AN ANALYSIS OF THE TSR-LSRB INTERACTION DURING AUTOINDUCER-2 CHEMOTAXIS IN ESCHERICHIA COLI

A Dissertation

by

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DOCTOR OF PHILOSOPHY

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Escherichia coli senses a variety of chemoeffectors. The Tsr chemoreceptor is known to mediate attractant response to serine. Previously, serine was the only attractant described for Tsr. Recent work suggests Tsr senses Autoinducer-2 as an attractant via a periplasmic binding protein. Autoinducer-2 is a general quorum sensing molecule recognized by Gram positive and Gram negative bacteria. In E. coli it is bound to LsrB during AI-2 uptake.

A testable model, based on the described interaction between MBP and the Tar chemoreceptor in E. coli, was generated. I first hypothesized that the shoulder regions of Tsr would associate with areas of both the amino and carboxyl domains of LsrB to elicit chemotaxis to AI-2. To test this, alanine-scanning mutations were generated in both Tsr and LsrB. Results suggest AI-2 is sensed through a direct interaction between Tsr and LsrB. I conclude residues Lys-147 and Glu-150 of Tsr and Asp-59, Asp-63 and Arg-252 of LsrB are critical for AI-2 chemotaxis.

I investigated the relationship between LsrB expression and AI-2 chemotaxis. I hypothesized altering expression of LsrB relative to chromosomal level would affect AI-2 chemotaxis. This was tested by growing cells to late phase, increasing LsrB present, and by inducing plasmid-borne LsrB. The results suggest increasing the LsrB present decreases AI-2 chemotaxis, though a clear conclusion cannot be drawn. I also explored the effect making lsr deletions would have on LsrB expression and AI-2 chemotaxis. I conclude there is little effect on AI-2 chemotaxis in these mutants. Lastly, I hypothe-
sized serine and AI-2 affect signaling to each other. This was tested by using Tsr variants deficient for serine chemotaxis to test AI-2 chemotaxis to establish that recognition of serine and AI-2 occurs independently at separate locations on Tsr. I introduced one ligand during log phase growth and tested chemotaxis response to the other ligand. Results indicate that serine and AI-2 bind at separate sites. However, AI-2 chemotaxis decreased when serine was present, though not vice versa. I conclude the response to serine is more robust since there are a limiting number of LsrB proteins present in the cell.
DEDICATION

I dedicate this dissertation to my wife, Ashlie Seely, for her endless patience and ceaseless motivation, and to my children, Astrid and Augustus Seely, for allowing me to fit lab time in between play times. I also dedicate this dissertation to my father and mother, Loran and Donna Seely, and to my brother, Philip Seely for their encouragement and support.
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Contributors

This work was overseen by a dissertation committee whose members were Professor Michael Manson (advisor) of the Department of Biology, Professor Steven Lockless of the Department of Biology, Professor Gregory Reinhart of the Department of Biochemistry and Biophysics, and Professor Arul Jayaraman of the Department of Chemical Engineering.

Contributions to this work were made by Dr. Sneha Jani of the Department of Biology, Dr. Manjunath Hegde, and Dr. Kuppan Gokulan of the Department of Health and Human Services in Little Rock, Arkansas. Dr. Kuppan Gokulan developed the LsrB-Tsr binding model. Dr. Manjunath Hegde constructed some of the LsrB mutants. Dr. Sneha Jani carried out the immunoblots and AI-2 uptake assay. She also constructed the lsr operon deletion mutants. All remaining work for this dissertation was completed by the student, under the advisement of Michael Manson of the Department of Biology. I was assisted by undergraduate researchers Joshua Zuniga, Paul Farrell and Attikos Hutras. The plasmids carrying the Tsr binding site mutations were a kind gift from Dr. Sandy Parkinson of the University of Utah.

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CHAPTER I
INTRODUCTION

*Escherichia coli* motility and environmental stimuli

*Escherichia coli* is a gram-negative bacterium of the family Enterobacteriaceae. It is rod-shaped and flagellated, measuring 2-4 μm in length and 0.7 μm in width. Its motility in liquid or semi-solid media is achieved through the use of 4-8 peritrichous flagella. When these flagella rotate counterclockwise, the flagellar filaments form a left-handed helical bundle that propels the cell forward in a run of several seconds (1). This run can be at speeds of up to ten times the cell body length per second (40μm/s), and either cell pole can act as leader (2). Cells switch from run to a very brief tumble when clockwise rotation of the flagella results in a chaotic, rapid reorientation of the cell due to dissociation of the flagellar bundle (1). Alternating between longer runs and shorter tumbles results in a three-dimensional random walk in a liquid environment (3).

The cell can modulate this run/tumble bias to respond to favorable or unfavorable environmental stimuli. Clockwise rotation of the flagellar motors is suppressed when a cell responds and moves to a favorable stimulus, allowing for a longer run in the direction of that stimulus, biasing the random walk when an attractant is present. In short, this biased random walk is comprised of the net movement up or down the attractant or repellent gradient (3) (Figure 1). Responding to these gradients of attractant or repellent is known as chemotaxis.
Figure 1. *E. coli* chemotaxis in an attractant gradient. Counter-clockwise flagellar rotation results in flagellar filaments forming a left-handed helical bundle, resulting in a run. Rotation clockwise dissociates the bundle and causes tumbling. Cell swims in a relatively straight-line run, interrupted by brief three-dimensional reorienting tumbles. This results in a biased three-dimensional random walk, as the cell suppresses the probability of tumbling and increases the periodicity of the runs as it migrates up the attractant gradient.
Four transmembrane methyl-accepting chemotaxis proteins (MCPs) are present in *E. coli* to mediate chemotaxis. These MCPs are located in polar clusters, typically in a large subpolar patch (2). They are named based on the molecules for which they are responsible for mediating response: taxis to *L*-serine and away from particular repellents (Tsr); taxis to *L*-aspartate and away from particular repellents (Tar); taxis to ribose and galactose (Trg); and taxis-associated protein for dipeptides (Tap) (4-10). A fifth receptor, Aer, mediates redox chemotaxis but does so in a methylation-independent manner (11-13). Serine and aspartate are the only substrates shown to bind directly to their specific chemoreceptors (14, 15). The other known substrates – maltose, ribose, galactose, glucose and dipeptides – require an initial binding to their respective periplasmic binding proteins before interacting with their specific MCPs. Additionally, some molecules are sensed as repellents by two of these chemoreceptors. Tsr senses both indole and *L*-leucine as repellents, while Tar senses nickel (Ni$^{2+}$) and some divalent cations such as cobalt (Co$^{2+}$) as repellents (6, 14, 16, 17).

**Structure and function of *E. coli* chemoreceptors**

Chemoreceptors exist in oligomeric units. Single receptor homodimers form trimers of dimers in the absence of other chemotaxis proteins, though these assemblies are stabilized by the scaffolding and kinase proteins at their cytoplasmic domain (18-21). These self-associated trimers of dimers localize to the poles of the cell in arrays (22-26) (Figure 2). These clusters contain at least a few of all four of the described chemoreceptors (Tsr, Tar, Trg and Tap) (27). These receptors interact in chemotaxis signaling, as
Figure 2. A patch of chemoreceptors embedded in the polar cell membrane. (a) An illustration depicting one lattice unit of chemoreceptors. The receptors are represented by blue circles, and are arranged in trimers of dimers. The CheA and CheW proteins are shown as red and purple rectangles, respectively. The P3 domain of CheA is represented by green circles. (b) An illustrated top view of a chemoreceptor array to show the hexagonal lattice. The cartoon illustrates the proposed assembly of an extended chemoreceptor array. CheA subunits P3 and P5, along with CheW are labeled in conjunction with the receptors to illustrate a core complex. Three core complexes are interconnected by CheA/CheW interactions to form a single lattice unit, which assembles further into larger receptor arrays (22, 23).
suggested by their close association in the receptor lattice (28-30). This is further supported by the finding that bound ligand can induce an amplified effect on signaling in receptor patches, up to 35 times more than expected of individual receptors alone (31). Some attractants can be sensed by *E. coli* at the sub-micromolar concentration level, and gradients can be sensed over five orders of magnitude (32, 33).

The amounts of Tsr and Tar, considered the major chemoreceptors, are present in approximately 3-5 fold greater abundance than Trg, Tap and Aer (34). The major chemoreceptors do not require the presence of another receptor in order to perform chemotaxis. This is not true of the minor chemoreceptors. Trg, Tap and Aer cannot perform chemotaxis alone, and each requires the presence of at least one of the major transducers before they can do so. Additionally, there is no requirement that a major chemoreceptor ligand be present as a prerequisite for the minor transducers to perform chemotaxis to their respective ligands (35, 36). These synergistic interactions serve to increase the sensitivity of the system with regard to chemoeffectector concentrations (37).

Each of the five separate functional domains of Tsr and Tar carries out a particular function (38, 39) (Figure 3). Ligands or binding proteins are sensed by the periplasmic domain, a four helix bundle (α1-4). In the studied cases, small molecule ligands bind at the interface of the two subunits while binding protein-ligand complexes bind across the apex of the dimer, interacting asymmetrically with both subunits (40-42). Based on recent work, the AI-2 binding protein LsrB is predicted to do the latter (43).

The transmembrane domain of the receptor is comprised of transmembrane helix one (TM1) and transmembrane helix two (TM2). This domain’s function is critical in
Figure 3. The functional domains of the chemoreceptor. The schematic illustration shows three-dimensional organization of the domains in relation to the receptor dimer in the cell membrane. Domains are distinct based on color and labeled in the center, and important features are labeled to the right.
transducing a signal across the cellular membrane. TM1 anchors the receptor in the membrane and stabilizes the homodimer. Bound ligand signals are transduced by TM2 across the membrane and into the cytoplasm. TM2 is directly linked to the receptor’s cytoplasmic region by a Histidine kinase, Adenylate cyclase, Methyl-accepting chemotaxis proteins, Phosphatase (HAMP) domain.

The HAMP domain exists as a four helix bundle. Each of the dimeric subunits consists of parallel, alpha helical amphipathic sequences (AS1 and AS2). Each dimer is connected by a flexible linker of fourteen residues (44-47). The cytoplasmic domain, an extended four helix coiled coil, is adjacent to the HAMP domain and consists of the dimeric adaptation and signaling domains, located just downstream of HAMP.

Spatial chemical gradients and the changes in chemoeffector concentrations can be recorded through a sensory adaptation system governed by reversible covalent modifications. Four to six glutamyl residues in this adaptation region can be methylated by the chemoreceptor-specific CheR methyltransferase or demethylated by the CheB methylsterase. These adaptation enzymes continuously maintain the methylation record through both phosphorylation of CheB by CheA, and through conformational changes that alter the propensities for methylation or demethylation. In other words, when an attractant is bound, methylation propensity is high, while demethylation is low; the opposite is true for unbound kinase-active receptors (48-50). This allows the cell to maintain a memory of its surroundings, particularly in regard to the chemoeffector gradient in which it may find itself. This is very useful in dynamic environments such as the human gastrointestinal (GI) tract.
A two-component phosphorelay pathway governs receptor activity and ultimately flagellar rotation. Prior to ligand-binding at its periplasmic domain, a chemoreceptor exists in a kinase-activating state that initiates a phosphorelay to the flagellar motors. At the distal tip of the kinase control domain's four helix bundle, are bound the kinase CheA and the scaffolding protein CheW. These Che proteins are necessary for the receptors to cluster in their polar patches (48, 49, 51).

The CheA histidine kinase, when bound to a chemoreceptor, is activated several hundred fold in the absence of chemoeffector gradients (52, 53). CheA uses ATP to autophosphorylate at a highly conserved His48 residue (54). This phosphate group can then be transferred to the Asp57 residue of the CheY response regulator protein, allowing CheY-P to bind with high affinity to the flagellar basal body, specifically the FliM motor protein (55-57). Upon sufficient CheY-P binding, one or more flagella will rotate clockwise (CW), leading to the characteristic tumble of the cell (58-61). Another substrate for CheA-P is the methylesterase CheB. Upon phosphorylation CheB-P will set about removing methyl groups from specific glutamyl residues of the adaptation domain involved in sensory adaptation (62).

**Stimulus input and signal transduction across the membrane**

At the periplasmic domain of the chemoreceptor, the input stimulus is first recognized. Direct ligand-binding, as in the case of L-aspartate with Tar or L-serine with Tsr, occurs at one of two rotationally symmetric sites at the homodimer interface. Subsequently, the chemoreceptor displays negative half-of-the-sites cooperativity to an ex-
treme degree, preventing a second, identical ligand molecule from entering and binding at the second site (63). Along with adaptive methylation, this negative cooperativity aides the receptors in returning to their pre-ligand bound signaling state during chemotaxis.

Ligand binding induces a conformational change of the chemoreceptor to the effect of a 20-degree rotation along the helical axes of the subunits, breaking the rotational symmetry of the homodimer (64, 65). This results in a slight downward displacement of helix 4, purportedly propagating the signal through TM2. This propagation through the transmembrane domain is not fully determined, but crosslinking studies with Tar suggest that attractant binding causes a piston-like, approximately 1.6 Ångstrom displacement of TM2 toward the HAMP domain, while methylation displaces it toward the periplasm in a return to the pre-ligand binding state (66-68).

Signal transduction passes through the transmembrane region and into the HAMP domain, where conformational changes link attractant binding at the periplasmic domain to kinase activity at the cytoplasmic domain. Here the evidence suggests that changes in stability of the four helix HAMP bundle are generated by helical sliding of TM2, which in turn modulates destabilization between AS2 of HAMP and the kinase-control region (69). Additional mutational and biochemical studies suggest that signal transduction is mediated through anti-symmetrical changes in helix-helix packing in the adaptation and cytoplasmic domains (70, 71). According to this predicted model, attractant binding information stabilizes the HAMP four helix bundle, thereby destabiliz-
ing the cytoplasmic domain and kinase control module, effectively turning off the ki-
nase.

When an attractant binds at the periplasmic receptor domain, CheA autophos-
phorylation activity is inhibited, reducing the amount of CheY-P present in the cyto-
plasm, inducing counter-clockwise (CCW) flagellar rotation. The reduction of CheY-P
is further enhanced through the activity of CheZ, a phosphatase that targets CheY-P, al-
lowing the cell to quickly reduce its CheY-P levels in response to attractant binding.
Bundles of CCW rotating flagella propel the cell in its characteristic run (72-76).

