ACYL-HOMOSERINE LACTONE BASED MODULATORS FOR RHLI, A QUORUM SENSING SIGNAL SYNTHASE IN PSEUDOMONAS AERUGINOSA

by

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A thesis
submitted in partial fulfillment
of the requirements for the degree of
Master of Science in Chemistry
Boise State University

December 2017
DEFENSE COMMITTEE AND FINAL READING APPROVALS

of the thesis submitted by

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Thesis Title: Acyl-Homoserine Lactone Based Modulators for RhlI, a Quorum Sensing Signal Synthase in Pseudomonas aeruginosa

Date of Final Oral Examination: 13 October 2017

The following individuals read and discussed the thesis submitted by student Daniel D. Shin, and they evaluated his presentation and response to questions during the final oral examination. They found that the student passed the final oral examination.

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The final reading approval of the thesis was granted by Rajesh Nagarajan, Ph.D., Chair of the Supervisory Committee. The thesis was approved by the Graduate College.
ACKNOWLEDGEMENTS

I would like to show my gratitude to Boise State University and the Department of Chemistry and Biochemistry for the opportunity of pursuing research. I thank Mila Lam, Levi Mitchell, John Taffin, and Neil Rexrode for their help with enzyme purification and small molecule synthesis. I have deep appreciation for Dr. Eric Brown, who was crucial in synthesizing small-molecules, and Dr. Shin Pu and Matthew Turner, who were pivotal in identifying the said compounds. The generous contribution of small molecule compounds from Dr. Helen Blackwell of University of Wisconsin made this work possible. My graduate committee members, Dr. Henry Charlier and Dr. Michael Callahan, provided the needed criticism to look at my work again from third and fourth angles so that the objective of this work became better focused. Most importantly, I would like to thank Dr. Rajesh Nagarajan for all the time, patience, concern, guidance, and dedication he has showed to me, not just as a student and a scientist, but as a human being.
ABSTRACT

Gram-negative bacteria use N-acyl-homoserine lactone (AHL) autoinducer based signal system, known as quorum sensing (QS), to modulate the gene expression for such traits as biofilm formation, toxin production, and antibiotic resistance. Therefore, there is great potential in pursuing quorum sensing inhibition (QSI) as a means of achieving antivirulence. *Pseudomonas aeruginosa*, an opportunistic pathogen commonly found in healthcare-related infections, use two LuxI/R type systems to regulate AHL-based quorum sensing: LasI/R and RhlI/R. LasI (initiator protein/signal synthase) and LasR (receptor) use 3-oxododecanoyl-L-homoserine lactone signal molecule while RhlI and RhlR use butanoyl-L-homoserine lactone autoinducer. Thus far, most of the studies have focused on inhibiting the Las system, in particular by using AHL signal analogs to interfere with signal-receptor binding. Recently, RhlI/R system has gained attention as potentially having greater effect in *P. aeruginosa* virulence. In this study, we have tested the effect of AHL analogs on RhlI, as product inhibitors with the goal of targeting both RhlI and RhlR for increased potency. Screening of compounds have revealed three variations to have the greatest effect on RhlI inhibition: longer/bulkier acyl-chain, D-stereocenter in the headgroup, and a less polar thiolactone head-group. Surprisingly, the addition of a carbonyl at the C3 position was found to activate the enzyme. Moreover, we measured kinetic constants of RhlI with various acyl-substrates and performed inhibition assays with inert acyl-substrate analogs to determine how RhlI activity changes to variations in the acyl-chain length. We found that the catalytic efficiency of acyl-
substrate and inhibition potency of the corresponding inert acyl-substrate analogs surges with increase in the length of the acyl-chain. These patterns suggest that long acyl-chains most likely bind to an alternate binding site with marked increase in both $k_{on}$ and $k_{off}$ rate constants. Our findings with AHL derivatives provide a basis for rational design of quorum sensing inhibitors to better combat *P. aeruginosa* bacterial infections.
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<td>CDC</td>
<td>Center for Disease Control and Prevention</td>
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<tr>
<td>PMSF</td>
<td>Phenylmethanesulfonyl fluoride</td>
</tr>
<tr>
<td>C4-CoA</td>
<td>Butanoyl-CoA</td>
</tr>
</tbody>
</table>
C6-CoA         Hexanoyl-CoA
C8-CoA         Octanoyl-CoA
C10-CoA        Decanoyl-CoA
C12-CoA        Dodecanoyl-CoA
DNase          Deoxyribonuclease I
RNase          Ribonuclease I
IPA            2-Propanol
B-PER          Bacterial protein extraction reagent
DMF            Dimethylformamide
DMSO           Dimethyl sulfoxide
HCl            Hydrochloric acid
IPTG           Isopropyl β-D-1-thiogalactopyranoside
MnSO₄          Manganese sulfate
TFA            Trifluoroacetic acid
MgCl₂          Magnesium chloride
kD             Kilodalton
Sfp            Surfactin-synthetase activating protein/phosphopantetheinyl transferase
UV-Vis         Ultraviolet-visible light
HPLC           High performance liquid chromatography
UHPLC          Ultra high performance liquid chromatography
QTOF           Quadrupole time-of-flight
ACN            Acetonitrile
Buffer A  50 mM Tris-HCl, pH 7.5, 0.2 M NaCl, 1 mM EDTA, 0.1 mM PMSF, 0.1 mM TLCK, 0.4 M sucrose, and 2.5% (v/v) glycerol

Psi  Pounds per square inch

OD$_{600}$  Optical density at 600 nm

w/v  weight to volume ratio

RT  Retention time

C6-ACP  Hexanoyl-ACP

C4-ACP  Butanoyl-ACP

IACP  Inert-ACP/alkyl-ACP

C4-IACP  Butyl-ACP

C6-IACP  Hexyl-ACP

C8-ACP  Octanoyl-ACP

C8-IACP  Octyl-ACP

C10-ACP  Decanoyl-ACP

C10-IACP  Decyl-ACP

C12-ACP  Dodecanoyl-ACP

NaHCO$_3$  Sodium bicarbonate

KHSO$_4$  Potassium bisulfate

HPLC-MS  High performance liquid chromatography mass spectrometry

MS  Mass spectrometry

LC-MS  Liquid chromatography-mass spectrometry
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ESI</td>
<td>Electrospray ionization</td>
</tr>
<tr>
<td>LC</td>
<td>Liquid chromatography</td>
</tr>
<tr>
<td>TLC</td>
<td>Thin layer chromatography</td>
</tr>
<tr>
<td>3-oxoC8-D-HSL</td>
<td>3-Oxooctanoyl-D-homoserine lactone</td>
</tr>
<tr>
<td>3-oxoC6-D-HSL</td>
<td>3-Oxohexanoyl-D-homoserine lactone</td>
</tr>
<tr>
<td>C12-D-thiolactone</td>
<td>Dodecanoyl-D-homocysteine thiolactone</td>
</tr>
<tr>
<td>3-oxoC12-D-thiolactone</td>
<td>3-Oxododecanoyl-D-homoserine lactone</td>
</tr>
<tr>
<td>C12-L-thiolactone</td>
<td>Dodecanoyl-L-homocysteine thiolactone</td>
</tr>
<tr>
<td>3-oxoC12-L-thiolactone</td>
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<td>3-oxoC6-D-thiolactone</td>
<td>3-Oxohexanoyl-D-homoserine lactone</td>
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<tr>
<td>3-oxoC10-D-thiolactone</td>
<td>3-Oxodecanoyl-D-homoserine lactone</td>
</tr>
<tr>
<td>C4-ICoA</td>
<td>Butyl-CoA</td>
</tr>
<tr>
<td>C6-ICoA</td>
<td>Hexyl-CoA</td>
</tr>
<tr>
<td>C8-ICoA</td>
<td>Octyl-CoA</td>
</tr>
</tbody>
</table>
CHAPTER ONE: INTRODUCTION

The Antibiotic Crisis

The modern age of medicine was made possible, in large parts, by the discovery and the proliferation of antibiotics. However, mankind has been slowly disarmed by the rise and the spread of antibiotic resistant superbugs. In the U.S. alone, antibiotic-resistant bacteria are responsible for at least two million infections, resulting in 23,000 lives and cost $35 billion annually.\(^1\) In response to this rise in antibiotic resistance, the White House has named it a threat to national security.\(^2\) Despite efforts to curb the tide, both CDC and WHO warn of a post-antibiotic age.\(^1,3\)

The rise of superbugs highlights a fundamental problem with antibiotics; the use of antibiotics is an all-or-nothing treatment in which all the drug-sensitive strains are eliminated while the resistant ones are untouched. This dichotomy places enormous selective pressure for resistant bacteria to propagate throughout the entire population. In a controlled environment, a strain of sensitive bacteria can be pressured to withstand ever-increasing concentrations of antibiotics in less than two-weeks.\(^4\) Even in the real-world setting, resistance often develops within a few years of the release of a new antibiotic (Figure 1).\(^5\) Therefore, drug companies are shying away from continuing to put resources into R&D of a drug that could become irrelevant soon after its commercialization (Figure 2).\(^6,7\) The resulting decrease in supply combined with the increase in demand have led to the investigation of novel methods to fight bacteria, such as predatory bacteria, bacteriophage, antimicrobial peptides, gene-editing enzymes, and quorum sensing.\(^8-13\) Of
these alternatives, most follow the pattern of antibiotics in that they kill sensitive strains, leaving behind resistant strains to flourish; however, quorum sensing (QS) provide a unique means of avoiding the pitfalls of antibiotics by targeting virulence rather than the organism directly.\textsuperscript{14}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{timeline.png}
\caption{Timeline of the effective-lifetime of antibiotics. The labels above the timeline indicates the year different antibiotics were released, whereas the year resistance to each antibiotic was first observed is shown below the timeline. In many cases, resistance developed within a few years of deployment of new antibiotics.}
\end{figure}
Figure 2. Decrease in new antibiotic development. In the heydays of antibiotic development in the 1980’s, more than 3 new drugs were released annually; however, in recent years, that number has been reduced to just two new antibiotics in five years.

Quorum Sensing

Once known mainly for their single-cellular life, bacteria are now known to communicate via chemical signals to approximate multicellular behavior in a process termed quorum sensing. The concentration of the signal molecules, called autoinducers, correlate to the number of bacteria in the area. Therefore, by monitoring the concentrations of the autoinducers, bacteria can detect the concentration of local bacteria population. Typically, when signal concentration is low, indicative of limited cell count, bacteria operate individually. However, once a certain concentration, or “quorum,” is reached, QS-controlled genes are turned on, activating group behaviors, which includes biofilm formation, virulence, antibiotic resistance, protease production, and siderophores. Therefore, theoretically, QS inhibitors could be used as combination drug with antibiotics since antibiotic resistance associated with QS inhibition (QSI) would be
reduced. Additionally, inhibiting quorum sensing should prevent virulent traits from emerging. Indeed, studies have shown QS-knockout strains to be incapable of colonial behaviors characteristic of pathogenicity. As an anti-virulence treatment, contrary to the lethal (to the bacteria) antibiotic therapy, QS inhibition (QSI) could potentially pave the path to an “evolution-proof” method of dealing with bacterial infections. It has been shown that QSI resistance does not lead to survival advantage. The study showed that since the majority of the population are unable to produce autoinducer signal molecules, the few resistant strains could not produce sufficient AHLs to reach “quorum” and induce QS-controlled gene expression; however, even if the resistance strains are able to express QS-controlled traits, many of which are group-beneficial, the benefits of those traits are shared with the inhibited strains, thus limiting any survival advantage QSI resistance could offer.

Quite contrary to the notion of being “simple” creatures, bacteria have complex communication system with different QS system for various types of bacteria. Gram-negative bacteria have LuxI (initiator/synthase) and LuxR (receptor) type proteins responsible for the synthesis and the uptake of N-acyl-homoserine lactone (AHL/AI-1) autoinducers (Figure 3; Table 1). In this communication system, AHL synthases release AHL autoinducers and release them to the environment. The binding of the signal with the designated receptor protein starts the upregulation of QS-controlled genes such as increased AHL synthase activity, biofilm formation, toxin production, and antibiotic resistance.

AHL synthases are bi-substrate enzymes that use S-adenosyl-L-methionine (SAM) and an acyl substrate (acyl-ACP or acyl-CoA) to synthesize AHL signal
molecule. Whereas SAM is a conserved substrate among various AHL synthases, preferential usage of an acyl substrate of certain acyl-chain moiety results in enzyme-specific signal molecules, allowing each species to speak its own unique “language,” or differentiated AHLs (Figure 4).\textsuperscript{21} Many AHL synthases have been shown to discriminate against non-native acyl-chain and only use a specific acyl substrate to produce enzyme-specific AHL signal, thus increasing the signal to noise ratio.\textsuperscript{22} Different receptor proteins likewise preferentially bind with their designated AHLs so that bacteria can conduct intra-specie communication without interference.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{quorum_sensing_system}
\caption{The typical quorum sensing system found in Gram-negative bacteria. A LuxI-type signal synthase produces AHL signal molecules which then binds to LuxR-type receptor proteins. The signal-receptor binding promotes the expression of QS-controlled genes, which includes further activation of the synthase.}
\end{figure}
Figure 4. **Species-specific signal molecules.** The different “languages” used by each Gram-negative bacteria species are dependent on the variations in the acyl-chain which is derived from the acyl substrate.

<table>
<thead>
<tr>
<th>Signal Type</th>
<th>Organism</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>AI-1/AHL</td>
<td>Gram-negative</td>
<td><img src="image" alt="Structure of AI-1/AHL" /></td>
</tr>
<tr>
<td>AIP</td>
<td>Gram-positive</td>
<td><img src="image" alt="Structure of AIP" /></td>
</tr>
<tr>
<td>AI-2</td>
<td>Universal</td>
<td><img src="image" alt="Structure of AI-2" /></td>
</tr>
</tbody>
</table>
Gram-positive bacteria, on the other hand, use short peptides, linear or cyclic, as QS signals (Table 1). As with Gram-negative QS system, the specificity comes from varying the signal molecules by modifying the peptide sequence and shape. Synthesized in the cytoplasm and actively transported out of the cell, these autoinducer peptides (AIP) bind to membrane-bound histidine kinase, leading to a phosphorylation cascade, activating QS regulator proteins for gene expression.

In addition to the species-specific AHL/AI-1 and AIP based QS for interspecies-communications, a universal signal-based QS system has also been discovered. This interspecies-communication QS is based on AI-2 signal molecule (Table 1). Like the AIP-based system, AI-2 binding event starts a phosphorylation cascade to regulate gene expression.

In light of the antibiotic crisis, of these three systems (AHL, AIP, AI-2), AHL-based QS found in human-pathogen-causing Gram-negative bacteria is of great interest. Four-main principles of targeting AHL communication system have been proposed: synthase inhibition, receptor inhibition, quorum-quenching enzymes, and AHL-sequestering antibodies. Most studies thus far have explored receptor inhibition; however, due to high affinity of native AHLs to AHL-receptor proteins, it is a difficult task to design a molecule that could out compete the binding of native autoinducer to LuxR-type proteins to disrupt QS. The progress towards developing synthase inhibitors, on the other hand, is hampered because most AHL synthases are yet to be characterized. Nevertheless, AHL synthase-knockout studies have led to elimination of virulence traits, supporting AHL synthase modulation as a means of QS control.
Efforts toward understanding the mechanism of AHL synthesis are important to design potent and selective AHL synthase inhibitors.

**Proposed AHL synthase mechanism**

AHL synthases are bi-ter enzymes that catalyzes the conversion of S-adenosyl-L-methionine (SAM) and acyl-acyl-carrier protein (acyl-ACP) or acyl-coenzyme A (acyl-CoA) to AHL autoinducer, 5′-deoxy-5′-methylthioadenosine (MTA), and holo-ACP or free-CoA (Figure 5). During catalysis, nucleophilic attack from the amine group of SAM to the carbonyl carbon of acyl-ACP cleaves the thioester bond, thereby releasing holo-ACP and transferring the fatty acid tail to SAM in the acylation half-reaction. In the lactonization reaction, SAM undergoes intramolecular ring closure, forming the lactone head-group and producing MTA side product.

**Figure 5.** The proposed mechanism of AHL-synthases. SAM is a conserved substrate amongst all AHL synthases. The variation comes from the R-group of the acyl substrate.

As with any other enzyme, AHL synthases can be inhibited by interference in the substrate binding, the catalysis, or the product release steps. However, each approach has its corresponding challenges. To disrupt substrate binding, substrate analogs would be the ideal starting point, yet SAM and acyl-ACP are commonly used in human enzymes,
SAM as a common methyl donor and acyl-ACP as a key player in fatty-acid biosynthesis.\textsuperscript{39, 40} Therefore, SAM or acyl-ACP based design of inhibitors have the potential to target all SAM or acyl-ACP using enzymes, thereby increasing the risk of unwanted side-effects. Moreover, how an AHL synthase preferentially bind to its native substrate over similarly shaped analogs, such as acyl-ACPs of different chain length, is not well understood, which impedes the design of substrate analogs that would out-compete the native substrates binding to the enzyme. Mechanism-based covalent inhibitors act by forming covalent bonds with certain active site residues; however, as most AHL synthases remain uncharacterized the active site residues to target remain unknown.

As holo-ACP and MTA are both relevant in human biology, holo-ACP in fatty acid synthesis and MTA in polyamine synthesis, product inhibitors based on those compounds carry the dangers of serious side-effects.\textsuperscript{41, 42} AHL-based inhibitors, on the other hand, has qualities suitable for pharmaceutical uses; AHL-analogs would have the potential of targeting both the synthase and the receptor, be QS-specific, have favorable cell-membrane diffusion characteristics, and have long shelf-life.\textsuperscript{30, 31, 33} There have been no reports of AHL-based modulators tested on AHL-synthases; however, studies reveal their great effect on AHL receptors, both as agonists and antagonists.\textsuperscript{43-45}

**RhII, QS signal synthase in *Pseudomonas aeruginosa***

*Pseudomonas aeruginosa* is a specie of common Gram-negative bacteria that can cause diseases in both animals and plants. It is categorized as opportunistic bacteria commonly associated with healthcare-associated infections, infecting 51,000 every year in American hospitals, killing 400.\textsuperscript{46} In humans, it can cause pneumonia, cystic fibrosis,
bacterial meningitis, septic shock, urinary tract infection, GI infection, and skin and soft tissue infections.\textsuperscript{6, 46, 47} \textit{P. aeruginosa} infections are becoming harder to treat due their increasing resistance to antibiotics, so much so that the CDC labeled it as a “serious threat” in their report on “Antibiotic Resistance Threats in the United States, 2013.”\textsuperscript{46}

The virulence and antibiotic resistance in \textit{P. aeruginosa} is controlled by its QS system comprised of AHL-quinolone system. Its AHL system is comprised of LasI/R and RhlI/R systems whereas the Pseudomonas quinolone signal (PQS) is under Pqs receptor (PqsR) control (Figure 6).\textsuperscript{48} The two AHL-based systems use very different signal molecules. LasI synthesizes 3-oxo-dodecanoyl-homoserine lactone (3OC12HSL) signal molecule, which binds with LasR to activate several QS-controlled genes, which includes the Rhl and Pqs system. The RhlI/R proteins, in contrast, use butanoyl-homoserine lactone (C4HSL) autoinducers as the signal molecule. Under normal conditions, LasI/R pair is thought to activate the Rhl and the Pqs system, though both the Rhl and the Pqs have been observed acting independent of the Las system under certain circumstances. Furthermore, Pqs generally activates Rhl whereas Rhl typically suppresses Pqs via mechanisms not completely understood. Studies have shown that inhibiting any of these three QS systems significantly reduces \textit{Pseudomonas aeruginosa} virulence. Studies thus far have focused primarily on Las-system inhibition and Rhl-specific inhibitors have yet to be studied in great detail. However, Rhl-system inhibition merit further examination for Rhl knockout strains strain to display pathogenic phenotype.\textsuperscript{49, 50}
Figure 6. QS system regulation in *P. aeruginosa*. LasI/R promotes both the Rhl and Pqs systems. Whereas Pqs induces Rhl activity, Rhl inhibits Pqs via unknown methods.

