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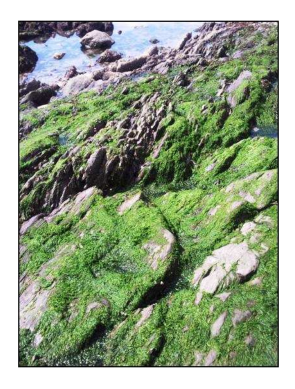
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The Signal Based Relationship between the Green Seaweed Ulva and its Indigenous Bacterial Community



Matthew S. Twigg M.Sc. B.Sc. (hon.)

Thesis submitted to the University of Nottingham for the degree of Doctor of Philosophy

December 2012

For Mum and Dad

"They say the sea is cold but the sea contains the hottest blood of all"

D. H. Lawrence

Declaration

Unless otherwise acknowledged the work presented in this thesis is my own. No part has been submitted for another degree in the University of Nottingham or any other institute of learning

Matthew S. Twigg M.Sc. B.Sc. (hon.)

December 2012

Abstract

This project has focused on the relationship between the green seaweed Ulva, commonly found in the intertidal zone of the UK coastline and its cognate bacterial community. It has previously been reported that motile Ulva zoospores are attracted to N-Acylhomoserine lactones (AHLs), signalling molecules utilised by Gram-negative bacteria in a density dependent form of cellular communication termed quorum sensing (QS) and produced by several biofilm dwelling species of marine bacteria. The species represented in the bacterial community associated with Ulva spp. were identified by generating a 16S rDNA phylogenetic clone library from bacterial DNA isolated from the surface of the seaweed. These data revealed that the majority of the population belonged to the Proteobacteria or Bacteroidetes phyla. In order to investigate whether QS signalling affected the rate of zoospore germination in addition to zoospore attraction, Ulva zoospores were settled and allowed to grow on synthetic AHLs, biofilms derived from AHL-producing model organisms and strains relevant to the Ulva epiphytic population which were shown to produce AHLs. Results from these experiments revealed that AHLs affected zoospore germination and the early growth of the Ulva germling as zoospores germinated and grown in the absence of AHLs were significantly longer than those germinated in the presence of AHLs. We therefore hypothesise that reduced germling growth in the presence of AHLs allows Ulva to obtain a healthy epiphytic bacterial community that is vital for the seaweed's later development. Further understanding of Ulva growth biology could have potential applications in preventing marine biofouling by this genus of seaweed.

This study progressed to characterise AHL production in a number of strains of Shewanella and Bacteroidetes bacteria, which, for differing reasons were deemed relevant to Ulva biology. Although data presented by this thesis showed AHL production in these bacterial groups, AHL synthase and response regulator sequences could not be identified in the published genome sequences from either Shewanella or the Bacteroidetes. This study also identified an AHL inactivating acylase enzyme in an environmental Shewanella isolate. This acylase, AacS, was shown to degrade a variety of synthetic AHLs and the AHLs produced by Yersinia pseudotuberculosis. This study has therefore increased the range of marine bacteria known to be producing AHLs, however the lack of AHL synthase and response regulator genes in the genomes of these bacteria leads to the conclusion that many marine bacteria possess novel, yet to be characterised AHL-mediated QS systems.

Finally, this study screened a number of extracts from marine microalgae for compounds that act as agonists or antagonists to AHL-mediated QS. Although no AHL mimics were identified data presented by this thesis showed extracts to affect the luminescence produced in lux-based AHL bio-reporters in the presence of exogenously added signal, affect a number of QS regulated phenotypes in marine pathogens and effect QS regulated genes in the human pathogen Pseudomonas aeruginosa. As such, we hypothesise that these microalgae have the ability to produce quorum-quenching compound(s). Further characterisation of quorum-quenching compound(s) produced by microalgae may be beneficial in the bio-control of pathogenic bacteria in aquaculture and may act as candidates for novel antibiotics.

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Acronyms and Abbreviations

Adv. Average

AHL N-acylhomoserine lactone

AQ 2-Alkyl-4-quinolones

ANOVA Analysis of Variance

BLAST Basic Local Alignment Search Tool

d.H₂O Distilled Water

CCAP Culture Collection of Algae and Protozoa

DCM Dichloromethane

DMSP Dimethylsulphoniopropionate

DMS Dimethylsulphide

DNA Deoxyribonucleic Acid

d.NTP Deoxyribonucleotide Triphosphate

DOTUR Distance-Based Operational Taxonomic Unit and Richness

EDTA Ethylenediaminetetraacetic Acid

FITC Fluorescein Isothiocyanate

HF High Fidelity

HPLC High Performance Liquid Chromatography

HSL Homoserine Lactone

LCMS Liquid Chromatography Mass Spectroscopy

m/v Mass to Volume

NCIMB National Collection of Industrial, food and Marine Bacteria

OD Optical Density

o/n Over Night **ORF** Open Reading Frame OTU Operational Taxonomic Unit **PBS Phosphate Buffer Solution PCR** Polymerase Chain Reaction PHQ 2-n-pentyl-4-quinolinol Ribosomal Deoxyribonucleic Acid rDNA **RLU** Relative Light Units RNA Ribonucleic Acid **RPM Rotations Per Minute SDS** Sodium Dodecyl Sulphate SE **Standard Error** Scanning Electron Micrograph **SEM** TAE Tris-acetate-EDTA TLC Thin Layer Chromatography Ultra Violet UV Wild Type wt Weight to Volume W/VVolume to Volume v/v5' 5 Prime (DNA Orientation) 3, 3 Prime (DNA Orientation)

List of Bacterial Signal Molecules

C4-HSL	N-butanoyl-L-homoserine lactone
3-oxo-C4-HSL	N-(3-oxobutanoyl)-L-homoserine lactone
3-OH-C4-HSL	N-(3-hydroxybutanoyl)-L-homoserine
C6-HSL	N-hexanoyl-L-homoserine lactone
3-oxo-C6-HSL	N-(3-oxhexananoyl)-L-homoserine lactone
3-OH-C6-HSL	N-(3-hydroxyhexanoyl)-L-homoserine lactone
C8-HSL	N-octanoyl-L-homoserine lactone
3-oxo-C8-HSL	N-(3-oxooctanoyl)-L-homoserine lactone
3-OH-C8-HSL	N-(3-hydroxyoctanoyl)-L-homoserine lactone
C10-HSL	N-decanoyl-L-homoserine lactone
3-oxo-C10-HSL	N-(3-oxodecanoyl)-L-homoserine lactone
3-OH-C10-HSL	N-(3-hydroxydecanoyl)-L-homoserine lactone
C12-HSL	N-dodecanoyl-L-homoserine lactone
3-oxo-C12-HSL	N-(3-oxododecanoyl)-L-homoserine lactone
3-OH-C12-HSL	N-(3-hydroxydodecanoyl)-L-homoserine lactone
C14-HSL	N-Tetradecanoyl-L-homoserine lactone
PQS	(Pseudomonas Quinolone Signal) 2-heptyl-3-hydroxy-4-quinolone
HHQ	2-heptyl-4-quinolone

Units of Measurement

Mass

kg Kilograms
g Grams
mg Milligrams

µg Micrograms
ng Nanograms

Length

m Metre
mm Millimetre
μm Micrometre
nm Nanometre

Substance Concentration

M Molar
mM Millimolar
μM Micromolar
nM Nanomolar

Amount of Substance

mol Moles

mmol Millimoles

μmol Micromoles

Volume

L Litre

ml Millilitre

μl Microlitre

Time

h Hours

min Minutes

s Seconds

Temperature

°C Degrees Celsius

Molecular Biology

bp Base pair

kb Kilobase pair

Da Dalton

kDa Kilo Dalton

Electrical Potential

kV Kilovolts

V Volts

Sound Frequency

MHz Megahertz

Chapter 1

General Introduction

1.1 Bacterial Quorum Sensing

1.1.1 The Discovery and General Overview of Bacterial Quorum Sensing

Since the foundation of the science of microbiology by Antoni Van Leeuwenhoek in 1664, the classical view of the bacterial world has been that of an autonomous, single celled organism. This individual bacterial cell reproduces via binary fission and strives for its survival within a particular environmental niche. The black and white picture presented by this view, is however completely incorrect. In order to survive, bacteria have developed means to recognise, co-operate or outcompete other prokaryotic and eukaryotic organisms within their local environment. One such method is facilitated by the production and secretion of chemical signalling molecules. These signalling molecules play a vital role in the microbial phenomenon known as 'quorum sensing' (QS) (reviewed by Fuqua et al., 1994).

The term 'quorum sensing' was first adopted in an review paper by Fuqua et al. in 1994 and is defined as being a population size-dependent form of cell-to-cell communication facilitated via the production and transduction of small signalling molecules (reveiws by Williams et al., 2007; Fuqua et al., 1994). Although the term quorum sensing was first adopted in 1994 the process of population-dependent bacterial communication was initially recognised through the study of bioluminescence in the marine bacterium Vibrio fischeri during the 1970's (Fuqua et al., 1994; Nealson et al., 1970). V. fischeri has the ability to emit bioluminescence, (Figure 1.1), due to the action of luciferase that oxidises reduced flavin and a long chain fatty aldehyde that is in turn provided by a fatty acid reductase (Hastings et al., 1985; Belas et al., 1982). The enzymatic components required for bacterial bioluminescence are transcribed from the lux operon. This operon consists of 7 genes; luxA and luxB which, when translated,

provide the two subunits of luciferase; the gene products of luxC, luxD and luxE provide the components of the fatty acid reductase and the gene products of luxG and luxH provide the reduced flavin substrate (Meighen, 1991). Nealson et al. (1970) observed that in V. fischeri lag phase and early log phase cultures, luciferase expression was repressed, however during mid log phase luciferase was produced and bioluminescence occured (Nealson et al., 1970). Nealson et al. (1970) also observed that once luciferase production was initiated the rise in bioluminescence increased at a greater rate than bacterial growth (Nealson et al., 1970). The initiation of luciferase transcription and subsequent rapid rise in bioluminescence was attributed to conditioning of the culture medium by some, then unknown, factor (Nealson et al., 1970). Further study into V. fischeri confirmed that culture conditioning initiated luciferase transcription, as the addition of cell free supernatant from late log phase cultures into fresh cultures resulted in the early induction of bioluminescence (Hastings and Nealson, 1977).

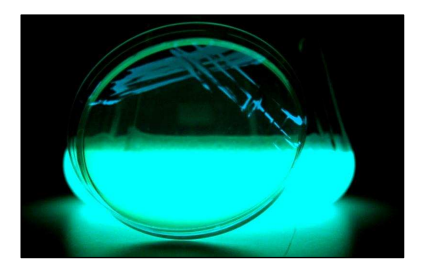


Figure 1.1. Bioluminescent stationary phase culture of V. fischeri. V. fischeri cultures are bioluminescent due to the actions of the enzyme luciferase, which is positively regulated via quorum sensing.

The molecule responsible for luciferase induction and therefore bioluminescence in V. fischeri was identified as N-(3-oxohexanoyl) homoserine lactone (Eberhard et al., 1981). This molecule is one of a larger family of small diffusible signal molecules known as N-acyl homoserine lactones (AHLs). Its discovery lead to the general paradigm of bacterial QS; that a bacterial cell or population of bacterial cells is able to produce and respond to the presence of molecules, more widely referred to as signalling molecules, in the extracellular environment, to bring about a population-wide change in behaviour (reveiwed by Fuqua et al., 1994). At low bacterial densities the concentration of signal molecules is low but as the bacterial density increases so does the concentration of the signal to a point where the concentration of signalling molecules passes a threshold and is detected by signal molecule response regulators. At this point the bacterial population is termed to be 'quorate', the signal is then transduced and target gene expression is triggered (Figure 1.2) (review by Bassler, 2002). In addition to regulating gene expression in the bacterial population, QS signal molecules may in some cases positively auto-regulate their own production leading to a further increased concentration of signal molecules when the bacterial population is quorate. This auto-regulation of cognate signal molecule production and the expression of other genes has been termed autoinduction, leading to QS signal molecules often being referred to as autoinducers (review by Miller and Bassler, 2001).

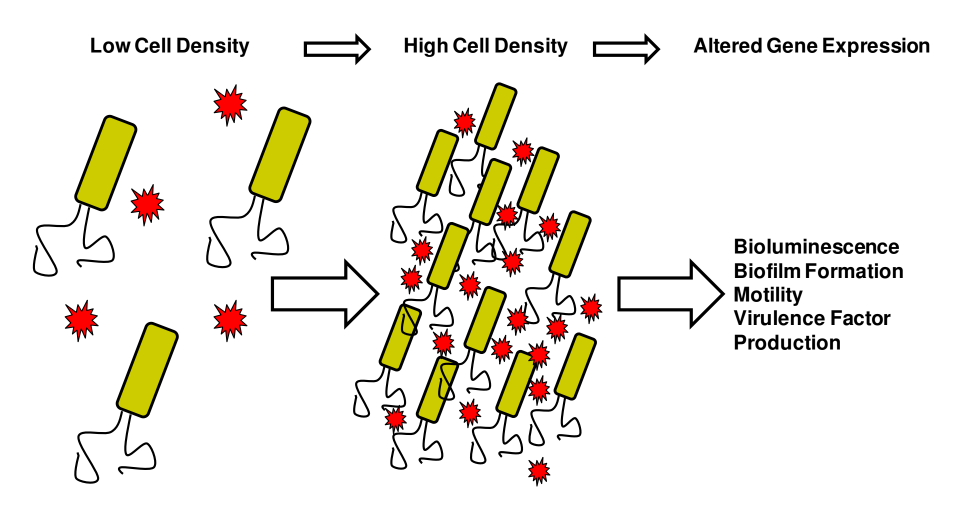


Figure 1.2. Illustration of bacterial quorum sensing. Bacteria can produce and export signalling molecules (red stars) to the extracellular environment. At low cell densities signal molecule concentration is low, as the cell density increases signal molecule concentration will also increase. At a specific threshold concentration signal molecules are detected by the bacteria effecting a change in the populations gene expression.

1.1.2 AHL-mediated Quorum Sensing

AHLs are by far the best characterised class of QS signal molecules. The chemical composition of an AHL autoinducer is that of a five-membered homoserine lactone ring N-acylated with a fatty acid side chain at α-position 1 (Eberhard et al., 1981). AHL acyl side chains range from 4 to 18 carbons in length, can either be saturated or un-saturated and may contain hydroxyl or oxy substitutions on the third carbon (Figure 1.3) (Marketon et al., 2002; Schaefer et al., 2000). The number of carbons in the acyl side chain is usually even however side chains with an odd number of carbons have been reported in different species including Rhizobium leguminosarum and Yersinia pseudotuberculosis (Ortori et al., 2007; Horng et al., 2002). In order to be biologically active, AHLs have to possess an acyl side chain of at least 4 carbons in length (Yates et al., 2002). This minimum side chain length requirement is due to the effect of pH on the homoserine lactone ring. Acyl side chains composed of 4 or more carbon atoms stabilise the structure of the AHL, maintaining biological activity (Chhabra et al., 2003; Yates et al., 2002).

Figure 1.3. General chemical structures of AHL signalling molecules. AHLs are composed of a homoserine lactone ring with an acyl fatty acid side chain ranging from 4 to 18 carbons in length. The acyl fatty acid side chains can be saturated or unsaturated and possess hydroxyl or oxy substitutions on the third carbon (Figure sourced from Williams et al., 2007).

AHLs are synthesised by signal synthase proteins that were first studied in the marine bacterium V. fischeri. The signal synthase protein in V. fischeri is LuxI, a protein with a relative mass of 25 kDa, produced as a result of the expression of the gene luxI (Devine et al., 1989; Engebrecht and Silverman, 1984). LuxI acts as an enzyme catalysing the formation of an amide bond between S-adenosylmethionine and 3-oxo-hexanoyl-CoA in order to produce 3-oxo-hexanoyl-HSL, the cognate AHL of V. fischeri (Eberhard et al., 1991). Other LuxI type AHL synthase enzymes carry out the same function but recognise different acylated-acyl carrier protein conjugates which provide the differing fatty

acid biosynthetic precursors required to produce the wide range of AHLs used by bacteria (Moré et al., 1996).

AHLs move to the extracellular environment either by free diffusion across the plasma membrane, or in the case of some long chain AHLs which have increased hydrophobicity, export to the extracellular environment is achieved via an efflux pump (Aendekerk et al., 2002; Kaplan and Greenberg, 1985). This is the case for Pseudomonas aeruginosa, which employs the MexAB-OprM pump for the export of 3-oxo-C12-HSL (Aendekerk et al., 2002).

When AHL concentration in the extracellular environment is sufficient to reach a threshold, LuxR, a twin domain 27 kDa protein binds the AHL at the amino terminus and the resulting complex binds DNA via a classic helix-turn-helix motif in the C-terminus (Fuqua et al., 1994; Slock et al., 1990; Engebrecht and Silverman, 1984). The luxR gene is found at the same locus as luxI within the V. fischeri lux operon (Devine et al., 1989). The AHL/LuxR complex has been shown to bind to palindromic DNA sequences, 18-20 bp in length upstream from the genes they regulate, termed lux boxes (Figure 1.4) (Gray et al., 1994).

Vibrio fischeri Bacterial Cell

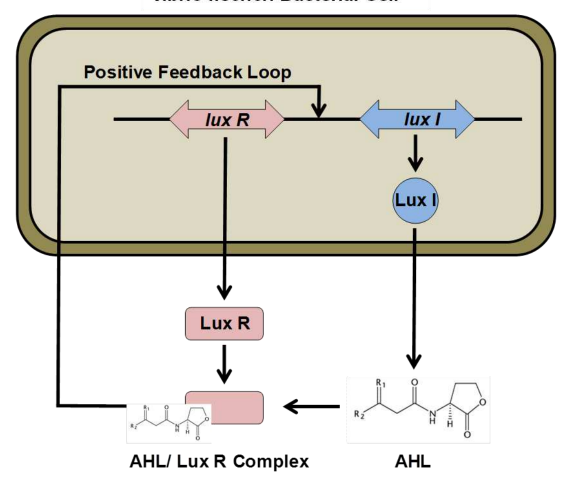


Figure 1.4. Schematic of the AHL-mediated QS system present in V. fischeri. LuxI produces AHL signalling molecules which are exported to the extracellular environment. LuxR acts as the AHL response regulator. AHL/LuxR complexes bind palendromic DNA sequences upstream of both luxI and genes regulated by AHL-mediated QS, altering gene expression and causing a positive feedback loop which increases AHL production. Many Gram-negative bacteria possess AHL-mediated QS systems.

Since the initial identification of N-(3-oxohexanoyl)-L-homoserine lactone as the autoinducer molecule in V. fischeri many other bacterial species have been reported to possess LuxR/I homologues capable of producing and transducing AHL molecules. AHL-mediated QS has also been proven to regulate a wide viariety of different phenotypes aside from bioluminescence. These include virulence, biofilm formation, siderophore production and motility

(reveiwed by Williams et al., 2007). A number of AHL-mediated QS systems belonging to different bacteria are listed alongside the phenotype(s) they regulate in Appendix 1. Although it is clear that AHLs are produced by a great number of bacterial species, all current research suggests that AHL-mediated quorum sensing is however restricted to Gram-negative bacteria only (reveiwed by Williams et al., 2007).

In addition to the LuxR/I systems described above and shown in Appendix 1, DNA sequencing of bacterial chromosomes has revealed the presence of a number of luxR genes unconnected with luxI-type AHL synthase genes (Patankar and Gonzalez, 2009). The resultant proteins expressed by these genes have been termed LuxR orphans which in contrast to the 'classical' LuxR homologue do not transduce AHLs produced by a cognate synthase but are capable of interacting with AHLs to regulate gene expression (Patankar and Gonzalez, 2009). LuxR orphans have been identified in bacterial species possessing a functional LuxR/I system and include; QscR in P. aeruginosa, ExpR and NesR in Sinorhizobium meliloti and VirR and ExpR2 in Erwinia sp. (Barnard and Salmond, 2007; Pellock et al., 2002; Chugani et al., 2001). Additionally there are a number of LuxR orphans in bacteria where there is no functional LuxI and therefore no cognate AHLs produced, these include; SdiA in Escherichia coli and Salmonella enterica and XccR in Xanthomonas campestris (Zhang et al., 2007; Ahmer, 2004; Ahmer et al., 1998). It is possible that LuxR orphans may enable bacteria to sense the presence of rival species in their proximal environment or gain an advantage by using exogenously produced AHLs to regulate their own gene expression without the metabolic burden of producing cognate AHLs. This hypothesis is supported by the observations of non AHL-producing bacteria such as E. coli possessing orphan LuxR homologues and of the P. aeruginosa LuxR orphan, QscR's, ability to detect 3-oxo-C10-HSL (Lee et al., 2006; Ahmer, 2004). Another hypothesis is that orphan LuxR homologues exist to form heterodimers with LuxR proteins linked to luxI orthologes. Transcriptional regulation as a result of LuxR heterodimerisation has been observed in Agrobacterium tumefaciens (Chai et al., 2001).

AHL synthesis and transduction is not only limited to LuxI and LuxR type proteins, a further family of proteins has also been shown to facilitate AHLmediated QS. Similar to the discovery of LuxI and LuxR, these proteins were also discovered during the study of bioluminescence in a marine bacterium, Vibrio harveyi. V. harveyi possesses an AHL synthase protein termed LuxM which catalyses the synthesis of its cognate AHL, 3-OH-C4-HSL (Bassler et al., 1993; Cao and Meighen, 1989). Structural analysis of LuxM showed it to have no homology to LuxI; however the two synthase genes catalyse AHL production using the same substrates (Hanzelka et al., 1999; Bassler et al., 1993). 3-OH-C4-HSL synthesised by LuxM is not transduced by a LuxR homologue, AHL detection and transduction is however carried out by a membrane bound two component sensor kinase protein termed LuxN (Freeman et al., 2000). At low cell densities and, therefore, low AHL concentration LuxN acts as a kinase and under such conditions phosphate is transferred to a cytoplasmic phosphotransferase protein LuxU and then onto the σ^{54} dependent activator LuxO. Phosphorylation of LuxO results in the repression of bioluminescence in V. harveyi (Freeman et al., 2000; Freeman and Bassler, 1999a; Freeman and Bassler, 1999b). At high cell

densities and therefore high AHL concentration, AHLs interact with LuxN switching its function from a kinase to a phosphatase, resulting in the loss of phosphate from the LuxU/O phosphor-relay cascade and, therefore, the loss of repression of bioluminescence (reveiwed by Cámara et al., 2002). LuxM/N-type QS systems homologous to the V. harveyi system have been identified in other marine Vibrio spp. including; V. fischeri (AinS/R) and V. anguillarium (VanM/N), where they occur along side LuxI/R-type QS systems (Milton et al., 2001; Hanzelka et al., 1999; Gilson et al., 1995).