The receptors can be reset to a pre-stimulus state following attractant binding.
This is achieved through methylation and demethylation of specific glutamyl residues
that serve as substrates for the CheR (methyltransferase) and CheB (methylesterase) pro-
teins that carry out adaptive methylation (77). These enzymes bind to the NWETF pen-
tapeptide at the carboxyl terminus of the major chemoreceptors (Tsr and Tar). The glu-
tamyl residues within the adaptation domain are reached due to a flexible coupling of the
pentapeptide to the four helix bundle. The minor chemoreceptors (Trg and Tap) lack the
pentapeptide at their carboxyl terminus, relying primarily on CheR and CheB enzymes
bound to Tsr and Tar within the mixed receptor patch to undergo adaptive methylation
(77). Adaptation enzymes interacting with one receptor can modify sites equivalent to
those on approximately 6 neighboring dimers (78). The flexible tether of the receptor
that binds CheR and CheB has been shown in molecular models to allow CheR to physi-

cally reach at least a few modification sites on eight of the neighboring receptors, indi-
cating so-called assistance neighborhoods for adaptation response (79). Adaptive meth-
ylation serves to convey an environmental memory to the cells, allowing them to sense dynamic changes in their surroundings. This is achieved in part because the kinetic rates of the enzymes responsible for adaptation are slower than the time-scale of chemoeffector binding, allowing the cells to compare current ligand occupancy at the periplasmic binding sites with methylation of the glutamyl residues resulting from previous ligand binding \((39, 80)\). This serves as the basis for a chemotaxis response. A more complete view of the signaling cascade and adaptation is represented in Figure 4.

**Quorum sensing and Autoinducer-2**

Population density is a key factor in regulating various functions in bacteria, including control of pathogenesis and recruitment to and maintenance of biofilms \((81, 82)\). Quorum sensing is the means by which bacteria can assess population density, and it relies on the recognition and determination of autoinducers (AIs). These compounds increase proportionally with population density, diffusing back into the cell upon reaching a certain concentration threshold, mediating the quorum sensing response. Some genes are induced by AIs, namely those that are involved in that specific AI’s production, creating a positive feedback loop. Typically, the cell density must reach \(10^8\) cells per mL or higher in order to trigger AI effects on the population.

Quorum sensing through AIs also allows bacteria to communicate both in an intra- and interspecies manner. General autoinducer synthesis genes have been found in approximately half of the sequenced bacterial genomes \((81, 83)\). While it was previously thought that intra-species communication and regulation were the most important
Figure 4. The chemoreceptor signaling cascade. Left receptor shows cascade effects following attractant ligand binding (red theme). CheA kinase is not phosphorylated, thus the phosphorelay of signal to the flagellar motor does not occur. Right receptor shows effects in the absence of attractant ligand (black theme). Inactive components are shown in white with color outline. Solid lines denote enzymatic reactions, while dashed lines denote binding interactions. Adaptation sites are shown as methylated (black circles) or unmethylated (white circles).
aspect of autoinducer production, as evidenced by the species specificity of particular AIs, newer studies have indicated cross talk between different species or genera of bacteria is equally important and can be achieved through the use of general AIs (84, 85). Current knowledge indicates bacteria can integrate signals from their environments to determine if the AI being sensed comes from itself, from other microbes present, or from human stress hormones (86). This ability would be particularly useful in the human GI tract.

Species-specific autoinducers (AI-1s) in Gram-negative bacteria are acyl homoserine lactones and modified peptides in Gram-positive bacteria. A separate class of AI is more generally recognized across many Gram-negative and Gram-positive species. Autoinducer-2 is an example of a general AI, and is used in conjunction with species-specific AIs in certain bacteria. For example, the marine bacterium *Vibrio harveyi* requires the presence of both a specific AI-1 and AI-2 to fully induce bioluminescence (87). This bioluminescence study was the first in which AI-2 was identified.

Following its identification, AI-2 and its synthase enzyme LuxS have been found to be widely distributed in different bacteria across the kingdom Eubacteria (83, 84, 88). This discovery helped shape the prediction that AI-2 is used for interspecies communication. AI-2 is synthesized by LuxS, a homodimeric Fe^{2+} metalloenzyme involved in the activated methyl cycle of bacteria (89-91). In this cycle, S-adenosylmethionine (SAM) is converted back to homocysteine, through an S-adenosylhomocysteine (SAH) intermediate, although it can also be converted to 4,5-dihydroxy-2,3-pentanedione (DPD) from S-adenosylribosylhomocysteine (SRH) by LuxS cleavage (84, 92, 93). This DPD mole-
cule is highly unstable and quickly converted to a precursor for AI-2 (93). Both the activated methyl cycle and DPD cyclization pathway are shown in Figure 5. This DPD biosynthetic pathway in *Vibrio harveyi* has been shown to be identical for many bacteria, including *E. coli* (92, 93).

Once the structure of AI-2 was determined through co-crystallization studies in *Vibrio harveyi*, it was determined that a borated DPD spontaneously cyclizes to form a furanosyl borate diester AI-2 compound, binding to its cognate receptor LuxP (94). A similar investigative approach was used in *Salmonella* to co-crystallize AI-2 with its cognate receptor, the Lsr (LuxS-regulated) binding protein, a product of the Lsr operon. The co-crystallization with its binding protein also revealed that *Salmonella* recognizes the chemically distinct, non-borated (2R,4S)-2-methyl-2,3,3,4-tetrahydroxytetrahydrofuran (R-THMF) form of AI-2 (95). Both the borated and non-borated forms exist in equilibrium with each other, though this can be influenced by the addition or removal of borate (81, 94).

**The Lsr operon and its role in AI-2 uptake and chemotaxis**

In *Salmonella enterica* serovar Typhimurium, AI-2 concentration is regulated by the seven Lsr operon gene products, including an ATP-binding cassette (ABC) transporter and AI-2 degradation proteins (96). In this system, AI-2 serves as a transcriptional activator, relieving LsrR repression of the *lsrACDBGE* operon. This increases expression of the AI-2 transport apparatus: ABC transporter proteins LsrA, LsrC and LsrD, as well as the periplasmic binding protein LsrB. Peak extracellular AI-2 accumu-
Figure 5. The synthesis pathways for DPD and AI-2. (a) The activated methyl cycle involves cleavage of S-ribosylhomocysteine (SRH) into homocysteine or dihydroxypentanedione (DPD), the precursor for AI-2. (b) DPD is highly unstable and spontaneously cyclizes in the presence of borate into borated and non-borated isomers that exist in equilibrium (95). Only the non-borated AI-2 (R-THMF) is sensed by *E. coli*. 
lation occurs during mid to late exponential growth and declines rapidly at the onset of stationary phase as a result of its import through the transporter into the cytoplasm (97). The \textit{lsrF}, \textit{lsrG} and \textit{lsrE} genes are required for processing and turnover of AI-2 once it reaches the cytoplasm (96). Two adjacent, but separately transcribed genes are \textit{lsrR} and \textit{lsrK}. The kinase product of \textit{lsrK} phosphorylates internalized AI-2, and it is this AI-2-P that de-represses the LsrR repressor and activates transcription of the other \textit{lsr} genes (Figure 6a) (96).

\textit{E. coli} possesses a homologous \textit{lsr} operon, the \textit{b1513} operon (Figure 6b), encoding homologous genes and functions (98). Because of this homology, the \textit{lsr} operon nomenclature will be used henceforth to describe \textit{E. coli} genes and gene products pertaining to the \textit{b1513} operon. The transport channel for AI-2 uptake is composed of LsrA, LsrC and LsrD and is located in the cytoplasmic membrane. Translated with a leader peptide, LsrB is exported to the periplasm via the Sec system. Here it is free to bind the extracellular AI-2 that remains in the periplasm following its export from the cell mostly via the TqsA protein (99). The remaining AI-2 molecules diffuse through outer membrane pores and accumulate externally, re-entering the periplasm via equilibration with the external environmental AI-2 when it reaches a concentration threshold.

Recent studies with AI-2 in \textit{E. coli} revealed that only the \textit{R-THMF} form is recognized as an attractant that stimulates chemotaxis, requiring both the Tsr chemoreceptor and the binding protein gene product of the Lsr operon, LsrB (43, 100). \textit{E. coli} LsrB closely resembles \textit{Salmonella} LsrB (Figure 7), to an 85\% sequence identity (101). When LsrB is deleted chromosomally, \textit{E.coli} does not respond to AI-2 in chemotaxis assays.
Figure 6. The *Salmonella lsr* operon and its homolog in *E. coli*. (a) The *lsr* locus in *Salmonella*. Transcription units are separated by black promoter boxes, and black arrows indicate direction of transcription for each unit. The small dark blue pentagons represent AI-2, which is made by LuxS and exported from the cell via the TqsA transporter (not shown). The Lsr ABC transporter reimports AI-2 into the cytoplasm, where it is phosphorylated by the LsrK kinase using ATP. Phosphorylated AI-2 relieves repression by LsrR, allowing expression of the *lsrACDBFG* operon. LsrF and LsrG degrade AI-2-P. The function of LsrE is not clear, and it is absent in *E. coli*. (b) The *b1513* operon of *E. coli* is a homolog of the *Salmonella lsr* operon. The *E. coli* gene designations are given inside the arrows, and the *Salmonella* homologs are listed beneath each arrow. Figure adapted from Xavier and Bassler (98).
Figure 7. *Salmonella* LsrB crystal structure. The cartoon depicts the crystal structure of *Salmonella* LsrB in complex with the R-THMF isomer of AI-2. The amino and carboxyl terminals are labeled, as is the AI-2 ligand. The two subunits are asymmetrical, entering the closed conformation only when AI-2 is bound. *E. coli* LsrB is 85% identical to *Salmonella* LsrB, with only a few amino acid residue differences, and the AI-2 binding sites are identical. Only the R-THMF isomer of AI-2 binds in such a manner to elicit an attractant chemotaxis response in *E. coli*. Deletion of LsrB abolishes chemotaxis to AI-2 in *E. coli*. PDB ID: 1TJY.
Figure 8. Proposed AI-2 chemotaxis pathway in *E. coli*. The R-THMF isomer of AI-2 is shown to the left (not to scale). The figure to the right shows the proposed pathway by which *E. coli* cells sense and respond to AI-2 (red pentagons) as a chemoattractant. The ABC transporter product of the *lsr* operon is shown in the cell membrane. AI-2-bound LsrB binds to the channel during AI-2 uptake as previously reported (98). Also present in the membrane is the Tsr chemoreceptor, complete with chemotaxis proteins for alternating flagellar rotation. This model proposes that LsrB can interact with the Tsr receptor to elicit an attractant response, inhibiting phosphorylation along the phosphorelay pathway and causing a CCW rotation of the flagella to propel the cell in a run.
Conversely, deletion of the ABC transporter components of the \textit{lsr} operon does not negatively affect AI-2 chemotaxis, suggesting that uptake of AI-2 to the cytoplasm is not necessary for chemotaxis to occur \((43)\). Only deletion of Tsr was shown to abolish AI-2 chemotaxis \((43)\). Thus, the study concluded that (i) both Tsr and LsrB were necessary for \textit{E. coli} to perform chemotaxis to AI-2; and (ii) uptake into the cell is not required to elicit an attractant response. It was hypothesized, based on this information, that AI-2-bound LsrB in the periplasm likely interacts with the periplasmic domain of Tsr to produce a chemotactic signal (Figure 8).

**Dissertation overview**

Previous work in the field of chemotaxis had never identified a binding protein interaction with the Tsr chemoreceptor. Only recently has a binding protein interaction for Tsr been discovered, although the details surrounding this interaction remained unclear \((43)\). The research undertaken and presented in this dissertation examines the interaction between Tsr and the AI-2-bound LsrB binding protein during AI-2 chemotaxis.

Chapter II describes my research in altering residues at the predicted interaction sites between Tsr and LsrB. The hypothesis was that individually altering select residues on both proteins would disrupt chemotaxis to AI-2 and give a better picture of the binding interaction and the critical residues involved in stabilizing the interaction to allow for signaling. The results suggest that the Lys-147 residue on the periplasmic shoulder regions of the Tsr homodimer are critical for AI-2 chemotaxis to occur. The results also suggest several residues in both the N-terminal and C-terminal domains of the LsrB monomer are important for this chemotaxis response, with Asp-59 in particular
showing a dominant negative phenotype for AI-2 chemotaxis when altered. All of the mutations generated were checked to make sure that that particular mutation did not prevent expression of the protein in an attempt to avoid false conclusions.

In Chapter III, my research attempted to correlate chemotaxis to AI-2 with various levels of LsrB expression. Plasmid-borne LsrB was induced with varying concentrations of IPTG to increase the amounts of the protein present in the cell during chemotaxis. It was determined that only about 23-25 copies of LsrB exists per cell under normal conditions, and that increasing the amount of LsrB present via induction drastically reduced chemotaxis to AI-2. Non-polar deletion mutants of the lsr operon were also used in attempt to determine how its components affect LsrB expression levels in vivo, and how these levels correspond to AI-2 chemotaxis.

Chapter IV describes my efforts in determining if (a) mutating residues crucial for the serine binding pocket also alter response to AI-2, and (b) if serine and AI-2 alter response to one another when the other is present at high concentrations. First, mutants for serine chemotaxis were exposed to a gradient of AI-2 and their response was determined. Results showed a lack of response to serine, but unimpaired response to AI-2. Additionally, mutations from Chapter II that reduced or abolished chemotaxis to AI-2 did not affect response to serine in those mutants. Second, exposing cells to high concentrations of one chemoeffector, then running chemotaxis assays to detect response to the other chemoeffector, gave an idea of the effect of the presence of one ligand on the other. Chapter V summarizes the experiments and discussion from my research and proposes future directions and experiments for this investigation.
CHAPTER II
EXAMINING THE TSR-LSRB BINDING INTERACTION THROUGH MUTATIONAL ANALYSIS

Introduction

Population density regulates many physiological activities and responses in bacteria, including formation of biofilms and expression of virulence factors. This regulation is accomplished through the use of quorum-sensing molecules that accumulate as population density increases. Autoinducer-2 (AI-2) is a general quorum-sensing molecule produced and secreted by numerous Gram-positive and Gram-negative bacteria. It is derived from the ribose moiety of S-adenosylhomocysteine following that molecule’s conversion of that to S-ribosylhomocysteine and subsequent cleavage by the LuxS enzyme into 4,5-dihydroxy-2,3-pentanedione (DPD). The unstable, linear DPD molecule spontaneously cyclizes into either (2S,4S)-2-methyl-2,3,3,4-tetraydroxytetrahydrofuran borate (S-THMF borate) or (2R,4S)-2-methyl-2,3,3,4-tetrahydroxytetrahydrofuran (R-THMF).

S-THMF borate regulates bioluminescence and virulence in Vibrio species, including Vibrio harveyi. A periplasmic binding protein of Vibrio, LuxP, binds this borated form of AI-2 with high affinity ($K_D$ of <1 µM) (102, 103). The R-THMF isomer of AI-2 is reported to bind to the LsrB periplasmic binding protein of Salmonella, E. coli and other enteric bacteria with an apparent $K_D$ of 160 µM (103). This reported binding affinity for R-THMF AI-2 to LsrB is somewhat conjectural, due to the extreme instabil-
ity of AI-2 in solution, and the equilibrium between the two naturally occurring THMF isomers.