**AHL synthase Kinetics**

**DCPIP Assay**

DCPIP colorimetric assay is a well-established assay used to determine AHL synthase activity by monitoring the enzyme-dependent rate of release of holo-ACP/CoA thiol product over time (Figure 7). DCPIP, in its oxidized form, is a blue compound that absorbs at 600 nm ($\varepsilon_{600} = 21,000 \, \text{M}^{-1}\text{cm}^{-1} = 2.1 \times 10^{-2} \, \mu\text{M}^{-1}\text{cm}^{-1}$). The holo-ACP thiol released upon acylation of SAM reduces the DCPIP dye to a colorless form, DCPIPH$_2$. By monitoring the change in the absorbance of DCPIP at 600 nm, the enzyme rate can be determined by the following equation:
\[ Abs_{600} = \varepsilon_{600} \cdot l \cdot c = (2.1 \times 10^{-2} \mu M^{-1})(cm^{-1})(c) \]  

\[ c = \frac{Abs_{600}}{\varepsilon_{600} \cdot l} = \frac{Abs_{600}}{\varepsilon_{600}} \cdot \left(\frac{1}{l} \cdot \frac{1}{c} \cdot \frac{1}{cm} \cdot \frac{1}{\mu M} \cdot \frac{1}{DCPIP \cdot cm} \right) = \frac{Abs_{600} \mu M DCPIP \cdot cm}{2.1 \times 10^{-2}} \]  

\[ rate = \frac{c}{t} = \left(\frac{Abs}{\varepsilon \cdot t} \right) \cdot \left(\frac{1}{\mu M DCPIP \cdot cm} \right) \cdot \left(\frac{2 \mu M thiol}{cm} \right) \cdot \left(\frac{1}{s} \right) \cdot \left(\frac{1}{1 \text{ min}} \right) \]  

**Figure 7.** DCPIP assay mechanism. DCPIP, in its oxidized form, is a blue compound that turns colorless upon reduction with two thiols released from acylation reaction in AHL synthesis. By monitoring the decrease of absorbance at 600 nm, AHL synthase rate can be determined.

**Enzyme Kinetics**

Initial rate of enzyme activity as a function of substrate concentration follows a hyperbolic curve (Figure 8). When the substrate is present in saturating concentrations, maximum initial rate is achieved, denoted by \( V_{max} \). The concentration of the substrate required to reach half-maximal velocity is termed \( K_m \).
This plot, also called the substrate-velocity curve, is commonly analyzed using the Michaelis-Menten equation. In the simplest situation where one substrate is converted to one product, the reaction can be summarized as:

\[ E + S \rightleftharpoons ES \rightarrow E + P \]

(4)

where \( E \) is the enzyme, \( S \) the substrate, \( ES \) the enzyme-substrate complex (Michaelis complex), \( P \) the product, and \( v \) the reaction rate. The rate of the reaction (\( v \)) can be described with the equation:

\[ V = \frac{V_{\text{max}}[S]}{[S] + K_m} \]

(5)

where \( V_{\text{max}} \) is the highest reaction rate that the enzyme can achieve at saturating substrate concentration, \( [S] \) the substrate concentration, and \( K_m \) the substrate concentration required for half-maximal rate. The \( V_{\text{max}} \) and \( K_m \) of an enzyme can be determined by measuring the initial enzyme rate with various substrate concentrations and fitting the resulting plot with the Michaelis-Menten equation. For easier interpretation, this hyperbolic curve can be linearized by taking the inverse of both sides of the equation,
thereby converting the Michaelis-Menten equation to the double-reciprocal or Lineweaver-Burke equation (Figure 9):

\[
\frac{1}{v} = \frac{K_m}{V_{max}} [S] + \frac{1}{V_{max}}
\]

(6)

where independent axis (x-axis) represents the substrate concentration and the dependent axis (y-axis), the inverse rate. In this form, a change in the \(V_{max}\) would lead to a shift in the y-intercept (intercept-effect) and a change in the \(\frac{K_m}{V_{max}}\) (inverse of catalytic efficiency, \(k_{cat}/K_m\)) ratio would cause a change in the slope (slope-effect).

![Typical Lineweaver-Burk plot](image)

**Figure 9.** Typical Lineweaver-Burk plot. Taking the inverse of the hyperbolic Michaelis-Menten plot linearizes the data. In the \(1/V\) vs \(1/[S]\) plot, the y-intercept corresponds to \(1/V_{max}\), the x-intercept to \(-1/K_m\), and the slope to \(K_m/V_{max}\).

However, AHL synthases are bi-ter enzymes (2 substrates, 3 products). The order of substrate binding and product release (the kinetic mechanism) can follow one of three patterns (Figure 10). 1) Ordered sequential mechanism: substrates bind to the enzyme in a specific order before the products are released in a defined sequence. 2) Random sequential mechanism: substrates all bind before the first product is released but the order of substrate addition or the product release, or both, is random. 3) Ping-pong sequential mechanism: the first product is released before the second substrate binds with the enzyme.
Figure 10. **Kinetic mechanism for bisubstrate enzyme.** (a) Ordered sequential: substrate binding occurs before product release and follow definite order. (b) Random sequential: substrate binding occurs before product release and follow random order. (c) Ping-pong: first product release step occurs before the second substrate addition.

Bi-substrate enzyme kinetics is represented by a more complicated Cleland equation as shown:

$$v = \frac{v_{max}[A][\theta]}{K_m^A + [A] + K_m^B + [B]}$$  \hspace{1cm} (7)

where $K_m^A$ is the Michaelis constant for substrate A at saturating concentrations of B, $K_m^B$ the Michaelis constant for substrate B at saturating concentrations of A, and K depends on the reaction type (sequential vs ping-pong). In a sequential mechanism, the Cleland equation becomes:

$$v = \frac{v_{max}[A][\theta]}{K_i^A K_m^B + K_m^A [A] + K_m^B [B] + [A][\theta]}$$  \hspace{1cm} (8)
where the $K_i^A$ is the dissociation constant for the EA complex in the absence of the second substrate, B. If the enzyme undergoes ping-pong mechanism, the K term drops and the Cleland equation becomes:

$$v = \frac{v_{\text{max}}[A][B]}{K_m^A[B] + K_m^B[A] + [A][B]}$$  \hspace{1cm} (9)$$

The double reciprocal of the Cleland equation for sequential mechanism is:

$$\frac{1}{v} = \frac{1}{v_{\text{max}}} \left[ (1 + \frac{K_m^B}{[B]}) + \frac{1}{[A]} \left( K_m^A + K_i^A \cdot \frac{K_m^B}{[B]} \right) \right]$$  \hspace{1cm} (10)$$

and for the ping-pong mechanism:

$$\frac{1}{v} = \frac{1}{v_{\text{max}}} \left[ (1 + \frac{K_m^B}{[B]}) + \frac{1}{[A]} (K_m^A) \right]$$ \hspace{1cm} (11)$$

at constant [B] and variable [A], the slope of sequential equation is:

$$\frac{1}{v_{\text{max}}} (K_m^A + K_i^A \cdot \frac{K_m^B}{[B]})$$ \hspace{1cm} (12)$$

which indicates that the slope depends on the concentration of substrate B (slope-effect).

For the ping-pong equation, the slope is:

$$\frac{K_m^A}{v_{\text{max}}}$$ \hspace{1cm} (13)$$

indicating that the slope is independent of the substrate B concentration (no slope-effect).

For both the sequential and the ping-pong mechanism equation, the intercept of this is:

$$\frac{1}{v_{\text{max}}} (1 + \frac{K_m^B}{[B]})$$ \hspace{1cm} (14)$$

showing that the intercept always depends on B concentration (intercept-effect). This dependence on [B] indicates that saturation with A does not cut off the effect of adding more B. Therefore, at saturating A concentration, the addition of B always increases the rate (increases the apparent $V_{\text{max}}$).

For sequential mechanism, since both the slope and the intercept are dependent on the fixed concentration of substrate B, the double reciprocal plot of $\frac{1}{v}$ vs $\frac{1}{[A]}$ at various
fixed [B] would feature a set of lines intersecting in quadrant II (Figure 11 a). For ping-pong mechanism, since only the intercept is affected by the concentration of B, the double reciprocal plot of $\frac{1}{V}$ vs $\frac{1}{[A]}$ at various fixed [B] would feature a set of parallel lines (Figure 11 b).

**Figure 11. Double reciprocal plot of sequential vs ping-pong reaction mechanism.** Enzyme reactions at various fixed B concentrations and varying A concentration are observed. (a) If A and B both bind before product release, sequential addition, both the slope and the intercept drop as [B] increases and the lines intersect in the second quadrant. (b) If the enzyme follows ping-pong mechanism, only the intercept decreases as [B] increases and the lines remain parallel.

Work by Greenberg showed that RhlI is an ordered bi-ter enzyme with SAM binding first followed by C4-ACP binding and the ordered release of holo-ACP, C4-HSL, and MTA (Figure 12). Therefore, RhlI reaction will follow equations 11 and 13 and the double reciprocal plot of $\frac{1}{V}$ vs $\frac{1}{[\text{SAM}]}$ at various fixed [C4-ACP] would show both slope and intercept-effects.

**Figure 12. Cleland diagram of RhlI catalyzed ordered bi-ter reaction.** “E” denotes free RhlI enzyme while “A,” “B,” “P,” “Q,” and “R” represents SAM, C4-ACP, holo-ACP, C4-HSL, and MTA, respectively.
However, to simplify the equation, if one of the substrates were to be at saturating concentrations (e.g. \([A]\gg [B], K_{m}^{A}\)), any term not including the \([A]\) term would drop out to form:

\[
V = \frac{V_{\text{max}}[A][B]}{K_{m}^{A}+[A][B]} \quad (15)
\]

then the \([A]\) term in the remaining equation would cancel out, simplifying the equation to:

\[
V = \frac{V_{\text{max}}[B]}{K_{m}^{B}+[B]} \quad (16)
\]

which is mathematically identical to the Michaelis-Menten equation. Therefore, by keeping one substrate, “A”, at a saturating concentration and varying the concentration of the other substrate, “B”, the apparent \(K_{m}\), \(V_{\text{max}}\), and \(k_{\text{cat}}\) values of the enzyme associated with “B” substrate can be determined, which will be referred to as “V vs [S]” kinetics. In this study, substrate “A” corresponds to SAM and substrate “B” corresponds to C4-ACP

**Enzyme Inhibition**

Inhibitors can decrease enzyme activity in one of three ways: competitive inhibition, uncompetitive inhibition, and mixed mode of inhibition (Figure 13). In competitive mode of inhibition, the inhibitor is competing with the substrate for the same enzyme form; in uncompetitive mode of inhibition, the inhibitor is not competing with the substrate and binds to a different enzyme form; and in mixed mode of inhibition, the inhibitor binds with both the same and different forms of the enzyme.
Figure 13. **Modes of inhibition.** (a) Competitive mode of inhibition: the inhibitor competes with the substrate for the same enzyme form. (b) Uncompetitive mode of inhibition: the inhibitor and the substrate bind to different enzyme forms. (c) Mixed mode of inhibition: the inhibitor binds to both the same and different enzyme forms. Noncompetitive inhibition is a special case of mixed mode when the $K_i$ and $K_{ii}$ values are equal.

The mode of inhibition can be determined by conducting the $V$ vs $[S]$ kinetics assay at different inhibitor concentration and the resulting rate curves will fit to the following form of the Michaelis-Menten equation:

$$V = \frac{V_{\text{max, app}}[S]}{K_m^{\text{app}} + [S]}$$  \hspace{1cm} (17)

If the inhibitor is targeting the same form of the enzyme to which the variable substrate binds, it would be inhibiting the enzyme competitively and mathematically, the $V_{\text{max, app}}$ and the $K_m^{\text{app}}$ would be:
\[ V_{\text{max}}^{\text{app}} = V_{\text{max}} \]  
\[ K_{m}^{\text{app}} = K_{m}(1 + [I]/K_{i}) \]

since \( K_{m}^{\text{app}} \) changes while \( V_{\text{max}}^{\text{app}} \) remains unaffected, according to the Lineweaver-Burk equation, the inhibition produces a slope-effect, denoted by \( K_{i} \) (binding affinity of the inhibitor to E form). Mechanistically, as substrate concentration increases, Le Chatlier’s principle predicts that the reaction would be pushed forward to form more EA complex, which would decrease the concentration of free enzyme, E. The decrease in E concentration would push the EI complex to revert to E + I. Therefore, the substrate can outcompete the inhibitor at high concentrations (closer to the y-axis of the double reciprocal plot) and reach \( V_{\text{max}} \) (no intercept-effect). However, the interference in binding increases the concentration of substrate for half-maximal rate (\( K_{m} \)), thus increasing the \( \frac{K_{m}}{V_{\text{max}}} \), or the slope (slope-effect). This inhibition pattern can be generalized to the following observation: if the inhibitor binds with the same form of the enzyme, “E,” as the variable substrate or binds with an enzyme form reversibly connected to “E” form, there is slope-effect.

If the inhibitor is targeting a different form of the enzyme as the variable substrate, it would be inhibiting the enzyme uncompetitively and the \( V_{\text{max}}^{\text{app}} \) and the \( K_{m}^{\text{app}} \) would be:

\[ V_{\text{max}}^{\text{app}} = \frac{V_{\text{max}}}{(1+[I]/K_{i})} \]  
\[ K_{m}^{\text{app}} = \frac{K_{m}}{(1+[I]/K_{i})} \]

since \( V_{\text{max}}^{\text{app}} \) changes, the Lineweaver-Burk equation dictates that this inhibition would cause an intercept-effect, denoted by \( K_{i} \) (binding affinity of the inhibitor to the ES form).
Moreover, as both the $K_m^{app}$ and the $V_{max}^{app}$ are changed by the same ratio, $(1 + \frac{[I]}{K_{ii}})$, the slope, $\frac{K_m}{V_{max}}$, remains the same. Mechanistically, since the inhibitor is binding to a different enzyme form than the substrate binding to, increasing the substrate concentration cannot overcome the inhibition; therefore, even at saturating substrate concentration (near y-axis), the apparent $V_{max}$ is lower than true $V_{max}$ (intercept-effect).

As noted above, slope-effect is only observed if the inhibitor and the variable substrate are binding to the same enzyme form or reversibly connected enzyme forms; in uncompetitive inhibition, the inhibitor binds to a different form of the enzyme and no slope-effect is observed. This inhibition pattern can be generalized as: if the inhibitor and the variable substrate bind with different forms of the enzyme, there is intercept-effect.

If the inhibitor is targeting both the same and a different form of the enzyme as the variable substrate, it would be inhibiting the enzyme via mixed model inhibition and the $V_{max}^{app}$ and the $K_m^{app}$ would be:

$$V_{max}^{app} = \frac{V_{max}}{(1 + \frac{[I]}{K_{ii}})}$$ (22)

$$K_m^{app} = \frac{K_m(1 + \frac{[I]}{K_{ii}})}{(1 + \frac{[I]}{K_{ii}})}$$ (23)

since $K_m^{app}$ and $V_{max}^{app}$ change by differing ratios, the Lineweaver-Burk equation indicates that the inhibition would cause both slope and intercept-effects, denoted by having both $K_{ia}$ and $K_{ii}$. In this case, since the inhibitor also binds with a different form of the enzyme, $V_{max}$ is affected and there is intercept-effect. Since the inhibitor and the substrate are binding to the same form of the enzyme, there is also slope-effect. In a special case of mixed model inhibition, in which the inhibitor binds to two different enzyme forms
equally well, the $K_{ii}$ value would equal $K_{is}$ value and the $K_{m}^{app}$ value would be equal to the $K_{m}$ value and only the $V_{max}$ would be affected.

Once the $V$ vs $[S]$ assays in varying $[I]$ is conducted, the data can be fitted to all the models and the best fit can be determined by Akaike’s information criterion method (AIC), which compares the scatter about the fit and the degrees of freedom associated with each model and assay. In this model, the different inhibition models are listed in order of increasing complexity in the following manner: competitive, noncompetitive, uncompetitive, and mixed model. The AIC model compares two inhibition patterns at a time, the simpler model (fewer parameters) assigned “model 1” and the more complex model (more parameters) assigned “model 2,” with the following equation:

$$\Delta AIC = N \cdot \ln \frac{SS2}{SS1} + 2DF = AIC_1 - AIC_2$$

(24)

in which $SS1$ and $SS2$ refers to sum of the square of the scatter about the fit in inhibition models 1 and 2 and $\Delta DF$, the difference in the degrees of freedom due to the parameters associated with the inhibition models. The simpler model is expected to have bigger/more scatter and higher degree of freedom; therefore, the $\ln \frac{SS2}{SS1}$ term is expected to be negative and the $\Delta DF$ term, positive. If the $\Delta AIC$ is negative, it would indicate that the scatter in the simpler model 1 was greater than expected thus model 2 would be a better fit, and if the $\Delta AIC$ is positive, the opposite would be true and model 1 would be a better fit. The overall probability of one model being the better fit over the other is determined by the following equation:

$$probability = \frac{\frac{1}{\frac{1}{e^{\Delta AIC}} + 1}}{\frac{1}{e^{\Delta AIC}} + 1}$$

(25)

By repeating the AIC comparison between each of the inhibition models, the best fit can be determined, thereby identifying the mode of inhibition.
Although assaying the enzyme with various fixed inhibitor concentration and variable substrate concentration is useful in determining the mode of inhibition and the binding affinity of the inhibitor to the enzyme, it is a laborious and time-consuming process. To quickly determine whether a compound is inhibiting, the enzyme is assayed at fixed substrate concentration and variable inhibitor concentration. The kinetics data from this test shows the potency of the inhibitor as the concentration of the inhibitor required to reduce the enzyme initial rate by half, or the IC\textsubscript{50} value; however, the IC\textsubscript{50} value alone cannot be used to determine the mechanistic mode of inhibition or the binding affinity of the inhibitor for the enzyme. The IC\textsubscript{50} value can be calculated by the following equation:

\[
V = \frac{V_{\text{max}} - V_{\text{min}}}{[I]^h + IC_{50}^h}
\]

in which \(V_{\text{max}}\) and \(V_{\text{min}}\) refers to the starting and the minimum activities, respectively; \([I]\), the inhibitor concentration; and \(h\), the hill constant (Figure 14).\textsuperscript{54}

**Figure 14.**  Representative IC\textsubscript{50} curve. The enzyme rate with no inhibitor is the baseline, denoted \(V_{\text{max}}\). The concentration of the inhibitor required to reach half \(V_{\text{max}}\) rate is IC\textsubscript{50}. 
Under competitive mode of inhibition, the $K_i$ and the IC$_{50}$ values are related by:

$$K_i = \frac{IC_{50}}{1 + \frac{S}{K_m}}$$  \hspace{1cm} (27)

under uncompetitive mode:

$$K_i = \frac{IC_{50}}{1 + \frac{K_m}{S}}$$  \hspace{1cm} (28)

and under noncompetitive mode:

$$K_i = IC_{50}$$  \hspace{1cm} (29)

However, if a compound were to bind at an allosteric site and alter the enzyme structure to promote product release (increase $k_{cat}$ and maybe decrease $K_m$) or substrate binding (decrease $K_m$), the enzymatic rate would increase. If the enzyme is assayed with fixed concentration of the substrate and variable concentration of the agonist, EC$_{50}$ is the concentration of the activator at which the enzyme rate is half way between the baseline and the maximum. and this agonistic effect would be modeled by the EC$_{50}$ equation:

$$V = V_{min} + \frac{[A]^h(V_{max} - V_{min})}{([A]^h + EC_{50}^h)}$$  \hspace{1cm} (30)

where $V_{min}$ and $V_{max}$ refers to the starting and the maximum rates, respectively; [A], the activator concentration; and h, the hill constant (Figure 15).
Representative EC<sub>50</sub> curve. The enzyme rate with no activator is the baseline and the maximum enzyme rate reached with addition of the activator is denoted \( V_{\text{max}} \). The concentration of the activator required to reach halfway between the baseline and \( V_{\text{max}} \) is EC<sub>50</sub>.

