Abstract: Chemoreceptor (also called methyl-accepting chemotaxis protein, MCP) is the leading signal protein in the chemotaxis signaling pathway. MCP senses and binds chemoeffectors, specifically, and transmits the sensed signal to downstream proteins of the chemotaxis signaling system. The genome of Agrobacterium fabrum (previously, tumefaciens) C58 predicts that a total of 20 genes can encode MCP, but only the MCP-encoding gene atu0514 is located inside the che operon. Hence, the identification of the exact function of atu0514-encoding chemoreceptor (here, named as MCP$_{514}$) will be very important for us to understand more deeply the chemotaxis signal transduction mechanism of A. fabrum. The deletion of atu0514 significantly decreased the chemotactic migration of A. fabrum in a swim plate. The test of atu0514-deletion mutant (Δ514) chemotaxis toward single chemicals showed that the deficiency of MCP$_{514}$ significantly weakened the chemotactic response of A. fabrum to four various chemicals, sucrose, valine, citric acid and acetosyringone (AS), but did not completely abolish the chemotactic response. MCP$_{514}$ was localized at cell poles although it lacks a transmembrane (TM) region and is predicted to be a cytoplasmic chemoreceptor. The replacement of residue Phe328 showed that the helical structure in the hairpin subdomain of MCP$_{514}$ is a direct determinant for the cellular localization of MCP$_{514}$. Single respective replacements of key residues indicated that residues Asn336 and Val353 play a key role in maintaining the chemotactic function of MCP$_{514}$.

Keywords: Agrobacterium fabrum; chemotaxis; chemoreceptor; cellular localization; methyl-accepting chemotaxis protein; chemoeffector

1. Introduction

Chemotaxis is an adaptive behavior of motile bacteria moving along the concentration gradient of chemoeffectors towards an optimal environment [1,2]. This adaptive behavior is regulated by a two-component system composed of histidine kinase CheA and chemotaxis response regulator protein CheY [3,4]. When a chemical attractant exists in the environment, methyl-accepting chemotaxis protein (MCP) can bind the attractant, and the ligand binding will change the conformation of MCP; then, through a coupling protein CheW, the kinase activity of CheA will be suppressed. The suppression of CheA activity will delay the transfer of the phosphoryl group to the response regulator protein CheY, resulting in the decrease in phosphorylated CheY [5,6]. The absence of phosphorylated CheY makes less change of the rotational direction of flagella and thus keeps bacteria swimming to the attractant. On the contrary, when the concentration of attractant goes down (or repellent goes up), MCP activates the kinase activity of CheA. Self-phosphorylated CheA passes the phosphoryl groups to CheY. Phosphorylated CheY will bind to the flagellar motor and change the rotational direction of flagella frequently, which causes cell tumbling and changes direction away from the adverse environment [7,8].

Many bacteria possess chemotactic behavior for a large number of chemicals, such as different nitrogen or carbon nutrient substances, different environmental pollutants, or different signal chemicals released by their ecological partners [9–13]. To sense the vast
chemoeffectors, bacteria must evolve an enormous number of different chemoreceptors. A typical chemoreceptor contains an N-terminal periplasmic ligand binding domain (LBD), a transmembrane (TM) helical region, a HAMP (existing in histidine kinases, adenylate cyclases, methyl-accepting chemotaxis proteins, and phosphatases) domain, and a C-terminal cytoplasmic signaling domain (SD), comprising a methyl-accepting (MA) subdomain, a flexible bundle (FB) subdomain, and a hairpin (HP) subdomain (Figure 1A) [7,14,15]. The hairpin subdomain interacts with the coupling protein CheW and the sensor kinase CheA. Chemoreceptors use LBD to recognize chemoeffectors. In order to recognize different chemicals, chemoreceptors have evolved many different types of LBDs [9]. According to the LBD and membrane topology, chemoreceptors can be divided into four classes, of which class IV is cytoplasmic (soluble) chemoreceptors and the rest are transmembrane chemoreceptors. Based on the length from the N-terminus to the MA domain, cytoplasmic chemoreceptors can be divided into two subclasses of IVa and IVb. The N-terminal domain of IVa has at least 108 amino acids, while the N-terminal domain of IVb is shorter than 108 amino acids [16,17]. Among 8384 chemoreceptors defined by the MA subdomain in the complete genome of the SMART database [18], 14% of the chemoreceptors are cytoplasmic chemoreceptors. The ligand-binding motif is generally predicted at the N-terminal of IVa subclass chemoreceptors. Based on the statistical analysis of 1129 chemoreceptors with LBD in the SMART database, 47% of LBD is PAS domain, such as AerC in A. brasilense [19]; CZB domain found in Helicobacter pylori TlpD [20], is the second most common LBD, accounting for 8% of all IVa chemoreceptors; protoglobin [21] is the third most common LBD, accounting for 7% of all IVa chemoreceptors; the remaining domains account for less than 1% [22].

