SYNTHETIC AGENTS FOR THE DERIVATIZATION OF N-ACYL HOMOSERINE LACTONES

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A dissertation submitted to the faculty of the University of North Carolina at Chapel Hill in partial fulfillment of the requirements for the degree of Doctor of Philosophy in the Department of Chemistry

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ABSTRACT

Michael A. Bertucci: Synthetic Agents for the Derivatization of N-Acyl Homoserine Lactones
(Under the direction of Michel R. Gagné, PhD)

Quorum sensing is a process of chemical communication in bacteria that enables bacteria to coordinate group function, including unified gene expression. As the quorum sensing mechanism is in part responsible for the virulent activity of many bacteria, inhibiting the communication pathway has the potential to lead to the development of novel antibiotics. The central focus of this dissertation is the investigation of strategies for chemically activating and modifying N-acyl homoserine lactones (AHLs), the agent of chemical communication in gram-negative bacteria. Probing the inherent reactivity of the AHLs is critical to understanding their evolved function while uncovering novel modes of inhibition and tools for chemical biology.

Initial studies began by utilizing small molecule thiourea and guanidine-based catalysts for accelerating ring-opening reactions of the AHL by amines. The supramolecular catalysts achieved ten and sixty-five fold increases in reaction rate, respectively, for the addition of piperidine to the characteristic α-amino-γ-butyrolactone moiety. This demonstrated the susceptibility of the chemical messengers to be activated for nucleophilic attack through appropriate arrangement of hydrogen bond donors and acceptors.

Further studies on the derivatization of the AHLs continued with the development of a hydrazine-mediated transamidation reaction under aqueous conditions. Experimental evidence supports the progression of the reaction via a condensation at the β-carbon of the
chemical messenger, followed by an intramolecular cyclization to cleave the AHL at the amide bond. The reaction is operable at physiological pH and is specific for the 3-oxo-AHL in the presence of other AHLs.

To rapidly identify a scaffold of non-covalent interactions to increase rate of the transamidation reaction, a reactive tagging assay was developed for high-throughput screening of hydrazine-containing peptide libraries. Combinatorial libraries were synthesized with designer Boc-protected hydrazines and screened against the AHL. Colorimetric hit development was achieved through a strained-cyclooctyne functionalized dye selecting peptides that react most readily with the chemical messenger. First generation library hits were resynthesized and screened for biofilm and quorum sensing inhibition. Further library development and screening holds the potential to identify peptides sequences and structures with the greatest propensity for rapidly functionalizing the 3-oxo-AHL.
ACKNOWLEDGEMENTS

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To the members of the UNC Men’s Club Volleyball team and the GPSF, we have certainly made lifelong friendships, and I am so appreciative for having the opportunities to compete and advocate with you while at Chapel Hill.

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LIST OF ABBREVIATIONS

% v/v  Percent volume per volume
°C  Degrees Celsius
2-Nal  2-naphthylalanine
3,5-diFPh  3,5-difluorophenylalanine
3-NO₂Phe  3-nitrophenylalanine
A  Alanine
Ac₂O  Acetic anhydride
AHL  N-acyl homoserine lactone
AIB  α-aminoisobutyric acid
Ala  Alanine
Arg  Arginine
Asn  Asparagine
Boc  t-Butyloxycarbonyl
BODIPY  Boron-dipyrrromethene
br  Broad
C  Cysteine
Cha  Cyclohexylalanine
Cit  Citrulline
COMU  (1-Cyano-2-ethoxy-2-oxoethylidenaminooxy)dimethylamino-morpholino-carbenium hexafluorophosphate
d  Doublet
DAB  2,4 diaminobutyric acid
DAP  2,3-diaminopropionic acid
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<th>Full Name</th>
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<td>DBU</td>
<td>1,8-Diazabicyclo[5.4.0]undec-7-ene</td>
</tr>
<tr>
<td>DCM</td>
<td>Dichloromethane</td>
</tr>
<tr>
<td>dd</td>
<td>Doublet of doublets</td>
</tr>
<tr>
<td>DFT</td>
<td>Density functional theory</td>
</tr>
<tr>
<td>DIEA</td>
<td>N,N-Diisopropylethylamine</td>
</tr>
<tr>
<td>DMF</td>
<td>N,N-Dimethylformamide</td>
</tr>
<tr>
<td>DMF-d&lt;sub&gt;7&lt;/sub&gt;</td>
<td>Deuterated N,N-dimethylformamide</td>
</tr>
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<td>DMSO</td>
<td>Dimethylsulfoxide</td>
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<td>DMSO-d&lt;sub&gt;6&lt;/sub&gt;</td>
<td>Deuterated dimethylsulfoxide</td>
</tr>
<tr>
<td>DNA</td>
<td>Deoxyribonucleic acid</td>
</tr>
<tr>
<td>dP</td>
<td>d-Proline</td>
</tr>
<tr>
<td>dPro</td>
<td>d-Proline</td>
</tr>
<tr>
<td>dt</td>
<td>Doublet of triplets</td>
</tr>
<tr>
<td>E</td>
<td>Glutamic acid</td>
</tr>
<tr>
<td>ee</td>
<td>Enantiomeric excess</td>
</tr>
<tr>
<td>eq</td>
<td>Equivalent</td>
</tr>
<tr>
<td>ESI</td>
<td>Electrospray ionization</td>
</tr>
<tr>
<td>Et&lt;sub&gt;2&lt;/sub&gt;O</td>
<td>Diethyl ether</td>
</tr>
<tr>
<td>EtOAc</td>
<td>Ethyl acetate</td>
</tr>
<tr>
<td>F</td>
<td>Phenylalanine</td>
</tr>
<tr>
<td>Fmoc</td>
<td>9-Fluorenlymethyloxycarbonyl</td>
</tr>
<tr>
<td>G</td>
<td>Glycine</td>
</tr>
<tr>
<td>g</td>
<td>Grams</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Full Form</td>
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<tr>
<td>--------------</td>
<td>-----------</td>
</tr>
<tr>
<td>GFP</td>
<td>Green fluorescent protein</td>
</tr>
<tr>
<td>Glu</td>
<td>Glutamic acid</td>
</tr>
<tr>
<td>Gly</td>
<td>Glycine</td>
</tr>
<tr>
<td>h</td>
<td>Hours</td>
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<tr>
<td>HBTU</td>
<td>O-(Benzotriazol-1-yl)-N,N,N',N'-tetramethyluronium hexafluorophosphate</td>
</tr>
<tr>
<td>His</td>
<td>Histidine</td>
</tr>
<tr>
<td>HOBt</td>
<td>Hydroxybenzotriazole</td>
</tr>
<tr>
<td>HPLC</td>
<td>High performance liquid chromatography</td>
</tr>
<tr>
<td>Ile</td>
<td>Isoleucine</td>
</tr>
<tr>
<td>ivDde</td>
<td>1-(4,4-Dimethyl-2,6-dioxocyclohexylidene)-3-methylbutyl</td>
</tr>
<tr>
<td>J</td>
<td>Coupling constant</td>
</tr>
<tr>
<td>K</td>
<td>Lysine</td>
</tr>
<tr>
<td>k</td>
<td>Rate constant</td>
</tr>
<tr>
<td>(k_{\text{app}})</td>
<td>Apparent rate constant</td>
</tr>
<tr>
<td>(k_{\text{rel}})</td>
<td>Relative rate</td>
</tr>
<tr>
<td>L</td>
<td>Leucine</td>
</tr>
<tr>
<td>LB</td>
<td>Luria-Bertani</td>
</tr>
<tr>
<td>LC-MS</td>
<td>Liquid chromatography-mass spectrometry</td>
</tr>
<tr>
<td>LC-TOF</td>
<td>Liquid chromatography time of flight</td>
</tr>
<tr>
<td>Leu</td>
<td>Leucine</td>
</tr>
<tr>
<td>LG</td>
<td>Leaving group</td>
</tr>
<tr>
<td>LUMO</td>
<td>Lowest unoccupied molecular orbital</td>
</tr>
<tr>
<td>Lys</td>
<td>Lysine</td>
</tr>
<tr>
<td>Symbol</td>
<td>Definition</td>
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<td>--------</td>
<td>------------------------------------</td>
</tr>
<tr>
<td>M</td>
<td>Molar</td>
</tr>
<tr>
<td>m</td>
<td>Multiplet</td>
</tr>
<tr>
<td>m/z</td>
<td>Mass to charge ratio</td>
</tr>
<tr>
<td>MALDI-MS</td>
<td>Matrix-assisted laser desorption/ionization mass spectrometry</td>
</tr>
<tr>
<td>MeCN</td>
<td>Acetonitrile</td>
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<tr>
<td>MeOH</td>
<td>Methanol</td>
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<tr>
<td>MS</td>
<td>Mass spectrometry</td>
</tr>
<tr>
<td>NMR</td>
<td>Nuclear magnetic resonance</td>
</tr>
<tr>
<td>O</td>
<td>Ornithine</td>
</tr>
<tr>
<td>OBOC</td>
<td>One-bead-one-compound</td>
</tr>
<tr>
<td>OD</td>
<td>Optical density</td>
</tr>
<tr>
<td>Orn</td>
<td>Ornithine</td>
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<tr>
<td>p</td>
<td>Pentet</td>
</tr>
<tr>
<td>P</td>
<td>Proline</td>
</tr>
<tr>
<td>PBS</td>
<td>Phosphate buffer solution</td>
</tr>
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<td>Abbreviation</td>
<td>Definition</td>
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<td>--------------</td>
<td>------------</td>
</tr>
<tr>
<td>Phe</td>
<td>Phenylalanine</td>
</tr>
<tr>
<td>Phg</td>
<td>Phenylglycine</td>
</tr>
<tr>
<td>ppm</td>
<td>Parts per million</td>
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<tr>
<td>Pro</td>
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</tr>
<tr>
<td>q</td>
<td>Quartet</td>
</tr>
<tr>
<td>QS</td>
<td>Quorum sensing</td>
</tr>
<tr>
<td>RB</td>
<td>Round bottom</td>
</tr>
<tr>
<td>ROH</td>
<td>An alcohol</td>
</tr>
<tr>
<td>RT</td>
<td>Room temperature</td>
</tr>
<tr>
<td>s</td>
<td>Seconds or singlet (when related to NMR)</td>
</tr>
<tr>
<td>Ser</td>
<td>Serine</td>
</tr>
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<td>SPPS</td>
<td>Solid phase peptide synthesis</td>
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<td>T</td>
<td>Threonine</td>
</tr>
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<td>t</td>
<td>Time or triplet (when related to NMR)</td>
</tr>
<tr>
<td>t_{50%}</td>
<td>Time to 50% conversion</td>
</tr>
<tr>
<td>TFA</td>
<td>Trifluoroacetic acid</td>
</tr>
<tr>
<td>TFE-d_3</td>
<td>Deuterated trifluoroacetic acid</td>
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<tr>
<td>THF</td>
<td>Tetrahydrofuran</td>
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<td>1,2,3,4-tetrahydro-6H-pyrimido[2,1-b]quinazolin-6-one</td>
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<td>TLC</td>
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<td>TS</td>
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<td>-------------------------------</td>
</tr>
<tr>
<td>Tyr</td>
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<td>U</td>
<td>α-aminoisobutyric acid</td>
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<td>UV</td>
<td>Ultraviolet</td>
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<td>Volts or valine (when related to amino acids)</td>
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<td>Val</td>
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<td>VEGF</td>
<td>Vascular endothelial growth factor</td>
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<td>W</td>
<td>Tryptophan</td>
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CHAPTER I: INTRODUCTION

A. Bacterial Quorum Sensing: Background and Significance

It was in the 1970s, while studying the bioluminescence of marine microbe *V. fischeri*, that Nealson and Hastings described an “adaptive function” of the bacteria that enabled it to regulate its luminescence at the colony level. In other words, an observed small colony of *V. fischeri* would not illuminate on a cell by cell basis, but rather simultaneously as a unit.¹ These observations by Nealson and Hastings laid the framework for understand a regulatory mechanism present in a wide-range of bacterial species that allows the organisms to participate in a process of unified gene expression. This process in now known as quorum sensing (QS).²,³

![Figure 1.1. Density-dependent regulation of gene expression in bacterial quorum sensing. The bacteria (green ovals) produce small molecule autoinducers (yellow triangles) to promote cell localization until the necessary quorate is reached to initiate gene transcription.](image)

The ability of bacteria to undergo unified gene expression through quorum sensing is dependent on the relative density of bacteria in a given environment. The bacteria must
reach a “quorum” or threshold concentration of cells in order to support the phenotype the colony seeks to express given its surroundings (Figure 1.1). Thus, the bacteria have evolved to develop a chemical communication mechanism to ensure that the necessary quorum can be met. The chemical signals, or autoinducers, that serve as this means of communication are continuously synthesized and released into the extracellular environment to be incorporated and recognized by a nearby bacterium of the same species. The increased dispersal of autoinducers enables more bacteria to “sense” the location of the forming consortia, cueing the individual cells to colonize. Upon binding to its cognate receptor, the signal-receptor complex initiates transcription of the gene encoded for the synthase of the signaling molecule. Through this autoinduction mechanism, the cell is directed to produce more of the same chemical signal to be released into the extracellular space. The bacteria gather until a threshold concentration is reached, enabling successful execution of a colony-wide event.

The chemical signal, mechanism of signal transduction, and resultant genes expressed via quorum sensing vary from species to species. Likewise, the density-dependent mechanism has been determined as the driving force for a wide variety of bacterial processes including bioluminescence, motility, conjugation, and virulence, each with the potential to benefit or harm its host.\(^4\) As noted above, the bioluminescence of \(V.fischeri\) illuminates the surrounding area for its host, the Hawaiian bobtailed squid, enabling it to traverse its marine landscape. In contrast, quorum sensing is the mechanism for the production of virulence factors in prevalent human pathogens such as \(P.aeruginosa\) and \(S.aureus\). The ability of bacteria to regulate group function by these means has been demonstrated to have major impacts in agriculture, manufacturing, and human health.\(^5\) Specifically, quorum sensing has been demonstrated to serve as a precursor to biofilm formation, an extracellular polymeric
substance secreted by bacteria that enhances their resistance to bactericidal treatments. Creation of this protective barrier of DNA, polysaccharide, and protein fragments is the source of pathogenicity in over 80% of bacterial infections.6,7 The study of bacterial quorum sensing has consequently expanded to not only understanding the molecular underpinnings of its regulation, but also to developing strategies to inhibit the pathway to prevent its virulent impact.

Further, inhibiting quorum sensing poses a unique benefit for the development of new antibiotics. Traditional antibiotics, such as vancomycin or linezolid, generally impede the proliferation of bacteria through inhibiting cell wall biosynthesis, transcription and translations mechanisms, or cellular metabolism.8 Each of these processes is required for cell survival, imposing an evolutionary pressure on the bacteria and encouraging the formation of resistance to such treatments over time. Though the quorum sensing pathway governs many of the bacteria’s infectious capabilities, prohibiting chemical communication between cells does not pose a direct threat to individual cell survival. Hence, therapies that efficiently quench the quorum sensing pathway have the potential to treat bacterial infections without rapidly promoting the development of bacterial resistance.9,10

B. N-Acyl Homoserine Lactones (AHLs) in Quorum Sensing (QS)

The identity and structure of the chemical signals responsible for regulating quorum sensing vary from autoinducing peptides (AIPs) in S. aureus,11 to diffusible fatty acids (DSFs) in B. cenocepacia.12 In the majority of gram-negative bacteria, small molecules known as N-acyl homoserine lactones (AHLs) are the agents of communication (Figure 1.2). AHLs have been identified as critical messengers in greater than fifty species of bacteria.4 Each member of the AHL family contains several structural similarities, including a
lipophilic hydrocarbon tail terminated with a characteristic α-amino-γ-butyrolactone. These similarities are matched by select sites of structural variability necessary for signal differentiation, including the presence of a β-hydroxyl (3-OH-AHL) or β-keto moiety (3-oxo-AHL) adjacent to the N-acyl functionality. As described further in Section IC and ID, the structural similarities and differences amongst AHLs has inspired the development of synthetic analogs to inhibit and study QS systems.

![Chemical structures](image-url)

**Figure 1.2.** Structural subfamilies of the known N-acyl-homoserine lactone autoinducers.

Many species of bacteria employ multiple AHLs to regulate different facets of their gene expression profile. In *P. aeruginosa*, the 3-oxo-C₁₂-AHL (1) and its resultant binding to LasR stimulates the production of a series of virulence factors in addition to the transcription of the RhlR receptor for recognition of the C₄-AHL (2) (Figure 1.3).

Subsequent binding of the C₄-AHL leads to production of a distinct set of virulence factors promoting *P. aeruginosa* pathogenicity. The collection of AHLs is unique to each species of bacteria, however multiple species employ the same individual autoinducer. This forwards the potential for crosstalk between gram-negative bacteria, adding an additional layer of complexity to the role of AHLs in the organism’s social behavior.
Figure 1.3. The hierarchal QS signaling pathway in *Pseudomonas aeruginosa*. The 3-oxo-C\(_{12}\)-AHL binds to its cognate receptor LasR to stimulate transcription of the 3-oxo-C\(_{12}\)-AHL synthase LasI and the receptor for the RhlR/RhlI pathway, mediated by the C\(_{4}\)-AHL. The corresponding signal-receptor complexes of both autoinducers encode for virulence factors responsible for the pathogenicity of the species. Adapted from Jimenez, et al.\(^{14}\)

Despite the commonality of *N*-acyl homoserine lactones in gram-negative quorum sensing, few details exist about the reactivity of this family of molecules. Yates and co-workers were the first to fully characterize the susceptibility of the AHLs to lactone hydrolysis in conditions of variable pH.\(^{17}\) Hydrolysis rates increased with elevated media alkalinity in addition to decreasing length of the AHL hydrocarbon tail, alluding to the function of the lipophilic moiety in stabilizing the parent AHL structure during extracellular transport (Scheme 1.1; Path A). Later, Janda and colleagues detected tetramic acids as a secondary product in 3-oxo-AHLs degradation pending media at or above pH 7.4. (Scheme 1.1; Path B) The corresponding acid derivatives were isolated, characterized, and demonstrated to possess bactericidal activity against a variety of gram-positive bacteria, thus
presenting an alternative role of the AHL as mode of chemical defense. These studies exemplify how understanding the innate reactivity of the AHLs can provide greater insight into their evolutionarily determined function.

![Scheme 1.1](image)

**Scheme 1.1.** Degradation pathways of the 3-oxo-AHLs under physiological conditions. Products corresponding to the hydrolysis of the lactone (Path A) detected in all AHLs while an intramolecular rearrangement to the tetramic acid (Path B) is only detected in the 3-oxo-AHLs.

Further understanding of AHL reactivity has been gained through the identification of quorum quenching enzymes in both bacteria and eukaryotes, divulging the chemical basis for natural defense against the N-acyl homoserine lactones. The majority of identified enzymes function either as lactonases, hydrolyzing the AHL lactone, or acylases, hydrolyzing the amide bond. The lactonase AiiA, originally isolated from *Bacillus*, is a dimeric zinc metalloenzyme determined to promote lactone hydrolysis via activation of a single water molecule after metal binding to a conserved protein sequence consisting of mostly histidine and aspartic acid residues. Multiple enzymes with similar conserved sequences have since been identified highlighting a commonality for the mechanism of AHL hydrolysis across multiple bacterial species. The promiscuous family of hydrolytic enzymes in eukaryotes known as paraoxonases (PON), characterized by a Ca\(^{2+}\)-His-His catalytic active site also demonstrated AHL lactonase activity. A large number of AHL acylases have also been
identified with PvdQ from *P. aeruginosa* being well understood due to the recent publication of a substrate-bound crystal structure. Degradation of the AHL proceeds via activation and attack of a nucleophilic serine reside at the amide with surrounding valine and asparagine residues to stabilize the developing negative charge in a oxyanion hole fashion (Figure 1.4). The study of quorum quenching enzymes has aided in the identified reasonable sites for non-enzymatic chemical modification to the AHLs as well as the types of non-covalent interactions that enable such a modification. These examples largely influenced our approach for developing peptide-based enzyme mimics for AHL degradation (see Chapter IV).