1.1.3 Alkyl Quinolone Signalling

A second method of signal induced gene expression regulation employed by Gram-negative bacteria is the use of alkyl quinolone (AQ) signal molecules. AQ signalling was first discovered within the opportunistic pathogen P. aeruginosa. P. aeruginosa possesses two AHL-mediated QS systems, LasR/I and RhlR/I (Latifi et al., 1995). These systems operate in a hierarchical manner, with the LasR/I system acting as a transcriptional regulator of the RhlR/I system (Latifi et al., 1996). P. aeruginosa also possesses another signal system unrelated to homoserine lactone signalling molecules, but instead relies on the 4-quinolone group of small molecules commonly associated with antibacterial functions (Deziel et al., 2005; Pesci et al., 1999). The specific quinolone identified as the signalling molecule in P. aeruginosa was 2-heptyl-3-hydroxy-4-quinolone; this molecule was given the generic name of Pseudomonas Quinolone Signal (PQS) (Figure 1.5) (Pesci et al., 1999). PQS is initially produced from chorismate that is converted to anthanylate by an anthralinate synthase enzyme encoded by phnA

and phnB (Essar et al., 1990). Anthralinate is then further converted to the direct precursor of PQS, 2-heptyl-4-quinolone (HHQ) by PqsA, B, C and D which are transcribed from the pqs operon (Gallagher et al., 2002). The final stage in PQS biosynthesis is the conversion of HHQ to PQS by PqsH (Deziel et al., 2004; Gallagher et al., 2002). HHQ is also believed to be a QS signalling molecule as it is exported to the extracellular environment where it can be taken up by other bacterial cells (Figure 1.5) (Deziel et al., 2004). The biosynthesis of PQS appears to be connected with a multidrug efflux pump, MexGHI-OpmD. This pump prevents the build up of anthranilate, which is toxic to the cell (Aendekerk et al., 2002).

Figure 1.5. Chemical structures of the alkyl quinolones produced and used as signalling molecules by P. aeruginosa. 2-heptyl-3-hydroxy-4-quinolone (PQS) and 2-heptyl-4-quinolone (HHQ) (Figure sourced from Fletcher et al., 2007).

PQS is connected to both of the AHL-mediated QS systems in P. aeruginosa as pqsH is transcriptionally regulated by the Las system, as is pqsR (also known as msvR), the master regulator of the pqs operon (Deziel et al., 2005; Wade et al., 2005). The association of PQS with the Rhl system was established

through the elastase gene, lasB. PQS acts as a regulator of lasB, but only in the presence of a functioning Rhl system. It was therefore shown that PQS regulated the Rhl system in regards to the transcriptional activation of lasB (Pesci et al., 1999). As the Las system regulates POS, which, in turn, regulates the Rhl system, it can be stated that PQS provides a regulatory bridge between the two AHLmediated QS systems in P. aeruginosa (McKnight et al., 2000). Other studies have demonstrated that PQS is mainly, but not exclusively, synthesised during the stationary phase of the growth cycle and, therefore, PQS has no direct connection with sensing cell density (Diggle et al., 2003). Its function in up-regulating Rhl is therefore hypothesised to be in order to up-regulate Rhl-controlled genes, which would reduce stress on the bacterial cell during stationary phase (Diggle et al., 2003; McKnight et al., 2000). Although no clear molecular mechanisms of how PQS achieves regulation have been defined it has been shown that the protein PqsE acts as an effector of PQS-dependent virulence factors as pqsE mutants have a parental POS profile but also have attenuated virulence (Diggle et al., 2003; Gallagher et al., 2002). It is clear that PQS is a prominent factor in P. aeruginosa virulence as mutants which lack the ability to produce PQS have been shown to have reduced virulence in a Caenorhabditis elegans nematode model, due to an impaired ability to mediate poisoning via the production of cyanide (Gallagher and Manoil, 2001). PQS is also known to be involved in iron sequestration and as such can trigger an iron starvation response (Bredenbruch et al., 2006).

Although quinolone based signalling has mainly been studied in P. aeruginosa, other bacterial species such as Burkholderia pseudomallei,

Pseudomonas fluorescens and the marine bacteria Pseudomonas bromoutilis and Alteromonas sp. have been shown to produce quinolones, similar in structure to PQS (Appendix 1) (Fletcher et al., 2007; Diggle et al., 2006; Long et al., 2003). Alteromonas sp. strain SWAT5 was isolated from marine snow and has been shown to produce alkyl quinolones, specifically 2-n-pentyl-4-quinolinol, (PHQ). These quinolones appear primarily to be involved in antibiosis, inhibiting the growth of phytoplankton and algae. However, due to PHQs structural similarities to PQS it could play a role in QS related signalling (Long et al., 2003).

1.1.4 Quorum Sensing in Gram-Positive Bacteria

In Gram-positive bacteria QS is facilitated by a number of different autoinducing peptides. These peptides are produced through the cleavage of oligopeptides and interact with two-component receptor kinases on the cell surface which in turn initiate phospho-transfer cascades, ultimately leading to response regulators which alter gene expression (reviewed by Reading and Sperandio, 2006). The agr autoinducer produced by Staphylococcus aureus is referred to as an autoinducing peptide (AIP) and is produced due to the actions of gene products transcribed from the agr operon. In S. aureus, AIP-mediated QS regulates toxin and protease secretion (reviewed by Novick, 2003). Other peptide mediated QS systems have been found in Gram-positive bacteria a number of which are shown in Appendix 1.

1.1.5 AI-2 and LuxS, a Universal Signalling Language?

AHL-mediated signalling is unique to Gram-negative bacteria. Gram-positive bacteria utilise small peptides as autoinducers to facilitate QS (Dunny and Leonard, 1997). There is, however, a QS system that may transcend this barrier. This system uses a small, diffusible furanosyl borate diester referred to as autoinducer-2 (AI-2) (Figure 1.6) (reviewed by Xavier and Bassler, 2003). As with LuxM/N-type AHL-mediated QS systems, AI-2-mediated QS was discovered through the study of a marine bacterium, V. harveyi (reviewed by Xavier and Bassler, 2003).

AI-2 is produced as a result of the actions of an iron containing S-ribosylhomocysteinase enzyme LuxS (Zhu et al., 2003; Schauder et al., 2001). AI-2 is produced during the activated methyl cycle in which S-adenosylmethionine (SAM) is converted to S-ribosylhomocysteine (SRH). LuxS acts upon SRH to produce the AI-2 precursor molecule 4,5-dihydroxy-2,3-pentanedione, which in turn spontaneously converts to AI-2 (Miller et al., 2004). Within V. harveyi AI-2 regulates bioluminescence alongside the LuxM/N AHL system discussed earlier (Schauder et al., 2001). After reaching threshold concentration AI-2 binds its cognate receptor, a periplasmic protein termed LuxP which in turn interacts with a hybrid sensor kinase protein LuxQ, the signal is then relayed through the LuxU, LuxO phospho-relay system in the same manner as the LuxM/N system (Schauder et al., 2001).

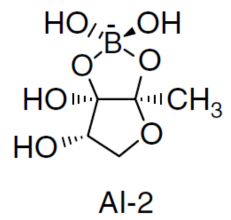


Figure 1.6. Chemical structure of AI-2 produced by Vibrio harveyi (Figure sourced from Williams et al., 2007).

In addition to the regulation of bioluminescence in V. harveyi, AI-2 mediated QS has been speculated to regulate gene expression in Vibrio cholera, E. coli and Salmonella typhurium (reviews by Xavier and Bassler, 2005; Miller et al., 2004; Cámara et al., 2002). The luxS gene is found in over 60 species of bacteria, from a multitude of different taxonomic groups including many Gram-positive bacteria and in many cases mutation of the luxS gene abolished AI-2 production. This has lead to the proposition that AI-2 is a universal bacterial signalling molecule (reviewed by Xavier and Bassler, 2003). In the last few years many studies detailing the effect of luxS deletion on an array of bacteria have been published (Stroeher et al., 2003; Lyon et al., 2001). However, it is difficult to interpret the resultant phenotypic changes as a defect in QS, or due to a defect in the activated methyl cycle of the cell, as most studies do not clearly separate the two. Sun et al. (2004) investigated the phylogenetic distribution of all genes involved in the synthesis of AI-2, as well as the corresponding lux genes making up the signal cascade necessary for the detection of AI-2 by analysing 138

complete genomes sequences (Sun et al., 2004). The signal transduction cascade for AI-2 was found to be restricted to Vibrio species. Indeed virulence factor production in the pathogenic marine bacterium Vibrio vulnificus is regulated by luxS (Sun et al., 2004; Kim et al., 2003). McDougald et al. (2003) demonstrated that AI-2 could induce starvation adaptation and stress response in the marine Vibrio, V. vulnificus and V. angustum S14. Signals produced from a range of other Vibrio species were also able to induce this response (McDougald et al., 2003). Thus, at least for Vibrio species AI-2 does function as a quorum sensing molecule, and interspecies AI-2 signalling between Vibrio species has been demonstrated. The role of LuxS and AI-2 with regards to bacterial signalling and QS therefore remains a hotly contested subject area.

1.2 Signal-Mediated Interactions

1.2.1 Inter and Intra-species Signalling

Outside laboratory conditions it is rare to find bacteria growing in single species cultures; for example in the marine environment a litre of surface seawater, on average contains 10⁹ bacterial cells from an estimated 1 million separate species (Curtis et al., 2002). In many environments, including the marine environment, bacteria are often found in multi-species biofilms as opposed to planktonic culture (Munn, 2004). A biofilm can be defined as an organised aggregate of bacterial cells adhered to a surface, often encased in a self produced extracellular polysaccharide matrix (Costerton et al., 1995). Bacteria growing in multi-species biofilms are able to detect and exploit signals produced by both members of their own species and members of different species within the

biofilm. Phenotypic response to exogenously produced signal was initially observed in Chromobacterium violaceium, which will produce a purple pigment, violacein, when exposed to C6-HSL produced by other bacteria (McClean et al., 1997). Another example is the ability of E. caratova to responded to exogenously produced AHLs using the AHL receptor ExpR2 (Sjöblom et al., 2006). Intraspecies signalling often mediates co-operative behaviour as is seen in the QS regulation of antimicrobial secondary metabolite production within a biofilm that disrupts the growth of competitive bacteria (Barnard et al., 2007). Examples of such behaviour include; the production of antimicrobial peptides by lactic acid bacteria such as Carnobacterium piscicola and Lactococcus lactis; production of phenazines by Pseudomonas chlororaphis; production of carbapenems by E. caratova; production of pyocyanin by P. aeruginosa and production of streptomycin by Streptomyces griseus (Barnard et al., 2007; Horinouchi, 2002; Quadri, 2002; Pessi and Haas, 2000; Pierson et al., 1994).

In contrast to intra-species signalling mediated co-operative behaviour, examples of competitive behaviour due to intra-species signalling have also been observed in biofilm communities. Naturally occurring P. aeruginosa QS mutants often gain advantages within Pseudomonas biofilms as they have the ability to either take advantage of signal without the imposed metabolic cost of producing signal, or fail to produce signal-mediated responce but take advantage of 'public goods' produced by other bacteria in response to signaling (reviewed by Diggle et al., 2007). Similar intra-species competitive behaviour has been observed in S. aureus. S. aureus strains can be grouped according to the type of autoinducing

peptides they produce; signal peptides produced by one group have been found to be inhibitory to strains from another group (Ji et al., 1997).

Signalling can also be utilised in a competitive or co-operative manner at the inter-species level. The use of QS to gain advantage over a different species has been observed in P. aeruginosa (An et al., 2006). When co-cultured with A. tumefaciens, AHL-producing P. aeruginosa effectively takes over the biofilm, this behaviour was not seen when co-culturing P. aeruginosa QS mutants with A. tumefaciens. It was therefore assumed that signalling was conferring a growth advantage to P. aeruginosa (An et al., 2006). This effect was seen to be reversed when co-culturing P. aeruginosa with indole producing E. coli strains as indole inhibits P. aeruginosa QS facilitating the survival of E. coli (Chu et al., 2012). In a mixed species biofilm, Burkholderia cenocepacia has the ability to detect and respond to the signals produced by P. aeruginosa. The detection of this signal was found to be unidirectional conferring B. cenocepacia with the ability to sense the presence of P. aeruginosa in a process termed 'QS eavesdropping' and is a clear example of inter-species competitive interaction (Riedel et al., 2001).

Examples of inter-species co-operation are less prominent in the literature than examples of competitive behaviour, however signal mediated inter species co-operation has been observed. Duan et al. (2003) showed an increase in lung damage caused by infection with P. aeruginosa co-cultured with avirulent oropharyngeal commensal strains in a rat lung model in comparison to infection with pure cultures of P. aeruginosa or oropharyngeal bacteria (Duan et al., 2003). Gene analysis showed that co-culturing up regulated many genes linked to

Pseudomonas virulence including many genes regulated by AI-2. As P. aeruginosa does not produce AI-2 this up regulation was attributed to AI-2 production by the oropharyngeal bacteria during the infection (Duan et al., 2003). In a Drosophila model, oropharyngeal bacteria, normally avirulent in the host, were shown to cause killing when co-cultured with P. aeruginosa (Sibley et al., 2008). This increase in virulence was again attributed to the up-regulation of virulence genes due to inter-species signalling (Sibley et al., 2008). A cooperative relationship modulated by QS signal molecules was also seen between Pseudomonas savastanoi, a pathogen of the olive tree (Olea europaea) and strains that form part of the plants epiphytic commensal population. AHLs produced by the epiphytic bacteria were able to restore virulence in QS mutants of P. savastanoi in an infection model. Additionally, disease caused by P. savastanoi was aggravated by the presence of the AHL-producing epiphytes (Hosni et al., 2011). In vivo the opportunistic pathogens P. aeruginosa and Burkholderia cepacia are often found together in mixed species biofilms (Riedel et al., 2001). In mixed species consortia formed in vitro from these two species B. cepacia demonstrates swarming motility, a phenotype not seen in pure culture (Venturi et al., 2010). In the same experiment B. cepacia was also shown to facilitate the entry of P. aeruginosa into an environment P. aeruginosa cannot tolerate individually (Venturi et al., 2010). Computer modelling attributed this co-operative behaviour to QS cross talk between the two species (Venturi et al., 2010).

1.2.2 Inter-Kingdom Signalling

In the majority of environments bacteria are in contact not only with other prokaryotic cells but also eukaryotic cells. These can be cells of a host that they are infecting, as part of a symbiotic relationship or cells co-inhabiting a particular environmental niche. As members of all biological kingdoms appear to possess the ability to carry out some form of signalling, a great deal of research has been focused on understanding if signalling mediates parasitic, symbiotic or commensal inter-kingdom interactions.

Examples of bacterial signalling mediating inter-kingdom co-operative behaviour have been observed in environments such as the human gut micro flora and the marine environment. Cell free supernatant from the probiotic strain Bacillus subtilis have been shown to stimulate the heat shock protein Hsp27 in CaCO2 colonic epithelial cells, protecting the epithelium from oxidative damage (Fujiya et al., 2007). The stimulation of hsp27 is thought to be mediated by QS signal peptides produced by the bacteria and additionally has the effect of providing the bacteria with a suitable environmental niche optimal for growth (Fujiya et al., 2007). Also in the gut the commensal organisms Bacteroides fragilis has been shown to produce a signal polysaccharide that has protective benefits to the host, suppressing harmful pro inflammatory cytokine responses to Helicobacter infection (Mazmanian et al., 2008). Although not an example of a non-bacterial organism responding to bacterial signalling molecules an example of inter-kingdom symbiotic behaviour involving bacterial QS can be found in the marine environment. This is the association between Vibrios and squid. As previously stated, a number of Vibrio species have the ability to bioluminesce

when growing in an increased population density. Squid species such as the bobtailed squid (Euprymna scalopies) have developed specific light organs that harbour populations of these bioluminescent bacteria. The bioluminescence provided by the bacteria gives the squid advantages with regards to either hunting for prey or avoiding predation. The bacteria are in turn provided with an ample source of nutrients from the animal enabling them to maintain a high population density, required for the facilitation of QS (Boettcher and Ruby, 1995).

Bacterial signalling molecules have also been shown to act as cues mediating responses from species outside the bacterial kingdom; such behaviour has been seen in the human body and in varying environmental niches. AHL signalling molecules such as 3-oxo-C12-HSL have been shown to effect human cancer cell lines through the regulation of Thymidylate synthase production (Dolnick et al., 2005). Additionally, AHLs produced by P. aeruginosa have been shown to mediate an inflammatory response in the Cystic Fibrosis lung, where there are specific receptors in host epithelial tissue that recognise these AHLs (Jahoor et al., 2008). The Alfalfa plant (Medicago sativa) also has the ability to detect AHL signalling molecules produced by bacterial plant pathogens and upregulate genes protecting against infection (Mathesius et al., 2003). The fresh water microalgae Chlamydomonas has been shown to produce riboflavin which may act as an AHL mimic (Teplitski et al., 2004). As such riboflavin has been shown to activate QS regulated genes in P. aeruginosa through activation of the LasR receptor (Teplitski et al., 2004). One of the more interesting examples of a eukaryotic organisms using AHL signalling molecules as cues to mediate behaviour has been observed in the green seaweed Ulva. AHLs have been shown to attract the plants motile zoospore during Ulva reproduction and, as the focus of this thesis, will be reviewed in section 1.4 (reviewed by Joint et al., 2007).

1.3 Quorum Sensing in the Marine Environment

1.3.1 Abundance of AHL-producing Bacteria within the Marine Environment

The marine environment is the largest ecosystem on the planet. Oceans and seas cover 71% of the Earth's surface, which amounts to a total area of $3.6\times10^8~\text{km}^2$ and an estimated total volume of $1.4\times10^{21}~\text{litres}$ (Munn, 2004). The marine environment is sub-divided into multiple zones depending on water column depth. Bacteria have been proven to be present in all of these zones but are most prevalent at the surface interface between water and atmosphere, the neuston (Munn, 2004). A study by Whitman et al. (1998) estimated that in the top 200 m of the worlds ocean there are 3.6×10²⁸ prokaryotic cells (Whitman et al., 1998). In spite of these vast numbers of prokaryotic life the marine environment can be considered a stressful environment as regards to microbial life; the overall nutrient concentration in seawater is relatively low, as is temperature, however salinity is high. As such, marine bacteria are highly adapted to the conditions found in the specific marine environmental niches they occupy (Munn, 2004). Owing to low nutrient conditions, physical processes within the oceans, viral lysis and the sheer size of the marine environment, bacterial populations do not generally increase over a population density of 10⁵ to 10⁶ cells ml⁻¹ (Munn, 2004). As QS is a population density-dependent process, the majority of QS regulated gene expression occurs at bacterial population with densities above approximately 10⁸ cells ml⁻¹ (Fuqua et al., 1996). Specialised niches do exist, however, which would promote the growth of dense microbial populations: the

colonisation of host animal light organs by the bioluminescent bacteria V. fischeri, and the role QS plays within this symbiosis is a well studied example (Lee and Ruby, 1994). AHL-producing bacteria are also found in abundance in the oceans and in a variety of different environments. These include; the aggregated organic and inorganic particles, commonly referred to as marine snow; algae; invertebrates such as corals and sponges and in biofouling communities (Tait et al., 2010; Tait et al., 2009; Mohamed et al., 2008; Wagner-Döbler et al., 2005; Taylor et al., 2004; Gram et al., 2002). Populations of between 10⁸ and 10⁹ cells ml⁻¹ have been found within marine snow and given that AHL-producing bacteria can be isolated from theses marine snow aggregates, this is a good indication that QS occurs within aggregated communities (Gram et al., 2002; Ploug, 2000). In addition to a significant number of marine Vibrio species shown to be producing QS signal molecules other signal molecule producing marine species include strains of Shewanella spp., Sulfitobacter spp., Flavobacterium spp., Glaciecola Tenacibaculum Roseobacter Flammevirga spp., spp., spp., Pseudoalteromonas spp. and Thalassomonas spp. (Romero et al., 2010; Tait et al., 2009; Huang et al., 2008; Milton, 2006). Interestingly, although QS signal molecule production has been shown to be abundant in marine bacteria, homologues of the genes currently implicated in both the synthesis and transduction of these signal molecules are absent in the genomes of many marine species, other than Vibrio spp. (Tait et al., 2009). This has lead to the prediction that there may be a number of yet to be discovered novel AHL-mediated QS systems in marine bacteria.

1.3.2 Constraints on AHL-mediated QS within the Marine Environment

The main constraints of AHL-mediated QS in the marine environment are AHL hydrolysis by seawater and inactivation of AHLs by bacteria possessing AHL inactivating enzymes. AHL signal molecules are known to undergo hydrolysis in an increased pH and/or temperature and acyl side chain length manner (Yates et al., 2002). AHLs with 3-oxo groups and short acyl side chains are particularly susceptible to hydrolysis but all AHLs will hydrolyse given suitable conditions. Seawater is typically pH 8.0-8.2 and therefore AHLs are not stable. In fact, AHL hydrolysis in seawater has been followed using HPLC; as expected AHL inactivation took place over a short time period, with AHLs hydrolysed by the alkaline pH of sea water. AHLs with shorter acyl side chains were inactivated faster than AHLs with longer side chains with AHL inactivation taking place according to first-order degridation kinetics (Figure 1. 7)(Hmelo and Van Mooy, 2009). Degridation rate coefficients for C6-HSL, 3-oxo-C6-HSL and 3-oxo-C8-HSL in natural seawater were 0.043 h^{-1} (± 0.003), 0.116 h^{-1} (± 0.005) and 0.148 h⁻¹ (± 0.002) (Hmelo and Van Mooy, 2009). Temperatures above 22°C also caused a marked increase in the hydrolysis of C6-HSL, 3-OH-C6-HSL and 3-oxo-C10-HSL in seawater (Hmelo and Van Mooy, 2009; Tait et al., 2005). Interestingly, natural seawater inactivated AHLs up to 57% faster than artificially prepared seawater. As protease K treatment of natural seawater reduced this rate of AHL inactivation to levels seen in artificially prepared seawater, it was proposed that natural sea water contains AHL-inactivating enzymes which increase the rate of AHL turnover beyond the level accounted for by hydrolysis due to high pH (Hmelo and Van Mooy, 2009).