Agrobacterium fabrum is a Gram-negative bacterium and induces crown gall tumor disease in most dicotyledonous plants by genetically transforming the host [23,24]. In the natural environment, A. fabrum is usually distributed around the rhizosphere of a plant, and chemotaxis is an important first process of its interaction with the host. It has two kinds of lifestyles; one is to survive in the soil environment independently and the other is symbiotic with the plant as a pathogen [25–27]. Only when it recognizes and senses the sugars, acidic pH or phenols released by the host can A. fabrum begin the infecting process by chemotaxis toward the wound site of the host [28]. In the late 1980s, chemotaxis of A. fabrum was preliminarily studied, proving that it can respond to sugars, amino acids and phenols released by the injured plant tissues [29–31], but the specific chemotaxis mechanism has been rarely studied [32]. The genome sequence of A. fabrum C58 predicts that it has only one chemotaxis gene cluster (che cluster). The gene organization of this gene cluster is shown in Figure 1B. This gene cluster contains the genes encoding most components of the chemotaxis system. Spaces between the adjacent genes are very short and some adjacent genes even share a few nucleotides. All the genes in this cluster are predicted to be controlled by the same upstream promoter and thus located on the same operon [12,33]. In addition to the atu0514 gene, the only gene annotated to encode MCP in this che operon, A. fabrum carries an additional 19 MCP-encoding genes, including 1 on the Ti plasmid, 1 on the At plasmid, 5 on the linear chromosome, and the remaining 12 on the circular chromosome [33]. The number of MCPs in A. fabrum C58 indicates its complex chemotaxis signal and strong environmental adaptability [34].

The A. fabrum atu0514 gene, located on the circular chromosome, is the only MCP-encoding gene in the che operon [12]. Unlike all other MCPs, MCP314 is co-expressed with all other core chemotaxis components. Therefore, it may be the most important chemoreceptor, and characterizing its function is very helpful for us to further understand the chemotaxis signal transduction mechanism of A. fabrum. In this study, we firstly constructed the atu0514-deletion mutant Δ514 and the complemented strain Δ514-C and tested the effect of atu0514-deficiency on the chemotactic response of A. fabrum C58. We also identified the key residues that affect the cellular localization and function of MCP314.
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chemotaxis signal transduction mechanism of A. fabrum. In this study, we firstly constructed the Δ514 and the complemented strain Δ514-C and tested the effect of Δ514-deficiency on the chemotactic response of A. fabrum C58. We also identified the key residues that affect the cellular localization and function of MCP514.

Figure 1. Topological structure of a typical MCP, gene organization of a A. fabrum che gene cluster and domain organization of MCP514. (A) Architectural model of a typical MCP. Cylinders represent α-helices. MA: methyl-accepting (or sensory adaptation) subdomain; FB: flexible bundle subdomain; HP: hairpin subdomain. (B) Gene organization of A. fabrum che gene cluster. Arrows represent genes. Above the arrows are the names and lengths of the genes. Numbers under the arrows indicate the nucleotide position of the gene border. (C) Predicted domain organization of MCP514. MA-H1: α-helix 1 of MA subdomain; MA-H2: α-helix 2 of MA subdomain; numbers under the domains indicate the amino acid position of the domain border.

2. Materials and Methods
2.1. Primers, Plasmids, Bacterial Strains and Growth Conditions

The primers, plasmids and bacterial strains used in this study are listed in Supplementary Tables S1 and S2. Lysogeny broth (LB) liquid or agar medium was used to grow E. coli at 37 °C [35]. A. fabrum was grown in MG/L or AB-sucrose liquid or agar medium at 28 °C [36,37]. Concentrations of ampicillin and kanamycin used for E. coli were 100 and 50 µg/mL, respectively. Concentrations of kanamycin and carbenicillin for A. fabrum were 100 µg/mL.

2.2. DNA Manipulations

DNA manipulations followed the standard molecular protocols [35]. Plasmid isolation was performed with the TIANprep Mini Plasmids Kit (TIANGEN BIOTECH Corporation, Beijing, China). PCR products obtained by Veriti 96-well cycler (Thermo Fisher Scientific Inc., Waltham, MA USA) and DNA fragments were purified from agarose gels by using
the TaKaRa MiniBEST Agarose Gel DNA Extraction Kit (TaKaRa Corporation, Dalian, China). Plasmids were transferred into *E. coli* competent cells by heat-shock [35] and into *A. fabrum* by the Eppendorf electroporation instrument Eporator® (Eppendorf AG, Hamburg, Germany) [36].