**Thesis objective**

The objective of this thesis is to determine how altering the acyl-chain, chirality, and the head-group polarity of native AHL signal molecule changes the activity of RhlI to determine the moieties of greatest important and understand the mechanism of their effect.

The native AHL product, butanoyl-L-homoserine lactone, was used as the control. The L-homoserine lactone headgroup alteration was the focus of the first generation of AHL analogs (compounds 1-12; Table 2). Modifications to the lactone ring (compounds: 2-4, 8-12), the chirality (compounds: 5, 12), and the tail-headgroup linkage (compounds: 6, 7, 10-12). Variations in the acyl-chains were introduced in the next generation of AHL analogs (Table 3) which was combined with the lessons learned from the first generation of AHL derivative to target headgroup hydrophobicity with thiolactone, cyclopentyl, and non-lactone compounds (compounds 8, 43-64, 76, 77; 4, 65-68; 2-3, 9-11, 69-73; 76-77; Table 4, 5), D vs L headgroup chirality (compounds 5, 28-36; 12, 41-42; 56-64, 75, 77;
Table 4), sulfonamide linkage (compounds 6, 37-40; 12, 41-42; Table 4), and acyl-chain length and substitution at the C3 position (Compounds 13-55, 57-68, 70-73, 78-82; Table 4, 5). The acyl-chain effects were further correlated by testing RhII activity with acyl-ACP substrates of varying acyl-chain moieties (compounds 83-87; Table 4; Figure 16) and testing the activity in the presence of alkyl-ACP (inactive acyl-ACP analogs; compounds 88-91; Table 6; Figure 16). Finally, the importance of the ACP moiety in binding was tested by observing RhII activity in the presence of alkyl-CoAs (compounds 92-94; Table 4; Figure 16).

This is the first study to test the effects of modified AHLs on AHL synthases. The work described in this thesis is the basis upon which future of rational inhibitor design can be based.

Table 2.  Structures of 1st generation AHL-based small molecules tested for RhII inhibition.
Table 3. Variations in the acyl-chains for 2nd generation of AHL-based small molecules tested for RhlI inhibition.

<table>
<thead>
<tr>
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Table 4. 2nd generation of AHL-based small molecules tested for RhII inhibition.\(^a\)

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\(\text{a = letter notations for R-groups refer to structures listed in Table 3}\)
Table 5. Nonlactone derivatives tested for RhII inhibition

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Table 6. Acyl-ACP, alkyl-ACP, and alkyl-CoA derivatives

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Figure 16. Structures of Acyl-ACP and CoA. (a) Apo-ACP is a relatively small protein made up of four helices. When apo-ACP is linked with a pantetheine linker, it is called holo-ACP. Acylated holo-ACPs are called acyl-ACPs. (b) Free coenzyme A consists of a nucleotide connected to the pantetheine linker. Acylated CoAs are called acyl-CoAs.
CHAPTER TWO: MATERIALS AND METHODS

Materials and Equipment

The following reagents were purchased from Sigma Aldrich: (S)-(+)–α-methoxyphenylacetic acid (MPA), 2-(N-morpholino)ethanesulfonic acid hydrate (MES), 2,2-dimethyl-1,3-dioxane-4,6-dione (Meldrum’s acid), 2,6-dichloroindophenol (DCPIP), 4-(2-hydroxyethyl)piperazine-1-ethanesulfonic acid (HEPES), ammonium sulfate, D-homoserine lactone hydrochloride, DL-homocysteine thiolactone, ethylenediaminetetraacetic acid (EDTA), glycerol, L-homoserine lactone hydrochloride, magnesium sulfate anhydrous (MgSO₄), maltose, N-(3-dimethylaminopropyl)-N′-ethylcarbodiimide hydrochloride (EDC), N-hydroxysuccinimide, Nα-tosyl-L-lysine chloromethyl ketone hydrochloride (TLCK), phenylmethanesulfonyl fluoride (PMSF), protamine sulfate, S-(5′-adenosyl)–L-methionine chloride (SAM), sucrose, toluene, trichloroacetic acid, coenzyme A free acid (CoA-SH), butanoyl-coenzyme A (C4-CoA), hexanoyl-coenzyme A (C6-CoA), octanoyl-coenzyme A (C8-CoA), decanoyl-coenzyme A (C10-CoA), dodecanoyl-coenzyme A (C12-CoA), deoxyribonuclease I (DNase), ribonuclease A (RNase), lysozyme, ampicillin, kanamycin, chloramphenicol, kanamycin, spectinomycin, and streptomycin. Thermo Fisher Scientific supplied the following: 2-propanol (IPA), acetonitrile, agar, ammonium acetate, bacterial protein extraction reagent (B-PER), chloroform, diethyl ether, dimethylformamide (DMF), dimethyl sulfoxide (DMSO), ethyl acetate, hydrochloric acid (HCl), imidazole, isopropyl β-D-1-thiogalactopyranoside (IPTG), Lennox broth, manganese sulfate (MnSO₄), methanol,
hexane, potassium carbonate, silica, sodium bicarbonate, tricine, and tris base. Butyric acid, hexanoic acid, octanoic acid, decanoic acid, dodecanoic acid, and trifluoroacetic acid (TFA) were obtained from Acros Organics. Alfa Aesar provided bromobutane, bromohexane, bromooctane, bromodecane, magnesium chloride (MgCl₂), and triethyl amine. The 3kD and 10kD spin filter columns and celite were purchased from EMD. Amylose resin, ethanol, and 0.22 µm sample filters were provided by NEB, Ultra Pure, and Costar, respectively. Dr. Peter Tipton (University of Missouri, Columbia), Dr. E. Peter Greenberg (University of Washington), and Dr. Michael Burkart (University of California-San Diego) provided clones for purifying RhlI, apo-ACP, and Sfp, respectively. Sfp was purified by Levi Mitchell and Nhu Lam (Both from Nagarajan lab, Boise State University). Various AHL analogs were supplied by Dr. Helen Blackwell (University of Wisconsin, Madison; compounds 1, 3-9, 12-55, 65-73), Dr. Eric Brown (Boise State University; compounds 2, 10, 11), and Neil Rexrode (Nagarajan lab, Boise State University; compounds 34, 36).

All UV-Vis spectrophotometric data was collected with Thermo Scientific Evolution 260 Bio UV-Vis spectrophotometer using Fisher 1 cm path length quartz cuvettes (14-385-928C). HPLC data was obtained with Thermo Scientific Dionex UltiMate 3000 UHPLC+ focused HPLC with Dionex UltiMate 3000 Automated Fraction Collector using Thermo Scientific Hypersil Gold C18 reverse-phase UHPLC column (25002-054630) or Thermo Scientific Hypersil Gold C18 reverse-phase preparative column (25005-159070). All mass spectrometry data was collected with Bruker maXis Quadrupole Time-of-Flight (QTOF) mass spectrometry and analyzed with the Bruker
Compass Data Analysis software. All the kinetics data were processed using GraphPad Prism 7.

**Methods**

**HPLC Methods**

HPLC and the fraction collector was used to isolate and collect alkyl-CoA and to monitor the synthesis and purity of acyl-ACP synthesis. Solvent “A” consisted of 25 mM ammonium acetate pH 5 solution. Solvent “B” is acetonitrile (ACN) + 0.1% trifluoroacetic acid (TFA), and solvent “C” is H₂O + 0.1% TFA. To isolated alkyl-CoA, the filtered sample was injected into a C18 reverse-phase preparatory column equilibrated with the initial solvent condition of 95.0% solvent A and 5.0% solvent B. The analyte was analyzed using a solvent gradient of 95.0% solvent A and 5.0% solvent B to 30.0% solvent A and 70.0% solvent B over 11 minutes at a flow rate of 3.0 mL/min (Table 7).

To monitor acyl-ACP synthesis, the analyte sample was injected into a C18 reverse-phase UHPLC column equilibrated with 25.0% solvent B and 75.0% solvent C. The acyl-ACP peaks were separated from apo-ACP peak with a solvent gradient of 25.0% solvent B and 75.0% solvent C to 75.0% solvent B and 25.0% solvent C over 10 minutes at a flow rate of 600 µL/min (Table 8). Due to the similar retention time for hexanoyl/hexyl-ACP and apo-ACP, a shallower gradient of 25.0% solvent B and 75.0% solvent C to 75.0% solvent B and 25.0% solvent C over 60 minutes at a flow rate of 200 µL/min was used (Table 9). The column and the sample loop were washed with methanol for 5 minutes between each run.
Table 7. Alkyl-CoA separation method

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<tr>
<th>Time (min)</th>
<th>Flow (mL/min)</th>
<th>%A&lt;sup&gt;b&lt;/sup&gt;</th>
<th>%B&lt;sup&gt;c&lt;/sup&gt;</th>
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<td>11.00</td>
<td>3.000</td>
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<sup>a</sup> Preparatory column  
<sup>b</sup> 25 mM ammonium acetate pH 5  
<sup>c</sup> acetonitrile (ACN) + 0.1% trifluoroacetic acid (TFA)  
<sup>d</sup> sample injection at Time 0.00

Table 8. ACP separation method

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Flow (mL/min)</th>
<th>%B&lt;sup&gt;b&lt;/sup&gt;</th>
<th>%C&lt;sup&gt;c&lt;/sup&gt;</th>
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<td>10.00</td>
<td>0.600</td>
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<sup>a</sup> UHPLC column  
<sup>b</sup> ACN + 0.1% TFA  
<sup>c</sup> H<sub>2</sub>O + 0.1% TFA  
<sup>d</sup> sample injection at Time 0.00

Table 9. C6-ACP separation method

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<tr>
<td>60.00</td>
<td>0.200</td>
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<td>25.0</td>
</tr>
</tbody>
</table>

<sup>a</sup> UHPLC column  
<sup>b</sup> ACN + 0.1% TFA  
<sup>c</sup> H<sub>2</sub>O + 0.1% TFA  
<sup>d</sup> sample injection at time 0.00

RhII Purification

RhII was purified via a previously described method with modifications.<sup>52</sup> *E. coli* with RhII plasmid was grown on an agar plate (20 g Lennox broth and 10 g agar per 1 L of medium) with ampicillin (100 µg/mL) for 12 hours at 37°C. A colony was isolated and was used to inoculate 25 mL of Lennox broth with ampicillin (100 µg/mL) and incubated
at 37°C with shaking (225 rpm) for 8-12 hours or until visible turbidity. The “mini-
growth” was then transferred over to 1 L of Lennox broth (20 g broth/L) with ampicillin
(100 µg/mL). The broth was incubated with shaking (225 rpm) at 37°C. When the OD_{600}
value reached 0.6-0.8, IPTG was added to 0.5 mM final concentration to promote protein
expression. The cell culture was incubated for 3 hours at room temperature. The growth
media was then spun down at 5,000 x g at 4°C for 15 minutes to collect cell paste. The
cell pellet was resuspended in “Buffer A,” which is composed of 200 mL of 50 mM Tris-
HCl, pH 7.5, containing 0.2 M NaCl, 1 mM ethylenediaminetetraacetic acid (EDTA), 0.1
mM phenylmethanesulfonyl fluoride (PMSF), 0.1 mM Nα-tosyl-L-lysine chloromethyl ketone hydrochloride (TLCK), 0.4 M sucrose, and 2.5% (v/v) glycerol. The resuspended
mixture was then lysed via sonication at 15,000 psi. The lysate was spun down for 40
minutes at 10,000 x g and at 4°C. Protamine sulfate was added to the supernatant to a
final concentration of 6 mg/g of cell pellet to cause nucleic acids to precipitate. The
nucleic acid precipitates were removed via centrifugation for 20 minutes at 10,000 x g
and at 4°C. The supernatant was then loaded onto an amylose column that has been
equilibrated with 5x bed volume with buffer A. The column was washed with 5x bed
volume with buffer A. RhlI was eluted out using buffer A with 10 mM maltose added.
The presence and purity of RhlI was checked with SDS-PAGE gel electrophoresis. The
protein sample was concentrated using 10 kD spin filter, the concentration checked via
UV-Vis spectrophotometry (ε_{280} = 107510 M^{-1}cm^{-1}), and stored in buffer A with 20%
glycerol at -80°C.
Apo-ACP Purification

Apo-ACP was purified with a well-established method. E. coli DK574 with pJT94 was grown on an agar plate (20 g Lennox broth and 10 g agar per 1 L of medium) with kanamycin (25 µg/mL), streptomycin (50 µg/mL), spectinomycin (50 µg/mL), and chloramphenicol (25 µg/mL) for 12 hours at 37°C. An isolated colony was used to inoculate 25 mL of Lennox broth with kanamycin (25 µg/mL), streptomycin (50 µg/mL), spectinomycin (50 µg/mL), and chloramphenicol (25 µg/mL) and incubated at 37°C with shaking (225 rpm) for 8-12 hours or until visible turbidity. The “mini-growth” was then transferred over to 1 L of Lennox broth (20 g broth/L) with kanamycin (25 µg/mL), streptomycin (50 µg/mL), spectinomycin (50 µg/mL), and chloramphenicol (25 µg/mL) and incubated at 37°C with stirring until OD$_{600}$ value reached 0.6-0.8. IPTG was then added to a final concentration of 1 mM to promote protein expression. The cell culture was incubated at 37°C for a further 3 hours then collected via centrifugation at 5,000 x g for 15 minutes. The cell paste was resuspended in 2 mL of B-PER reagent, 1 mL of lysozyme (2 mg/mL) 20 μL each of DNase (1 mg/mL) and RNase (1 mg/mL) and 25 μL of phenylmethylsulfonyl fluoride (13 mg/750 μL 2-propanol) to lyse the cells and their nucleic acids. The lysate mixture was incubated with gentle shaking at room temperature for 20 minutes. The lysate was spun down for 30 minutes at 20,000 x g and at 4°C. The supernatant was collected and was added MgCl$_2$ and MnSO$_4$ to final concentrations of 25 mM and 1.2 mM, respectively and incubated at 37°C for 4 hours. Extraneous proteins were precipitated by slow addition of 2-propanol to 50% initial volume while on ice. The precipitates were removed via centrifugation at 14,000 x g for 30 minutes. The supernatant was stirred with 2 g of DE52 diaminoethyl cellulose resin
overnight at 4°C. The mixture was packed in to a column and washed with 10 x bed volume with 10 mM lithium 4-morpholineethane-sulfonate (MES) pH 6.1 and 0.25 mM LiCl. The protein was eluted out using 10 mM lithium MES pH 6.1 and 0.5 M LiCl. The presence and purity of apo-ACP was checked via Tris/Tricine SDS-PAGE gel electrophoresis. Apo-ACP samples were polled and the protein was precipitated with the addition of 0.02% (0.2 mg/mL) sodium deoxycholate and 5% (50 mg/mL) trichloroacetate (w/v). The mixture was incubated at 37°C with gentle shaking for 30 minutes. The mixture was then spun at 21,000 x g for 30 minutes to collect ACP pellet. The ACP pellet was resuspended in 60 mL of 0.5 M Tris-HCl pH 8.0 and concentrated using 3 kD spin filter. The concentration was determined via UV-Vis spectrophotometry ($\varepsilon_{280} = 1490 \text{ M}^{-1}\text{cm}^{-1}$), and stored in 10 mM MES pH 6.1 + 20 % glycerol at -80 °C.

**Alkyl-CoA Synthesis**

To a solution of coenzyme A, free acid (CoA-SH; 50 mg, 65.1 µmol) and alkyl bromide (120.2 µmol) in 2mL of 1:1 water:DMF, potassium carbonate was added to pH 8-9 (Figure 17). The reaction was stirred under nitrogen overnight and the completion of the reaction was tested by checking for the reduction of DCPIP by unreacted CoA-SH. The reaction mixture was diluted with water to a final volume of 5 mL and extracted with 5 mL of diethyl ether. The aqueous mixture was filtered with 0.2 µm filter and the alkyl-CoA was isolated and collected with semi-prep HPLC using the Alkyl-CoA separation method (Table 5). Organic solvent was removed via evaporation by a gentle stream of nitrogen through the product solution and the aqueous solution was lyophilized to yield alkyl-CoA powder.
Figure 17. Synthesis of alkyl-CoA. Alkyl bromide was reacted with free-CoA in basic solution to produce alkyl-CoA.

92. Butyl-Coenzyme A. ESI-TOF: expected m/z [M + H\(^+\)] = 824.1851, observed [M + H\(^+\)] = 824.1835. (Appendix Figures A14)

93. Hexyl-Coenzyme A. ESI-TOF: expected m/z [M + H\(^+\)] = 852.2164, observed [M + H\(^+\)] = 852.2147. (Appendix Figures A15)

94. Octyl-Coenzyme A. ESI-TOF: expected m/z [M + H\(^+\)] = 880.2477, observed [M + H\(^+\)] = 880.2444. (Appendix Figures A16)

95. Decyl-Coenzyme A. ESI-TOF: expected m/z [M + H\(^+\)] = 908.2790, observed [M + H\(^+\)] = 908.2733. (Appendix Figures A17)

Alkyl-/Acyl-ACP synthesis

Alkyl-/acyl-pantetheine was transferred from CoA to apo-ACP via phosphopanetheinyl transferase (Sfp) catalyzed reaction (Figure 18).\(^{22}\) The reaction mixture consisted of 50 mM Tris-HCl pH 6.8, 10 mM magnesium chloride, 600 µM apo-ACP, 750 µM alkyl-/acyl-CoA, and 10 µM Sfp, with the CoA-substrate being the last to be added. Alkyl-/acyl-CoAs with aliphatic chains of 6 and fewer carbons were added all at once. Alkyl-/acyl-CoAs with aliphatic chains of 8 or more carbons were added in three equal portions every 15 minutes to prevent CoA precipitating and crashing out of the predominantly aqueous reaction mixture. The reaction was incubated at 37 °C and checked via HPLC every 30 minutes for completion using ACP separation method (Table 6). However, C6-ACP separation method was used to check hexyl- and hexanoyl-ACP synthesis since C6-ACP, C6-IACP, and apo-ACP have similar retention time (Table 7). For completion, reactions lasted 2 hours for C4-ACP and C4-IACP, between 4 and 5
hours for C6-ACP and C6-IACP, and 6 to 7 hours for C8- to C12-ACP and IACPs. Upon completion, ammonium sulfate was added to the reaction to 75% saturation and kept at 4 °C for 1 hour to precipitate Sfp.\textsuperscript{56} The Sfp precipitates were pelleted by centrifugation at 15,000 x g for 15 minutes. The supernatant was desalted and concentrated by 3 kD spin filter spun at 5,000 x g at 4 °C. The concentration was determined via UV-Vis spectrophotometry (ε\textsubscript{280} = 1490 M\textsuperscript{-1}cm\textsuperscript{-1}), and stored in 10 mM MES pH 6.1 + 20 % glycerol at -80 °C.

![Diagram](image)

**Figure 18.** *Sfp* catalyzed acyl-pantetheine transfer reaction. *Sfp* catalyzes the transfer of pantetheine linker from a CoA to apo-ACP to produce alkyl-acyl-ACP.