**C. Derivatizing AHL Autoinducers for QS Inhibition**

Since the discovery and characterization of the AHLs, the family of autoinducers has served as a scaffold for the development of QS inhibitors. The largest effort over the last twenty years has focused on the creation of small molecules capable of inhibition through competitive receptor binding. Thousands of compounds have been synthesized in library format and screened to offer valuable structure-activity relationships for signal-receptor affinities. Modifications to the AHL framework include heteroatom substitutions at the 3-
oxo position and lactone,\textsuperscript{25,26} alteration of the lactone structure and size,\textsuperscript{27–29} and arylation of the hydrophobic tail.\textsuperscript{4,30,31} (Figure 1.5a). Amongst the array of analogs synthesized, several displayed notable antagonism and agonism of the QS pathway at low micromolar potencies in species that employ the native AHLs for communication, including \textit{P. aeruginosa}, \textit{A. tumefaciens}, and \textit{V. fischeri}. The aggregate data from these screens has led to the establishment of structural guidelines for receptor specific and cross-species analog affinity, forwarding a better understanding of messenger-receptor binding and the creation of more potent analogs.\textsuperscript{13}

\textbf{Figure 1.5.} Manipulation of the AHL to promote QS Inhibition. a) Select examples of AHL analogs and corresponding IC\textsubscript{50} values for LasR inhibition in \textit{P. aeruginosa}.\textsuperscript{27,7} b) AHL sequestration by cyclodextrins, depicting prodigiosin production (\textbullet) and cell growth (\textcircled{O}) in the presence compared to the absence of cyclodextrin (▲, △). The reduction in prodigiosin levels indicating a reduction in viable AHLs in solution for QS regulation.\textsuperscript{32}

Advances in quorum sensing inhibition have also been afforded in directly impeding the function of the AHL by employing chemical methods to reduce the concentration of the active messenger \textit{in situ}. Janda and coworkers synthesized a small number of AHL-inspired haptens to select for monoclonal antibodies capable of degrading the native ligand. The selected antibody served to catalytically hydrolyze the lactone, presenting the first example
Taking advantage of the inherent lipophilicity of AHLs, Kato and colleagues found that the addition of immobilized cyclodextrins to cellular assays reduced the in solution concentration of the chemical messengers (Figure 1.5b). The resultant cyclodextrin-AHL complex afforded attenuated production of the secondary metabolite prodigiosin in the human pathogen *S. marcescens* and was hypothesized to gain added stability through ionic interactions between the signal and its acceptor. These approaches capitalize on the intrinsic functionality and reactivity of the autoinducer. Further, the use of cyclodextrins and catalytic antibodies demonstrate the effectiveness of enzyme mimics in quorum quenching, taking advantage of select non-covalent interactions to forward AHL recognition and QS inhibition.

**D. Derivatizing AHL Autoinducers for Chemical Biology**

The dynamic and complex nature of quorum sensing regulatory pathways has presented difficulties in identifying the biomolecular events that take place from autoinducer binding to gene transcription. Likewise, as more examples of AHL mediated crosstalk between species and eukaryotes surface, great interest lies in developing tools to elucidate environmental factors and the molecular architecture that support these interactions. To generate new chemical biology strategies to study QS, the parent structure of the AHL has again served as a template. Meijler and coworkers completed the synthesis of a 3-oxo-C$_{12}$ AHL analog dually functionalized with a diazirine moiety and terminal alkyne (Figure 1.6a). The unique construct maintained the functional requirements for receptor binding while enabling light-mediated carbene insertion at the diazirine functionality to establish a covalent linkage between the AHL analog and its receptor. Through Huisgen cycloaddition with an azide-functionalized fluorescent tag, the signal-receptor complex was visualized in an *in vitro*
The Blackwell group, by derivatizing the 3-oxo-C₁₂ AHL with a terminal amine, was able to bind and isolate the QscR receptor in *P. aeruginosa* through affinity chromatography. Both methods could serve as valuable tools for elucidating AHL-receptor binding, including identification of receptors in other species that enable the crosstalk through AHL recognition.

**Figure 1.6.** Advances in LasR labeling in *P. aeruginosa* by AHL analog synthesis and *in situ* derivatization of the autoinducer. a) upon binding to LasR, a light-mediated carbene insertion covalently links the AHL to the receptor allowing subsequent fluorescent tag attachment via Huisgen cycloaddition to observe the receptor bound signal. b) covalent attachment of an isothiocyanate 3-oxo-C₁₂-AHL analog is exposed to an aminoxyl-BODIPY derivative forming the corresponding oxime and allowing *in vivo* visualization of LasR.

Further advances in chemical biology have also been realized through the derivatization of the native AHL *in situ*. Meijler and coworkers covalently attached a 3-oxo-
C_{12}-AHL derivative to its cognate receptor in \textit{P. aeruginosa}, LasR. Taking advantage of the carbonyl reactivity that is uniquely characteristic of the 3-oxo-AHLs, an aminoxy functionalized BODIPY dye was then successfully linked to the bound autoinducer (Figure 1.6b). The resultant oxime enabled live cell labeling of LasR for \textit{in vivo} visualization of the location of the receptor within the bacteria.\textsuperscript{38} Such technologies that exploit the unique reactivity of the AHL could be a gateway to tagging and visualizing the autoinducers while active in their role as quorum sensing modulators.

\textbf{E. Statement of Research}

Reflected in this dissertation, research efforts have focused on expanding the suite of chemical based strategies for modifying the N-acyl-homoserine lactones. Inspired by the progress made in derivatizing the AHLs \textit{in situ} for QS inhibition and chemical biology, our approach has centered on exploiting the reactivity of the autoinducers to expose covalent and non-covalent interactions that can promote their degradation. As detailed in the following chapters, the use of amine nucleophiles has led to modes of AHL functionalization that focus on both the \(\gamma\)-butyrolactone found in all AHLs and the ketone characteristic of the 3-oxo-AHLs. The supramolecular interactions provided by biomimetic catalysts, both small molecule and peptide-based, were assessed for their ability to activate the AHL for subsequent covalent modification. Ultimately, our approach has enabled AHL derivatization reactions to advance from catalyzing aminolysis reactions of the AHL in organic solvents to the development of a high-throughput reactive tagging assay for identifying peptides capable of rapid derivatization of the 3-oxo-AHL under physiological conditions. The knowledge uncovered regarding intrinsic AHL reactive and the technologies developed in these
investigations contribute to the ongoing effort to understand virulence mechanisms of quorum sensing while creating new strategies to inhibit them.
REFERENCES


CHAPTER II: THIOUREA AND GUANIDINE CATALYZED AMINOLYSIS OF N-ACYL HOMOSERINE LACTONES

A. Introduction

Chemists have been long inspired by the efficiency and specificity by which nature carries out biochemical processes. Consequently, much time as been devoted to artificially recreating these pathways through synthetic chemistry. The intricate array of hydrogen bonding interactions and proton transfers that in part enable enzyme function has spawned the development of specifically crafted hydrogen bond accepting and donating organocatalysts.\(^1\) Mimicking the supramolecular interactions found in nature, these small molecule catalysts have been developed to improve rates and stereoselectivities of a variety of organic transformations. Two such hydrogen bonding compounds that have garnered wide use are thiourea and guanidine catalysts.

i. Thiourea Catalysis

The thiourea catalyst was first introduced by Jacobsen and coworkers who showcased its ability to accelerate asymmetric Strecker reactions.\(^2\) Further study of the Strecker system revealed that the thiourea operates through a mechanism of dual hydrogen bond donation.\(^3\) (Figure 2.1). The acidity of the thiourea moiety makes it successful in coordinating Lewis basic functionalities such as carbonyls, anions, nitrates, and sulfonic acids.\(^4\) The generality of its method of activation has allowed the thiourea to be employed in Pictet-Spangler reactions\(^5\), Mannich reactions\(^6,7\), Diels-Alder reactions\(^8\), and Michael additions\(^9-11\).
Figure 2.1. Transition state model of Jacobsen’s thiourea catalyst for asymmetric Strecker reactions (minor enantiomer). Hydrogen bond donation of the thiourea to the cyanide nucleophile provided a mechanistic rationale for the observed rates and product stereoselectivities. Adapted from Zuend, et al.12

Advancements in the field of thiourea catalysis have been forwarded through the use of bifunctional derivatives of the catalyst. These catalysts have been appended to impart an element of catalyst chirality or provide additional sites for non-covalent interactions. Incorporation of a chiral hydrogen bond acceptor in close proximity to the thiourea has been demonstrated to enhance nucleophilic addition reactions in both rate and enantioselectivity (Figure 2.2a). For example, Takemoto and colleagues utilized chiral bifunctional thiourea catalysts in the Michael addition of 1,3 diketones to nitroolefins (Figure 2.2b).11 The use of the $N,N$-dimethylcyclohexyl functionalized catalyst 3 proved to be the most effective, rapidly affording addition products with up to 99% ee. Through mechanistic studies, the tertiary amine was determined to play a critical role in the activation of the Michael donor serving as a hydrogen bond acceptor to the incoming enol nucleophile.
Figure 2.2 – Bifunctional thiourea catalysis with Takemoto’s catalyst 3. a) dual activation by catalyst 3 as described by Hedrick\textsuperscript{13} in the transition state of nucleophilic additions to L-lactide. Similar modes of activation have been described in the catalyst’s use in b) Michael additions to nitroolefins\textsuperscript{11}, c) polymerizations of cyclic esters\textsuperscript{13}, and d) kinetic resolutions of azlactones\textsuperscript{14}.

Carbonyl–based substrates, specifically esters and lactones, are also amenable to activation by thioureas. A racemic version of Takemoto’s catalyst was employed by Hedrick and coworkers in the polymerization of cyclic esters (Figure 2.2c).\textsuperscript{13} Through increasing the electrophilicity of the ester, the bifunctional thiourea effectively advanced rates of polymerization with only 5 mol % catalyst loading. Comparable rates of polymerization could not be obtained when the thiourea and dimethylamino moiety were screened.
independently, supporting the necessity of the catalyst’s bifunctionality for reactivity.

Similar modes of catalyst activation were described by Berkessel and colleagues in their use of thioureas for the kinetic resolution of azlactones (Figure 2.2d).\textsuperscript{14}

\begin{center}
\begin{tabular}{lll}
\textbf{Scheme 2.1.} Thiourea catalysis under aqueous conditions. The catalysts have been employed for a) Diels-Alder reactions\textsuperscript{15} and b) enantioselective Michael additions to nitroolefins.\textsuperscript{10} & \\
\end{tabular}
\end{center}

Transformations mediated by thiourea catalysis are most readily conducted in nonpolar, aprotic solvents in order to optimize the hydrogen bonding events necessary for substrate activation. However, several examples in the literature cite successful thiourea-catalyzed transformations in water. Schreiner and Wittkopp witnessed moderate rate accelerations in the Diels-Alder cycloadditions when employing a trifluoromethylated catalyst in a 9:1 water:tert-butanol mixture, observing an 11\% increase in yield after 1 h (Scheme 2.1a). The sustained substrate activation was hypothesized to occur due to the hydrophobicity of the catalyst and reagents, promoting aggregation of the reaction components in aqueous media.\textsuperscript{8} Xiao and coworkers employed bifunctional proline-thiourea catalysts for the asymmetric addition of ketones to nitroolefins in water (Scheme 2.1b).
Though catalysis was largely mediated by the addition of 10 mol% benzoic acid, ee’s of 79-99% were obtained across the catalyst and substrate scope, suggesting necessary engagement of the thiourea during the progression of the reaction.\textsuperscript{10} The potential for the thiourea catalyst to maintain its function of supramolecular activation in water makes it a viable candidate for screening in biological relevant systems.

\textbf{ii. Guanidine Catalysis}

With arginine as the most prevalent example, guanidines are a motif integral to natural chemical processes. The functionality has been detected in a variety of natural products from single-celled prokaryotes to higher order eukaryotes.\textsuperscript{16} The uniquely strong basicity of guanidines renders them protonated across a wide range of pH’s, making them highly capable of anion binding and hydrogen bonding.\textsuperscript{17} Enzymes that involve arginine such as inosine 5’-monophosphate dehydrogenase and fumarate reductase, proceed by mechanisms that involve the amino acid participating in multiple hydrogen bonding interactions or serving as general base.\textsuperscript{18}

\begin{figure}[h]
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\caption{Comparison of properties of cyclic guanidine TBD and acyclic TMG.\textsuperscript{19} The ring strain in TBD inhibits the delocalization of electrons across the guanidine core, increasing its basicity. The bicycle also restricts C=N isomerization and C-N free rotation to disfavor inactive forms of the catalyst as seen with TMG. The active forms of each catalyst are highlighted in green. TMG = 1,1,3,3-tetramethylguanidine.}
\end{figure}
The biological relevance of guanidine has inspired its use in organocatalysis. The functionality has been demonstrated as a suitable catalyst for aldol reactions, carbonylations, methylations, silylations, and conjugate additions.\textsuperscript{20} The most frequently employed guanidines are bicyclic guanidines, such as 1,5,7-triazabicyclo[4.4.0]dec-5-ene or TBD, due to the increase in basicity provided by associated ring strain and transition state planarity relative to their acyclic counterparts. Further, the bicyclic structure preorganizes the catalyst into the active isomer necessary for optimal substrate interaction with the basic guanidine core (Figure 2.3).\textsuperscript{19} These organocatalysts were originally thought to operate solely through general base or hydrogen bonding mechanisms, however instances in which the guanidine serves as nucleophilic catalyst have been well documented.\textsuperscript{20,21}

\begin{center}
\includegraphics[width=\textwidth]{scheme22.png}
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\textbf{Scheme 2.2.} Transition states in the revised mechanism of TBD-catalyzed cyclic ester polymerizations. TBD operates through a dual hydrogen bonding mechanism, simultaneously activating both the ester and alcohol (\textbf{TS1}). Stabilization of the resultant oxyanion and hydrogen bonding to oxygen adjacent to the carbonyl (\textbf{TS2}) enables rapid generation of the acyclic product.

Guanidine catalysts have also been demonstrated to be effective in accelerating transesterifications and acyl transfer reactions by greater than two orders of magnitude over
the uncatalyzed reaction. The mechanism by which these guanidine catalyzed transformations proceed has been debated, being shown to vary depending on the reaction and substrate. Waymouth and colleagues employed TBD as a catalyst for the polymerization of cyclic esters.\textsuperscript{22,23} Identification of an \(N\)-acyl intermediate by \(^1\text{H}\) NMR led to the conclusion that the increase in polymerization rates was due to TBD serving as a nucleophilic catalyst. DFT calculations by Simon and Goodman on polymerizations of lactones challenged the proposed mechanism, suggesting a lower energy hydrogen bonding pathway. This mechanism depicts the bifunctionality of the catalyst, activating both the incoming nucleophile and substrate, prior to an ultimate proton transfer from the catalyst to the formed tetrahedral intermediate to afford the ring-opening product (Scheme 2.2).

Further, \(\beta\)-butyrolactone was not amenable to TBD catalysis but still detectably formed what was deemed a non-productive \(N\)-acyl intermediate.\textsuperscript{24} The hydrogen bonding mechanism was later confirmed by Waymouth in an independent set of computational experiments.\textsuperscript{25}

\begin{scheme}
\begin{align*}
\text{a)} & \quad \text{TBD-catalyzed amidation of benzyl acetate. Waymouth and colleagues determined catalysis to occur through a nucleophilic mechanism involving first-order formation and decay of an \(N\)-acylated TBD intermediate.} \\
\text{b)} & \quad \text{\(N\)-acylated TBD is less energetically stable than the corresponding TBO analog, supporting the increased reactivity of TBD.}
\end{align*}
\end{scheme}

In contrast, the ability of TBD to function as a nucleophilic catalyst has been demonstrated in acyl transfer reactions with acyclic esters. Waymouth proved such a
mechanism in the amidation of benzyl acetate by observing the reaction to proceed with first order kinetics in both the generation and decay of the corresponding acetylated-TBD intermediate by $^1$H NMR (Scheme 2.3a). The importance of the structure, namely the N-C=N dihedral angle for reactivity of the formed intermediate, was also highlighted as the rates of reaction decreased on substituting the less spatially constrained 1,4,6-triazabicyclo[3.3.0]oct-4-ene (TBO) for TBD.$^{26}$ A twist in the dihedral angle of the acyl-TBD species was determined by DFT calculations, supporting its relatively increased reactivity (Scheme 2.3b). Though variable in its mode of catalysis, TBD is unique in its ability to activate lactone-based substrates for addition reactions by modes inspired by nature.

iii. Rationale for AHL aminolysis

Our investigations into fundamental AHL reactivity and activation began by targeting the lactone of the chemical messenger. Modification of this functionality is a pathway for the deactivation of the signal in nature (see Chapter I) and is a more chemically labile target than the N-acyl bond. The $\gamma$-butyrolactone is also present across all known N-acyl homoserine lactones, holding the potential for our developed methodology to be applicable across multiple gram-negative species.

Initially, we tested the susceptibility of the AHL to modification by transesterification. By employing general base catalysts such as imidazole and N-methylimidazole, the acyclic derivative of the AHL was obtained.$^{27-29}$ However, the reaction required solvent quantities of the transesterification agent, in this case methanol. Moreover, in monitoring the reaction by analytical HPLC, it was evident that a solvent dependent equilibrium between the methyl ester and the lactone was established. The resistance of the
autoinducers to progress fully to the methanolysis product indicated that the system strongly favors the reverse reaction and any obtained AHL methyl ester will be readily susceptible to recyclization (Figure 2.4).

![Reaction Scheme](image)

**Figure 2.4.** Equilibrium established between the 3-oxo-C_{12}-AHL (1) and its methyl ester. In methanol, 78% of the methyl ester was obtained after 75 hours in the presence of imidazole. Upon introduction of equimolar amounts of acetonitrile (red line), recyclization from the methyl ester to the lactone was observed.

The observed dynamics of AHL transesterification supported the necessity of forming a more thermodynamically stable acyclic product to irreversibly modify the autoinducer. Consequently, we investigated amine nucleophiles for AHL lactonolysis, hypothesizing that the resultant amide from such a transformation would essentially negate the contribution of the reverse reaction (recyclization). Despite the energy gained in the formation of the amide, the relatively low ring strain of five membered lactones make them resistant to modification.
by a nucleophile.\textsuperscript{30} Attempted additions to $\gamma$-butyrolactone based substrates have required high temperatures, high amine equivalents, and long reaction times. Hence, Lewis acids, such as AlCl$_3$, and other catalysts have often been employed to increase rates and conversions.\textsuperscript{31,32}

\textbf{Scheme 2.4.} Aminolysis of the C$_6$-AHL with piperidine.

To probe the ability of the AHL to be modified by aminolysis, the C$_6$-AHL (4) was combined with 20 eq. of piperidine in MeCN (Scheme 2.4). The reaction reached complete conversion to the corresponding aminolysis product (5) after 4 days at RT. Despite the sluggish rates of conversion, the product was isolated with no detected tendency for cycloreversion.

\textbf{Figure 2.5.} Proposed mode of AHL activation by a thiourea catalyst.

As an irreversible modification of the AHL had been achieved, we sought to discover a means of catalyzing this transformation to increase reaction rates and divulge functionality capable of activating the AHL as a substrate. Previous research on thioureas and guanidines
as efficient catalysts for carbonyl additions (see Introduction) made them attractive
candidates for screening for AHL aminolysis reactions. We envisioned that both catalysts
could serve as hydrogen bond donors to lower the LUMO at the carbonyl and increase the
electrophilicity of the lactone, thus activating it for acyl transfer (Figure 2.5). Both catalysts
also have the potential to be readily modified to impose elements of bifunctional activation
and synthetically tuned to alter properties of acidity and steric constraint for the purpose of
optimizing catalytic efficiency.

B. Results and Discussion

i. Thiourea-catalyzed AHL aminolysis

As an initial proof of concept for our proposed mode of activation, thiourea catalyst
7a was synthesized by addition of cyclohexylamine to the corresponding isothiocyanate and
added to the reaction of piperidine and the C6-AHL. Utilizing 10 mol % of the catalyst,
complete conversion was obtained in 19 h, equating to approximately six times the rate of the
uncatalyzed reaction in MeCN.
Table 2.1 Thiourea catalyst scope and percent yields.

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<th>Catalyst</th>
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With the initial success of catalyst 7a, we synthesized a small library of thioureas to determine if such mild rates of catalysis could be improved. Nine potential organocatalysts were synthesized and can be compared based on their potential for multimodal interactions with the substrate and nucleophile (Table 2.1). Catalysts 7a and 7b are monofunctional, relying solely on the thiourea for substrate activation. The remaining catalysts each contain an element of bifunctionality with the potential to further activate the AHL or nucleophile.
Catalysts 7c, 7d, 7e, and 3 are all appended with tertiary amines capable of serving as hydrogen bond acceptors to assist in enhancing the nucleophilicity of piperidine.