2.3. Mutagenesis and Complementation of Atu0514

Based on the principle of homologous recombination, we used the pEX18Km-derived gene replacement plasmids to construct the corresponding gene deletion mutants [38,39]. Plasmid pEX18Km carries both a positive selection marker (kanamycin resistance, K<sub>mR</sub>) and counterselectable marker (suicide gene sacB) and cannot be replicated in *A. fabrum*. The positive selection marker allowed to select the transformants, in which the whole plasmid was integrated into the genome by the first homologous recombination. The counterselectable marker allowed to counterselect the transformants, in which both the target DNA fragment and the undesirable plasmid backbone were deleted from the genome by the second homologous recombination. The combined utilization of selectable and counterselectable markers can precisely generate an unmarked mutant without any undesirable DNA fragment (Figure S1A). *A. fabrum* wild type C58 was used to construct the atu0514-deletion mutant Δ514. *A. fabrum* cheW<sub>1</sub>-cheW<sub>2</sub> double-deletion mutant Δw [40] was used to construct atu0514-cheW<sub>1</sub>-cheW<sub>2</sub> triple-deletion mutant Δ514Δw. The desirable mutant was screened using PCR (Figure S1B) and verified by sequencing. The DNA fragment encoding amino acids 14–453 of MCP<sub>514</sub> was precisely deleted in both Δ514 and Δ514Δw mutants (Figure S1C). The complementation of MCP<sub>514</sub> in the atu0514-deletion mutant was fulfilled by the introduction of plasmid expressing MCP<sub>514</sub> (or its variants). Gene fusions with the egfp as well as deletion constructs were created by overlap extension PCR, as described by Higuchi [41]. These egfp-fused genes were cloned into the modified vector pUCA19 to generate the plasmids expressing eGFP-fused proteins. When these eGFP-fused proteins were expressed in *A. fabrum* cells, the cellular localizations of these eGFP-fused proteins could be observed by using Zeiss confocal microscope LSM 880 NLO (Carl Zeiss AG, Oberkochen, Germany).

2.4. Chemotaxis Assays

The procedure of capillary assay was followed as described by Adler in 1973 [42], with minor modifications. *A. fabrum* cells were harvested from mid-log-phase culture by centrifugation at 4000 rpm for 3 min at room temperature (25 °C) and suspended in chemotaxis buffer (0.1 mmol/L EDTA, 10 mmol/L KH<sub>2</sub>PO<sub>4</sub>, pH 7.0) to an OD<sub>600nm</sub> of 0.1. *A. fabrum* cell suspension (300 µL) was used to make a bacterial pond. The capillary tube was sealed at one end and filled with attractant at peak concentration dissolved in the chemotaxis buffer. The open end of the capillary tube was inserted into the bacterial pond and incubated for 1 h at room temperature (25 °C), then, the solution in the capillary tube was expelled and completely transferred into 1 mL of AB-sucrose medium. Dilutions were plated in duplicate on MG/L plates and incubated for 2 days at 28 °C. The colonies in the plates were counted and represented the number of cells attracted to the capillary tube.

The procedure of swim agar plate assay was followed that described by Merritt [43], with minor modifications. The tested strains were inoculated in AB-sucrose liquid medium and grown to the middle log phase, and then, the OD<sub>600nm</sub> was adjusted to 0.6 by using AB-sucrose liquid medium. A total of 3 µL of bacterial suspension was dropped onto an AB-sucrose swim plate containing 0.2% agar, and 5 replicates were set. After incubation at 28 °C for 36–48 h, the diameter of the bacterial colony circle was measured, and the data were statistically analyzed.

2.5. Bacterial Two-Hybrid Analyses

The bacterial two-hybrid system from the Stratagene® (Agilent Technologies Inc., Santa Clara, CA, USA) was used for testing protein–protein interactions. All the operations were conducted according to the manual. The open read frame (ORF) of atu0514 was
inserted into the bait plasmid pBT to express λcI-MCP_{514} (bait) fusion protein. ORFs of cheW_1 and cheW_2 were inserted to the target plasmid pTRG to express CheW_1 (target)–RNAP and CheW_2 (target)–RNAP fusion proteins, respectively. The interaction between bait (MCP_{514}) and target (CheW_1 or CheW_2) will take λcI and RNAP together to induce the expression of β-galactosidase, and thus, the bacterial colonies growing on plates containing 80 µg/mL X-gal will be blue. Otherwise, the bacterial colonies will be of normal color. Galactosidase activity was determined by the method of Miller [44].