\textbf{83-87. Butanoyl-, Hexanoyl-, Octanoyl-, Decanoyl-, and Dodecanoyl-ACP.} Mass confirmed in previous work from Nagarajan laboratory.\textsuperscript{22}

\textbf{88. Butyl-ACP.} ESI-TOF: expected mass = 8904.6, observed mass = 8904.3.

(Appendix Figures A10)

\textbf{89. Hexyl-ACP.} ESI-TOF: expected mass = 8932.6, observed mass = 8932.4.

(Appendix Figures A11)

\textbf{90. Octyl-ACP.} ESI-TOF: expected mass = 8960.7, observed mass = 8960.4.

(Appendix Figures A12)

\textbf{91. Decyl-ACP.} ESI-TOF: expected mass = 8988.7, observed mass = 8988.4.

(Appendix Figures A13)
Purification of D-homocysteine thiolactone

Enantiomerically pure sample of D-homocysteine thiolactone was obtained using a published method with modification (Figure 19).57 A solution of DL-homocysteine thiolactone (768.15 mg; 5.0 mmole), triethylamine (N(Et)_3; 1.2 mL), N-hydroxysuccinimide (690.5 mg; 6 mmole), S-(+)-2-methoxyphenylacetic acid (MPA; 1.0 g; 6.0 mmole), and 1-ethyl-3-(3-dimethylaminopropyl)carbodiimide HCl (EDC; 1.150 g; 6 mmole) in 100 mL chloroform was stirred overnight at 4°C. This reaction resulted in the synthesis of L-homocysteine thiolactone-containing (3S,2’S)-3-(2′-methoxy-2′-phenyl-)acetamido-2-thiophenone and D-homocysteine thiolactone-containing (3R,2’S)-3-(2′-methoxy-2′-phenyl-)acetamido-2-thiophenone. The solution was then washed with water (50 mL), NaHCO₃ (5%, 20 mL), HCl (2M, 20 mL), and brine (saturated, 40 mL). After drying with MgSO₄, the solvent was removed under low pressure. The two diastereomers were separated by silica gel column chromatography using hexane-ethyl acetate solution (7:3). The more polar, D-thiolactone-containing isomers were isolated and refluxed in ethanol-4M HCl (2:1; 15 mL) solution overnight. Ethanol was removed with rotary evaporation. Water was added to the resulting solution to make it 10 mL and washed with toluene (10 mL x 3). The aqueous layer was isolated and the solvent was removed under high vacuum. The resulting D-thiolactone-HCl mixture was recrystallized with anhydrous 2-propanol.
Figure 19. Purification of pure D-homocysteine thiolactone. A racemic mixture was acylated with MPA in an EDC-coupled reaction to form diastereomers with distinct polarity. The two diastereomers separated via silica gel column chromatography. D-Homocysteine thiolactone was obtained by removing the MPA by refluxing the compound in HCl.

**D-Homocysteine thiolactone HCl.** (comparable to literature value\(^57\)) m.p. 176-178°C \([\alpha]_D^{25} -36.3\) (0.05 mg/mL in H\(_2\)O). \(^1\)H NMR (600 MHz, D\(_2\)O): \(\delta\) 2.10 (1H, m, 4\(\alpha\)-H), 2.62 (1H, m, 4\(\beta\)-H), 3.33 (1H, m, 5\(\alpha\)-H), 3.24 (1H, m, 5\(\beta\)-H), 4.29 (1H, s, 3-H). (Appendix Figure B1)

**Synthesis of N-acyl-D-homocysteine thiolactones**

The D-thiolactone head-group was acylated using a previously published protocol with modification (Figure 20).\(^43\) The thiolactone head-group (25 mg; 162.7 µmole) was stirred overnight (12 hours) at room temperature with DCC (33.6 mg; 162.7 µmole), N-hydroxysuccinimide (18.7 mg; 162.7 µmole), triethylamine (50 µL) in acetonitrile (10 mL) with the appropriate carboxylic acid (162.7 µmole: butyric acid: 14.9 µL; hexanoic
acid: 20.6 µL; octanoic acid: 25.8 µL; decanoic acid: 28 mg; dodecanoic acid: 32.6 mg).

The resulting mixture was cooled to 4 °C for an hour and then filtered with celite to remove \(N,N^\prime\)-dicyclohexylurea precipitate. The filtrate solution was washed with water, HCl (2M), NaHCO\(_3\) (saturated), and brine (saturated) (3x10 mL each). After drying with MgSO\(_4\), and removing the solvent under low pressure, the product was purified with a silica gel column using 7:3 hexane: ethyl acetate eluent.

Figure 20. Acylation of D-thiolactone headgroup. D-thiolactone headgroup was acylated by a DCC-coupled reaction in acetonitrile. Urea byproduct was removed via filtration and excess carboxylic acid was removed via silica gel column chromatography.

56. N-Butanoyl-D-homocysteine thiolactone. \(^1\)H NMR (600 MHz, CDCl\(_3\)): \(\delta\) 0.93 (3H, t, \(J = 7.3\) Hz, CH\(_3\)), 1.65 (2H, m, CH\(_2\)), 1.90 (1H, m, -lac), 2.20, (2H, dt, \(J = 7.3, 3.9\) Hz, CH\(_2\)), 2.93 (1H, m, -lac), 3.23 (1H, ddd, \(J = 11.4, 4.5, 1.0\) Hz, -lac), 3.33 (1H, ddd, \(J = 11.7, 11.7, 5.1\) Hz, -lac), 4.49 (1H, ddd, \(J = 12.9, 6.4, 6.4\) Hz, -lac), 5.92 (1H, s, NH); \(^{13}\)C NMR (CDCl\(_3\), 600 MHz): \(\delta\) 13.9, 19.2, 27.8, 32.4, 38.5, 59.7, 173.7, 205.8; ESI-TOF: expected m/z [M + H\(^+\)] = 188.0734, observed [M + H\(^+\)] = 188.0763. (Appendix Figures A1, B2-B6)

57. N-Hexanoyl-D-homocysteine thiolactone. \(^1\)H NMR (600 MHz, CDCl\(_3\)): \(\delta\) 0.87 (3H, t, CH\(_3\)), 1.29 (4H, m CH\(_2\)), 1.64 (2H, m, CH\(_2\)), 1.89 (1H, m, -lac), 2.22, (2H, dt, \(J = 7.5, 3.5\) Hz, CH\(_2\)), 2.97 (1H, m, -lac), 3.24 (1H, ddd, \(J = 11.4, 6.9, 1.1\) Hz, -lac), 3.34 (1H, ddd, \(J = 11.7, 11.7, 5.3\) Hz, -lac), 4.48 (1H, ddd, \(J = 12.8, 6.4, 6.4\) Hz, -lac), 5.83 (1H, s, NH); \(^{13}\)C NMR (CDCl\(_3\), 600 MHz): \(\delta\) 13.9, 22.4, 25.2, 27.6, 31.4, 32.2, 36.4, 59.6,
173.7, 205.6; ESI-TOF: expected m/z [M + H⁺] = 216.1047, observed [M + H⁺] = 216.1099. (Appendix Figures A2, B7-B9)

58. N-Octanoyl-D-homocysteine thiolactone. ¹H NMR (600 MHz, CDCl₃): δ 0.86 (3H, t, J = 6.9 Hz, CH₃), 1.26 (8H, m CH₂), 1.62 (2H, m, CH₂), 1.88 (1H, m, -lac), 2.22, (2H, dt, J = 7.5, 3.5 Hz, CH₂), 2.96 (1H, m, -lac), 3.23 (1H, ddd, J = 11.3, 4.4, 0.9 Hz, -lac), 3.34 (1H, ddd, J = 11.7, 11.7, 5.2 Hz, -lac), 4.48 (1H, ddd, J = 12.9, 6.4, 6.4 Hz, -lac), 5.82 (1H, s, NH); ¹³C NMR (CDCl₃, 600 MHz): δ 14.3, 22.8, 25.7, 27.8, 29.4, 29.9, 31.9, 32.5, 36.7, 59.8, 173.9, 205.8; ESI-TOF: expected m/z [M + H⁺] = 244.1360, observed [M + H⁺] = 244.1405. (Appendix Figures A3, B10-B14)

59. N-Decanoyl-D-homocysteine thiolactone. ¹H NMR (600 MHz, CDCl₃): δ 0.85 (3H, t, J = 6.9 Hz, CH₃), 1.25 (12H, m CH₂), 1.61 (2H, m, CH₂), 1.87(1H, m, -lac), 2.21, (2H, dt, J = 7.4, 3.6 Hz, CH₂), 2.95 (1H, m, -lac), 3.23 (1H, ddd, J = 11.4, 4.5, 1.1 Hz, -lac), 3.34 (1H, ddd, J = 11.5, 11.5, 5.1 Hz, -lac), 4.48 (1H, ddd, J = 12.7, 6.3, 6.3 Hz, -lac), 5.86 (1H, s, NH); ¹³C NMR (CDCl₃, 600 MHz): δ 14.0, 22.6, 25.5, 27.6, 29.2, 29.3, 29.4, 31.8, 32.2, 33.9, 36.4, 59.5, 173.6, 205.5; ESI-TOF: expected m/z [M + H⁺] = 272.1673, observed [M + H⁺] = 272.1720. (Appendix Figures A4, B15-B19)

60. N-Dodecanoyl-D-homocysteine thiolactone. ¹H NMR (600 MHz, CDCl₃): δ 0.86 (3H, t, J = 6.8 Hz, CH₃), 1.25 (16H, m CH₂), 1.62 (2H, m, CH₂), 1.88(1H, m, -lac), 2.21, (2H, dt, J = 7.5, 3.4 Hz, CH₂), 2.97 (1H, m, -lac), 3.23 (1H, ddd, J = 11.2, 4.4, 0.8 Hz, -lac), 3.34 (1H, ddd, J = 11.7, 11.7, 5.3 Hz, -lac), 4.47 (1H, ddd, J = 12.8, 6.2, 6.2 Hz, -lac), 5.81 (1H, s, NH); ¹³C NMR (CDCl₃, 600 MHz): δ 14.1, 22.6, 24.9, 25.5, 27.6, 29.2, 29.3, 29.4, 29.6, 31.9, 32.2, 33.9, 36.4, 59.6, 173.6, 205.5; ESI-TOF: expected m/z [M + H⁺] = 300.1986, observed [M + H⁺] = 300.2042. (Appendix Figures A5, B20-B24)
Synthesis of \(N\)-(3-Oxoacyl)-D-homocysteine thiolactones

The synthesis of \(N\)-(3-oxoacyl)-D-homocysteine thiolactones has been developed previously (Figure 21).\(^{58}\) A solution of Meldrum’s acid (300.0 mg; 2.1 mmole), DCC (515.4 mg; 2.5 mmole), 4-(dimethylamino)pyridine (279.7 mg; 2.3 mmole), and the appropriate carboxylic acid (2.1 mmole: butyric acid: 191.0 µL; hexanoic acid: 262.8 µL; octanoic acid: 329.9 µL; decanoic acid: 358.6 mg) in dichloromethane (10 mL) was stirred at room temperature overnight. The resulting mixture was cooled to 4 °C for an hour and then filtered with celite to remove \(N,N'\)-dicyclohexylurea precipitate. The filtrate was washed with HCl (2M, 20 mL). After drying with MgSO\(_4\), the solvent was removed under low pressure. A solution of D-homocysteine thiolactone HCl (30.0 mg; 0.2 mmole), triethylamine (35 µL) and the appropriate acylated Meldrum’s acid (0.2 mmole) in acetonitrile (10 mL) was prepared. The solution was stirred at room temperature for 1 hour and then refluxed overnight. The solvent was removed under low pressure and the product was then dissolved in ethyl acetate (10 mL). The solution was washed with NaHCO\(_3\) (saturated), KHSO\(_4\) (1M), and brine (saturated) (10 mL x 3 each). After drying with MgSO\(_4\), and removing the solvent under low pressure, the product was purified with a silica gel column using 7:3 hexane: ethyl acetate eluent.
Figure 21. Synthesis and purification of 3oxoacyl-D-thiolactones. Carboxylic acid was attached to Meldrum’s acid via DCC-coupled reaction. The acylated Meldrum’s acid was reacted with thiolactone headgroup under heat. This reaction produced 3oxoacyl-thiolactone product and small amounts of a side product, which was removed via silica gel column chromatography.

61. N-(3-Oxohexanoyl)-D-homocysteine thiolactone. $^1$H NMR (600 MHz, CDCl$_3$): $\delta$ 0.91 (3H, t, $J = 7.4$ Hz, CH$_3$), 1.61 (2H, m, CH$_2$), 1.99 (1H, m, -lac), 2.49, (2H, t, $J = 7.4$ Hz, CH$_2$), 2.83 (1H, m, -lac), 3.24 (1H, ddd, $J = 11.5$, 6.9, 1.0 Hz, -lac), 3.33 (1H, ddd, $J = 11.7$, 11.7, 5.2 Hz, -lac), 3.43 (2H, s, CH$_2$) 4.56 (1H, ddd, $J = 12.9$, 6.8, 6.8 Hz, -lac), 7.43 (1H, s, NH); $^{13}$C NMR (CDCl$_3$, 600 MHz): $\delta$ 13.7, 17.1, 27.7, 31.8, 46.0, 48.6, 59.5, 166.4, 204.7, 206.6; ESI-TOF: expected m/z [M + H$^+$] = 230.0845, observed [M + H$^+$] = 230.0896. (Appendix Figures A6, B25-B29)

62. N-(3-Oxooctanoyl)-D-homocysteine thiolactone. $^1$H NMR (600 MHz, CDCl$_3$): $\delta$ 0.86 (3H, t, $J = 7.1$ Hz, CH$_3$), 1.26 (4H, m CH$_2$), 1.56 (2H, m, CH$_2$), 1.99 (1H, m, -lac), 2.50, (2H, t, $J = 7.4$ Hz, CH$_2$), 2.82 (1H, m, -lac), 3.23 (1H, ddd, $J = 11.4$, 7.0, 1.1 Hz, -lac), 3.33 (1H, ddd, $J = 11.7$, 11.7, 5.2 Hz, -lac), 3.43 (2H, s, CH$_2$), 4.56 (1H, ddd, $J = 13.0$, 6.6, 6.6 Hz, -lac), 7.45 (1H, s, NH); $^{13}$C NMR (CDCl$_3$, 600 MHz): $\delta$ 14.0, 22.6, 23.3, 27.7, 31.3, 31.7, 44.1, 48.6, 59.5, 166.4, 204.7, 206.7; ESI-TOF: expected m/z [M + H$^+$] = 244.1360, observed [M + H$^+$] = 244.1433. (Appendix Figures A7, B30-B34)
63. **N-(3-Oxodecanoyl)-D-homocysteine thiolactone.** $^1$H NMR (600 MHz, CDCl$_3$): δ 0.84 (3H, t, $J = 7.1$ Hz, CH$_3$), 1.24 (8H, m CH$_2$), 1.55 (2H, m, CH$_2$), 1.99 (1H, m, -lac), 2.49 (2H, t, $J = 7.4$ Hz, CH$_2$), 2.81 (1H, m, -lac), 3.23 (1H, ddd, $J = 11.4$, 7.1, 1.0 Hz, -lac), 3.32 (1H, ddd, $J = 11.8$, 11.8, 5.3 Hz, -lac), 3.42 (2H, s, CH$_2$), 4.56 (1H, ddd, $J = 12.9$, 6.6, 6.6 Hz, -lac), 7.46 (1H, s, NH); $^{13}$C NMR (CDCl$_3$, 600 MHz): δ 14.2, 22.8, 23.6, 27.7, 29.14, 29.17, 31.7, 31.8, 44.0, 48.6, 59.4, 166.5, 204.7, 206.6; ESI-TOF: expected m/z [M + H$^+$] = 286.1471, observed [M + H$^+$] = 286.1472. (Appendix Figures A8, B35-B39)

64. **N-(3-Oxododecanoyl)-D-homocysteine thiolactone.** $^1$H NMR (600 MHz, CDCl$_3$): δ 0.85 (3H, t, $J = 7.1$ Hz, CH$_3$), 1.24 (12H, m CH$_2$), 1.56 (2H, m, CH$_2$), 1.99 (1H, m, -lac), 2.50, (2H, t, $J = 7.3$ Hz, CH$_2$), 2.83 (1H, m, -lac), 3.24 (1H, ddd, $J = 11.2$, 6.9, 1.0 Hz, -lac), 3.33 (1H, ddd, $J = 11.7$, 11.7, 5.2 Hz, -lac), 3.43 (2H, s, CH$_2$), 4.56 (1H, ddd, $J = 12.9$, 6.7, 6.7 Hz, -lac), 7.45 (1H, s, NH); $^{13}$C NMR (CDCl$_3$, 600 MHz): δ 14.3, 22.9, 23.6, 27.7, 29.2, 29.4, 29.5, 29.6, 31.8, 32.1, 44.1, 48.5, 59.4, 166.4, 204.7, 206.7; ESI-TOF: expected m/z [M + H$^+$] = 314.1784, observed [M + H$^+$] = 314.1791. (Appendix Figures A9, B40-B14)

**Mass Spectrometry**

Molecular mass of ACP derivatives was determined by high performance liquid chromatography mass spectrometry (HPLC-MS) using a high resolution Quadrupole Time of Flight (QTOF) instrument with electrospray ionization (ESI). The ESI source was operated at positive ion mode, the nebulizer pressure at 1.2 bar, nitrogen drying gas flow at 8 L/min, drying temperature at 200 °C, and the voltage of the capillary and the end-plate offset to 3000 V to -500 V. The mass range was set from 250 to 2900 m/z and
low concentration ESI tuning mix (Agilent Technologies, Santa Clara, California) was used to calibrate the instrument in the mass range. Ten microliters of samples were injected onto a Phenomenex Kinetex XB-C18 column (100 x 2.1 mm, 2.6μm) (Phenomenex Corporation, Torrance, California) at a flow rate of 0.3 mL/min followed by a simple linear gradient for sample desalting and separation. The initial eluent was 98% mobile phase A (99.9% water, 0.1% formic acid) and 2% B (99.9% acetonitrile, 0.1% formic acid) for 5 min and then mobile phase B was increased to 50% in 25 min. LC eluent was diverted to the waste during the first five minutes of the gradient to eliminate salts in the sample buffer.

Small molecule samples were prepared in concentrations of 0.5 mg/mL in methanol. Molecular mass of AHL analogs were determined with the instrument described above with direct sample injection via ESI inlet. The ESI source was operated at positive ion mode, the nebulizer pressure at 0.4 bar, nitrogen drying gas flow at 4 L/min, drying temperature at 200 °C, and the voltage of the capillary and the end-plate offset to 3000 V to -500 V. The mass range was set from 80 to 800 m/z and sodium formate was used to calibrate the instrument in this mass range.

The collected data was analyzed with the Bruker Compass Data Analysis software and the observed m/z values were compared to the theoretical monoisotopic mass calculated by Bruker Compass IsotopePattern software.

**Kinetics Assays**

**Determination of Background rate**

A colorimetric DCPIP indirect assay was used to determine RhlI activity. Assay mixtures composed of HEPES buffer (100 mM, pH 7.3), SAM (300 μM), butanoyl-ACP
(30 µM), DCPIP (30 µM), and sufficient nanopure water for a total volume of 100 µL were tested, with DCPIP being added the last. Upon the addition of DCPIP, the absorbance of the mixture was observed at 600 nm for 30 minutes. The background rate was calculated by DCPIP reduction ($\varepsilon_{600} = 21,000 \text{ M}^{-1}\text{cm}^{-1}$; see Chapter 1, equations 1-3). The time range at which the background rate was 5% of the lowest enzyme rate observed was used as the incubation time.

