Research article

Characterization of the nodulation plasmid encoded chemoreceptor gene mcpG from Rhizobium leguminosarum

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Abstract

Background: In general, chemotaxis in Rhizobium has not been well characterized. Methyl accepting chemotaxis proteins are sensory proteins important in chemotaxis of numerous bacteria, but their involvement in Rhizobium chemotaxis is unclear and merits further investigation.

Results: A putative methyl accepting chemotaxis protein gene (mcpG) of Rhizobium leguminosarum VF39SM was isolated and characterized. The gene was found to reside on the nodulation plasmid, pRleVF39d. The predicted mcpG ORF displayed motifs common to known methyl-accepting chemotaxis proteins, such as two transmembrane domains and high homology to the conserved methylation and signaling domains of well-characterized MCPs. Phenotypic analysis of mcpG mutants using swarm plates did not identify ligands for this putative receptor. Additionally, gene knockouts of mcpG did not affect a mutant strain’s ability to compete for nodulation with the wild type. Notably, mcpG was found to be plasmid-encoded in all strains of R. leguminosarum and R. etli examined, though it was found on the nodulation plasmid only in a minority of strains.

Conclusions: Based on sequence homology R. leguminosarum mcpG gene codes for a methyl accepting chemotaxis protein. The gene is plasmid localized in numerous Rhizobium spp. Although localized to the sym plasmid of VF39SM mcpG does not appear to participate in early nodulation events. A ligand for McpG remains to be found. Apparent McpG orthologs appear in a diverse range of proteobacteria. Identification and characterization of mcpG adds to the family of mcp genes already identified in this organism.

Background

Using a process termed chemotaxis, motile bacteria are capable of sensing their external environment and responding appropriately by moving towards increasing concentrations of nutrients and away from increasing concentrations of toxic compounds. The chemotaxis signaling pathway has been well characterized due to extensive studies using the model organism Escherichia coli, as well as detailed molecular characterization in numerous taxonomically distinct bacteria [1,2]. Methyl accepting chemotaxis proteins (MCPs) play key roles in the chemotactic response of many bacteria. These sensor proteins have N-terminal domains that detect attractants and repellents. When a ligand binds to a MCP, information about the external environment is transmitted from the MCP to a two component signal transduction system (CheA/CheY) inside the cell. The effect is alteration of swimming behavior by changing the direction of flagellar rotation or, in some cases, the speed of flagellar rotation, resulting ultimately in movement towards higher...
MCPs contain highly conserved structural domains; typically there are two transmembrane domains, a highly conserved signaling domain involved with CheA interaction, and two domains for methylation of glutamate residues [5]. MCP homologues have been discovered in a wide range of bacteria [[6,7] for review]. However, the function of these proteins in chemotaxis in many cases is only predicted through homology to the archetypal MCPs from *E. coli* and consequently the ligands and the precise role of these putative MCPS in chemotactic signaling have yet to be determined. Despite detailed characterization of the signaling pathways used in bacterial chemotaxis, the ecological significance of chemotaxis in bacteria remains relatively uncharacterized.

Chemotaxis has long been suggested to play a role in the early events of nodulation between rhizobia and host legumes [8–16]. Many legume root exudate compounds are chemoattractants for rhizobia. Both *Rhizobium leguminosarum* and *Sinorhizobium meliloti* exhibit chemotaxis towards the flavonoid compounds that induce symbiotic nodulation (*nod*) genes [13,15]. During infection the *Rhizobium* site of entry is commonly the tip of a developing root hair [17]. The legume plant may enhance infection by directing the rhizobia to the proper site of infection through secretion of chemoattractants. Chemotaxis towards plant root exudates could amplify *nod* gene induction by stimulating movement towards increasing inducer concentrations present at root surfaces [15].

In many species of rhizobia, symbiotic nitrogen fixation genes are localized on large plasmids, commonly referred to as sym plasmids (pSyms) or nodulation plasmids. In addition to these plasmids, most rhizobia harbor multiple large plasmids, some of which are still cryptic, and others whose function has only recently been elucidated [18–25]. It has been shown that these cryptic plasmids may play a role in rhizosphere competitive fitness [[19,20,24,25], Hynes unpublished]. Genes with homology to MCP-encoding genes have been previously reported on plasmids in strains from *Rhizobium leguminosarum* [16,26], *Rhizobium* sp. NGR234 [27] and *Sinorhizobium meliloti* [28]. In one instance, mutation of a plasmid residing *mcp* gene, *mcpp* from *R. leguminosarum* biovar *viciae* VF39SM, resulted in loss of ability to compete against wild type in the formation of nodules on Trapper peas [16]. The presence and significance of plasmid encoded *mcp* genes in rhizobia remains relatively unstudied. Studying sym plasmid localized *mcp* genes is of particular interest as co-localization with nodulation genes may suggest a role for nodulation related chemotaxis. This work characterizes *mcpG*, a sym plasmid encoded *mcp* gene, isolated from *R. leguminosarum* biovar *viciae* VF39SM.

