemoreceptors with a C-terminal pentapeptide. More than 50% of P. atrosepticum SCRI1043 chemoreceptors contain pentapeptides, a number that is well above the bacterial average of 10% [9], suggesting that pentapeptide function is important for signaling. However, CheB function does not require pentapeptide binding, and future studies are necessary to elucidate to what degree CheR function requires pentapeptide recognition.

4. Materials and Methods

4.1. Bacterial Strains and Growth Conditions

Bacterial strains used in this study are listed in Table 1. *P. atrosepticum* SCRI1043 and its derivative strains were routinely grown at 30 °C in Luria broth (5-g/L yeast extract, 10-g/L bacto tryptone, and 5-g/L NaCl) or minimal medium (0.41-mM MgSO₄, 7.56-mM (NH₄)₂SO₄, 40-mM K₂HPO₄, and 15-mM KH₂PO₄) supplemented with 0.2% (w/v) glucose as the carbon source. *E. coli* strains were grown at 37 °C in LB. *E. coli* DH5 α was used as a host for gene cloning. Media for the propagation of *E. coli* β2163 were supplemented with 300-mM 2,6-diaminopimelic acid. When appropriate, antibiotics were used at the following final concentrations (in μ g mL⁻¹): kanamycin, 50, tetracycline, 10, streptomycin, 50, and ampicillin, 100. Sucrose was added to a final concentration of 10% (w/v) when required to select derivatives that underwent a second crossover event during marker exchange mutagenesis.

Strains and Plasmids	Genotype or Relevant Characteristics ^a	Reference		
Strains				
Escherichia coli BL21(DE3)	F^- ompT gal dcm lon hsd $S_B(r_B^-m_B^-)$ λ(DE3 (lacI lacUV5-T7p07 ind1 sam7 nin5)) (malB ⁺) _{K-12} (λ ^S)	[66]		
E. coli BL21-AI	F- ompT hsdS _B ($r_B^-m_B^-$) gal dcm araB::T7RNAP-tetA	Invitrogen		
E. coli DH5α	F^- endA1 glnV44 thi-1 recA1 relA1 gyrA96 deoR nupG purB20 φ80dlacZΔM15 Δ(lacZYA-argF)U169, hsdR17($r_K^-m_K^+$), λ^-	[67]		
E. coli CC118λpir	araD Δ(ara, leu) ΔlacZ74 phoA20 galK thi-1 rspE rpoB argE recA1 λpir	[68]		
E. coli β2163	F- RP4-2-Tc::Mu Δ <i>dapA</i> ::(<i>erm-pir</i>); Km ^R , Em ^R	[69]		
Pectobacterium atrosepticum SCRI1043	Wild type strain	[32]		
P. atrosepticum SCRI1043 ∆cheB	SCRI1043 in-frame deletion mutant of cheB	This study		
P. atrosepticum SCRI1043 ∆cheA	SCRI1043 deletion mutant of <i>cheA</i> ; Km ^R	This study		

Table 1. Strains and plasmids used in this study.

Strains and Plasmids	Genotype or Relevant Characteristics ^a	Reference			
Plasmids					
pET28b(+)	Protein expression plasmid; Km ^R	Merckmillipore (Kenilworth, NJ, USA			
pET28b-CheB ₁	Km ^R ; pET28b(+) derivative containing <i>P. aeruginosa cheB1</i> (PA1459)	This study			
pET28b-CheB ₂	Km ^R ; pET28b(+) derivative containing <i>P. aeruginosa cheB2</i> (PA0173)	This study			
pET28b-CheB ₂ D55E	Km ^R ; pET28b(+) derivative containing <i>P. aeruginosa cheB2</i> D55E mutant				
pET28b-CheB ₃	Km ^R ; pET28b(+) derivative containing <i>P. aeruginosa cheB3</i> (PA3703)	This study			
pET28b-CheB ₄	Km ^R ; pET28b(+) derivative containing <i>P. aeruginosa cheB4</i> (PA0414)	This study			
pET28b-CheB_Pec	Km ^R ; pET28b(+) derivative containing <i>P. atrosepticum cheB</i> (ECA1693)	This study			
pUC18Not	Ap ^R ; identical to pUC18 but with two NotI sites flanking pUC18 polylinker	[68]			
pUC18Not_∆cheB	Ap ^R ; 1.5-kb PCR product containing a 954 bp in frame deletion of <i>cheB</i> (ECA1693) of SCRI1043 inserted into the SphI/NdeI sites of pUC18Not	This study			
pUC18Not_∆cheA	Ap ^R ; 1.5-kb PCR product containing a 1561 bp deletion of <i>cheA</i> (ECA1689) of SCRI1043 inserted into the EcoRI/HindIII sites of pUC18Not	This study			
p34S-Km3	Km ^R , Ap ^R ; <i>Km3</i> antibiotic cassette	[70]			
pUC18Not_AcheA-km3	Ap ^R , Km ^R ; 0.96-kb BamHI fragment containing <i>km3</i> cassette of p34S-Km3 was inserted into BamHI site of Δ <i>cheA</i> in pUC18Not_ΔcheA	This study			
pKNG101	Sm ^R ; oriR6K mob sacBR	[71]			
pKNG101_∆cheB	Sm ^R , Km ^R ; 1.5 kb NotI fragment of pUC18Not_ΔcheB was cloned at the same site in pKNG101	This study			
pKNG101_∆cheA-km3	Sm ^R , Km ^R ; 2.4-kb NotI fragment of pUC18Not_∆cheA-Km3 was cloned at the same site in pKNG101	This study			

Table 1. Cont.

^a Ap, ampicillin, Em, erythromycin, Km, kanamycin, Sm, streptomycin, and Tc, tetracycline.