AI-2 is synthesized by LuxS and secreted into the periplasmic space, partly by the activity of TqsA efflux transport. Once in the periplasm, it can passively diffuse into the external environment to be sensed by neighboring cells. The ability of cells to take up AI-2 relies on the products of the *lsr* operon. The LsrB periplasmic binding protein binds AI-2 and interacts with an ABC transporter composed of a heterodimer of the two membrane-spanning LsrCD proteins and the cytoplasmic kinase LsrA.

The LsrK kinase phosphorylates the DPD intermediate of AI-2, creating phospho-DPD (DPD-P). This phosphorylated molecule binds the repressor of the *lsr* operon, LsrR, relieving repression and inducing transcription of *lsrACDBFG*. Two downstream gene products in this operon, LsrF and LsrG, degrade DPD-P, completing the regulatory cycle of AI-2 uptake and processing. This is an unconventional cycle, as AI-2 induces expression of factors needed both for its uptake and those required for its destruction.

Recently, it was shown that AI-2 is a potent chemoattractant for *E. coli* strain RP437, requiring the activity of both the LsrB periplasmic binding protein and the Tsr chemoreceptor (43). Import of AI-2 into the cytoplasm is not required for its being sensed as an attractant. A mutant lacking the ABC transporter protein LsrC was still able to perform normal chemotaxis to AI-2. Because of this finding, it was proposed that AI-2-bound LsrB interacts directly with the periplasmic domain of the Tsr chemoreceptor dimer. There is precedent for this type of interaction between binding protein and chemoreceptor, as evidenced with the *E. coli* aspartate receptor (Tar) and maltose bind-
ing protein (MBP) \((104, 105)\). Periplasmic maltose must first bind to MBP, causing a conformational shift of MBP that can interact with Tar. Maltose-loaded MBP interacts with both subunits across the Tar dimer, involving certain residues on the surfaces of MBP and Tar \((104, 106)\). This is also true of ribose and galactose binding proteins with Trg and DppA with Tap \((5, 7, 8, 10)\).

Based on the published Tar-MBP interaction, a computerized docking simulation was performed of the x-ray crystallographic structure of *Salmonella* LsrB, in its ligand-bound conformation \((95)\), with the x-ray crystallographic structure of the ligand-free periplasmic domain of *E.coli* Tsr \((107)\). In the Tar-MBP model, the ligand-free Tar was oriented to the most likely fit of MBP using SPOCK software \((108, 109)\), then refined based on the experimental data.

Likewise, the Tsr-LsrB simulation was carried out using this software to obtain a testable model. Next, site-directed alanine scanning mutagenesis was used to probe the amino acid residues predicted to be at the sites of interaction between the two proteins. Tsr variants generated retained wild-type function for serine chemotaxis, and LsrB mutant proteins remained stable, and maybe functional for AI-2 uptake. The binding site for R-THMF was not affected by these residue changes.

The results reported here support the computer docking model for the LsrB-Tsr interaction and identify residues on both proteins that are critical for their interaction during binding and signaling, ostensibly without affecting the ability of LsrB to carry out its previously described AI-2 binding and transport functions.
Materials and methods

Bacterial strains, plasmids and media. Strain CV1, equivalent to strain RP437 (110), was used as the wild-type E. coli strain for chemotaxis. All relevant strains and plasmids are listed in Table 1. Plasmids pCA24N was used to express the wild-type and mutant Tsr and LsrB proteins at near-chromosomal levels. Additionally, plasmid pBAD18 was used to express lsrB from the arabinose-inducible pBAD promoter (111). Mutations were introduced into tsr and lsrB using the standard protocol for site-directed mutagenesis (Stratagene). Tryptone broth (TB; 10 g/L tryptone and 8 g/L NaCl) was used to culture cells at 30°C for all assays (112). Chemotaxis buffer (CB; 1X phosphate-buffered saline, 0.1 mM EDTA, 0.01 mM L-methionine and 10 mM DL-lactate) was used to wash cells and deliver attractants. AB medium contained 2 g/L casamino acids, 12.3 g/L magnesium sulfate heptahydrate, 17.5 g/L NaCl, 1 mM L-arginine, 1 % v/v glycerol and 10 mM potassium phosphate buffer, pH 7.0 (113). Chemically synthesized DPD (114) (17.57 mM) was purchased from the research group of Rita Ventura at the Insituto de Tecnologia Química e Biolóigica (ITQB-UNL) in Oeiras, Portugal. L-serine was purchased from Fisher Scientific (Fairlawn, NJ). Vibrio harveyi TL26 was a kind gift from Bonnie Bassler, Princeton University.

Measuring chemotaxis to attractants. Capillary assays, first developed by Adler et al. (115) were used to quantify the response to AI-2. Overnight cultures of cells expressing Tsr or LsrB mutant proteins from pCA24N were grown at 30°C in TB supplemented with streptomycin (50 µg/mL) and chloramphenicol (35 µg/mL). Cultures were back-diluted to a turbidity of ~0.050 at O.D.600 in 25 mL TB without antibiotic and then
<table>
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<th>Strain or plasmid</th>
<th>Genotype</th>
<th>Resistance</th>
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<td>Wild-type chemotaxis</td>
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<tr>
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<td>CV1 Δtsr</td>
<td>Strep</td>
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<td>CV5 +pCA24N-&lt;i&gt;tsr&lt;/i&gt;</td>
<td>Strep/Cm</td>
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<td>TL26</td>
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<td>Amp</td>
<td>This study&lt;sup&gt;h&lt;/sup&gt;</td>
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<sup>a</sup> Strep – streptomycin; Kan – kanamycin; Cm – chloramphenicol; Amp – ampicillin

<sup>b</sup> Contains Δ<sup>tsr9101</sup> introduced via phage transduction

<sup>c</sup> Constructed by transforming strain CV5 with pCA24N-<i>tsr</i>

<sup>d</sup> Constructed by removing Kan cassette (120) from MJ101(43)

<sup>e</sup> Constructed by removing the kanamycin resistance cassette from <i>lsrB</i> in MJ101 using FLP recombinase. This removal was selected for through lack of growth on Kan media and PCR was used to verify its removal. Phage transduction was then used to add the kanamycin resistance cassette to disrupt luxS. This was selected for through growth on Kan media and verified through PCR, then Kan cassette was removed using FLP recombinase.

<sup>f</sup> Disrupted for sensing AHLs and CAI-1. It senses only exogenously added AI-2.

<sup>g</sup> This plasmid was transformed into MJ101 and selected for through Amp-resistance. The recombinase recognizes flanking sites of the Kan-resistance cassette and removes it from the DNA. The plasmid is then cured by growth at 43°C, restricting its temperature sensitive origin of replication. The resulting strain is now both Kan- and Amp-sensitive.

<sup>h</sup> Constructed by cloning CV1 chromosomal <i>lsrB</i> into the pBAD18 vector.
Figure 9. The capillary assay. Cell chamber ponds are created by placing a C-shaped plastic gasket onto a glass microscope slide, then adding a square, glass coverslip on top. Washed cells are loaded into the chamber to create the pond. 1 mm capillaries loaded with chemoeffector are inserted into the chamber pond and incubated for 45 minutes to allow diffusion and chemotaxis response. Capillaries are then removed and their contents expelled into dilution buffer for plating. Overnight incubation of plates allows for colonies to be counted and the number of cells accumulated inside the capillary can be calculated (115).
grown to mid-logarithmic phase at an \( \text{O.D.}_{600} \approx 0.500 \). Expression of Tsr from AS2002 was induced with 100 \( \mu \)M IPTG (121) at \( \text{O.D.}_{600} \approx 0.300 \) and allowed to continue growth to \( \approx 0.500 \). Expression of LsrB from AS2014 was not induced. Cells expressing plasmid pBAD-\( \text{lsrB} \) were prepared similar to cells expressing pCA24N-\( \text{lsrB} \) with the exception that expression of pBAD-\( \text{lsrB} \) was induced with 0.2\% \( \text{L} \)-arabinose at the time of back dilution from overnight culture. Cell cultures of 5 mL were centrifuged at 600 x g for 10 minutes to separate the soft pellet from the growth media. The pellet was gently resuspended in an equal volume of CB and incubated for fifteen minutes of recovery time in a 30°C rotor set to low speed. The resuspended cells were used for the assay. Motility of the resuspended cells was confirmed through phase contrast microscopy.

Capillary assays used plastic gaskets sandwiched between a glass microscope slide and glass coverslip to create the pond chamber (Figure 9). About one sixth of the gasket was excised to provide an insertion point for both cells and capillaries into the chamber. A 1 microliter volume capillary was flame-sealed at one end and lowered into a solution containing the desired concentration of attractant to be tested. The CB-buffered attractant was pulled several millimeters into the capillary.

Capillaries contained either CB alone or CB supplemented with the different concentrations of \( \text{L} \)-serine or AI-2. The cells were inserted into the chamber ponds via gentle pipetting to avoid bubbles or flagellar shearing. Capillaries were then inserted through the opening of the chamber and cells were incubated at 30°C for 45 minutes. Diffusion of the chemoeffector from the open end of the capillary into the pond created a gradient that could be sensed by the cells (115). These cells migrated up the gradient
and into the capillary mouth if the attractant signal was strong enough. Once incubation finished, the capillaries were removed and their contents expelled into CB for dilution. Dilutions were plated on nutrient agar supplemented with appropriate antibiotic and allowed to grow at 37°C overnight. Colonies were counted and the number of cells that entered the capillaries was calculated. All data was normalized to CB background.

Computer generated docking model and residue selection. The MBP-Tar model (122) was used as a guide to generate a computerized docking model for LsrB-Tsr. SPOCK software (108) was used to align amino acid residues on the surface of AI-2-bound LsrB (95) (Protein Data Bank ID=1TJY) believed to be in contact with Tsr, based on the crystal structure for E. coli Tsr (107) (PDB ID=2D4U). Also taken into account were unpublished findings that replacements at particular residues in LsrB altered chemotaxis to AI-2. Care was taken to ensure these residues were oriented for close contact with Tsr. Molecular dynamics calculations were done by using AMBER (123) software to simulate docking, following the removal of all overlap between the Tsr and LsrB structures. A distance-dependent dielectric field, at an initial temperature of 300 Kelvin with a time step of 0.001 ps, was used to carry out all simulations, with the default force field suggested by the AMBER software.

Site-directed mutagenesis of tsr and lsrB. Point mutations were made in plasmid-borne tsr and lsrB genes using a site-directed PCR mutagenesis protocol closely following the protocol suggested by the Stratagene QuikChange™ II Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA). Amino acid residues in α-helix 3-4 (Gly144-Glu150) of the subunit 1 and 1’ “shoulder” regions of the Tsr dimer were targeted and individual-
ly changed to alanine or other residues using primers designed according to well established principles for mutagenesis (124). Amino acid residues Leu56-Pro65 of β-strand 2, and Asp249-Glu256 of α-helix 8 of LsrB were targeted for individual replacement using the same guidelines. Primers were synthetic oligonucleotides (Integrated DNA Technologies™) containing the desired mutation, each complementary to opposite strands of the pCA24N-\textit{tsr}, pCA24N-\textit{lsrB} and pBAD-\textit{lsrB} vectors. These primers were extended by the Pfu Ultra DNA polymerase during temperature cycling over multiple PCR cycles. Following PCR, the product is treated with the Dpn I endonuclease to eliminate methylated and hemi-methylated DNA (the parental template). Because the \textit{tsr} and \textit{lsrB} genes cloned into pCA24N and pBAD18 were derived from \textit{E. coli}, this Dpn I digest selects for mutated synthesized DNA because plasmid DNA isolated from our \textit{E. coli} strains is methylated. Mutated \textit{tsr} and \textit{lsrB} genes were then subjected to multiple rounds of PCR sequencing to confirm that only the desired point mutations were present. Plasmids containing desired mutant genes were then transformed into their competent cells (CV\textsubscript{1} Δ\textit{tsr} for \textit{tsr} and CV\textsubscript{1} Δ\textit{lsrB} for \textit{lsrB}) and frozen.

\textit{Creation of the ΔlsrB/ΔluxS strain AS2014.} Strain MJ101 (\textit{lsrB}Ω\textit{Kan}\textsuperscript{r}) was transformed with pCP20 (FLP recombinase/Cm\textsuperscript{r}) (117). Transformants were allowed to grow at plasmid’s permissive temperature (30°C) to constitutively express FLP recombinase gene product. FLP recombinase removes the kanamycin cassette during growth phase of transformants to leave a clean, non-polar deletion of the targeted gene, a result confirmed by failure of the transformants to grow on kanamycin media (50 μg/mL) during replica plating. Transformants were then grown at restrictive temperature of pCP20
(43°C) to cure the cells of the plasmid. Plasmid curing was confirmed by replica plating cells onto media supplemented with chloramphenicol (35 µg/mL) and their subsequent failure to grow. The resulting strain MJ101 was now ΔlsrB (no Kan cassette).

P1 phage transduction was performed to transduce the kanamycin cassette from a CV1 (luxSΩKan') strain to the ΔlsrB strain. The transductants were grown on media supplemented with kanamycin (50 µg/mL). Isolated transductants were subjected to PCR with primers for the kanamycin cassette gene, the lsrB gene and the luxS gene. Following PCR the products were run on a 1% agarose gel supplemented with ethidium bromide, then viewed under ultraviolet light to confirm insertion of the kanamycin cassette and removal of the luxS gene. Removal of the kanamycin cassette disrupting luxS was done as explained above. Once these results were confirmed, strain AS2014 (ΔlsrB/ΔluxS) was made competent and ready for use.

*Immunoblot analysis for LsrB expression.* Overnight cultures of CV1 ΔlsrB containing pCA24N expressing LsrB protein were diluted into 500 mL TB and grown at 30°C with shaking to O.D.₆₀₀ = 0.500. Cultures were centrifuged and the pellet was frozen at -80°C before resuspension in 25 mL lysis buffer (5 mM DTT, 50 mM Tris (pH 8.0), 300 mM NaCl, 1 mM PMSF). Resuspended pellets were then sonicated for 8 minutes (15 seconds ON / 45 seconds OFF, 37% amplitude). Centrifugation at ultra-high speed followed before the supernatant was collected in lysis buffer. This was then concentrated to 1 mL and quantified using the BCA assay. PAGE was performed on these whole cell protein preps (20 µL/well) in the presence of 10 µg DTT on 10 % Mini-PROTEAN TGX® Stain-Free™ Protein Gels (Bio-Rad). The separated proteins were
transferred onto a low-fluorescence PVDF membrane (Thermo-Fisher Scientific) at 100 volts for 1 hr using the Mini Trans-Blot® Electrophoretic Transfer Cell (Bio-Rad). The membrane was placed in blocking buffer (3% non-fat dry milk; 10 mM Tris pH 7.5, 100 mM NaCl, 0.1% Tween 20 (TBST)) and incubated with agitation for 1 hour at room temperature. It was then transferred into primary antibody solution (1:2500 Protein A purified anti-LsrB antibody in blocking buffer) and incubated with agitation for 1 hr at room temperature, followed by thorough washing in TBST (3x, 5 mins each). The membrane was then transferred into secondary antibody solution (1:25,000 anti-goat HRP (Sigma A5420) in blocking buffer) and incubated with agitation for 1 hr at room temperature, followed by thorough washing in TBST (3x, 5 mins each). The membrane was developed with Clarity™ ECL Blotting Substrate (Bio-rad) and imaged using the ChemiDoc™ Touch Imaging System (Bio-Rad). ImageLab™ software (Bio-rad) was used for total protein normalization and estimation of relative LsrB levels.