**Results**

*mcpG* Cloning and Gene Characterization

*mcpG* was identified in a *R. leguminosarum* VF39SM cosmid library using a DNA probe derived from the highly conserved structure of MCPs. The restriction map of *mcpG* was drawn to scale relative to the translation product of a typical *mcp* ORF (TM, transmembrane domain; K1 and R1, methylation regions; HCSD, highly conserved signaling domain). The site of insertion of the Sp omega fragment and subsequent deletion in *mcpG* is also shown. Restriction enzymes are as follows: B, *Bam*HI; Bc, *Bcl*I; P, *Pst*I; S, *Sal*I; and X, *Xho*I.
conserved signaling domain of *R. leguminosarum* VF39SM mcpD [16]. The mcpG gene was subcloned from a cosmid as a BamHI fragment. Figure 1 provides a restriction map of mcpG. The entire mcpG coding region plus adjacent DNA regions was sequenced using a combination of subclones and use of appropriate primers. The mcpG gene sequence has been deposited in GenBank under the accession number AF141674. The open reading frame of mcpG is predicted to be 654 aa, with a molecular mass of 68.48 kDa. Transmembrane domains are predicted in McpG using two separate transmembrane helix prediction programs. TopPred II [29] predicts two transmembrane domains (TMD), the first spanning aa residues 12 to 32 and the second from aa 196 to 216. TM-HMM [30] also predicts two transmembrane regions similar in location to TopPred II; first TMD from aa 7 to 29 and the second from aa 194 to 216. These predictions suggest the N-terminal domain of McpG is located in the periplasmic space and likely functions as a sensor domain. The software predictions are in good agreement with the secondary structure of a typical MCP protein, whereby two transmembrane domains flank the N-terminal domain of the protein (Figure 1).

The C-terminal domain of McpG contains strong sequence homology to the methylation, and signaling domains of well characterized MCP proteins. Figure 2 shows an alignment of the C-terminal domains of McpG and the *E. coli* MCP Tsr. Results from a BLASTP alignment search of the GenBank database show that mcpG ORF exhibits the greatest similarity to putative MCPs from *Agrobacterium tumefaciens* C58 (McpU, Genbank accession AAK87918) and *S. meliloti* (McpY, Genbank accession AAG37852). However, a BLASTP alignment search using only the sequence flanked by the transmembrane domains (aa residues 34–190), presumed to be the sensor domain of McpG, yields different results. Figure 3 shows the alignment of the sensor region to ORFs of probable MCPs submitted to the Genbank database from genome sequencing projects (*Pseudomonas syringae*, http://genome.ornl.gov/microbial/psyr/; *Rhodopirillum rubrum*, http://genome.ornl.gov/microbial/rrub/; *Xanthomonas campestris*, http://cancer.lbi.ic.unicamp.br/xanthomonas/). A MCP sensor region detects attractants and repellents, directly or indirectly; therefore genes that are potential orthologs are those with significant homologies in the N-terminal sensor regions. To date, BLASTP searches of the sensor regions from the previously characterized *R. legu-
minosarum VF39SM MCPs [16] have not detected orthologs in any bacteria, including members of the Rhizobiaceae family (data not shown).

The genome sequencing of *R. leguminosarum* bv. *viciae* 3841 is ongoing at the Welcome Trust Sanger Institute http://www.sanger.ac.uk/Projects/R_leguminosarum. Searching the database for sequence homology to *mcpG* yields a single gene that is 99% identical to the *mcpG* sequence. In strain 3841 a gene precedes the *mcpG* homologue with homology to a family of monooxygenases. The highest homology (80% identical) is to an alkanesulfonate monooxygenase from *A. tumefaciens* C58 (Genbank accession AAL44243). Downstream of the *mcpG* homologue is a gene coding for a monocarboxylate permease (Genbank accession AJ421944) that is required for optimal growth with alanine as a sole carbon and nitrogen source [31]. Our sequencing results on upstream and downstream regions of *mcpG* from VF39 show that the gene organization in this strain is the same as in 3841.