**Determination of Kinetic Constants**

Butanoyl- (C4), hexanoyl- (C6), octanoyl- (C8), decanoyl- (C10), and dodecanoyl- (C12) ACPs were tested (compounds 83-87; chapter 1, Table 3). The reaction mixture was composed of RhlI (0.3 µM or 0.9 µM), DCPIP (300 µM), SAM (300 µM), HEPES buffer (100 mM, pH 7.3), varying concentrations of acyl-ACP (2-200 µM), and nanopure water for a total of 100 µL reaction volume. The RhlI concentration was maintained at 0.3 µM when working with C4-ACP native substrate and kept to 0.9 µM when non-native substrates were tested. All components, sans enzyme, were incubated together for 10 minutes prior to initiation by RhlI. The absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to RhlI rate (equations 1-3, chapter 1) and fitted to Michaelis-Menten equation (equation 5, chapter 1) to determine the kinetic constants. Each run was conducted in triplicates and the spread used to determine error with GraphPad Prism 7.

**IC$_{50}$/EC$_{50}$ determination**

Since AHLs are mostly hydrophobic, AHL derivatives were first dissolved in DMSO. The IC$_{50}$ value of DMSO was determined to identify the appropriate volume of
DMSO to use. The reaction mixture was composed of RhlI (0.3 µM), DCPIP (300 µM), SAM (300 µM), C4-ACP (14 µM), HEPES buffer (100 mM, pH 7.3), varying concentrations of DMSO (10% – 25% total volume), and nanopure water for a total of 100 µL reaction volume. All components, sans enzyme, were incubated together for 10 minutes prior to initiation by RhlI. After enzyme addition, the absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to enzyme rate (equations 1-3, chapter 1) to determine the appropriate volume of DMSO to use for RhlI enzyme assay.

The effects of various AHL analogs were tested by measuring their IC$_{50}$ or EC$_{50}$ values via DCPIP colorimetric assay. The reaction mixture was composed of RhlI (0.3 µM), DCPIP (300 µM), SAM (300 µM), C4-ACP (14 µM), HEPES buffer (100 mM, pH 7.3), varying concentrations (0-2 mM) of AHL analogs in DMSO (10% of total reaction volume as determined by above experiment; see Appendix Figure C17), and nanopure water for a total of 100 µL reaction volume. All components, sans enzyme, were incubated together for 10 minutes prior to initiation by RhlI. After enzyme addition, the absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to RhlI rate (equations 1-3, chapter 1) and fitted to IC$_{50}$ or EC$_{50}$ equations depending on the effect (equations 26 and 30, chapter 1). Each run was conducted in triplicates and the spread used to determine error with GraphPad Prism 7.

The effects of IACP$s$ and ICoA$s$ were tested by measuring their IC$_{50}$ values via DCPIP colorimetric assay. The reaction mixture was composed of RhlI (0.3 µM), DCPIP (300 µM), SAM (300 µM), C4-ACP (14 µM), HEPES buffer (100 mM, pH 7.3), varying
concentrations (0 - 1 mM) of IACP or ICoA, and nanopure water for a total of 100 µL reaction volume. All components, sans enzyme, were incubated together for 10 minutes prior to initiation by RhII. After enzyme addition, the absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to RhII rate (equations 1-3, chapter 1) and fitted to IC₅₀ equation (equation 26, chapter 1) with GraphPad Prism 7.

**Time-dependent IC₅₀ test**

The inhibitory effects as a function of time was determined by incubating RhII with varying concentration of an inhibitor for 0, 10, 30, and 60 minutes. The reaction mixture was composed of RhII (0.3 µM), DCPIP (300 µM), SAM (300 µM), C4-ACP (14 µM), HEPES buffer (100 mM, pH 7.3), varying concentrations (10 µM - 2 mM) of the inhibitor. All components, sans RhII and the inhibitor, were incubated together for 10 minutes prior to initiation by RhII-inhibitor mixture. Upon reaction initiation, the absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to RhII rate (equations 1-3, chapter 1) and fitted to IC₅₀ or EC₅₀ equations depending on the effect (equations 26 and 30, chapter 1). Each run was conducted in triplicates and the spread used to determine error with GraphPad Prism 7.

**Mode of Inhibition tests**

The activity of RhII as a function of C4-ACP concentration was determined under varying amounts of an inhibitor. The inhibitor concentration was determined from the IC₅₀ test, choosing two concentrations below, two above and one run with zero inhibitor. Each reaction mixture contained RhII (0.3 µM), DCPIP (300 µM), SAM (300 µM),
HEPES buffer (100 mM, pH 7.3), and varying concentrations of C4-ACP (2-20 μM) and of the inhibitor (0-400 μM). The concentration of C4-ACP was varied while the inhibitor concentration constant. All components, sans enzyme, were incubated together for 10 minutes prior to initiation by RhII. The absorbance was monitored for 300 seconds at 600 nm. The decrease in DCPIP absorbance in the 100 to 200 second range was converted to RhII rate (equations 1-3, chapter 1) and fitted to modified Michaelis-Menten equation using $V_{\text{max}}^{\text{app}}$ and $K_m^{\text{app}}$ to determine the apparent kinetic constants (equation 17, chapter 1). After determining the mode of inhibition, the $V_{\text{max}}^{\text{app}}$ and $K_m^{\text{app}}$ values were used to calculate the inhibitor binding affinity, $K_i$, value (competitive mode of inhibition: equations 18, 19; uncompetitive mode of inhibition: equations 20, 21; and mixed or noncompetitive mode of inhibition: equations 22, 23; see chapter 1). Each data point was collected in single run and fitted to various inhibition models and the best model was determined by comparing each fit using Akaike’s method (AIC) (equations 24 and 25, chapter 1).
CHAPTER THREE: RESULTS AND DISCUSSION

Enzyme Purification

RhlI purification

Four hours of growth was required for the RhlI-containing strains in 1 L medium to reach OD$_{600}$ value between 0.6 and 0.8 for sufficient cell density. Physical lysis via sonication and the subsequent centrifugation resulted in clear dull-yellow lysate. The RhlI-containing plasmid also codes for maltose binding protein (MBP) for use with amylose affinity column. The RhlI and MBP pair has a combined molecular weight of 65.5 kD, which was supported by the analysis of SDS-PAGE gel (Figure 22)

![Figure 22. SDS-PAGE gel of RhlI protein isolated using amylose chromatography. Lane 1: EZ prestained protein ladder; Lane 2: RhlI column load run-through; Lane 3: load wash; Lanes 4-8: Buffer A (50 mM Tris-HCl, pH 7.5, 0.2 M NaCl, 1 mM EDTA, 0.1 mM PMSF, 0.1 mM TLCK, 0.4 M sucrose, and 2.5% (v/v) glycerol) +10 mM maltose elution fractions 1-5. RhlI + MBP has a combined molecular weight of 65.5 kD. The 100 kD ladder is the first one from the top, followed by 30 kD marker. The thick bands in between the 100 and 30 kD marker, much closer to the 100 kD marker, are presumed to be containing purified RhlI.](image-url)
Apo-ACP Purification

The cell culture reached optimal cell density indicated by OD$_{600}$ = 0.6-0.8 within 4 hours of growth. Chemical lysis via B-PER, DNase, RNase, Lysozyme, and PMSF followed by centrifugation was effective in producing clear dull-yellow lysate. Upon addition of MgCl$_2$ and MnSO$_4$ to activate ACP-hydrolase, co-coded with apo-ACP gene to convert holo-ACP to apo-ACP, the solution turned cloudy. A clear lysate was again achieved after precipitating ACP-hydrolase using 2-propanol followed by centrifugation. This solution was run through DE52 diaminoethyl cellulose anion exchange column for purification. SDS-PAGE gel confirmed the isolation of apo-ACP at ~9 kD (Figure 23).

Figure 23. SDS-PAGE gel of apo-ACP isolated with anion exchange chromatography. Lane 1: EZ prestained protein ladder; Lane 2: Crude apo-ACP; Lane 3: load wash; Lanes 4-10: 10 mM lithium MES pH 6.1 + 0.5 M LiCl elution fractions 1, 3, 5, 7, 9, 11. The second-from-the-bottom ladder corresponds to 10 kD. The protein bands aligned with the 10 kD marker are presumed to be apo-ACP (8.6 kD).

Alkyl-CoA Synthesis

In alkyl-CoA synthesis reaction, free-Coenzyme A (limiting reagent) is alkylated with alkyl-bromides. As described in Chapters 1 and 2, DCPIP turns colorless upon reaction with thiols. Therefore, upon adding a sample of the reaction mixture to an aliquot of DCPIP, if no color change is observed, it would indicate that free-CoA, a thiol,
was completely consumed and that the reaction went to completion. The alkyl-CoA (inactive-CoA; ICoA) was isolated via HPLC using the Alkyl-CoA separation method described in Chapter 2 and the fractions were collected using an automated fraction collector. Since the compounds are run through a reverse-phase column, compounds with longer alkyl-chain (more nonpolar) have longer retention time (RT) as compared to ICoAs with shorter, less hydrophobic, alkyl-chains (Figure 24).

![Figure 24. Elution time of various alkyl-CoAs.](image)

**Figure 24. Elution time of various alkyl-CoAs.** The reverse-phase preparatory column has higher affinity for more hydrophobic compounds, causing compounds with longer alkyl chain, thus greater hydrophobicity, to have greater retention time.

**Acyl/Alkyl-ACP Synthesis**

Phosphopantetheinyl transferase, or Sfp, catalyzes the transfer of acyl-/alkyl-pantetheine from CoA nucleotide to ACP protein. Apo-ACP (limiting reagent) was reacted with acyl-/alkyl-CoA in the presence of Sfp. Due to precipitation of highly hydrophobic CoAs with acyl-/alkyl- chain of ten carbons or longer, C10-CoA and C12-CoA were added to the reaction in portions over a 90-minute period to avoid having the CoA crash out of the predominantly aqueous solvent. The synthesis of acyl-/alkyl-ACPs were monitored by HPLC using the ACP separation method as described in Chapter 2. Using the standard ACP separation method, apo-ACP elutes out at 7.8 minutes. The
addition of pantetheine linker greatly reduces its overall hydrophobicity, causing holo- and butanoyl-ACPs to have shorter RT than apo-ACP whereas acyl/alkyl-ACPs with eight or longer carbon chain nonpolar enough to have longer RT than apo-ACP (Figure 25a). Whereas apo-ACP elutes out at 7.8 minutes, butanoyl-, octanoyl-, decanoyl-, and dodecanoyl-ACPs elute out at 7.3, 8.0, 8.3, and 8.7 minutes, respectively. However, hexyl- and hexanoyl-ACPs have nearly identical RT as apo-ACP, necessitating separation using C6-ACP separation method with shallower solvent gradient (Figure 25b). In this method, apo-ACP elution time is 34 minutes, compared with 33 minutes for hexanoyl-ACP. There is virtually no difference in the elution time between an acyl-ACP and its alkyl-ACP counterpart (e.g. both butanoyl-ACP and butyl-ACP elutes out at 7.3 minutes). The reaction was deemed complete when the limiting reagent, apo-ACP, peak at 7.8 min was completely depleted, which occurred in 3 hours or less. However, the batch was deemed unusable if there was a significant peak present at 6.5 minutes, corresponding to holo-ACP contamination. The solution became cloudy when ammonium sulfate was added to precipitate Sfp. The precipitated transferase was then removed with centrifugation. The clear lysate was run through 3 kD spin filtration column to remove ammonium sulfate, CoA byproduct, and excess acyl-/alkyl-CoA. This filtration process was repeated until the peak at 280 nm (corresponding to ACP) was at least 10% greater than the peak at 260 nm (corresponding to CoA) as determined by the UV-vis spectrophotometer.
Figure 25. **Elution time of various acyl-/alkyl-ACPs.** The more hydrophobic compounds have higher affinity to the reverse-phase UHPLC column and have longer retention time. The addition of the pantetheine linker greatly reduces the hydrophobicity of ACP, (a) causing acyl-/alkyl-ACPs with four carbon chains or shorter to elute out before apo-ACP and those with chains of eight carbons or longer to elute out after apo-ACP. (b) Hexanoyl-/hexyl-ACP has nearly identical retention time as apo-ACP thus requiring the solvent gradient to shift over a longer time-period at a lower flowrate.

**Small Molecule Synthesis**

**D-Homocysteine Thiolactone (Figure 19, Chapter 2)**

Stereoisomerically pure D-homocysteine thiolactone was purified from a racemic mixture of DL-homocysteine thiolactone by acylating the thiolactone headgroup with S-(+)-2-methoxyphenylacetic acid in a EDC-coupled reaction to produce two diastereomers with distinct differences in polarity that could be separated with column chromatography. The EDC-coupled reaction produced white crystalline product easily soluble in chloroform but not so much in the 7:3 hexane: ethyl acetate eluting solvent, thus the
diastereomer mixture was dissolved in minimal amount of chloroform and then loaded to the silica gel column. The separation of the diastereomers via silica gel column was confirmed with thin layer chromatography (TLC) with UV active silica with the 7:3 hexane: ethyl acetate eluent. Because there is only a small difference in the polarities of the diastereomers, there were significant number of fractions with both products. The fractions with the both diastereomers were collected and run through the silica gel column again to fully isolate the desired product. In addition to comparing the polarity of the diastereomers via elution time, the purity of the diastereomers was further confirmed by comparing the melting point of each compound; (3′S,2′S)-3-(2′-methoxy-2′-phenyl-acetamido-2-thiophenone (containing L-homocysteine thiolactone) had a melting point of 160-164°C (literature value: 161-163°C) whereas (3′R,2′S)-3-(2′-methoxy-2′-phenyl-acetamido-2-thiophenone (containing D-homocysteine thiolactone) had a melting point of 110-115°C (literature value: 109-111°C). Once the D-homocysteine thiolactone head-group was isolated and recrystallized in 2-propanol, its stereo purity was confirmed by optical rotation measurement $[\alpha]_D^{25}$ -36.3 (0.05 mg/mL in H$_2$O) (literature value: -21.7$^{57}$).

**N-Acyl-D-Homocysteine Thiolactone (Figure 20, Chapter 2)**

The DCC-coupled reaction resulted in colorless mixture with white cloudy urea precipitate, which was easily removed via filtration with celite. Butyric acid was sufficiently polar to yield clean N-Butanoyl-D-homocysteine thiolactone through a series of aqueous washes with water, HCl (2M; 3x10 mL), NaHCO$_3$ (saturated; 3x10 mL), and brine (saturated; 3x10 mL). However, carboxylic acids with tails of six carbons or longer were not polar enough to be removed by aqueous washes alone, requiring purification via
silica gel column chromatography. The polarity between the N-acyl-D-homocysteine thiolactones and their corresponding carboxylic acids were close enough that two runs through the column were required to obtain a pure product.

**N-(3-Oxoacyl)-D-Homocysteine Thiolactone (Figure 21, Chapter 2)**

Meldrum’s acid acylation reaction resulted in a colorless mixture with white cloudy urea precipitate, which was removed via celite filtration, which yielded a pale-yellow flake-like product which was stored at -20 °C to protect the temperature sensitive Meldrum’s acid. Refluxing the acylated Meldrum’s acid with the D-homocysteine thiolactone head-group opened the Meldrum’s acid ring and attached the 3-oxoacyl tail to the headgroup. The polarity of the N-(3-Oxoacyl)-D-homocysteine thiolactones and their corresponding carboxylic acids were close enough that two runs through the silica gel column were required to obtain a pure product.

**Spectral Data**

See Appendix for all Mass Spectrometry (Appendix A), NMR (Appendix B), and UV-vis spectrophotometer (Appendix C) spectra.

**AHL Analog Kinetics**

**Background rate**

As reported previously, non-specific reduction of DCPIP is a major limitation of the DCPIP colorimetric assay, which is compounded by significant levels of contamination in commercially available SAM-Cl. To circumvent the issue, the reaction mixture, except the enzyme, was incubated with DCPIP and the decrease in absorbance at 600 nm was observed. The rate of decrease of absorbance flattens around 600 s (10 min) and the background rate in the 600-900 s range is equivalent of 0.013
µM/min, about 5% of the lowest enzyme rate observed in this project (Figure 26).

Therefore, to minimize background rate interference, the reaction mixture was incubated with DCPIP for 10 min before initiation with the enzyme.

![Absorbance over time graph](image)

**Figure 26.** DCPIP background rate progress curve. The 100 µL reaction mixture was composed of HEPES buffer (100 mM, pH 7.3), SAM (300 µM), butanoyl-ACP (30 µM), DCPIP (30 µM), and water. The absorbance at 600 nm was observed for 1200 s (20 min). The background rate flattens out at 600 s (10 min).

The Effects of AHL analogs on RhlI enzymatic rate

Previous studies have shown that RhlI activity is unaffected by butanoyl-homoserine lactone (C4-HSL; compound 1), its native N-acyl-homoserine lactone (AHL) product. To explore the components of the AHL structure that could be modified to affect RhlI activity, the first set of AHL analogs were designed to test the effects of alterations to the headgroup on RhlI enzymatic rate (Figure 27). The L-homoserine lactone headgroup was modified in the lactone ring (compounds: 2-4, 8-12), the chirality (compounds: 5, 12), and the tail-headgroup linkage (compounds: 6, 7, 10-12). While most derivatives failed to effect RhlI, a change in the chirality and linkage (compounds 5 and 12) in AHL inhibited RhlI. Although analog 8, the thiolactone derivative, did not inhibit RhlI, work from our collaborators have found thiolactone analogs to have
significant antagonistic and agonistic effects on QS receptor proteins; therefore, we were interested in exploring thiolactone analogs in an effort to discover compounds that could inhibit both RhII (AHL synthase) and RhIR (AHL receptor) simultaneously.\textsuperscript{43} The initial study and previous findings prompted us to further expand the analog libraries with alterations to the headgroup stereocenter, sulfonamide linkage between the headgroup and the aliphatic chain, and thiolactone ring headgroup in the search for better RhII inhibitors. (for kinetics data, see Appendix, Figure C1)

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<td>IC\textsubscript{50}: 170.9 ± 98.0 µM</td>
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**Figure 27.** Initial set of AHL-analogs and their effects. The initial rates of RhII with 300 µM of SAM and 14 µM of C4-ACP in the presence of 0-20 mM of the AHL analogs were observed. Only the chiral and linkage changes (compounds 5 and 12) caused inhibition. (Appendix Figure C1)

**Acyl-L-homoserine lactone (L-HSL)**

In many QS receptor studies, modifications to the acyl-chain had significant antagonistic and agonistic effects on the receptor.\textsuperscript{43,59} To determine whether the same is true for RhII, the native headgroup, L-HSL, was acylated with various acyl, 3-oxoacyl, and 3-hydroxyacyl-chains (compounds 1,13-27; Figure 28). Regardless of the
modification to the hydrocarbon chain, none of the sixteen analogs with the L-HSL headgroup moiety inhibited RhlI. (Appendix Figure C2)

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**Figure 28.** Effects of acyl-L-homoserine lactones, 3-oxoacyl- and 3-hydroxyacyl-L-homoserine lactones on RhlI initial rate. The initial rates of RhlI in the presence of 0-2 mM of acyl-L-homoserine lactones of various acyl-chain lengths were observed. Acyl-L-homoserine lactones with chains between 4 and 16 carbons long did not inhibit RhlI. (Appendix Figure C2)

Acyl-D-homoserine lactone (D-HSL)

While none of the eight acyl-L-HSLs (compounds: 1, 13-19) inhibited RhlI, two analogs out of seven acyl-D-HSLs were found to inhibit the enzyme: compounds 5, IC$_{50}$: 687.7 ± 90.7 µM, and compound 31, IC$_{50}$: 20.2 ± 10.1 µM (Figure 29). Interestingly,
while compound 5 has a butanoyl tail, which is the native acyl-chain for the Rhl QS system, compound 31 has a phenylbutanoyl chain, which should be too large to bind to RhlI acyl-chain binding pocket. (Appendix Figure C3)

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<td>31</td>
<td>20.2 ± 10.1 µM</td>
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Figure 29. **Effects of acyl-D-homoserine lactones.** The initial rates of RhlI in the presence of 0-2 mM of acyl-D-homoserine lactones of various acyl-chain lengths were observed. Of these compounds, acyl-D-homoserine lactones with butanoyl and butyl-phenyl chain (compounds 5 and 31) inhibited RhlI, with the longer/bulkier compound 31 having lower IC<sub>50</sub> value (20.2 ± 10.1 µM vs. 687.7 ± 90.7 µM). (Appendix Figure C3)

3-oxoacyl-D-HSL

However, the more interesting pattern emerges with 3-oxoacyl-D-HSL derivatives (Figure 30). While compound 35, medium chain-length analog (3-oxoC8-D-HSL), inhibited RhlI (IC<sub>50</sub>: 282.0 ± 34.1 µM), a shorter chain derivative, compound 34 (3-oxoC6-D-HSL), activated RhlI (EC<sub>50</sub>: 224.4 ± 55 µM). This discovery of an activator, suggests that the small molecule modulators are perhaps binding to an allosteric or nonspecific binding site. The presence of a nonspecific binding site could explain how compound 31, with its large tail, binds with and inhibits RhlI. (Appendix Figure C4)
Figure 30. Effects of 3-oxoacyl-D-homoserine lactones. The initial rates of RhlI in the presence of 0-2 mM of 3-oxoacyl-D-homoserine lactones of various acyl-chain lengths were observed. Of these compounds, acyl-D-homoserine lactones with hexanoyl chain (compound 34) activated RhlI initial rate (EC\textsubscript{50}: 224.4 ± 55 µM) whereas the one with octanoyl chain (compound 35) inhibited RhlI (IC\textsubscript{50}: 282.0 ± 34.1 µM) and the one with dodecanoyl chain did not affect RhlI enzyme rate. (Appendix Figure C4)

Acyl-sulfonamide-DL-HSL.