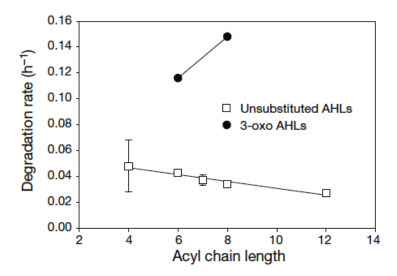


Figure 1.7. Degradation rate coefficients (± **SE**) **of various AHLs in natural seawater.** AHLs with shorter acyl side chains are inactivated at an increased rate in comparrsion to long chain AHLs. AHLs with 3-oxo substitutions are inactivated at an significantly increased rate to non-substituted AHLs (Figure sourced from Hmelo and Van Mooy, 2009).

Many marine bacteria have been shown to produce AHLs with long acyl side chains. This could be an artefact of rapid hydrolysis of short chain AHLs in the marine environment (Gram et al., 2002). Diffusion of AHLs from marine biofilms into the seawater is also effected by acyl side chain length with long chain AHLs possessing a slower rate of diffusion in comparison with AHLs with short acyl side chains (Tait et al., 2005). Rapid AHL inactivation due to the pH of seawater most likely constrains QS in the marine environment to either the close proximal environment to the cell producing the signal or to areas that are protected from seawater pH such as marine biofilms. In fact the majority of marine bacteria shown to be engaging in AHL-mediated QS have been isolated from either biofilm communities or aggregated particulates such as marine snow (Tait et al., 2009; Gram et al., 2002).

The evidence that AHL inactivation in non-sterile natural seawater takes place at an increased rate in comparison to sterile artificially produced seawater due to the actions of AHL inactivating enzymes poses the question of how widespread enzymatic AHL inactivation may be in the marine environment (Hmelo and Van Mooy, 2009). There are two broad classes of AHL inactivating enzymes; AHL lactonases and AHL acylases (Dong and Zhang, 2005). Lactonase enzymes inactivate AHLs by disrupting the homoserine lactone ring by hydrolysis of the lactone bond producing an N-acyl homoserine analogue (Figure 1.8) (Dong and Zhang, 2005). The first AHL lactonase to be identified, AiiA, was isolated from a Bacillus sp. strain found in soil samples (Dong et al., 2000). Since this discovery AHL lactonases have been identified in P. aeruginosa, Microbacterium testaceum, Ochrobactrum sp., and Rhodococcus sp. (Mei et al., 2010; Wang et al., 2010; Schipper et al., 2009; Uroz et al., 2008). Acylase enzymes however, inactivate AHLs by cleaving the acyl side chain from the homoserine lactone ring by hydrolysing the amide linkage producing a fatty acid and homoserine lactone (Figure 1.8) (Dong and Zhang, 2005). AHL acylases have been found to be produced by P. aeruginosa, Streptomyces sp. and Comamonas sp. (Uroz et al., 2007; Park et al., 2005; Huang et al., 2003). AHL inactivating enzymes have now been found to be produced by a wide range of marine bacteria including isolates from the Alphaproteobacteria (Hyphomonas sp.), Gammaproteobacteria (Shewanella sp. and Alteromonas sp.), Firmicutes (Oceanobacillus Actinobacteria sp.), (Rhodococcus erythropolis) and Bacteriodetes (Tenacibaculum discolour) phyla (Romero et al., 2011; Tait et al., 2009; Morohoshi et al., 2008).

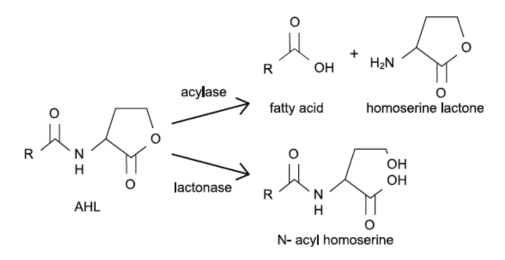


Figure 1.8. Enzymic mechanisms of AHL inactivation. AHL inactivating enzymes either act by hydrolysis of the homoserine lactone ring or by cleaving the acyl side chain from the homoserine lactone ring (Figure sourced from Defoirdt et al., 2004).

1.3.3 Milky Seas Phenomenon

Despite the low bacterial densities found within the water column, and the instability of AHLs in seawater, AHL-producing bacteria have been isolated from the open ocean (Bruhn et al., 2007). The function of AHL-mediated QS within these bacteria is currently unknown. However, QS within the marine environment has been shown to produce spectacular effects on a large scale. A phenomenon termed 'milky seas' has been recorded in shipping logs and fictional literature for over a hundred years (Nealson and Hastings, 2006). Milky seas are large patches of sustained luminescence on the surface of the ocean. This sustained luminescence differs from that caused by dinoflagellate blooms which 'sparkle' only when agitated or disturbed. The largest recorded instance of milky sea activity spanned an area of approximately 14,300 Km², (roughly the size of the US state Connecticut), and was detected using U.S. defence satellite sensor systems (Miller et al., 2005). Water samples were taken from areas of milky seas

and shown to contain V. harveyi bacteria in association with the microalga Phaeocystis (Lapota et al., 1988). Based on knowledge of V. harveyi and its interactions with microalgae a hypothesis for the milky sea phenomenon was put forward. The milky seas are thought to be attributed to the breakdown of algal blooms which causes a temporary, but substantial, increase in nutrient availability on the surface of the ocean. V. harveyi associated with the algae take advantage of the increased nutrient concentration, growing to a population size where signal molecule concentration reaches near sufficient amounts to activate luciferase gene expression, causing luminescence (Nealson and Hastings, 2006). The actions of grazing marine organisms preying on these bacterial and algal blooms does however reduce the bacterial densities to levels below that required for the sustained bioluminescence seen during milky sea events. This has lead to the question of whether microalgae to which the V. harveyi bacteria are associated with produce molecules that mimic AHL signalling molecules and therefore activate bioluminescence. Molecules that have this ability to mimic AHLs have been shown to be produced by the freshwater microalgal species Chlamydomonas reinhardtii where they can effect bacterial QS (Teplitski et al., 2004).

1.4 Ulva

1.4.1 The Genus Ulva

Species of seaweed belonging to the the genus Ulva are part of the plant phyla Chlorophyta (Green seaweed) first described by Carl Linnaeus in 1753 (Linné and Salvius, 1753). Species of Ulva are colloquially referred to as sea cabbage, gutleaf or grass kelp due to their spread leaf or tubular leaf morphology (Figure 1.9) (Burrow, 2001). According to the current World Register of Marine Species, there are 130 separate species of Ulva globally, 11 of which are indigenous to the UK coast line where they are found colonising the rocky shore environment within the intertidal zone (Guiry, 2012; Burrow, 2001). In older publications many species of Ulva are often classified as Enteromorpha spp., however these two Chlorophyta genera were amalgamated in 2003 (Hayden, 2003).



Figure 1.9. Examples of different species of Ulva endemic to the UK coastline. From top left; Ulva compressa, Ulva intestinalis, Ulva linza and Ulva lactuca.

Species of Ulva are economically important as they significantly contribute to marine biofouling; the colonisation of manmade marine surfaces such as sea walls and ships' hulls by algae and marine animals (Munn, 2004; Callow, 1986). Ulva growth on ships' hulls has the effect of causing hydrodynamic drag in the water which has the effect of increasing fuel usage by up to 10% (Munn, 2004; Callow and Callow, 2002). The US navy estimates this loss of fuel efficiency due to biofouling to cost over \$1 billion per year (Schultz et al., 2000). In addition to increasing hydrodynamic drag on ships hulls Ulva fouling can block waste effluent and cooling pipes (Figure 1.10) (Munn, 2004; Callow and Callow, 2002).



Figure 1.10. An example of Ulva spp. biofouling. Ulva spp. colonisation causing a blockage of an effluent pipe (figure sourced from Callow and Callow, 2002).

1.4.2 Ulva Zoospore Settlement

Ulva spp. reproduce either sexually by the production of biflagellate gametes or asexually through the production of quadriflagellate zoospores (Callow et al., 1997; Miyake, 1931). Ulva zoospores are naked, pear shaped, up to $7 \mu m$ in length and $5 \mu m$ in diameter (Figure 1.11) (Callow et al., 1997).

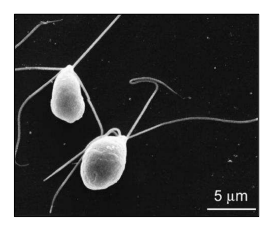


Figure 1.11. SEM of Ulva zoospores. Ulva zoospores are pear shaped and possess 4 flagella which confer motility (figure sourced from Callow and Callow, 2006).

In the UK Ulva spore release tends to take place during spring and summer, approximately 1-2 h after the high tide, potentially using photoperiod and temperature as sporulation cues (reveiwed by Maggs and Callow, 2001). During sporulation zoospore release rate has been estimated at 5.3 X 10⁵ spores per plant per day from the fertile tip of the Ulva thallus, the released zoospores will then rapidly settle on available substrata, germinate and grow into an adult plant (Callow et al., 2002; Maggs and Callow, 2001). The driving signal dictating that Ulva zoospores will settle on the surface of the rocks at the bottom of intertidal zone rock pools is light. Ulva zoospores are negatively phototactic and will therefore swim away from a light source (Callow and Callow, 2000). In the

natural environment, this negative phototaxis allows the zoospores to swim away from the surface of the seawater to the substrata at the bottom of rock pools (Callow and Callow, 2000). Ulva zoospore settlement to surface substrata then involves a transition from a free-swimming state to a non-motile adhered state (Callow et al., 1997). Adherence is a two stage process; primary adhesion to a surface is temporary, involving the zoospore testing the surface for optimal conditions (Callow et al., 1997). Upon selection of an optimum surface the zoospore secretes an N-linked, polydisperse glycoprotein from Golgi bodies in the apical end of the spore which cross links with cell wall matrix components forming a physical connection between the spore, adhesive and substratum (Callow and Callow, 2000). This process is referred to as secondary adherence (Callow and Callow, 2000). The strength of the adhesive holding the zoospore onto the surface has shown to increase with time. After 4 hours contact a pressure of up to 250 kPa is required to quantitatively dislodge the settled zoospores. Put in practical terms, to dislodge zoospores attached to the non-treated hull of a ship, the ship would need to travel at a speed of 42 knots in order to generate sufficient flow conditions. Such speeds are beyond the range of most vessels (Finlay et al., 2002). Observations of secondary zoospore adherence have shown that physical properties of a surface will influence where Ulva zoospores form permanent attachments, these properties include surface topology and surface chemistry (Callow et al., 2002; Callow et al., 2000). Ulva zoospores have been shown to preferably settle on rougher surfaces. Synthetic surfaces with micro valleys and pillars encourage increased zoospore settlement in comparison to a flat surface. Settlement was most pronounced where the diameters of the surface topography were identical to the dimensions of the zoospore. The reason for preferential selection of a rough surface is due to settlement on such surfaces being more energetically favourable (Callow et al., 2002). The wettability of a surface will also affect Ulva zoospore settlement (Callow et al., 2000). Surfaces with an increased advancing water contact angle (θ_{AW}) have a decreased wettability and are therefore hydrophobic (Zisman, 1964). Using surfaces of varied wettability formed from polymers with differing percentages of monomers containing either hydrophilic hydroxyl groups or hydrophobic methyl groups Callow et al. 2000 showed that as a surface becomes less wettable the number of zoospores preferentially settling on the surface increases (Figure 1.12) (Callow et al., 2000).

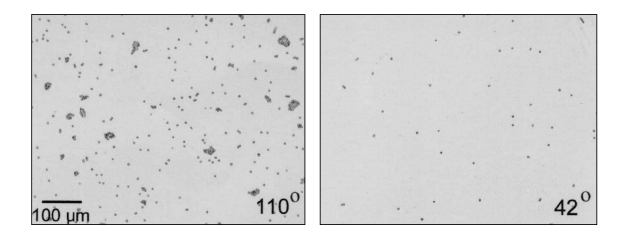


Figure 1.12. Effect of wettability on Ulva zoospore settlement. Increased zoospore settlement was observed on hydrophobic surfaces with high θ_{AW} (110°) in comparison to hydrophilic surfaces with low θ_{AW} (42°) (figure sourced from Callow et al., 2000).

Aside from the physical properties of a surface another key factor in primary Ulva zoospore settlement is the presence of bacteria which grow as biofilms on marine surfaces. Studies showed that Ulva zoospores were preferentially attracted to substrata covered with mixed species bacterial biofilms (Dillon et al., 1989). Additionally Joint et al. (2000) showed that zoospore settlement increases as bacterial density within the biofilm is increased, with

zoospore preferentially settling on microcolony structures within the biofilm (Joint et al., 2000). Patel et al., (2003) isolated bacteria from the surface of rocks colonised by Ulva spp. and showed that the bacteria present fell into three main groups; Gammaproteobacteria, Alphaproteobacteria and the Bacteroidetes (previously called the Cytophaga-Flavobacteria-Bacteroid group) (Patel et al., 2003). After producing single species biofilms of each strain, zoospore settlement assays showed zoospores predominantly settled on biofilms composed of bacteria composed of Vibrio spp. and Shewanella spp., part of the Gammaproteobacteria phylum (Patel et al., 2003). However, as this study was a culture based study it is not a completely accurate sample of the bacterial population growing on and around Ulva spp.. Tait et al., (2009) further assessed the bacterial population associated with Ulva spp. by producing a clone library of 16S rDNA from scrapings from of rocks colonised by Ulva spp.. This showed that the population was dominated by the Rhodobacteraceae, a family of Alphaproteobacteria and the Flavobacteriaceae, a family of Bacteroidetes (Tait et al., 2009).

The possibility that the zoospores were responding to AHL signals produced by the biofilm was tested using the fish pathogen Vibrio anguillarum as a model (Joint et al., 2002). V. anguillarum possesses a LuxR/I homologue termed VanR/I and a LuxM/N homologue termed VanM/N (Milton et al., 2001; Milton et al., 1997). V.anguillarum strains mutated to be defective in AHL production, (V.anguillarum NB10 ΔvanM, ΔvanI and ΔvanMI), were used in zoospore settlement assays. Increased zoospore settlement was evident on biofilms of the AHL-producing V. anguillarium wt and ΔvanI biofilms but not on biofilms of the ΔvanM (Joint et al., 2002). The VanM/N system exerts

hierarchical control over the VanR/I system, therefore the ΔvanI mutant only lacks the ability to produce 3-oxo-C10-HSL. Whereas, mutation of the VanM synthase, (Δ vanM), and mutation of both synthase genes, (Δ vanMI), also prevents 3-OH-C6-HSL production, completely rendering V. anguillarum AHL deficient (Milton, 2006; Milton et al., 2001; Milton et al., 1997). The evidence that zoospores do not settle on biofilms composed of the ΔvanM mutant but do settle on the Δ vanI mutant suggests that the zoospores are responding to the presence of AHLs regulated by the VanM/N system. Zoospore settlement was also disrupted in biofilms composed of the V. anguillarum ΔvanMI double AHL synthase mutant (Joint et al., 2002). As mutations in vanI/M may have impacted on multiple phenotypes within V. anguillarium which may have affected zoospore settlement, E. coli biofilms expressing recombinant vanI and vanM were subjected to zoospore settlement assays. Spores preferentially settled on the recombinant strains over wt E. coli biofilms, which do not produce AHLs (Joint et al., 2002). Additionally, synthetic AHLs were tested for their ability to enhance zoospore settlement. AHLs with acyl side chains of C6 to C14 were shown to increase zoospore settlement (Joint et al., 2002).

Tait et al (2005) repeated the zoospore attraction assays using the model organism V. anguillarum. In this study a wt strain (NB10), vanM mutant strain (NB10 ΔvanM) and a strain which expressed the AHL lactonase AiiA (NB10 aiiA) were tested for their ability to attract zoospores. Zoospores were shown selectively to settle on the wt but not on the QS mutant or the lactonase expressing strains (Tait et al., 2005). Zoospores were also shown to be attracted to biofilms composed of AHL-producing Aeromonas hydrophila and E. coli

expressing a range of recombinant AHL synthases. The link between signal molecule production and zoospore attraction was further investigated by analysis of zoospore response to specific AHLs. Zoospores were shown to be attracted to all biofilms that produced AHLs; however, increased settlement was seen in biofilms producing longer chain AHLs. Biofilms producing AHLs with hydroxyl or oxy groups did not cause significant differences in zoospore attraction in comparison to biofilms producing un-substituted AHLs (Tait et al., 2005). Increased settlement of zoospores on biofilms with longer chain AHLs is not surprising as longer chain AHLs are less likely to be hydrolysed by the alkaline pH of seawater, which would prevent zoospore attraction (Hmelo and Van Mooy, 2009; Tait et al., 2005). Continuous production of AHLs in a natural biofilm would also be a promoting factor for zoospore attraction (Tait et al., 2005). Tait et al, (2005) also showed that the zoospores are attracted to specific parts of the biofilm, namely the bacterial microcolonies (Figure 1.13). Using Gfp AHL biosensors, these microcolonies were shown to be areas of high AHL production, further confirming the hypothesis that Ulva zoospore were attracted to AHLs (Tait et al., 2005).

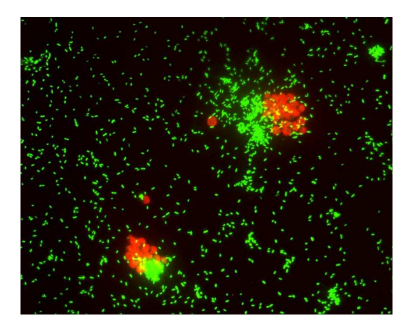


Figure 1.13. AHL production positively effects Ulva zoospore settlement. Ulva zoospores (red) were seen to preferentially settle on bacterial microcolonies within the biofilm, (dense green assemblages), where AHL concentration is at its greatest (figure sourced from Tait et al., 2005).

The initial work carried out by Joint et al., (2002) and Tait et al., (2005) connecting Ulva zoospore settlement with QS was carried out using a model marine organism V. anguillarium (Tait et al., 2005; Joint et al., 2002). Tait et al., (2009) also showed that AHLs are produced by bacteria growing on and around Ulva. Specifically AHL production was identified in bacterial species not previously known to be producers of signal molecules; these include species of Sulfitobacter sp., Glaciecola sp., Marinobacterium sp. and Shewanella sp. AHLs extracted from these bacteria and identified by mass spectroscopy were shown to possess acyl side chains of varying lengths containing both oxy and hydroxyl substitutions (Tait et al., 2009). Shewanella isolates were shown to produce AHLs during the late exponential stage of growth, around 12 h post inoculation. In stationary phase, (>18 h post inoculation), long chain AHLs could no longer be detected indicating the expression of an AHL degrading enzyme by this species.

Turnover of AHLs was found to influence Ulva zoospore attraction to a Shewanella biofilm. Zoospores were attracted to biofilms of medium density where AHL concentration is highest and not to high density biofilms where signal molecule degradation had taken place causing reduced concentration of AHL (Tait et al., 2009).

The process of how zoospores become associated with bacterial microcolonies producing AHLs was initially thought to be a chemotactic response, but this appears not to be the case. Wheeler et al. (2006) used V. anguillarium biofilms and synthetic AHLs and showed that signal molecules affect the swimming speeds of the zoospores. Zoospores exposed to wt V. anguillarum NB10 biofilms were shown to have reduced swimming speeds compared to those exposed to a biofilm composed of V. anguillarum NB10 ΔvanM and this result was replicated when zoospores were exposed to synthetic AHLs (Figure 1.14) (Wheeler et al., 2006). The presence of AHLs on a surface clearly caused a reduction is zoospore swim speed which had the effect of keeping the zoospores in close proximity to the AHL source, in turn promoting zoospore settlement. Chemokinesis was most profound when zoospores are exposed to longer chain AHLs, with 3-oxo-C12 HSL causing the greatest reduction in zoospores swim speed (Wheeler et al., 2006). Wheeler et al. (2006) proposed a model for zoospore settlement which states negative phototaxis is the driving factor to force zoospores down through the water column and that AHLs act as short range locator cues, modulating zoospore motility for favourable surface selection (Figure 1.14) (Wheeler et al., 2006).

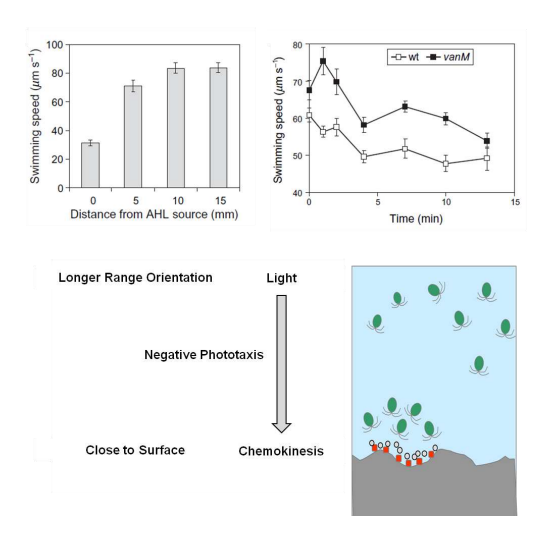


Figure 1.14. AHLs affect Ulva zoospore swim speed. Ulva zoospore swim speed was seen to increase as the distance from a synthetic AHL source increased (Top left) (sourced from Wheeler et al., 2006). Zoospores exposed to AHL producing V. anguillarium wt biofilms had reduced swim speed in comparison to zoospore exposed to AHL deficient V. anguillarium $\Delta vanM$ biofilms (Top right) (sourced from Wheeler et al., 2006). Diagram demonstrating the major cues which act on Ulva zoospore determining settlement (Bottom) (Provided by Dr. Karen Tait).