In addition to the conserved methylation and signaling domains found in McpG (figure 2a) another conserved domain likely exists. Searching the NCBI Conserved Domain Database http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml revealed that the *mcpG* ORF has significant homology to the HAMP motif (figure 2b). The HAMP domain is a gene coding for a monooxygenase (Genbank accession AJ421944) that is required for optimal growth with alanine as a sole carbon and nitrogen source [31]. Our sequencing results on upstream and downstream regions of *mcpG* from VF39 show that the gene organization in this strain is the same as in 3841.

Plasmid localization and distribution of *mcpG* in Rhizobium spp

A DNA probe from the ca. 500 bp *Pst*I fragment of *mcpG* was used to search for *mcpG* homologues in other *Rhizobium* spp. The probe originates from the 3′ region of the gene, thereby lacking the highly conserved region of *mcp* genes, and is therefore unique to *mcpG* (Fig. 1); the fact that this probe did not hybridize to other *mcp* genes was confirmed by Southern blots to total DNA of VF39, which gave only one band as expected, in spite of the presence of at least 17 *mcp* gene homologues in this strain [16]. As evidenced by lanes A-C in Figure 4, *mcpG* homologues are found on plasmids from a number of *R. leguminosarum* strains.
species representing all three biovars of this species. Strains 3841 and T3CB, which are both derived from the same parent strain, carry the mcpG gene on the nodulation plasmid pRL10JI. The remaining strains represented in Figure 4 harbor mcpG on cryptic plasmids. Based on the published descriptions of these strains and our unpublished results, the pSym in each of these strains is: the largest plasmid in strain 14–2 (lane F), the smallest plasmid in strain 8002 (lane H) and the second smallest plasmid in strain GF160 (lane J). Strain 8401 (lane I) is 8002 cured of its pSym, and the identity of the pSym in strain 162Y10 is not known, though it is not one of the two smallest comigrating bands, one of which is hybridizing to our probe. mcpG is also found on plasmids in Rhizobium etli strains: Brazil5, F8, and CFN42 (data not shown). In strain CFN42, the plasmid carrying the mcpG homologue is pRetCFN42c, which is not the pSym [19].

Insertional mutagenesis of mcpG

Mutagenesis of mcpG was accomplished by disrupting the ORF through insertion of a Sp\(^{e}\) cassette within the coding region of mcpG. Figure 1 illustrates the location of the gene disruption relative to the predicted ORF. The mutated gene was introduced into VF39SM via double recombination using the strategy of Quandt and Hynes [34]. Southern blots of genomic DNA from putative mcpG mutants confirmed the replacement of the wild type gene with the insertionally disrupted mutant gene (data not shown).

The mcpG mutant was screened for altered chemotactic response to a number of carbon sources. Of particular interest were carbon sources whose catabolic genes reside on pReVF39d, the plasmid that carries mcpG. These carbon sources are: adonitol, alanine, hydroxy-L-proline, and trigonelline [[24,35], Hynes unpublished]. On swarm plates, the mcpG mutant maintained the same chemotactic response as wild type to these particular carbon sources (data not shown). Additionally, the chemotactic behavior of the mcpG mutant to a wide range of other carbon sources (various sugars and amino acids) was identical to the wild type (data not shown). The chemotactic phenotype of the mcpG mutants towards alanine, lactate, and pyruvate, which are known substrates of the monocarboxylate transporter which is encoded by the mct gene located adjacent to mcpG [31] was also unaltered.

The effect of a mcpG mutation on the ability of VF39SM to nodulate competitively was tested using a nodulation competition assay. Figure 5 shows the results of the competition experiment. Two independently isolated mcpG mutants were unaffected in their ability to compete for nodulation against the wild type strain VF39SM.