*AI-2 uptake assays with Vibrio harveyi.* Overnight cultures of LsrB variants were diluted into 15 mL of TB at 30°C with shaking to O.D.\textsubscript{600} = 0.500. Cultures were then distributed into separate tubes, AI-2 was added, and the samples were incubated at 30°C with shaking. Samples were collected every 30 min for a total of 10 hrs. Samples were centrifuged and the supernatants collected and frozen for use in the *Vibrio harveyi* TL26 bioluminescence assay. This reporter strain is disrupted for signaling pathways for acyl-homoserine lactones (AHLs) and the alpha-hydroxyketone (CAI-1), as well as for synthesis of AI-2 (ΔluxS); therefore, it responds only to exogenously added AI-2 and cannot produce AI-2 on its own. TL26 was diluted 1:5000 and 90 µL was added to each
reaction containing 10 µL of thawed samples. Bioluminescence was then measured every hour until the background began increasing. Fold induction in luminescence was measured as Relative Light Units (RLU) of sample / RLU of medium.

**Confirmation of LsrB variant response from a separate vector.** Because of the absence of tight control of the $P_{Ts}$ promoter in pCA24N, and a short C-terminal extension, key lsrB variants were constructed in an expression vector with tighter promoter control and no C-terminal extension. Site-directed mutagenesis was utilized as described previously to introduce point mutations D59A, V60A, T61A, D63A, R252A and E256A into pBAD-lsrB. These mutant proteins were then expressed through addition of 0.2% by volume L-arabinose to back diluted cultures growing at 30°C. Capillary assays were performed as described previously to measure chemotaxis to AI-2 gradients.

**Results**

**Generation of the Tsr-LsrB docking model.** Using the SPOCK and AMBER software programs, the best predicted interaction between the periplasmic domain of Tsr and the AI-2-bound LsrB binding protein was determined. This interaction and regions of interest are shown in Figure 10.

**Mutagenesis of tsr.** The effect of mutating individual residues from Gly144 to Glu150 to Ala in the “shoulder” regions of Tsr subunits 1 and 1’ was tested. These amino acid residues and their relative positions are shown highlighted in red in Figure 11. The rest of the Tsr protein remains unchanged, including the published serine binding sites.
Figure 10. The Tsr-LsrB docking model. (a) The published docking model for the *E.coli* aspartate chemoreceptor (Tar) with ligand-bound maltose binding protein (MBP) (122). The upper image shows the likely conformation the proteins adopt during interaction. MBP is shown in blue, with maltose bound at its center. The periplasmic domain of Tar is colored by subunit. The lower image shows the “tightness” of the Tar-MBP interaction. Yellow arrows indicate critical binding sites. (b) The proposed docking model for Tsr-LsrB based on the SPOCK and AMBER software employed (unpublished). The upper image shows the likely interaction dynamics, while the lower image speculates on the strength of the binding. Yellow arrows indicate predicted critical residues that strengthen the binding.
Figure 11. Tsr residues mutated based on docking model. (a) The cartoon structure of the dimerized Tsr periplasmic domain is shown. This model depicts the closed binding conformation of the receptor when serine is bound, and the monomeric subunits are defined by color. (b) The backbone structure taken from the image in (a) indicates amino acid residues predicted to be in closest contact with LsrB by highlighting them in red. This indicated portion extends from Gly144 to Glu150 of subunits 1 and 1’. Individual residues were subjected to site-directed mutagenesis to alanine or other amino acids. The serine binding pocket is unaffected by these mutations. Structures adapted from PDB ID: 2D4U (107).
Reviewing the components necessary for E. coli cells to mediate AI-2 chemotaxis.

To quantify the AI-2 response and compare it to that of L-serine, capillary assays were performed (115). CV1 and CV4 (Δtar) cells accumulated in capillaries containing either L-serine or AI-2. Cells that were Δtsr (CV5) failed to accumulate in either capillaries containing L-serine or AI-2, confirming the original finding that the Tsr receptor is necessary to mediate AI-2 chemotaxis (Figure 12). Further confirmation of earlier findings that LsrB is required for AI-2 chemotaxis was supplied by the failure of MJ101 (lsr-BΩKan') cells to accumulate in capillaries containing AI-2, although they maintained a normal response to L-serine (Figure 13). Additionally, cells deleted for one of the membrane-bound transporter proteins (lsrCΩKan') maintained the ability to respond to both L-serine and AI-2 (Figure 13).

The ability of tsr variants to mediate chemotaxis to AI-2. To quantify AI-2 response, and compare it to that of L-serine, capillary assays were again performed with the tsr variants generated from the docking model. The plasmid bearing wild-type tsr set the standard for comparison, while the CV5 (Δtsr) strain served as the negative control. The point mutations were introduced into strain AS2002, and expression of the mutant proteins was induced with 100 μM IPTG. Four of the point mutants, G144A, G146A, I148A and N149A (position 145 is an Ala), showed little to no variation in AI-2 response, with dose response curves similar to that of the wild-type tsr. The E150A point mutant was noticeably impaired for response to AI-2, though not to a significant degree. The K147A point mutant showed little to no accumulation in the capillary in the presence of AI-2 (Figure 14). The response to 10 mM serine was not inhibited with any of
Figure 12. Response of WT and variant cells to chemoattractant. Shown are the normalized values (buffer-only controls subtracted) of CV1 (WT), CV4 (Δtar) and CV5 (Δtsr) cells responding to capillaries containing either L-serine or AI-2. Incubation of cells occurred for 45 minutes at 30°C. Trials were performed in triplicate with three biological replicates each. Cells deleted for the Tsr chemoreceptor failed to respond significantly to either attractant. Cells deleted for the Tar chemoreceptor responded to gradients of AI-2 at near wild-type levels.
Figure 13. Response to chemoeffector by \textit{lsr} operon variants. Shown are normalized values of \(\Delta\text{lsrB} (\text{lsrB}\Omega\text{Kan}^\text{r})\) and \(\Delta\text{lsrC} (\text{lsrC}\Omega\text{Kan}^\text{r})\) cells responding to either \(L\)-serine or AI-2. Cells were incubated at 30°C for 45 minutes. Trials were performed in triplicate for three biological replicates. Cells lacking the AI-2 binding protein LsrB were unable to respond to a gradient of AI-2. Cells lacking one of the AI-2 transport channel proteins LsrC were able to respond normally to gradients of AI-2.
the mutant proteins except for a 10% decrease with E150A (Figure 15). Glu-150 is the residue closest to the serine-binding site.

The amino acid residues Lys147 and Glu150 were selected for charge reversal point mutation to further investigate their importance. The Ile148 residue was also selected as a control since its dose response curve showed near wild-type level response in the alanine scanning mutant testing. K147E, I148E and E150K were introduced separately into AS2002 and tested in the capillary assay for response to L-serine and AI-2. The K147E variant showed a complete lack of response to AI-2. The I148E variant responded with a near wild-type response curve, whereas the E150K variant gave a slightly weaker response than its E150A counterpart – lower than the wild-type response but not significantly (Figure 16). Again, serine response was not affected by these mutations.

The ability of lsrB variants in the N-terminal region (β-strand-2) to mediate chemotaxis to AI-2. Capillaries assays were used to measure the chemotactic ability of lsrB variants to AI-2. The docking model was again consulted to determine which amino acid residues in the N-terminal region of LsrB should be subjected to alanine-scanning mutagenesis. The plasmid bearing wild-type LsrB was used as the standard for comparison for AI-2 chemotaxis. MJ101 (lsrBΩKan5) was the negative control for chemotactic response to AI-2, showing no response in the assay. Several alanine-scanning variants in β-strand-2 showed little to no difference in dose response to AI-2 from the capillary assay. These included L56A, G57A, V58A, Y62A, G64A and P65A. The most severe dose response alteration was exhibited by the D59A variant, with elim-
Figure 14. Tsr alanine-scanning mutant response to AI-2. Background accumulations in buffer-only capillaries were between 7,000 and 15,000. Shown are the normalized values (buffer-only controls subtracted) of CV5+pCA24N-tsr (AS2002), CV5 (Δtsr) and AS2002 tsr point mutant cells responding to capillaries containing AI-2. Capillary assays were performed over a 45 minute incubation at 30°C in a slide warming chamber. Trials were performed in triplicate for three biological replicates. Tsr variants were expressed from pCA24N with 100 µM IPTG induction. The K147A variant accumulated only slightly more than the negative control. Also reduced for AI-2 chemotaxis was the E150A Tsr variant. All other point mutants responded at near wild-type levels.
Figure 15. Tsr alanine-scanning mutant response to 10 mM L-serine. Background accumulations in buffer-only capillaries were between 7,000 and 15,000. Assays were carried out in triplicate for three biological replicates. Cells were incubated for 45 minutes at 30°C. Variants were induced at 100 µM IPTG from plasmid pCA24N. Capillary assays were performed over a 45 minute incubation period at 30°C.
Figure 16. Tsr charge reversal mutant response to AI-2. Background accumulations in buffer-only capillaries were between 7,000 and 15,000. Charge reversals of three selected residues were carried out and variants were induced from plasmid pCA24N with 100 μM IPTG. K147E loses any ability to mediate AI-2 chemotaxis. E150K response to AI-2 is relatively unchanged from E150A. Capillary assays were performed over a 45 minute incubation period at 30°C. Trials were carried out in triplicate for three biological replicates.
ination of the dose response to varying degrees in the V60A, T61A and D63A variants (Figure 17).

*The ability of lsrB variants in the C-terminal region (α-helix-8) to mediate chemotaxis to AI-2.* Amino acids in α-helix-8 of the LsrB C-terminus were subjected to alanine-scanning mutagenesis based on the docking model. These variant responses were again compared to the chemotactic response of wild-type LsrB expressed from plasmid pCA24N, and negative control MJ101. Based on the results of these capillary assays, only the N249A mutant retained a wild-type dose response curve. The R252A and E256A variants showed significantly altered response compared to wild-type, and the R252A/E256A double mutant failed to respond (Figure 18). Charge reversal mutations were then introduced at these positions both singly and in a tandem. Both the R252E and E256R variants showed a severe alteration in the AI-2 response compared to wild-type, with the double mutant (R252E/E256R) losing all ability to respond to AI-2 in the capillary assay (Figure 19).

*Ability of LsrB variants to transport AI-2 in uptake assay.* Each of the LsrB variants was tested for its ability to participate in AI-2 uptake using the *Vibrio harveyi* TL26 bioluminescence assay. All variants in β-strand-2 showed a decrease in bioluminescence similar to wild-type LsrB between the 3 to 4 hour time points (Figure 20). AI-2 uptake was not hindered in the D59A, V60A, T61A or D63A AI-2 chemotaxis mutants. Likewise, variants in α-helix-8 were not hindered in their ability to transport AI-2 relative to wild-type (Figure 21). Only double mutants T61A/D63A and R252A/E256A showed
Figure 17. LsrB β-strand 2 variant response to AI-2. Background accumulations in buffer-only capillaries were between 7,000 and 15,000. Most variants retain a wild-type dose response curve. Variants D59A, V60A, T61A and D63A, all show a drastically different dose response to a range of AI-2. The D59A variant loses nearly all of its ability to respond to AI-2 via chemotaxis. No induction was necessary as uninduced plasmid-born LsrB levels were similar to those of chromosomally produced LsrB. Capillary assays were performed over a 45 minute incubation period at 30°C. Trials were carried out in triplicate with three biological replicates.
Figure 18. LsrB α-helix 8 variant response to AI-2. Background accumulations in buffer-only capillaries were between 7,000 and 15,000. Most variants retain a wild-type dose response curve. Notable variants include R252A, E256A and the R252A/E256A double mutant, all of which show a lowered dose response to a range of AI-2. No induction was necessary as uninduced plasmid-born LsrB levels were similar to those of chromosomally produced LsrB. Capillary assays were performed over a 45 minute incubation period at 30°C. Trials were carried out in triplicate with three biological replicates.
Figure 19. LsrB α-helix 8 charge reversal variant response to AI-2. Shown are the selected point mutants that were subjected to charge reversal. No induction from pCA24N was necessary since LsrB levels were similar to chromosomal levels. Background accumulations ranged between 7,000 to 10,000 cells in buffer only. Cells were incubated at 30°C for 45 minutes. Trials were carried out in triplicate for three biological replicates. Charge reversal at R252 and E256 severely reduced AI-2 chemotaxis at higher concentrations and eliminated it at lower. A charge reversal double mutant completely lost ability to mediate AI-2 chemotaxis.
Figure 20. Ability of LsrB β-strand 2 variants to uptake AI-2. Shown is the fold induction in luminescence in *Vibrio harveyi* TL26 following addition of *E. coli* supernatant from various growth time points. Bioluminescence is measured in relative light units (RLU) and is defined as RLU of the sample / RLU of the culture medium. All LsrB β-strand 2 point mutants show the same reduction in luminescence profile as that observed with wild-type (~2 to 4 hours). As a negative control, the ΔlsrB mutant continued to induce bioluminescence out to nearly 8 hours.
Figure 21. Ability of LsrB α-helix 8 variants to uptake AI-2. Shown is the fold induction in luminescence in *Vibrio harveyi* TL26 following addition of *E. coli* supernatant from various growth time points. Bioluminescence is measured in relative light units (RLU) and is defined as RLU of the sample / RLU of the culture medium. All LsrB α-helix 8 point mutants show the same reduction in luminescence profile as that observed with wild-type (~2 to 4 hours). As a negative control, the ΔlsrB mutant continued to induce bioluminescence out to nearly 8 hours.
any difference in uptake, with a delay of approximately one hour, though uptake still occurred (Figure 22).

*Ability of key lsrB variants to respond to only exogenously sensed AI-2.* To ensure mutant response was to AI-2 sensed from the outside environment, plasmids carrying key *lsrB* variants were transformed into cells deleted for the *luxS* gene. LuxS is involved in the production of AI-2 from S-adenosylhomocysteine, and deleting it removes the cell’s capability of producing any endogenous AI-2. This serves the purpose of confirming that certain LsrB variants are not simply being overwhelmed by endogenously produced AI-2 within the cytoplasm and being sequestered away to give the negative chemotaxis result. In a Δ*luxS* background, D59A, T61A, D63A and R252A showed no AI-2 chemotaxis improvement in the capillary assay when compared to strains carrying intact *luxS*. Additionally, wild-type cells deleted for *luxS* were still able to mediate chemotaxis to AI-2 (Figure 23).