Tetraalkylammonium 7f could provide stability to the developing anionic transition-state characteristic of amine additions to carbonyl. Dithiourea catalysts 7g and 7h possess two thioureas that may act individually on two AHLs or synergistically on a single molecule of substrate by creating a more extensive surface of available hydrogen bond donors. The 3,5-bis(trifluoromethyl)phenyl group was retained in all catalysts due to its ability to inductively increase the acidity of the thiourea and provide additional stabilization with a uniquely polar ortho-CH bond.34

![Diagram of AHL-Catalyst Interaction via 1H NMR](image)

**Figure 2.6.** Evidence of an AHL-thiourea interaction by $^1$H NMR. A downfield shift in the N-H resonances of 10 mol % of catalyst 7a is observed upon addition of the C$_6$-AHL.
The catalysts were then screened for activity in the same piperidine/C$_6$-AHL model system as described above by $^1$H NMR, monitoring the integration of the $\alpha$-proton resonance over time. Prior to the addition of piperidine, an observable shift was detected in the proton resonances of the thiourea catalyst in the presence of the AHL (Figure 2.6). This confirms the hypothesized interaction between the thiourea and the homoserine lactone.

**Table 2.2** Relative rates of thiourea catalyzed aminolysis in CD$_3$CN and DMF-d$_7$.

![Reaction Scheme](image)

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>$k_{rel}$ CD$_3$CN</th>
<th>$k_{rel}$ DMF-d$_7$</th>
</tr>
</thead>
<tbody>
<tr>
<td>7a</td>
<td>4.5</td>
<td>3.4</td>
</tr>
<tr>
<td>7b</td>
<td>5.6</td>
<td>3.6</td>
</tr>
<tr>
<td>7c</td>
<td>4.3</td>
<td>3.4</td>
</tr>
<tr>
<td>7d</td>
<td>4.4</td>
<td>3.3</td>
</tr>
<tr>
<td>7e</td>
<td>5.5</td>
<td>3.2</td>
</tr>
<tr>
<td>7f</td>
<td>4.5</td>
<td>-</td>
</tr>
<tr>
<td>3</td>
<td>8.2</td>
<td>-</td>
</tr>
<tr>
<td>7g</td>
<td>7.9</td>
<td>5.2</td>
</tr>
<tr>
<td>7h</td>
<td>10.0</td>
<td>5.3</td>
</tr>
</tbody>
</table>

*The reactions were monitored until 50% conversion ($t_{50}$) was reached*

The entire library of catalysts proved active in enhancing rates of AHL aminolysis (Table 2.2). Ring-opening rates up to 8.2 times the uncatalyzed reaction were achieved among the monofunctional and bifunctional catalysts screened with dithiourea catalyst 7h reaching an increase of an order of magnitude relative to the control. However, in comparing the activity of the monofunctional and bifunctional catalysts, those catalysts which include secondary sites of activation served to provide little to no advantage as the rates obtained were similar. Takemoto’s catalyst (3) generated the aminolysis product at twice the rate of
any monofunctional thioureas, supporting the involvement of the $N,N,\text{-dimethylamino}$ group in catalysis. This mild acceleration relative to use of the same catalyst in previously published systems$^{11,13,14}$ may be due to the inherent weakness of the interaction of the tertiary amine with the nucleophile, namely the N-H---N hydrogen bond. In the case of catalysts 7c – 7f, the lack of conformational constraints to position the bifunctionality at the catalytic active site is one possible explanation for the observed rates.

**Table 2.3.** Relative rates of aminolysis of different AHLs in CD$_3$CN$^{a}$

<table>
<thead>
<tr>
<th>AHL</th>
<th>Structure</th>
<th>$k_{rel, 7a}$</th>
<th>$k_{rel, 7h}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td><img src="image1" alt="Structure 1" /></td>
<td>2.7</td>
<td>4.0</td>
</tr>
<tr>
<td>2</td>
<td><img src="image2" alt="Structure 2" /></td>
<td>4.1</td>
<td>7.3</td>
</tr>
<tr>
<td>4</td>
<td><img src="image4" alt="Structure 4" /></td>
<td>4.5</td>
<td>10</td>
</tr>
</tbody>
</table>

$^{a}$with piperidine as the amine nucleophile in 20-fold excess.

Next, the transferability of the thiourea catalysts across the AHL family was assessed. Inspired by the autoinducers responsible for quorum sensing in *P. aeruginosa*, we screened both catalyst 7a and 7h against the 3-oxo-C$_{12}$ (1) and C$_4$-AHL (2) (Table 2.3). The C$_4$-AHL proceeded in a similar fashion to the C$_6$-AHL. The 3-oxo-C$_{12}$-AHL showed a marked decrease in its susceptibility to thiourea catalysis achieving rates of only four times the control reaction with catalyst 7h. The 3-oxo-C$_{12}$-AHL was also sparingly soluble in CD$_3$CN and substrate aggregation could negatively impact the interaction of the autoinducer with the catalyst. In addition, the extra carbonyl moiety characteristic of the 3-oxo-AHL could be serving as a site for competitive catalyst binding.
With the ability of AHL aminolysis to be accelerated by H-bonding catalysts established, the impact of solvent background reactivity and catalysis was tested (Table 2.4). First, catalysts 7a and 7h were screened in aprotic solvents with a wide range of polarity. Background reaction rates markedly increased as solvent polarity was increased, however catalysis was still observed. Notably, the impact of the thioureas was evident in DMF-d$_7$, a solvent in which competitive hydrogen bonding was expected to deactivate the catalyst.

Table 2.4. Relative rates of C$_6$-AHL aminolysis in the presence of catalysts 7a and 7h in aprotic solvents and polar protic solvent mixtures

<table>
<thead>
<tr>
<th>Entry</th>
<th>Solvent (% v/v)</th>
<th>$t_{50}%$ uncat ($\times 10^3$ s)</th>
<th>$k_{rel}$ 7a</th>
<th>$k_{rel}$ 7h</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>C$_6$D$_6$</td>
<td>5.1</td>
<td>2.0</td>
<td>4.3</td>
</tr>
<tr>
<td>2</td>
<td>CD$_2$Cl$_2$</td>
<td>3.6</td>
<td>1.0</td>
<td>1.1</td>
</tr>
<tr>
<td>3</td>
<td>CD$_3$CN</td>
<td>35.5</td>
<td>4.5</td>
<td>10</td>
</tr>
<tr>
<td>4</td>
<td>DMF-d$_7$</td>
<td>100.7</td>
<td>3.4</td>
<td>5.3</td>
</tr>
<tr>
<td>5</td>
<td>DMSO-d$_6$</td>
<td>123.2</td>
<td>1.9</td>
<td>2.2</td>
</tr>
<tr>
<td>6</td>
<td>10% MeOH-d$_4$/CD$_3$CN</td>
<td>14.3</td>
<td>0.9</td>
<td>1.5</td>
</tr>
<tr>
<td>7</td>
<td>10% D$_2$O/ DMSO-d$_6$</td>
<td>21.0</td>
<td>1.3</td>
<td>1.0</td>
</tr>
<tr>
<td>8</td>
<td>10% TFE-d$_3$/CD$_3$CN</td>
<td>9.7</td>
<td>1.1</td>
<td>1.0</td>
</tr>
<tr>
<td>9</td>
<td>10% D$_2$O/CD$_3$CN</td>
<td>4.2</td>
<td>1.0</td>
<td>0.9</td>
</tr>
</tbody>
</table>

*with piperidine as the amine nucleophile in 20-fold excess; $b_{t_{50}\%}$ represents the reaction time to reach 50% conversion

Next, with the goal of determining if catalysis would be sustained under physiological conditions, the catalysts were screened in a series of protic co-solvents. A significant increase in the uncatalyzed reaction rate was observed with 10% v/v of a protic additive, supporting a component of protic solvent assistance in the aminolysis reaction. Further, the impact of the thiourea catalysts under these conditions was negated. Unlike the previously
discussed examples of thiourea catalysis in water\textsuperscript{8,10}, the presence of protic additives inhibited the supramolecular events necessary for catalysis in our system.

**ii. Guanidine-catalyzed AHL aminolysis**

In an effort to improve upon rates obtained in our thiourea screens, we undertook a short study of the impact of guanidines on enhancing AHL reactivity. With its success in catalyzing transesterification and acyl transfer reactions, TBD was selected for an initial screen in the C\textsubscript{6}-AHL/piperidine control reaction described above (Scheme 2.4). Upon addition of 10 mol % TBD, complete conversion to the aminolysis product was achieved in 1.5 h or 65 times faster than the control reaction. Notably, these rates are 6.5 times faster than our most successful thiourea catalyst 7h.

\begin{center}
\begin{tikzpicture}
\node at (0,0) {\includegraphics[width=0.8\textwidth]{scheme2_5.png}};
\end{tikzpicture}
\end{center}

**Scheme 2.5.** Generation of N-acylated TBD intermediate (9) at variable equilibration times. Preforming the N-acyl intermediate inhibits formation of AHL aminolysis product. Relative t\textsubscript{50\%} values were determined by \textsuperscript{1}H NMR.

The significant rate enhancement compared to the previously screened thiourea catalysts made further study of the TBD system critical for understanding the mode of AHL activation. As noted in beginning of Chapter 2, TBD has been proposed to serve as both a nucleophilic catalyst and hydrogen bond donor depending on the reaction and substrate. To
assess the possibility of catalysis by a nucleophilic pathway, the C₆-AHL (5) was combined with a stoichiometric amount of TBD in the absence of piperidine to test for the formation of an N-acyl intermediate (9) by ¹H NMR. Observable shifts in the catalyst and substrate proton resonances were detected and the corresponding intermediate was confirmed by 2D NMR spectroscopy. However, upon addition of piperidine to a solution containing the preformed intermediate, the rates of conversion to aminolysis product 6 dramatically decreased (Scheme 2.5), suggesting that formation of the N-acyl intermediate was catalytically unproductive. If the acylation of TBD was a key step in the catalytic cycle, the intermediate would have readily dissipated in the presence of a nucleophile. Hence, similar to Goodman’s observations in the transesterification of β-butyrolactone, the aminolysis of the AHL was likely not proceeding through a nucleophilic pathway and more reasonably by acting through a hydrogen bond mediated mechanism.²⁴

To demonstrate the importance of hydrogen bond donation in the TBD-catalyzed aminolysis of the autoinducer, a series of amidine and isothiourea catalysts (Table 2.5) were synthesized and screened. Both amidines and isothioureas have been employed in acyl transfer catalysis but are incapable of serving as hydrogen bond donors.³⁵–³⁹ When combined with the C₆-AHL in the presence of piperidine, no increase in reaction rate was afforded relative to the control. The same decrease in activity was observed in screening MTBD (10b), in which methylation of the guanidine expunges the potential of the catalyst to participate hydrogen bond donation to the substrate. Both examples corroborate the function of TBD as a hydrogen bond donor and not a nucleophilic catalyst in our system.
Table 2.5. Relative rates of amidine and isothiourea-mediated C₆-AHL aminolysis with piperidine in CD₃CN.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>k_{rel}</th>
<th>Catalyst</th>
<th>k_{rel}</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>64.8</td>
<td>10c</td>
<td>1.1</td>
</tr>
<tr>
<td>10a</td>
<td>2.0</td>
<td>10d</td>
<td>1.1</td>
</tr>
<tr>
<td>10b</td>
<td>1.5</td>
<td>10e</td>
<td>1.1</td>
</tr>
</tbody>
</table>

Establishing the importance of TBD as a hydrogen bond donor, we sought to assess the interplay of the basicity of the guanidine in the increasing aminolysis rates in addition to its potential for bifunctional activation. An equimolar amount of trifluoroacetic acid was titrated into a reaction mixture containing the catalyst and substrate to fully protonate the guanidine and eliminate secondary modes of activation beyond hydrogen bond donation (Scheme 2.6). Upon the addition of piperidine, the resulting guanidinium (11) was inactive as rates obtained were similar to the uncatalyzed reaction. The inactivity of the protonated guanidine demonstrates that TBD is not acting solely through a mechanism of hydrogen bond donation and does employ a secondary mode of activation in the derivatization of the QS autoinducer.

Scheme 2.6. Relative rates of aminolysis upon TBD protonation. An equivalent of trifluoroacetic acid (TFA) was added to form the corresponding guanidinium ion (11). The loss of catalytic activity indicates that TBD does not exclusively serve as an H-bond donor.

Table 2.6. Relative rates of C₆-AHL aminolysis with decreasing catalyst basicity in CD₃CN.
To further explore the importance of basicity to catalyst function, catalysts 12a and 12b were obtained and tested due to their reduced pKa values relative to TBD. When each was added in 10 mol % quantities to the control reaction of C₆-AHL with piperidine, attenuation of catalytic rates was observed (Table 2.6). The decrease in relative rates of catalysis directly correlates to the decrease of the pKa of the catalyst compared to TBD. Thus, the basicity of TBD is critical in achieving accelerated rates of AHL aminolysis.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Structure</th>
<th>pKa⁶</th>
<th>k_rel</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td><img src="image1" alt="Structure" /></td>
<td>26.0</td>
<td>64.8</td>
</tr>
<tr>
<td>12a</td>
<td><img src="image2" alt="Structure" /></td>
<td>(12.5)</td>
<td>2.0</td>
</tr>
<tr>
<td>12b</td>
<td><img src="image3" alt="Structure" /></td>
<td>(5.3)</td>
<td>1.5</td>
</tr>
</tbody>
</table>

⁶pKa in CD₃CN; parenthesis indicate pKa in MeOH-d₄/D₂O

Figure 2.7. Bifunctional activation of the AHL and amine nucleophile by TBD.

The collection of results from this short study led us to hypothesize that the TBD is acting through a bifunctional model of activation, similar to that described by Waymouth and colleagues.²⁶ As the contribution of hydrogen bond donation to the catalytic efficiency of TBD has been proven, the neighboring basic site could reasonably be serving as a hydrogen bond acceptor to activate piperidine during nucleophilic attack (Figure 2.7). This dual mode
of activation is likely supported by the uniquely small dihedral angle of TBD that permits both donor and acceptor functions of the catalyst to work in tandem. Further mechanistic investigation and catalyst development is necessary to confirm these mechanistic details, but the decision was made to not to complete these studies in lieu of as to pursuing novel modes of activating and modifying the AHLs under aqueous conditions (Chapter 3 & 4).

C. Conclusions

By probing the reactivity of the characteristic γ-butyrolactone of the AHL family of quorum sensing autoinducers, their susceptibility to ring-opening by amine nucleophiles was demonstrated. In contrast to attempts at transesterification of the lactone, complete conversion was obtained through the formation of a more thermodynamically stable, amidated product. The introduction of thiourea catalysts demonstrated that the incorporation of hydrogen bond donors can increase the electrophilicity of the autoinducer for subsequent addition. Employing a guanidine catalyst, TBD, in a similar fashion reinforced substrate activation by hydrogen bond donation while achieving greater conversion rates. From our experimental evidence, it is reasonable to attribute this increase in rate to the elevated basicity of TBD and complimentary structure of the guanidine core that supports an effective bifunctional catalytic pathway for lactone aminolysis.

This is the first time small molecule organocatalysts have been employed to activate the QS autoinducers and have provided valuable information into the non-covalent interactions that advance their derivatization. Though the catalytic activity of both the thiourea and guanidine catalysts was not sustained in protic environments, these interactions relate to those observed in enzymatic catalytic sites that have evolved for degradation of the signaling molecule. Moving away from small molecule catalysts towards peptide-based
enzyme mimics may provide the opportunity to exploit our findings about AHL reactivity in an aqueous milieu.

D. Experimental

i. General Reagents and Catalyst Synthesis

Commercial reagents and catalysts, including TBD (8), MTBD (10b), DBU (10a), and THQPO (12b) were obtained from Sigma-Aldrich and Fisher Scientific. Column chromatography of all compounds, except catalyst 7f, was completed using Fisher 230-400 Mesh Grade 60 silica gel. Catalyst 7f was purified using a Waters semi-prep HPLC/UV-Vis with an XBridge 5μM C18 column as detailed in the procedure below. NMR solvents were purchased from Cambridge Isotopes Laboratories. Product characterization and kinetic experiments were completed on either a 600 MHz or 500 MHz Bruker NMR spectrometer. All related coupling constants are reported in Hertz and shifts in ppm. Elemental analyses (C, H and N) were completed by Robertson Microlit Laboratories, Inc. High resolution mass spectrometry (HRMS) was completed by the Mass Spectrometry Laboratory at the School of Chemical Sciences, University of Illinois using ESI on a Waters Q-ToF Ultima mass spectrometer.

General Procedure for Thiourea Catalyst Synthesis Preparation

To a 25-mL Schlenk flask flushed with N2, 3,5-bis(trifluoromethyl)phenyl isothiocyanate 6 (0.336 mL, 1.84 mmol) was added and then solvated in benzene (1 mL). The corresponding amine (1.84 mmol) was added slowly to the flask. The walls of flask were washed with benzene (1 mL) and the reaction was allowed to stir at room temperature for 3 h. The reaction progress was monitored by TLC. The reaction mixture was concentrated under
vacuum and redissolved in DCM. The product was purified by column chromatography on silica gel with a gradient mobile phase [DCM to MeOH/DCM (1:9)].

1-(3,5-bis(trifluoromethyl)phenyl)-3-cyclohexylthiourea (7a)

\[
\begin{align*}
\delta_H (\text{DMSO}-d_6, 500 \text{ MHz}) & \quad 9.88 (1\text{H}, \text{s}), 8.24 (2\text{H}, \text{s}), 8.18 (1\text{H}, \text{s}), 7.72 (1\text{H}, \text{s}), 4.12 (1\text{H}, \text{br s}), 1.92 (2\text{H}, \text{m}), 1.71 (2\text{H}, \text{m}), 1.58 (1\text{H}, \text{m}), 1.35 – 1.29 (5\text{H}, \text{m}); \\
\delta_C (\text{DMSO}-d_6, 125 \text{ MHz}) & \quad 179.1, 141.9, 130.0 (q, J_{CF} = 33.9 \text{ Hz}), 123.2 (q, J_{CF} = 272.7 \text{ Hz}), 121.6, 115.7, 52.2, 31.5, 25.0, 24.3; \\
\text{HRMS (ESI) calcd for C}_{15}H_{17}N_2F_6S: 371.1017, \text{ found: 371.1014 (M + H)}. 
\end{align*}
\]

1-(3,5-bis(trifluoromethyl)phenyl)-3-propylthiourea (7b)

\[
\begin{align*}
\delta_H (\text{DMSO}-d_6, 500 \text{ MHz}) & \quad 10.00 (1\text{H}, \text{s}), 8.23 (3\text{H}, \text{br s}), 7.69 (1\text{H}, \text{s}), 7.72 (1\text{H}, \text{s}), 3.40 (2\text{H}, \text{m}), 1.56 (2\text{H}, \text{dt, } J = 7.5 \text{ Hz}), 0.89 (3\text{H}, \text{t, } J = 7.5 \text{ Hz}); \\
\delta_C (\text{DMSO}-d_6, 125 \text{ MHz}) & \quad 180.2, 141.7, 129.8 (q, J_{CF} = 32.7 \text{ Hz}), 122.9 (q, J_{CF} = 272.9 \text{ Hz}), 121.4, 115.5, 45.3, 21.4, 11.0; \\
\text{HRMS (ESI) calcd for C}_{12}H_{13}N_2F_6S: 331.0712, \text{ found: 331.0704 (M + H)}; \\
\text{Anal. Calc. for C}_{12}H_{13}N_2F_6S: C, 43.64; H, 3.66; N, 8.48; \text{ found: C, 43.56, H, 3.66, N, 8.35}. 
\end{align*}
\]

1-(3,5-bis(trifluoromethyl)phenyl)-3-(2-(dimethylamino)ethyl)thiourea (7c)
δ\textsubscript{H} (DMSO-d\textsubscript{6}, 500 MHz) 10.27 (1H, s), 8.26 (2H, s), 8.10 (1H, s), 7.72 (1H, s), 3.56 (2H, br q), 2.44 (br t, 2H), 2.19 (6H, s); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 150 MHz) 180.0, 141.9, 130.2 (q, \(J_{CF} = 33.2\) Hz), 123.2 (q, \(J = 273.1\) Hz), 121.3, 115.8, 56.8, 44.9, 41.7; HRMS (ESI) calcd for C\textsubscript{13}H\textsubscript{16}N\textsubscript{3}F\textsubscript{6}S: 360.0696, found: 360.0692 (M + H).