Discussion

The sequence homology of the mcpG ORF to known MCP proteins and its secondary structure strongly suggest that mcpG codes for a methyl accepting chemotaxis protein. Notably, sequence data from regions surrounding mcpG suggest that, as is the case in R. leguminosarum 3841, putative oxygenase genes flank mcpG and a monocarboxylate transport gene is in close proximity [31]. However alanine, lactate and pyruvate chemotaxis was unaffected in swarm plate assays with a mcpG mutant strain. In fact, swarm plate analysis with numerous carbon sources did not reveal a mutant phenotype. Given the large number of compounds that are chemoattractants for rhizobia [36] and the metabolic diversity of Rhizobium [37,38] identify-
ing a ligand for McpG could be difficult. Swarm plate analysis of mutant strains created from using previously isolated VF39SM mcp genes failed to reveal mutant phenotypes except in the case of a mutation in mcpB which resulted in impairment of a chemotactic response to a variety of carbon sources on swarm plates [16]. Metabolism of the test compound is a prerequisite for the swarm assay, so it is possible that a mutant phenotype is hidden in swarm plate analysis by an overriding redox chemotactic response. Redox chemotaxis has been observed in members of the alpha-proteobacteria family; particularly in Azospirillum brasilense where energy taxis is the dominant chemotactic behavior [39], but also in Rhodobacter sphaeroides [40]. The capability of redox chemotaxis in R. leguminosarum VF39SM is being investigated and an alternative chemotaxis assay that does not require metabolism of the chemoattractant and therefore eliminates overriding effects of any redox chemotaxis is currently being optimized.

Finding putative orthologs of mcpG in a diverse range of proteobacteria suggests that the Mcp senses a compound(s) found in the habitats of all these bacteria. All of these organisms have been isolated from soil environments and both X. campestris and P. syringae are known to colonize the rhizosphere of plant species. Interestingly, no mcpG orthologs are found in the genome sequences of Agrobacterium tumefaciens, M. loti, or S. meliloti. Based on the annotated database of X. campestris, a family of 19 mcp genes may exist; the genome sequences of P. syringae, and R. rubrum, to date, are drafts and therefore it is premature to conclude the size of the mcp gene families in these species. The large mcp gene family of X. campestris is very similar to that reported for R. leguminosarum VF39SM [16].

The results from the nodulation competition assay suggest mcpG does not play a role in the early events of plant infection. Previous work with mcp genes from VF39SM has shown that at least 2 mcps (mcpB and mcpC) do contribute to the strain's ability to compete for infection sites on pea plants [16]. Although mcpG is not needed during early infection events it may offer a competitive advantage in the rhizosphere.

Figure 5
Results of nodulation competition assay between wild type VF39SM and the two mcpG- strains, HLG1 and HLG2. The ratios are expressed as VF39SM:HLG1 or HLG2. The graph indicates that the % of nodules occupied by each mutant strain was in accordance to the initial inoculation used. The Chi-square test was used to confirm there was no statistically significant difference between the initial inoculum % and the recovery % from nodules.
Plasmids in *Rhizobium* sp. carry important genetic determinants, such as genes required for symbiotic nitrogen fixation, bacteriocin production, and catabolism of various carbon sources [18, 21–25, 41, 42]. The plasmid profiles of rhizobia vary greatly, both in size and number of plasmids. The origin and evolution of these plasmids is relatively unknown. Interestingly, in all strains tested, only *R. leguminosarum* VF39SM and *R. leguminosarum* 3841 had *mcpG* on the sym plasmid; in the remaining strains *mcpG* was found on non-sym plasmids. Genes for adonitol metabolism have been localized to plasmids in each of the strains used in Figure 4[24]. Notably, the adonitol catabolism genes appear to localize to the same plasmid as *mcpG* in all the tested strains. The localization of *mcpG* to different plasmids in a number of *Rhizobium* spp. may prove useful in supplementing ongoing studies regarding plasmid origin and evolution in *Rhizobium*.

The ecological significance of plasmid encoded *mcp* genes remains relatively uncharacterized. VF39SM contains at least 4 plasmid encoded *mcp* genes [16, 35, this study]. Additionally the pSym-encoded *mcp* gene isolated in a *R. leguminosarum* bv. *vicae* strain by Brito et al. [26] is different from these 4 genes. Notably, searching the DNA sequence of *Sinorhizobium meliloti* 1021 [28] for homology to *mcp* genes reveals only one plasmid encoded *mcp* gene. This gene resides on pSymA while no ORFs with homology to MCPs are found on pSymB. Both pSymA and pSymB are extremely large plasmids, and, similarly to *R. leguminosarum* plasmids, contain genes for catabolism of many substrates, yet they do not harbor *mcp* genes to the same extent. This contrast suggests that the ecological role of plasmid encoded *mcp* genes in *R. leguminosarum* may be distinctive and significant.