*Complementation of lsrB variants in a wild-type background.* In an attempt to determine if any of the LsrB variants deficient for AI-2 chemotaxis could be complemented by the presence of wild-type protein, I transformed these variants into wild-type CV1 cells and tested their ability to respond to AI-2 in the capillary assay. Protein levels of chromosomally-encoded and plasmid-encoded wild-type LsrB were already found to be nearly identical, so induction with IPTG was not necessary. Interestingly, the dose response of the LsrB mutant D59A was not restored to wild-type level when the wild-type LsrB protein was present (Figure 24). The V60A variant dose response became similar to wild-type, while the dose responses of the other variants tested fell somewhere
Figure 22. Ability of *E.coli* cells carrying an LsrB double mutant to uptake AI-2. Shown is the fold induction in luminescence in *Vibrio harveyi* TL26 following addition of *E. coli* supernatant from various growth time points. Bioluminescence is measured in relative light units (RLU) and is defined as RLU of the sample / RLU of the culture medium. Both the T61/D63A and R252A/E256A double mutant show a delay in quenching of bioluminescence in *V. harveyi* of approximately one hour relative to wild-type.
Figure 23. AI-2 chemotaxis by LsrB variants in a \( \Delta luxS \) background. Key LsrB variants were expressed without induction from plasmid pCA24N in cells carrying a chromosomal deletion for the \( luxS \) gene (strain AS2014). Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
Figure 24. AI-2 chemotaxis by LsrB variants in a CV1 background. Key LsrB variants were expressed without induction from plasmid pCA24N in cells carrying a chromosomal wild-type copy of LsrB. Response to AI-2 gradients was rescued to some degree for most variants, with V60A being rescued to nearly wild-type levels. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
in between. These results suggest the D59A mutant may exhibit a dominant-negative phenotype, a condition not unlike that described in previous studies with MBP and Tar (122).

**Ability of pBAD-LsrB variants to mediate response to AI-2 gradients.** LsrB variants expressed under control of the $P_{BAD}$ promoter were exposed to serine and AI-2 gradients in the capillary assay. The accumulation of $lsrB$ variant cells above background levels was within 97% of the response measured for those same over-expressed mutants carrying the C-terminal extension in pCA24N-$lsrB$ (Figure 25). There was no significant variation in AI-2 response in the presence or absence of the C-terminal extension.

**Discussion**

The docking of ligand-bound maltose binding protein (MBP) to Tar has been investigated extensively (105, 122, 125-127). The resulting docking model has provided an excellent point of reference for our own proposed docking model of LsrB with Tsr. Because of the large degree of sequence and structural similarity between Tsr and Tar, orienting a binding protein for potential interaction with Tsr can be patterned somewhat after the MBP-Tar model. Using this published model a proposed docking model for the interaction of ligand-bound LsrB and the periplasmic domain of Tsr was generated (Figure 10). Residues of interest were highlighted for genetic analysis. Figure 26 illustrates the location of the amino acid residues of interest in both the N-terminal and C-terminal regions of the ligand-bound conformation of LsrB. Figure 11 indicates residues of particular interest in the shoulder regions of the Tsr homodimer.
Figure 25. Response to AI-2 by pBAD-*lsr*B variants. Key LsrB variants were cloned into pBAD. Mutants were induced with 0.2% arabinose for full induction. Each key alanine substitution showed the same chemotaxis to AI-2 as it did when expressed from pCA24N. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
Figure 26. Structure of *E. coli* LsrB with residues of interest. This structure of the LsrB binding protein is adapted from the published LsrB of *Salmonella typhimurium* (95). The β-strand 2 residues D53-P65 are highlighted in red in the N-terminus, while residues N249, R252 and E256 of α-helix 8 are highlighted in red in the C-terminus of the protein. Each of these residues was chosen due to its proximity and likelihood of binding calculated from the generated Tsr-LsrB docking model.
Alanine scanning mutagenesis of both proteins revealed particular amino acid residues that are critical for AI-2 chemotaxis to occur. In Tsr the positively charged lysine at position 147 and negatively charged glutamate at position 150 must be present for E. coli cells to perform chemotaxis to AI-2. Mutating these residues to alanine greatly depresses chemotaxis to AI-2, but has little effect on response to serine, leading us to conclude these alterations likely do not change the overall structure or function of the Tsr protein, only its ability to interact with LsrB and signal. The fact that these are charged residues led us to surmise that electrostatic forces were likely at work during the interaction with LsrB. I tested this theory by introducing charge reversals at these positions. The result was a complete elimination of AI-2 response in cells carrying the K147E mutation, and a further attenuation of response in cells with the E150K mutation. Reversing the charge at 147 likely resulted in electrostatic repulsion between the two proteins, causing the failure of the cells to respond to AI-2. Because Tsr is a homodimer, lysine-147 is present in both shoulder regions and likely serves as an anchoring point for the interaction with LsrB during chemotaxis.

In LsrB the critical amino acids identified during the alanine-scanning mutagenesis were a mix of charged and non-polar residues. Asp-59, Val-60, Thr-61, Asp-63, Arg-252 and Glu-256 each seem to play a role in supporting chemotaxis to AI-2. Changing each of these to Ala negatively affected AI-2 response to some degree, with D59A showing the greatest deficiency. The T61/D63 and R252A/E256 double mutants were removed from consideration due to the delay they showed in transporting AI-2 into the cell. This delay compared to wild-type is a likely indicator that these proteins either
couldn’t bind AI-2 well, or couldn’t interact with the LsrACD transport channel to import AI-2. Based on the location of the negatively charged aspartate residues (D59 and D63) in β-strand 2, they are likely candidates to electrostatically interact with the positively charged Lys-147 of Tsr on one homodimer shoulder. In α-helix 8 is where the negatively charged Glu-256 possibly interacts with the other Lys-147 of the opposite shoulder in Tsr. Due to their proximity, mutating the Val-60 and Thr-61 residues in β-strand 2 of LsrB may affect the positioning of the Asp-59 and/or Asp-63 residues needed for interaction with Tsr. Perhaps the alteration of these LsrB residues disrupts the tight interaction with Tsr, resulting in an inability to propagate a signal through transmembrane helix-2 of the chemoreceptor. Additionally, the positively charged Arg-256 in α-helix 8 may stabilize the LsrB-Tsr docking by interacting with the aforementioned Glu-150 of Tsr. These results lend credibility to our proposed docking model and elucidate the nature of the protein-protein interaction taking place during chemotaxis to AI-2.

Of obvious note is the wide range of expression levels detected in the LsrB mutants expressed from pCA24N (data not shown). While these results are curious, they likely do not affect AI-2 chemotaxis. The fact that each of the LsrB mutants mentioned here maintains its ability to import AI-2 into the cell, according to the *Vibrio harveyi* bioluminescence assay, indicates the reduction in AI-2 chemotaxis is not simply the result of altering the export and subsequent periplasmic concentration of LsrB with the C-terminal extension. In other words, they perform their native function at wild-type levels, thus the mutations affect only chemotaxis. This is further supported by the fact that
LsrB variants expressed from the pBAD plasmid respond to AI-2 at identical levels of those from the pCA24N plasmid.

While the protein expression levels are curious, it is certainly possible that AI-2 chemotaxis is not directly correlated to LsrB expression in *E. coli*. Further investigation into this issue will be discussed in Chapter V. Overall, it appears that docking of LsrB to Tsr and signaling during AI-2 chemotaxis relies primarily on tight interaction with the N-terminal domain, specifically β-strand 2 of LsrB. The C-terminal domain, specifically α-helix 8, likely serves to further stabilize the interaction and facilitate signaling.

Furthermore, the ability of the cells to produce their own AI-2 through the activity of LuxS doesn’t appear to affect their response to extracellular AI-2. Key LsrB mutants expressed in a LuxS deletion strain showed no variation in AI-2 response compared to strains with intact LuxS. Production of AI-2 in the cytoplasm and its subsequent export apparently does not inhibit those cells from recognizing and responding to environmental AI-2. This may be due to the fact that by the time cells are producing enough AI-2 themselves, the chemotaxis response to environmental AI-2 has already occurred.

Lastly, when attempts to complement mutant LsrB proteins with copies of wild-type LsrB were made, the D59A and D63A variants failed to complement. The reason for this is unclear, though perhaps they represent dominant negative phenotypes, where the mutant proteins are able to dock with Tsr but are unable to elicit signal propagation through the chemoreceptor. If this is true, the interaction is still tight enough to block wild-type LsrB from complementing the mutation. The investigation of these mutants will be further discussed in Chapter V.
CHAPTER III
CORRELATING LSRB EXPRESSION LEVELS WITH AI-2 CHEMOTAXIS

Introduction

The expression of the periplasmic LsrB protein is regulated within the *lsr* operon of *E. coli*. When population density is low, and a quorum sensing threshold of AI-2 is not achieved in the environment, the *lsr* operon is repressed by the LsrR protein. Following sufficient addition of AI-2 to the local environment, this LsrR repression is relieved by interaction with phosphorylated AI-2 (AI-2-P). Thus, the operon produces LsrB, which in turn binds periplasmic AI-2 and transports it into the cytoplasm via the LsrACD transporter, resulting in the presence of more AI-2-P to relieve repression of the operon. It is not until the last components of the operon (LsrFG) are made in sufficient quantities that AI-2-P is deconstructed and recycled, returning the operon to a repressed state (in concert with waning levels of environmental AI-2 presumably).

As I have shown in the previous chapter, wild-type levels of chromosomal LsrB are sufficient to elicit a chemotaxis response to the *R*-THMF form of AI-2 in *E. coli* (Figure 12). I wanted to determine (a) how much LsrB was produced in cells at various growth phases and its correlation to AI-2 chemotaxis, and (b) what affect stepwise induction of LsrB expression from a plasmid had on the ability of the cells to perform chemotaxis to AI-2. For the first question, cells were grown to increasing optical densities before protein levels and chemotaxis response were measured. Log phase growth in *E. coli* is considered to begin when cells reach an O.D.<sub>600</sub> just below 0.3; it progresses to
mid-logarithmic growth near 0.5; and enters late logarithmic growth between 0.7 and 0.9. During early log growth, cells would likely produce little AI-2, as the components of the \textit{lsr} operon are not being highly expressed. During mid-logarithmic growth, cells would begin expressing more of the LsrB binding proteins in order to transport AI-2 into the cells, leading to more expression of the \textit{lsr} operon. Once they reach later log growth, cells should have high amounts of AI-2 present.

In the second scenario, an induction series was performed on plasmid-borne LsrB and response to AI-2 was measured in the capillary assay. I found that increasing LsrB within the cells correlated negatively with AI-2 chemotaxis, though there was little effect on serine chemotaxis, and optimal response to AI-2 as a chemoattractant occurred during mid-log phase growth.

Furthermore, I hypothesized that LsrB expression would change when different components of the \textit{lsr} operon were independently deleted. For instance, I expected more LsrB to be present when LsrR, the operon repressor, was deleted. Conversely, I surmised that deleting components of the ABC transporter LsrACD or the AI-2 degradation pathway LsrFG would have little to no effect on expression of LsrB. I also tested the effect of deleting LuxS and TqsA on LsrB expression. Likewise, I investigated whether deletions of these different genes affected chemotaxis to AI-2, and if these effects could be correlated to LsrB expression in these mutants. Lastly, I tested effect on AI-2 chemotaxis in cells at various growth phases when the \textit{luxS} gene was deleted. Deletion of the LuxS enzyme ensures the cells are not producing endogenous AI-2, allowing us to ex-
amine whether accumulation of endogenous AI-2 at earlier and later phases of growth influenced response to exogenous AI-2 added to the medium.

I found there were few variations in LsrB expression in the individual deletion mutants. I also found there was little to no difference in the AI-2 dose response curves of the individual deletion mutants, whether the component deleted was involved in synthesis of AI-2, export of AI-2, or even import of AI-2. Additionally, I saw a decrease in AI-2 response at later growth phases compared to mid-log phase growth, and observed strains carrying the luxS deletion varied little in chemotaxis response from their counterparts able to produce their own AI-2. Overall, expression of LsrB did not seem to correlate to AI-2 chemotaxis.

Materials and methods

Bacterial strains, plasmids and media. Wild-type strain CV1 was used for the growth phase tests. Deletion mutant strains were derived from CV1 using phage transduction to disrupt target genes with a kanamycin resistance cassette before its removal as previously described (43). Cassette was then removed by transforming cells with the pCP20 plasmid encoding an FLP recombinase (117). The pCP20 plasmid was removed through a temperature-sensitive growth restriction and PCR was used to verify a clean gene deletion with removal of the kanamycin resistance cassette. Table 2 lists the target genes deleted for use in this study. IPTG-inducible plasmid pCA24N was employed to produce stepwise increases in LsrB expression in strain MJ101. All strains were grown in tryptone broth media (10 g/L tryptone; 8 g/L NaCl) supplemented with antibiotic
where appropriate. IPTG was added to inducible strains when optical density reached 0.300 at wavelength 600 nm.

*Measuring chemotaxis to AI-2 and serine at different growth phases.* Strain CV1 was grown overnight in TB supplemented with streptomycin (50 µg/mL), then back-diluted into four separate flasks to a turbidity of approximately 0.05 at O.D.₆₀₀ in 25 mL tryptone broth without antibiotic. Cells were then grown at 30°C with shaking to O.D₆₀₀ = 0.3, 0.5, 0.7 and 0.9 respectively. All densities were either concentrated or diluted to O.D₆₀₀ = 0.5 with chemotaxis buffer for uniform density upon addition to the cell reservoirs. Motility was confirmed through microscopy. Capillary assays (115) were used to quantify response to AI-2 and L-serine at the four different growth phases. Centrifugation at 600 x g for 10 minutes separated the cell pellet from the growth media. After removal of the supernatant, cells were gently resuspended in an equal volume of chemotaxis buffer and given fifteen minutes of recovery time in a 30°C rotor. Motility was again confirmed through microscopy before use in the capillary assay. Cells were exposed to gradients of 1 mM, 100 µM, 10 µM and 1 µM AI-2 in chemotaxis buffer.