4.2. Generation of Protein Expression Plasmids

Plasmids used in this study are listed in Table 1. Genes encoding *P. aeruginosa* PAO1 CheB₁ (PA1459), CheB₂ (PA0173), CheB₃ (PA3703), CheB₄ (PA0414), and *P. atrosepticum* SCRI1043 CheB (ECA1693) were amplified by PCR using the oligonucleotides indicated in Table S3 and genomic DNA as the template. The latter PCR product was digested with NheI and SaII, whereas the remaining products were digested with NdeI and BamHI. The resulting DNA fragments were cloned into pET28b(+) linearized with the respective endonucleases. The generated plasmids were verified by DNA sequencing.

4.3. Site-Directed Mutagenesis

The Hemsley method [72] was used to generate the CheB₂ D55E mutant. The pair of overlapping mutagenic primers pET28_CheB₂_D55E_f and pET28_CheB₂_D55E_r (Table S3) were used to amplify the entire plasmid pET28b-CheB₂ using the *Pfu* Turbo DNA polymerase (Agilent Technologies, Santa Clara, CA, USA). Following the elimination of template DNA by a digestion with DpnI, the resulting mixture was transformed into *E. coli* DH5 α , and colonies were selected on LB agar plates supplemented with kanamycin. Plasmid inserts and flanking regions were sequenced.

4.4. Protein Overexpression and Purification

Plasmids for the overexpression of the wt and mutant CheB proteins of PAO1 were transformed into E. coli BL21 (DE3). Alternatively, pET28b-CheB_Pec was transformed into E. coli BL21-AITM. The resulting strains were grown under continuous stirring (200 rpm) at 30 °C in 2-L Erlenmeyer flasks containing 500 mL of LB medium supplemented with 50-µg/mL kanamycin. At an OD_{660nm} of 0.6, protein expression was induced by the addition of 0.1-mM isopropyl β-D-1-thiogalactopyranoside. In addition, L-arabinose was added to E. coli BL21-AITM cultures for a final concentration of 0.2% (w/v). Growth was continued at 16 °C overnight prior to cell harvest by centrifugation at 10,000× g for 30 min. Cell pellets for the purification of CheB proteins of *P. aeruginosa* were resuspended in buffer A (20-mM Tris/HCl, 500-mM NaCl, 5% (v/v) glycerol, 10-mM imidazole, 0.1-mM EDTA, and 5-mM β -mercaptoethanol, pH 8.0), whereas pellets for the purification of CheB_Pec were resuspended in buffer B (20-mM Tris/HCl, 150-mM NaCl, 10-mM imidazole, 0.1-mM EDTA, 10% (v/v) glycerol, and 10-mM β -mercaptoethanol, pH 8.0). Subsequently, cells were broken by French press treatment at 62.5 lb/in². After centrifugation at $20,000 \times g$ for 30 min, supernatants were loaded onto 5-mL HisTrap HP columns (Amersham Biosciences, Little Chalfont, UK) equilibrated with the corresponding buffer A or B and eluted with an imidazole gradient of 40–500 mM in the corresponding buffer. When necessary, the His-tag was removed by treatment with bovine thrombin (Sigma-Aldrich, Buchs, Switzerland) at 2 U/mL and 16 °C for 2 h. For crystallization, CheB_Pec was dialyzed into 5-mM Tris/HCl, 5-mM Pipes, 5-mM Mes, 2-mM dithiothreitol, 150-mM NaCl, and 10% (v/v) glycerol, pH 7.4 and purified by size-exclusion chromatography using a HiPrepTM 26/60 SephacrylTM S200 HR gel filtration column (GE Healthcare, Chicago, IL, USA) at a flow rate of 1 mL/min. All proteins were purified at 4 °C. Purified proteins were dialyzed overnight into the corresponding analysis buffers for immediate analysis.

4.5. Isothermal Titration Calorimetry (ITC)

All experiments were conducted on a VP microcalorimeter (Microcal, Amherst, MA, USA) at 25 °C. For the analysis of the CheB homologs of *P. aeruginosa*, 15–40 μ M of proteins (dialyzed into 5-mM Tris/HCl, 5-mM Pipes, and 5-mM Mes, pH 7.0) were placed into the sample cell and titrated with 1–7-mM solution of the GWEEF peptide (synthesized by Biomedal S.L., Seville, Spain). For the analysis of CheB_Pec, the protein was dialyzed into 5-mM Tris/HCl, 5-mM PIPES, 5-mM MES, 10% (*v*/*v*) glycerol, 2-mM dithiothreitol, 150-mM NaCl, and 0.1-mM EDTA, pH 7.4, adjusted to 15–50 μ M and titrated with 4.8–14.4- μ L aliquots of 1–5-mM peptide solutions (synthesized by GenScript[®], Piscataway, NJ, USA). All ligand solutions were prepared in dialysis buffer immediately before use. The mean enthalpies measured from the injection of the peptide into the buffer were subtracted from the raw titration data prior to data analysis with the "one binding site model" of the MicroCal version of ORIGIN.

4.6. Chemoreceptor Sequence Analysis

Sequences were retrieved from the MIST 3.0 database [73], transmembrane regions identified using DAS [74], and ligand-binding domains annotated according to Pfam [75]. Pentapeptides at chemoreceptors were identified as reported in [8]. These peptides matched the xZxxZ motif (where Z represents either F, W, or Y) and were separated from the chemoreceptor signaling domain by a linker sequence of at least 10 amino acids.

4.7. Derivatization of CheB_Pec by Beryllium Fluoride

A modified version of the protocol described in [45] was employed. Briefly, 0.1-M BeSO₄, 10-mM NaF, and 10-mM MgCl₂ (all final concentrations) were added to CheB_Pec dialyzed into 5-mM Tris/HCl, 5-mM PIPES, 5-mM MES, 10% (v/v) glycerol, 2-mM dithiothreitol, 150-mM NaCl, and 0.1-mM EDTA, pH 7.4. The resulting mixture was incubated at 25 °C for 3 h.