**Conclusions**

Based on sequence homology the *mcpG* ORF codes for a methyl accepting chemotaxis protein, and appears to be homologous to an MCP found in a diverse group of proteobacteria. The ligand for *mcpG* remains to be discovered and, in terms of early nodulation events, *mcpG* does not contribute to nodulation efficacy. *mcpG* is found widely distributed amongst *Rhizobium* spp., but is located on different types of plasmids in different strains. Isolation and characterization of this gene adds to the family of previously described *mcp* genes in *R. leguminosarum* [16, 26, 35].

**Table 1: Bacterial Strains and Plasmids**

| Bacterial Strain or Plasmid | Relevant Characteristics | Source or Reference |
|-----------------------------|--------------------------|---------------------|
| *E. coli* DH5α              | *EndA1, hsdR17, supE44, thi-1, recA1, gyrA96, relA1,* • (argF-lacZYA), U169. β 80flacZ ΔM15 | Life Technologies |
| S17-1                       | Spβ. RP4 tra region, mobilizer strain | [51] |
| *Rhizobium leguminosarum*   |                          |                     |
| VF39SM                      | biovar vicieae, Smr'     | [52] |
| HLG1                        | VF39SM, mcpG: ΩSp, Smr', Spβ | This work |
| HLG2                        | VF39SM cured of pReVF39d | [48] |
| LRS39401                    | VF39SM cured of pReVF39f | [48] |
| T3CB                        | biovar vicieae, 1062 derivative, | [53] |
| W14-2                       | biovar trifolii, Smr'    | [18] |
| 8002                        | biovar phaseoli          | [54] |
| 8401                        | 8002 cured of pSym plasmid | [54] |
| 0F60                        | biovar vicieae, Smr',    | [55] |
| 3841                        | biovar vicieae, JB300 derivative | [56] |
| 162Y10                      | biovar trifolii          | [57] |
| **Plasmids**                |                          |                     |
| pBSISK(+)                   | Cloning vector, Ap'      | Stratagene |
| pCYΩSp                      | Sp resistance gene from pH45Ω [58] flanked by MCS of p1918 with a MCS composed of inverted repeats of the pUC19 MCS, Ap' | [35] |
| p1918                       | delivery vector for gene replacement, Gm' | [34] |
| pJQ200mp18                  | 6 kb BamHI VF39SM genomic fragment containing mcpG cloned into pBSISK(+) | This work |
| pKC42                       | Bcl fragment with mcpG ORF cloned into pJQ200mp18 | This work |
| pCYG1                       | Bcl fragment with mcpG ORF cloned into pJQ200mp18 | This work |
| pCYG2                       | Derivative of pCYG1 with ΩSp cassette cloned into mcpG gene | This work |

Plasmids in *Rhizobium* sp. carry important genetic determinants, such as genes required for symbiotic nitrogen fixation, bacteriocin production, and catabolism of various carbon sources [18, 21–25, 41, 42]. The plasmid profiles of rhizobia vary greatly, both in size and number of plasmids. The origin and evolution of these plasmids is relatively unknown. Interestingly, in all strains tested, only *R. leguminosarum* VF39SM and *R. leguminosarum* 3841 had *mcpG* on the sym plasmid; in the remaining strains *mcpG* was found on non-sym plasmids. Genes for adonitol metabolism have been localized to plasmids in each of the strains used in Figure 4[24]. Notably, the adonitol catabolism genes appear to localize to the same plasmid as *mcpG* in all the tested strains. The localization of *mcpG* to different plasmids in a number of *Rhizobium* spp. may prove useful in supplementing ongoing studies regarding plasmid origin and evolution in *Rhizobium*.
Methods

Bacterial strains, plasmids and growth conditions
The bacterial strains and plasmids used in this study are listed in Table 1. R. leguminosarum strains were routinely cultured on TY medium [43] at 30°C while E. coli strains were cultured on LB medium [44] at 37°C. Chemotactic response of R. leguminosarum to specific carbon compounds was assayed using swarm media containing Vincent's minimal medium [45] 0.15% agarose and a sole carbon source as the potential chemoattractant [16]. The concentration of the carbon source was 1 mM. Carbon sources were purchased from Sigma-Aldrich (ON, Canada). When necessary, Rhizobium strains were cultured in media containing antibiotics at the following concentrations: spectinomycin, 500 µg/ml; streptomycin, 500 µg/ml. E. coli was cultured with the following antibiotic concentration when required: ampicillin, 100 µg/ml; and spectinomycin, 100 µg/ml. Antibiotics were obtained from Sigma-Aldrich (ON, Canada).