2.6. Fluorescence Microscopy

For microscopy observation, agrobacterial cells from the mid-log-phase cultures were added to the center of the slides. A coverslip was placed on top of the culture droplet. The edges of the coverslip were sealed with acrylic polymer to prevent drying. *A. fabrum* cells were visualized by a Zeiss LSM 880 NLO system (Carl Zeiss AG, Oberkochen, Germany) using an Ar laser (excitation wavelength of 488 nm and emission wavelength of 500 to 550 nm) and a ×100 oil immersion objective. The images were analyzed and edited using ZEN lite (Blue edition), version 3.2 (Carl Zeiss AG, Oberkochen, Germany).

2.7. Statistical Analysis

The quantitative data shown in this study were the means with the standard deviations (SDs), which were derived from at least three independent experiments conducted in triplicate. Differences among bacterial strains were compared using one-way analysis of variance (ANOVA), followed by the Tukey test for multiple comparisons. The statistical analysis was conducted using Microsoft Office Excel’s data analysis tool (2019 version) (Microsoft Corporation, Redmond, WA, USA).

3. Results

3.1. MCP_{514} Is a Cytoplasmic Chemoreceptor, but Localized at Cell Poles

By SMART analysis, MCP_{514} has a total of 514 amino acids and carries three conserved domains, protoglobin domain, HAMP domain and cytoplasmic signal domain (SD), in the order from N-terminal to C-terminal (Figure 1C). The protoglobin domain of MCP_{514} shares 13.51% sequence identity with the LBD of HEMAT, a cytoplasmic chemoreceptor from *B. subtilis* [21], through the SWISS-Model homology search. According to Alexandre’s heptapeptide classification, cytoplasmic signal domain (SD) belongs to the 36H family [45]. The two best-studied MCPs of the 36H family are Tsr and Tar proteins of *E. coli*. Further analysis on the secondary structure of MCP_{514} by SOSUI [46] and SPLIT [47] shows that MCP_{514} does not contain the hydrophobic domain, indicating that MCP_{514} does not possess a transmembrane (TM) region and belongs to an IVa cytoplasmic chemoreceptor.

Transmembrane chemoreceptors are mainly localized at cell poles, but the cytoplasmic chemoreceptors have a wider cellular localization mode, ranging from co-localization with transmembrane chemoreceptor arrays to a diffuse cytoplasmic distribution [48]. The localization of some cytoplasmic chemoreceptors is associated with the physiology and life cycle of bacteria [22]. According to previous classification [9], MCP_{514} should be classified into the category of the cytoplasmic chemoreceptor due to the lack of a transmembrane region, but we do not know the cellular localization of MCP_{514}. To observe the cellular localization of MCP_{514}, the enhanced green fluorescent protein (eGFP) was fused to the N-terminus of MCP_{514} because only the N-terminally GFP-tagged MCP_{514} was functional [49]. This eGFP-MCP_{514} fusion protein was expressed in the MCP_{514}-deficient strain Δ514. Figure 2A showed that the eGFP-MCP_{514} fusion protein is localized at the poles of the *A. fabrum* cell. However, the eGFP-MCP_{514} fusion protein is distributed in the whole cell of *E. coli* (Figure 2B), verifying that MCP_{514} lacks a transmembrane domain. These data also imply that the polar localization of MCP_{514} in *A. fabrum* cell requires the assistance of other *A. fabrum* proteins.
chemicals. However, the chemotaxis of strain completely disappear, indicating that MCP 514 is not the receptor directly recognizing these substances but affects the chemotaxis ability of bacteria [42]. The chemotactic responses of wildtype C58, mutant Δatu0514 showed that the deficiency of MCP 514 significantly attenuates the overall chemotactic response of A. fabrum to nutrient substances. The deletion of the cheA gene will completely abolish the chemotactic response of A. fabrum, and thus, the CheA-deficient mutant Δa was used as a control of chemotaxis deficiency [40]. As shown in Figure 3, the deficiency of MCP 514 significantly attenuates the overall chemotactic response of A. fabrum to nutrient substances. The complementation of MCP 514 by the introduction of MCP 514-expressing plasmid can fully restore the chemotactic response of the MCP 514-deficient mutant to the level of the wildtype, confirming the role of MCP 514 in the chemotactic response. These results also provided evidence that the deletion of atu0514 did not have a polar effect on the rest of the che operon. Previous research showed that A. fabrum has chemotaxis toward sugars, amino acids, organic acids and phenols [29,30]. The traditional capillary assay is an effective method for quantifying the chemotaxis ability of bacteria [42]. The chemotactic responses of wildtype C58, mutant Δ514 and complemented strain Δ514-C to four various chemicals (1 µmol/L sucrose, 1 mmol/L valine, 1 mmol/L citric acid and 0.1 µmol/L acetylpiridineone (AS)) were measured by using the traditional capillary method. As shown in Figure 4, the deletion of atu0514 significantly weakens the chemotaxis of A. fabrum toward these four different types of chemicals. However, the chemotaxis of strain Δ514 toward these chemicals does not completely disappear, indicating that MCP 514 is not the receptor directly recognizing these substances but affects the chemotaxis efficiency in other ways.