Chromosomal mutants of SCRI1043 were constructed by homologous recombination using a derivative plasmid of the suicide vector pKNG101. These plasmids were confirmed by DNA sequencing and carried deletion mutant alleles for the replacement of wild-type genes in the chromosome. In all cases, plasmids for mutagenesis were transferred to *P. atrosepticum* SRCI1043 by biparental conjugation using *E. coli* β 2163. The plasmids for the construction of the deletion mutants were generated by amplifying the up- and downstream flanking regions of the gene to be mutated. The resulting PCR products were digested with the enzymes specified in Table 1 and ligated in a three-way ligation into pUC18Not, producing plasmids pUC18Not_ Δ cheB and pUC18Not_ Δ cheA. Subsequently, the kanamycin resistance cassette Km3 from the plasmid p34S-km3 was inserted into the BamHI of pUC18Not_ Δ cheA, resulting in plasmid pUC18Not_ Δ cheA-Km3. The Δ cheB and Δ cheA-km3 deletion constructs were then subcloned into the marker exchange vector pKNG101 using NotI. Mutant strains defective in *cheA* and *cheB* were generated using plasmids pKNG101_ Δ cheA, respectively.

4.9. Quantitative Capillarity Chemotaxis Assays

Overnight cultures of *P. atrosepticum* were grown at 30 °C in minimal medium. At an OD₆₆₀ of 0.35–0.4, the cultures were washed twice with chemotaxis buffer (50-mM K₂HPO₄/KH₂PO₄, 20- μ M EDTA, and 0.05% (*v*/*v*) glycerol, pH 7.0) and diluted to an OD₆₆₀ of 0.1 in the same buffer. Subsequently, 230 μ L of the resulting bacterial suspension were place into the wells of 96-well plates. One-microliter capillary tubes (P1424, Microcaps; Drummond Scientific, Broomall, PA, USA) were heat-sealed at one end and filled with either the chemotaxis buffer (negative control) or chemotaxis buffer containing casamino acids. The capillaries were immersed into the bacterial suspensions at their open ends. After 30 min at room temperature, the capillaries were removed from the bacterial suspensions, rinsed with sterile water, and the content expelled into 1 mL of minimal medium salts. Serial dilutions were plated onto minimal medium supplemented with 15-mM glucose as the carbon source. The number of colony-forming units was determined after incubation at 30 °C for 36 h. In all cases, data were corrected with the number of cells that swam into buffer-containing capillaries.

4.10. Crystallization and Resolution of the Three-Dimensional Structure of CheB_Pec

CheB_Pec was dialyzed into 5-mM Tris/HCl, 150-mM NaCl, 2-mM dithiothreitol, and 10% (v/v) glycerol, pH 7.4 and concentrated to 2.35 mg/mL using 3-kDa cut-off centricon concentrators (Merckmillipore (Kenilworth, NJ, USA). Crystallization conditions were screened using the capillary counter-diffusion technique, and a set of conditions prepared ad hoc that were reported in [76]. CheB_Pec was loaded into capillaries of 0.2-mm inner diameters, and crystals appeared in several conditions, namely C4 (1.25-M sodium citrate and 0.1-M Na/HEPES, pH 7.5); C5 (1.7-M ammonium sulphate, 3.5% (*w*/*v*) PEG 400, and 0.1-M Na/HEPES, pH 7.5); and C7 (2.0-M ammonium sulphate and 0.1-M Tris/HCl, pH 8.0 and 8.5). Crystals were extracted from the capillary and equilibrated in the mother solution supplemented with 15% (v/v) glycerol. Individual crystals were placed into LithoLoops (Molecular Dimensions, Portobello, Sheffield, UK), flash-frozen in liquid nitrogen and stored until data collection at the Xaloc beamline of the Spanish Synchrotron Radiation Source Alba. Several full datasets were obtained and automatically indexed, reduced, and scaled using the default data processing with EDNA [77] within the MXCuBE [78] data collection interface. The automatically determined space groups, I422 or F422, did not permit phasing using molecular replacement. Data were manually inspected, indexed, and merged with iMOSFLM [79] in space group I4 and scaled and reduced using Aimless [80] of the CCP4 program suite [81]. The structure was determined by molecular replacement in Phaser [82] using a homology model [83] based on the CheB structure of S. enterica sv. Typhimurium (PDB ID 1a20) and after lowering the symmetry of the space group to I4. Five polypeptide chains were placed into the unit cell giving rise to a Matthews coefficient [84] of 2.85 and

a water content of 57%. Refinement was done by phenix.refine of the PHENIX suite [85], including titration-libration-screw (TLS) parameterization [86]. Cycles of manual building and inspection were done in Coot [87]. The final refined model was verified with Procheck [88], Molprobity [89], and PDBe validation servers [90]. Table S4 summarizes the crystallographic data statistics and final model characteristics. The coordinates and the experimental structure factors for the CheB methylesterase from *P. atrosepticum* SCRI1043 were deposited at the Protein Data Bank with ID 6ymz.