*Measuring chemotaxis to attractants in the deletion strains.* Capillary assays (115) were chosen to quantify the chemotaxis responses to AI-2. Overnight cultures of cells deleted for various AI-2 pathway genes were grown at 30°C in TB supplemented with streptomycin (50 µg/mL). Cultures were back-diluted to a turbidity of ~0.05 at O.D.₆₀₀ in 25 mL TB without antibiotic and grown to mid-logarithmic phase at an O.D₆₀₀ = ~0.5. Cells were then centrifuged at 600 x g for 10 minutes to separate the soft pellet from the growth media. The supernatant was removed and the pellet then gently
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<th>Strain</th>
<th>Target Gene</th>
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<td>Escherichia coli</td>
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<tr>
<td>CV1&lt;sup&gt;a&lt;/sup&gt;</td>
<td>&lt;i&gt;lsrA&lt;/i&gt;</td>
<td>Kinase for LsrACD transporter</td>
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<td>Periplasmic binding protein for AI-2</td>
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<td>&lt;i&gt;luxS&lt;/i&gt;</td>
<td>Synthesis of AI-2 precursor DPD</td>
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<td>&lt;i&gt;tqsA&lt;/i&gt;</td>
<td>Export of AI-2 from the cell</td>
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<sup>a</sup>Kanamycin resistance cassettes were transduced to disrupt individual genes and then removed by FLP recombinase via temperature sensitive plasmid pCP20 (117). Deletion of the gene of interest and removal of the kanamycin cassette were confirmed through PCR.
resuspended in chemotaxis buffer as described previously. The cells were given fifteen minutes of recovery time in a 30°C rotor set to low speed before use. Motility of washed cells was confirmed through phase contrast microscopy before their addition to the cell reservoirs for exposure to attractant-laden capillaries.

**Immunoblots for LsrB expression.** Whole cell protein preps were performed. Overnight cultures were diluted into 500 mL TB and grown at 30°C with shaking to O.D.<sub>600</sub> = 0.5. Cultures were centrifuged and the pellet was frozen at -80°C before being resuspended in 25 mL lysis buffer (5 mM DTT, 50 mM Tris (pH 8.0), 300 mM NaCl, 1 mM PMSF). Resuspended pellets were then sonicated for 8 minutes (15 seconds ON / 45 seconds OFF, 37% amplitude). Centrifugation at ultra-high speed followed before the supernatant was collected in lysis buffer. This was then concentrated to 1 mL and quantified using the BCA assay. Polyacrylamide gel electrophoresis was used with purified LsrB antibody to compare LsrB levels of growth phase variants and mutants to wild-type LsrB expressed from either the chromosome or the pCA24N plasmid.

**IPTG induction series.** Strain CV1 with the lsrB deletion, carrying the pCA24N-lsrB plasmid, was grown overnight in TB supplemented with chloramphenicol (35 µg/mL) to select for the plasmid. The overnight was back-diluted into 25 mL of TB and grown at 30°C with shaking (250 rpm) to O.D.<sub>600</sub> = 0.3. At this point, varying concentrations of IPTG (purchased from Fisher Scientific) were added to the culture to induce lsrB expression. Concentrations of IPTG used in the series included 0 µM, 1 µM, 10 µM, 100 µM and 1000 µM. Cells were then returned to growth at 30°C until reaching an
O.D. of 0.5. The cultures were then prepared for use in the capillary assay as described previously, and capillary assays were performed to measure chemotaxis ability to AI-2. Western blots were carried out as previously described to measure LsrB expression with regard to increasing concentrations of IPTG.

**Results**

*Growth phase effects on AI-2 chemotaxis in wild-type cells.* Growth of CV1 cells to varying phases showed obvious effects on AI-2 chemotaxis. All growth phases (O.D. = 0.3, 0.5, 0.7 and 0.9) were tested at an O.D. = 0.5 for uniform testing in the capillary assay. Because there is some evidence that *E.coli* cells begin to accumulate endogenously produced AI-2 in early log-phase growth (Sneha Jani, unpublished), I tested from this point to late log phase growth. At O.D. = 0.3 the dose response never saw an accumulation of more than 21,000 colony-forming units (CFUs) at any of the tested concentrations of AI-2. This represented only 22% the response of cells at that same concentration during mid log phase growth, with the accumulation even lower at other points along the dose response curve. Background accumulation of CFUs was also nearly 3 times lower at this growth phase compared to mid log, despite the densities being identical at the time of the assay. At O.D. = 0.7, peak accumulation reached 33,000 CFUs (1 mM AI-2), approximately 35% of the response at the same concentration in mid log phase. At O.D. = 0.9 the maximum response was observed at 1 mM AI-2 and numbered 17,500 CFUs. This was calculated to be 18% of the CFUs accumulated at mid log phase (Figure 27).
Figure 27. Response of log phase growth variants to AI-2 gradients. Cells were grown to appropriate optical density, washed, and then resuspended in chemotaxis buffer to a final O.D.₆₀₀ = 0.500. Capillary assays were incubated at 30°C for 45 minutes, and data points represent three biological replicates with each concentration done in triplicate.
**Growth phase effects on L-serine chemotaxis in wild-type cells.** Effects on chemotaxis to L-serine across the various growth phases tested were less extreme. However, at O.D.\(_{600} = 0.3\) the serine dose response curve was noticeably depressed compared to mid log phase (Figure 28). Peak accumulation to serine at mid log phase occurred at 1 mM (97,000 CFUs), but showed just 71,000 CFUs in cells grown to O.D.\(_{600} = 0.3\), 73% of the accumulation at 0.5. The dose response curves for the remaining densities (0.7 and 0.9) were nearly identical to the mid log phase dose response curve (Figure 28).

**Growth phase effects on AI-2 chemotaxis in ΔluxS cells.** When exposed to exogenous AI-2 in a ΔluxS background, early log phase cells (O.D.\(_{600} = 0.3\)) and late log phase cells (O.D.\(_{600} = 0.7\) and 0.9) all showed peak accumulation at 1 mM, though none garnered much more than 30,000 CFUs, approximately one-third the response by mid log phase cells at the same concentration (Figure 29). Response by these same growth phase variants to L-serine (Figure 30) generated a dose response curve nearly identical to that previously shown for wild-type cells.

**IPTG induction of pCA24N-lsrB and its effect on LsrB expression and AI-2 chemotaxis.** CV1 (ΔlsrB) was transformed with pCA24N-lsrB and induced with varying concentrations of IPTG. Western blots indicated that the amount of LsrB produced varied with increasing concentration of IPTG added during early log phase cell growth (Figure 31). At 1 and 10 µM IPTG induction, the amount of LsrB was actually less than uninduced. The increases in LsrB levels were significant at 100 µM IPTG, producing nearly 3 times that of uninduced. At 1 mM IPTG, LsrB expression was approximately 6 times that of un-induced wild-type.
Figure 28. Response of log phase growth variants to serine gradients. Cells were grown to appropriate optical density, washed, then resuspended in chemotaxis buffer to a final O.D.⁶₀₀ = 0.500. Peak accumulation occurred at 1 mM serine concentration. At an earlier stage of growth, E.coli CV1 cells responded at 73% of the response at the same serine concentration along the dose response curve at mid log phase (0.500). Later log stages of growth were at or near mid log phase dose response. Capillary assays were incubated at 30°C for 45 minutes, and data points represent three biological replicates with each concentration done in triplicate.
Figure 29. Response of ΔluxS log phase growth variants to AI-2 gradients. Cells were grown to appropriate optical density, washed, then resuspended in chemotaxis buffer to a final O.D.₆₀₀ = 0.500. Peak accumulation occurred at 1 mM AI-2 concentration. Capillary assays were incubated at 30°C for 45 minutes, and data points represent three biological replicates with each concentration done in triplicate.
Figure 30. Response of ∆luxS log phase growth variants to serine gradients. Cells were grown to appropriate optical density, washed, then resuspended in chemotaxis buffer to a final O.D.₆₀₀ = 0.500. Peak accumulation occurred at 1 mM serine concentration. At an earlier stage of growth, cells responded at 73% of the response at the same serine concentration along the dose response curve at mid log phase (0.500). Later log stages of growth were at or near mid log phase dose response. Capillary assays were incubated at 30°C for 45 minutes, and data points represent three biological replicates with each concentration done in triplicate.
Figure 31. Western blots for IPTG induction of LsrB expression. Plasmid pCA24N was used to express LsrB with varying amounts of IPTG induction during growth. Induction ranged from 0 µM to 1000 µM IPTG, and whole cell extracts were prepared. Samples were detected with anti-LsrB antibody and LsrB levels were calculated based on chemiluminescence intensity. (a) Western blot image shows little LsrB production at 1 and 10 µM IPTG induction, while bands are significantly darker for 100 and 1000 µM IPTG induction compared to uninduced samples from the same plasmid. (b) LsrB levels relative to undistributed samples were estimated. At 1 and 10 µM IPTG induction, LsrB levels are less than 50% of uninduced amount. At 100 µM IPTG induction, there is approximately 3 times as much LsrB produced compared to uninduced. At 1000 µM IPTG induction, there is approximately 6 times as much LsrB produced compared to uninduced sample.
The chemotaxis assays for these inductions exhibited a negative correlation between amount of LsrB induced and ability of the cells to respond to AI-2 (Figure 32). The dose response curves were similar between the un-induced and 1 µM induction strains, though they shifted significantly beyond 1 µM induction. Without induction, cells carrying plasmid-borne LsrB accumulated approximately 96,000 colony-forming units (CFUs) compared to approximately 95,000 CFUs of the 1µM induction strain at 1 mM AI-2. They decreased similarly along the same dose response curve with decreasing AI-2 concentration. The difference in dose response at 10 µM IPTG was significantly lower, with no single accumulation garnering more than 78% of the un-induced value. The results were even more drastic at 100 µM and 1 mM IPTG induction, with no single response point for either garnering more than 33% and 4%, respectively, of un-induced response. As a control, 100 mM L-serine was used in the capillary assay as well. All of the induction strains accumulated CFUs of a similar number as the un-induced strain.

**Effect of single gene knockouts on LsrB expression.** The LsrB gene product is the fourth gene in the *lsr* operon to be transcribed during expression, preceded by the components of the ABC transporter – LsrACD. The expression of LsrB in the absence of both upstream and downstream flanking genes in the *lsr* operon was investigated. Figure 33 shows the relative LsrB protein levels in the various deletion mutant backgrounds. The amount of LsrB produced was noticeably higher than wild-type in the Δ*lsrR* knockout and slightly lower in the Δ*lsrK* and Δ*luxS* knockouts.

**Effect of single gene knockouts on AI-2 chemotaxis.** In addition to investigating
Figure 32. IPTG induction series response to AI-2. LsrB was expressed from pCA24N using various amounts of IPTG (0 to 1000 µM) during early log phase growth, then exposed to AI-2 gradients after growth to mid log phase. Serine response was not altered to any significant degree by overexpression of LsrB in this manner.
Figure 33. Expression of LsrB in select deletion mutants. Strain CV1 was deleted for each of the components of the *lsr* operon, as well as genes involving AI-2 synthesis (*luxS*) and export (*tqsA*). Western blots were performed on whole cell extracts of deletion mutants using anti-LsrB antibody. Most knockouts showed wild-type levels of LsrB. Knockouts for *lsrK* and *luxS* showed reduced expression of LsrB relative to wild-type. Knockouts for *lsrG* and *lsrR* showed increased expression. Deleting the repressor LsrR boosted LsrB expression the most of any of the deletion mutants.
how the LsrB levels within the cell are affected when flanking \textit{lsr} operon genes are deleted, I also tested the ability of these knockout mutants to perform AI-2 chemotaxis in the capillary assay. The wild-type strain produced its maximum response at 1 mM AI-2, with 95,000 CFUs, then decreased to 86,000 CFUs at 100 \(\mu\)M, 66,000 CFUs at 10 \(\mu\)M and 6,000 at 1 \(\mu\)M. Most of the deletion mutants in the primary transcription unit of the \textit{lsr} operon (\textit{lsrACDBFG}) produced similar dose response curves to AI-2 compared to wild-type \textit{E.coli} (Figures 34 and 35). Only the \(\Delta\text{lsrD}\) strain showed a higher response at each point along the dose response curve compared to wild-type, yet even that increase was minimal at no more than 10\%. The other \textit{lsr} operon transcription unit knockouts, consisting of \(\Delta\text{lsrR}\) and \(\Delta\text{lsrK}\), along with both \(\Delta\text{luxS}\) and \(\Delta\text{tqsA}\) were observed for AI-2 chemotaxis. Only the \(\Delta\text{lsrK}\) mutant deviated from wild-type along the dose response curve (Figure 35). Though similar to wild-type with 92,000 CFUs at 1 mM AI-2, it dropped to approximately 84\% of wild-type at 100 \(\mu\)M AI-2 and only 71\% at 10 \(\mu\)M AI-2 before bottoming out at 1 \(\mu\)M AI-2. The \(\Delta\text{lsrR}\) mutant actually showed slightly higher dose response than wild-type, though not significant at any point along the curve.

**Discussion**

The expression of LsrB is regulated by the \textit{lsr} operon and the presence of AI-2. When present at sufficient concentration in the periplasm, AI-2 is bound to LsrB which aids in importing AI-2 into the cytoplasm via the membrane transporter comprised of LsrA, LsrC and LsrD. Once internalized, AI-2 is phosphorylated by the kinase LsrK. It then binds to the \textit{lsr} operon repressor LsrR, relieving repression of \textit{lsrACDBFG}. LsrF
Figure 34. Response to AI-2 by lsr operon knockouts. Components of the \textit{lsrACDBFG} operon were deleted individually in CV1, then tested for ability to respond to AI-2 gradients. All knockouts were able to perform chemotaxis at wild-type levels to AI-2, the exception being the \textit{lsrB} deletion. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
Figure 35. Response to AI-2 by *lsr*-associated knockouts. Regulatory genes *lsrK* and *lsrR*, as well as AI-2 synthesis protein LuxS and AI-2 exporter TqsA were deleted individually in CV1, then tested for ability to respond to AI-2 gradients. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
and LsrG are involved in breaking down phosphorylated AI-2 so it doesn’t accumulate to high levels in the cytoplasm. Because of the feedback loop created by the presence of AI-2, altering any one of these components should have an effect on the levels of LsrB in the cell, and likewise the ability of the cells to respond to AI-2 gradients via chemotaxis.