3.3. Both CheW1 and CheW2 Interact with MCP514 but Do Not Affect the Cellular Localization of MCP514

It is known that CheW couples CheA to chemoreceptors and forms stable ternary signaling complexes with chemoreceptors and CheA [3]. To confirm the role of MCP514 in the chemotaxis signal transduction pathway of A. fabrum, the bacterial two-hybrid system was used to test the interaction between MCP514 and two CheWs. As shown in Figure 5A, the colors of the colonies representing MCP514/CheW1 and MCP514/CheW2 interaction are bluer than that of the negative control. The quantification of β-galactosidase activity also showed that the β-galactosidase activities of the colonies expressing these two tested protein pairs (MCP514/CheW1 and MCP514/CheW2) are significantly higher than that of

![Figure 2. Cellular localization of MCP514. Plasmid expressing the eGFP–MCP514 fusion protein was introduced into A. fabrum MCP514-deficient mutant Δ514 (A) and E. coli strain DH5a (B), respectively. Bacterial cells were grown to middle-log phase and observed by using confocal laser-scanning microscopy.](image)
the negative control (Figure 5B). This indicates that MCP$_{514}$ protein interacted with both CheW$_1$ and CheW$_2$ proteins.

Figure 3. Effect of MCP$_{514}$ deficiency on the chemotactic response of _A. fabrum_. The tested _A. fabrum_ strains were grown to middle-log phase. Bacterial cells were collected and resuspended to the same OD$_{600nm}$ (0.6). Equal amounts of cells from these cell suspensions were inoculated on the swim plates. The plates were incubated at 28 °C for 2 days. (A) Typical colonies of these tested _A. fabrum_ strains. (B) The swim-ring diameters of these tested _A. fabrum_ strains on the swim plate. The data represent the means ± SDs from five independent experiments in triplicate. The bars paired with two asterisks “**” represent that they are statistically different at $p < 0.01$ via the one-way ANOVA, followed by Tukey test. C58, _A. fabrum_ wildtype C58; Δ514, MCP$_{514}$-deficient mutant; Δ514-C, Δ514 mutant complemented with MCP$_{514}$ through plasmid; Δα, CheA-deficient mutant (a control of chemotaxis deficiency).

Figure 4. Effect of MCP$_{514}$ deficiency on the chemotactic responses of _A. fabrum_ to sucrose (A), valine (B), acetosyringone (C), and citric acid (D). The tested _A. fabrum_ strains were grown to middle-log phase. Bacterial cells were collected and resuspended to the same OD$_{600nm}$ (0.6). Equal amounts of cells from these cell suspensions were inoculated on the swim plates. The plates were incubated at 28 °C for 2 days. (A) Typical colonies of these tested _A. fabrum_ strains. (B) The swim-ring diameters of these tested _A. fabrum_ strains on the swim plate. The data represent the means ± SDs from five independent experiments in triplicate. The bars paired with two asterisks “**” represent that they are statistically different at $p < 0.01$ via the one-way ANOVA, followed by Tukey test. C58, _A. fabrum_ wildtype C58; Δ514, MCP$_{514}$-deficient mutant; Δ514-C, Δ514 mutant complemented with MCP$_{514}$ through plasmid; Δα, CheA-deficient mutant (a control of chemotaxis deficiency).
(B), citric acid (C) and acetosyringone (D). A. fabrum cells in the middle-log phase were collected, washed and then adjusted to an OD_{600nm} of 0.1 with chemotaxis buffer. Capillary tubes containing chemotaxis buffer with 10^{-6} mol/L sucrose, 10^{-3} mol/L valine, 10^{-3} mol/L citric acid, or 10^{-2} mol/L acetosyringone were inserted into the cell suspensions for 1 h at room temperature (25 °C). Cells migrating to the capillary tube were counted by using colony count. The attracted cells are equal to the cells in the capillary tube with attractant minus the cells in the capillary tube without attractant. The data represent the means ± SDs from three independent experiments with triplicate. The bars paired with “*”, “**” and “***” marks represent that they are different in a statistical manner at \( p < 0.05, 0.01 \) and 0.001, respectively, via the one-way ANOVA, followed by Tukey test. The strain names in the horizontal axis are the same as in Figure 3.