Supplementary Materials: Supplementary Materials can be found at http://www.mdpi.com/1422-0067/21/22/ 8459/s1. Figure S1. Sequence alignment of CheB from *E. coli* K-12 substrain MG1655 and *S. enterica* serovar Typhimurium str. LT-2. Figure S2. Microcalorimetric binding studies of the CheB₂ D55E mutant with the McpB-derived pentapeptide GWEEF. Figure S3. Alignment of the linker sequences of the 19 chemoreceptors from *Pectobacterium atrosepticum* containing terminal pentapeptides. Figure S4. Secondary structure prediction of the C-terminal segment of pentapeptide containing chemoreceptors of *P. atrosepticum*. Figure S5. Analysis of the structure of *P. atrosepticum* CheB. Structural superimposition of chain A onto the remaining 4 chains of the asymmetric unit. Figure S6. B-factors of C α atoms of *P. atrosepticum* chain B. Figure S7. Structural superimposition of chain A of *P. atrosepticum* CheB with CheB of *S. enterica* sv. Typhimurium (pdb ID 1A2O). Table S1. *P. atrosepticum* SCRI1043 chemoreceptors with a C-terminal pentapeptide. Table S2. Structural alignment of chains A to E of *P. atrosepticum* CheB amongst each other and with related structures. Table S3. Oligonucleotides used in this study. Table S4. Data collection and refinement statistics of the 3D structure of *P. atrosepticum* CheB.

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Abbreviations

CheB_Pec	CheB of <i>P. atrosepticum</i> SCRI1043
Chp	chemosensory pili
4HB	four helix bundle
HBM	helical bimodular
ITC	isothermal titration calorimetry
LBD	ligand binding domain
Wsp	wrinkly spreader phenotype

References

- 1. Wuichet, K.; Zhulin, I.B. Origins and diversification of a complex signal transduction system in prokaryotes. *Sci. Signal.* **2010**, *3*, ra50. [CrossRef] [PubMed]
- Hickman, J.W.; Tifrea, D.F.; Harwood, C.S. A chemosensory system that regulates biofilm formation through modulation of cyclic diguanylate levels. *Proc. Natl. Acad. Sci. USA* 2005, 102, 14422–14427. [CrossRef] [PubMed]
- Whitchurch, C.B.; Leech, A.J.; Young, M.D.; Kennedy, D.; Sargent, J.L.; Bertrand, J.J.; Semmler, A.B.; Mellick, A.S.; Martin, P.R.; Alm, R.A.; et al. Characterization of a complex chemosensory signal transduction system which controls twitching motility in *Pseudomonas aeruginosa*. *Mol. Microbiol.* 2004, 52, 873–893. [CrossRef] [PubMed]
- 4. Parkinson, J.S.; Hazelbauer, G.L.; Falke, J.J. Signaling and sensory adaptation in *Escherichia coli* chemoreceptors: 2015 update. *Trends Microbiol.* **2015**, 23, 257–266. [CrossRef] [PubMed]
- 5. Bi, S.; Sourjik, V. Stimulus sensing and signal processing in bacterial chemotaxis. *Curr. Opin. Microbiol.* **2018**, 45, 22–29. [CrossRef] [PubMed]

- 6. Li, M.; Hazelbauer, G.L. Methyltransferase CheR binds to its chemoreceptor substrates independent of their signaling conformation yet modifies them differentially. *Protein Sci.* **2020**, *29*, 443–454. [CrossRef] [PubMed]
- 7. Bartelli, N.L.; Hazelbauer, G.L. Direct evidence that the carboxyl-terminal sequence of a bacterial chemoreceptor is an unstructured linker and enzyme tether. *Protein Sci.* **2011**, *20*, 1856–1866. [CrossRef]
- 8. Perez, E.; Stock, A.M. Characterization of the *Thermotoga maritima* chemotaxis methylation system that lacks pentapeptide-dependent methyltransferase CheR:MCP tethering. *Mol. Microbiol.* **2007**, *63*, 363–378. [CrossRef]
- 9. Ortega, A.; Krell, T. Chemoreceptors with C-terminal pentapeptides for CheR and CheB binding are abundant in bacteria that maintain host interactions. *Comput. Struct. Biotechnol. J.* **2020**, *18*, 1947–1955. [CrossRef]
- 10. Wu, J.; Li, J.; Li, G.; Long, D.G.; Weis, R.M. The receptor binding site for the methyltransferase of bacterial chemotaxis is distinct from the sites of methylation. *Biochemistry* **1996**, *35*, 4984–4993. [CrossRef]