1-(3,5-bis(trifluoromethyl)phenyl)-3-(3-(dimethylamino)propyl)thiourea (7d)

δ\textsubscript{H} (DMSO-d\textsubscript{6}, 600 MHz) δ10.09 (1H, s), 8.33 (1H, s), 8.23 (2H, s), 7.70 (1H, s), 3.51 (2H, br s), 2.27 (2H, t, \(J = 6.60\) Hz), 2.12 (6H, s), 1.69 (2H, p, \(J = 6.60\) Hz); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 150 MHz) δ180.3, 141.9, 130.2 (q, \(J_{CF} = 33.2\) Hz), 123.2 (q, \(J_{CF} = 273.1\) Hz), 121.8, 115.9, 56.8, 45.0, 42.6, 25.9; HRMS (ESI) calcd for C\textsubscript{14}H\textsubscript{18}N\textsubscript{3}F\textsubscript{6}S: 374.1126, found: 374.1126 (M + H).

1-(2-(1H-imidazol-4-yl)ethyl)-3-(3,5-bis(trifluoromethyl)phenyl)thiourea (7e)

δ\textsubscript{H} (DMSO-d\textsubscript{6}, 500 MHz) 11.90 (1H, br s), 10.21 (1H, s), 8.27 (3H, br s), 7.72 (1H, s), 7.57 (1H, s), 6.88 (1H, s), 3.76 (2H, br q), 2.81 (2H, t, \(J = 7.5\) Hz); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 125 MHz) 180.2, 141.8, 134.4, 130.0 (q, \(J_{CF} = 32.7\) Hz), 128.2, 123.1 (q, \(J_{CF} = 272.9\) Hz), 121.6, 115.8,
43.8, 26.0; HRMS (ESI) calcd for C\textsubscript{14}H\textsubscript{13}N\textsubscript{4}F\textsubscript{6}S: 383.0765, found: 383.0772 (M + H); Anal. Calc. for C\textsubscript{14}H\textsubscript{13}N\textsubscript{4}F\textsubscript{6}S: C, 43.98, H, 3.16, N, 14.65; found: C, 44.01, H, 3.20, N, 14.61.

1-(3,5-bis(trifluoromethyl)phenyl)-3-((1R,2R)-2-(dimethylamino)cyclohexyl)thiourea (3)

\[
\begin{align*}
\text{δ}_H (\text{DMSO-d}_6, 600 \text{ MHz}) & 10.02 (1H, s), 8.23 (1H, s), 8.17 (2H, s), 7.69 (1H, s), 4.10 (1H, br s), 2.54 (1H, s), 2.22 (7H, s), 1.85 (1H, br s), 1.76 (1H, br s), 1.64 (1H, d, J = 10.2 \text{ Hz}), 1.32 – 1.09 (4H, m); \\
\text{δ}_C (\text{DMSO-d}_6, 125 \text{ MHz}) & 179.0, 142.4, 130.8 (q, J_{CF} = 27.5), 123.7 (q, J_{CF} = 226 \text{ Hz}), 121.4, 115.9, 65.5, 55.7, 32.1, 25.1, 25.0, 21.5; \\
\text{MS (ESI)} & \text{calcd for C}_{17}H_{21}N_3F_6S: 414.13; \text{found: 414.17 (M+H)}
\end{align*}
\]

**Synthesis of Catalyst 7f**

Catalyst 7f was synthesized through a modified version of the general procedure above. In a 2-mL eppendorf tube, (2-aminoethyl)trimethylammonium chloride hydrochloride (50 mg, 0.286 mmol) was combined with N,N-diisopropylethylamine (0.050 mL, 0.286 mmol). Then, 6 (0.052 mL, 0.286 mmol) was then added to the tube. The reaction mixture was vortexed for 10 min in the eppendorf tube. The reaction mixture was quenched by adding 2 mL of MeOH to the tube. The reaction mixture was concentrated and redissolved in distilled water. The resultant aqueous solution was filtered and diluted to prepare for purification. The product was purified by semi-prep HPLC with a gradient mobile phase [MeCN/H\textsubscript{2}O (1:4) to MeCN/H\textsubscript{2}O (1:1)]. Notably, a small percentage of trifluoroacetic acid was retained by the catalyst after purification and is present in the \textsuperscript{13}C NMR. Given the large
excess of piperidine present in the catalyst screening, this trace acid did not significantly affect the rate of AHL aminolysis.

2-(3-(3,5-bis(trifluoromethyl)phenyl)thioureido)-N,N,N-trimethylethanaminium (7f)

\[
\text{δ}_H (\text{DMSO-d}_6, 500 \text{ MHz}) 10.78 (1\text{H, br s}), 8.80 (1\text{H, s}), 8.26 (2\text{H, s}), 7.80 (1\text{H, s}), 3.99 (2\text{H, m}), 3.54 (1\text{H, t}, J = 6.50), 3.14 (9\text{H, s}); \text{δ}_C (\text{CD}_3\text{CN-d}_3, 125 \text{ MHz}) 182.9, 160.8 (q, J = 35.2), 142.4, 131.5 (q, J_{CF} = 32.7 \text{ Hz}), 124.1 (q, J_{CF} = 271.6 \text{ Hz}), 123.5, 64.8, 54.1, 38.7; \text{HRMS (ESI) calcd for C}_{14}H_{18}N_{3}F_{6}S: 374.1126, found: 374.1120 (M + H).}

Synthesis of Catalysts 7g and 7h

Catalysts 7g and 7h were synthesized following the general procedure but reacting 2 eq (3.68 mmol) of 6 with 1 eq (1.84 mmol) of a diamine. Ethylenediamine was the diamine chosen to construct 7g and 1,3 diaminopropane to make 7h.

1,1'-(ethane-1,2-diyl)bis(3-(3,5-bis(trifluoromethyl)phenyl)thiourea) (7g)

\[
\text{δ}_H (\text{DMSO-d}_6, 500 \text{ MHz}) 10.14 (2\text{H, s}), 8.34 (2\text{H, s}), 8.22 (4\text{H, s}), 7.72 (2\text{H, s}), 3.77 (4\text{H, br s}); \text{δ}_C (\text{DMSO-d}_6, 125 \text{ MHz}) 180.7, 141.4, 129.9 (q, J_{CF} = 32.7 \text{ Hz}), 122.9 (q, J_{CF} = 272.9}
Hz), 122.0, 116.0, 42.5; HRMS (ESI) calcd for C$_{20}$H$_{15}$N$_4$F$_{12}$S$_2$: 603.0547, found: 603.0546 (M + H); Anal. Calc. for C$_{20}$H$_{15}$N$_4$F$_{12}$S$_2$: C, 39.87; H, 2.34; N, 9.30; found: C, 40.15; H, 2.47; N, 9.06.

1,1’-(propane-1,3-diyl)bis(3-(3,5-bis(trifluoromethyl)phenyl)thiourea) (7h)

\[
\begin{align*}
\delta_H (\text{DMSO-}d_6, 600 \text{ MHz}) & \quad 10.09 (2H, s), 8.26 (2H, s), 8.22 (4H, s), 7.70 (2H, s), 3.57 (4H, s), 1.89 (2H, p, J = 6.6 \text{ Hz}); \\
\delta_C (\text{DMF-}d_7, 125 \text{ MHz}) & \quad 181.6, 142.7, 131.1 (q, J_{CF} = 32.7 \text{ Hz}), 123.8 (q, J_{CF} = 271.5 \text{ Hz}), 122.6, 116.5, 42.1, 28.3; \\
\text{HRMS (ESI) calcd for C}_{21}\text{H}_{17}\text{N}_4\text{F}_{12}\text{S}_2 & \quad 617.0703, \text{ found 617.0699 (M + H); Anal. Calc. for C}_{21}\text{H}_{17}\text{N}_4\text{F}_{12}\text{S}_2: C, 40.91; H, 2.62; N, 9.09, \text{ found: C, 40.64; H, 2.58; N, 8.86.}
\end{align*}
\]

Synthesis of Catalyst 10e

To a N$_2$ flushed flask, 6 (1.00 mL, 8.4 mmol) was dissolved in 20 mL of THF followed by the addition of 3-aminopropan-1-ol (0.61 mL, 8.0 mmol). The reaction mixture was allowed to stir at RT for 5 hours and then concentrated under vacuum. The reaction mixture was redissolved in concentrated HCl and stirred at 80°C for 4h. After cooling the reaction to room temperature, 15 mL of 2.5 M NaOH was added and allowed to stir for ten minutes. The resultant reaction mixture was vacuum filtered to afford the desired product.

(Z)-N-(3,5-bis(trifluoromethyl)phenyl)-1,3-thiazinan-2-imine (10e)
51% yield. $\delta_H$ (CD$_3$CN, 600 MHz) 7.67 (2H, s), 7.57 (1H, s), 3.48 (2H, t), 3.09 (2H, t), 2.02 (2H, m); $\delta_C$ (CD$_3$CN, 125 MHz) 152.7, 148.1, 131.4 (q, $J_{CF} = 27.5$ Hz), 123.6 (q, $J_{CF} = 225.0$ Hz), 121.85, 115.0, 42.6, 26.9, 21.9; HRMS (ESI) calcd for C$_{12}$H$_{10}$F$_6$N$_2$S 329.0542, found 329.0549 (M + H);

**Synthesis of Catalysts 10c and 10d**

Catalysts 10c and 10d are known compounds and were synthesized according to known procedures. Spectral data below is included for completeness.

**2,3,6,7-tetrahydro-5H-thiazolo[3,2-a]pyrimidine (10c)**

\[
\begin{align*}
\delta_H & (\text{CD}_3\text{CN}, 600 \text{ MHz}) 3.48 (2H, J = 7.2 \text{ Hz}, \text{t}), 3.28 (1H, J = 5.4 \text{ Hz}, \text{t}), 3.22 (2H, J = 6.0 \text{ Hz}, \text{t}), 3.10 (2H, J = 6.6 \text{ Hz}, \text{t}), 1.80 (2H, \text{m}); \\
\delta_C & (\text{CD}_3\text{CN}, 125 \text{ MHz}) 160.0, 54.3, 45.5, 45.1, 26.4, 21.2.
\end{align*}
\]


\[
\begin{align*}
\end{align*}
\]
δ_H (CD_3CN, 600 MHz) 7.34 (1H, d), 7.23 (1H, td, J = 9.6, 0.6 Hz, 1H), 6.99 (1H, td, J = 9.0, 0.6 Hz, 1H), 6.84 (1H, J = 9.6 Hz), 3.78 (2H, t), 3.46 (2H, t), 1.97 (2H, m); δ_C (CD_3CN, 125 MHz) 157.3, 141.4, 126.1, 122.5, 121.9, 121.7, 107.5, 45.7, 42.3, 20.0.

**Synthesis of Catalyst 12a**

LiAlH_4 (106 mg, 2.80 mmol) was added to an oven-dried round bottom flask and suspended in dry Et_2O. Catalyst 12b (225 mg, 1.12 mmol) was added to the slurry and the reaction mixture was refluxed for 20 h under N_2. The reaction mixture was then cooled and placed on ice. With stirring, 0.11 mL of H_2O was added slowly to the mixture followed by 0.22 mL of 10% NaOH and then 0.33 mL of H_2O to quench excess reducing agent. The mixture was allowed to stir for ten minutes and then vacuum filtered. The filter cake was added to 25 mL of stirring chloroform and heated to 40°C. The reaction mixture was allowed to stir for 10 minutes to dissolve the desired product. Following a second vacuum filtration, the filtrate was collected and concentrated to yield the desired product.

1,3,4,6-tetrahydro-2H-pyrimido[2,1-b]quinazoline (12a)

Yield 70%. δ_H (CD_3CN, 400 MHz) 6.97 (1H, J = 7.2 Hz, t), 6.83 (1H, J = 7.2 Hz, d), 6.67 (1H, J = 7.2 Hz, t), 6.57 (2H, J = 7.6 Hz, d), 4.24 (2H, s), 3.21 (1H, J = 5.6, t), 3.11 (1H, J = 6.0, t), 1.97 (2H, m); HRMS (ESI) calcd for C_{11}H_{12}N_3O 202.0976, found 202.0975 (M + H);
The C₄, C₆, and 3-oxo-C₁₂ AHLs are known compounds and were synthesized following previously published procedures.⁴⁰–⁴²

**N-Hexanoyl-L-homoserine Lactone (4)**

\[
\text{\includegraphics[width=0.2\textwidth]{image}}
\]

63.8% Yield; \(\delta_H (\text{DMSO-d}_6, 500 \text{ MHz})\) 8.30 (1H, s), 4.49 (1H, m), 4.31 (1H, td, \(J_I = 9.0 \text{ Hz}, J_2 = 1.5 \text{ Hz}\)), 4.17 (1H, m), 2.35 (1H, m), 2.13–2.06 (3H, m), 1.47 (2H, m, \(J = 7.5 \text{ Hz}\)), 1.26–1.21 (4H, m), 0.84 (3H, t, \(J = 7.0 \text{ Hz}\)); \(\delta_C (\text{DMSO-d}_6, 125 \text{ MHz})\) 175.9, 172.7, 65.7, 48.3, 35.5, 31.2, 28.7, 25.2, 22.3, 14.3; MS (ESI) calcd for C₁₀H₁₇NO₃: 200.13, found: 200.12 (M + H).

**N-Butyryl-L-homoserine Lactone (2)**

\[
\text{\includegraphics[width=0.2\textwidth]{image}}
\]

56.1% Yield; \(\delta_H (\text{DMSO-d}_6, 500 \text{ MHz})\) 8.30 (1H, s), 4.51 (1H, m), 4.32 (1H, td, \(J_I = 8.5 \text{ Hz}, J_2 = 2.0 \text{ Hz}\)), 4.18 (1H, m), 2.36 (1H, m), 2.14–2.05 (3H, m), 1.59 (2H, m, \(J = 7.5 \text{ Hz}\)), 0.84 (3H, t, \(J = 7.5 \text{ Hz}\)); \(\delta_C (\text{DMSO-d}_6, 150 \text{ MHz})\) 175.3, 172.0, 65.1, 47.7, 36.9, 28.2, 18.5, 13.4; MS (ESI) calcd for C₈H₁₅NO₃: 172.10, found: 172.06 (M + H).

**N-(3-Oxododecanoyl)-L-homoserine Lactone (1)**

\[
\text{\includegraphics[width=0.2\textwidth]{image}}
\]
45.2% Yield; $\delta_H$ (CDCl$_3$, 600 MHz) 7.75 (1H, d, $J = 6.6$ Hz), 4.62 (1H, m), 4.48 (1H, td, $J_1 = 9.0$ Hz, $J_2 = 1.2$Hz), 4.29 (1H, m), 3.48 (2H, s), 2.73 (1H, m), 2.54 (2H, t, $J = 7.2$ Hz), 2.27 (1H, m), 1.58 (2H, p, $J = 7.2$ Hz), 1.31-1.26 (12H, m), 0.88 (3H, t, $J = 6.6$ Hz); $\delta_C$ (CDCl$_3$, 150 MHz) 206.5, 175.0, 166.6, 65.9, 49.0, 48.3, 43.8, 31.8, 29.6, 29.4, 29.3, 29.2, 29.0, 23.4, 22.7, 14.1; MS (ESI) calcd for C$_{16}$H$_{27}$NO$_4$: 298.20, found: 298.18 (M + H).

ii. Catalyst Screening by $^1$H NMR

*General Procedure for Catalytic Experiments by $^1$H NMR*

In a 4-mL scintillation vial, 4 (10.0 mg, 0.0502 mmol) and the thiourea/guanidinium catalyst being screened (0.00502 mmol) were combined. The catalyst and substrate were then dissolved in 1 mL of the deuterated solvent of choice. As an internal standard, 1,4-difluorobenzene (5.16 μL, 0.00502 mmol) was added to the reaction mixture. Piperidine (99.0 μL, 0.1004 mmol) was transferred into the vial and the reaction mixture was vortexed and quickly transferred to an NMR tube for analysis. $^1$H NMR experiments were acquired in a sequence with a spectra being acquired between every 376 – 669 s depending on the experiment. The integration of the α-proton resonance of the lactone was monitored over time to track the conversion of starting material to product.

(S)-N-[(4-hydroxy-1-oxo-1-(piperidin-1-yl)butan-2-yl)hexanamide (5)

![Chemical Structure](image)

$\delta_H$ (CD$_3$CN-d$_3$, 500 MHz) 6.86 (1H, s), 4.92 (1H, m), 3.58 (1H, dd, $J_1 = 2.5$ Hz, $J_2 = 5.0$ Hz), 3.53-3.41 (6H, m), 2.21-2.17 (3H, m), 1.81 (1H, m), 1.62 (2H, m), 1.56 (4H, m), 1.50-1.45
(3H, m), 1.32-1.26 (4H, m), 0.88 (3H, t, \( J = 7.0 \) Hz); \( \delta \text{C} \) (CD\(_3\)CN-d\(_3\), 125 MHz) 173.8, 170.0, 57.8, 46.4, 46.2, 43.0, 36.2, 36.0, 31.5, 26.4, 25.7, 25.6, 24.5, 22.5, 13.6; MS (ESI) calcd for C\(_{15}\)H\(_{28}\)N\(_2\)O\(_3\): 285.22, found: 285.20 (M + H); Anal. Calc. for C\(_{15}\)H\(_{28}\)N\(_2\)O\(_3\): C, 63.35; H, 9.92; N, 9.85; found: C, 63.34; H, 9.76; N, 9.80.
REFERENCES


12. Zuend, S. J. & Jacobsen, E. N. Mechanism of Amido-Thiourea Catalyzed Enantioselective Imine Hydrocyanation: Transition State Stabilization via Multiple Non-


CHAPTER III: HYDRAZINE MEDIATED TRANSAMIDATION OF 3-OXO-N-ACYL HOMOSERINE LACTONES

A. Introduction

i. Amines as Nucleophiles in Water

Reactions that employ amines as nucleophiles in water are central to all life processes. Enzyme-mediated amine additions, namely through acyl transfer reactions, are pervasive in cellular and molecular biology. Amide bond formation, the key linkage in protein biosynthesis, is achieved by the addition of the $N$-terminus of a tRNA-tethered amino acid via acyl transfer to a growing polypeptide chain.\textsuperscript{1} Posttranslational modifications to histone lysine residues are mediated by a series of transferases that catalyze acyl transfer reactions of the amine side chain with their respective coenzymes. These modifications, such as lysine methylation, acetylation, and ubiquitination, have been shown to be critical for DNA processing and gene regulation while also being linked to a variety of human diseases.\textsuperscript{2,3}
Figure 3.1. Comparison of lysine pKa values in solution and in the active site of histone lysine monomethyltransferase SET7/9. The presence of the sulfonium ion has been computationally demonstrated to effectively reduce lysine's pKa from 10.8 to 8.3. Participation of local amino acids in hydrogen bonding lysine resultantly increases its nucleophilicity and enables transfer of the methyl group to lysine from S-adenosylmethionine (SAM).\textsuperscript{3,4}

Harnessing the nucleophilicity of amines for acyl transfer under physiological conditions in a non-enzymatic environment is not trivial. Part of the difficulty associated with achieving these transformations synthetically lies in the interdependence of nucleophilicity and basicity. In general, amines with higher pKa values are more nucleophilic. However, the basicity of the amines renders them almost fully protonated when well-solvated at physiological pH. The effective concentration of the neutral amine in solution resultantly decreases, attenuating its capacity to serve as a nucleophile.\textsuperscript{5,6} In an enzyme, nucleophilicity is modulated by both the hydrophobicity of the catalytic site and the presence of ordered hydrogen bond networks, often forwarded by suppression in amine pKa.\textsuperscript{7} (Figure 3.1). Thus, amine nucleophilicity in water is highly dependent on the amine selected and local environment in which the reaction proceeds.
Despite this setback, synthetic chemists have developed methodologies to elicit amine nucleophilicity for organic transformations in water. Intramolecular additions have proven to be an effective strategy for promoting the reaction of amine nucleophiles despite the extant equilibrium favoring the corresponding ammonium species. Installing such an element of preorganization increases the effective concentration of the neutral amine at the electrophilic site, thus decreasing entropic barriers to reaction. Such principles have been implemented to advance synthesis of peptide sequences beyond fifty residues through the process of native chemical ligation (NCL) (Figure 3.2). Two peptide fragments, one containing a C-terminal thioester fragment (13) and the other containing an N-terminal cysteine (14), react to form the corresponding thioester (15). The formation of the thioester is rapid and reversible at neutral pH. However, the pendant α-amino group of the cysteine is positioned to undergo an intramolecular $S\rightarrow N$ acyl transfer through a favorable five membered transition state. The
result is the ligation product (16) of the two shorter peptide fragments through the irreversible formation of a native peptide bond. NCL has been effectively employed to perform ligations on the time scale of hours at micromolar concentrations, with the amine of the N-terminal cysteine greater than 90% protonated at physiological pH.\(^9\) Similar modes of exploiting amine nucleophilicity in water are present in the mechanisms of both expressed protein ligation and intein chemistry.\(^{10,11}\)