Unlike the pattern found with acyl-HSLs, RhlI inhibitors were found from both L and D sulfonamide derivatives (Figure 31). Compound 12, a sulfonamide-D-HSL analog, was found to be a more potent inhibitor than compound 37, a sulfonamide-L-HSL analog, with IC\textsubscript{50} values of 170.9 ± 98.0 µM vs. 345.3 ± 79.3 µM, respectively. However, no other variations in the aliphatic chain with sulfonamide linkage was found to inhibit RhlI. (Appendix Figure C5)

Figure 31. Effects of L and D sulfonamides. The initial rates of RhlI in the presence of 0-2 mM of L and D sulfonamide-homoserine lactones of various chain lengths were observed. The compounds with the short chains, butylsulfonamide-L-homoserine lactone (compound 37) and propylsulfonamide-D-homoserine lactone (compound 12), inhibited RhlI with IC\textsubscript{50} values of 345.3 ± 79.3 µM and 170.9 ± 98.0 µM, respectively. (Appendix Figure C5)
Acyl- and 3-oxoacyl-L-homocysteine thiolactone

Although compound 8 did not inhibit RhII, expansion of the hydrocarbon tail library yielded two inhibitors: compounds 46 and 55 with IC₅₀ values of 387.3 ± 88.6 µM and 621.2 ± 49.5 µM, respectively (Figure 32). Again, contrary to initial expectations, it is the long-chain analogs that inhibit RhII while the short-chain derivatives fail to effect RhII activity. Furthermore, this is the first category of compounds with a L-stereocenter headgroup in which multiple inhibitors were discovered. (Appendix Figure C6)

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<td>49</td>
<td>50</td>
</tr>
<tr>
<td>51</td>
<td>52</td>
<td></td>
</tr>
<tr>
<td>53</td>
<td>54</td>
<td>55</td>
</tr>
</tbody>
</table>

**Figure 32. Effects of acyl- and 3-oxoacyl-L-homocysteine thiolactones.** The initial rates of RhII in the presence of 0-2 mM of acyl- and 3-oxoacyl-L-homocysteine thiolactones of various acyl-chain lengths were observed. Of these compounds, only C12-L-homocysteine thiolactone and 3-oxo-C12-L-homocysteine thiolactone (compounds 46 and 55), the ones with the longest straight chain, inhibited RhII. Of these two compounds, one with the acyl-chain had lower IC₅₀ value than the one with the 3-oxoacyl-chain (IC₅₀: 387.3 ± 88.6 µM vs. 621.2 ± 49.5 µM). (Appendix Figure C6)
Acyl- and 3-oxoacyl-D-thiolactone

Following the pattern found with L- and D-HSL derivatives, D-thiolactone analogs produced much more potent inhibitors than L-thiolactones (Figure 33). Compared with IC$_{50}$ value of 387.3 ± 88.6 µM for compound 46 (C12-L-thiolactone), compound 60 (C12 D-thiolactone) had IC$_{50}$ value of 11.4 ± 1.5 µM, a 34-fold decrease. And whereas compound 55 (3-oxoC12-L-thiolactone) had IC$_{50}$ value of 621.2 ± 49.5 µM, compound 64 (3-oxoC12-D-thiolactone) was found to have a much lower IC$_{50}$ value 127.4 ± 70.2 µM, close to 5-fold decrease. However, the most interesting phenomena occurred with short and medium-chain derivatives. As seen with 3-oxoacyl-D-HSLs, which were found to be activating with a shorter chain and inhibiting with a longer chain, 3-oxoC6, 3-oxoC8, and 3-oxoC10-D-thiolactones (compounds 61-63) activated while 3-oxo C12-D-thiolactone (compound 64) inhibited RhlI. Interestingly, as the chain lengthened from 3-oxoC6 to 3-oxoC8 and 3-oxoC10, the activation potency increased indicated by EC$_{50}$ values decreasing 1506 ± 109.9 µM, 854.5 ± 74.4 µM, and 57.1 ± 22.7 µM. (Appendix Figure C7)
Figure 33. Effects of acyl- and 3-oxoacyl-D-homocysteine thiolactones. The initial rates of RhlI in the presence of 0-2 mM of acyl- and 3-oxoacyl-D-homocysteine thiolactones of various acyl-chain lengths were observed. Of these compounds, dodecanoyl-D-homocysteine thiolactone and 3-oxo-dodecanoyl-L-homocysteine thiolactone (compounds 60 and 64), the ones with the longest straight chain, inhibited RhlI. As with their L-counterpart, one with the acyl-chain had lower IC$_{50}$ value than the one with the 3-oxoacyl-chain (IC$_{50}$: 11.4 ± 1.5 µM vs. 127.4 ± 70.2 µM). The shorter-chain 3-oxoacyl-D-homocysteine thiolactones (compounds 61-63) activated RhlI activity, with the EC$_{50}$ values decreasing as the carbon chain lengthened. (Appendix Figure C7)

**Acyl-Cyclopentanamide**

Since thiolactone derivatives inhibited and activated RhlI to a greater degree than lactone analogs, the effects of cyclopentyl headgroup was tested to explore further increase in headgroup hydrophobicity (compounds 4, 65-68; Figure 34). However, no cyclopentyl derivatives effected RhlI activity. (Appendix Figure C8)
Figure 34. Effects of acyl-cyclopentanamide on RhlII activity. The initial rates of RhlII-catalyzed C4-HSL synthesis were similar with and without acyl-cyclopentanamides, suggesting no inhibitory effect of this class of molecules on RhlII activity. (Appendix Figure C8)

Non-lactone derivatives

While cyclopentyl derivatives tested the effects of nonpolar headgroups, various non-lactone analogs were designed to test headgroups of various configurations and hydrophilicity (compounds 2, 3, 9-11, 69-73; Figure 35). None of these non-lactone compounds inhibited RhlII. (Appendix Figure C9)
Figure 35. **Effects of non-lactone AHL analogs on RhlI activity.** The initial rates of RhlI in the presence of 0–2 mM of various non-lactone AHL analogs were observed and none of them inhibited RhlI. (Appendix Figure C9)

**Headgroup vs Tail chain effects**

Thus far, only lactone and thiolactone derivatives modulated RhlI rate (activation and inhibition), and the variation in the acyl-chain enhanced the effect. To check if the inhibitory effects observed for lactone and thiolactone derivatives described above was caused by nonspecific binding of headgroup or fatty acid to the synthase enzyme, the initial rate of RhlI was observed in the presence of the headgroup (L-HSL, D-HSL, L-thiolactone, and D-thiolactone; compounds 74-77), the fatty acid chain tail (butyric, hexanoic, octanoic, decanoic, and dodecanoic acid; compounds 78-82), or both (D-thiolactone + dodecanoic acid, compounds 77 + 82, corresponding to compound 60; Figure 36). None of these kinetic assays resulted in the inhibition of RhlI, suggesting that both the headgroup and the aliphatic tail moieties must be covalently linked to each other to observe the inhibition/activation effects described above. (Appendix Figure C10)
the initial rates of RhlI in the presence of 0-2 mM of L and D homoserine lactone and homocysteine thiolactone headgroups and various carboxylic acid tail groups were observed. Furthermore, the initial rates of RhlI were observed in the presence of 0-2 mM of both compounds 77 and 82, which was analogous to compound 60 which did inhibit RhlI. None of these compounds inhibited RhlI, indicating that neither the acyl-chain nor the headgroup alone has sufficient binding affinity to RhlI to cause inhibition. (Appendix Figure C10)

Determining the mode of inhibition

As described in chapter 1, RhlI is an ordered bi-ter enzyme with C4-ACP binding second and C4-L-HSL released second (Figure 37). AHL analogs are expected to compete for C4-L-HSL binding site. As such, C4-ACP and AHL analogs bind to different RhlI forms (C4-ACP to EA form and analogs to ER form), which would cause intercept-effect in a double reciprocal plot (Chapter 1, p. 21). Furthermore, C4-ACP binding and C4-L-HSL release are separated by a product release step in both the forward and reverse directions; therefore, C4-ACP binding and AHL analog binding are irreversibly connected, which would be manifested by a lack of slope-effect in a double
reciprocal plot (Chapter 1, p. 20). Intercept-effect without slope-effect would produce a set of parallel lines, indicative of uncompetitive inhibition.

**Figure 37. Cleland diagram of RhlI catalyzed reaction.** “E” denotes RhlI while “A,” “B,” “P,” “Q,” and “R” represents SAM, C4-ACP, holo-ACP, C4-L-HSL, and MTA, respectively.

The Lineweaver-Burk plot of initial RhlI rate versus variable C4-ACP concentrations at various fixed C12-D-thiolactone (Figure 38a) or 3-oxoC12-D-thiolactone (Figure 38b) concentrations show a set of parallel lines, indicative of uncompetitive-mode of inhibition which supports the initial expectation that AHL analogs are binding to C4-L-HSL binding site (ER enzyme conformation in Figure 31 above). Further analysis using the Akaike’s method (AIC; see equations 24, 25), confirms that all the inhibition data is best fitted by uncompetitive inhibition model (Table 10). Moreover, as predicted by the trend in IC₅₀ values, C12-D-thiolactone (Kᵢ: 86.2 ± 9.6 µM) has higher binding affinity than 3-oxoC12-D-thiolactone (Kᵢ: 431.6 ± 30.8 µM) (Table 11).
Double Reciprocal Plot of RhlI activity with varying C4-ACP concentrations and various fixed AHL analog concentrations. Double reciprocal of the initial rate of RhlI against C4-ACP concentration in the presence of various fixed concentrations of (a) compound 60 (C12-D-thiolactone) and (b) compound 64 (3-oxoC12-D-thiolactone) was plotted. The fixed AHL analog concentrations were chosen to be 0, below the IC50 value, around the IC50 value, and two above the IC50 value. While keeping the AHL analog concentration fixed, C4-ACP concentration was varied from 2 to 20 µM. The inverse of the initial rate was plotted against inverse of the C4-ACP concentration which revealed a set of parallel lines, indicative of uncompetitive mode of inhibition.

Table 10. Determining best fit model for the mode of inhibition using AIC

<table>
<thead>
<tr>
<th>Analog</th>
<th>Best fit</th>
<th>U vs C</th>
<th>U vs M</th>
<th>U vs N</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>Uncompetitive</td>
<td>&gt;99.99% U</td>
<td>78.81% U</td>
<td>95.84% U</td>
</tr>
<tr>
<td>64</td>
<td>Uncompetitive</td>
<td>&gt;99.99% U</td>
<td>54.69% U</td>
<td>68.30% U</td>
</tr>
</tbody>
</table>

a = Uncompetitive mode of inhibition  
b = Competitive mode of inhibition  
c = Mixed mode of inhibition  
d = Noncompetitive mode of inhibition

Table 11. Effect of AHL analogs on RhlI initial enzyme rate with variable C4-ACP concentration.

<table>
<thead>
<tr>
<th>Analog</th>
<th>IC50 (µM)</th>
<th>Ki (µM)</th>
<th>Mode of Inhibition</th>
</tr>
</thead>
<tbody>
<tr>
<td>60</td>
<td>11.4 ± 1.5</td>
<td>86.2 ± 9.6</td>
<td>Uncompetitive</td>
</tr>
<tr>
<td>64</td>
<td>127.4 ± 70.2</td>
<td>431.6 ± 30.8</td>
<td>Uncompetitive</td>
</tr>
</tbody>
</table>

AHL analog Trends

Chain length effect

RhlI inhibition was primarily observed with D-HSL, L-thiolactone, and D-thiolactone headgroups. With each headgroup, a short-chain derivative resulted in no or
weak inhibition while a long or bulky-chain analog displayed much more potent inhibitory effects (Table 12). Moreover, if the short-chain analog does inhibit, it achieves greater maximum inhibition (maximum % inhibition). This pattern of long-chain derivatives having lower IC50 is highly surprising. The acyl-chain moiety is expected to bind in the small acyl-chain binding pocket of RhII, specific for a butanoyl chain of the C4-ACP native substrate, used by its native substrate/product. In the absence of the crystal structure of RhII, this phenomenon can be explained by several hypotheses: (1) both the k_on and k_off values could be higher for longer chains or (2) the longer acyl-chains are binding to a nonspecific or an alternate acyl-chain binding site with higher affinity than the acyl-chain binding pocket. This acyl-chain length pattern of longer chains better inhibiting RhII can be further studied by analyzing RhII enzymatic rate with nonnative acyl-ACP substrates and the inhibition patterns of inhibitors of various hydrocarbon tail length. Acyl carrier protein (ACP) engulfs the acyl-chain until the proteins binds to the appropriate enzyme active site and then releases the acyl-chain. Therefore, any acyl-chain binding site must be specific and close to ACP binding site. Long-chain acyl-ACP substrates undergo RhII catalysis with decreasing K_m as the acyl-chain length is increased (see “Determining Kinetic Constants with various Acyl-ACPs” section below). Similar pattern (decrease in K_i and IC50 with increase in acyl-chain length) was observed with longer chain alkyl-ACPs and alkyl-CoAs, thus ruling out inhibition due to nonspecific binding for long-chain analogs. Therefore, the acyl-chain must bind at or somewhere close to the acyl-chain binding pocket in RhII.
Table 12. Trends with variations in the acyl-chain length/size.

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt; (µM)</th>
<th>% Inhibition</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt; Trends&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Molecule 5] 5</td>
<td>687.7 ± 90.7</td>
<td>40%</td>
<td>Weaker</td>
</tr>
<tr>
<td>![Molecule 31] 31</td>
<td>20.2 ± 10.1</td>
<td>25%</td>
<td>Stronger</td>
</tr>
<tr>
<td>![Molecule 43] 43</td>
<td>None</td>
<td>0%</td>
<td>No Inhibition</td>
</tr>
<tr>
<td>![Molecule 46] 46</td>
<td>387.3 ± 88.6</td>
<td>60%</td>
<td>Weak Inhibition</td>
</tr>
<tr>
<td>![Molecule 53] 53</td>
<td>None</td>
<td>0%</td>
<td>No Inhibition</td>
</tr>
<tr>
<td>![Molecule 55] 55</td>
<td>621.2 ± 49.5</td>
<td>60%</td>
<td>Weak Inhibition</td>
</tr>
<tr>
<td>![Molecule 56] 56</td>
<td>None</td>
<td>0%</td>
<td>No Inhibition</td>
</tr>
<tr>
<td>![Molecule 60] 60</td>
<td>11.4 ± 1.5</td>
<td>40%</td>
<td>Weak Inhibition</td>
</tr>
</tbody>
</table>

<sup>a</sup> = On a relative scale

Acyl-chain vs 3-oxoacyl-chain effects

A very interesting pattern develops with 3-oxoacyl-chain derivatives (Table 13). Compared with their acyl-chain counterparts, 3-oxoacyl analogs have much less inhibitory characteristics, indicated by their significantly higher IC<sub>50</sub> values (621.2 ± 49.5 µM vs 387.3 ± 88.6 µM for compounds 55 and 46, respectively; and 127.4 ± 70.2 µM vs 11.4 ± 1.5 µM for compounds 64 and 60, respectively). Furthermore, varying the chain
length significantly alters its behavior. As discussed above, shortening the chain seems to decrease the inhibitory characteristics of the analog. The decrease in the inhibitory characteristic due to having a carbonyl at the C3 position combined with the short-chain effect appears to have a synergistic result of activating RhlI activity. While 3-oxoC8-D-HSL (compound 35) inhibited RhlI with IC$_{50}$ value of 282.0 ± 34.1 µM, shortening the 3-oxoacyl-chain to 3-oxoC6 chain (compound 34) caused the derivative to activate RhlI with EC$_{50}$ value of 224.4 ± 55 µM. The same pattern holds true with 3-oxoacyl-D-thiolactone analogs. While 3-oxoC12-D-thiolactone, the long-chain derivative, inhibited RhlI with IC$_{50}$ value of 127.4 ± 70.2 µM, shorter chain analogs activated RhlI. However, until the long-chain effect caused 3-oxoC12-D-thiolactone (compound 64) to be an inhibitor, lengthening the chain from 3-oxoC6-D-thiolactone (compound 61) to 3-oxoC8-D-thiolactone (compound 62) and then to 3-oxoC10-D-thiolactone (compound 63) heightened RhlI activation, shown by decreasing EC$_{50}$ values of 1506 ± 109.9 µM, 854.5 ± 74.4 µM, and 57.1 ± 22.7 µM, respectively. Although the EC$_{50}$ values decrease as the chain length increase, the maximum activation also decrease as the chain lengthens. Activation is usually indicative of a presence of an allosteric site. Moreover, if the 3-oxoacyl-chains bind to and occupy the RhlI acyl-chain binding pocket, it is inconceivable that the acyl-chain from C4-ACP can also bind to the acyl-chain binding pocket, further supporting hypothesis 2.
Table 13. Patterns in varying effects of acyl- and 3-oxoacyl-chain derivatives.