- 11. Le Moual, H.; Quang, T.; Koshland, D.E., Jr. Methylation of the *Escherichia coli* chemotaxis receptors: Intraand interdimer mechanisms. *Biochemistry* **1997**, *36*, 13441–13448. [CrossRef] [PubMed]
- Li, M.; Hazelbauer, G.L. Adaptational assistance in clusters of bacterial chemoreceptors. *Mol. Microbiol.* 2005, 56, 1617–1626. [CrossRef] [PubMed]
- Garcia-Fontana, C.; Lugo, A.C.; Krell, T. Specificity of the CheR2 methyltransferase in *Pseudomonas aeruginosa* is directed by a C-terminal pentapeptide in the McpB chemoreceptor. *Sci. Signal.* 2014, 7, ra34. [CrossRef] [PubMed]
- Djordjevic, S.; Stock, A.M. Chemotaxis receptor recognition by protein methyltransferase CheR. *Nat. Struct. Biol.* 1998, 5, 446–450. [CrossRef]
- 15. Barnakov, A.N.; Barnakova, L.A.; Hazelbauer, G.L. Efficient adaptational demethylation of chemoreceptors requires the same enzyme-docking site as efficient methylation. *Proc. Natl. Acad. Sci. USA* **1999**, *96*, 10667–10672. [CrossRef]
- Barnakov, A.N.; Barnakova, L.A.; Hazelbauer, G.L. Location of the receptor-interaction site on CheB, the methylesterase response regulator of bacterial chemotaxis. *J. Biol. Chem.* 2001, 276, 32984–32989. [CrossRef]
- 17. Barnakov, A.N.; Barnakova, L.A.; Hazelbauer, G.L. Allosteric enhancement of adaptational demethylation by a carboxyl-terminal sequence on chemoreceptors. *J. Biol. Chem.* **2002**, 277, 42151–42156. [CrossRef]
- 18. Li, M.; Hazelbauer, G.L. Cellular stoichiometry of the components of the chemotaxis signaling complex. *J. Bacteriol.* **2004**, *186*, 3687–3694. [CrossRef]
- Li, M.; Hazelbauer, G.L. The carboxyl-terminal linker is important for chemoreceptor function. *Mol. Microbiol.* 2006, 60, 469–479. [CrossRef]
- 20. Djordjevic, S.; Goudreau, P.N.; Xu, Q.; Stock, A.M.; West, A.H. Structural basis for methylesterase CheB regulation by a phosphorylation-activated domain. *Proc. Natl. Acad. Sci. USA* **1998**, *95*, 1381–1386. [CrossRef]
- 21. Anand, G.S.; Goudreau, P.N.; Stock, A.M. Activation of methylesterase CheB: Evidence of a dual role for the regulatory domain. *Biochemistry* **1998**, *37*, 14038–14047. [CrossRef] [PubMed]
- 22. Anand, G.S.; Stock, A.M. Kinetic basis for the stimulatory effect of phosphorylation on the methylesterase activity of CheB. *Biochemistry* **2002**, *41*, 6752–6760. [CrossRef] [PubMed]
- 23. West, A.H.; Martinez-Hackert, E.; Stock, A.M. Crystal structure of the catalytic domain of the chemotaxis receptor methylesterase, CheB. J. Mol. Biol. 1995, 250, 276–290. [CrossRef]
- 24. Okumura, H.; Nishiyama, S.; Sasaki, A.; Homma, M.; Kawagishi, I. Chemotactic adaptation is altered by changes in the carboxy-terminal sequence conserved among the major methyl-accepting chemoreceptors. *J. Bacteriol.* **1998**, *180*, 1862–1868. [CrossRef] [PubMed]
- 25. Li, J.; Li, G.; Weis, R.M. The serine chemoreceptor from *Escherichia coli* is methylated through an inter-dimer process. *Biochemistry* **1997**, *36*, 11851–11857. [CrossRef]
- 26. Matilla, M.A.; Krell, T. Chemoreceptor-based signal sensing. Curr. Opin. Biotechnol. 2017, 45, 8–14. [CrossRef]
- 27. Corral-Lugo, A.; Matilla, M.A.; Martin-Mora, D.; Jimenez, S.H.; Torres, N.M.; Kato, J.; Hida, A.; Oku, S.; Conejero-Muriel, M.; Gavira, J.A.; et al. High-Affinity Chemotaxis to Histamine Mediated by the TlpQ Chemoreceptor of the Human Pathogen *Pseudomonas aeruginosa. mBio* **2018**, *9*, e01894-18. [CrossRef]
- 28. Martin-Mora, D.; Ortega, A.; Matilla, M.A.; Martinez-Rodriguez, S.; Gavira, J.A.; Krell, T. The Molecular Mechanism of Nitrate Chemotaxis via Direct Ligand Binding to the PilJ Domain of McpN. *mBio* **2019**, *10*, e02334-18. [CrossRef]