The nucleophilicity of an amine in water is also altered by its neighboring atom. Influence of the \(\alpha\)-effect, or the presence of an adjacent heteroatom containing at least one pair of non-bonded electrons to the nucleophilic site, enhances nucleophilicity without a significantly modifying basicity.\(^{12,13}\) The adjacent lone pair of electrons destabilizes the ground state of the amine through electron repulsion, increasing its general reactivity.\(^{14}\) The presence of the electron withdrawing heteroatom leads to greater desolvation of the unbonded electrons on nitrogen, enhancing its propensity for addition to an electrophile. Even when partially protonated at physiological pH, these amines still retain a high nucleophilicity parameter. Hydroxylamine and hydrazine are two such examples of amines influenced by the \(\alpha\)-effect.\(^6\)

### ii. Applications of Hydrazine as a Nucleophile in Medicinal Chemistry and Chemical Biology

The enhanced nucleophilicity of hydrazine and hydroxylamine has become a valuable tool for both medicinal chemistry and chemical biology. Both amines react with aldehydes and ketones to form the corresponding hydrazone and oxime in water (Scheme 3.1). As aldehydes and ketones are absent from cell surfaces, hydrazone and oxime formation is an example of a bioorthogonal reaction or an organic transformation that proceeds without
interfering with biological processes. Consequently, the chemoselective condensation can be employed in vitro and in vivo to selectively conjugate biomolecules, label cellular targets, and engineer immunotherapies.\textsuperscript{15–18}

**Scheme 3.1.** Oxime and hydrazone formation in the presence of an aldehyde or ketone.

In a seminal work incorporating hydrazone formation into a biological setting for imaging, Heitzmann & Richards employed hydrazine functionalized biotin molecules as conjugates to membrane-bound glycoproteins for visualization of the cell membranes by electron microscopy.\textsuperscript{19} More recently, hydrazinonicotinic acid has been employed for labeling cancer cells. Two notable examples include use of the hydrazine to conjugate RGD peptide derivatives, known for selectively binding $\alpha_\beta$-integrins on cancerous cells, to $^{18}\text{F}$ radiolabels and $^{99m}\text{Tc}$ to VEGF, an overexpressed growth factor in tumors (Figure 3.3a).\textsuperscript{20,21} Hydrazone formation for the incorporation of fluorescent tags has been similarly employed for visualizing biochemical pathways and binding events in both prokaryotic and eukaryotic cellular environments.\textsuperscript{16,22–24}
Figure 3.3. Select applications of hydrazone formation in medicinal chemistry and chemical biology. a) A radiolabeled RGDyK peptide conjugate is prepared by hydrazone formation (blue) between hydrazinonicotinic acid and an $^{18}$F-functionalized benzaldehyde for in vivo tumor imaging.\textsuperscript{20} b) Ricin toxin A is selectively conjugated to Jurkat cells presenting a ketone-functionalized ManLev derivative through hydrazone formation (blue), resulting in cell death.\textsuperscript{25}

The chemoselectivity of the hydrazone ligation for biomolecular conjugation also allows for the engineered modification of cellular function (Figure 3.3b). This was demonstrated by Bertozzi and colleagues through use of a biotinylated hydrazide for recognition of manufactured ketone-containing N-Levulinoylmannosamine (ManLev) derivatives on T-lymphocytes. An avidin-ricin toxin conjugate was then introduced to bind
biotin for direct delivery of the toxin to the cell surface, leading to cell death in the sugar presenting lymphocytes. These results demonstrated that use of hydrazine derived conjugates can enable new strategies in drug delivery and stimulating immunological recognition and response.\textsuperscript{25}

\textbf{Scheme 3.2.} Concerted mechanism of hydrazone formation. Protonation of the tetrahedral intermediate (17) under acidic conditions enhances leaving group ability (\textsuperscript{`}OH vs. H\textsubscript{2}O) and thus the rate of hydrazone formation (18).

While the application of the reactions is broad, oxime and hydrazone formation have their innate limitations. Considering the mechanism of the reaction in conditions where nucleophilic attack at the carbonyl carbon is fast, the rate-determining step is protonation of the tetrahedral intermediate 17 to eliminate water and form the corresponding C=N condensation product 18 (Scheme 3.2). Thus, both oxime and hydrazone formation are accelerated under acidic conditions. This impedes the rates of the condensations at physiological pH, requiring hours or even days to reach maximum conversion. As a result, a large excess of either the amine or carbonyl are generally required at millimolar concentrations for achieving practical rates of bioconjugation.\textsuperscript{26} A study conducted by Dirksen and Dawson compared the rates of both ligation reactions. Hydrazone formation was shown to proceed at rates 20-fold faster than oxime formation, supporting the use of hydrazines to reduce reaction times.\textsuperscript{27} Further, phenylhydrazines were demonstrated to undergo hydrazone formation more rapidly than aliphatic hydrazine derivatives.\textsuperscript{28,29}

The poor kinetics of hydrazone and oxime formation were greatly improved when the amenability of both reactions to nucleophilic catalysis by aniline was introduced by Dirksen.
and Dawson in 2006. Peptide ligation rates were shown to improve more than 400 times that of the uncatalyzed reaction at physiological pH.\textsuperscript{26,30} Since then, successive generations of aniline-based catalysts, including anthranilic acids, phosphonates, and phenylenediamines have been introduced to improve reaction rates, catalyst loadings, and catalyst solubility.\textsuperscript{31–33}

In most cases, superstoichimetric quantities of the catalyst are still necessary to afford significant increases in ligation rates. Crisalli and Kool offered an exception by optimizing the nucleophilic and electrophilic partners for hydrazone formation to eliminate the need of a nucleophilic catalyst. Through the use 2-carboxylphenylhydrazine (19) and quinoline-8-carboxaldehyde (20), greater than 90\% hydrazone formation was achieved in under 2h at pH 7.4 (Scheme 3.3). These increased rates are attributed to both optimized electronics, specifically an electron poor aldehyde, and the presence of acidic and basic functionality to catalyze the breakdown of the formed tetrahedral intermediate.\textsuperscript{28} While these rates are highly compatible for applications in chemical biology, the substrate specificities currently limit broad employment of the reactivity.

\begin{center}
\begin{tikzpicture}
  \node (a) at (0,0) {\includegraphics[width=0.8\textwidth]{hydrazone}};
  \node at (1.5, -1.5) {\textbf{Scheme 3.3.} Rapid hydrazone formation in the absence of aniline derivatives. The use of pyridyl and carboxyl groups intramolecularly catalyzes conversion from the tetrahedral intermediate to the hydrazone at rates greater than other common bioorthogonal reactions, such as strain-promoted azide-alkyne cycloadditions.\textsuperscript{28}}
\end{tikzpicture}
\end{center}

Though great progress has been made in achieving rate enhancement, one underlying drawback of hydrazone and oxime formation is the reversibility of the ligation. Both
Hydrazones and oximes have been experimentally proven to be more stable than simple imines, but are still prone to hydrolysis. The resultant ligation products have half-lives that vary from days to hours, making the chemistry less desirable when irreversible conjugations are required. In the following chapter, a hydrazine-mediate method for the selective degradation of 3-oxo-AHLs is described. The irreversibility of the transformation and its ability to proceed in both acidic and neutral media has the potential to circumvent current drawbacks of hydrazone ligation while derivatizing a critical quorum sensing target.

**B. Results and Discussion**

**i. Aminolysis of the AHL under Aqueous Conditions**

In Chapter I, the ability to amidate the γ-butyrolactone of the AHL was demonstrated in organic solvents. To progress toward derivatization of the AHL in conditions more relevant to the natural environment of the autoinducer, we sought to conduct the ring opening reaction under aqueous conditions. However, the nucleophile employed during our catalytic experiments, piperidine, has a pKa of 11.2. This renders the nucleophile inactive at physiological pH as it lies almost exclusively in its protonated form. As a result, the control reaction of piperidine and the C$_6$-AHL presented in Chapter I failed to proceed with identical amine and substrate concentrations in pH 7 buffer.

To identify amines capable of rapid aminolysis under aqueous conditions, a competitive screen of potential nucleophiles against the AHL was prepared. Further, the screen would enable the aminolysis rates to be compared to the background rate of lactone hydrolysis. The selection of amines for the screen was focused to those with a pKa at or below that of piperidine (Table 3.1). Understanding the varied nucleophilicity of each of the amines, we hypothesized that the content of neutral amine present in solution would largely
dictate reactivity with the AHL.\textsuperscript{36,37} A cocktail of amines was prepared by combining six amines in an equimolar ratio in PBS buffer. The pH of the solution was adjusted with HCl, and then combined with the autoinducer. The reaction progress was monitored by LC-MS, allowing for separation, identification, and relative quantification of the formed aminolysis products.

**Table 3.1.** Product masses in C\textsubscript{6}-AHL aminolysis competitive screen.

<table>
<thead>
<tr>
<th>Amine</th>
<th>Observed [M+H]\textsuperscript{a}</th>
<th>pKa\textsuperscript{b}</th>
<th>Amine</th>
<th>Observed [M+H]\textsuperscript{a}</th>
<th>pKa\textsuperscript{b}</th>
</tr>
</thead>
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<tr>
<td>\textsuperscript{+}HN-NH\textsubscript{2}</td>
<td>232.1</td>
<td>8.1</td>
<td>\textsuperscript{+}NH\textsubscript{3}</td>
<td>-</td>
<td>10.7</td>
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<tr>
<td>\textsuperscript{-}HO-NH\textsubscript{2}</td>
<td>233.1</td>
<td>6.0</td>
<td>\textsuperscript{-}NH\textsubscript{2}</td>
<td>-</td>
<td>4.6</td>
</tr>
<tr>
<td>\textsuperscript{+}O-NH\textsubscript{2}</td>
<td>-</td>
<td>8.4</td>
<td>\textsuperscript{+}NH\textsubscript{2}</td>
<td>-</td>
<td>5.0, 9.7</td>
</tr>
<tr>
<td>\textsuperscript{-}NH\textsubscript{2}</td>
<td>-</td>
<td>11.2</td>
<td>\textsuperscript{+}H\textsubscript{2}N-NH\textsubscript{3}</td>
<td>260.1</td>
<td>6.9, 10.0</td>
</tr>
<tr>
<td>\textsuperscript{+}NH\textsubscript{2}</td>
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<td>11.2</td>
<td>\textsuperscript{-}NH\textsubscript{3}</td>
<td>-</td>
<td>7.9</td>
</tr>
<tr>
<td>\textsuperscript{-}NH\textsubscript{3}</td>
<td>-</td>
<td>11.2</td>
<td>\textsuperscript{+}H\textsubscript{3}N\textsuperscript{-}O\textsuperscript{-}</td>
<td>-</td>
<td>10.8</td>
</tr>
<tr>
<td>AHL hydrolysis (21)</td>
<td>218.2</td>
<td>-</td>
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<td></td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{a}reference to C\textsubscript{6}-AHL (5) observed mass of 200.1 m/z [M+H]; \textsuperscript{b}pKa’s obtained from reported values

A series of aminolysis products were detected by mass spectrometry, proving our desired reactivity. Upon consumption of the AHL, products corresponding to the addition of hydrazine, hydroxylamine, and ethylenediamine to the AHL were observed, with the product corresponding to the addition of hydrazine having the greatest LC yield. Each of these amines has a pKa of 8.12 or lower, displaying the impact of protonation state on the amine’s participation in the acyl transfer reaction. The effectiveness of hydrazine and hydroxylamine
can be attributed to enhanced nucleophilicity by the α-effect (see Introduction). At 20 eq. of hydrazine, formation of the aminolysis product outcompeted background hydrolysis of the AHL (21). This provides promise for using hydrazine to achieve more rapid degradation of the autoinducer than the naturally imposed hydrolysis of the signaling molecule during its extracellular transport.

Hydrazine, hydroxylamine, and ethylenediamine were all screened independently to validate relative rates of conversion in the absence of the other amines contained in the initial cocktail. Each proceeded with similar relative rates of conversion as observed in the competitive screen, disproving any contribution by synergistic effects. As the competitive screen was designed simply to identify the most effective amines for AHL aminolysis, assessment of nucleophilicity beyond the three most readily detected was not pursued.

**ii. Hydrazine-mediated Transamidation of 3-oxo-AHLs**

As hydrazine proved to be the standout nucleophile from our aqueous competitive screen, we sought to test it with the 3-oxo variant of the AHL. As displayed in the competitive screens with the C₆-AHL, both aminolysis and hydrolysis of the lactone of the autoinducer were expected to occur. However, the additional carbonyl functionality in this AHL subfamily serves as a secondary site for nucleophilic attack to form the corresponding hydrazone by condensation. Hydrazone formation would also uniquely position the α-amine proximal to the N-acyl moiety of the AHL and we hypothesized that an additional lytic pathway could be achieved in cleaving the amide bond (Scheme 3.4).
Scheme 3.4. Proposed route to 3-oxo-AHL amide cleavage through hydrazone formation and subsequent transamidation.

With multiple pathways of degradation identified, the 3-oxo-C₈-AHL (22) was synthesized as model substrate for screening due to its miscibility in water. The autoinducer was combined with 10 eq. of hydrazine buffered at pH 7 to determine the distribution of products by LC-MS. Upon consumption of the AHL, the major product generated had a mass of 155.1 m/z [M+H], indicating cleavage of the parent AHL structure (242.1 m/z [M+H]). The truncated mass was identified to be hydroxypyrazole 23. The total mass chromatogram contained only trace amounts of both the aminolysis and hydrolysis product relative to the cleavage product, divulging that the observed transformation is both a novel and dominant pathway to reduce the active form of the autoinducer in situ.

Further modification of reaction conditions by decreasing the amine equivalents demonstrated that the hydroxypyrazole can be generated as the major product with only stoichiometric quantities of hydrazine. This is in contrast to the 20 eq. of hydrazine required for lactone aminolysis to out compete rates of background hydrolysis. Decreasing the pH of the reaction media to 4.5 to mitigate hydrolysis of the lactone allowed for exclusive formation of the hydroxypyrazole product (Figure 3.4). The reaction was replicated on a larger scale to allow for isolation and characterization of the hydroxypyrazole by ¹H and ¹³C NMR.
Figure 3.4. Total ion chromatogram of hydrazine-mediated AHL transamidation. The trace displays clean conversion of the 3-oxo-C₈ AHL (22) (1.0 mM) to the corresponding hydroxypyrrole (23) product at pH 4.5 in NaOAc buffer.

Recalling our initial hypothesis, condensation of hydrazine onto the 3-oxo-position of the AHL followed by transamidation through intramolecular cyclization presents a viable pathway for the generation of the 5-hydroxypyrrole product (Scheme 3.4). The predicted 5-membered cyclic transition state for nucleophilic attack is reminiscent of native chemical ligation, an already well-established method for amide bond formation under aqueous conditions. The aromaticity of the resultant pyrazole product likely provides an additional thermodynamic driving force to enable the observed AHL cleavage. The hydrazone (256.2 m/z [M+H]), which is the putative intermediate in the transformation, was only observed in trace amounts by LC-MS during the course of the reaction. The hydrazone was found to build up more significantly when substituted hydrazines (Section iv) and hydrazine-containing peptides were employed (Chapter IV).
iii. Reaction Selectivity and pH Dependence

As described in Chapter 1, the communication networks in gram-negative bacteria that are regulated by quorum sensing often employ identical N-acyl homoserine lactones across different species. In addition, multiple AHLs are often responsible for the regulation of gene expression in an individual species (Figure 1.3). A reaction that selectively modifies a single class of AHLs could give insight into the importance of a singular autoinducer in influencing a colony phenotype or permit selective inhibition of quorum-based communication in a multi-species environment. To determine the ability of hydrazine to selectively react with 3-oxo-AHLs in the presence of other AHLs, a competition experiment containing a representative member of each AHL structural subfamily was conducted. The C₆, 3-OH-C₈, and 3-oxo-C₈-AHLs were combined and subjected to a single equivalent of hydrazine at pH 4.5 (Figure 3.5). The reaction was monitored by HPLC and unambiguously demonstrated that the 3-oxo-C₈-AHL was the only species consumed amongst the other autoinducers. Accordingly, the resultant 5-hydroxypyrazole transamidation product was the only species to be amplified in the trace over time. This selective depletion of the 3-oxo-AHL supports the importance of hydrazone formation in the transamidation mechanism as the non-keto functionalized AHLs proved unreactive. The exclusive generation of the hydroxypyrazole also reinforces the kinetic dominance of the condensation-cyclization pathway compared to AHL modification by lactone aminolysis.
Figure 3.5. AHL selectivity in hydrazine-mediated transamidation. The 3-oxo-C₈-AHL (22) is cleaved in the presence of the 3-OH-C₈-AHL (24) and the C₆-AHL (4) to form hydroxypyrazole (23). The overlaid green, red, and blue lines on the HPLC trace represent injections made at 0 h, 14 h, and 44 h, respectively.

To investigate the impact of pH on the rate of transamidation, the 3-oxo-C₈ AHL was screened in a series of NaOAc and PBS buffers of variable pH. Four reactions were conducted from pH 4.5 to 7.4 with 5 eq. of hydrazine and monitored by HPLC. The autoinducer was converted to hydroxypyrmaole most efficiently at pH 7.4, with a relative rate of AHL consumption that was almost five times the rate at pH 4.5 (Figure 3.6). Importantly, the background rates of AHL hydrolysis also increase with increased pH and significant hydrolysis product was observed at pH 7.4. Hence, we determined pH 6.5 to be an optimal pH for further reaction development.
Figure 3.6. pH dependence on 3-oxo-C₈ AHL rates of consumption with hydrazine. Reaction progress was monitored by analytical HPLC at 210 nm with a 5:1 ratio of hydrazine to AHL.

By the known mechanism of hydrazone formation, the rate-determining step of the reaction is protonation of the tetrahedral intermediate. Consequently, hydrazone formation is known to be acid catalyzed (Figure 3.6). The reversal of the anticipated pH dependence on reaction rate distinguishes the contribution of both the condensation and cyclization to the kinetics of the reaction. Hydrazone formation slows as the solution becomes more basic, decreasing k₁ (Scheme 3.4). However, the effective concentration of the neutral α-amine of the hydrazone is greater at increased pH, enhancing its nucleophilicity. This increases the contribution of k₂ to the overall reaction, reflected in the increased rate of pyrazole formation at higher pH. Thus, it is reasonable that intramolecular attack of the amino imine onto the amide to yield the hydroxypyrazole is rate-determining in AHL transamidation.

iv. Reaction Scope

After characterizing the selectivity and pH dependence of the hydrazine-mediated 3-oxo-AHL transamidation reaction, a series of commercially available hydrazines were
screened to probe substituent effects on rate and reactivity. The selected hydrazines (Table 3.2) were tested individually in 5-fold excess in the presence of the 3-oxo-C₈ AHL at pH 6.5.

**Table 3.2.** Transamidation of the 3-oxo-C₈ AHL in the presence of hydrazine derivatives.

![Chemical reaction diagram](image)

<table>
<thead>
<tr>
<th>Entry</th>
<th>R</th>
<th>% Conversion at 30 hᵃ</th>
<th>Entry</th>
<th>R</th>
<th>% Conversion at 30 hᵃ</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>AHL hydrolysis</td>
<td>28.7</td>
<td>5</td>
<td>CH₃</td>
<td>60.2</td>
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<tr>
<td>2</td>
<td>O</td>
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<td>HO-CH₂</td>
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</tr>
<tr>
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<td>O</td>
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<td>7</td>
<td>H</td>
<td>96.7</td>
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<tr>
<td>4</td>
<td>O</td>
<td>43.0</td>
<td>8</td>
<td>O-C₆H₄</td>
<td>100</td>
</tr>
</tbody>
</table>

ᵃPercent conversion determined by monitoring the integration of 22 over time on an analytical HPLC at 210nm

![Conversion of 3-oxo-C₈ AHL with Hydrazine Derivatives](image)

**Figure 3.7.** Percent conversion of the 3-oxo-C₈ AHL (22) in the presence of hydrazine derivatives. The numbers displayed in the legend correspond to the entries listed in Table 3.2.
The nature of the hydrazine employed had a significant impact on rate and the relative content of hydrazone and hydroxypyrazole product. Alkylated derivatives of hydrazine (Table 3.2, Entries 4-6) were noticeably less reactive with the AHL, but still possess the capability to cleave at the N-acyl moiety as the corresponding pyrazole fragment masses were detected by LC-MS. Rates also decreased with increased substituent size and pKa, factors that would expectedly attenuate amine nucleophilicity under aqueous conditions. Hydrazides (Table 3.2, Entries 2-3), which are common functionalities chosen for hydrazone formation in bioorthogonal chemistry, proved essentially unreactive with the autoinducers, yielding only trace amounts of the corresponding hydrazone. Failure of the hydrazides to cyclize can be attributed to the decreased nucleophilicity of the α-amine when in direct resonance with a carbonyl. Delocalization of the non-bonded α-amine electrons in stabilized amide resonance forms decreases the propensity for addition to the AHL.