1.5 Project Overview and General Experimental Aims

This project has set about expanding our knowledge of bacterial/algal interactions in the marine environment. The interactions investigated by this project involve bacterial QS signal molecules and utilise both the macroalga Ulva and a number of species of microalgae to highlight the cross kingdom biological

implications of bacterial signalling in the marine environment. As such, the primary experimental aim of this thesis was to further address the relationship between the green seaweed Ulva and its indigenous epiphytic bacterial community. This was be achieved by isolating and characterising key microbial groups colonising the surface of Ulva spp. and investigating the extent of AHL-mediated signalling and QS within these groups. The importance of AHL signalling molecules to the germination process of Ulva zoospores was also be investigated, as well as examining whether microalgae produced compounds that either stimulated or inhibited AHL-mediated bacterial signalling.

This thesis contains the following results chapters:

- Chapter 3 characterises the composition of Ulva's epiphytic bacterial community, the signalling molecules produced by these bacteria and experimentally investigates the hypothesis that AHLs effect Ulva zoospore germination.
- Chapter 4 is an investigation of AHL signalling, QS and AHL degradation in two groups of bacteria found to be prominent in the Ulva epiphytic community, namely the Bacteroidetes and Shewanella.
- **Chapter 5** investigates whether microalgae produces compounds which could interefere with QS-mediated responses in bacteria.

Chapter 2

Materials and Methods

2.1 Culture Media and Growth Conditions

2.1.1 Culture Media

The media used to culture bacteria and microalgae is listed below. Solid media plates were routinely made with 1.5% w/v agar (Oxoid No.1), semi solid LB agar were made using 0.3% w/v agar (Oxoid No.1). Media was sterilised by autoclaving at 121°C for 15 min. Seawater for marine media was obtained from the Western English Channel and filtered through a 0.2 µm nitrocellulose filter prior to use.

- Actinomycete Isolation Media consisted of 22 g premixed Actinomycete
 Isolation Media (Difco) in 1 L d.H₂O.
- F/2 medium consisted of 1 ml 75 g L⁻¹ NaNO₃ solution, 1 ml 5 g L⁻¹ NaH₂PO₄ H₂O solution in 1 L 0.2 μm filtered seawater, autoclave. Post autoclave 1 ml F/2 trace metal solution (Appendix 2) and 0.5 ml F/2 vitamin solution (Appendix 1) was added (Guillard and Ryther, 1963).
- Luria Bertani medium (LB) consisted of 10 g tryptone (Oxoid), 5 g yeast extract (Oxoid) and 10 g NaCl in 1 L of d.H₂O (Sambrook and Russell, 2001).
- Luria-Bertani Lennox (Ylb) broth 10 g tryptone (Oxoid), 5 g yeast extract
 (Oxoid) and 5 g NaCl in 1 L of d.H₂O (Lennox, 1955)
- Marine medium (MB) consisted of either 47.4 g premixed marine broth
 (Difco) in 1 L of d.H₂O or 5 g peptone (Difco) and 1 g yeast extract
 (Oxoid) in 1 L 0.2 μm filtered seawater with no further additive (Zobell, 1941).

- R2A medium consisted of 18.2 g premixed R2A media (Difco) in 1 L d.H₂O.
- Seawater medium, made with 0.2 µm filtered seawater and 1.5% agar
 (Oxoid No1) medium was not supplemented with further vitamins or nutrients.
- Terrific Broth (TB) consisted of 12 g tryptone, 24 g yeast extract 4 ml 100% glycerol and 100 ml TB phosphate buffer (0.17 M KH₂PO₄ and 0.72 M K₂HPO₄) in 1 L d.H₂O (Sambrook and Russell, 2001).
- Tryptone Soy Broth (TSB) consisted of 30 g premixed TSB media (Oxoid)
 in 1 L d.H₂O.
- SOC Media consisted of 20 g tryptone, 5 g yeast extract and 0.5 g NaCl in 950 ml d.H₂O. After desolving these components, 10 ml of 250 mM KCl and 5 ml MgCl₂ were added and the volume adjusted to 980 ml with d.H₂O prior to autoclaving. Post autoclaving 20 ml 1 M sterile glucose solution was added to the media (Sambrook and Russell, 2001).

2.1.2 Culture Media Supplements

Media was supplemented when needed with the following antibiotics and chemicals at the working concentrations shown. Antibiotic and chemical stocks were prepared in accordance with (Sambrook and Russell, 2001), filter sterilised where appropriate and stored as per manufacturer's instructions: ampicillin (Amp) at 50-100 µg ml⁻¹, carbenacillin (Cb) at 100 µg ml⁻¹, chloramphenicol (Cm) at 30 µg ml⁻¹, gentamycin (Gm) 10 µg ml⁻¹, tetracycline (Tet) 10-20 µg ml⁻¹, isopropyl-

1-thio-β-D-galactopyranoside (IPTG) at 64 μg ml⁻¹ and 5-bromo-4-chloro-3 indolyl β-D-galactoside (X-Gal) at 64 μg ml⁻¹ (Sigma-Aldrich).

2.1.3 Bacterial Growth Conditions

Unless otherwise stated all marine bacteria, C. violaceum and Yersinia pseudotuberculosis strains were routinely cultured at 30°C. E. coli and P. aeruginosa strains were cultured at 37°C. All Bacterial cultures grown in liquid broth were agitated during growth at 200 rpm to allow adequate mixing of the culture and aeration during growth. Unless otherwise stated all bacterial liquid cultures were grown in 10 ml volumes of culture media. Bacterial growth was monitored by measuring the absorbance of a culture at an optical density (OD) of 600 nm using a 67 Series Spectrophotometer (Jenway).

2.2 Bacteria Strains and Plasmids

2.2.1 Bacterial Strains

During the course of this study, a number of bacterial strains were isolated from the marine environment or obtained from the culture collections at PML and the NCIMB. These strains are described in the results chapters of this thesis. Non-marine and marine strains used in various assays as either vectors, bio-reporters or model organisms are listed in Table

Table 2.1. Bacterial stains used in this study for cloning, maintaining vectors and as bio-reports

Strain	Description	Reference/ Source
Aeromonas hydrophila		
AH1-N	Wild type Aeromonas hydrophila strain	(Swift et al., 1999)
T 11		
E. coli		(2
DH5α	supE44 ΔlacU169 (Φ80 lacZ ΔM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1	(Sambrook and Russell, 2001)
JM109	recA1 supE44 endA1 hsdR17 gypA96 relA1 thi D (lac-proAB)	(Yanisch-Perron et al., 1985)
BL21	hsdS gal (λdts 857 ind1 Sam7 nin5 lacUV5-T7 gene 1)	(Sambrook and Russell, 2001)
C. violaceum		
CV026	cviI::mini-Tn5 derivative of ATCC 31532, Km ^R , AHL ⁻	(McClean et al., 1997)
P. aeruginosa		
PAO1	Wild-type, Nottingham strain	(Stover et al., 2000)
PAO1 rhlI::lux	Pseudomonas rhlI::lux fusion bio-reporter	Dr. James Lazenby, Uni of Nottingham
PAO1 lasI::lux	Pseudomonas lasI::lux fusion bio-reporter	Dr. James Lazenby, Uni of Nottingham
PAO1 rhlA::lux	Pseudomonas rhl::lux fusion bio-reporter	Dr. James Lazenby, Uni of Nottingham
PAO1 lasB::lux	Pseudomonas lasB::lux fusion bio-reporter	Dr. James Lazenby, Uni of Nottingham
Vibrio Strains		
V. anguillarium NB10	Wild type, serotype 01, clinical isolate from the Gulf of Bothnia	(Norqvist et al., 1989)
V. anguillarium	Double AHL synthase mutant	(Tait et al., 2009)
NB10 ΔvanI vanM		
V. coralliilyticus	Wild type Vibrio species isolated from diseased Pocillopora damicornis	(Tait et al., 2010)
LMG20984T		
V. tubiashii NCIMB 1337	Wild type Vibrio species isolated from Crassostrea virginica	(Tubiash et al., 1970)
Y. pseudotubercolosis		
YPIII	Y. pseudotuberculosis wild type strain	(Rosqvist et al., 1988)

2.2.2 Plasmids

Table 2.2. Plasmids used in this study. All plasmids were stored in d.H₂O at -20°C.

Plasmid	Description	Reference/ Source
pBBRIMCS-5	Broad host range vector containing lacZ, Cm ^r and a multiple cloning site.	(Kovach et al., 1994)
pCOLD	Protein expression vector	(Hayashi and Kojima, 2008)
pCOLDaacS	pCOLD expression vector harbouring aacS ORF as an Nde I/Eco RI fragment in frame with cspA prmotor and his tag	This study
pET3a	Cloning vector used as wt control in E. coli Ulva zoospore germination assay	(Tait et al., 2005)
pETVanI2	Plasmid expressing V. angullarium AHL synthase VanI	(Tait et al., 2005)
pGEM T Easy	Cloning vector with an f1 origin of replication containing lacZ, Amp ^r and a multiple cloning site.	Promega
pGEM::aiiA	pGEM vector containing Cm ^r and the AHL lactonase aiiA from Erwinia carotovora.	S. Atkinson, Uni of Nottingham
pME6000	Shuttle vector with a pBBR1 origin of replication containing Tet ^r and lacZ.	(Maurhofer et al., 1998)
pMH655	Plasmid containing Gm ^r and the Quorum Sensing Inhibitor System (QSIS).	(Rasmussen et al., 2005)
pMT01	pBBRIMCS1-5 vector containing aiiA from pSU18::aiiA.	This study
pMW47.1	Plasmid expressing P. aeruginosa AHL synthase RhlI	(Latifi et al., 1995)
pSB536	N-AHL bio-reporter composed of a pUC18 derived plasmid containing Amp ^r and a fusion of AyhR and lux promotor	(Swift et al., 1997)
	from Vibrio fisheri to the lux operon from Photorhabdus luminescens.	
pSB401	N-AHL bio-reporter composed of a pACYC184 derived plasmid containing Tet ^r and a fusion of luxR and lux	(Winson et al., 1998)
	promotor from Vibrio fisheri to the lux operon from Photorhabdus luminescens.	
pSB1142	N-AHL bio-reporter composed of a pACYC184 derived plasmid containing Tet ^r and a fusion of lasR and lasI	(Winson et al., 1998)
	promotor from Pseudomonas aeruginosa to the lux operon from Photorhabdus luminescens.	
pT7T3	Cloning vector used as wt control in E. coli Ulva zoospore germination assay	(Tait et al., 2005)
pT7T3luxI	Plasmid expressing Vibrio fisheri AHL synthase LuxI	(Tait et al., 2005)

2.3 Isolation of Marine Bacteria from Ulva spp.

Marine bacteria were isolated from rocks colonised by Ulva spp., the Ulva holdfast-rock interface and from the thallus of wild Ulva spp. Bacterial isolates were all obtained from either Wembury beach, Devon, UK (50°19'00'' N 4°05'03'' W), or Polzeath beach, Cornwall, UK (50°34'39'' N 4°55'03'' W).

Isolation of strains from the rocks colonised by Ulva spp. and from the Ulva holdfast/rock interface was carried out in accordance with the method described by Tait et al. 2009 taking scrapings from the rocks and/or Ulva holdfast, plating onto seawater agar (described in Section 2.1.1), and incubating for 15 days at 15°C. Resultant single colonies were then isolated onto MB agar (Tait et al. 2009).

Isolation of strains from the Ulva thallus was carried out by adapting the methodology described by Patel et al. (2003). Intialy Ulva thallus tissue was vortexted in 30 ml strerile Phosphate Buffer Solution (PBS) for approximately 3 min. The resultant PBS supernatant was decanted, serially diluted in sterile PBS to a range of dilutions factoring between 10⁻³ and 10⁻⁷ and plated onto either seawater agar, MB agar, Actinomycete Isolation Agar or R2A agar, (described in Section 2.1.1). The resultant isolation plates were cultured for 72 h at 30°C, before single colonies were indervidually isolated on to MB agar (Patel et al. 2003).

2.4 Preparation and Manipulation of DNA

2.4.1 Preparation of Chromosomal DNA

Chromosomal DNA from bacterial cell pellets obtained by centrifugation of stationary phase cultures or from cells collected from the surface of mature Ulva thallus material was extracted either using the DNeasy Blood and Tissue Kit (Qiagen) or Wizard Genomic DNA Purification Kit (Promega). Chromosomal DNA extracts using both kits were performed as per manufacturer's instructions. Final elution of DNA was in sterile molecular grade d.H₂O (Sigma-Aldrich) at 50°C.

2.4.2 Plasmid Extraction

Plasmid DNA was extracted from stationary phase cultures using a Qiaquick Mini Prep Kit (Qiagen) used as per manufacturer's instructions. Final elution of DNA was in sterile molecular grade d.H₂O (Sigma-Aldrich) at 50°C.

2.4.3 Cleaning of PCR Product DNA

DNA from PCR products was cleaned using either a Qiagen PCR Purification Kit (Qiagen) or a Wizard DNA Clean-Up System (Promega) used as per manufacturer's instructions. Final elution of DNA was carried out using sterile molecular grade d.H₂O (Sigma-Aldrich) at 50°C.

2.4.4 Restriction Digest

Restriction digests of plasmid DNA was carried out on ice in a total volume of 20 μ l using 0.2-1 μ g of DNA, 1 μ l of restriction endonuclease, restriction buffer at 1 X concentration and the remaining volume made up with sterile molecular grade d.H₂O. Restriction digests were incubated at 37°C for 2-3 h, unless otherwise stated. Restriction endonucleases were purchased with appropriate restriction buffer from Promega or New England Biolabs and were stored in accordance with the manufactures instructions at -20°C.

2.4.5 DNA Analysis by Agarose Gel Electrophoresis

DNA samples were analysed by electrophoresis using agarose gels composed of 0.8-1.5% ultra pure agarose (Invitrogen), in 1 X TAE buffer, (80 mM Tris-acetate, 19 mM EDTA at pH 7.6-7.8). To visualise DNA fragments ethidium bromide was added to the melted agarose gels at a final concentration of 10 µgml⁻¹. Prior to loading onto the gel 6 X loading buffer (Promega) was added to DNA samples. To judge DNA fragment sizes appropriate DNA ladder in 6 X loading buffer (Promega) was added to gels alongside samples. Gels were run in horizontal gel apparatus (Fisher Scientific, Bio-Rad) in 1 X TAE buffer at 90-120 V depending on the volume of the gel. DNA samples were visualised on the gels using a UV transluminator (UVP).

2.4.6 DNA Purification from Agarose Gels

DNA was purified from agarose gels by cutting the required fragments from the gel. Fragments were then extracted and cleaned using a Qiagen Gel Extraction Kit (Qiagen), in accordance with manufacturer's instructions. Final elution of DNA was carried out in sterile molecular grade d.H₂O (Sigma-Aldrich) at 50°C.

2.4.7 DNA Quantification

Quantification of DNA was carried out by micro spectrophotometry measuring absorbance at 260 nm using a NanoDrop 2000 (Thermo Scientific).

2.5 PCR

2.5.1 PCR Primers

PCR primers used in this study are listed in table 2.3. PCR primers used in this study were routinely designed 'by eye' looking at DNA sequence data. The degenerate primers used to amplify putative Shewanella AHL acylase genes were designed from the analysis of genomic DNA sequence located up-stream and down-stream of the putative AHL acylase ORFs (Sbal687_3701 and Sput200_0855 respectively) annotated in the genomes of Shewanella baltica OS678 (CP002383) and Shewanella putrafaciens. 200 (CP002457). The primer MTaacF was designed from conserved sequence in both genomes located 10 bp up-stream from the putative AHL acylase ORF start codon, includes the start codon plus the starting 6 bp of the putative AHL acylase ORF. The primer MTaacR was designed from conserved sequence in both genomes located 104 bp down-stream from the putative AHL acylase ORF stop codon. All primers were checked

for secondary structures and primer dimmer formation using the primer analysis software provided on the Eurofins oligonucleotide orders webpage. Oligonucleotide primers were synthesised either by Sigma-Aldrich or Eurofins and stored at -20°C in molecular grade water at a concentration of 100 μ M. Primers were diluted to a working concentration of 10 μ M prior to being added to PCR reaction mixes.

Table 2.3. List of PCR primers used in this study. Primer sequences are listed using the IUPAC 1-letter code abbreviations.

Primer	Sequence (5'-3')	Anneal	Additional Information	Reference
		Temp.		
96bfm	GAGTTTGATYHTGGCTCAG	53	Used to type marine bacteria. Amplified 16S rDNA corresponding to positions 9-1512 of the E. coli 16S rDNA sequence	(Muhling et al., 2008)
1152uR	ACGGHTACCTTGTTACGACTT	53	Used to type marine bacteria. Amplified 16S rDNA corresponding to positions 9-1512 of the E. coli 16S rDNA sequence	(Muhling et al., 2008)
341F	CCTACGGGAGGCAGCAG	53	Used to amplify 16s rDNA to produce clone library. Amplified 16S rDNA corresponding to positions 341-926 of the E. coli 16S rDNA sequence	(Muyzer et al., 1993)
907R	CCGTCAATTCMTTTGAGTTT	53	Used to amplify 16s rDNA to produce clone library. Amplified 16S rDNA corresponding to positions 341-926 of the E. coli 16S rDNA sequence	(Muyzer et al., 1993)
MTaacF	GATAATAATGAAATTC	43	Used to amplify Shewanella AHL acylase genes (see Section 2.5.1 text).	This study
MTaacR	TGTTAAWTTTTWACAAKTRYRT	43	Used to amplify Shewanella AHL acylase genes (see Section 2.5.1 text).	This study
MRIaacF	CTCCACCTACCGAACCTGAA	53	Upstream primer designed to amplify aac gene in Shewanella onediensis MR1	(Tait et al., 2009)
MRIaacR	TGCAGCATCAACTCAGTGGT	53	Downstream primer designed to amplify aac gene in Shewanella onediensis MR1	(Tait et al., 2009)
BlaacF	CATGGTGCTAGGTAATC	53	B1aac clone Sequencing Primer	This study
BlaacR	GCTGTCCAGCCTCTG	53	B1aac clone Sequencing Primer	This study
A1aacF	GCCGCGTATCGTTGG	53	Alaac clone Sequencing Primer	This study
A1aacR	CAGTGATAGCTGTCCAGC	53	A1aac clone Sequencing Primer	This study
NdeIB1aac-F	TTTTTTTTCATATGAAATTCAA CAAACTCGCGATCGCTATGGG	53	Forward primer used to amplify B1aac ORF which engineers Nde I site (highlighted in bold)	This study
B1aacEcoRI-R	TTTTTTTTCTGCAGTTATGGTT TTTGTAGTGTCAGCTCAGTCGTCG	53	Reverse primer used to amplify B1aac ORF which engineers Eco RI site (highlighted in bold)	This study
M13F	GTAAAACGACGCCAGT		General DNA sequencing primer	(Sambrook and Russell, 2001)
M13R	GGAAACAGCTATGACCATG		General DNA sequencing primer	(Sambrook and Russell, 2001)

2.5.2 General PCR Protocol Using Non-Proofreading DNA Polymerase

Amplification of DNA using a non-proofreading DNA polymerase was carried out using GoTaq DNA Polymerase (Promega). Reactions were made as per manufactures instructions to a total volume of 50 μl and contained a final concentration of 1 X GoTaq Flexi Buffer (Promega), 1 mM MgCl₂, 0.2 mM dNTPs, 1 μM forward and reverse primers, approximately 250 ng chromosomal DNA or 1 μl of boiled bacterial colony suspended in d.H₂O and 1 unit of GoTaq DNA polymerase. Amplification was carried out in a LabCycler thermocycler (SensoQuest) using an initial de-naturation step of 96°C for 2 min followed by 35 cycles of 95°C for 1 min, primer anneal temperature for 30 s and 72°C for 1 min per 1 kb of template sequence. Finally 1 step of 72°C for 10 min was used to remove the polymerase from the DNA.

2.5.3 General PCR Protocol Using Proofreading Enzyme DNA Polymerase

When required a proofreading DNA polymerase was used to carry out PCR, the enzyme used was Phusion High-Fidelity DNA Polymerase (NEB). Reactions were made to a total volume of 50 µl as per manufacturer's instructions and contained a final concentration of 1 X Phusion HF Buffer (NEB), 10 mM dNTPs, 0.5 µM forward and reverse primers, approximately 250 ng chromosomal DNA or 1 µl of boiled bacterial colony suspended in d.H₂O, 3% DMSO and 1 unit Phusion High-Fidelity DNA Polymerase. Amplification was carried out in a LabCycler thermocycler (SensoQuest) using an initial de-naturation step of 98°C for 30 s followed by 30-35 cycles of 98°C for 10 s, primer annealing temperature for 30 s and 72°C for 30 s per 1 kb of template sequence. Finally 1 step of 72°C for 5 min was used to remove the polymerase from the DNA.

2.6 Cloning

2.6.1 DNA Ligation

Purified DNA was ligated into vectors in a 1:3, 1:5 or 1:10 ratio of vector to insert depending on the concentration of each. 1-2 μ l T4 DNA Ligase (Promega) and 1 X DNA ligase buffer (Promega) was used for all ligation reactions. Total volume of ligation was between 10-20 μ l DNA ligations were carried out by initially cooling the reaction to 1°C followed by overnight incubation at 18°C.