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt;/EC&lt;sub&gt;50&lt;/sub&gt; (µM)</th>
<th>% Inhibition/Activation</th>
<th>IC&lt;sub&gt;50&lt;/sub&gt; and EC&lt;sub&gt;50&lt;/sub&gt; Trends&lt;sup&gt;a&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Compound 34" /></td>
<td>EC&lt;sub&gt;50&lt;/sub&gt;: 224.4 ± 55 100% A</td>
<td></td>
<td>Activation</td>
</tr>
<tr>
<td><img src="image" alt="Compound 35" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 282.0 ± 34.1 60% I</td>
<td></td>
<td>Inhibition</td>
</tr>
<tr>
<td><img src="image" alt="Compound 55" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 621.2 ± 49.5 60% I</td>
<td></td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td><img src="image" alt="Compound 46" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 387.3 ± 88.6 60% I</td>
<td></td>
<td>Stronger Inhibition</td>
</tr>
<tr>
<td><img src="image" alt="Compound 64" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 127.4 ± 70.2 50% I</td>
<td></td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td><img src="image" alt="Compound 60" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 11.4 ± 1.5 30% I</td>
<td></td>
<td>Stronger Inhibition</td>
</tr>
<tr>
<td><img src="image" alt="Compound 61" /></td>
<td>EC&lt;sub&gt;50&lt;/sub&gt;: 1506 ± 109.9 100% A</td>
<td></td>
<td>Weak Activation</td>
</tr>
<tr>
<td><img src="image" alt="Compound 62" /></td>
<td>EC&lt;sub&gt;50&lt;/sub&gt;: 854.5 ± 74.4 60% A</td>
<td></td>
<td>Strong Activation</td>
</tr>
<tr>
<td><img src="image" alt="Compound 63" /></td>
<td>EC&lt;sub&gt;50&lt;/sub&gt;: 57.1 ± 22.7 40% A</td>
<td></td>
<td></td>
</tr>
<tr>
<td><img src="image" alt="Compound 64" /></td>
<td>IC&lt;sub&gt;50&lt;/sub&gt;: 127.4 ± 70.2 50% I</td>
<td></td>
<td>Inhibition</td>
</tr>
</tbody>
</table>

<sup>a</sup> On a relative scale
Headgroup chirality effect

The D-stereocenter derivatives were found to have much greater effect on RhlI enzymatic rate than their L-stereocenter counterparts (Table 14). While the native AHL product, C4-L-HSL, did not inhibit RhlI, change of the headgroup chirality to C4-D-HSL transformed it to a weak inhibitor. Whereas both C12-L-thiolactone (compound 60) and 3-oxoC12-L-thiolactone (compound 64) both inhibited RhlI, the IC\textsubscript{50} value dropped by 34-fold and 5-fold, respectively for their corresponding D-stereocenter counter parts (compounds 46 vs 60 and 55 vs 64).

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC\textsubscript{50} (µM)</th>
<th>% Inhibition</th>
<th>IC\textsubscript{50} Trends\textsuperscript{a}</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>None</td>
<td>0%</td>
<td>No Inhibition</td>
</tr>
<tr>
<td>5</td>
<td>687.7 ± 90.7</td>
<td>40%</td>
<td>Weak Inhibition</td>
</tr>
<tr>
<td>46</td>
<td>387.3 ± 88.6</td>
<td>60%</td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td>60</td>
<td>11.4 ± 1.5</td>
<td>30%</td>
<td>Stronger Inhibition</td>
</tr>
<tr>
<td>55</td>
<td>621.2 ± 49.5</td>
<td>60%</td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td>64</td>
<td>127.4 ± 70.2</td>
<td>50%</td>
<td>Stronger Inhibition</td>
</tr>
</tbody>
</table>

\textsuperscript{a} On a relative scale
Headgroup hydrophilicity effect

Reducing the hydrophilicity of the headgroup by replacing homoserine lactone with homocysteine thiolactone head made a marked increase in the magnitude of the effect on RhlI inhibition and activation (Table 15). While no L-HSL derivatives inhibited RhlI, C12-L-thiolactone and 3-oxo-C12-L-thiolactone were shown to be inhibitors of the AHL synthase. With D-HSL derivatives, $IC_{50}$ values ranged from 687.7 to 20.2 µM (compounds 5 and 31), which was significantly reduced to 127.4 to 11.4 µM (compounds 64 and 60) by replacing D-HSL with a more hydrophobic D-thiolactone headgroup.
Table 15. The effect of headgroup hydrophobicity on RhlI inhibition.

<table>
<thead>
<tr>
<th>Compound</th>
<th>IC50 (µM)</th>
<th>% Inhibition</th>
<th>IC50 Trends^a</th>
</tr>
</thead>
<tbody>
<tr>
<td>All 3-oxoacyl-/3-hydroxyacyl/acyl-L-HSL</td>
<td>621.2 ± 49.5</td>
<td>60%</td>
<td>No Inhibition</td>
</tr>
<tr>
<td>55</td>
<td>387.3 ± 88.6</td>
<td>60%</td>
<td>Stronger Inhibition</td>
</tr>
<tr>
<td>46</td>
<td>687.7 ± 90.7</td>
<td>60%</td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td>5</td>
<td>282.0 ± 34.1</td>
<td>60%</td>
<td>Weaker Inhibition</td>
</tr>
<tr>
<td>35</td>
<td>20.2 ± 10.1</td>
<td>25%</td>
<td></td>
</tr>
<tr>
<td>31</td>
<td>127.4 ± 70.2</td>
<td>60%</td>
<td></td>
</tr>
<tr>
<td>64</td>
<td>11.4 ± 1.5</td>
<td>60%</td>
<td>Stronger Inhibition</td>
</tr>
<tr>
<td>60</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

^a = On a relative scale

Determining Kinetic Constants with various Acyl-ACPs

To determine the effect of the acyl-chain variation on acyl-ACP substrate catalytic efficiencies, RhlI activity was assayed with butanoyl-ACP (compound 83; Chapter 1, Table 6), the native substrate, and then with four long-chain substrates: C6-ACP, C8-ACP, C10-ACP, and C12-ACP (compounds 84-87). The reaction setup was
based on published reaction conditions for previous studies on RhII with its native C4-ACP substrate. The acyl-ACP was the variable substrate while SAM was kept to 300 µM (Figure 39). As expected, the highest initial enzyme rate was achieved with the native substrate, C4-ACP, with a $k_{cat}$ value of $3 \pm 0.3 \text{ min}^{-1}$. This was followed by $k_{cat}$ values of $0.86 \pm 0.03 \text{ min}^{-1}$, $0.64 \pm 0.02 \text{ min}^{-1}$, $0.39 \pm 0.01 \text{ min}^{-1}$, and $0.61 \pm 0.01 \text{ min}^{-1}$ for C6-, C8-, C10-, and C12-ACPs, respectively. The $K_m$ value for the native product was found to be $7 \pm 2 \mu M$. Typically, non-native substrates have higher $K_m$ value than that of the native substrate. However, contrary to expectation, the $K_m$ value decreased as the acyl-chain lengthened: $1.2 \pm 0.2 \mu M$ for C6-ACP, $0.21 \pm 0.06 \mu M$ for C8-ACP, $0.16 \pm 0.03 \mu M$ for C10-ACP, and $0.26 \pm 0.05 \mu M$ for C12-ACP. Although both $K_m$ and $k_{cat}$ values both decreased, $K_m$ dropped more precipitously, causing the catalytic efficiency, determined by $k_{cat}/K_m$, to be much greater for the non-native substrate than that of the native substrate as summarized in Table 16 (Figure 40).
Figure 39. **Substrate-velocity curves of RhlI with native and nonspecific acyl-ACP substrates.** RhlI initial rates as a function of acyl-ACP substrate concentration at fixed 300 µM SAM. (a) varying [C4-ACP] and 0.3 µM RhlI, (b) varying [C6-ACP] and 0.9 µM RhlI, (c) varying [C8-ACP] and 0.9 µM RhlI, (d) varying [C10-ACP] and 0.9 µM RhlI, and (e) varying [C12-ACP] and 0.9 µM RhlI. As the acyl-chain length of acyl-ACP increases, $V_{max}$ is reached at lower acyl substrate concentrations, indicative of decreasing $K_m$ values.
**Figure 40.** Trends in k_{cat}, K_{m}, and k_{cat}/K_{m} values of RhlI with various acyl substrates. As the acyl-chain deviates further from the native substrate, both the (a) k_{cat} and the (b) K_{m} values decrease. However, K_{m} drops much faster than k_{cat}, which leads to (c) the catalytic efficiency, k_{cat}/K_{m} to, rise with longer acyl moiety.

**Table 16.** RhlI initial enzyme rate with various acyl-ACP substrates

<table>
<thead>
<tr>
<th>Acyl-ACP</th>
<th>[RhlI] (µM)</th>
<th>k_{cat} (min^{-1})</th>
<th>K_{m} (µM)</th>
<th>k_{cat}/K_{m} (µM^{-1} min^{-1})</th>
<th>k_{cat}/K_{m} (Relative)^a</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4-ACP</td>
<td>0.3</td>
<td>2.6 ± 0.3</td>
<td>7 ± 2</td>
<td>0.36 ± 0.01</td>
<td>100.0</td>
</tr>
<tr>
<td>C6-ACP</td>
<td>0.9</td>
<td>0.86 ± 0.02</td>
<td>1.2 ± 0.2</td>
<td>0.719 ± 0.003</td>
<td>197.4</td>
</tr>
<tr>
<td>C8-ACP</td>
<td>0.9</td>
<td>0.66 ± 0.02</td>
<td>0.40 ± 0.06</td>
<td>1.655 ± 0.006</td>
<td>454.1</td>
</tr>
<tr>
<td>C10-ACP</td>
<td>0.9</td>
<td>0.378 ± 0.009</td>
<td>0.14 ± 0.04</td>
<td>2.71 ± 0.02</td>
<td>743.2</td>
</tr>
<tr>
<td>C12-ACP</td>
<td>0.9</td>
<td>0.60 ± 0.01</td>
<td>0.26 ± 0.03</td>
<td>2.324 ± 0.006</td>
<td>637.6</td>
</tr>
</tbody>
</table>

\[ a = \left(\frac{k_{cat}}{K_{m}}\right)/0.36 \]

Crystal structures have shown many AHL synthases to have similar structure, all having specific ACP binding site, SAM binding site, and acyl-chain binding pocket.\textsuperscript{60-63} The acyl-chain binding pocket is easily recognizable by its V-cleft shape, which has a definite spatial limitation to only accommodate acyl-chains of specific size. Moreover, previous studies from the Nagarajan laboratory have shown that variations in the acyl-chain from the native substrate significantly reduce the enzymatic rate of AHL synthases.\textsuperscript{22, 60} This pattern of behavior allows the bacteria to discriminate against the synthesis of wrong signals, conserve resources, and increase the signal-to-noise ratio of its own signal molecules. Therefore, the higher catalytic efficiency non-native acyl substrate with RhlI poses a serious conundrum. One possible cause for this phenomenon
could be with the ACP protein. *Pseudomonas aeruginosa* genome codes for three
different acyl carrier proteins: ACP1, ACPP, and ACP3.\(^{51}\) RhlI has been shown to react
with C4-ACP1 and C4-ACPP substrates with comparable catalytic efficiency but with
much greater catalytic efficiency than *E. coli* ACP-using substrate, which, in turn, has
much higher catalytic efficiency than C4-ACP3. However, this work was conducted
using *E. coli* ACP since *P. aeruginosa* ACPS and *E. coli* ACP share similar sequences
and to provide a common reference point to compare various AHL synthases (Figure 41).
Mair Churchill’s work with LasI (another AHL synthase found in *P. aeruginosa*) showed
that when LasI is expressed in *E. coli* with access to *E. coli* ACP only, the amount of
non-native AHL products is significantly higher than when LasI is expressed in *P.
aeruginosa* (Figure 42).\(^64\) Additionally, alkyl-CoA inhibition test (see *Alkyl-CoA
inhibition* section below) also seems to indicate that the ACP moiety plays an important
role in binding with RhlI. Therefore, the pattern of higher catalytic efficiency with
longer-chain acyl-ACP could be an artifact of using a non-native ACP and could be
removed by using *P. aeruginosa* ACP. Nevertheless, regardless of the ACP-effect, that
RhlI can accommodate long acyl-chain is surprising. As noted above, this phenomenon
can be explained if longer acyl-chains have higher \(k_{\text{off}}\) rate (hypothesis 1) or if RhlI has
an allosteric site with equal or higher affinity for acyl-chains of various length than the
acyl-chain binding pocket (hypothesis 2). Under hypothesis 1, the increasing catalytic
efficiencies with longer acyl-chains could be due to higher \(k_{\text{on}}\) rate while the decreasing
\(k_{\text{cat}}\) rate could be caused by higher \(k_{\text{off}}\) value. If hypothesis 2 is correct, it is possible that
the allosteric site promotes binding while the acyl-chain binding pocket promotes
reaction catalysis, analogous to kinetic vs. thermodynamic controls of reaction. The
feasibility of these proposals was examined by analyzing the inhibition patterns of inhibitors of various hydrocarbon tail length.

The effect of hydrocarbon chain length was tested using alkyl-ACPs, also known as inert-ACP or IACP. The removal of the carbonyl group from acyl-ACP converts the
highly reactive thioester bond to relatively nonreactive thioether bond, making alkyl-ACPs inactive analogs of acyl-ACP (Figure 43). The IC$_{50}$ values of butyl-, hexyl-, octyl-, and decyl-ACPs (compounds 88-91; Chapter 1, Table 6) with RhII were determined to be 9.9 ± 4 µM, 0.74 ± 0.4 µM, 0.068 ± 0.02 µM, and 0.102 ± 0.04 µM, respectively (Table 17; Figure 44). Although the IC$_{50}$ values decrease as the alkyl chain lengthens, the maximum inhibition (indicated by %Rate) also decrease from 50% inhibition with C4-IACP to 20% inhibition with all other IACPs (Table 17). This pattern of partial inhibition suggests the presence of less active form of enzyme, indicative of allosteric inhibition.

![Figure 43. Designing inactive acyl-ACP analog.](image)

(a) In acyl-ACP, the carbonyl group is a part of unstable thioester bond and creates a zone of electronegativity suitable for nucleophilic attack. However, (b) the removal of the carbonyl group in alkyl-ACP forms relatively stable thioether bond unfavorable to nucleophilic attack, thus forming inert-ACP (IACP) analog of the acyl substrate.

### Table 17. Effect of IACP on RhII initial enzyme rate.

<table>
<thead>
<tr>
<th>IACP</th>
<th>IC$_{50}$ (µM)</th>
<th>% Inhibition</th>
<th>$K_i$ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4-IACP</td>
<td>9.9 ± 4</td>
<td>50</td>
<td>15.9 ± 0.9</td>
</tr>
<tr>
<td>C6-IACP</td>
<td>0.74 ± 0.4</td>
<td>20</td>
<td>10.6 ± 1.1</td>
</tr>
<tr>
<td>C8-IACP</td>
<td>0.058 ± 0.02</td>
<td>20</td>
<td>6.5 ± 0.7</td>
</tr>
<tr>
<td>C10-IACP</td>
<td>0.102 ± 0.04</td>
<td>20</td>
<td>4.8 ± 0.3</td>
</tr>
</tbody>
</table>
IC₅₀ test of various IACPs. The initial rate of RhlI with 300 µM of SAM and 14 µM of C4-ACP in the presence of varying concentrations of (a) C4-, (b) C6-, (c) C8-, and (d) C10-IACPs were determined. Whereas C4-IACP, the native substrate analog, achieved the maximum inhibition (~50% inhibition compared to ~20% for all others), other IACPs reached minimum activity at much lower inhibitor concentration.

Given that these four IACPs all inhibit RhlI, the mode of inhibition of these IACPs could shed light to how RhlI copes with longer acyl-chains. Competitive inhibition would definitively indicate that the IACPs are competing with butanoyl-ACP for the same binding site and that RhlI acyl-chain binding pocket can accommodate longer chains. On the contrary, uncompetitive mode of inhibition could be a result of the IACP binding after AHL departs from the active site (hypothesis 1) or it could signify that the IACP is not competing for the acyl-chain binding pocket and support the hypothesis of longer chains binding to nonspecific site. Similarly, noncompetitive inhibition could be a result of the inhibitor-free enzyme (EI) complex formed before
E: SAM complex formation (hypothesis 1) or if the inhibitor binding to the hypothetical alternate/allosteric acyl-chain binding site (hypothesis 2). Plotting the double reciprocal (Lineweaver-Burk plot) of initial RhII rate versus variable C4-ACP concentrations at various fixed IACP concentrations show a set of lines intersecting each other at the x-axis, indicative of noncompetitive-mode of inhibition (Figure 45). Further analysis using the Akaike’s method (AIC; see equations 24, 25), confirms that all the inhibition data is best fitted by noncompetitive inhibition model (Table 16). This finding is especially surprising for C4 IACP, since its similarity with the native substrate, C4-ACP, led to the prediction that would competitively inhibit RhII activity. As predicted by the IC50 values, Ki values decrease as the alkyl-chain lengthened (Table 18). If the inhibition is due to alkyl-ACP binding to free RhII before SAM binding with RhII, the drop in the overall inhibition with longer alkyl chains could be due to higher k_{off} rate while the decrease in Ki (Ki = k_{off}/k_{on}) could be a result of long-chain ACPs having even higher k_{on} rate. However, the noncompetitive mode of inhibition and the drop in both Ki and overall inhibition could be due to hydrocarbon chains binding with equal or higher affinity to an alternative acyl-chain binding sites than to the acyl-chain binding pocket. Although the alternative pocket is yet to be identified, inhibition data with IACPs and ICoAs (see “Alkyl-CoA Inhibition” section below) suggest that the acyl-chain binding pocket of the alternative binding site would most likely be close enough or overlap with the native acyl-chain pocket in RhII.
Figure 45. **Double Reciprocal Plot of RhII activity with varying C4-CP concentrations and various fixed IACP concentrations.** Double reciprocal of initial rate of RhII activity vs. C4-ACP concentration in the presence of various fixed concentrations of (a) C4-IACP, (b) C6-IACP, (c) C8-IACP, and (d) C10-IACP was plotted. The fixed IACP concentrations were chosen to be 0, below the IC₅₀ value, around the IC₅₀ value, and two above the IC₅₀ value. While keeping the IACP concentration fixed, C4-ACP concentration was varied from 2 to 20 µM. The inverse of the initial rate was plotted against inverse of the C4-ACP concentration which revealed a set of lines intersecting near the x-axis, indicative of noncompetitive mode of inhibition.

Table 18. **Determining best fit model for the mode of inhibition using AIC**

<table>
<thead>
<tr>
<th>Analog</th>
<th>Mode</th>
<th>N⁰ vs Cᵇ</th>
<th>N vs Uᶜ</th>
<th>N vs Mᵈ</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4-IACP</td>
<td>Noncompetitive</td>
<td>&gt;99.99 N</td>
<td>99.99 N</td>
<td>67.00 N</td>
</tr>
<tr>
<td>C6-IACP</td>
<td>Noncompetitive</td>
<td>99.87 N</td>
<td>91.03 N</td>
<td>74.41 N</td>
</tr>
<tr>
<td>C8-IACP</td>
<td>Noncompetitive</td>
<td>99.87 N</td>
<td>82.48 N</td>
<td>75.70 N</td>
</tr>
<tr>
<td>C10-IACP</td>
<td>Noncompetitive</td>
<td>&gt;99.99 N</td>
<td>96.94 N</td>
<td>75.72 N</td>
</tr>
</tbody>
</table>

a = Noncompetitive mode of inhibition  
b = Competitive mode of inhibition  
c = Uncompetitive mode of inhibition  
d = Mixed mode of inhibition

A pure noncompetitive mode of inhibition could be a result of nonspecific, non-mechanistic enzyme deactivation. To determine whether IACP inhibition is a specific
effect, RhlI was incubated with C4-IACP and C8-IACP for 0 to 60 min and the IC$_{50}$ was taken at various time points. Preincubation of RhlI with IACP did not significantly alter the inhibitory effects of IACPs, indicating that the inhibition was due to specific binding of IACP to RhlI (Figure 46).