By comparison, phenylhydrazine (Table 3.2, Entry 8) proved to be the most efficient hydrazine in derivatizing the AHL, reaching 82% conversion after 2.5 h. The rapid depletion of the AHL was driven by formation of the corresponding hydrazone, the dominant product detected by LC-MS. From 2.5 to 30 h, the concentration of the AHL steadily declined as the hydrazone proceeded to cyclize to the N-phenylhydroxypyrazole. Complete conversion of the AHL was obtained, proving that phenylhydrazine is a competent nucleophile for transamidation. Thus, the known increased ligation rates provided by phenylhydrazine translated to 3-oxo-AHL derivatization, while obviating the susceptibility of resultant hydrazone to hydrolysis by coupling its formation with an irreversible amide cleavage.\(^\text{34}\)
C. Conclusions

To uncover new chemical strategies for N-acyl homoserine lactone functionalization under physiological conditions, the susceptibility of AHLs towards amine additions was probed in aqueous solutions. Aminolysis of the γ-butyrolactone was achieved by the employment of α-effect nucleophiles, such as hydroxylamine or hydrazine. With the proper excess of amine, the amidated product proved to be more kinetically favorable than AHL hydrolysis.

By strategically exploiting their unique β-ketoamide moiety, the first example of a non-enzymatic amide cleavage of 3-oxo-AHLs in aqueous media using hydrazine and hydrazine derivatives was reported. Through detection of transient hydrazone masses by LC-MS, we hypothesize that reversible association of the hydrazine to the AHL enables irreversible, intramolecular acyl transfer to form the observed hydroxylpyrazole product. The low activation energy of the 5-membered transition state for hydrazone cyclization coupled with the thermodynamic driving force provided by the aromaticity of the pyrazole product are each important in overcoming the high barrier for amide cleavage in water.

Hydrazine-mediated transamidation presents a new platform for the study and manipulation of quorum sensing pathways that contain 3-oxo-AHLs, such as in *P. aeruginosa*. The irreversible transformation proceeds most readily at physiological pH. Moreover, the selectivity of hydrazine for the 3-oxo-AHL was demonstrated in the presence of other known AHL substructures. This provides precedence for the use of hydrazine derivatives for the functionalization of the autoinducer in a multispecies environment and in quorum sensing circuits that employ multiple AHLs. As rates of transamidation are improved, the methodology could be an alternative route to quorum quenching by reducing
the extracellular concentration of the signaling molecule in situ. Our strategy to optimize the 3-oxo-AHL transamidation reaction through the generation of hydrazine-containing peptide libraries is discussed in Chapter IV.

D. Experimental

i. General Reagents and Synthesis

Commercial reagents and solvents, including hydrazine derivatives, were purchased from Sigma-Aldrich and Fisher Scientific. Hydrazine competition experiments were monitored on an Agilent 1200 Series LC-MS using a 2.7 μM HALO C18 column. Reaction kinetics were monitored on an HP 1100 Series analytical HPLC/UV-Vis using a 2.7 μM HALO C18 column at 210 nm. Hydroxypyrazole 23 was purified using a Waters semi-prep HPLC/UV-Vis with a Waters 4.2 μM C18 column. Product characterization (1H and 13C) was completed on a 500 or 600 MHz Bruker NMR spectrometer. All related coupling constants are reported in Hertz and shifts in ppm. NMR solvents were purchased from Cambridge Isotopes Laboratories. High resolution mass spectra were obtained on an Agilent Accurate LC-TOF Mass Spectrometer (Agilent Series 6220) operating in positive ion mode with an electrospray ionization source (fragmentor = 175 V). The data was analyzed using an Agilent MassHunter Workstation Software, Qualitative Analysis (V. B.02.00).

General Procedure for the Characterization of Hydroxypyrazole Products

To a 20-mL scintillation vial, 22 (12.1 mg, 0.0501 mmol) was added to 10.0 mL of 200 mM PBS buffer at pH 7.4 and vortexed. The hydrazine derivative (0.250 mmol) was then added to the reaction mixture. The reaction was allowed to stir at RT and reaction progress was monitored by LC-MS. Upon complete conversion to the hydroxypyrazole product, the
reaction mixture was directly purified by semi-prep HPLC under neutral conditions.

Separations were performed in a mobile phase of 20 to 50% B over 20 minutes where A = 100% water, 0.1% methanol and B = 100% acetonitrile. Products were then confirmed by HRMS and $^1$H, $^{13}$C NMR where possible. Several of the hydrazines screened did not progress to the corresponding hydroxypyrazole products by LC-MS (Table 3.2, Entries 2-4) and therefore could not be isolated for mass analysis.

**3-pentyl-1H-pyrazol-5-ol**

\[
\text{\begin{picture}(100,100)
\put(28,35){\includegraphics[width=3cm]{pyrazol.png}}
\end{picture}}
\]

$\delta_{\text{H}}$ (DMSO-d$_6$, 600 MHz) 11.2 (1H, s), 9.26 (1H, s), 5.21 (1H, s), 2.41 (2H, t, $J = 6.6$ Hz), 1.52 (2H, p, $J = 7.8$ Hz), 1.29 (4H, m), 0.86 (3H, t $J = 6.6$ Hz); $\delta_{\text{C}}$ (DMSO-d$_6$, 150 MHz) 161.4, 144.7, 88.3, 31.3, 28.8, 26.1, 22.3, 14.4; HRMS (ESI) calcd for C$_8$H$_{14}$N$_2$O: 155.1179, found: 155.1181 (M + H).

**1-methyl-3-pentyl-1H-pyrazol-5-ol**

\[
\text{\begin{picture}(100,100)
\put(28,35){\includegraphics[width=3cm]{pyrazol.png}}
\end{picture}}
\]

$\delta_{\text{H}}$ (DMSO-d$_6$, 600 MHz) 5.36 (1H, s), 3.48 (3H, s), 2.42 (2H, t, $J = 6.0$ Hz), 1.54 (2H, p, $J = 7.2$ Hz) 1.27 (4H, m), 0.86 (3H, t $J = 7.2$ Hz); $\delta_{\text{C}}$ (DMSO-d$_6$, 150 MHz) 150.5, 87.3, 32.6, 31.3, 28.5, 27.6, 22.3, 14.3; HRMS (ESI) calcd for C$_9$H$_{16}$N$_2$O: 169.1335, found: 169.1336 (M + H).

**1-(2-hydroxyethyl)-3-pentyl-1H-pyrazol-5-ol**
δ\textsubscript{H} (DMSO-d\textsubscript{6}, 600 MHz) 5.37 (1H, s), 3.88 (2H, t, \(J = 6.0\) Hz), 3.63 (2H, t, \(J = 6.0\) Hz), 2.43 (2H, t, \(J = 7.8\) Hz), 1.54 (2H, p, \(J = 7.2\) Hz) 1.30 (4H, m), 0.87 (3H, t, \(J = 6.6\) Hz); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 150 MHz) 150.5, 87.2, 59.5, 48.2, 31.3, 28.5, 27.6, 22.3, 14.3; HRMS (ESI) calcd for C\textsubscript{10}H\textsubscript{18}N\textsubscript{2}O\textsubscript{2}: 199.1441, found: 199.1446 (M + H).

3-pentyl-1-phenyl-1H-pyrazol-5-ol*

δ\textsubscript{H} (DMSO-d\textsubscript{6}, 600 MHz) 11.43 (1H, s), 7.71 (2H, d, \(J = 7.8\) Hz), 7.42 (2H, t, \(J = 7.8\) Hz), 7.21 (1H, t, \(J = 7.2\) Hz), 5.38 (1H, s), 2.44 (2H, t, \(J = 7.8\) Hz), 1.58 (2H, m), 1.32 (4H, m), 0.88 (3H, t); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 150 MHz) 162.2, 153.4, 139.5, 129.2, 125.4, 121.0, 87.0, 31.5, 28.8, 25.8, 22.4, 14.4; HRMS (ESI) calcd for C\textsubscript{14}H\textsubscript{18}N\textsubscript{2}O: 231.1492, found: 231.1495 (M + H). *the keto tautomer (minor) was also observed by NMR but only the resonances for the enol are reported.

General Procedure for N-Acyl Homoserine Lactone (AHL) Synthesis

The 3-OH-C\textsubscript{8} AHL and 3-oxo-C\textsubscript{8} AHL are known compounds and were synthesized following previously published procedures.\textsuperscript{39,40}

N-(3-hydroxyoctanoyl)-L-homoserine Lactone (24)
51.3% Yield (racemate); δ\textsubscript{H} (CDCl\textsubscript{3}, 600 MHz) 6.85-6.80 (1H, br d), 4.56 (1H, m), 4.47 (1H, m), 4.28 (1H, m), 4.00 (1H, br s), 3.38-3.31 (1H, br d), 2.76 (1H, m), 2.44 (1H, m), 2.33 (1H, m), 2.22 (1H, m), 1.43 (1H, m), 1.32 (2H, m), 1.29 (5H, m), 0.88 (3H, t); δ\textsubscript{C} (CDCl\textsubscript{3}, 150 MHz) 175.7, 173.0, 68.6, 66.2, 49.1, 42.6, 36.9, 31.9, 30.0, 25.1, 22.6, 14.0; MS (ESI) calcd for C\textsubscript{12}H\textsubscript{21}NO\textsubscript{4}: 244.2, found: 244.2 (M + H).

\textit{N-(3-Oxooctanoyl)-L-homoserine Lactone (22)}

δ\textsubscript{H} (CD\textsubscript{3}CN, 500 MHz) 7.21 (1H, s), 4.48 (1H, m), 4.35 (1H, td, \textit{J} \textsubscript{1} = 7.5 Hz, \textit{J} \textsubscript{2} = 1.5 Hz), 4.20 (1H, m), 3.34 (2H, s), 2.50-2.46 (3H, m), 2.22 (1H, m), 1.52 (2H, p, \textit{J} = 7.0 Hz), 1.31-1.20 (4H, m), 0.87 (3H, t, \textit{J} = 7.0 Hz); δ\textsubscript{C} (DMSO-d\textsubscript{6}, 150 MHz) 210.0, 175.5, 166.7, 65.8, 50.7, 48.6, 42.3, 31.1, 28.7, 23.0, 22.4, 14.3; MS (ESI) calcd for C\textsubscript{12}H\textsubscript{19}NO\textsubscript{4}: 242.1, found: 242.1 (M + H).

\textbf{ii. Competition and Kinetics Experiments}

\textit{Procedure for Amine Competition Experiment}

In a 20-mL scintillation vial, hydrazine (3.16 μL, 100 μmol), hydroxylamine (6.13 μL, 50% v/v in water, 100 μmol), morpholine (3.16 μL, 100 μmol), piperidine (3.16 μL, 100 μmol), pyrrolidine (3.16 μL, 100 μmol), and azetidine (3.16 μL, 100 μmol) were combined and dissolved in 10.0 mL of PBS. While stirring, the pH of the amine cocktail was adjusted to 7.04 by titration with concentrated HCl. The C\textsubscript{6}-AHL (4) (1.00 mg, 5.02 μmol) was then
added to the reaction mixture. The reaction was monitored by LC-MS with automated injection every hour. Separations were performed in a mobile phase of 15% B over 12 minutes where A = 95:5 H₂O:acetonitrile, 0.1% formic acid and B = 0:100 water:acetonitrile, 0.1% formic acid. The presence and relative distribution of aminolysis products was determined by comparing the integration of the total ion chromatogram (TIC) peaks generated for each product. The same procedure was repeated for the other six amines displayed in Table 3.1.

*Procedure for AHL Competition Experiment*

In a 20-mL scintillation vial, 4 (1.99 mg, 10.0 μmol), 24 (2.43 mg, 10.0 μmol), and 22 (2.41 mg, 10.0 μmol) were all combined and dissolved in 10.0 mL of NaOAc buffer at pH 4.5. Hydrazine (0.32 μL, 10. μmol) was then added to the reaction mixture. The reaction progress was monitored by analytical HPLC with automated injection every 40 minutes. Separations were performed in a mobile phase of 25 to 45% B over 25 minutes where A = 95:5 water:acetonitrile and B = 0:100 water:acetonitrile. After 44 h, the products were analyzed by LC-MS.

*General Procedure for Comparative Kinetics (pH Dependence & Hydrazine Derivatives)*

Stock solutions of both the 3-oxo-C₈ AHL (22) (2.0 mM) and the corresponding hydrazine (100 mM) were prepared in distilled water and 400 mM PBS or NaOAc buffer, respectively. The hydrazine was then diluted 10x in buffer to a final concentration of 10 mM. In a 2.0-mL vial, 500 μL of both the AHL and hydrazine stock solutions were combined. The reaction progress was monitored by analytical HPLC with automated injection every 160 minutes.
Separations were performed in a mobile phase of 30 to 50% B for phenylhydrazine and the hydrazide derivatives (Table 3.2, Entries 2, 3, and 8) and a mobile phase of 5 to 40% B over 22 minutes for the remaining hydrazines. Percent conversions were obtained by integration of the peak corresponding to the AHL after each injection with concentrations determined by a prepared standard curve. Products were analyzed by LC-MS.
REFERENCES


CHAPTER IV: DEVELOPMENT OF HYDRAZINE-CONTAINING PEPTIDES FOR 3-OXO-AHL DERIVATIZATION THROUGH COMBINATORIAL METHODS

A. Introduction

i. Synthetic Peptides as Enzyme Mimics

Enzymes are the envy of the synthetic chemist. Their exquisite selectivity and efficiency in carrying out chemical transformations is of the highest standard, with single molecule affinities and rates reaching greater than $10^9$ that of the uncatalyzed reaction.¹ Thus, efforts to recreate enzymatic activity via synthetic mimics have been undertaken to better understand the chemical basis of enzyme function and to harness their reactivity to advance organic transformations and intervene in biological processes.² The diversity of functional enzyme mimics includes small molecule catalysts (Chapter II), macromolecular assemblies,³ and catalytic antibodies.⁴

At the core of enzyme catalysis is the sequence of amino acids that promotes the observed enzyme function. The connectivity and spatial arrangement of the amino acids advantageously presents the corresponding functional groups to mediate substrate binding and modification. The resultant enzyme active site is a coordinated array of covalent and non-covalent interactions, such as hydrogen bonding, ion pairing, acylation, and proton transfer.⁵ Despite the size and complexity of enzymes, only a few of the amino acid residues are active participants in catalysis, inspiring chemists to consider minimalist structures to afford comparable reactivity. Synthesizing peptides, or short sequences of amino acids
ranging from 2 – 50 residues in length, has become a strategy for reconstituting these interactions.  

Scheme 4.1. Kinetic resolution of secondary alcohols by Miller’s octapeptide 25. The rigidity of the β-hairpin of the peptide’s secondary structure along with the diversity of its primary sequence has been demonstrated to lead to the increase acylation rates of the (S,S) enantiomer.

In organic solvents, peptides have been employed as asymmetric catalysts to enhance rates and selectivity of several well-known organic transformations, such as Aldol reactions, Michael additions, and oxidations.  

The use of N-methylhistidine-containing β-hairpin mimics for acyl transfer catalysis was realized by Miller and colleagues, in the kinetic resolution of secondary alcohols (Scheme 4.1). Through the comparison of both tetra- and octapeptide derivatives, increased sequence length improved the rigidity of the catalyst, realized in a 50-fold enhancement in selectivity. This selectivity was attenuated in Pro-AIB and stapled β-hairpin analogs, displaying the importance of the peptide secondary structure and cross-strand hydrogen bonding for the preferential acylation of the (S,S) enantiomer.
Derivatives of the \(N\)-methylhistidine catalysts provided similar enzyme-like substrate-peptide interactions for site-specific acylation reactions and alcohol desymmetrizations.\(^{10-12}\)

![Scheme 4.2](image)

**Scheme 4.2.** Peptide enzyme mimics for retro-aldol additions by lysine-mediated imine formation. In a study conducted by Tanaka *et al.*, the displayed poly-lysine sequence provided the greatest aldolase activity.\(^{13}\)

The non-covalent interactions critical to peptide folding are amplified when in organic solvents, providing an advantage in decreasing peptide length while still observing appreciable rate accelerations and product selectivity. Employing synthetic peptides in aqueous media often requires larger or uniquely well-folded peptide assemblies due to the presence of competitive solvent interactions.\(^{14}\) However, several efficient constructs have still proven quite successful in replicating nucleophilic enzyme-like pathways. Baltzer and coworkers designed a helix-loop-helix dimer capable of accelerated p-nitrophenyl ester hydrolysis up to three orders of magnitude greater than the non-peptide acyl transfer catalyst \(N\)-methylimidazole. The reactivity of the dimer was attributed to a cooperative general acid and nucleophilic catalysis by two neighboring histidine residues.\(^{15}\) A second generation of
the peptide improved hydrolysis rates by the incorporation of arginine residues to promote stabilization of the developing negative charge of the N-acyl histidine species.\textsuperscript{16}

Barbas and coworkers employed a 24-residue poly-lysine peptide sequence to mimic aldolase activity through imine formation.\textsuperscript{13} Peptides that contained greater helical content achieved rates up to 1900 times greater than the uncatalyzed reaction, rationalized by formation of an amphipathic surface leading to an increase in lysine nucleophilicity (Scheme 4.2). Using the known structures of the miniature proteins, Allemann and colleagues have rationally engineered catalysts that elicit esterase and decarboxylase-like activity.\textsuperscript{17,18} In the esterase study, site-specific incorporation of two histidine residues on the α-helical portion of bovine pancreatic polypeptide create a catalytic HisH\textsuperscript{+}-His motif for p-nitrophenyl fumarate cleavage (Scheme 4.3). The assembled “active site” was proven to be further supported by ionic interactions of neighboring aspartic acid and arginine residues, displaying enzyme-like behavior even on a solvent-exposed surface.

\textbf{Scheme 4.3.} Miniature proteins for ester hydrolysis. Hypothesized mechanism of esterase activity of a dihistidine modified bovine pancreatic peptide displaying the functional coordination of ionic and hydrogen bonding interactions in hydrolysis of nitrophenyl esters. Adapted from Nicoll and Allemann.\textsuperscript{18}
Our approach is to channel the enzyme-like characteristics of peptides to develop a hydrazine-containing peptide for rapid 3-oxo-AHL transamidation (Chapter III). The peptide would serve as a biologically relevant scaffold to reduce the concentration of the active autoinducer, inhibiting the bacterial quorum sensing mechanism. By establishing a productive network of non-covalent interactions, we hypothesize that the peptide could increase rates of transamidation though such mechanisms as enhancing the nucleophilicity of the hydrazine moiety or promoting localization of the AHL to the reactive surface (Figure 4.1). As the required sequence and structure for attaining the envisioned active site is difficult to rationally predict, we chose to employ combinatorial chemistry to design and identify our target peptides.

**Figure 4.1.** Peptide enzyme mimics for AHL transamidation. The hypothetical active site displays how non-covalent interactions could advance the 3-oxo-AHL transamidation. The presence of lipophilic (green), anion (red), and basic (blue), and other residues along the peptide scaffold could aid in AHL localization and enhancing the reactivity of the hydrazine or ketone.

**ii. Methods of Combinatorial Synthesis and Screening of Peptide Libraries**

Beyond their structural parallels, peptides are attractive candidates for the development of enzyme mimics due their facile and robust synthesis. In addition, they offer an immense diversity of sequence space that can be probed using both natural and unnatural
amino acids. To take advantage of these two characteristics, such peptides are often synthesized in library format using combinatorial methods and then screened in a high-throughput fashion to probe the reactivity of the peptide with the target substrate.

**Figure 4.2.** Split and pool synthesis of peptide libraries. Variable residues (X and Y) can be natural or unnatural amino acids introduced to increase library diversity.