Our approach to investigate the correlation between abundance of LsrB and chemotaxis response was three-fold. First, I checked to see if cells naturally produce more LsrB by growing them to later log phase. Our western blots for protein levels confirmed expression of LsrB increases with growth phase, with approximately 300 copies per cell present during mid logarithmic growth (S. Jani and M. Manson, unpublished), so I checked the chemotaxis ability of these growth variants in the capillary assay. By growing wild-type cells to different densities and introducing them in the chemotaxis assays at a uniform density, I restricted the variables to growth phase rather than growth phase and density combined. Contrary to our hypothesis, allowing more LsrB production had a negative effect on chemotaxis to AI-2. Optimal AI-2 response occurred at mid log phase and decreased significantly with later phases of growth. I considered this may be due to the accumulation of endogenous AI-2 in the periplasm, so I repeated the assays with cells unable to produce their own AI-2 (ΔluxS). These cells would be exposed only to exogenous AI-2. The fact that these luxS mutants responded to the same degree as wild-type cells at each growth phase leads us to believe endogenous AI-2 does not contribute to the reduced chemotaxis response. This lack of robust response could also be due to the fact that cells are less motile and less chemotactic during late log to
stationary phase as compared to mid log phase (128, 129). Quantitative studies have also shown that flagellar synthesis and mean run speed peak during mid log phase then decrease significantly as cells progress to late log and stationary phase (130). In our particular strain (CV1) it has been observed that a roughly inverse relationship exists between run speed and tumble frequency during later stages of growth, and that it is more difficult to achieve full adaptation of a ligand-bound chemoreceptor during this timeframe (131). Indeed, I found that response to L-serine was reduced as growth phase progressed. A combination of these factors, in addition to various other global changes that occur as growth progresses, may make it difficult to accurately characterize chemotaxis at later stages of growth.

The second approach involved deleting components of the uptake and regulation pathway, including components of the transporter lsrACD and the kinase and repressor lsrK and lsrR, respectively. This series also included luxS and TqsA, a membrane protein involved in export of newly synthesized AI-2 from the periplasm. When protein levels of LsrB were measured through western blotting, the amount of LsrB in the ΔlsrR knockout was nearly six times the amount of expression as wild-type. This did not dramatically increase the chemotaxis response to AI-2 in ΔlsrR, however, as its dose response was nearly identical to wild-type. On the other end of the expression spectrum, LsrB was expressed at only 20% of wild-type levels in ΔlsrK and only 10% of wild-type in ΔluxS. Neither of these deficiencies appeared to affect AI-2 chemotaxis, however, as the dose response to AI-2 in these deletion mutants remained virtually identical to wild-type, suggesting relatively few LsrB proteins can mediate a strong response to AI-2.
A third approach involved inducing LsrB from a plasmid promoter. Under $P_{TAC}$ control LsrB was overexpressed to a large degree, yet this had a negative impact on AI-2 chemotaxis as measured in the capillary assay. Western blots showed increasing levels of LsrB with increasing induction (with the exception of 1 and 10 µM IPTG induction), but with those increases came less chemotaxis to AI-2. This was puzzling since the limiting step of AI-2 chemotaxis seemed to be the amount of available LsrB to bind AI-2 and interact with Tsr. These results implied a negative correlation between amount of periplasmic LsrB and chemotaxis to AI-2. It remains unclear why expression of LsrB actually decreased at lower levels of IPTG induction (1 and 10 µM) from the plasmid.

These results further solidify our hypothesis that optimal AI-2 chemotaxis occurs when LsrB copy level is at our just below 300 molecules per cell at mid log phase, and that the binding between Tsr and LsrB must be very tight to elicit such a strong response despite so few molecules present. Further studies are required to fully understand the correlation between LsrB expression levels and chemotaxis to AI-2. These ideas will be discussed in Chapter V.
CHAPTER IV
DETERMINING THE EFFECT ON CHEMOTAXIS WHEN SERINE AND AI-2 ARE PRESENT SIMULTANEOUSLY

Introduction

A mechanism for simultaneous sensing of two chemoeffectors by the same chemoreceptor has been described previously. The Tar receptor can mediate chemotaxis to two independently binding ligands: L-aspartate and substrate-loaded maltose binding protein (MBP) (126). Saturating concentrations of one ligand can and do affect response to the opposite ligand. A more detailed analysis revealed that when forced to signal through the same Tar subunit, aspartate and maltose competed for signaling.

Serine is a known, potent attractant for E.coli during chemotaxis. Its receptor is one of the more abundant receptors present in the polar chemoreceptor signaling patches of E. coli cells (34). The serine chemoreceptor (Tsr) binds L-serine directly at a known binding site consisting of a pocket formed by an arginine 69 residue on one Tsr monomer and a threonine 156 on the other (107, 132, 133). Sensing of AI-2 should not involve these binding pockets and does not occur through direct ligand-binding. Based on the docking model generated for the proposed interaction between ligand-bound LsrB and Tsr, AI-2 is sensed indirectly by Tsr through LsrB docking at its periplasmic shoulder regions.

To ensure sensing of AI-2 does not in some way involve intact serine binding sites, I used the known binding site mutants to test the cells’ ability to respond to extra-
cellular AI-2. If the cells respond similar to wild-type, I surmise signaling for AI-2 neither involves direct sensing at the binding pockets, nor requires intact binding pockets comprised of Arg-69 and Thr-156 to transduce a response to AI-2.

Using the Tar-MBP model as a starting point, I hypothesized that high concentrations of one ligand could possibly affect the ability of the cells to respond to the other ligand in the capillary assay. The serine chemoreceptor is one of the more abundant MCPs in *E. coli* cells, with Tar numbering nearly 15,000 at mid-log growth (34, 134), but those cells contain only about 300 copies of LsrB (S. Jani and M. Manson, unpublished) at mid-log phase growth. Thus, I believed that high concentrations of AI-2 would have little effect on the ability of cells to respond to serine. Conversely, I hypothesized that high concentrations of serine would render the cells blind to AI-2 gradients. In other words, I looked for a shift near half maximal effective concentration (EC₅₀) of response when the second ligand was present.

**Materials and methods**

*Bacterial strains, plasmids and media.* Strain CV1 was used as our wild-type strain. Strain CV5 (Δ*tsr*) and CV1 (Δ*lsrB*) were derived from CV1 as previously described. Plasmid pPA114, a relative of pKG116 (18, 135) carrying serine binding site mutations in *tsr* under sodium salicylate control, was used to express Tsr variants (136). The pPA114 plasmid was induced with 0.6 µM sodium salicylate. All strains were grown in tryptone broth media (10 g/L tryptone; 8 g/L NaCl) supplemented with antibiotic where appropriate.
Measuring chemotaxis to attractants by mutants for serine chemotaxis. Capillary assays (114) were employed to quantify the ability of mutants for serine chemotaxis to respond to varying concentrations of AI-2. Overnight cultures were grown at 30°C in TB supplemented with streptomycin (50 μg/mL) and chloramphenicol (30 μg/mL) for selection of the pPA114 plasmid. Back dilutions were made to a turbidity of ~0.05 at O.D.\textsubscript{600} in 25 mL TB without antibiotic. Cells were grown to mid-logarithmic phase (O.D.\textsubscript{600} = ~0.5) and centrifuged at 600 x g for 10 minutes to isolate them from the growth media. Once the supernatant was removed, an equal volume of chemotaxis buffer was used to resuspend the cells. They were given 15 minutes incubation at 30°C in a low speed test tube roller for recovery. Motility was confirmed through direct observation with phase contrast microscopy.

Measuring chemotaxis to attractants in the presence of second ligand. Capillary assays (115) were chosen to quantify the response to the chosen chemoattractant. Overnight cultures of cells were grown at 30°C in TB supplemented with streptomycin (50 μg/mL). Cultures were back-diluted to a turbidity of ~0.05 at O.D.\textsubscript{600} in 25 mL TB. A 100 μM concentration of the second ligand was also added to the growth media at this time. When testing the response to AI-2 in the capillary assay, 100 μM L-serine was added; when testing for response to serine, 100 μM AI-2 was added. Cells were then grown to mid-logarithmic phase at an O.D.\textsubscript{600} = ~0.5. Cells were then centrifuged at 600 x g for 10 minutes to separate the cells from the media. The supernatant was removed and the pellet gently resuspended in chemotaxis buffer containing 100 μM of the second ligand as described above. The cells were given fifteen minutes of recovery time in a
Figure 36. Response to L-serine by Tsr binding site knockout mutants. Cells carrying substitutions at the major and minor serine binding sites of the Tsr receptor were exposed to 10 mM L-serine and their response was measured in the capillary assay. Mutant Tsr proteins were expressed from sodium salicylate-inducible plasmid pPA114 (136). Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
30°C rotor at low speed. Motility of cells was then confirmed through phase contrast microscopy. Second ligand was also added to dilutions of target ligand to ensure constant exposure during the capillary assay.

Results

*Ability of cells mutated for serine binding site to respond to AI-2.* Previous genetic studies have shown that mutations of the residues forming the serine-binding pocket between the Tsr subunits of its periplasmic domain negatively affect chemotaxis to L-serine (137-139). I confirmed this L-serine chemotaxis deficiency in the capillary assay. Cells carrying the R69E mutation (minor binding site) accumulated at less than 1% of the response by wild-type CV1 cells; cells carrying the T156K mutation (major binding site) showed only 3.5% wild-type response; binding pocket double mutants (R69E/T156K) accumulated to only 0.6% wild-type levels (Figure 36). These binding pocket mutants, however, were not affected for chemotaxis to AI-2 (Figure 37). Each of the mutants tested accumulated to within 97% or better of the wild-type dose response curve for AI-2 in the capillary assay. Cells lacking either Tsr or LsrB failed to accumulate significantly to an AI-2 gradient.

*Effects of second ligand on response to target ligand.* When cells exposed to saturating concentrations of AI-2 during growth were exposed to L-serine in the capillary assay, the dose response to serine (normalized to background control) was 105,000 cells at 1 mM (92% serine only); 72,000 cells at 100 µM (85% serine only); 31,000 cells at 10 µM (62% serine only); and 1,500 cells at 1 µM (35% serine only) (Figure 38). The
Figure 37. Response to AI-2 by serine binding site mutants. Mutations were selected based on previous findings of their affecting chemotaxis to serine. Mutant Tsr proteins were expressed from sodium salicylate-inducible plasmid pPA114 (136). Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
Figure 38. Response to serine by cells grown with AI-2. Wild-type CV1 cells were grown in the presence of 100 µM AI-2, then continuously exposed to this concentration of AI-2 during the capillary assay. AI-2 was also present in the capillaries containing the L-serine concentrations for dose response. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
background only control (saturating AI-2 present) averaged 9,500 cells accumulated, an amount similar to chemotaxis buffer lacking saturating ligand. Response to AI-2 in the presence of saturating serine concentration (100 µM), again normalized to background, was quite different. At 1 mM AI-2 exposure 43,000 cells accumulated (44% AI-2 only); at 100 µM exposure 7,000 cells accumulated (8% AI-2 only); at 10 µM exposure 1,500 cells accumulated (2% AI-2 only); and at 1 µM exposure 800 cells accumulated (14% AI-2 only) (Figure 39). Controls of background only (saturating L-serine present) averaged 60,000 cells accumulated, whereas the background only response when no saturating chemoattractant was present averaged only 12,000 cells.

Discussion

Each of the methyl accepting chemotaxis proteins (MCPs) of *E. coli* interact with a cognate binding protein carrying a particular ligand. Some also sense ligands through direct binding interactions. These interactions take place at the periplasmic region of the MCPs and the signal is transduced across the membrane to the distal tip of the chemoreceptor. In cases where multiple ligands are recognized by a single chemoreceptor, as in the case with both aspartate and maltose in Tar, independent binding sites exist for the different ligands and have been genetically analyzed (41, 125).

Our recent work showed that AI-2 is sensed through *E. coli* Tsr with the aid of the AI-2 binding protein LsrB (Chapter II, Figures 8 and 13). I surmised that the known ligands of Tsr utilize distinct binding sites of the receptor’s periplasmic domain. I reasoned, simply, that if distinct sites existed for serine and AI-2 binding, a disruption in the
Figure 39. Response to Al-2 by cells grown with serine. Wild-type CV1 cells were grown in the presence of 100 µM L-serine, then continuously exposed to this concentration of serine during the capillary assay. Serine was also present in the capillaries containing the Al-2 concentrations for dose response tests. Capillary assays were performed at 30°C for 45 minutes. Each trial represents three biological replicates carried out in triplicate.
ability of the cells to respond to serine should not affect chemotaxis toward AI-2, or vice versa. To elucidate this I tested for AI-2 chemotaxis in Tsr receptors that are unable to bind and respond to serine. I already knew *E. coli* carrying Tsr mutations making them deficient for AI-2 chemotaxis were still able to respond to serine (Figure 15). Quantifying response to AI-2 in serine chemotaxis negative cells would allow us to quickly confirm this hypothesis.

Our results here support this hypothesis. Mutations in the serine binding pockets of the Tsr homodimer, Arg-69 and Thr-156, did not affect response to varying concentrations of AI-2, as these mutants responded to this ligand just as well as wild-type. This was not a surprise, however, as our docking model suggested AI-2 cannot directly bind to Tsr, and the binding protein LsrB interacts with Tsr at its periplasmic shoulder regions.

A second approach involved exposing Tsr to hyper-physiological concentrations of one ligand prior to introducing the other. Much as in the case of Tar with aspartate and maltose binding protein, I believed exposure to and binding of one ligand might preclude binding and or response to the other. Therefore, cells were grown in the presence of a high concentration of serine before use in chemotaxis assays, and serine was even added to the chemotaxis buffer solution to ensure constant exposure over the course of the assays. The same was done using AI-2 as the second ligand. Due to the abundance of Tsr in mid-logarithmic growth phase cells, and the relatively paltry amount of LsrB in comparison (roughly 300 molecules per cell) at this same growth phase, I hypothesized
response to serine would not be affected by the presence of AI-2, but that response to AI-2 gradients would be nearly impossible.

The results gathered here seem to support this specific hypothesis, though testing various background concentrations of the second ligand is necessary to make broader conclusions. Response to serine was largely unaffected by the presence of exogenous AI-2, likely because at mid log phase cells produce an abundance of Tsr chemoreceptor in comparison to AI-2 binding proteins. With so many receptors for serine and so few LsrB proteins present to bind the added AI-2, cells have little problem recognizing serine gradients and responding accordingly. On the other hand, when serine is present at high concentrations it renders the *E. coli* cells blind to gradients of AI-2. Even if all of the approximately 300 molecules of LsrB were loaded with AI-2, they would be unable to occupy all of the available Tsr receptors.

With so little LsrB present in wild-type cells, the window for response to AI-2 appears to be very particular. Perhaps overexpression of LsrB to levels closer to those of Tsr would allow for a level playing field in terms of opportunity for binding, although this does not align with our observations from Chapter III, where increasing levels of LsrB leads to decreased chemotaxis response to AI-2. Further experiments along this line of thinking shall be discussed in Chapter V.
CHAPTER V
SUMMARY AND DISCUSSION

Summary

Until recently, a periplasmic binding protein had not been identified for the Tsr chemoreceptor in *E. coli*. The other MCPs (Tar, Tap, Trg) and the interactions with their cognate binding proteins have been studied extensively. When the identification of a periplasmic binding protein for Tsr was proposed, a genetic analysis of the proposed interaction, and a correlation between protein expression and chemotaxis became the goal of this thesis work. In this dissertation I address the accuracy of the proposed docking model for the interaction between *E.coli* Tsr and AI-2-bound LsrB through site-directed mutagenesis, and I attempt to correlate chemotaxis to AI-2 with expression of LsrB and interference by the presence of a potential competing ligand.