**Figure 46.** **Time-dependency of IACP inhibition.** RhlI was preincubated with varying concentrations of (a) C4- and (b) C8-IACPs for 0 to 60 minutes. The AHL synthase reaction was initiated by the addition of RhlI + IACP mixture to reaction mixtures consisting of 300 µM SAM and 14 µM C4-ACP. With both C4- and C8-IACP, the variations in RhlI activity remained under 5% as incubation time changed from 0 to 60 minutes, eliminating the possibility of nonspecific, nonmechanistic inhibition of RhlI by IACPs.

**Alkyl-CoA Inhibition**

To differentiate the contribution of the hydrocarbon tail and the ACP moieties in binding affinity with RhlI, inhibition of enzyme activity with butyl-, hexyl-, and octyl-CoAs (C4-, C6-, and C8-ICoAs; compounds 92-94, respectively; Chapter 1, Table 3) were investigated in greater detail. RhlI initial rate was observed at a fixed C4-ACP concentration and variable ICoA concentrations (Figure 47). The IC$_{50}$ values of the ICoAs were calculated from the data (Table 19). The IC$_{50}$ values of ICoAs are about one hundred times greater than the IC$_{50}$ values of corresponding IACPs: 72.7 ± 13 vs 0.74 ± 0.4 µM for C6-ICoA and C6-IACP, respectively; and 5.1 ± 0.8 vs 0.058 ± 0.02 µM for C8-ICoA and C8-IACP, respectively. However, the sub-10 µM IC$_{50}$ values for ICoAs
and IACPs are significantly lower than most of the IC50’s obtained with AHL analogs and no inhibition achieved with fatty acid tail alone, suggesting that ACP-pantetheine (holo-ACP) moieties contribute significantly to the potency of the binding affinity. (Appendix Figure C15)

Figure 47. IC50 test of various ICoAs. The initial rate of RhlI with 300 µM of SAM and 14 µM of C4-ACP in the presence of varying concentrations of (a) C6- and (b) C8-ICoAs were determined.

<table>
<thead>
<tr>
<th>ICoA</th>
<th>IC50 (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C4-ICoA</td>
<td>None up to 1 mM</td>
</tr>
<tr>
<td>C6-ICoA</td>
<td>72.7 ± 13</td>
</tr>
<tr>
<td>C8-ICoA</td>
<td>5.1 ± 0.8</td>
</tr>
</tbody>
</table>

Conclusion

This project represents the first effort to use AHL-based small molecules as modulators of AHL-synthases.

We hypothesize that acyl-homoserine lactone based inhibitors would be quorum sensing-specific inhibitors binding to both acyl-chain binding pocket and SAM binding site. By varying headgroup polarity and chirality along with acyl-chain size and
substitution, we have identified key characteristics to improve inhibition and activation (Figure 48). A long or bulky acyl-chain with headgroup of increased hydrophobicity and D-stereocenter is required to inhibit RhII. A slightly more hydrophobic headgroup with D-stereocenter acylated with 3-oxoacyl-chain of medium-long length (6-10 carbons long) yielded the best activators of RhII. We also found that the fatty acid tail group and the headgroup cannot independently inhibit RhII activity. Second generation of AHL-based modulators of RhII activity could include bicyclic headgroups with various substitutions to better fine-tune the levels of hydrophobicity and probe yet-to-be-identified characteristics of the headgroup binding sites.

**Figure 48. Moieties of interest for improved AHL-based RhII modulators.**
Inhibitors (a): hydrophobic headgroup with D-stereocenter acylated with long chain; Activators (b): long 3-oxoacyl-chain attached to hydrophobic headgroup with D-stereocenter.

The discovery of AHL-based small molecule inhibitors of RhII is promising. Since antibiotic resistance is often a QS-controlled phenotype, QS inhibitor with limited potency could be used as a combination drug to reduce antibiotic resistance to further the
usability of currently available drugs. Furthermore, if AHL analogs could target both RhlI and RhlR, its inhibition potency could be significantly heightened to make it a viable QS inhibitor. Therefore, despite the failure of the current library of AHL analogs to inhibit RhlI with sub-micromolar IC$_{50}$, AHL-based modulators of quorum sensing merit further research.

However, focusing strictly on AHL synthase inhibition, AHL-analogs were heavily outcompeted. Although targeting acyl-chain and SAM binding sites with AHL analogs allow for QS-specific modulation, fatty-acid, IACP, and ICoA inhibition data suggest that much greater binding energy is associated with the pantetheine and ACP binding sites (Figure 49). To maximize both specificity and potency, the next generation of AHL synthase inhibitors could be designed to target SAM binding site with ACP or pantetheine binding sites.

**Figure 49. Specificity vs. Potency in targeting RhlI.** Acyl-ACP and SAM are commonly used substrates in human enzymes; therefore, (a) by targeting both acyl-chain binding pocket and SAM binding site, AHL-synthase-specific inhibitor could be designed. However, this study has demonstrated that (b) the pantetheine and ACP binding sites need to be targeted for more potent inhibition of RhlI.
We proposed two hypotheses to explain the trends we see with aliphatic-chain length and RhlI activity. As the acyl-chain length increases, the IC$_{50}$, $K_i$, and percent inhibition all decrease for AHL-based RhlI inhibitors while $K_m$ and $k_{\text{cat}}$ decreases for corresponding acyl-ACP substrates. The decrease in $K_m$ was much steeper than the decrease in $k_{\text{cat}}$ for longer-chain acyl-ACP substrates, leading to an increase in catalytic efficiencies for longer-chain non-native substrates relative to C4-ACP. In addition, as the alkyl-chain length increases, inhibition increase as observed by significant decreases in IC$_{50}$ and $K_i$ values for the longer-chain analogs; however, the percent inhibition also decreased. This acyl-chain pattern could be due to higher $k_{\text{off}}$ rate and even higher $k_{\text{on}}$ rate for longer-chain compounds binding to RhlI. Higher $k_{\text{off}}$ rate would cause less potent effect (lower overall inhibition) while higher $k_{\text{on}}$ rate would cause the $K_i$ value to decrease, indicative of higher binding affinity. However, this trend could be caused by the presence of an alternative acyl-chain binding site (or allosteric site). That all the inhibitors discovered in this study were partial inhibitors suggests allosteric inhibition. This proposal of a possible alternate acyl-chain binding site is further supported by the presence of activators. If the 3-oxoacyl-chains of the activators were to bind with the acyl-chain binding pocket, as indicated by hypothesis 1, it would inhibit C4-ACP binding; however, under hypothesis 2, the 3-oxoacyl-chains would bind with the alternate site. The binding could cause changes to RhlI tertiary structure and form a structural configuration more favorable for catalysis to occur. Furthermore, hypothesis 2 raises another question: how could long-chain derivates binding to one allosteric site cause both activating and inhibiting behaviors? We hypothesize that these effects could be due to either a) the carbonyl at the C3 position alters the mode of binding for activators...
compared with inhibitors; or (b) there could be more than one allosteric site, one for inhibition and another for activation. Co-crystal structures of activators and inhibitors complexed with RhII should provide light on some of these unanswered questions. Although we have not found a submicromolar inhibitor for AHL synthase in this thesis yet, our research efforts on AHL derivatives as potential quorum sensing modulators should open new doors to develop quorum sensing specific inhibitors for pathogenic bacteria. In conclusion, the combination of scarcity of antibacterials in the drug pipeline, increasing resistance to antibiotics, and favorable pharmaceutical qualities of AHL derivatives as novel antivirulent molecules merit further research in this area.
REFERENCES


48. Welsh, M. A.; Eibergen, N. R.; Moore, J. D.; Blackwell, H. E., Small molecule disruption of quorum sensing cross-regulation in pseudomonas aeruginosa causes


54. Equation: [Inhibitor] vs. response -- Variable slope.  

55. Equation: [Agonist] vs. response -- Variable slope.  


Mass Spectra

Figure A1. Mass Spectrum of Compound 56
\[ \text{C}_8\text{H}_{13}\text{NO}_2\text{S} \]
Expected m/z [M + H$^+$]: 188.0734, observed: 188.0763; relative mass error: 15.28 ppm;
Expected m/z [M + Na$^+$]: 210.0554, observed: 210.0589; relative mass error: 16.81 ppm

Figure A2. Mass Spectrum of Compound 57
\[ \text{C}_{10}\text{H}_{17}\text{NO}_2\text{S} \]
Expected m/z [M + H$^+$]: 216.1047, observed: 216.1099; relative mass error: 23.94 ppm;
Expected m/z [M + Na$^+$]: 238.0867, observed: 238.0942; relative mass error: 31.63 ppm
Figure A3. Mass spectrum of Compound 58

\[ \text{C}_{12}\text{H}_{21}\text{NO}_2\text{S} \]

Expected m/z [M + H\(^+\)]: 244.1360, observed: 244.1405; relative mass error: 18.32 ppm;
Expected m/z [M + Na\(^+\)]: 266.1180, observed: 266.1232; relative mass error: 19.66 ppm;
Expected m/z [M + K\(^+\)]: 282.0919, observed: 282.0962; relative mass error: 15.21 ppm

Figure A4. Mass Spectrum of Compound 59

\[ \text{C}_{14}\text{H}_{25}\text{NO}_2\text{S} \]

Expected m/z [M + H\(^+\)]: 272.1673, observed: 272.1720; relative mass error: 17.17 ppm;
Expected m/z [M + Na\(^+\)]: 294.1493, observed: 294.1555; relative mass error: 21.18 ppm;
Expected m/z [M + K\(^+\)]: 310.1291, observed: 310.1291; relative mass error: 19.00 ppm
Figure A5. Mass Spectrum of Compound 60

C₁₀H₁₅NO₃S

Expected m/z [M + H⁺]: 230.0845, observed: 230.0896; relative mass error: 21.99 ppm;
Expected m/z [M + Na⁺]: 252.0665, observed: 252.0729; relative mass error: 25.46 ppm;
Expected m/z [M + K⁺]: 268.0404, observed: 268.0452; relative mass error: 17.83 ppm
Figure A7. Mass Spectrum of Compound 62

$C_{12}H_{19}NO_3S$

Expected m/z [M + H$^+$]: 258.1158, observed: 258.1162; relative mass error: 1.39 ppm;
Expected m/z [M + Na$^+$]: 280.0978, observed: 280.1008; relative mass error: 10.77 ppm;
Expected m/z [M + K$^+$]: 296.0717, observed: 296.0723; relative mass error: 1.95 ppm

Figure A8. Mass Spectrum of Compound 63

$C_{14}H_{23}NO_3S$

Expected m/z [M + H$^+$]: 286.1471, observed: 286.1472; relative mass error: 0.21 ppm;
Expected m/z [M + Na$^+$]: 308.1291, observed: 308.1320; relative mass error: 9.46 ppm;
Expected m/z [M + K$^+$]: 324.1030, observed: 324.1035; relative mass error: 1.47 ppm
Figure A9. Mass Spectrum of Compound 64

C₁₆H₂₇NO₃S

Expected m/z [M + H⁺]: 314.1784, observed: 314.1791; relative mass error: 2.10 ppm;
Expected m/z [M + Na⁺]: 336.1604, observed: 336.1633; relative mass error: 8.68 ppm;
Expected m/z [M + K⁺]: 352.1343, observed: 352.1354; relative mass error: 3.06 ppm

Figure A10. Mass Spectrum of Compound 88

Expected m/z [M + 5H⁺]: 1781.9, observed: 1781.7;
Expected m/z [M + 6H⁺]: 1485.1, observed: 1484.9;
Expected m/z [M + 7H⁺]: 1273.1, observed: 1273.1;
Expected m/z [M + 8H⁺]: 1114.1, observed: 1113.9;
Expected m/z [M + 9H⁺]: 990.4, observed: 990.3;
Expected m/z [M + 10H⁺]: 891.5, observed: 891.4
Figure A11. Mass Spectrum of Compound 89
Expected m/z [M + 5H⁺]: 1787.5, observed: 1787.5;
Expected m/z [M + 6H⁺]: 1489.8, observed: 1489.6;
Expected m/z [M + 7H⁺]: 1277.1, observed: 1277.1;
Expected m/z [M + 8H⁺]: 1117.6, observed: 1117.6;
Expected m/z [M + 9H⁺]: 993.5, observed: 993.5;
Expected m/z [M + 10H⁺]: 894.3, observed: 894.2

Figure A12. Mass Spectrum of Compound 90
Expected m/z [M + 5H⁺]: 1793.1, observed: 1792.9;
Expected m/z [M + 6H⁺]: 1494.5, observed: 1494.2;
Expected m/z [M + 7H⁺]: 1281.1, observed: 1281.1;
Expected m/z [M + 8H⁺]: 1121.1, observed: 1120.9;
Expected m/z [M + 9H⁺]: 996.6, observed: 996.5;
Expected m/z [M + 10H⁺]: 897.1, observed: 897.0
Figure A13. Mass Spectrum of Compound 91

Expected m/z [M + 5H⁺]: 1798.7, observed: 1798.5;
Expected m/z [M + 6H⁺]: 1499.1, observed: 1499.1;
Expected m/z [M + 7H⁺]: 1285.1, observed: 1285.1;
Expected m/z [M + 8H⁺]: 1124.6, observed: 1124.6;
Expected m/z [M + 9H⁺]: 999.8, observed: 999.7;
Expected m/z [M + 10H⁺]: 899.9, observed: 899.8

Figure A14. Mass Spectrum of Compound 92

\[ \text{C}_{25}\text{H}_{44}\text{N}_7\text{O}_{16}\text{P}_3\text{S} \]

Expected m/z [M + H⁺]: 824.1851, observed: 824.1835; relative mass error: -1.92 ppm;
Expected m/z [M + Na⁺]: 846.1670, observed: 846.1657; relative mass error: -1.57 ppm;
Expected m/z [M + 2Na⁺ - H⁺]: 868.1490, observed: 868.1472; relative mass error: -2.04 ppm
Figure A15. Mass Spectrum of Compound 93

C_{27}H_{48}N_{16}O_{16}P_{3}S

Expected m/z [M + H\(^{+}\)]: 852.2164, observed: 852.2147; relative mass error: -1.98 ppm;
Expected m/z [M + Na\(^{+}\)]: 874.1983, observed: 874.1953; relative mass error: -3.47 ppm;
Expected m/z [M + 2H\(^{+}\)]: 426.6118, observed: 426.6105; relative mass error: -3.12 ppm

Figure A16. Mass Spectrum of Compound 94

C_{20}H_{52}N_{16}O_{16}P_{3}S

Expected m/z [M + H\(^{+}\)]: 880.2477, observed: 880.2444; relative mass error: -3.73 ppm;
Expected m/z [M + K\(^{+}\)]: 918.2036, observed: 918.1978; relative mass error: -6.28 ppm;
Expected m/z [M + 2H\(^{+}\)]: 440.6275, observed: 440.6259; relative mass error: -3.59 ppm
Figure A17. Mass Spectrum of Compound 95

C_{31}H_{56}N_{7}O_{16}P_{3}S

Expected m/z [M + H⁺]: 908.2790, observed: 908.2733; relative mass error: -6.26 ppm;
Expected m/z [M + 2H⁺]: 454.6431, observed: 454.6401; relative mass error: -6.67 ppm;
Expected m/z [M + H⁺ + Na⁺]: 465.6345, observed: 465.6300; relative mass error: -9.62 ppm
APPENDIX B
NMR spectra

Figure B1. D-homocysteine thiolactone $^1$H NMR

Figure B2. Compound 56 $^1$H NMR
Figure B3. Compound 56 COSY NMR

Figure B4. Compound 56 HSQC NMR
Figure B5. Compound 56 HMBC NMR

Figure B6. Compound 56 $^{13}$C NMR
Figure B7. Compound 57 $^1$H NMR

Figure B8. Compound 57 COZY NMR
Figure B9. Compound 57 $^{13}$C NMR

Figure B10. Compound 58 $^1$H NMR
Figure B11. Compound 58 COSY NMR

Figure B12. Compound 58 HSQC NMR
Figure B13.  Compound 58 HMBC NMR

Figure B14.  Compound 58 $^{13}$C NMR
Figure B15. Compound 59 $^1$H NMR

Figure B16. Compound 59 COSY NMR
Figure B17. Compound 59 HSQC NMR

Figure B18. Compound 59 HMBC NMR
Figure B19. Compound 59 $^{13}$C NMR

Figure B20. Compound 60 $^1$H NMR
Figure B21. Compound 60 COSY NMR

Figure B22. Compound 60 HSQC NMR
Figure B23.  Compound 60 HMBC NMR

Figure B24.  Compound 60 $^{13}$C NMR
Figure B25. Compound 61 $^1$H NMR

Figure B26. Compound 61 COSY NMR
Figure B27. Compound 61 HSQC NMR

Figure B28. Compound 61 HMBC NMR

Figure B29. Compound 61 $^{13}$C NMR
Figure B30. Compound 62 $^1$H NMR

Figure B31. Compound 62 COSY NMR
Figure B32. Compound 62 HSQC NMR

Figure B33. Compound 62 HMBC NMR

Figure B34. Compound 62 $^{13}$C NMR
Figure B35. Compound 63 $^1$H NMR

Figure B36. Compound 63 COSY NMR
Figure B37. Compound 63 HSQC NMR

Figure B38. Compound 63 HMBC NMR

Figure B39. Compound 64 $^{13}$C NMR
Figure B40.  Compound 64 $^1$H NMR

Figure B41.  Compound 64 COSY NMR
Figure B42. Compound 64 HSQC NMR

Figure B43. Compound 64 HMBC NMR

Figure B44. Compound 64 $^{13}$C NMR
APPENDIX C
Figure C1. IC₅₀ of First generation of AHL analogs.
(Corresponds to Figure 27, Chapter 3)
Figure C2. IC50 of L-HSL derivatives.
(Corresponds to Figure 28, Chapter 3)
Figure C3.  **IC$_{50}$ of acyl-D-HSL analogs.**  
(Corresponds to Figure 29, Chapter 3)

Figure C4.  **IC$_{50}$ and EC$_{50}$ of 3-oxoacyl-D-HSL analogs.**  
(Corresponds to Figure 30, Chapter 3)
Figure C5. IC$_{50}$ of DL-sulfonamide analog.
(Corresponds to Figure 31, Chapter 3)
Figure C6. IC₅₀ of acyl- and 3-oxoacyl-L-homocysteine thiolactones. (Corresponds to Figure 32, Chapter 3)
Figure C7. IC\textsubscript{50} and EC\textsubscript{50} of acyl- and 3\textit{oxo}acyl-D-homocysteine thiolactones.
(Corresponds to Figure 33, Chapter 3)

Figure C8. IC\textsubscript{50} of cyclopentyl derivatives.
(Corresponds to Figure 34, Chapter 3)
Figure C9. IC$_{50}$ of non-lactone AHL analogs.
(Corresponds to Figure 35, Chapter 3)
Figure C10. IC$_{50}$ of headgroup and tail moieties.
(Corresponds to Figure 36, Chapter 3)
Figure C11. Substrate-velocity curve of RhlI with various acyl-ACP substrates.

Figure C12. IC_{50} of various alkyl-ACPs.
(a) C4-IACP, (b) C6-IACP, (c) C8-IACP, (d) C10-IACP
Figure C13. RhII Inhibition assays with various alkyl-ACPs.
(a) C4-IACP, (b) C6-IACP, (c) C8-IACP, (d) C10-IACP

Figure C14. Double reciprocal plot for RhII inhibition with various alkyl-ACPs.
(a) C4-IACP, (b) C6-IACP, (c) C8-IACP, (d) C10-IACP
Figure C15. IC50 of IACPs at different time points
C4-IACP and (b) C8-IACP

Figure C16. IC50 of ICoA derivatives.

Figure C17. DMSO Inhibition
RhII activity was not inhibited up to 10% DMSO (by volume)