Split and pool synthesis is one widely used strategy for the rapid construction large peptide libraries (Figure 4.2).\(^ {19,20} \) The technique is derived from the universally employed Merrifield method for solid-phase peptide synthesis (SPPS) that allows for the stepwise construction of amino acid sequences on a polymeric resin or bead.\(^ {21,22} \) With the peptides bound to the solid support, the resin can be apportioned at any point during the synthesis into separate reaction vessels, allowing for the introduction of variable amino acids into the sequence. After coupling, the beads are pooled into a single reaction vessel, generating a library with a size equal to the number of variable residues incorporated. Subsequent
splitting, coupling, and pooling of the beads allows for a multiplicative increase in the size of the peptide library with minimal synthetic effort. Despite repeated separation and recombination of the resin, each bead retains up to $10^{13}$ copies of a single peptide sequence, also known as the one-bead-one-compound (OBOC) concept. Consequently, an entire library of peptides can then be screened while assessing the reactivity of just a single library member.\(^{23}\)

**Scheme 4.4.** Reactive tagging strategy for colorimetric identification of peptides for acyl transfer catalysis. Covalent attachment and removal of the dye from the peptide through the formed N-acyl intermediate colorizes ($k_1$) and decolorizes ($k_2$) the resin. The beads that most rapidly change color were resynthesized for solution phase screening for acyl transfer catalysts. Assay developed by Bezer et al.\(^{24}\)

High-throughput screening strategies are utilized in conjunction with split and pool synthesis for efficient identification of the most reactive sequences or “hits” in a given OBOC peptide library. Colorimetric reactive tagging assays have become a valuable tool for the completion of such screens as the selected sequences can be identified by a color change in the resin. Covalent capture of a dye-tagged substrate has been employed in identifying peptides for small molecule catalysis in acyl transfer reactions,\(^{25-27}\) aldol additions,\(^{28}\) and oxime-based protein bioconjugations.\(^{29}\) The Gagné and Waters groups have recently employed the strategy for the identification of histidine-containing peptides for acyl transfer catalysis.\(^{24}\) Disperse red functionalized activated esters were synthesized and reacted with solid phase helical peptide libraries to monitor rates of acylation and deacylation at the histidine residue. The peptides capable of rapid acylation ($k_1$) colorized most readily while
those susceptible to deacylation ($k_2$) quickly decolored in the presence of a weak nucleophile (ROH) (Scheme 4.4). Through optimization of a hit obtained in the $k_1$ screen, the assay proved fruitful in identifying a catalyst displaying a rate of acyl transfer 2800 times the background reaction in solution.

![Scheme 4.4](image)

**Figure 4.3.** Reactive tagging scheme for identifying hydrazine-containing peptides for AHL transamidation.

We envision that a similar method can be employed for the high-throughput screening of hydrazine-containing peptides for AHL derivatization (Figure 4.3). The peptide libraries will be constructed using split and pool synthesis, with the inclusion of a photolabile linker between the resin and the peptide sequence to allow for light-mediated cleavage of the peptide from the solid support. By development of a dye-tagged AHL, the substrate can then be screened with the peptide library, leading to a color change upon the reaction of the peptide and the autoinducer. The darkest colored beads will then be manually selected and cleaved from the resin under UV light. The liberated peptide “hits” can then be identified by MALDI-MS. The designed library screening protocol allows for visual identification of the most reactive peptides for AHL transamidation. Upon determining peptides capable of the
transformation, their structure and sequence can then be analyzed to deduce important amino acids and through-space interactions that afford the observed reactivity.

B. Results and Discussion

i. Preparation of Hydrazine-Containing Peptides

To successfully generate peptide libraries for screening in our proposed reactive tagging assay, a method for facile incorporation of hydrazine into a peptide sequence needed to be developed. Inspired by the widespread use of thiol additions to maleimides in bioconjugation chemistry, we sought to determine if conjugate addition to an unnatural maleimide residue would be viable for rapid installation of hydrazine. A series of test peptides (26a, 26b, 26c) were synthesized with 3-maleimidopropionic acid included at the N-terminal residue of each sequence (Figure 4.4a). While on solid support, the peptides were then shaken in a 2% v/v solution of hydrazine in DMF. Upon cleaving the peptides from resin, a mass corresponding to the desired succinimoyl-hydrazine (27) was detected by MALDI-MS. However, two other masses corresponding to a second addition of hydrazine to the maleimide (28) and complete cleavage of the maleimide residue from the sequence (29) were also observed (Figure 4.4b). Stepwise optimization of the conjugate addition, including decreasing hydrazine equivalence, decreasing reaction times, and altering the solvent, were made in order to disfavor the propensity of the nucleophile for overaddition. The presence of both side products was greatly reduced through use of 1% v/v solution of hydrazine in ethanol.
Figure 4.4. Conjugate addition of hydrazine to maleimide-functionalized peptides and corresponding side products. a) test peptides synthesized for optimizing hydrazine addition conditions, where A=alanine and U=α-aminoisobutyric acid b) products observed, as determined by MALDI-MS, upon the reaction of hydrazine with the test peptides.

As an initial proof of concept for the peptide’s ability to undergo transamidation with the quorum sensing autoinducers, the succinimoyl-hydrazine derivative of peptide 26a was combined with 3-oxo-C₈-AHL (22), in 50 mM NaOAc buffer. After 1h, the corresponding hydroxypyrazole product was observed by MALDI-MS. Detection of the peptide-pyrazole adduct validated the basic reactivity necessary for pursuing our combinatorial approach.

However, the presence of overaddition and cleavage products persisted in subsequent screens of the control peptides. Also, the degree to which the side products formed was observed to vary based on peptide sequence. This differential reactivity could inadvertently bias the eventual peptide library screening.
As a result, we turned to synthesizing a series of Boc-protected hydrazines that could be installed on the peptide via standard Fmoc-SPPS conditions. This would ensure the uniform distribution of the nucleophilic residue amongst the library members while allowing for liberation of the hydrazine-functionality during global acid deprotection of the remaining amino acid side chains. As hydrazine proved to be too reactive to yield a single Michael adduct, tert-butyl carbazate was employed to simultaneously decrease the nucleophilicity of the Michael donor while installing the desired Boc-protecting group for SPPS. The Boc-protected succinimoyl-hydrazine (30) was isolated as a single product in 90% yield (Figure 4.5a). To enhance the diversity of the hydrazines to include in the library screens, Boc-protected phenylhydrazine (31) and hydrazinoacetic acid (32) were also prepared. Phenylhydrazine proved to be the most reactive hydrazine in the small molecule screens with the 3-oxo-C_{8}-AHL (Chapter III) and hydrazinoacetic acid, though electronically similar to the succinimoyl-hydrazine, provides a much shorter linker length between the peptide and the hydrazine residue. This could prove important for optimizing the non-covalent interactions provided by the peptide to achieve the desired transamidation rate enhancement.
Figure 4.5. Synthesis of peptide libraries using Boc-protected hydrazines. a) Boc-protected hydrazine derivatives synthesized for use in AHL transamidation library design b) Strategy for site-selective hydrazine incorporation at the N-terminus of a peptide library or through an orthogonally DAP(ivDde) residue, where Xaa represents any amino acid.

As in our initial test peptide screens, the carboxylic acid moiety of each Boc-protected hydrazine allows it to be easily coupled to the N-terminus of a peptide during library synthesis. However, accessing a wider variety of library members would require a method for incorporating the hydrazine residue at any location along the peptide sequence. To address this issue, Fmoc-2,3-diaminopropionic acid (DAP) with an orthogonal ivDde side chain protecting group was employed (Figure 4.5b). The unnatural amino-serine analog is introduced as any other amino acid during SPPS. At the completion of the synthesis, the ivDde group is selectively removed and a Boc-protected hydrazine is coupled to the resultant free amine. Site-specifically introduction of hydrazine at the DAP residue lengthens the distance of the hydrazine from the core of the peptide, but increases the primary and secondary structures that can be evaluated by our high-throughput screen.
ii. First Generation Library Synthesis

Our strategy for the development of peptide libraries for selective and rapid derivatization of the 3-oxo AHLs involves the use of minimalist peptide sequences that could present a surface of functionality to both attract the substrate and enhance the reactivity of the hydrazine. Thus, in designing our first generation of libraries, short sequences that have been previously determined to display secondary structure in water were employed. Short poly-Ala-AIB peptides (Table 4.1, L1) have demonstrated helical propensity, generally existing as a mixture of α/3_{10}-helix based on the present ratio of AIB:Ala residues. In addition to its lipophilicity serving as a potential advantage in recruiting the AHL to the peptide surface, the library was constructed to modify a single amino acid adjacent to the hydrazine to determine any steric or electronic implications the substitution may have on reactivity. A β-hairpin inspired library, including a dPro-Gly turn motif, was also constructed to include variable residues cross-strand to the hydrazine to observe their impact on foldedness and AHL reactivity (Table 4.1, L2). Finally, another poly-Ala-AIB sequence was appended with a type II’ dPro-Pro turn followed by three variable residues in hopes of directing the variable residues back towards the hydrazine while presenting a novel surface for peptide-substrate interaction (Table 4.1, L3).

Table 4.1. First generation of peptide libraries for AHL transamidation.
A variety of natural and unnatural amino acids were incorporated in the libraries to provide a diversity of non-covalent interactions. Anionic, cationic, aromatic, lipophilic, hydrogen bonding, and conformation inducing residues were inserted non-selectively at the variable sites chosen in each library design.

### iii. Development of a Colorimetric Screening Method

With synthesis of our first generation libraries completed, development of a colorimetric tag was necessary to visualize the most reactive library members. As proposed in Figure 4.3, the employment of a dye-functionalized AHL analog as the substrate for the
screen would lead to a color change in the peptides that most rapidly converted the AHL to the corresponding hydroxypyrazole products. In an effort to prepare an AHL-dye conjugate capable of screening under aqueous conditions, 5(6)-carboxyfluorescein was coupled to an amine-functionalized AHL derivative. The resultant fluorescein-AHL (33) was obtained in milligram quantities after purification by semi-prep HPLC. However, preliminary control screens of our first generation libraries with the fluorescein-AHL led to a series of false positive “hits”, promoting resin coloration in the presence of protected peptide libraries and the resin alone. Screening of alternative non-AHL functionalized water-soluble dyes, including Orange G and Aniline Blue, also led to resin coloration. Non-specific interactions between peptide library members and colorimetric indicators in water, especially due to hydrophobic effects, are not uncommon in solid-phase combinatorial screens. \(^{35,36}\) We hypothesized that the fluorescein-AHL was not sufficiently miscible at the low millimolar concentrations being utilized for library screening, leading to its affinity for both the protected peptides and the resin.

![Figure 4.6.](image)

**Figure 4.6.** Non-specific interactions of the fluorescein-AHL. Without the presence of a hydrazine-containing peptide, coloration was observed upon agitating acylated Tentagel resin in a buffered solution of 33.
**Figure 4.7.** AHL transamidation hit identification without a colorimetric tag. Library hits were identified by detection of shift in mass corresponding to the hydroxypyrazole adduct of the 3-oxo-C₁₂-AHL and the peptide.

As an alternative strategy, screening of **L1**, **L2**, and **L3** was completed without the fluorescein-functionalized AHL. The resin-bound libraries were shaken with a buffered solution in the presence of the parent 3-oxo-C₁₂-AHL containing 2% DMSO for enhanced AHL solubility. The screen was terminated by cleaving the entire library from resin to observe a shift in mass (+178.1 m/z) by MALDI-MS corresponding to the formation of the peptide-AHL conjugates (Figure 4.7). We hypothesized that by limiting the screening time, only the most reactive peptides in the library would react to form the hydroxypyrazole product. The strategy proved beneficial in identifying a series of hits from our first generation libraries (Section iv). However, employment of the method uncovered a few inherit limitations. The library composition had to be restricted to peptide sequences with a molecular weight range no greater than 178 mass units, the mass of the hydroxypyrazole adduct, to avoid overlap of the tagged and untagged peptides in the mass spectrum. In addition, signal intensity is known to be dependent on the gas phase protonation affinities of each amino acid, requiring careful interpretation of library hits containing basic residues like lysine or arginine.⁳⁷
Figure 4.8. Revised screening protocol for hydrazine-containing peptide libraries via strain-promoted Huisgen cycloaddition.

To obviate these drawbacks, we considered a third modification to our screening protocol that would expose the library to a functionalized-AHL derivative under aqueous conditions followed by a chemoselective reaction of the AHL analog to develop the screen in organic media (Figure 4.8). This revised two-step assay would remove the non-specific interactions observed between the dye and the peptide library members while also allowing the synthesis of larger and more diverse libraries. Considering chemical processes orthogonal to hydrazone formation, the Huisgen cycloaddition presented a viable and efficient method for selectively linking the dye to the peptide-bound AHL. Consequently, an azido-functionalized 3-oxo-AHL (34) was prepared in addition to a complementary strained-cyclooctyne disperse red derivative (35). The azido-AHL was prepared to structurally mimic the native 3-oxo autoinducer in P. aeruginosa, the 3-oxo-C_{12} AHL (1). Disperse red was
selected due to its frequent use in screening peptide catalyst libraries in organic solvents, including its previous uses in the Gagné Lab.\textsuperscript{24,25,38} Modification of the dye with a dibenzocyclooctyne moiety, originally developed by the Boons group,\textsuperscript{39} allows for rapid conversion to the desired cycloadduct without the use of a copper catalyst, an additive which would inherently discolor the resin. As a control reaction, \textbf{34} and \textbf{35} were combined in equimolar amounts in acetonitrile. Maximum conversion to the desired triazole-containing product \textbf{36} was observed by mass after 4 hours (Scheme 4.5). Further, when the two-step screen was performed with unmodified resin, the dye washed cleanly from the solid support proving that the non-specific interactions observed in our initial screening method had been avoided. This protocol was used to validate the hits in \textbf{L3} and will be used to screen all subsequent generations of peptide libraries.

![Scheme 4.5.](image)

**Scheme 4.5.** Formation of the AHL-dye conjugate, Solution phase control reaction of the azido-AHL (2.0 mM) and dye-functionalized cyclooctyne (2.0 mM) for chemoselective formation of \textbf{36}.

**iv. Library Hits and Solution Phase Screening**

During the iterative redesign of our colorimetric assay, the first generation libraries were tested to identify sequences capable of rapid 3-oxo-AHL transamidation. Each library was screened in buffer at two pH values (4.5, 6.5) with two autoinducers (3-oxo-C\textsubscript{8}-AHL, 3-oxo-C\textsubscript{12}-AHL). Each screen displayed only mild differentiation amongst library members, resulting in a large collection of pyrazole-tagged sequences. The peptides selected as hits displayed the highest relative intensities by MALDI-MS across multiple screens (Table 4.2).
Table 4.2. Hit sequences from the first generation libraries.

<table>
<thead>
<tr>
<th>Library</th>
<th>Library Hits</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1A</td>
<td>ShUAUAAUAUA-NH₂</td>
</tr>
<tr>
<td>L1B</td>
<td>PhHUAUAAUA-NH₂</td>
</tr>
<tr>
<td>L2</td>
<td>ShVWWdPGORV(2-Nal)-NH₂</td>
</tr>
<tr>
<td>L3</td>
<td>Ac-W(Cha)(Phg)PdPUAAUAUAA-NH₂</td>
</tr>
</tbody>
</table>

As AIB is known to increase helix propensity in water, the selection of the amino acid in L1 suggests that inducing greater conformational stability in the peptide may enhance its reactivity. The hits from L2 and L3 contain both lipophilic and aromatic residues, perhaps leading to more rapid AHL transamidation through hydrophobic interactions between the autoinducer and the peptide.

After identification in solid-phase screen, the hits from each library were resynthesized for solution phase screening with the 3-oxo-AHL. Solution phase screens were completed to determine the ability of each peptide to promote AHL derivatization in the absence of the resin and to compare the relative transamidation rates of peptides from
different libraries. As controls to validate hit selection, an additional non-hit peptide from each library was synthesized and tested. The peptides were combined with the 3-oxo-C₈-AHL in pH 6.5 buffer and the disappearance of the AHL was monitored by analytical HPLC (Table 4.3). All the sequences tested were capable of AHL transamidation as the mass of peptide-hydroxypyrazole adduct was detected by LC-MS in each experiment. Supporting our proposed reaction pathway, the putative hydrazone product was observed to form and dissipate as the reaction progressed to yield the hydroxypyrazole product (Figure 4.9). The phenylhydrazine-functionalized peptide most efficiently decreased the concentration of AHL in solution, consistent with our previously conducted small molecule screens (Chapter III). However, the relative rates of AHL consumption achieved by each peptide did not display a significant improvement in the rates provided by hydrazine alone, suggesting the presence of the peptide in our initial design does not largely influence the transamidation reaction. The selected variable residues also proved to be only mildly influential to reaction rates as demonstrated by the similarity in $k_{rel}$ between the peptides selected as hits and the non-hit sequences synthesized.

**Table 4.3.** Relative rates of AHL consumption of hits and non-hits resynthesized from the first generation libraries.
Figure 4.9. Representative HPLC trace for determining rates of AHL consumption in solution phase. In the above trace, the hit from L1A is combined with 40 and monitored from 0 to 14 h. The hydrazone and its isomers are initially visible before being consumed to produce the corresponding pyrazole product.

The minimal differentiation observed in the relative rates of our first generation libraries reflects the minimal differentiation observed during the solid-phase screens. The reactivity of the peptides also parallels the transamidation rates without the peptide. These results suggest that our initial libraries simply do not provide the necessary network of non-
covalent interactions to generate a pseudo-enzymatic environment for promoting AHL derivatization. Strategies to enhance the influence of the peptide on rate could include designing sequence with more well-folded and characterized secondary structures, such as introducing structurally reinforcing covalent crosslinks (i.e. – disulfide bonds, lactam formation) or conducting site-specific mutagenesis of miniature proteins. This would enable a more focused selection of variable sites and residues with a greater probability of those residues engaging at the active site of AHL transamidation. Based on the interpretation of our preliminary results above, the design of new generations of hydrazine-containing peptide libraries is currently ongoing.

**v. Biofilm and Quorum Sensing Inhibition Screening**

Despite the sluggish rates of our first generation library hits, we sought to advance our peptides to screening in cell-based assays to assess the potential of the 3-oxo-AHL transamidation for both biofilm and quorum sensing inhibition. Standard conditions for bacterial culture require the addition of amino acid and carbon sources to buffer in order to support bacterial growth and proliferation. To determine if our reaction would be sustained in bacterial growth media, peptide 47 was screened with the 3-oxo-C₈ AHL (22) in the presence of increasing concentrations of Luria-Bertani (LB) broth. (Figure 4.10). In all instances, conversion to the corresponding hydrazone and hydroxypyrazole products was observed by HPLC. The reaction proved equally robust when screened in PBS buffer containing 0.3 mg/mL tryptic soy broth and 0.01% w/v glucose, a potentially competitive electrophile for the hydrazine-containing peptide. Establishing the ability of the transamidation reaction to proceed in the presence of bacterial growth sources was a necessary precursor to conducting cell-based assays. With the variety of media tested, these
experiments also showcase the selectivity of the peptide for the 3-oxo AHL in an environment of diverse biomolecular functionality, a critical parameter for quorum sensing probes and inhibitors.

![Diagram of peptide 47](image)

Figure 4.10. AHL transamidation in bacterial growth media. The disappearance of 1 (1.0 mM) is monitored upon the addition of peptide 47 (1.0 mM) in varied concentrations of LB broth. Conversion to the pyrazole product was confirmed by LC-MS.

As discussed in Chapter I, quorum sensing has been demonstrated to regulate biofilm production, the most prevalent mechanism for pathogenicity in *P. aeruginosa*. The sustained reactivity of peptide 47 in bacterial growth media encouraged screening of the first generation of libraries for biofilm inhibition. Effective derivitization of the primary autoinducer in *P. aeruginosa*, the 3-oxo-C₁₂ AHL, would impede the bacteria’s ability to reach the required quorate to regulate biofilm formation.⁴⁰ Peptides ⁴⁰, ⁴¹, ⁴₃, and ⁴₄ from the first generation library screens were prepared by serial dilution in 25% LB broth. Stock
solutions of diphenylsulfide (DPS), a known inhibitor of biofilm formation were also prepared as a positive control for comparison to the peptide-containing wells.\textsuperscript{41} The peptides and DPS were added to a 24-well plate containing \textit{P.aeruginosa} PAK and incubated for 18 - 24h. As expected, DPS prevented biofilm growth at both 1 mM and 100 μM, validating our conditions for the screen. Both peptides 40 and 41 inhibited biofilm formation at 1 mM, while peptides 43 and 44 inhibited at a reduced concentration of 100 uM. Peptide concentrations above 1 mM prohibited all cell growth, establishing an upper limit for the use of hydrazine-containing peptides in biofilm inhibition due to their cytotoxicity. Control experiments were also completed with 30 and 31 without the Boc-protection to determine if the hydrazine alone was causing reduction in the biofilm. Notably, the hydrazines prevented all planktonic growth at concentrations as low as 100 μM, proving more cytotoxic alone than when attached to the peptide. The reduced toxicity is valuable when considering the hydrazine-containing peptides as future candidates for \textit{in vivo} testing.

\begin{table}
\begin{tabular}{|c|c|}
\hline
\textbf{Peptide} & \textbf{MICs} \\
\hline
40 & 1 mM \\
41 & 1 mM \\
43 & 100 μM \\
44 & 100 μM \\
\hline
\end{tabular}
\end{table}
Figure 4.11. Biofilm inhibition assay with hydrazine-containing peptides. a) Well-plate displaying the biofilm reduction in the wells containing 43 and 44 compared to the control without a peptide (circled red); a clear well indicates concentrations at which all planktonic growth was inhibited. b) The corresponding minimum inhibitory concentrations (MICs) for biofilm formation of the screened peptides.