The results from Chapter II suggest that the shoulder regions of the Tsr homodimer, and residues in β-strand 2 and α-helix 8 of LsrB are critical for the interaction of Tsr and substrate-bound LsrB during AI-2 chemotaxis in *E. coli*. The findings in Chapter II support the proposed docking model developed and discussed in that chapter. Results from Chapter III indicate that LsrB is present in low copy number (~300 molecules per cell) at mid log phase compared to the high-abundance chemoreceptors Tsr and Tar (~15,000 molecules per cell), and that overexpression of LsrB does not directly correlate to AI-2 chemotaxis. Results here also showed that deleting various components of the *lsr* operon and production pathways, while slightly altering the expression levels of
LsrB, had little effect on AI-2 chemotaxis. Only when the periplasmic binding protein for AI-2 (LsrB) was deleted was chemotaxis negatively affected. Results from Chapter IV indicate that the presence of high concentrations of L-serine in the environment virtually blind the cells to the presence of AI-2, though the opposite is not true. A high concentration of AI-2 does little to reduce chemotaxis to serine in mid-log phase cells. Here, it was also confirmed that the serine binding pockets of Tsr are not necessary for dose response to AI-2. In this chapter, I discuss the likely mechanism of interaction between Tsr and LsrB, and explore the possible relationship between cellular levels of AI-2, LsrB and chemotaxis. I also suggest potential experiments to elucidate this relationship, including biochemical studies involving investigation of binding affinity between Tsr and LsrB.

Discussion

A docking model for the Tsr-LsrB interaction. A docking model for the interaction between AI-2 bound LsrB and the Tar chemoreceptor was proposed and tested in Chapter II. This model was based on the published model generated from studies of the Tar chemoreceptor and maltose binding protein (MBP), and an established experimental approach was available (105). Single amino acid residues were shown to be vital for maltose chemotaxis to occur. Mutation of these residues negatively affected response to maltose, presumably by disrupting the Tar-MBP interaction.

The results in this chapter support our proposed docking model and identify key residues in both proteins that must be present for AI-2 chemotaxis to take place. Alanine
substitutions in Tsr at Lys-147 and Glu-150 reduced AI-2 chemotaxis response in the capillary assay. The Lys-147-Ala mutant lost nearly all ability to respond to AI-2 gradients. The Glu-150-Ala mutant was significantly reduced for AI-2 chemotaxis, though it was nowhere near as depressed as the Lys-147-Ala mutant. In LsrB alanine substitutions at Asp-59, Val-60, Thr-61, Asp-63, Arg-252 and Glu-256 reduced AI-2 chemotaxis to varying degrees. I will discuss the likely role of each of these residues in the Tsr-LsrB interaction.

Cells expressing the Tsr variant Lys-147-Ala failed to mediate more than 20% of the wild-type AI-2 chemotaxis response. Lysine is a positively charged residue capable of forming a salt bridge with residues of the opposite charge in close proximity. The docking model intimates a very tight coupling between Tsr and LsrB, conditions conducive for electrostatic interactions to predominate. Seeing as how the most critical residues in LsrB for AI-2 chemotaxis appear to be Asp-59 and Asp-63, positively charged amino acids, a salt bridge between Lys-147 in Tsr and Asp-59 or Asp-63 in LsrB is possibly a stabilizing force for this docking, and furthermore may be crucial for eliciting a signal after binding. Because the Lys-147 is present in the other half of the Tsr homodimer, this positive charge may also attract the Glu-256 of the LsrB C-terminal domain to form a second salt bridge, further stabilizing the interaction.

The Glu-150 of Tsr potentially forms a salt bridge with the Arg-252 of the LsrB C-terminal domain, adding additional support to the interaction. Taken together with the other likely salt bridges formed, the predominant stabilization forces behind the docking of Tsr to LsrB seem to be electrostatic. When these forces are unperturbed, the two pro-
teins could come together tightly enough to elicit a strong chemotaxis response despite a large disparity between the number of Tsr and LsrB molecules present in each cell. Such a tight interaction suggests a low dissociation constant and high affinity between Tsr and LsrB, much higher than the binding affinity of Tar for MBP. Further biochemical studies are necessary to make firm conclusions in this area.

The remaining residues of interest likely contribute to the overall stability of the binding. The LsrB Thr-61, a polar residue, likely forms a supporting hydrogen bond with a residue in Tsr or a water molecule. Water molecules have been shown to play roles in ligand-protein interaction. Why the hydrophobic Val-60 in LsrB’s N-terminal domain is critical remains uncertain. Substituting alanine for valine is simply swapping two hydrophobic amino acids, though valine is typically buried to a greater degree, and changing it to alanine possibly disrupts the conformation of LsrB enough to prohibit the Asp-59 to Lys-147 salt bridge from forming without affecting the affinity of LsrB for the LsrACD transport channel during AI-2 uptake.

None of these critical single amino acid mutations affect the ability of LsrB to bind AI-2 in the periplasm and transport it back into the cell via the LsrACD transport apparatus. Bioluminescent assays measuring AI-2 uptake in *Vibrio harveyi* show each of the point mutants perform AI-2 uptake along the same time frame as wild-type LsrB cells. Therefore, I conclude these residues are critical for the Tsr-LsrB interaction, and play seemingly no role in normal *lsr* operon function. These findings ensure the negative effects on response to AI-2 result from disrupting chemotaxis rather than protein expression.
Additional mutations outside the proposed areas of interaction in LsrB and Tsr, and their effects on AI-2 chemotaxis, are needed as controls to support our docking model. If mutations in areas of the proteins away from the interaction sites do not alter AI-2 chemotaxis, it would lend support to the nature of our proposed docking interaction.

The failure of wild-type LsrB to rescue the negative phenotype for AI-2 chemotaxis resulting from the Asp-59-Ala and Asp-63-Ala point mutations remains an interesting result. This finding of a potential dominant-negative phenotype requires further consideration. It may result from a stronger binding of this mutant LsrB with Tsr, precluding binding of wild-type LsrB to the chemoreceptor. If this is the case, notably it does not seem to inhibit response to serine. This scenario suggests the Asp-59 and Asp-63 mutants are AI-2 signaling mutants rather than binding mutants. A separate scenario for this effect may be dimerization of mutant and wild-type LsrB proteins, though AI-2 uptake remained consistent with wild-type monomeric LsrB in the bioluminescence assay. Complementation studies in which residues in Tsr are mutated, beginning with Lys-147, could reverse this dominant negative phenotype to some degree. Exchanging charges between Tsr and LsrB (Lys-147-Glu and Asp-59/63-Arg, respectively) could potentially counteract the disruption of salt bridge formation.

A second possibility is that the LsrB protein could be locked into a closed conformation resulting from this point mutation. While it is unlikely a surface mutation would have this effect, dominant negative phenotypes have been observed for maltose chemotaxis in *E. coli* and were the result of crosslinking event that locked MBP into its
substrate-free/closed conformation (140). Perhaps introducing inter-domain cysteine-cysteine crosslinks into wild-type LsrB could be used to test this. If the dominant negative phenotype is the result of a closed conformation mutation, the dose response curve for this crosslinking mutant could look similar to the D59A or D63A mutants.

*The amount of LsrB present does not correlate directly with chemotaxis to AI-2.*

As mentioned in Chapter I, the expression of LsrB from the *lsrACDBFG* operon is regulated by the presence of phosphorylated AI-2 (AI-2-P). When a threshold of environmental AI-2 is reached and it accumulates within the periplasm, it is imported into the cell via an ABC transporter. Some amount of LsrB is present to bind the AI-2 and interact with the LsrACD transport channel. Once imported, AI-2 is phosphorylated by the kinase LsrK, at which point it can bind to LsrR and relieve repression of the *lsr* operon. When this occurs, more transport channel proteins are expressed (LsrACD) and more LsrB is produced, allowing more AI-2 to be taken into the cell. Growing cells to later log phase allows up to six times as much LsrB to be present compared to the approximately 300 molecules per cell during mid logarithmic growth.

Our initial hypothesis was that cells might respond to increasingly higher levels of AI-2 if more LsrB was available for binding and interacting with Tsr. Not only was this assumption incorrect, the response actually decreased with increasing LsrB levels at later phase growth. This result was confirmed with overexpression from an LsrB-bearing plasmid. To us, this suggests an optimal window for response to AI-2, namely at mid log phase cell growth and roughly 300 copies of LsrB per cell. This suggests a scenario where *E. coli* cells growing and dividing in the gastrointestinal tract receive
quorum sensing information via AI-2 concentration and respond quickly, taking them to areas of higher cell density where nutrients may be more readily available, such as a pyre’s patch. Once in these areas, the increasing levels of LsrB allow for import and degradation of AI-2 by the lsrFG gene products whose expression is also induced as repression is relieved, thus returning the lsr operon to its pre AI-2 threshold state.

Due to the numerous global changes occurring during later phase growth (128, 130), it is difficult to make direct correlations between protein levels and chemotaxis response. As cells move into later stages of the growth curve, swim speed and motility decrease in general, and frequency of tumbling increases. Perhaps even the changing profile of the high abundance chemoreceptors Tsr and Tar at later log growth (141) makes chemotaxis to attractants, AI-2 among them, difficult to measure.

*Chemotaxis to AI-2 is reduced in the presence of serine.* Studies involving simultaneous exposure to gradients of L-aspartate and maltose for the *E. coli* Tar chemoreceptor have suggested a preference by cells for one attractant over another. Under conditions where the ligands were forced to bind and signal through the same Tar subunit, saturating concentrations of maltose only moderately inhibited aspartate chemotaxis, while the opposite showed a more drastic effect. When saturating aspartate was present at the time the cells were exposed to maltose gradients, the response to maltose was virtually eliminated (126). The ligands were characterized as competing and aspartate was the winner. This competition did not occur when ligands were forced into signaling through opposite subunits of Tar.
My studies with background concentrations of serine and AI-2 have suggested these two ligands can signal through the same subunit as well. Disrupting the binding pocket for serine at either Arg-69 or Thr-156 did not interfere with AI-2 chemotaxis. Likewise, mutants deficient in AI-2 chemotaxis still responded at wild-type levels to serine gradients. When both attractants were present at the same time, however, the cells were unable to mount a dose response to AI-2. When saturating levels of serine remained constant throughout the exposure to AI-2, cells failed to perform any significant response to the latter in the capillary assay. On the other hand, when AI-2 was the saturating ligand at constant exposure, it did not reduce the chemotaxis response to serine relative to wild-type.

It is already apparent that LsrB is present at lower copy number per cell during mid logarithmic growth compared to Tsr, and this may be the central reason for the lack of response to AI-2 when serine is present. If the cells have more opportunity to interact with serine due to the abundance of Tsr, they may become blind to AI-2 gradients at higher serine concentrations. My results lean toward this conclusion. With saturating AI-2 present, at 1 mM serine the response was within 93% of wild-type in the absence of competing ligand; at 100 µM serine the response fell to 85% of no competitor; and at 10 µM serine it dropped to 63% of wild-type without AI-2 present (Figure 38). This suggests that only when levels of serine are low enough will cells begin to recognize the presence of AI-2 in the environment.

In the gastrointestinal tract of humans, *E.coli* does well to respond to serine as a priority to AI-2 since serine is a nutrient source for the cells. Autoinducer-2 serves as a
signaling molecule, not a nutrient source; therefore, in this case it is in the cells’ best interest to respond to a nutrient source over a non-nutrient source, especially considering that stable biofilms are not likely to appear in the dynamic gut. Perhaps this is why so few copies of LsrB are present in the cells at a given time relative to Tsr or Tar. When it serves the best interest of the cells to migrate away from low nutrient areas toward higher population densities – a potential indicator of plentiful nutrients – AI-2-laden LsrB has little competition for binding to Tsr and eliciting a chemotactic response. Perhaps the grand balance of a healthy microbiome within the human gut is partially achieved through regulated response to nutrient sources and signaling molecules.

**Future Studies**

My work here has provided a docking model for the interaction of AI-2-bound LsrB with Tsr during AI-2 chemotaxis in *E. coli*. It has also shown there is no direct correlation between increasing levels of LsrB and chemotaxis to AI-2. Furthermore, it has indicated that the presence of a high concentration of L-serine can reduce the ability of the cells to perform chemotaxis to AI-2. However, there are certain aspects of AI-2 chemotaxis that remain unanswered.

First, it can be determined whether serine and AI-2 compete to signal through the same subunit of Tsr. Tsr can exist as a heterodimer of wild-type and mutant proteins when point mutants and wild-type versions of Tsr are expressed from compatible plasmids within the same cell. These heterodimers will allow serine to bind in only one conformation, causing it to signal through a particular subunit. If mutant LsrB proteins can
be forced to interact with Tsr and signal through that same subunit, it might be determined if serine and AI-2 signals compete for signaling. This same scenario was studied with aspartate and MBP in Tar, and it was determined that when forced to signal through the same subunit of Tar, competition did occur (122).

Second, we can elucidate the effects of the presence of secondary attractants on response to either serine or AI-2. Varied background concentrations of either serine or AI-2 would give a better picture of the amount of secondary ligand necessary to inhibit response to the first. Up to this point, only 100 µM of either serine or AI-2 was used as background, but testing higher and lower concentrations would give us a better idea of the EC₅₀ for each. Furthermore on this point, expressing LsrB to levels higher than the 300 copies produced by the chromosome at mid-log phase would clarify the ability of serine at 100 µM to reduce AI-2 chemotaxis. By expressing extra LsrB from an inducible plasmid, perhaps AI-2 chemotaxis would not be affected by the presence of serine. Even more useful would be the use of tethered cell assays or microfluidic assays that test the response of cells to multiple ligands in real time. The capillary assay gives only a measure of the end result of exposure to chemoeffectors.

Next, what roles do the aspartates in LsrB at positions 59 and 63 play during chemotaxis? My results suggest both are critical to the ability of cells to respond to gradients of AI-2. It is unclear if they help stabilize the Tsr-LsrB interaction, or if they allow for signaling through Tsr to occur post-binding. It would be useful to see the chemotaxis response of a D59A/D63A double mutant to determine if the charge or the position is the important aspect.
Additionally, determining the binding affinity of LsrB for Tsr is important. Due to its unstable nature, AI-2 has been an issue here. According to our hypothesis, it must be present and bound to LsrB to get a closed conformation of LsrB that can interact with Tsr. This is a problem for most biochemical assays that can help determine binding affinity. It is also a problem for obtaining a co-crystal structure of LsrB-Tsr. If we could obtain either a stable form of AI-2, or a closed conformation of LsrB that interacts with Tsr, it would make it possible to carry out the necessary biochemical assays for determining binding affinity and the nature of the interaction. Assays such as isothermal titration calorimetry, or equilibrium gradient ultracentrifugation could be employed.
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