2.6.2 Preparation of Electro-competent Bacteria

E. coli were made electro-competent by growing seed cultures to a mid log phase growth point (OD_{600} of between 0.6 and 0.8). Cells were then washed 3 times in sterile ice-cold 10% glycerol at 1 X, 0.5 X and 0.1 X volume respectively at 4°C. After washing cells were re-suspended in 50 μ l of sterile ice-cold 10% glycerol and stored at -80°C in 20 μ l aliquots (Sambrook and Russell, 2001). Electro-competent Shewanella cells were prepared by washing mid log phase culture 3 times 1 X, 0.5 X and 0.033 X volume of sterile ice-cold 1 M D-Sorbitol followed by re-suspending in 20 μ l sterile ice-cold 1 M D-Sorbitol (Myers and Myers, 1997).

2.6.3 Transformation of Bacteria via Electroporation

Prior to electroporation DNA was dialysed using 0.025 μ m millipore filters (Millipore Corporation). Electroporation was carried out in 2 mm electroporation cuvettes using 20 μ l of competent cells and 1-10 μ l of DNA. An electroporation pulse

of 2.5 kV was delivered using the BioRad Gene Pulsar connected to a BioRad pulse controller (BioRad Laboratories). The competent cells were recovered in 1 ml of either LB or SOC media and incubated for 1 h at 37°C before being plated onto LB media supplemented with appropriate antibiotics, X-Gal and IPTG (Sambrook and Russell, 2001). Electroporation of Shewanella was carried out in accordance with Myers and Mayers 1997 with using the same equipment as detailed above, recovery of electroporated cells was carried out at 30°C using media appropriate for the strain (Myers and Myers, 1997).

2.6.4 Chemical Transformation of E. coli

Chemically competent E. coli JM109 cells were purchased from Promega, stored and used in accordance with the manufactures instructions. 1-10 µl of DNA was added on ice to 40 µl aliquots of chemically competent E. coli JM109 cells. DNA and Cells were incubated on ice for 20 min, heat shocked at 42°C for 3 seconds and returned to ice for a further 2 min. Transformations were recovered in 1 ml of either LB or SOC media and incubated for 1 h at 37°C before being plated onto LB media supplemented with appropriate antibiotics, X-Gal and IPTG (Sambrook and Russell, 2001).

2.6.5. 16S rDNA Clone Library Construction

Epiphytic bacteria were obtained from the surfaces of Ulva thallus material collected from Wembury beach, UK (50°19′00′′N 4°05′03′′W) by prolonged vortexing in sterile phosphate buffered solution. Bacteria were pelleted by centrifugation and chromosomal DNA was extracted as described in Section 2.4.1. Using the extracted

chromosomal DNA as a template,16S rDNA corresponding to nucleotides 341-926 of the E. coli 16S rDNA sequence was amplified via PCR (see Section 2.5.2) using primers 341F and 907R (see Table 2.3). The amplified 16S rDNA was cloned into the pGEM T easy vector and transformed into E. coli DH5a via electroporation (see Sections 2.6.1 and 2.6.3). A total of 96 clones were selected for sequencing using the M13F and M13R universal sequencing primers (see Section 2.7.1).

2.6.6. Shewanella Genomic Libary Constuction and QSIS Screening

Shewanella strain P3 was grown to stationary phase and chromosomal DNA was extracted as described in section 2.4.1. P3 chromosomal DNA was digested separately using Eco RI, Hind III and Bam HI (Promega) for 12h at 37°C as described in section 2.4.4. The digested chromosomal DNA was ligated into shuttle vector pME6000 digested with the corresponding restriction enzyme (see Section 2.6.1). Ligations were transformed into competent E. coli DH5α pMH655 via electroporation (see Section 2.6.3). Subsequent transformant colonies were screened for AHL degrading genes by picking individual colonies onto LB media containing 15% sucrose, 20 μM synthetic 3-oxo-C12-HSL, ampicillin 50 μg ml⁻¹ and tetracycline 10 μg ml⁻¹ and then onto LB media containing ampicillin 50 μg ml⁻¹ and tetracycline 10 μg ml⁻¹ (adapted from method used by Rasmussen et al. 2005). Colonies that showed the same level of growth on media containing AHLs and sucrose as on media where AHLs and sucrose were omitted were selected for further study and therefore assayed for AHL degrading activity using the microtitre plate method detailed in section 2.9.5.

2.7 DNA Sequencing and Sequence Analysis Techniques

2.7.1 DNA Sequencing

Sequencing reactions were carried out at the University of Nottingham DNA Sequencing Facility, Queens Medical Centre, Nottingham. The non-radioactive Taq Dye Primer Cycle sequencing kit was used according to the manufacturers recommendations (Applied Biosystems), and the resulting sequencing reactions were analysed by 8% polyacrylamide gel electrophoresis using an Applied Biosystems 373A automated sequencer.

2.7.2 DNA Sequence Analysis

Initial DNA sequence analysis was carried out using DNA Baser (Heracle BioSoft) and Lasergene version 7.0 software package (DNAStar). Comparisons of DNA and protein sequences were performed using the Basic Local Alignment Search Tool (BLAST) algorithms available the NCBI's at web page (http://blast.ncbi.nlm.nih.gov/Blast.cgi). Phylogenetic analysis was conducted out using the method previously used by Tait et al. 2009. Alignment of clone library 16S rDNA sequences and the subsequent formation of phylogenetic trees was carried out using the Molecular Evolutionary Genetics Analysis (MEGA) tool (Tamura et al., 2011). The 16S rDNA alignment was carried out using a ClustalW alignment algorithm and the subsequent phylogenetic tree was produced from a neighbour-joining and bootstrap analysis performed with 1000 replications at a sequence similarity cut off of 97% (Tait et al. 2009). The species richness and the comparison of the Ulva thallus 16S rDNA clone library to the rocky shore and seawater bacterial populations was carried out

using the Distance-Based Operational Taxonomic Unit and Richness (DOTUR) program (Schloss and Handelsmen 2005).

2.7.3 Protein Sequence Analysis

Consensus sequences of LuxI-type AHL synthase, LuxR-type AHL response regulator, LuxM-type AHL synthase, LuxN type AHL response regulator and AHL inactivating enzymes were produced from alignments of published protein sequences imported from the NCBI protein database (See Appendices 4-8). Protein alignments were carried out on the MegAlign program, part of the Lasergene version 7.0 software package (DNAStar) using a ClustalW alignment algorithm. The resultant majority sequence was exported from MegAlign to the SeqBuilder, program part of the Lasergene version 7.0 software package (DNAStar) and used as consensus sequences in BLAST search of both Shewanella and Bacteroidetes genomes using the NCBI Genomes BLAST program.

Resultant hits from the BLAST searches were analysed for conserved structural/functional motifs using the Pfam sequence search facility provided by the Sanger Institute (http://pfam.sanger.ac.uk/) (Punta et al. 2012).

2.8 Protein Expression and Analysis

2.8.1 Protein Expression

Putative Shewanella AHL degrading enzymes were expressed using the pCOLD expression vector (Hayashi and Kojima, 2008). The ORF of aacS was amplified from a pGEMaacS template with primers NdeIB1aac-F and B1aacEcoRI-R using a proof reading DNA polymerase (NEB). The resultant PCR product was cleaned and digested with Nde I and Eco RI using the appropriate buffer for double digest, (Promega). The Nde I/ Eco RI aacS fragment was cloned into the pCOLD expression vector ensuring that it was in frame with the His tag and cspA promoter present on the pCOLD vector. The resultant pCOLDaacS construct was transformed into E. coli BL21 and grown to stationary phase in LB cb₁₀₀ at 37°C. BL21 pCOLDaacS and BL21 pCOLD cultures were used separately to seed 50 ml TB cb₁₀₀ to an OD₆₀₀ of 0.05 which were grown to an OD₆₀₀ of 0.4 at 37°C, left to stand at 15°C for 30 min before continuing to culture, with shaking, at 15°C for 24 h. Successful protein expression was indicated by a twofold reduction in the growth of BL21 pCOLDaacS, as measured by OD₆₀₀, in comparison with BL21 pCOLD (Hayashi and Kojima, 2008).

2.8.2 Cell Lysis and Protein Extraction

Cell lysis was achieved by sonication. Cultures were centrifuged at 10,000 rpm for 3 min to pellet the cells. Cells were re-suspended in sterile PBS and sonicated on ice at a frequency of 10-12 MHz for 10 min in repeated cycles 10 s sonication followed by 20 s on ice (Sambrook and Russell, 2001).

Unlysed cells were removed by centrifugation at 5,000 rpm for 5 min at 4° C. 500 µl of the resultant supernatant was removed and centrifuged at 13,000 rpm for 30 min at 4° C to pellet inclusion bodies and membranes. The soluble fraction was transferred to a fresh microtube and the insoluble fraction re-suspended in 1 ml sterile PBS both were re-centrifuged at 13,000 rpm for 30 min at 4° C. Protein fractions were prepared for analysis using sample buffer composed of 60 µl β mercaptoethanol to 1ml bromophenol blue. An equal volume sample buffer was added to the soluble fraction and the insoluble fraction re-suspended in 1 ml sample buffer, both were boiled for 5 min prior to analysis via SDS-PAGE (Sambrook and Russell, 2001).

2.8.3 SDS-Polyacrylamide Gel Electrophoresis

Proteins were analysed using a 12% polyacrylamide gel. Gels were made individually using glass casting plates and casting frames (Bio-Rad). Resolving gels were made from 4 ml 30% (w/v) acrylamide/bis (37.5:1); 2.5 ml 1.5 M Tris-HCl at pH 8.8; 100 μl 10% (w/v) SDS and 3.35 ml d.H₂O, set with; 100 μl 10% (w/v) ammonium persulfate and 15 μl TEMED. Stacking gels were made from 830 μl 30% (w/v) acrylamide/bis (37.5:1); 620 μl 1.5 M Tris-HCl at pH 6.8; 50 μl 10% (w/v) SDS and 2.5 ml d.H₂O, set with; 50 μl 10% (w/v) ammonium persulfate and 5 μl TEMED. Samples were loaded onto the gel along with appropriate protein size markers and run in 1 X SDS running buffer composed of 14 g Glycine, 3 g Tris and 1 g SDS in 1 L d.H₂O. All SDS PAGE was carried out in a Mini-Protean Tetra Electrophoresis System (Bio-Rad) and gels were run for 80 min at 150 v (Sambrook and Russell, 2001). Post electrophoresis gels were washed 3 X in d.H₂O and stained for 1 h with Simply Blue SafeStain (Invitrogen). Gels were de-stained on a shaking platform o/n in 100 ml H₂O.

2.9 Extraction and Analysis of AHLs

2.9.1 Production of Synthetic AHLs

Synthetic AHLs were produced by Alex Thurman at the School of Molecular Medical Science, University of Nottingham. AHL stocks were suspended in 1 ml HPLC grade methanol and stored at -20°C. Synthetic AHLs produced for this study and their abbreviations are listed at the front of this thesis.

2.9.2 Extraction of AHLs from Culture Supernatant

AHLs were extracted from culture supernatants using a method adapted from (McClean et al., 1997). Bacteria were grown to stationary phase in 20 ml cultures and centrifuged to pellet cells. Culture supernatants were decanted and acidified to pH 2.0 using 1 M HCl. AHLs were extracted from the acidified culture supernatant using 0.5 vol dichloromethane. The solvent layer was removed and dried either under nitrogen gas or via rotary evaporation or using a vacuum RC1022 centrifuge (Thermo Scientific). Extractions were then reconstituted in 2 ml acetonitrile, re-dried and stored as dried extracts at -20°C.

2.9.3 AHL Detection Using Bio-reporters

Detection of AHLs using bio-reporters was carried out using either a Petri dish based assay or a microtitre plate based assay using cell free supernatant extract prepared as described above. AHL bio-reporters used were the lux-based E. coli pSB536, pSB401 and pSB1142 and the pigment-based CV026 (Winson et al., 1998; McClean et al., 1997; Swift et al., 1997). The Petri dish based assay was conducted

using the method as described in McClean et al. 1997. Luminescence as a result of bioreporter activation was recorded using a luminographer (Hamamatsu); activation of
CV026 produced a purple pigment (McClean et al., 1997). Microtitre plate based AHL
detection assays were carried in accordance with the method described by Tait et al.
(2009) using the lux-based bio reporters E. coli pSB536, pSB401 and pSB1142
(Winson et al., 1998; Swift et al., 1997). A black clear bottomed 96 well microtitre
plate (Greiner Bio One) was used for all assays with 50-100 µl of cell free supernatant
extract dried to each well assayed. Bio-reporter activation was either measured at a set
point after 3 h incubation at 37°C with luminescence being measured at 450 nm using a
Berthold MITHRAS microtitre plate reader, or tracked over a 12 h incubation period
with bioluminescence and OD being measured every 30 min using a Infinite 200 PRO
series microtitre plate reader (Tecan). Intensity of bio-luminescence was calculated in
Relative Light Units (RLU) (Tait et al., 2009).

2.9.4 AHL Detection via LCMS

AHL detection in bacterial extracts via Liquid Chromatogarphy Mass Spectroscopy Liquid Chromatography was conducted by Mary Bruce at the Department for Molecular Medical Sciences at The University of Nottingham. Liquid Chromatography was carried out using the Agilent 1200 series HPLC, comprising degasser, binary pump, column heater and autosampler using the methodology as previously described by Yates et al. 2002 (Yates et al., 2002). Mass Spectroscopy was carried out using the Bruker HCT Plus ion trap in multiple reaction mode (MRM) and Hystar software, ions were introduced using positive ion electrospray from the Agilent HPLC system. Using the Smartfrag option on the software, the trap was set to isolate from full scan and then fragment ions at m/z 172.1 and 190.1. The ion charge control

was used to prevent charge overload in the trap. The instrument was optimised using the smart parameter setting for m/z 172. The monitored mass range was 50 - 250 m/z. Data analysis was carried out using the Bruker Data Analysis version 3.3 package. Extracted ion chromatograms (EIC) of m/z 102.1 and 172.3 were produced from the positive ion MSMS of m/z 172.1 and 190.1 respectively. Retention times and peak spectra were matched to the 1 μ M standard (x6) injected at the beginning of each method set. Injections of sample solvent were also monitored to assess carryover (Ortori et al., 2007).

2.9.5 AHL Inactivation Assay

AHL inactivating activity was measured using the lux-based bio-reporters E. coli pSB536, pSB401 and pSB1142 (Winson et al., 1998; Swift et al., 1997). Strains being assayed were grown overnight to stationary phase along with a non AHL-inactivating control. Cultures were then diluted to an OD_{600} of 0.01 in media seeded with appropriate AHL at a concentration of 1.24 μ M and grown to stationary phase. Cultures were extracted in either DCM or acidified ethyl acetate as described in section 2.8.2 and residual AHL concentration measured using an appropriate bio-reporter using the microtitre plate assay as described in section 2.8.3.

2.10 Microalgae Assays

2.10.1 Microalgal Strains and Growth Conditions

In order to investigate microalgal/bacterial interactions three separate axenic cultures of microalgae were acquired from the Culture Collection of Algae and

Protozoa (CCAP, Oban, UK). These cultures were; Nannochloropsis oculata (Eugstigmatophyte) (CCAP 849/1); Tetraselmis suecica (Prasinophyte) (CCAP 66/8) and Isocrysis galbana (Haptophyte/ Prymnesiophyceae (CCAP 927/1). All three species were routinely cultured from a starting inoculate of 10 ml of stationary phase culture in 1 L F/2 media, (Guillard and Ryther, 1963). In order to maintain an axenic state all microalgal cultures were supplemented with an antibiotic cocktail of ampicillin (200 μg ml⁻¹), streptomycin (100 μg ml⁻¹) and kanamycin (50 μg ml⁻¹). In addition, all three cultures were viewed under a microscope at 100 X magnification, confirming that all three cultures lacked any bacterial contamination. During growth microalgal culture were incubated at 18°C in the close proximity to a fluorescent white light source with a 16 h light, 8 h dark cycle. Microalgal growth was monitored by measuring the absorbance of a culture at an Optical Density (OD) of 600 nm using a 67 Series Spectrophotometer (Jenway).

2.10.2 Extraction of Microalgal Cultures

QS mimics and QS inhibitory compound(s) were obtained by solvent extraction of microalgal cultures. Solvent extraction was carried out 2 X using 0.5 vol dichloromethane in accordance with the method previously detailed in Section 2.9.2 for the extraction of AHLs from the cell free supernatants of bacterial cultures. For experiments using the lux-based E.coli AHL bio-reporters, separate extracts from individual cultures were used. Experiments using the P. aeruginosa transcriptional fusion bio-reporters and marine bacteria protease assays used pooled extracts from 3 X 1 L microalgal cultures. This allowed a greater number of experiments to be performed and overcame any inconsistency in the resultant quorum-quenching activity between extracts from separate cultures.

2.10.3 AHL Mimic Assay

The presence AHL mimic compounds in the extracts from microalgal cultures was assayed for using the microtitre plate assay, based on the method described by Tait et al. (2009) for the detection of AHLs in marine bacterial culture extracts and is outlined previously (Section 2.9.3) (Tait et al., 2009). A volume 30 µl of microalgal culture extracts was used in all assays, co-cultured with 300 µl of the lux-based AHL bio-reporters E. coli pSB536, pSB401 and pSB1142 (Winson et al., 1998; Swift et al., 1997).

2.10.4 Quorum-quenching Activity in Microalgae

Quorum-quenching activity by microalgal species was measured using the lux-based AHL bio-reporters E. coli pSB536, pSB401 and pSB1142 (Winson et al., 1998; Swift et al., 1997) and lux-based P. aeruginosa transcriptional fusion reporters PAO1 lasI::lux, PAO1 RhII::lux, PAO1 LasB::lux and PAO1 RhIA::lux (un-published). All microalgal quorum-quenching assays were carried out using a 96 well microtitre plates based technique adapted from the AHL detection assay, (Tait et al., 2009), described in section 2.9.3. When using the E. coli AHL bio-reporters 1.24 µM C4-HSL, C6-HSL and 3-oxo-C12-HSL were dried to wells of black clear-bottomed 96 well microtitre plate (Greiner Bio One). In addition to the AHLs a volume of solvent extract from the microalgal species representative of 20 ml original culture or an equal volume of solvent extract from F/2 medium and HPLC grade solvent was also dried to the 96 well microtitre plate. When using the P. aeruginosa transcriptional fusion bio-reporters the synthetic AHLs were negated from the assay as these reporters have the ability to produce both C4-HSL and 3-oxo-C12-HSL, the volume of microalgal extract, F/2

medium extract and solvent control was however the same (a volume representing 20 ml original culture). In all assays 300 µl of appropriate bio-reporter was added to each well. Each experimental condition was assayed in triplicate. Assay plates were incubated at 37°C in an Infinite Pro200 microtitre plate reader (Tecan) over a 12 h period. Absorbance at OD₆₀₀ and bio-reporter luminescence at 100 nm in each well was measured every 30 min. Bio-reporter activity over the 12 h period was recorded as a function of luminescence (RLU)/OD₆₀₀, these measurements were used to calculate total bio-reporter luminescence over the 12 h period from each extract assayed.

2.10.5 Fractionation

In order to initiate the process of identifying compound(s) responsible for observed quorum-quenching activity by microalgae, extracts from stationary phase microalgal cultures were fractionated. Fractionation was performed by initially resuspending the concentrated pooled extract from 3 X 1 L cultures of each microalgal species in 1 ml methanol. The re-suspended extract was then diluted 10⁻¹ in a volume of 10 ml methanol. Subsequently the diluted extracts were applied to Oasis HLB solid phase extraction cartridges (Waters) by running the extracts through the cartridge. The extract was allowed to elute through the cartridges by gravity and the run-through collected. A methanol:d.H₂O dilution series was prepared, (10% methanol – 100% methanol, in 10% increments, 10 ml volumes). Each methanol fraction was applied to the HLB solid phase extraction cartridges, in order, stating with the 100% methanol fraction. Each of the 10 resultant fractions eluted from the cartridges were concentrated via rotary evaporation, re-suspended in 1 ml 100% methanol and assayed with the P.

aeruginosa transcriptional fusion bio-reporters using the method described previously in Section 2.10.3.

2.10.6 Protease Assay

The fractionated extracts from stationary phase microalgae cultures were assayed for effect on protease production in the marine pathogens Vibrio anguillarium, Vibrio coralliilyticus, Vibrio tubiashii and Aeromonas hydrophila (See table 2.1). 10 ml o/n cultures of each bacterium were diluted 1:50 and 150 µl of diluted culture were added to 15 µl of each solvent fraction that had been previously air-dried to remove all traces of methanol within microtubes. The subsequent cultures were grown for 16 h, centrifuged and the supernatant used in protease activity assays, carried out, black clear bottomed 96 well microtitre plate (Greiner Bio One) using the Protease Fluorescent Detection Kit (Sigma), as per the manufactures instructions. Controls included media blank and solvent negative control. Protease activity was calculated as the quantity (ng) of trypsin released by the degradation of casein labelled with fluorescein isothiocyanate (FITC). Fluorescence was measured as per as per the manufactures instructions (Protease Fluorescent Detection Kit - Sigma) using a Infinite Pro200 microtitre plate reader (Tecan) at 485 nm excitation and 535 nm emission.

2.11 Ulva Zoospore Germination Assay

2.11.1 Biofilm Growth

All biofilms were grown on 25 mm X 25 mm glass cover slides within biofilm incubation chambers. Cover slides were placed vertically in holders within the biofilm

incubation chambers, submerged in 300 ml filtered seawater and sterilised. Bacteria were grown to stationary phase in appropriate culture media, re-suspended in sterile filtered seawater and adjusted to an OD_{600} of 1.0 prior to inoculation of sterile biofilm incubation chambers. Biofilm incubation chambers were inoculated separately with 0.5, 1, 2 and 4 ml volumes of bacteria. Marine biofilms were grown for 48-72 h at room temperature ($\approx 22^{\circ}$ C). Biofilms of E. coli strains were grown in 70% filtered sea water for 24 h in order to reduce osmotic stress on the bacteria by salinity (Tait et al., 2005).

2.11.2 Biofilm Density Measurement

Biofilm density was determined as per the method described by Joint et al. 2002 with microscope image analysis at 40 X magnification, using a Reichert Jung Polyvar microscope and an Optronics Magna Fire SP camera. Image ProPlus Version 5 imaging software was used to measure the area of bacteria within the field of view. The density of bacteria per unit area of slide (L) was determined by L = x/A, where x is the total area covered by bacteria and A is the area of the total field of view (Joint et al., 2002).