DNA manipulation and sequencing
Restriction enzymes and modifying enzymes were purchased from Life Technologies (ON, Canada) and used according to the manufacturer's instructions.

Plasmid profiles of Rhizobium strains were visualized on agarose gels using a modified Eckhardt procedure [46] described by Hynes et al. [47] and modified by Hynes and McGregor [48]. Probe labeling, southern blots, and detection procedures were performed using the non-radioactive DIG labeling and detection system as specified per the manufacturer's instructions (Roche Biochemicals, Laval, PQ, Canada).

DNA sequencing of the mcpG gene was accomplished using both subcloning and primer walking approaches. Contigs were assembled using DNASIS (Hitachi Software Engineering, CA, USA). Primers for sequencing were designed using the Oligo software application (National Biociences Inc., MN, USA) and synthesized by Operon (CA, USA). Sequencing was performed by the UC DNA Services sequencing facilities (University of Calgary, AB, Canada). Template DNA was prepared according to the facility's specifications. Sequence characterization was performed using DNASIS (Hitachi Software Engineering, CA, USA). Sequence alignments were performed using the BLAST [49] and clustalW [50] programs. To predict membrane spanning regions in the mcpG ORF the programs TopPred II and TMHMM were used [30,31].

Mutagenesis of mcpG was accomplished using an insertional mutagenesis strategy. A 2.1 Kb Bgl II fragment containing the mcpG ORF was cloned into the BamHI site of pQ200mp18. A spectinomycin resistance gene cassette was excised from p1918::Sp using Sall. The cassette was inserted into the mcpG gene through an XhoI digestion of pQ200mp18::mcpG. The resultant vector had a 341 bp deletion and a Sp cassette inserted within the mcpG ORF. The disrupted gene was introduced into VF39SM via double recombination using a protocol described by Quandt and Hynes [34]. Correct gene replacement was confirmed by Southern hybridization.

Nodulation competition experiments
Trapper peas (Pisum sativum) were surface sterilized by washing the seeds in 50% Sodium Hypochlorite for 5 minutes, followed by a second wash in 70% ethanol. Following the washes the seeds were rinsed 3 times in sterile distilled water. The seeds were germinated by placing them on water agar plates (12.5 g agar for 1 litre of distilled water) and incubating them at room temperature in the dark for 3 days. Seedlings were transferred to modified magenta jars that were designed to resemble Leonard Jars [24,45]. The peas were grown in a vermiculite substrate.

Once the peas were transferred to the magenta jars the seedlings were co-inoculated with VF39SM and a mcpG mutant strain in approximate 9:1, 1:1, and 1:9 ratios. The exact ratios were confirmed by performing viable plate counts on the inoculant cultures. The inoculated peas were then grown for 4 weeks, after which the nodules were harvested, and surface sterilized by washing them in a 20% solution of bleach for 5 min, followed by a 5 min wash in 70% ethanol. The nodules were then rinsed twice in sterile distilled water. Surface sterilized nodules were placed individually in microfuge tubes containing 50 µl of sterile distilled H2O and crushed using inoculating sticks. 5 µl of the macerate was spotted in duplicate onto TY plates containing Sm, and TY plates containing Sm and Sp to distinguish which strain had formed the nodule, the wild-type or the mcpG mutant strain. For each competition experiment set >50 nodules were sampled.

List of abbreviations
aa, amino acid; Ap, ampicillin; Kb, kilobase pairs; ORF, open reading frame; Sm, streptomycin; Gm, gentamicin; Sp, spectinomycin.

Authors’ Contributions
CKY drafted the manuscript, designed and executed the strategy for making the mcpG mutants, did the work on mutant chemotactic phenotypes, completed the sequencing, conducted Southern hybridizations and participated in the nodulation competition experiments. KTC subcloned and sequenced mcpG. KLDB conducted the nodulation competition experiments. MFH was involved in subcloning experiments, sequence assembly and bioinformatics, and performed the Eckhardt gels, as well as some of the Southern Blots. As principal investigator in the lab,
MFH also provided intellectual guidance for the entire the study, and was involved in writing the paper.
All authors read and approved the final manuscript.

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