Both CheW1 and CheW2 interact with MCP514 and most of the ternary MCP-CheW-CheA complexes are localized in the cell poles. However, MCP514 lacks a transmembrane domain. It is unknown whether the cellular localization of MCP514 is dependent on the ternary MCP–CheW–CheA complex. Therefore, we tested the effects of the CheW deficiency on the cellular localization of MCP514. To determine whether CheW affects the cellular localization of MCP514, a plasmid expressing eGFP–MCP514 fusion protein was transferred into the \( atu0514\)–cheW1–cheW2 triple deletion mutant \( \Delta514\Deltaw \). Fluorescence observation showed that MCP514 in the CheW-deficient strain is still localized at cell poles, indicating that CheW deficiency does not affect the polar localization of MCP514 (Figure 6).

### 3.4. Helical Structure of Hairpin Subdomain Is Required for the Cellular Localization of MCP514

Due to the lack of a transmembrane domain in MCP514 and the evidence that CheW deficiency does not affect the polar localization of MCP514, it is most likely that MCP514 is localized in the cell poles via interacting with other MCPs. Results from E. coli MCPs showed that the hairpin subdomain of MCP is a coiled-coil of two antiparallel helices with a ‘U-turn’ and two hairpin subdomains form a supercoiled four-helical bundle, which makes MCP form homodimeric molecules [30]. The dimers of different MCPs can form mixed trimers of dimers via the interactions between their highly conserved helical bundle, and several residues play key roles in the formation of trimer (Figure S1) [51–53].

![Figure 5](image-url)
were fused to eGFP and expressed as the eGFP-MCP514 variant in the MCP514-deficient mutant. Residue Phe328 in the hairpin subdomain of MCP514 was replaced by Ala, Pro or Trp to generate three MCP514 variants, MCP514F328A, MCP514F328P and MCP514F328W.

To determine if the hairpin subdomain of MCP514 affects the cellular localization of MCP514, key residue Phe328 in the hairpin subdomain was changed to Ala, Pro or Trp. These three single-residue-substituted MCP514 variants were fused to eGFP, respectively. Fluorescence observation showed that the substitution of Phe328 for Pro causes the diffusion in the cytoplasm (Figure 7C), whereas the replacement of Phe328 by Ala or Trp does not affect the polar location of MCP514 (Figure 7A,B). Proline is a constraint on the formation of helix. Replacement of Phe328 by Pro will destroy the helical structure of the hairpin subdomain. These results demonstrated that the helical structure of the hairpin subdomain is required for the cellular localization of MCP514.

![Figure 6](image)

**Figure 6.** Cellular localization of MCP514 in CheW-deficient *A. faebrum* mutant. Plasmid expressing eGFP-MCP514 fusion protein was introduced into the atu051–cheW1–cheW2 triple-deletion mutant Δ514Awt. Cells were grown to the middle-log phase and observed by using confocal laser-scanning microscopy.

To further determine the key residues of MCP514 that are involved in the cellular localization of MCP514, we aligned the sequence of MCP514 with the sequences of Tsr and Tar from *E. coli* [55] and chose Phe328, Asn336, Glu340, Arg343 and Val353 of MCP514 as the target residues of the chemotactic function of MCP514.

![Figure 7](image)

**Figure 7.** Effects of the substitution of key residue Phe328 on the cellular localization of MCP514. Key residue Phe328 in the hairpin subdomain of MCP514 was replaced by Ala, Pro or Trp to generate three MCP514 variants, MCP514F328A (A), MCP514F328P (B) and MCP514F328W (C). Three MCP514 variants were fused to eGFP and expressed as the eGFP-MCP514 variant in the MCP514-deficient mutant Δ514, respectively. *A. faebrum* mutant Δ514 cells expressing these eGFP-MCP514 variant fusion proteins were grown to the middle-log phase and observed by using confocal laser-scanning microscopy.