To confirm that the biofilm inhibition observed was a direct consequence of QS inhibition, the peptides were screened using a wild-type reporter strain of *P. aeruginosa* PAO1. The strain is transfected with a plasmid encoding for the production of green fluorescent protein (GFP) upon initiation of the QS mechanism. The mutant was cultured in full-strength LB broth and subjected to peptides 40-46 at a final concentration of 1.0 mM. By comparing the relative activity of the wild-type alone, no peptide succeeded in significantly inhibiting the quorum sensing mechanism. Consequently, a correlation between the observed potency of the peptides in biofilm inhibition with their ability to inhibit the QS mechanism cannot be made from our initial results.

Figure 4.12 – Quorum sensing inhibition with hydrazine-containing peptides. Incubation of the first generation of hydrazine-containing peptides with a *P. aeruginosa* PAO1 reporter strain displayed no decrease in relative activity over the 12 h assay.
As determined by the initial spike in relative activity of the wild-type, quorum sensing was initiated in the mutant in less than two hours. Considering the rates of AHL transamidation observed in the solution phase screens of the peptide hits, derivatization of the 3-oxo-C_{12}-AHL is likely to be occurring too slowly to impact the quorum sensing pathway. Developing peptides capable of derivatizing the AHL on a similar time scale as quorum sensing initiation would increase the probability of inhibiting the pathway. Optimization of the growth conditions, including reducing the concentration of the growth media or decreasing the initial optical density of the bacteria, would also be beneficial for observing peptide-based quorum quenching in future screens.

C. Conclusions

The immense diversity and functional flexibility provided by synthetic peptides make them a powerful class of molecules for designing enzyme mimics. In this work, the utility of peptides as a scaffold for promoting our unique hydrazine-mediated AHL transamidation reaction was explored. Through the use of combinatorial chemistry and the power of split and pool peptide synthesis, 127 peptides in four independent OBOC libraries were screened. Library synthesis was facilitated by the employment of Boc-protected hydrazines in conjunction with the use of an orthogonally protected DAP(ivDde) residue for site-selective introduction of the nucleophile along the peptide sequence. Through a series of design modifications, a novel colorimetric reactive tagging assay was developed for high-throughput screening of the hydrazine-containing libraries while eliminating the propensity for non-specific peptide-dye interactions.

Both on solid-support and in solution, the selected peptide hits unequivocally displayed the capability of derivatizing 3-oxo-AHLs. Resynthesis and screening of selected
sequences demonstrated that the reactivity of the peptides was sustained in solution, even in
the presence of bacterial carbon sources such as LB broth and glucose. The primary
sequence proved to only mildly influence transamidation reaction rates in the first generation
libraries, as relative rates similar to hydrazine alone were obtained. The efficiency of
hydroxypyrazole formation was largely determined by the hydrazine incorporated, evidenced
by comparing 41 (phenylhydrazine peptide) to the other sequences screened. We
hypothesize that the modest reactivity of the peptides led to their inability to induce quorum
sensing inhibition in P.aeruginosa PAO1. The key to increasing these reaction rates comes
in identifying sequences in which the non-covalent interactions of the peptide variable
residues participate at the site of AHL transamidation. We are confident that introducing a
greater element of rational design in the selection of our libraries while continuing to increase
the diversity of variable residues in each sequence will enable the identification of an
authentic enzyme-like hit for AHL derivatization.

Interestingly, peptides 40, 41, 43 and 44 were qualitatively observed to inhibit biofilm
formation at minimum concentrations of 100 μM. Our results encourage further screens of
hydrazine-containing peptides to discover greater potencies for P.aeruginosa biofilm
inhibition and determine sequence and structure-activity relationships. However, the
mechanism of observed biofilm reduction is undetermined and cannot yet be correlated with
QS inhibition without corroborating evidence for the gfp-reporter strain assay.
Figure 4.13. Developed protocol for peptide library design and screening. The above schematic depicts our developed procedure for the stepwise discovery of hydrazine-containing peptides for QS and biofilm inhibition through combinatorial peptide library design and colorimetric screening.

In summary, a functional protocol for screening hydrazine-containing peptides for quorum sensing and biofilm inhibition through a high-throughput reactive tagging assay has been established (Figure 4.12). Through iterations of library design, synthesis, and screening, the libraries can be sequentially modified to approach the most efficient scaffolds supporting AHL transamidation. Noting the relatively small number of peptides that were investigated in our first generation libraries, the sequence space remaining to be explored is vast and efforts in developing new generations of hydrazine-containing peptide libraries are currently underway.

D. Experimental

i. General Reagents and Synthesis

Commercial reagents and solvents were purchased from Sigma-Aldrich and Fisher Scientific. Resins for peptide synthesis were purchased from Peptides International and the Fmoc-photolabile linker was obtained from Advanced Chemtech. LB broth for the bacterial
cell assays was obtained from BD Biosciences. *P. aeruginosa* PAK was donated by the Schoenfisch Lab at the University of North Carolina at Chapel Hill, while *P. aeruginosa* PAO1 and the corresponding plasmid for *gfp* production (pGJB5 in *E. coli* DH5α) was obtained from the Whiteley Lab at the University of Texas at Austin. Peptide libraries were analyzed for hits by MALDI-FT on IonSpec spectrometer using 2,5-dihydroxybenzoic acid as the matrix. Resynthesized peptides were purified using a Waters semi-prep HPLC/UV-Vis with a Waters 4.2μM C18 column. Relative rates of peptide transamidation were obtained on an HP 1100 Series analytical HPLC/UV-Vis using a 2.7μM HALO C18 column at 210 nm with products confirmed by mass on an Agilent 1200 Series LC-MS using a 2.7μM HALO C18 column. Small molecule product characterization (1H and 13C) was completed on a 400 or 600 MHz Bruker NMR spectrometer. All related coupling constants are reported in Hertz and shifts in ppm. NMR solvents were purchased from Cambridge Isotopes Laboratories. High resolution mass spectra were obtained on an Agilent Accurate LC-TOF Mass Spectrometer (Agilent Series 6220) operating in positive ion mode with an electrospray ionization source (fragmentor = 175 V). The data was analyzed using an Agilent MassHunter Workstation Software, Qualitative Analysis (V. B.02.00).

**Synthesis of Boc-Protected Hydrazines**

Boc-protected hydrazine 32 was purchased commercially and hydrazine 31 is a known compounds and was synthesized following previously published procedures. For the synthesis of Boc-protected hydrazine 30, 3-maleimidopropionic acid (1.69 g, 0.0100 mol) was dissolved in 20 mL of EtOH. With stirring, tert-butyl carbazate (1.32 g, 0.0100 mol) was dissolved in 5 mL of EtOH and added dropwise to the reaction mixture. The reaction
mixture was allowed to heat at 60°C for 8 h. The reaction mixture was concentrated under vacuum to yield 30.

4-(2-(tert-butoxycarbonyl)hydrazinyl)benzoic acid (31)

\[
\begin{align*}
\text{O} & \quad \text{N} \\
\text{H} & \quad \text{N} \\
\text{O} & \quad \text{N} \\
\text{H} & \quad \text{O} \\
\text{H} & \quad \text{OH} \\
\text{H} & \quad \text{H}
\end{align*}
\]

99.0% yield; \( \delta_H \) (DMSO-d\(_6\), 400 MHz) 12.2 (1H, s), 8.92 (1H, s), 8.24 (1H, s), 7.72 (2H, d, \( J = 8.8 \) Hz), 6.63 (2H, d, \( J = 8.8 \) Hz), 1.41 (9H, s); MS (ESI) calcd for C\(_{12}\)H\(_{16}\)N\(_2\)O\(_4\): 253.11, found: 253.12 (M + H).

4-(2-(tert-butoxycarbonyl)hydrazinyl)benzoic acid (30)

\[
\begin{align*}
\text{O} & \quad \text{N} \\
\text{H} & \quad \text{N} \\
\text{O} & \quad \text{N} \\
\text{H} & \quad \text{O} \\
\text{H} & \quad \text{OH} \\
\text{H} & \quad \text{H}
\end{align*}
\]

72.0% yield; \( \delta_H \) (DMSO-d\(_6\), 600 MHz) 12.4 (1H, s), 8.36 (1H, s), 5.09 (1H, s), 3.95 (1H, m), 3.55 (1H, m), 2.81 (1H, dd, \( J_1 = 8.4 \) Hz, \( J_2 = 9.6 \) Hz), 2.51 – 2.46 (3H, m), 1.39 (9H, s); \( \delta_C \) (DMSO-d\(_6\), 150 MHz) 176.5, 175.9, 172.4, 156.8, 79.3, 57.6, 34.4, 34.3, 32.0, 28.6; MS (ESI) calcd for C\(_{12}\)H\(_{19}\)N\(_3\)O\(_6\): 302.1347, found: 302.1343 (M + H).

*Synthesis of Fluorescein-AHL*
In a 20-mL scintillation vial, 48 (150 mg, 0.352 mmol) was dissolved in 2 mL of DCM and combined with 2 mL of TFA. After 20 min, the solvent and acid were removed under a stream of \( \text{N}_2 \) gas to yield the free amine. The deprotected AHL, 5(6)-carboxyfluorescein (139 mg, 0.369 mmol), and DIEA (0.121 mL, 0.739 mmol) were all combined and dissolved in DMF (4 mL). COMU (158 mg, 0.369 mmol) was then added and the reaction mixture was allowed to stir at 0°C for 1h. The reaction mixture was then warmed to room temperature and allowed to stir for an additional 3h. The contents of the flask was then diluted in EtOAc (50 mL) and extracted with 1M HCl (2 x 10 mL), 1M NaHCO\(_3\) (2 x 10 mL), and washed with brine (1 x 15 mL). The organic layer was collected, dried over NaSO\(_4\), filtered, and concentrated. The concentrated organic layer was redissolved in 40% MeCN in water and purified by reverse-phase HPLC to isolate the 5-carboxyfluorescein regioisomer (33). The Boc-protected AHL 48 was prepared according to previous published procedures.\(^{43}\)

\[(S)-N-(11,13-dioxo-13-((2-oxotetrahydrofuran-3-yl)amino)tridecyl)-3',6'-dihydroxy-3-oxo-3H-spiro[isobenzofuran-1,9'-xanthene]-6-carboxamide (33)\]
2.1% Yield; δ\textsubscript{H} (CDCl\textsubscript{3}, 600 MHz) 8.06 (2H, m), 7.54 (1H, s), 7.25 (1H, br s), 7.15 (1H, s) 6.77 (2H, d, J = 2.4 Hz), 6.73 (2H, d, J = 8.4 Hz), 6.62 (2H, dd, J\textsubscript{1} = 2.4 Hz, J\textsubscript{2} = 8.4 Hz), 4.53 (1H, m), 4.40 (1H, td, J\textsubscript{1} = 7.2 Hz, J\textsubscript{2} = 1.8 Hz), 4.24 (1H, m), 3.38 (2H, s), 3.29 (2H, m), 2.53 (3H, m), 2.26 (1H, m), 1.51 (4H, m), 1.53 – 1.25 (13H, m); δ\textsubscript{C} (CD\textsubscript{3}CN, 150 MHz); MS (ESI) calcd for C\textsubscript{38}H\textsubscript{40}N\textsubscript{2}O\textsubscript{10}: 685.2759, found: 685.2756 (M + H).

**Synthesis of Azido-AHL**

Azido-AHL 34 is a known compounds and was synthesized following previously published procedures.\textsuperscript{43}

**(S)-13-azido-3-oxo-N-(2-oxotetrahydrofuran-3-yl)tridecanamide (34)**

17.7% Overall Yield (3 steps); δ\textsubscript{H} (CDCl\textsubscript{3}, 600 MHz) 7.29 (1H, s), 4.60 (1H, m), 4.47 (1H, td, J\textsubscript{1} = 7.6 Hz, J\textsubscript{2} = 1.6 Hz), 4.29 (1H, m), 3.48 (2H, s), 3.26 (2H, t, J = 6.8 Hz), 2.75 (1H, m), 2.54 (2H, t, J = 7.2 Hz), 2.26 (1H, m), 2.22 (1H, m), 1.60 (4H, m), 1.39-1.21 (12H, m); MS (ESI) calcd for C\textsubscript{17}H\textsubscript{28}N\textsubscript{4}O\textsubscript{4}: 353.21, found: 353.25 (M + H).

**Synthesis of Strained-Cyclooctyne Disperse-Red Dye**
In a 10-mL RB flask, 49 (70.8 mg, 0.190 mmol), 50 (50.0 mg, 0.181 mmol), and DIEA (0.066 mL, 0.3801 mmol) were all combined and dissolved in DMF (2 mL). COMU (81.4 mg, 0.190 mmol) was then added and the reaction mixture was allowed to stir at 0°C for 1 h. The reaction mixture was then warmed to room temperature and allowed to stir for an additional 3 h. The contents of the flask were then diluted in EtOAc (25 mL) and extracted with 1 M HCl (2 x 5 mL), 1 M NaHCO$_3$ (2 x 5 mL), and washed with brine (1 x 5 mL). The organic layer was collected, dried over Na$_2$SO$_4$, filtered, and concentrated. The concentrated organic layer was redissolved in DCM and purified by column chromatography. Compound 38 was synthesized by previously reported procedure$^{25}$ and 39 was purchased commercially.

![Compound 35](image)

73.7% Yield; $\delta_H$ (CDCl$_3$, 600 MHz) 8.32 (2H, d, $J = 9$ Hz), 7.92 (3H, m), 7.69 (1H, d, $J = 7.6$ Hz), 7.42 – 7.27 (7H, m), 7.03 (1H, t, $J = 5.6$ Hz), 6.76 (2H, d, $J = 9$ Hz), 5.15 (1H, d, $J = 13.8$ Hz), 3.86 (2H, q, $J = 18.6$ Hz), 3.69 (1H, d, $J = 14.0$ Hz), 3.65 – 3.52 (6H, m), 3.39 (1H, m), 3.21 (1H, m), 2.48 (1H, m), 1.95 (1H, m), 1.26 (3H, t, $J = 7.1$ Hz); MS (ESI) calcd for C$_{36}$H$_{34}$N$_6$O$_5$: 631.2663, found: 631.2666 (M + H).

### ii. Peptide Library Synthesis and Screening

*General Procedure for Synthesis of Peptide Libraries*
Peptides were synthesized by hand using standard Fmoc-SPPS on TentaGel S NH₂ resin (0.29 meq/g loading). As the first coupling in each library synthesis, the Fmoc-photolabile linker (2 eq) was attached to the resin by agitation with N₂ in a 0.2 M solution of HBTU/HOBt in DMF (2 eq) and DIEA (4 eq) for 2 h. All subsequent couplings were completed with a 4:4:8-fold excess of amino acid:0.2 M HBTU/HOBt:DIEA in DMF with a coupling time of 1.5 h. After coupling, the resin was washed sequentially with DMF (3x), DCM (3x), and DMF (2x). Fmoc-deprotection was completed by subjecting the resin to a 20% piperdine/DMF solution (3 x 30 min) followed by the same washing sequence as the coupling reaction. Completed sequences were acetyl capped at the N-terminus with a solution of 5% acetic anhydride and 6% lutidine in DMF (2 x 30 min) followed by the same washing sequence. For peptides containing DAP(ivDde), the resin was agitated in a 2% hydrazine/DMF solution (6 x 2 min) with a DMF wash (5x) in between each cycle to remove the ivDde protecting group. The desired Boc-protected hydrazine was then coupled onto the resin using the same ratios of HBTU/HOBt and DIEA as previous. The peptides were then dried under N₂ and stored in the freezer until screening.

**General Procedure for Screening of Peptide Libraries**

The protected peptide library (25 mg, 0.0070 mmol) was added to a 25-mL plastic fritted syringe and deprotected by shaking on an orbital shaker with 95:2.5:2.5 trifluoroacetic acid:triisopropylsilane:water for 2 h. The deprotected resin was washed thoroughly with water (3x), MeOH (3x), DMF (3x), and DCM (3x). For non-colorimetric screens of L1, L2, and L3, the 3-oxo-C₁₂ AHL (2.08 mg, 0.0070 mmol) was dissolved in a 2% solution of DMSO in PBS buffer and added to the resin. The resin was allowed to shake in buffer for 1
h, with reaction times varying in subsequent screens based on the intensity of the tagged products observed mass spectrometry. The resin was then washed and transferred to a 2-mL Eppendorf tube containing MeOH (0.5 – 1.0 mL). The Eppendorf tube was placed under a UV light (365 nm) for at least 3 h with stirring to cleave the photolinker and solubilize the peptide library. The peptide solution was then analyzed by MALDI-MS. For colorimetric screens of L3, etc. the azido-AHL 34 (1.97 mg, 0.0070 mmol) was dissolved in a 2% solution of DMSO in PBS buffer and added to the resin. The resin was shaken, washed, and then combined with dye 35 (4.4 mg, 0.007 mmol) in 3.5 mL of MeCN. The resin was again shaken in the present of the dye for 4 h, washed with MeCN (2x), DMF (3x), and DCM (3x) and plated on a plastic petri dish. The resin was inspected under a confocal microscope and the darkest colored beads were manually picked for cleavage. The “hits” were transferred to a MeOH containing Eppendorf tube, placed under UV-light, and then analyzed by MALDI-MS.

Synthesis of Peptides for Solution Phase Screening

The peptides were synthesized by the same procedure utilized for peptide library synthesis but on CLEAR Amide resin (0.35 meq/g). The completed sequences were cleaved from the resin by shaking with 95:2.5:2.5 trifluoroacetic acid:triisopropylsilane:water for 2 h. The solution was then flushed from the resin into a 50 mL Falcon tube and concentrated under N₂. The concentrated acid solution was then precipitated in 50 mL of diethyl ether and centrifuged to pellet the peptide. The ether was decanted and the crude peptide was again dried under N₂. The peptide was then purified by semi-prep HPLC in an MeCN:H₂O mobile phase with 0.1% trifluoroacetic acid and lyophilized to dryness.
**Solution Phase Peptide Screening**

The purified peptide (0.0010 mmol) was added to a solution of the 3-oxo-C₈-AHL (0.24 mg, 0.0010 mmol) in PBS at pH = 6.5. The reagents were combined at time = 0 ± 5 min and placed on an analytical HPLC autosampler for injection every 135 minutes. Separations were performed in a mobile phase of 25 to 50% B over 25 minutes. Relative rates were obtained calculating the decrease in AHL concentration over time.

**iii. QS and Biofilm Inhibition Assays**

**Biofilm Inhibition Assay**

An overnight culture of *P. aeruginosa* PAK was grown in 100% LB broth in PBS at 37°C on a rotary shaker. The bacteria was then reinoculated in 25% LB at pH 6.5 and grown to an OD₆₀₀ = 0.25. The peptides (0.015 mmol) were dissolved in DMSO (15 μL) followed by the addition of 25% LB PBS at pH = 6.5 to a final reach a final peptide concentration of 100 mM. Serial dilutions to 10 mM and 1.0 μM were prepared from a stock of 5% DMSO/95% LB PBS solution. In a 24-well plate, 900 μL of the bacterial culture was combined with 100 μL of the peptide solutions and incubated for 18 h. Biofilm inhibition was qualitatively assessed by visual inspection to determine the presence of a pellicle biofilm.

**QS Inhibition Assay**

Tested peptides (0.010 mmol) were dissolved in DMSO (5 μL) and followed by the addition of 100% LB PBS at pH 6.5 to reach a final concentration of 20 mM. An overnight culture of *P. aeruginosa* PAO1 (*PrsaL-gfp*) was diluted to an OD₆₀₀ of 0.05 in the same broth containing...
buffer. The culture (190 μL) was then transferred to a Greiner 96-well flat bottom plate followed by the addition of the peptide (10 μL) to their corresponding wells. The resultant GFP expression was monitored by fluorescence plate reader over 12 h. The fluorescence intensities recorded were normalized for cell density to obtain the reported relative activity of each peptide.
REFERENCES