2.11.3 Ulva Zoospore Release

Ulva zoospore release was carried out in accordance with methodology of Callow et al. 1997. Mature Ulva spp. thallus material was collected from the rocky shore environment at Wembury beach, Devon, UK (50°19'00''N 4°05'03''W), at 1-2 h post high water during the period of a spring tide. The Ulva thallus material was dried between paper towels over night at a constant temperature of 15°C. After drying the thallus apical tips were cut into sterile filtered sea water on ice and agitated to stimulate

Ulva zoospore release. After which the thallus material was discarded and the zoospore suspension was exposed to an overhead light source for approximately 15 min in order to select for healthy, active zoospores by exploiting their negative photo-tactic phenotype. Excess sea water and non active zoospores were discarded and the zoospore suspension containing the active zoospore was diluted to an OD_{600} of 0.5 using sterile filtered sea water (Callow et al., 1997).

2.11.4 Ulva Zoospore Germination Assay

15 ml of the final Ulva zoospore suspension was used to inoculate biofilms or sterile glass slides within 90 mm Petri dishes. Ulva zoospores were settled onto the glass slides or biofilms in the dark for 1 h (Tait et al., 2005). Post settlement zoospore slides were transferred from the 90 mm Petri dish to sterile 60 mm Petri dishes and submerged in 10 ml sterile filtered seawater, 70% sterile filtered sea water was used with zoospore assays using E. coli biofilms. For assays using synthetic AHLs, the appropriate AHL was added to the separate dishes at final concentrations of 0.5, 5, 10, 20, 30, 40 and 50 μM. Zoospore slides were incubated at 18°C in proximity to a light source with a 16 h light, 8 h dark cycle for 24 and 48 h. At the 24 and 48 h time points zoospore slides were removed to sterile 60 mm Petri dishes, fixed with 2% (v/v) glutaraldehyde and stained with dilute carbol fuschin.

Ulva zoospore slides were viewed at 10 X magnification using a Reichert Jung Polyvar microscope with attached Optronics Magna Fire SP camera. The lengths of 300 Ulva zoospores and germlings were measured from randomly selected images of each

experimental condition using Image ProPlus Version 5 imaging software. Measurements were exported to MS Excel and average zoospore/germling length was calculated. In all Ulva germination experiments a successfully germinated Ulva zoospore was defined as having a length equal to or greater than $15 \, \mu m$.

2.11.5 Statistical Analysis

Statistical analysis of Ulva zoospore germination assays was carried out by performing ANOVA and PERMANOVA tests using statistical analysis software packages Minitab 16 (Minitab) and PRIMER 6 (PRIMER-E).

Chapter 3

The Effect of Bacterial Signal

Molecules on Ulva Zoospore

Germination

3.1 Introduction

3.1.1 Effects of Marine Bacteria on Ulva Zoospore Settlement

Ulva spp reproduce either via the production of gametes or the production of motile quadriflagellate zoospores (reveiwed by Maggs and Callow, 2001). The Ulva zoospore body is pear shaped, approximately 5-8 µm in length and 5 µm in diameter at the widest point (Callow et al., 2002). Ulva spp sporulation takes place during the summer period and in most instances approximately 1-2 h after the high tide. During sporulation Ulva zoospores are released from the apical tips of the Ulva thallus, zoospores then proceed to adhere to surfaces in their proximal environment (Callow et al., 2000). Adherence of zoospores to a surface is a two stage process. Initial adhesion to a surface is temporary and involves the zoospore testing the surface for optimal conditions followed by the secretion of glycoprotein from Golgi bodies in the apical end of the zoospore. Upon identifying an optimum surface, secondary adherence is permanent (Finlay et al., 2002; Callow et al., 2000). Following permanent adhesion Ulva zoospores germinate and mature into an adult plant (Callow et al., 2000). Ulva zoospores utilise their flagella in order to travel from the site of sporulation on the Ulva thallus to the surface they eventually adhere too. Flagella driven motility in Ulva zoospores is primarily governed by negative phototaxis, ensuring that the zoospores are oriented and move towards a solid substrate such as the rock surface which provides optimal substratum for adherence and germination (Callow and Callow, 2000). Factors acting as important cues for the selection of an optimal surface for zoospore adherence include the topology of the surface, wettability, the surface chemistry and the presence of a bacterial biofilm (Callow et al., 2002; Callow and Callow, 2000; Dillon et al., 1989).

Further investigation of the relationship between Ulva zoospore settlement and bacterial biofilms revealed that Ulva zoospores preferentially settle on bacteria producing AHL signal molecules (Tait et al., 2009; Tait et al., 2005; Joint et al., 2002). In investigating the method by which AHLs cause preferential settlement of Ulva zoospores Wheeler et al. (2006) demonstrated that in the presence of AHLs zoospore motility is significantly reduced, causing zoospores to cluster and eventually settle at areas of increased AHL concentration (Wheeler et al., 2006). In addition to AHLs having an effect on the settlement of Ulva zoospores, AHLs have also been shown to affect sporulation in other species of algae. High concentrations of C4-HSL were shown to up-regulate sporulation in Acrochaetium spp. This effect was inhibited by a number of halogenated furanone compounds, AHL structural homologues produced by algae such as Delisea pulchra (Weinberger et al., 2007; Manefield et al., 1999).

3.1.2 Bacterial Population Associated with the Rocky Shore Environment

Ulva spp. grow on the rocks present in the intertidal zone of the UK coastline which is an environment heavily colonised by marine bacteria which form biofilms on the surfaces of these rocks (Munn, 2004). A 16S rDNA phylogenetic analysis of the intertidal rocky shore environment has revealed the presence of a diverse microbial population (Tait et al. 2009). The rocky shore population was found to be dominated by Alphaproteobacteria and Flavobacteria; however other bacterial groups such as the Sphingobacteria, Deltaproteobacteria, Gammaproteobacteria, Bacilli, Cyanobacteria and Verrucomicrobiae were also shown to be present (Figure 3.1). It was possible to isolate bacteria from a number of these groups, (Flavobacteria, Deltaproterobacteria, Gammaproteobacteria and Bacilli) and grow them in laboratory culture, however

culture based isolations did not reflect the population as defined by molecular methods (Tait et al., 2009).

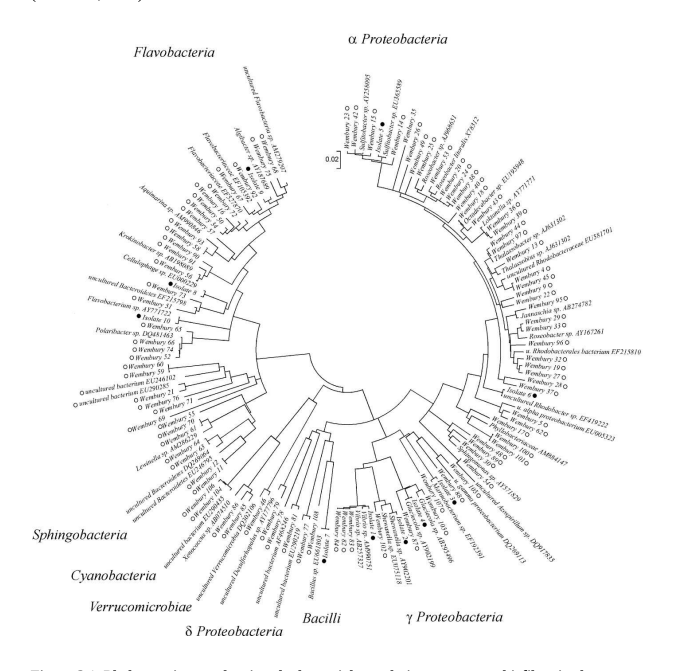


Figure 3.1. Phylogenetic tree showing the bacterial population present on biofilms in the rocky shore environment where Ulva spp. are found growing. The phylogenetic tree was produced from an alignment of 16S sequences from clones (⋄) and isolates (♠) taken from rocky shore biofilms. The reference strain sequences were obtained from Genbank (Figure sourced from Tait et al., 2009).

3.1.3 Bacterial Effects on Ulva spp. Growth and Morphology

In addition to their effect on Ulva zoospore settlement, bacteria appear to have a profound impact on the germination of the Ulva zoospore and growth of the Ulva plantlet. In 1980, Provasoli and Pintner showed that when grown axenically, Ulva lactuca assumed a different morphology to when grown in the presence of marine bacteria. The morphology observed in axenic culture is an atypical 'pin cushion' growth as opposed to wild type foliaceous growth (Provasoli and Pintner, 1980). This phenomenon of aberrant morphological differences in U. luctuca cultures when grown axenically has additionally been observed in Ulva linza, Ulva pertusa and Ulva compressa (Marshall et al., 2006; Nakanishi et al., 1996). Adding back bacterial strains to axenic U. linza cultures has been proven to restore wild type morphology (Figure 3.2).

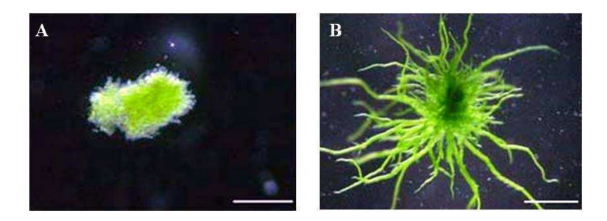


Figure 3.2. Images comparing the aberrant morphology associated with axenic Ulva culture and the wild type morphology. (A) 28 day old axenic culture of U. linza, (B) 28 day old initially axenic culture of U. linza inoculated with marine bacterial strain UC19 (Cytophaga sp.). The scale bars represent 1 mm (Figure sourced from Marshall et al., 2006).

Of 1555 bacterial strains isolated from U. pertusa, 676 strains (41%) showed morphogenesis activity (Nakanishi et al., 1996). In U. linza, a number of different strains isolated from the algae not only induced morphological change in axenic cultures but also increased the relative growth rate of the plant (Marshall et al., 2006). From 38 unique isolates, 5 affected both morphology and growth of U. linza when added to axenic cultures. Phenotypic analysis of marine bacterial strains that affected either morphology or growth of U. linza showed that they grouped to the Proteobacteria, Bacteroidetes and Gram positive cocci, and that the bacterial effect on U. linza growth was independent of bacterial phylogeny (Marshall et al., 2006). The effect of bacteria on the morphology of green alga is not only restricted to the genus Ulva. Monostroma oxyspermum cannot differentiate without the presence of a bacterial population and approximately 1% of 1159 bacterial strains isolated from green alga were shown to induce morphological changes in M. oxyspermum (Matsuo et al., 2003; Tatewaki et al., 1983).

As yet there is no scientifically proven mechanism established as to why bacteria affect the morphology of Ulva spp. and other green algae; however there are reports of marine bacteria producing plant growth regulators and vitamins which may affect the morphological differentiation of algae (Croft et al., 2006; Maruyama et al., 1986). A primary example of a plant hormone produced by marine epiphytic bacteria that influences growth morphology is thallusin, shown to effect Ulva intestinalis, U. pertusa and M. oxyspermum (Matsuo et al., 2005). Additionally other algal growth hormones such as cytokinin-type hormones, auxin-type hormones and indole-3-acetic acid have been shown to be produced by marine bacteria (Bradley, 1991; Maruyama et al., 1986). Other hypotheses include bacteria being responsible for supplying nitrogen

to the algae based upon isolates from green alga possessing the nitrogenase gene nifH and/or bacteria being responsible for the turnover of plant hormones affecting growth and morphology (Ashen and Goff, 2000; Chisholm et al., 1996).

3.1.4 Experimental Aims

Previous work has focused on the bacterial community present in the environment where Ulva spp. are found growing, and the relationship between Ulva spp. zoospore settlement and bacterial signalling molecules. This study attempted to define the bacterial population present on the surface of the mature Ulva thallus via phylogenetic analysis of a 16S rDNA clone library and isolate strains of marine bacteria, representative of this population from the Ulva thallus. Marine bacteria representative of the Ulva spp. cognate population were then screened for the production of both AHL and AHQ type signal molecules in order to assess if these bacteria were actively engaged in signalling. Based upon previous work which showed that AHLs affect Ulva spp. zoospore settlement and that marine bacteria have a profound effect on Ulva spp. growth morphology we hypothesised that bacterial signal molecules may not only affect Ulva zoospore settlement but also Ulva zoospore germination and the early growth of the Ulva germling. This hypothesis was tested by exposing germinating Ulva zoospores to bacterial biofilms composed of strains indigenous to the Ulva spp. bacterial population, biofilms of transgenic E. coli which expressed various AHL synthase genes, and exogenously adding synthetic AHLs to zoospores settled on sterile glass slides prior to germination.

3.2 Results

3.2.1 16S rDNA Clone Library of the Ulva spp. Bacterial Population

To determine the bacterial population associated with the surface of Ulva, a clone library of 16S rDNA was constructed and 76 clones sequenced, aligned and compared with the Genbank database using the BLAST program. A phylogenetic tree resulting from the analysis of this alignment shows the species richness present on the Ulva thallus to be dominated by the phyla Proteobacteria (65.79%) and Bacteroidetes (34.21%) (Figure 3.3). At class level, the proteobacterial population was dominated by Alphaproteobacteria, which had a relative abundance of 39.47%, with many clones clustering to the Rhodobacteraceae family (34.42% of total library). The remaining proteobacterial clones all clustered with known Gammaproteobacteria (22.36%) and Epsilonproteobacteria (3.94%), with gammaproteobacterial sequences clustering to the Alteromonadaceae. Vibrionaceae. Oceanospirillales and Chromatiales. The Bacteroidetes population was dominated by the Flavobacteria and Sphingobacteria classes, which had relative abundances of 14.47% and 19.74% of the total clone library respectively.

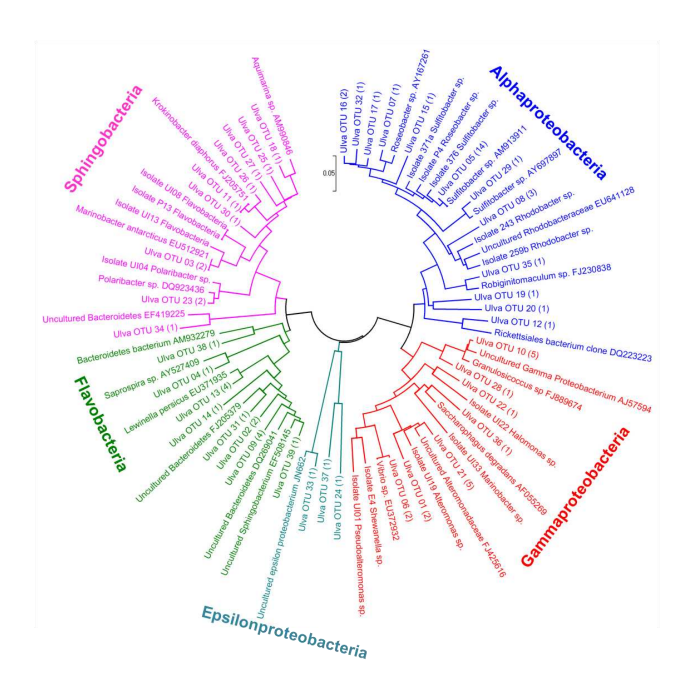
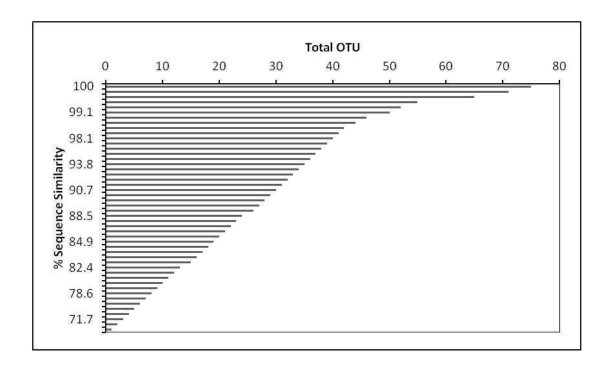


Figure 3.3. Phylogenetic tree of the Ulva spp. thallus bacterial population. The tree resulted from a sequence alignment of 16S rDNA from bacterial clones and isolates obtained from the Ulva thallus at 97% sequence similarity. The reference strains were taken from Genbank. In brackets is the number of clones in each Operational Taxonomic Unit (OTU). The tree topology is based on neighbour-joining and bootstrap analysis was performed with 1000 replications.

The Distance-Based Operational Taxonomic Unit and Richness (DOTUR) program (Schloss and Handelsman, 2005), showed that reducing the percentage sequence similarity when analysing the 16S rDNA clone library reduced the total number of OTUs present in the clone library (Figure 3.4).



% Sequence Similarity	Total OTU
100%	75
99%	50
97%	39
95%	36
90%	27

Figure 3.4. DOTUR analysis of Ulva spp. thallus 16S clone library. Graph displaying the number of OTUs in the 16S rDNA Ulva spp. thallus clone library against the percentage sequence similarly between clones. A reduction in sequence similarity reduced the total OTU in the clone library. The table below the graph displays OTU at the standard cut-off points of 99%, 97%, 95% and 90% sequence similarity.

The top five most abundant OTUs in the Ulva spp. thallus clone library were identified by comparison to the Genbank genomic sequence database via BLAST

analysis. The phylogeny and Genbank accession number of hits identified as having the strongest homology to each OTU are listed in Table 3.1.

Table 3.1. Identity of most abundant OTUs in Ulva spp. thallus clone library. Table displaying BLAST hits in the Genbank database with the greatest homology to the top 5 most abundant OTUs in the Ulva spp. thallus 16S clone library.

OTU Number	Number of clones in each OTU	Phylogeny	Genbank Accession Number
Ulva OTU 5	14	Sulfitobacter sp	AM913911
Ulva OTU 10	5	Uncultured Alteromonadaceae	FJ425616
Ulva OTU 21	5	Uncultured	EU005276
		Gammaproteobacteria	
Ulva OTU 9	4	Uncultured Sphingobacteria	FN433448
Ulva OTU 13	4	Lewinella agarilytica	AM286229

3.2.2 Comparison of Ulva Thallus Population to Rocky Shore and Sea Water Column Population

The 16S rDNA clone library of the Ulva thallus bacterial population was compared to two other similar phylogenetic analyses. One, a 16S rDNA clone library detailing the bacterial population of the rocky shore habitat at Wembury Bay colonised by Ulva spp. carried out in June 2007 (Tait et al., 2009). The other was a 16S rRNA tagged 454 pyrosequenceing data set, also carried out in June 2007, detailing the bacterial population present in the seawater column at the L4 sample site in the Western English Channel, 50°15′N, 04°13′W (Gilbert et al., 2010; Gilbert et al., 2009). The percentage relative abundances of bacterial phyla were compared between the three populations, and it was found that in each population the Proteobacteria and Bacteroidetes phyla dominanted, however the percentage relative abundances of these phyla differed between each population. The Ulva thallus and rocky shore populations showed a greater percentage abundance of Bacteroidetes clones than the water column

population. However, the abundance of Cyanobacteria in the water column was much greater than in the Ulva thallus population, where no Cyanobacteria clones were seen and in the rocky shore population, where Cyanobacteria relative abundance was only 2.19%. In addition to these larger differences in phylum abundance, the Ulva thallus population lacked the low abundances of other bacterial phyla such as Fusobacteria, Fermicutes and Verrucomicrobia which were present in both the rocky shore population and the water column population (Figure 3.5).

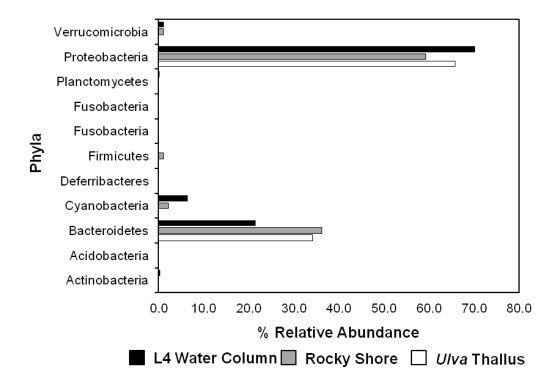
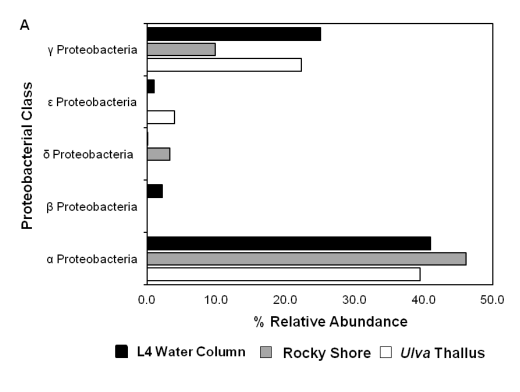
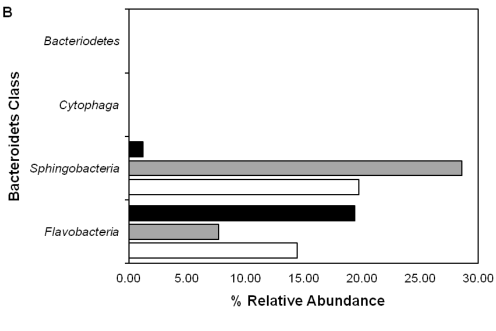


Figure 3.5. Relative abundance of bacterial phyla in three separate marine environmental niches. Comparison of percentage relative abundance of bacterial phyla present in the clone libraries representing the Ulva thallus, rocky shore (Tait et al., 2009) and L4 seawater water column (Gilbert et al., 2009) bacterial populations.

The two dominating phyla, Proteobacteria and Bacteroidetes, in the three populations were further dissected to compare percentage relative abundances at class level. As in the Ulva thallus population the Alphaproteobacteria and

Gammaproteobacteria were the dominant two classes of Proteobacteria present in the rocky shore (46.15% and 9.89% respectively) and water column populations (41.03% and 25.09% respectively), with the Alphaproteobacteria being the most dominant class of Proteobacteria in all three populations. However, the rocky shore clone library contained a greater fraction of Alphaproteobacteria. The rocky shore and water column also contained Betaproteobacteria and Deltaproteobacteria; these were not seen in the Ulva thallus population (Figure 3.6A). The two classes present in the Bacteroidetes phyla in all three populations were the Flavobacteria and the Sphingobacteria; however the percentage relative abundances of each class differed dramatically between each population (Figure 3.6B). The ratio of Sphingobacteria: Flavobacteria differed markedly in each data set: Ulva thallus 37:63; rocky shore 79:21 and water column 6:94.