**3.5. Two Key Residues of Hairpin Subdomain Play a Key Role in Maintaining the Chemotactic Function of MCP514**

The cellular localization of MCP514 is dependent on the hairpin subdomain, which is involved in the interactions between different MCPs, as well as the interactions with the CheA and CheW [54]. The chemotactic signal is collaboratively transduced by the mixed MCP teams, and all signals from different MCPs will converge to CheA [51]. It is reasonable that residues involving in the interaction between MCPs may affect the chemotactic function of MCP514.
site-directed mutation (Figure S2). Five residues were respectively replaced by alanine to generate five single residue-substituted MCP\textsubscript{514} variants. These MCP\textsubscript{514} variants were expressed in the MCP\textsubscript{514}-deficient mutant (Δ514) by the introduction of the MCP\textsubscript{514} variant-expressing plasmid, respectively. Colonies of the MCP\textsubscript{514}-deficient mutant expressing various MCP\textsubscript{514} variants are shown in Figure 8A. The diameters of these tested strain colonies were used to quantify the effects of various MCP\textsubscript{514} variants on the chemotactic response of \textit{A. fabrum} [43]. As shown in Figure 8, two single residue-substituted MCP\textsubscript{514} variants, MCP\textsubscript{514}\textsuperscript{N336A} and MCP\textsubscript{514}\textsuperscript{V353A}, are unable to restore the chemotactic response of the MCP\textsubscript{514}-deficient mutant to the level of the wildtype, demonstrating that residues Asn336 and Val353 play a key role in maintaining the chemotactic function of MCP\textsubscript{514}.

![Figure 8](image_url)

**Figure 8.** Effects of MCP\textsubscript{514} variants on the chemotactic response of \textit{A. fabrum}. The test procedure was the same as described in Figure 3. (A) Typical colonies of these tested \textit{A. fabrum} strains. (B) The swim-ring diameters of these tested \textit{A. fabrum} strains on the swim plate. The data represent the means ± SDs from five independent experiments in triplicate. The bars paired with ~~~*, ~~~**, ~~~*** and ~~~**** marks represent that they are different in a statistical manner at \( p < 0.05, 0.01, 0.001 \) and 0.0001, respectively, via the one-way ANOVA, followed by Tukey test. C58, \textit{A. fabrum} wildtype C58 strain; Δ514, MCP\textsubscript{514} deficient mutant; Δ514-C, Δ514 mutant complemented with native MCP\textsubscript{514}; MCP\textsubscript{514}\textsuperscript{F328A}, MCP\textsubscript{514}\textsuperscript{N336A}, MCP\textsubscript{514}\textsuperscript{E340A}, MCP\textsubscript{514}\textsuperscript{K343A} and MCP\textsubscript{514}\textsuperscript{V353A} represent Δ514 mutant complemented with the corresponding single residue-substituted MCP\textsubscript{514} variants, respectively; Δa, CheA deficient mutant.

4. Discussion

In natural environments, \textit{A. fabrum} is usually distributed around the rhizosphere of a plant [24]. Chemotaxis is the important initial step for \textit{A. fabrum} to infect the plant host [56]. Only when \textit{A. fabrum} correctly recognizes and responds to the chemical signals released by the plant host can it contact the host plant, infect the host and start the tumorigenic processes [27,57–59]. MCPs are the first components of the chemotaxis system. The recognition of chemicals by these proteins is the initial stage of chemotaxis signaling transduction. Adaptive modification of the conserved glutamate domain of the MCP signal domain ensures that they are highly sensitive to different concentrations of chemoeffectors [60]. Although \textit{A. fabrum} C58 contains 20 MCP-encoding genes, only one MCP-encoding gene (atu0514) is in the che operon [12]. It is rational that the MCP\textsubscript{514} encoded by the atu0514 gene may play a unique role in the signaling transduction of chemotaxis.

Our results show that MCP\textsubscript{514} deficiency significantly affects not only the overall chemotactic response of \textit{A. fabrum} to nutrient substances (Figure 3) but also the chemotaxis toward four various types of chemicals (Figure 4). The ligand binding domain (LBD) of MCP\textsubscript{514} is predicted to be a protoglobin, which cannot bind these four tested types of chemicals (Figure 1B). MCPs with different LBDs can form mixed trimers of dimers, and thousands of MCPs form a receptor cluster. The receptor cluster comprised of MCPs with different detection specificities collaboratively transduces the chemotactic signal in a team signaling model [51]. The effects of MCP\textsubscript{514} deficiency on the chemotaxis of \textit{A. fabrum} toward four various types of chemicals demonstrate that MCP\textsubscript{514} is a very important...
member of the chemoreceptor signaling team, and the absence of MCP\textsubscript{514} will affect the signaling efficiency of the whole chemoreceptor signaling team, explicating the reason of the \textit{atu0514} gene locating the \textit{che} operon.