- 29. Luu, R.A.; Kootstra, J.D.; Nesteryuk, V.; Brunton, C.N.; Parales, J.V.; Ditty, J.L.; Parales, R.E. Integration of chemotaxis, transport and catabolism in *Pseudomonas putida* and identification of the aromatic acid chemoreceptor PcaY. *Mol. Microbiol.* **2015**, *96*, 134–147. [CrossRef]
- 30. Compton, K.K.; Hildreth, S.B.; Helm, R.F.; Scharf, B.E. *Sinorhizobium meliloti* Chemoreceptor McpV Senses Short-Chain Carboxylates via Direct Binding. *J. Bacteriol.* **2018**, 200, e00519-18. [CrossRef]
- Stover, C.K.; Pham, X.Q.; Erwin, A.L.; Mizoguchi, S.D.; Warrener, P.; Hickey, M.J.; Brinkman, F.S.; Hufnagle, W.O.; Kowalik, D.J.; Lagrou, M.; et al. Complete genome sequence of *Pseudomonas aeruginosa* PAO1, an opportunistic pathogen. *Nature* 2000, 406, 959–964. [CrossRef]
- Bell, K.S.; Sebaihia, M.; Pritchard, L.; Holden, M.T.; Hyman, L.J.; Holeva, M.C.; Thomson, N.R.; Bentley, S.D.; Churcher, L.J.; Mungall, K.; et al. Genome sequence of the enterobacterial phytopathogen *Erwinia carotovora* subsp. *atroseptica* and characterization of virulence factors. *Proc. Natl. Acad. Sci. USA* 2004, 101, 11105–11110. [CrossRef] [PubMed]
- Ortega, D.R.; Fleetwood, A.D.; Krell, T.; Harwood, C.S.; Jensen, G.J.; Zhulin, I.B. Assigning chemoreceptors to chemosensory pathways in *Pseudomonas aeruginosa*. *Proc. Natl. Acad. Sci. USA* 2017, 114, 12809–12814. [CrossRef] [PubMed]
- 34. Kato, J.; Nakamura, T.; Kuroda, A.; Ohtake, H. Cloning and characterization of chemotaxis genes in *Pseudomonas aeruginosa. Biosci. Biotechnol. Biochem.* **1999**, *63*, 155–161. [CrossRef] [PubMed]
- Masduki, A.; Nakamura, J.; Ohga, T.; Umezaki, R.; Kato, J.; Ohtake, H. Isolation and characterization of chemotaxis mutants and genes of *Pseudomonas aeruginosa*. J. Bacteriol. 1995, 177, 948–952. [CrossRef] [PubMed]
- Fulcher, N.B.; Holliday, P.M.; Klem, E.; Cann, M.J.; Wolfgang, M.C. The *Pseudomonas aeruginosa* Chp chemosensory system regulates intracellular cAMP levels by modulating adenylate cyclase activity. *Mol. Microbiol.* 2010, *76*, 889–904. [CrossRef] [PubMed]
- 37. Orillard, E.; Watts, K.J. Deciphering the Che2 chemosensory pathway and the roles of individual Che2 proteins from *Pseudomonas aeruginosa*. *Mol. Microbiol.* **2020**. [CrossRef]
- 38. Ferrandez, A.; Hawkins, A.C.; Summerfield, D.T.; Harwood, C.S. Cluster II che genes from *Pseudomonas aeruginosa* are required for an optimal chemotactic response. *J. Bacteriol.* **2002**, *184*, 4374–4383. [CrossRef]
- Garvis, S.; Munder, A.; Ball, G.; de Bentzmann, S.; Wiehlmann, L.; Ewbank, J.J.; Tummler, B.; Filloux, A. *Caenorhabditis elegans* semi-automated liquid screen reveals a specialized role for the chemotaxis gene *cheB2* in *Pseudomonas aeruginosa* virulence. *PLoS Pathog.* 2009, *5*, e1000540. [CrossRef]
- Starkey, M.; Hickman, J.H.; Ma, L.; Zhang, N.; De Long, S.; Hinz, A.; Palacios, S.; Manoil, C.; Kirisits, M.J.; Starner, T.D.; et al. *Pseudomonas aeruginosa* rugose small-colony variants have adaptations that likely promote persistence in the cystic fibrosis lung. *J. Bacteriol.* 2009, 191, 3492–3503. [CrossRef]
- 41. Caiazza, N.C.; Merritt, J.H.; Brothers, K.M.; O'Toole, G.A. Inverse regulation of biofilm formation and swarming motility by *Pseudomonas aeruginosa* PA14. *J. Bacteriol.* **2007**, *189*, 3603–3612. [CrossRef] [PubMed]
- Mansfield, J.; Genin, S.; Magori, S.; Citovsky, V.; Sriariyanum, M.; Ronald, P.; Dow, M.; Verdier, V.; Beer, S.V.; Machado, M.A.; et al. Top 10 plant pathogenic bacteria in molecular plant pathology. *Mol. Plant. Pathol.* 2012, 13, 614–629. [CrossRef] [PubMed]
- Toth, I.K.; Bell, K.S.; Holeva, M.C.; Birch, P.R. Soft rot erwiniae: From genes to genomes. *Mol. Plant. Pathol.* 2003, 4, 17–30. [CrossRef] [PubMed]
- 44. Porter, S.L.; Armitage, J.P. Phosphotransfer in *Rhodobacter sphaeroides* chemotaxis. *J. Mol. Biol.* **2002**, 324, 35–45. [CrossRef]
- 45. Guhaniyogi, J.; Robinson, V.L.; Stock, A.M. Crystal structures of beryllium fluoride-free and beryllium fluoride-bound CheY in complex with the conserved C-terminal peptide of CheZ reveal dual binding modes specific to CheY conformation. *J. Mol. Biol.* **2006**, *359*, 624–645. [CrossRef] [PubMed]
- Rapun-Araiz, B.; Haag, A.F.; De Cesare, V.; Gil, C.; Dorado-Morales, P.; Penades, J.R.; Lasa, I. Systematic Reconstruction of the Complete Two-Component Sensorial Network in *Staphylococcus aureus*. *mSystems* 2020. [CrossRef] [PubMed]
- 47. Collins, K.D.; Lacal, J.; Ottemann, K.M. Internal sense of direction: Sensing and signaling from cytoplasmic chemoreceptors. *Microbiol. Mol. Biol. Rev.* 2014, *78*, 672–684. [CrossRef]
- 48. Waterhouse, A.; Bertoni, M.; Bienert, S.; Studer, G.; Tauriello, G.; Gumienny, R.; Heer, F.T.; de Beer, T.A.P.; Rempfer, C.; Bordoli, L.; et al. SWISS-MODEL: Homology modelling of protein structures and complexes. *Nucleic Acids Res.* **2018**, *46*, W296–W303. [CrossRef]