L4 Water Column

Figure 3.6. Relative abundance of bacterial classes in three separate marine environmental niches. Comparison of percentage relative abundance of proteobacterial classes (**A**) and Bacteroidetes classes (**B**) present in the clone libraries representing the Ulva thallus, rocky shore (Tait et al., 2009) and L4 seawater water column (Gilbert et al., 2009) bacterial populations.

■ Rocky Shore □ Ulva Thallus

3.2.3 Isolation and Typing of Marine Bacteria

In order to gain a further insight into the relationship Ulva has with the bacterial community that colonises the algae, strains were selected that represented the Ulva thallus epiphytic population as identified by the 16S clone library, (see Section 3.2.1). Bacterial strains were isolated from the thallus of wild Ulva spp. colonising rocks at Wembury beach, Devon in September 2008 and from the holdfast of wild Ulva spp. colonising rocks at New Polzeath Beach, Cornwall in September 2008. A total number of 46 isolates were obtained. Some of these isolates possessed the same colony morphologies and were therefore grouped accordingly. From the 40 isolates with distinct colony morphology, representative strains were selected and characterised by phylogenetic typing of their 16S rDNA gene. The majority of the strains isolated directly from the thallus surface and holdfast of Ulva spp. belonged to the Gammaproteobacteria, however strains of Alphaproteobacteria, Flavobacteria and Gram-positive Actinobacteria were also identified (Table 3.2). Bacteria from the PML culture collection were also added to this collection, these strains were isolated from the rocks colonised by Ulva spp. at Wembury beach in January 2001. Strains obtained from the PML culture collection were also characterised by phylogenetic typing of their 16S rDNA to verify phylogeny and how they related to the Ulva thallus clone library.

Table 3.2. Phylogenetic identification of marine bacterial strains based on their 16S rDNA.

Table listing the identity and phylogeny of strains taken from the PML culture collection (PML CC) and those isolated from Ulva thallus and holdfast tissue.

Strain	Identification	Phylogeny (Phylum, Class)	Genbank Accession Number	% Sequence Identity	Ref
I259b	Rhodobacter sp.	Proteobacteria, Alphaproteobacteria	N/A	N/A	PML CC
1243	Rhodobacter sp.	Proteobacteria, Alphaproteobacteria	N/A	N/A	PML CC
I371a	Sulfitobacter sp. KMUT3	Proteobacteria, Alphaproteobacteria	AB583769	98%	PML CC
1376	Sulfitobacter sp. KMUT 3	Proteobacteria, Alphaproteobacteria	AB583769	99%	PML CC
P4	Roseobacter denitrificans NBRC15277	Proteobacteria, Alphaproteobacteria	DQ915623	99%	PML CC
E4	Shewanella sp. SCSA3	Proteobacteria, Gammaproteobacteria	AM884345	97%	PML CC
483	Sulfitobacter sp. KMUT3	Proteobacteria, Alphaproteobacteria	AB583769	98%	PML CC
P13	Favobacterium sp. BSw21403	Bacteroidetes, Flavobacteria	FJ748511	98%	PML CC
UI01	Pseudoalteromonas sp. DJLY29	Proteobacteria, Gammaproteobacteria	EU169492	99%	This Study
UI08	Cellulophaga sp. BSw21403	Bacteroidetes, Flavobacteria	FJ748511	98%	This Study
UI09	Winogradskyella eximia KMM	Bacteroidetes, Flavobacteria	AY521225	98%	This Study
UI18	Rhodobacteraceae bacterium ROS8	Proteobacteria, Alphaproteobacteria	AY841782	99%	This Study
UI13	Cellulophaga sp. RE2-13	Bacteroidetes, Flavobacteria	AF539758	98%	This Study
UI36	Aeromicrobium sp DR8	Actinobacteria, Actinobacteria	FJ464983	98%	This Study
UI20	Pseudomonas sp. LD12	Proteobacteria, Gammaproteobacteria	AM913883	99%	This Study
UI11	Pseudoalteromonas sp. 8	Proteobacteria, Gammaproteobacteria	DQ642811	99%	This Study
UI19	Alteromonas sp. MA112	Proteobacteria, Gammaproteobacteria	AB491743	99%	This Study
UI33	Marinobacter sp. SCSWD16	Proteobacteria, Gammaproteobacteria	FJ461454	99%	This Study
UI39	Marinobacter sp. H96B3	Proteobacteria, Gammaproteobacteria	FJ746575	99%	This Study
UI22	Halomonas sp. NAH1	Proteobacteria, Gammaproteobacteria	EU239362	99%	This Study
UI12	Marinobacter flavimaris CJHH25	Proteobacteria, Gammaproteobacteria	EU169559	99%	This Study
UI04	Polaribacter sp. CS05	Proteobacteria, Gammaproteobacteria	EU477168	99%	This Study
UI32	Marinobacter sp. QJWW107	Proteobacteria, Gammaproteobacteria	FJ384492	99%	This Study
RUBI01	Kocuria rhizophila ES_145con	Actinobacteria, Actinobacteria	EU934094	99%	This Study
RUBI02	Vibrio rumoiensis LAR3	Proteobacteria, Gammaproteobacteria	DQ530292	99%	This Study
RUBI03	Paracoccus sp. jx9	Proteobacteria, Alphaproteobacteria	FJ539115	100%	This Study
RUBI04	Kocuria sp. BBN2C-02d	Actinobacteria, Actinobacteria	FJ357623	99%	This Study
RUBI05	Vibrio rumoiensis LAR03	Proteobacteria, Gammaproteobacteria	DQ530292	98%	This Study
RUBI06	Micrococcus sp. LZXC21	Actinobacteria, Actinobacteria	DQ659067	99%	This Study

3.2.4 QS Signal Molecule Characterisation using AHL Bio-reporters

The Gram-negative strains isolated from the Ulva thallus and the Ulva holdfast rock interface in this study were assayed for AHL signal molecule production using AHL bio-reporter strains. The lux-based E. coli JM109 bio-reporters harbouring pSB536 for the detection of C4-HSL, pSB401 for the detection of C6-HSL and C8-HSL and pSB1142 for the detection of C10-HSL and C12-HSL preferentially were used (Winson et al., 1998; Swift et al., 1997). The initial method employed to assay signal molecule production involved streaking the test strain against the bio-reporter on agar plates composed of two separate culture media, one capable of supporting the bio-reporter and the second capable of supporting the test strain. This method proved to be inconclusive for all strains assayed with the exception of those isolated from the holdfast rock interface. Of the six strains isolated from this interface one, RUBI03 (Paracoccus sp.) showed activation of the JM109 pSB401 bio-reporter, indicating the production of either C6-HSL or C8-HSL.

As the T-streak method proved inconclusive for strains isolated from the Ulva thallus a 96 well plate assay was adopted to screen acidified cell free supernatant extracts obtained from these strains for AHL production using the E. coli AHL bioreporters listed previously. Table 3.3 lists the strains assayed using the 96 plate well technique and shows which bio-reporters were activated by each strain. In the majority of strains where activation of bio-reporters was seen, pSB536 was found to be the most prevalently activated reporter. As JM019 pSB536 is activated by the short chain C4-HSL, activation of this bio-reporter by cell free supernatant extracts indicates production of this signal molecule. Strain UI19 (Alteromonas sp.), RUBI03

(Paracoccus sp.), 483 (Sulfitobacter sp.) and E4 (Shewanella sp.) showed activation of the pSB401 bio-reporter indicating the production of either C6-HSL or C8-HSL.

Table 3.3. Table listing marine bacterial strains assayed for AHL production via 96 well plate method. Strains activating the bio-reporter suggesting AHL production are marked with a +, Strains marked with - denotes no activation of the bio-reporter.

Strain	Identity	Bio-re	porter Ac	tivation
		pSB536	pSB401	pSB1075
UI01	Pseudoalteromonas sp.	-	-	-
UI08	Cellulophaga sp.	+	-	-
UI09	Winogradskyella eximia	-	-	-
UI18	Rhodobacteraceae bacterium	-	-	-
UI13	Cellulophaga sp.	+	-	-
UI20	Pseudomonas sp.	+	-	-
UI11	Pseudoalteromonas sp.	-	-	-
UI19	Alteromonas sp	+	+	-
UI39	Marinobacter sp.	-	-	-
UI33	Marinobacter sp.	+	-	-
UI22	Halomonas sp.	+	-	-
UI12	Marinobacter flavimaris	-	-	-
UI04	Polaribacter sp.	+	-	-
UI32	Marinobacter sp.	-	-	-
RUBI02	Vibrio rumoiensis	-	-	-
RUBI03	Paracoccus sp.	+	+	-
RUBI06	Vibrio rumoiensis	-	-	-
E4	Shewanella sp.	+	-	+
483	Sulfitobacter sp.	-	+	-

3.2.5 QS Signal Molecule Characterisation using LCMS

The screening of marine bacteria directly isolated from Ulva for signal molecule production using bio-reporters showed that a number of strains were producing AHLs. This method of screening however did not provide data relating to specific AHL production for each strain. In order to gain more insight into the AHLs produced by bacteria that colonise Ulva, Liquid Chromatography coupled with Mass Spectroscopy (LCMS) was utilised. Acidified cell free supernatant extracts of strains isolated from Ulva and strains selected from the PML culture collection that were present in the Ulva clone library were analysed for the presence of individual AHLs using non quantitative LCMS. This detects un-substituted, 3-oxy-substituted and 3-hydroxyl-substituted homoserine lactones with fatty acid side chains ranging from 4 to 14 carbons in length. Extracts from acidified sterile marine broth and methanol were used as negative controls (Table 3.4). The results of the LCMS screen confirmed the AHL profiles which had previously been seen using bio-reporter based assays: bacteria associated with Ulva produce a wide range of AHL signal molecules. Particularly prevalent were C4-HSL and AHLs with fatty acid side chains 8 carbons in length, both un-substituted and substituted with either 3-oxy or 3-hydroxyl groups. Examples of compound mass spectra showing 3-oxo-C8-HSL in extracts of I376 (Sulfitobacter sp.) and C4-HSL in UI20 (Pseudomonas sp.) are displayed in Figure 3.7.

Table 3.4. Table showing AHLs produced by strains representative of Ulva's epiphytic bacterial community as identified by non-quantitative LCMS analysis. Boxes in the table shaded grey denotes production of that AHL by the strain.

Strain	Identity	AHL					
		Un-substituted Series (X-HSL)					
		C4	C6	C8	C10	C12	C14
RUBI03	Parracoccus sp.						
UI20	Pseudomonas sp.						
UI19	Alteromonas sp.						
UI33	Marinobacter sp.						
I371a	Sulphitobacter sp.						
243	Roseobacter sp.						
I376	Sulfitobacter sp.						
P13	Flavobacterium sp.						
UI08	Cellulophaga sp.						
UI13	Cellulophaga sp.						
438	Roseobacter sp.						
	•	3	3-oxo Gr	oup Seri	ies (3-oxo	-X-HSL	<u>, </u>
		C4	C6	C8	C10	C12	C14
RUBI03	Parracoccus sp.						
UI20	Pseudomonas sp.						
UI19	Alteromonas sp.						
UI33	Marinobacter sp.						
I371a	Sulphitobacter sp.						
243	Roseobacter sp.						
I376	Sulfitobacter sp.						
P13	Flavobacterium sp.						
UI08	Cellulophaga sp.						
UI13	Cellulophaga sp.						
438	Roseobacter sp.						
		3-h	ydroxyl	Group S	eries (3-0	ОН-Х-Н	SL)
		C4	C6	C8	C10	C12	C14
RUBI03	Parracoccus sp.						
UI20	Pseudomonas sp.						
UI19	Alteromonas sp.						
UI33	Marinobacter sp.						
I371a	Sulphitobacter sp.						
243	Roseobacter sp.						
I376	Sulfitobacter sp.						
P13	Flavobacterium sp.						
UI08	Cellulophaga sp.						
UI13	Cellulophaga sp.						
438	Roseobacter sp.						

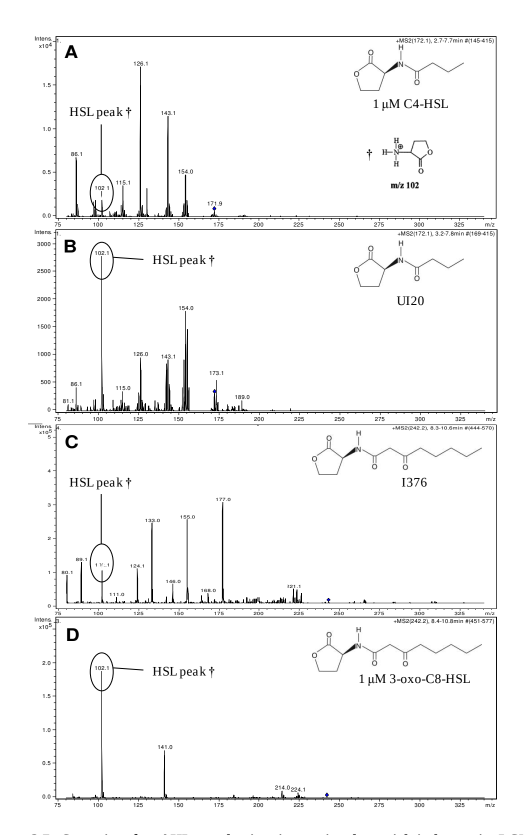


Figure 3.7. Screening for AHL production in marine bacterial isolates via LCMS. Examples of product ion spectra from the breakdown of nominal precursor ion of m/z 172 (C4-HSL), 1 μ M C4-HSL standard (A), UI20 (Pseudomonas sp.) (B) and of nominal precursor ion m/z 172 (C4-HSL), I376 (Sulfitobacter sp.) and 1 μ M 3-oxo-C8-HSL standard (D). The peak representing in daughter ion m/z 102.1 (homoserine lactone) highlighted in all spectra.

In addition to assaying for AHL production the same strains were assayed for the production of AQ type signal molecules using non-quantitative LCMS. The AQs assayed were the PQS type quinolones with alkyl chains of 7, 9 and 11 carbons; HHQ type quinolones with alkyl chains of 7, 9 and 11 carbons and N-oxide AHQs with alkyl chains of 7 and 9 carbons. LCMS analysis failed to show any strong evidence of AQ production in the marine bacterial stains assayed (data not shown).

3.2.6 Ulva spp. Zoospore Germination Response when Exposed to V. anguillarium Biofilms

V. anguillarium produces 3-oxo-C10-HSL, C6-HSL and 3-hydroxy-C6-HSL directed by the AHL synthase homologues vanI and vanM, (Milton et al., 2001; Milton et al., 1997), and was utilised in the initial experiments exploring the relationship between AHL-producing bacteria and Ulva zoospore settlement (Tait et al., 2005; Joint et al., 2002). V. anguillarium is, therefore, a viable model organism to investigate the effect of AHL-producing biofilms on Ulva zoospore germination. As the amount of AHLs produced depends on biofilm density, the effect on Ulva germination on biofilms of varying density was compared. Repeated observations after 72 h incubation, using length equal to or greater than 15 μm to define a successfully germinated zoospore, Ulva spp. germlings grown on AHL expressing V. anguillarium wt biofilms had reduced average lengths when compared to Ulva spp. germlings grown on biofilms composed of a V. anguillarium ΔvanI/M mutant (Figure 3.8). As the zoospores were incubated on the V. anguillarium biofilms for 72 h prior to being fixed and stained for image analysis, 100% of the settled Ulva spp. zoospores had germinated and in many cases it was difficult to select and measure individual Ulva germlings as the biofilm

slides were crowded with germinated zoospores. As a result of this observation it was decided to reduce the zoospore incubation time to 48 h and add a sample point at 24 h for future experiments, in order to better assay the period of zoospore germination.

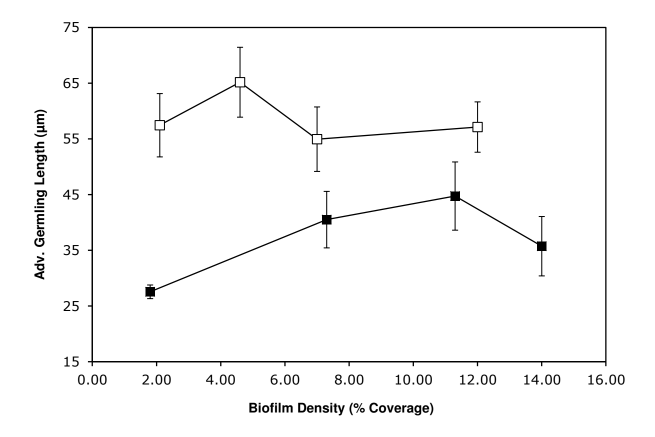


Figure 3.8. Effect of V. anguillarum biofilms on Ulva zoospore germination. Average length of Ulva spp. germlings grown on biofilms composed of V. anguillarum wt (\blacksquare) and V. anguillarium $\Delta \text{vanI/M}$ (\square) of varying bacterial density. Error bars represent 95% confidence intervals.

3.2.7 Ulva Zoospore Germination Response when Exposed to Biofilms Composed of Marine Bacteria Indigenous to Ulva's Bacterial Population

Although the germling growth on biofilms of V. anguillarium wt was reduced when compared to growth on biofilms of the V. anguillarium $\Delta vanI/M$ mutant, this strain is not known to be associated with the mature algae. Similar experiments were

therefore conducted with bacteria isolated from the Ulva thallus. The 16S clone library of bacteria associated with Ulva spp. showed both Sulfitobacter spp. (an alphaproteobacterial species) and Uncultured Alteromonadales to be highly abundant within the population (Section 3.2.1). On the basis of these findings two strains; 483 and E4 were selected for repeated experiments investigating zoospore response to Ulva's indigenous bacteria. Strain 483 is a strain of Sulfitobacter and E4 a strain of Shewanella, a member of the Alteromonadale order. Strain 483 produces C8-HSL and E4 produces C4-HSL and C8-HSL as shown by activations of AHL bio-reporters pSB536 and pSB401, (Table 3.3). Both 483 and E4 were transformed with the AHL lactonase aiiA from Bacillus sp. 240B1, cloned into the broad host vector pBBRIMCS-1 (pMT01). Successful transformation with the pMT01 plasmid rendered cognate AHLs biologically inert as aiiA hydrolyses the homoserine lactone ring. This was confirmed as both 483 pMT01 and E4 pMT01 failed to activate lux-based AHL bioreporters pSB536 and pSB401 (data not shown). Biofilms of strains 483 and E4 transformed with empty pBBRIMCS-1 vector which produced AHLs and 483 and E4 transformed with pMT01 were grown from a range of starting inoculums on glass cover slides. Ulva zoospores were settled onto the biofilms, incubated as described previously and sampled at 24 h and 48 h.

The varying volumes of inoculate used to grow biofilms of Shewanella sp. E4 and Sulfitobacter sp. 483 produced biofilms with bacterial densities ranging from approximately 35 to 75% coverage, with no major difference in growth observed between strains carrying the pBBRIMCS-1 or pMT01 plasmid.

Repeated observations showed that at the 24 h sample point there was little observable difference in average germling length when zoospores were exposed to the AHL-producing Shewanella sp. E4 pBBRIMCS-1 biofilms in comparison to biofilms composed of Shewanella sp. E4 pMT01 which did not produce AHLs (Figure 3.9). However, at higher biofilm densities average germling length was increased on Shewanella sp. E4 pMT01 in comparison to Shewanella sp. E4 pBBRIMCS-1 (Figure 3.9). In contrast, the percentage of zoospores that had germinated at the 24 h sample point was increased on biofilms composed of Shewanella sp. E4 pMT01 (Table 3.5). At the 48 h sample point, average germling length was greatly increased on biofilms composed of Shewanella sp. E4 pMT01 in comparison to Shewanella sp. E4 pBBRIMCS-1 biofilms (Figure 3.9). As at 24 h, percentage germination of zoospores was increased on Shewanella sp. E4 pMT01 biofilms in comparison to Shewanella sp. E4 pBBRIMCS-1 biofilms at the 48 h sample point (Table 3.5).

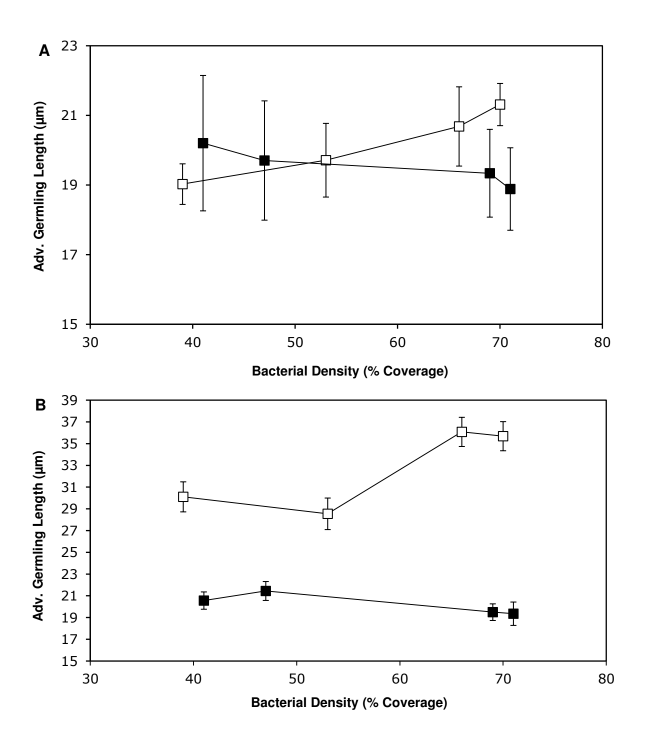


Figure 3.9. Effect of Shewanella biofilms on Ulva zoospore germling growth. Average germling length of zoospores exposed to biofilms composed of AHL-producing Shewanella sp. E4 pBBRIMCS-1 (■) and non AHL-producing Shewanella sp. E4 pMT01 (□) at 24 h incubation (**A**) and 48 h incubation (**B**). Error bars represent 95% confidence intervals.