Based on the sequenced genomes, 43\% of archaeal and 14\% of bacterial MCPs lack a transmembrane (TM) region. These TM-lacking MCPs are classified as cytoplasmic chemoreceptors. Unlike the transmembrane chemoreceptors, which are mainly located at the pole of the cell, cytoplasmic MCPs adopt more exotic locations [22]. Some cytoplasmic chemoreceptors are polarized at one end of the cell, such as, the HemAt of \textit{B. subtilis} and the IcpA of \textit{S. meliloti} [21,61]; other cytoplasmic chemoreceptors have both polar and diffuse states, for example, the AerC of \textit{A. brasilense} was diffused in cytoplasm under an oxygen-rich environment but located in the cell polar under symbiotic nitrogen-fixing state, which helps cells adapt to hypoxic environments [19]. There are also some bacteria whose cytoplasmic chemoreceptors formed clusters in the cytoplasm, for example, the TlpC and TlpT of \textit{R. sphaeroides} are located in the center of the cell by forming cytoplasmic clusters [62]. It is believed that the subcellular localization and distribution of the cytoplasmic chemoreceptor arrays are associated with the life cycle of bacteria [22]. Our results show that MCP\textsubscript{514} localizes cell poles. Although both CheW\textsubscript{1} and CheW\textsubscript{2} interact with MCP\textsubscript{514}, respectively, (Figure 5), the cellular localization of MCP\textsubscript{514} is independent of CheW (Figure 6), which is consistent with the previous results obtained from \textit{E. coli}, in which trimers of homodimers of MCPs form clusters of MCPs and in turn recruit CheA/CheW to assemble MCP-CheW-CheA ternary complexes [52,54]. In combination with the previous studies on other cytoplasmic MCPs [22,62], our results could support that this TM-lacking chemoreceptor, MCP\textsubscript{514}, localizes the cell poles through interacting with other transmembrane chemoreceptors, although the experimental evidence may be required to demonstrate the interaction of MCP\textsubscript{514} with other MCPs.

According to the previous studies on Tsr and Tar of \textit{E. coli}, the hairpin (HP) subdomain of MCP is not only the region forming homodimeric MCP but also the contacting sites for the formation of mixed trimers as well as the assembly of MCP-CheW-CheA ternary complexes [51,55]. Five conserved amino acid residues in the two antiparallel helices of \textit{E. coli} Tsr (Phe373, Asn381, Glu385, Arg388 and Val353) are important for the cluster formation and function of MCP [51]. Hydrophobic interactions between the helices contribute to the main trimer packing forces [55]. Five corresponding residues of MCP\textsubscript{514} conducted the single residue of the respective substitution. Phe328 of MCP\textsubscript{514} is in the key site of the helical structure of the HP subdomain. The replacement of Phe328 by the helix destroyer, proline, results in MCP\textsubscript{514} diffusing in the cytoplasm. Replacement of Phe328 by alanine does not affect both the cellular location and chemotactic function of MCP\textsubscript{514}. These results demonstrate that Phe328 is not the direct determinant to the chemotactic function of MCP\textsubscript{514}, but indirectly affects MCP\textsubscript{514} function through stabilizing the helical structure. Amongst the five tested MCP\textsubscript{514} variants, three variants have the full function of the wildtype, which is slightly different from the results of \textit{E. coli} Tsr [51].

5. Conclusions

MCP\textsubscript{514} is localized at cell poles, although it lacks a transmembrane region. The cellular localization of MCP\textsubscript{514} is independent of the assembly of MCP-CheW-CheA ternary complex but dependent on the interaction with other MCPs through the hairpin (HP) subdomain. Compared with other MCPs, MCP\textsubscript{514} plays a superior role in maintaining the signaling efficiency of the whole chemoreceptor signaling team. The helical structure of the hairpin subdomain is the prerequisite of MCP\textsubscript{514} cellular localization and functioning. The molecular mechanism of MCP\textsubscript{514} interacting with other MCPs and functioning in the chemotaxis signaling is similar to that of \textit{E. coli} MCPs, although the roles of some key residues of MCP\textsubscript{514} in the transducing signal are slightly different from those of \textit{E. coli} Tsr.
Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/microorganisms9091923/s1, Figure S1: Construction procedure, screening and verification of atu0514-deficient mutants. Figure S2: Sequence alignment of three A. fabrum MCPs and two E. coli MCPs. Table S1: Bacterial strains and plasmids used in this study. Table S2: Primers used in this study

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