- Taylor, B.L.; Zhulin, I.B.; Johnson, M.S. Aerotaxis and other energy-sensing behavior in bacteria. *Annu. Rev. Microbiol.* **1999**, 53, 103–128. [CrossRef]
- 50. Upadhyay, A.A.; Fleetwood, A.D.; Adebali, O.; Finn, R.D.; Zhulin, I.B. Cache Domains That are Homologous to, but Different from PAS Domains Comprise the Largest Superfamily of Extracellular Sensors in Prokaryotes. *PLoS Comput. Biol.* **2016**, *12*, e1004862. [CrossRef]
- 51. Matilla, M.A.; Mora, D.M.; Krell, T. The use of Isothermal Titration Calorimetry to unravel chemotactic signaling mechanisms. *Environ. Microbiol.* **2020**, *22*, 3005–3019. [CrossRef] [PubMed]
- 52. Cho, H.; Wang, W.; Kim, R.; Yokota, H.; Damo, S.; Kim, S.H.; Wemmer, D.; Kustu, S.; Yan, D. BeF(3)(-) acts as a phosphate analog in proteins phosphorylated on aspartate: Structure of a BeF(3)(-) complex with phosphoserine phosphatase. *Proc. Natl. Acad. Sci. USA* **2001**, *98*, 8525–8530. [CrossRef] [PubMed]
- 53. Laskowski, R.A.; Jablonska, J.; Pravda, L.; Varekova, R.S.; Thornton, J.M. PDBsum: Structural summaries of PDB entries. *Protein Sci.* 2018, 27, 129–134. [CrossRef] [PubMed]
- Sun, Z.; Liu, Q.; Qu, G.; Feng, Y.; Reetz, M.T. Utility of B-Factors in Protein Science: Interpreting Rigidity, Flexibility, and Internal Motion and Engineering Thermostability. *Chem. Rev.* 2019, 119, 1626–1665. [CrossRef] [PubMed]
- Combet, C.; Blanchet, C.; Geourjon, C.; Deleage, G. NPS@: Network protein sequence analysis. *Trends Biochem. Sci.* 2000, 25, 147–150. [CrossRef]
- Garcia-Fontana, C.; Vilchez, J.I.; Gonzalez-Requena, M.; Gonzalez-Lopez, J.; Krell, T.; Matilla, M.A.; Manzanera, M. The involvement of McpB chemoreceptor from *Pseudomonas aeruginosa* PAO1 in virulence. *Sci. Rep.* 2019, *9*, 13166. [CrossRef] [PubMed]
- Ortega, D.R.; Yang, W.; Subramanian, P.; Mann, P.; Kjaer, A.; Chen, S.; Watts, K.J.; Pirbadian, S.; Collins, D.A.; Kooger, R.; et al. Repurposing a chemosensory macromolecular machine. *Nat. Commun.* 2020, *11*, 2041. [CrossRef]
- 58. McClune, C.J.; Alvarez-Buylla, A.; Voigt, C.A.; Laub, M.T. Engineering orthogonal signalling pathways reveals the sparse occupancy of sequence space. *Nature* **2019**, 574, 702–706. [CrossRef]
- 59. Matilla, M.A.; Krell, T. The effect of bacterial chemotaxis on host infection and pathogenicity. *FEMS Microbiol. Rev.* **2018**, 42. [CrossRef]
- Lacal, J.; Garcia-Fontana, C.; Munoz-Martinez, F.; Ramos, J.L.; Krell, T. Sensing of environmental signals: Classification of chemoreceptors according to the size of their ligand binding regions. *Environ. Microbiol.* 2010, 12, 2873–2884. [CrossRef]
- Ortega, A.; Zhulin, I.B.; Krell, T. Sensory Repertoire of Bacterial Chemoreceptors. *Microbiol. Mol. Biol. Rev.* 2017, *81*, e00033-17. [CrossRef] [PubMed]
- Hida, A.; Oku, S.; Nakashimada, Y.; Tajima, T.; Kato, J. Identification of boric acid as a novel chemoattractant and elucidation of its chemoreceptor in *Ralstonia pseudosolanacearum* Ps29. *Sci. Rep.* 2017, 7, 8609. [CrossRef] [PubMed]
- 63. Rico-Jimenez, M.; Reyes-Darias, J.A.; Ortega, A.; Diez Pena, A.I.; Morel, B.; Krell, T. Two different mechanisms mediate chemotaxis to inorganic phosphate in *Pseudomonas aeruginosa*. *Sci. Rep.* **2016**, *6*, 28967. [CrossRef] [PubMed]
- 64. Huang, Z.; Ni, B.; Jiang, C.Y.; Wu, Y.F.; He, Y.Z.; Parales, R.E.; Liu, S.J. Direct sensing and signal transduction during bacterial chemotaxis toward aromatic compounds in *Comamonas testosteroni*. *Mol. Microbiol.* **2016**, 101, 224–237. [CrossRef] [PubMed]
- 65. Gardina, P.J.; Bormans, A.F.; Manson, M.D. A mechanism for simultaneous sensing of aspartate and maltose by the Tar chemoreceptor of *Escherichia coli*. *Mol. Microbiol*. **1998**, 29, 1147–1154. [CrossRef] [PubMed]
- 66. Jeong, H.; Barbe, V.; Lee, C.H.; Vallenet, D.; Yu, D.S.; Choi, S.H.; Couloux, A.; Lee, S.W.; Yoon, S.H.; Cattolico, L.; et al. Genome sequences of *Escherichia coli* B strains REL606 and BL21(DE3). *J. Mol. Biol.* **2009**, 394, 644–652. [CrossRef]
- 67. Woodcock, D.M.; Crowther, P.J.; Doherty, J.; Jefferson, S.; DeCruz, E.; Noyer-Weidner, M.; Smith, S.S.; Michael, M.Z.; Graham, M.W. Quantitative evaluation of *Escherichia coli* host strains for tolerance to cytosine methylation in plasmid and phage recombinants. *Nucleic Acids Res.* **1989**, *17*, 3469–3478. [CrossRef]
- Herrero, M.; de Lorenzo, V.; Timmis, K.N. Transposon vectors containing non-antibiotic resistance selection markers for cloning and stable chromosomal insertion of foreign genes in gram-negative bacteria. *J. Bacteriol.* 1990, 172, 6557–6567. [CrossRef]

- 69. Demarre, G.; Guerout, A.M.; Matsumoto-Mashimo, C.; Rowe-Magnus, D.A.; Marliere, P.; Mazel, D. A new family of mobilizable suicide plasmids based on broad host range R388 plasmid (IncW) and RP4 plasmid (IncPalpha) conjugative machineries and their cognate *Escherichia coli* host strains. *Res. Microbiol.* **2005**, 156, 245–255. [CrossRef]
- 70. Dennis, J.J.; Zylstra, G.J. Plasposons: Modular self-cloning minitransposon derivatives for rapid genetic analysis of gram-negative bacterial genomes. *Appl. Environ. Microbiol.* **1998**, *64*, 2710–2715. [CrossRef]
- 71. Kaniga, K.; Delor, I.; Cornelis, G.R. A wide-host-range suicide vector for improving reverse genetics in gram-negative bacteria: Inactivation of the blaA gene of *Yersinia enterocolitica*. *Gene* **1991**, *109*, 137–141. [CrossRef]
- 72. Hemsley, A.; Arnheim, N.; Toney, M.D.; Cortopassi, G.; Galas, D.J. A simple method for site-directed mutagenesis using the polymerase chain reaction. *Nucleic Acids Res.* **1989**, *17*, 6545–6551. [CrossRef] [PubMed]
- 73. Gumerov, V.M.; Ortega, D.R.; Adebali, O.; Ulrich, L.E.; Zhulin, I.B. MiST 3.0: An updated microbial signal transduction database with an emphasis on chemosensory systems. *Nucleic Acids Res.* **2019**, *48*, D459–D464. [CrossRef] [PubMed]
- 74. Cserzo, M.; Wallin, E.; Simon, I.; von Heijne, G.; Elofsson, A. Prediction of transmembrane alpha-helices in prokaryotic membrane proteins: The dense alignment surface method. *Protein Eng.* **1997**, *10*, 673–676. [CrossRef]
- 75. El-Gebali, S.; Mistry, J.; Bateman, A.; Eddy, S.R.; Luciani, A.; Potter, S.C.; Qureshi, M.; Richardson, L.J.; Salazar, G.A.; Smart, A.; et al. The Pfam protein families database in 2019. *Nucleic Acids Res.* **2019**, *47*, D427–D432. [CrossRef]
- 76. González-Ramírez, L.A.; Ruiz-Martínez, C.R.; Estremera-Andújar, R.A.; Nieves-Marrero, C.A.; García-Caballero, A.; Gavira, J.A.; López-Garriga, J.; García-Ruiz, J.M. Efficient Screening Methodology for Protein Crystallization Based on the Counter-Diffusion Technique. *Cryst. Growth Des.* 2017, *17*, 6780–6786. [CrossRef]
- 77. Incardona, M.-F.; Bourenkov, G.P.; Levik, K.; Pieritz, R.A.; Popov, A.N.; Svensson, O. EDNA: A framework for plugin-based applications applied to X-ray experiment online data analysis. *J. Synchr. Rad.* **2009**, *16*, 872–879. [CrossRef]
- 78. Gabadinho, J.; Beteva, A.; Guijarro, M.; Rey-Bakaikoa, V.; Spruce, D.; Bowler, M.W.; Brockhauser, S.; Flot, D.; Gordon, E.J.; Hall, D.R.; et al. MxCuBE: A synchrotron beamline control environment customized for macromolecular crystallography experiments. *J. Synchr. Rad.* 2010, *17*, 700–707. [CrossRef]
- 79. Battye, T.G.G.; Kontogiannis, L.; Johnson, O.; Powell, H.R.; Leslie, A.G.W. iMOSFLM: A new graphical interface for diffraction-image processing with MOSFLM. *Acta Crystallogr. D* **2011**, *67*, 271–281. [CrossRef]
- 80. Evans, P.R.; Murshudov, G.N. How good are my data and what is the resolution? *Acta Crystallogr. D* 2013, 69, 1204–1214. [CrossRef]
- Winn, M.D.; Ballard, C.C.; Cowtan, K.D.; Dodson, E.J.; Emsley, P.; Evans, P.R.; Keegan, R.M.; Krissinel, E.B.; Leslie, A.G.W.; McCoy, A.; et al. Overview of the CCP4 suite and current developments. *Acta Crystallogr. D* 2011, 67, 235–242. [CrossRef] [PubMed]
- 82. Bunkoczi, G.; Echols, N.; McCoy, A.J.; Oeffner, R.D.; Adams, P.D.; Read, R.J. Phaser.MRage: Automated molecular replacement. *Acta Crystallogr. D* 2013, *69*, 2276–2286. [CrossRef] [PubMed]
- 83. Kelley, L.A.; Mezulis, S.; Yates, C.M.; Wass, M.N.; Sternberg, M.J.E. The Phyre2 web portal for protein modeling, prediction and analysis. *Nat. Prot.* **2015**, *10*, 845–858. [CrossRef]
- 84. Matthews, B.W. Solvent content of protein crystals. J. Mol. Biol. 1968, 33, 491-497. [CrossRef]
- 85. Afonine, P.V.; Grosse-Kunstleve, R.W.; Echols, N.; Headd, J.J.; Moriarty, N.W.; Mustyakimov, M.; Terwilliger, T.C.; Urzhumtsev, A.; Zwart, P.H.; Adams, P.D. Towards automated crystallographic structure refinement with phenix.refine. *Acta Crystallogr. D* **2012**, *68*, 352–367. [CrossRef]
- 86. Painter, J.; Merritt, E.A. Optimal description of a protein structure in terms of multiple groups undergoing TLS motion. *Acta Crystallogr. D* **2006**, *62*, 439–450. [CrossRef]
- 87. Emsley, P.; Lohkamp, B.; Scott, W.G.; Cowtan, K. Features and development of Coot. *Acta Crystallogr. D* 2010, 66, 486–501. [CrossRef]
- 88. Laskowski, R.A.; MacArthur, M.W.; Moss, D.S.; Thornton, J.M. PROCHECK: A program to check the stereochemical quality of protein structures. *J. Appl. Crystallogr.* **1993**, *26*, 283–291. [CrossRef]

- 89. Chen, V.B.; Arendall, W.B., III; Headd, J.J.; Keedy, D.A.; Immormino, R.M.; Kapral, G.J.; Murray, L.W.; Richardson, J.S.; Richardson, D.C. MolProbity: All-atom structure validation for macromolecular crystallography. *Acta Crystallogr. D* **2010**, *66*, 12–21. [CrossRef]
- 90. Berman, H.; Henrick, K.; Nakamura, H. Announcing the worldwide Protein Data Bank. *Nat. Struct. Biol.* **2003**, *10*, 980. [CrossRef]

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