Table 3.5. Ulva zoospore percentage germination in the presence of Shewanella biofilms. Table comparing percentage of Ulva zoospores that germinated when exposed to biofilms of increasing inoculate volume composed of non AHL-producing Shewanella sp. E4 pMT01 and AHL-producing Shewanella sp. E4 pBBRIMCS (E4 wt) at both 24 h and 48 h incubation.

			Percentage Ulva Zoospore Germination (24 h)		Germination Germination		nation
		Strain	E4 pMT01	E4 wt	E4 pMT01	E4 wt	
ıme		0.5	48	10	76	56	
Volu	(ml)	1	27	9	77	65	
Inoculum Volume	n)	2	44	13	87	51	
Ino		4	73	18	87	47	

Repeated observations also showed that at both 24 h and 48 h sample points average germling length was increased on biofilms composed of non AHL-producing Sulfitobacter sp. 483 pMT01 in comparison to AHL-producing Sulfitobacter sp. 483 pBBRIMCS-1 biofilms (Figure 3.10). This pattern was also observed with regards to percentage germination of Ulva zoospores with germination being greater on Sulfitobacter sp. 483 pMT01 biofilms in comparison to Sulfitobacter sp. 483 pBBRIMCS-1 biofilms at both 24 h and 48 h sample points (Table 3.6).

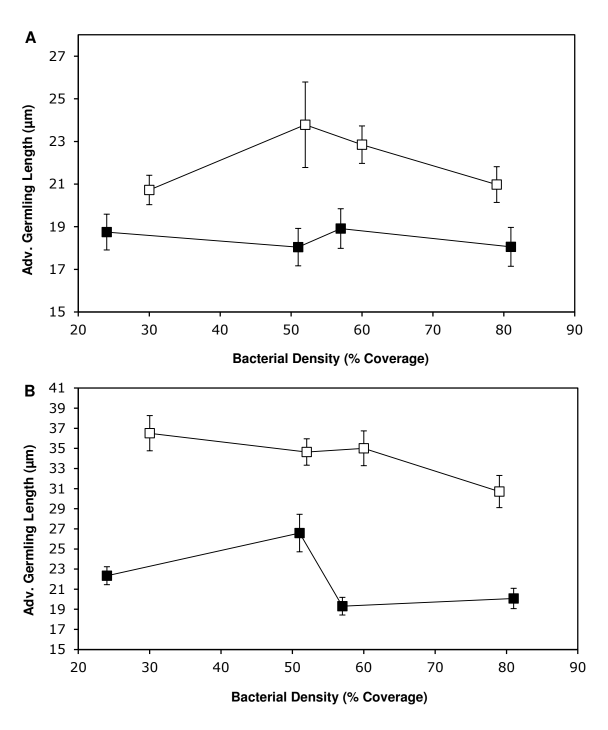


Figure 3.10. Effect of Sulfitobacter biofilms on Ulva zoospore germling growth. Average germling length of zoospores exposed to biofilms composed of AHL-producing Sulfitobacter sp. 483 pBBRIMCS-1 (■) and non AHL-producing Sulfitobacter sp. 483 pM101 (□) at 24 h incubation (**A**) and 48 h incubation (**B**). Error bars represent 95% confidence intervals.

Table 3.6. Ulva zoospore percentage germination in the presence of Sulfitobacter biofilms. Table comparing percentage of Ulva zoospores that germinated when exposed to biofilms of increasing inoculate volume composed of non AHL-producing Sulfitobacter sp. 483 pMT01

and AHL-producing Sulfitobacter sp. 483 pBBRIMCS (483 wt) at both 24 h and 48 h

incubation.

			Percentage Ulva Zoospore Germination (24 h)		Percentage U Germi (48	_
		Strain	483 pMT01	483 wt	483 pMT01	483 wt
ıme		0.5	57	21	80	75
Volu	(ml)	1	35	18	85	60
Inoculum Volume	n)	2	64	17	80	60
Ino		4	56	14	76	61

3.2.8 Ulva Zoospore Germination Response when Exposed to Transgenic E. coli Biofilms

The introduction of AiiA into Shewanella sp. E4 and Sulfitobacter sp. 483 may have altered phenotypes other than AHL production that may have affected zoospore germination. In order to overcome this problem and to expand the observations of the relationship between AHLs and Ulva zoospore germination seen with exposure to biofilms of indigenous marine bacteria, Ulva zoospore germination was assayed using biofilms composed of transgenic E. coli strains which expressed various recombinant AHL synthase genes. These AHL synthases were rhll from P. aeruginosa (directing the production of C4-HSL), luxI from V. fischeri (directing the production of 3-oxo-C6-HSL) and vanI from V. anguillarum (directing the production of 3-oxo-C10-HSL) (Hanzelka et al., 1997; Milton et al., 1997; Parsek et al., 1997). For each transgenic E. coli strain, zoospore germination was compared to E. coli possessing vector plasmids

without recombinant AHL synthase homologues. These strains provided a constant supply of AHLs, facilitating a more stable assay for monitoring the response of Ulva zoospores to AHLs. Owing to the detrimental effect of the osmotic pressure of seawater on the growth and survival of E. coli biofilms, zoospore slides were incubated for a reduced time of 24 h in 70% sterile filtered seawater prior to being fixed and stained. At 24 h incubation, there was a small yet significant decrease in average germling length when Ulva zoospores were settled and grown on biofilms composed of E. coli expressing rhlI and vanI, (one-way ANOVA P < 0.01). Although the response to E. coli expressing luxI was not found to be significant, (one-way ANOVA P > 0.01), a modest decrease in the presence of this biofilm was also apparent (Figure 3.11). Marked reductions in the percentage of germinated Ulva zoospores were however observed on biofilms of all AHL expressing transgenic E. coli strains at 24 h, 33%, 19% and 22% respectively in the rhlI, luxI and vanI expressing stains in comparison to 78% 73% and 60% on the biofilms of the respective AHL deficient vector control stains (Table 3.7).

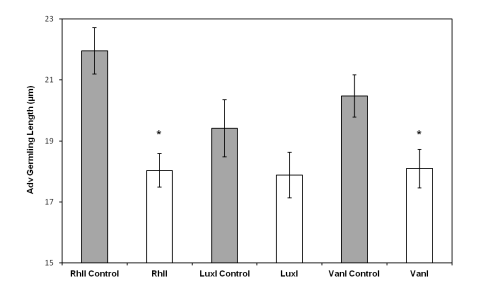


Figure 3.11. Effect of transgenic E. coli biofilms on Ulva germling growth. Average Ulva germling length when exposed to transgenic E. coli biofilms expressing recombinant AHL synthase homologues in comparison to biofilms composed of non AHL-producing E. coli controls at 24 h incubation. Error bars represent 95% confidence intervals, * marks values significantly different to controls (one-way ANOVA P < 0.01).

Table 3.7 Ulva zoospore percentage germination in the presence of transgenic E. coli biofilms. Table comparing percentage of Ulva zoospores that germinated when exposed to biofilms composed of AHL synthase expressing E. coli and wt E. coli controls at 24 incubation.

	AHL Expressing	Control
rhlI (C4-HSL)	33	78
luxI (3-oxo-C6-HSL)	19	73
vanI (3-oxo-C10-HSL)	22	60

3.2.9 Ulva Zoospore Germination in the Presence of Synthetic AHLs

Previously, increased Ulva zoospore settlement has been seen on agarose films containing synthetic AHLs with fatty acid side chains ranging from 6 to 14 carbons in length (Tait et al., 2005). In addition, many of the bacteria associated with Ulva have been shown to produce a wide range of signalling molecules. Having identified AHL-producing biofilms to inhibit the early growth of Ulva germlings, this study

investigated the relationship between Ulva zoospore germination and synthetic AHL signalling molecules. Ulva zoospores were first settled onto sterile glass surfaces, exposed to set concentrations of exogenous C4-HSL, C6-HSL, C8-HSL, C10-HSL and C12-HSL and incubated for a period of 24 h and 48 h under the condition described in section 3.2.6.

Under all experimental conditions both the average length of Ulva germlings and percentage of germination of Ulva zoospores was greater after 48 h incubation compared to 24 h incubation. Two-way PERMANOVA analysis to determine the effect of AHL concentration and incubation time on germling length revealed a significant effect of AHL concentration for all AHLs assayed (P < 0.01). However, germling length in different AHL concentrations was not the same after 24 h and 48 h of incubation (P > 0.01). For clarity, only the results for the 48 h time point are displayed here, however the results from the 24 h time point are discussed within the main text and can be found in Appendix 3.

At the 24 h sample point there was a significant decrease in average germling length at 0.5 μ M C4-HSL in comparison to when no exogenous C4-HSL was added (one-way ANOVA P < 0.01). However, no significant difference in average germling length was observed at any other C4-HSL concentration in comparison to when no exogenous AHL was added (one-way ANOVA P > 0.01). At the 48 h sample point there was a significant reduction in average germling length at a C4-HSL concentration of 5 μ M in comparison to when no exogenous C4-HSL is presen, (one-way ANOVA P < 0.01) (Figure 3.12). However, there was no significant difference in average germling length

at 0.5 μ M and 50 μ M C4-HSL in comparison to when no exogenous AHL was added (one-way ANOVA P > 0.01). (Figure 3.12).

At the 24 h sample point there was a significant decrease in average germling length in the presence of C6-HSL at 5 μ M in comparison to when no exogenous C6-HSL was added (one-way ANOVA P < 0.01). No significant differences in germling length observed at any other C6-HSL concentration in comparison to when no exogenous C6-HSL was present (one-way ANOVA P > 0.01). At the 48 h sample point there was a significant reduction in average germling length in the presence of 50 μ M C6-HSL compared to when no exogenous C6-HSL was present (one-way ANOVA P < 0.05) (Figure 3.12). No significant difference in average germling length was seen at 0.5 μ M and 5 μ M C6-HSL in comparison to when no exogenous C6-HSL was present (one-way ANOVA P > 0.01) (Figure 3.12).

At the 24 h sample point there was a significant increase in average germling length in the presence of 50 μ M C8-HSL in comparison to when no exogenous C8-HSL was present (one-way ANOVA P < 0.01). However, no significant differences in germling length were observed at any other C8-HSL concentration in comparrsion to when no exogenous C8-HSL was present (one-way ANOVA P > 0.01). At the 48 h sample point there was also a significant increase in average zoospore length in the presence of 50 μ M C8-HSL (one-way ANOVA P < 0.01) (Figure 3.12). No significant difference in average germling length was seen at 0.5 μ M and 5 μ M C8-HSL in comparrsion to when no exogenous C8-HSL was present (one-way ANOVA P > 0.01) (Figure 3.12).

At the 24 h sample point there was no significant difference in average germling length in the presence of C10-HSL at any concentration compared to when no exogenous C10-HSL was present (one-way ANOVA P > 0.01). At the 48 h sample point there was a significant decrease in average germling length at 50 μ M C10-HSL compared to when no exogenous C10-HSL was present, however there was an increase in germling length at 5 μ M (one-way ANOVA P < 0.01) (Figure 3.12). No significant difference in average germling length was seen at 0.5 μ M C10-HSL compared to when no exogenous C10-HSL was present (one-way ANOVA P > 0.01) (Figure 3.12).

Repeated observations at the 24 h sample point showed average germling length was significantly increased at 0.5 μ M C12-HSL and significantly decreased at 50 μ M C12-HSL in comparison to where no exogenous C12-HSL was present (one-way ANOVA P < 0.01). At the 48 h sample point there was a significant decrease in average germling length at 50 μ M C12-HSL in comparison to where no exogenous C12-HSL was present (one-way ANOVA P < 0.01) (Figure 3.12).

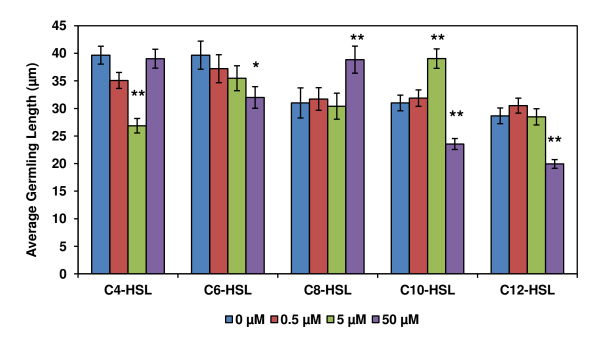


Figure 3.12. Effect of exogenously added synthetic AHLs on Ulva germling growth. Average Ulva germling length when exposed to set concentrations of various AHLs at 48 h incubation. Error bars represent 95% confidence intervals and asterisks show those values that differed significantly from that of 0 μ M AHL (one-way ANOVA * = P < 0.05, ** = p < 0.01).

As with Ulva zoospore germination assays using bacterial biofilms the percentage germination of zoospore was recorded at the 24 h and 48 h sample points in the presence of all synthetic AHLs assayed. At the 24 h sample point, percentage zoospore germination was reduced at 0.5 μ M C4-HSL and 50 μ M C4-HSL in comparison to when no C4-HSL was present (18%, 20% and 34% respectively). However it was increased at 5 μ M C4-HSL (45%) in comparison to when no C4-HSL was present (34%) (Table 3.8). At the 48 h sample point, percentage germination of Ulva zoospores was reduced when exposed to C4-HSL at all concentrations assayed with the greatest reduction in percentage germination at 5 μ M C4-HSL in comparison to when no C4-HSL was present (61% and 86% respectively) (Table 3.8).

At the 24 h sample point percentage zoospore germination was reduced in the presence of C6-HSL compared to when no exogenous C6-HSL was present, particularly at 0.5 μ M and 50 μ M C6-HSL where percentage germination was 11% and 14% respectively compared to 35% at 0 μ M C6-HSL (Table 3.8). As with the 24 h sample the percentage zoospore germination at the 48 h sample point was also reduced when exogenous C6-HSL was present, with the greatest reduction seen at 5 μ M C6-HSL in comparison to when no C6-HSL was present (78% and 86% respectively) (Table 3.8).

At both the 24 h and 48 h sample points the percentage zoospore germination was increased when exogenous C8-HSL was present in comparison to when no exogenous C8-HSL was present (Table 3.8). The exception to this trend was where zoospore germination was recoreded in the presence of 5 μM C8-HSL at the 48 h sample point where percentage germination was slightly reduced comparison to when no C8-HSL was present (78% and 80% respectively) (Table 3.8).

At the 24 h sample point there was an increase in percentage zoospore germination at 0.5 μ M and 5 μ M C10-HSL compared to when no exogenous C10-HSL was present (28%, 23% and 21% respectively). However, at 50 μ M C10-HSL percentage germination was reduced compared to when no exogenous C10-HSL was present (17% and 21% respectively) (Table 3.8). At the 48 h sample point percentage germination was increased at 0.5 μ M and 5 μ M C10-HSL with a substantial fall in germination at 50 μ M C10-HSL of 64% compared to 80% when no exogenous C10-HSL was present (Table 3.8).

At the 24 h sample point percentage zoospore germination was notably reduced at 50 μ M C12-HSL; 17% germination at 50 μ M C12-HSL compared to 64% germination at 0 μ M C12-HSL (Table 3.8). At 48 h, increased percentage zoospore germination was seen at 0.5 μ M C12-HSL, germination was then substantially reduced below the level seen when no exogenous C12-HSL was present. At C12-HSL concentrations of 5 μ M and 50 μ M, 70% and 46% respectively compared to 80% when no C12-HSL was present (Table 3.8).

Table 3.8. Ulva zoospore percentage germination in the presence of exogenously added synthetic AHLs. Table showing the percentage Ulva zoospore germination in the presence of set concentrations of various AHLs at both 24 h and 48 h incubation.

	Percentage Ulva Zoospore Germination (24 h)			Pei		Ilva Zoos ination Bh)	spore	
AHL Conc.	0	0.5	5	50	0	0.5	5	50
(μ M)								
C4-HSL	34	18	45	20	86	82	61	84
C6-HSL	35	11	30	14	86	85	78	80
C8-HSL	21	38	35	38	80	81	78	88
C10-HSL	21	28	23	17	80	80	89	64
C12-HSL	64	78	72	17	69	83	79	46

As C12-HSL was the most stable synthetic AHL in seawater due its increased acyl side chain, experiments assayed for effect on Ulva zoospore germination were repeated using C12-HSL concentrations ranging between 5 μ M and 50 μ M. With the notable exception of at 20 μ M, no significant difference in average germling length was observed at any concentration of C12-HSL in comparison to when no exogenous C12-HSL was present at the 24 h sample point (one-way ANOVA P > 0.001) (data not shown). However, at the 48 h sample point average germling length was significantly reduced

when C12-HSL was present at all concentrations assayed compared to when no exogenous C12-HSL was present (one-way ANOVA P < 0.001) (Figure 3.13).

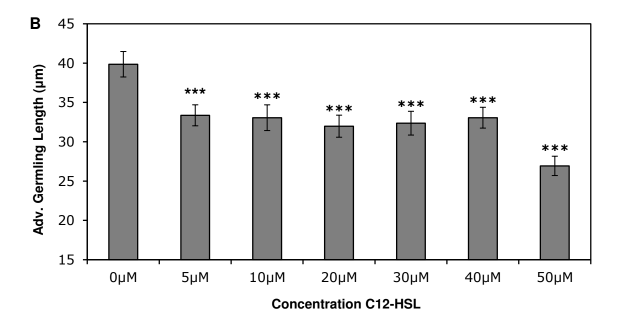


Figure 3.13. Effect of exogenously added synthetic C12-HSL on Ulva germling growth. Average Ulva germling length when exposed to a range of C12-HSL concentrations at 48 h incubation. Error bars represent 95% confidence intervals and asterisks show those values that differed significantly from that of 0 μ M AHL (one-way ANOVA =P < 0.001).

The percentage of germinated Ulva zoospores was also recorded for each concentration of C12-HSL at both the 24 h and 48 h sample points. At the 24 h sample point the percentage of germinated zoospores was reduced at all C12-HSL concentrations, with the exception of 40 μ M (39%), compared to when no exogenous C12-HSL was present (35%) (Table 3.9). At the 48 h sample point percentage germination was reduced in the presence of C12-HSL at all concentrations in comparison to when no exogenous AHL was present; particularly at 50 μ M C12-HSL where zoospore germination was at 65% in comparison to 86% when no exogenous C12-HSL was present (Table 3.9).

Table 3.9. Ulva zoospore percentage germination in the presence of exogenously added synthetic C12-HSL. Table showing the percentage Ulva zoospore germination in the presence of set concentrations of C12-HSL at both 24 h and 48 h incubation.

C12-HSL Conc. (µM)	Percentage Ulva Zoospore Germination (24 h)	Percentage Ulva Zoospore Germination (48 h)
0	35	86
5	29	78
10	27	71
20	23	77
30	25	78
40	39	80
50	27	65

3.3. Discussion

3.3.1 The Bacterial Population Associated with Ulva

This study has identified the epiphytic bacterial community residing on the surface of the macroalgae Ulva that was found growing in the intertidal rocky shore environment of southwest region of the UK coastline. The population present on the surface of the Ulva thallus is composed of bacteria from the phyla Proteobacteria and Bacteroidetes. The dominant classes of bacteria are Alphaproteobacteria, Gammaproteobacteria, Flavobacteria and Sphingobacteria. The bacterial population present on the Ulva thallus was found to be highly similar to the population present within biofilms found on the rocks colonised by Ulva (Tait et al., 2009). The same dominant groups were represented in both populations; however, the relative abundance of each class differed between the populations seen in the two environments.

The epiphytic population present on Ulva, identified by this study was also remarkable similar to the epiphytic bacterial population present on the surface of Ulva australis, an antipodeanian member of the Ulvacean family (Burke et al., 2011). Using a large number of 16S rRNA clones Burke et al. (2011) showed the U. australis population to be composed of Proteobacteria and Bacteroidetes at the phylum level, dominated by Alphaproteobacteria, Gammaproteobacteria and Flavobacteria. At the species level this study showed a significantly large amount of variability between clone libraries from different individual U. astralis samples collected in the same area (Burke et al., 2011). This dominance of Proteobacteria and Bacteroidetes observed by both this study and the study into the epiphytic population of U. australis is replicated in the epiphytic communities present on other algal species such as Laminaria

hyperborean, Laminaria saccharinam, (Bengtsson et al., 2010, Staufenberger et al., 2008) and endophytic communities of Bryopsis sp. (Hollants et al., 2011)

The Ulva thallus population identified by this study was also similar to the bacterial population observed in the water column of the Western English Channel; however as with the rocky shore population (Tait et al., 2009), the water column population shows greater bacterial diversity and an increased range of bacterial phyla (Gilbert et al., 2009). Variability between the Ulva epiphytic population and the surrounding seawater bacterial population was also observed in U. australis (Burke et al., 2011). The similarities between the Ulva thallus population and the rocky shore and water column populations suggests Ulva acquires its bacterial population from the rocks on which it colonises and the seawater to which it is exposed during the tidal cycle. However, the difference in three populations supports a hypothesis that some bacteria are better adapted for epiphytic growth on a biotic surface. The main difference between the Ulva thallus population and the rocky shore was that the relative abundance of the Sphingobacteria class of Bacteroidetes is much higher in the rocky shore population. The sphingobacterial population in both the Ulva thallus and rocky shore populations is in turn much greater than that found in the water column population. The difference between Sphingobacterial abundance in the Ulva thallus, the rocky shore and the water column is expected as costal environments tend to possess a higher abundance of Bacteroidetes in comparison with the open ocean (Alonso et al., 2007). Additionally, Sphingobacteria are also commonly found in areas of reduced salinity such as the intertidal zone (Alonso et al., 2007). Another factor that would explain the difference between the Ulva thallus population and that of the seawater is the increased nutrient availability present on the Ulva thallus; algal species exude a number of nutrients such as organic carbon compounds that bolster microbial growth compared to the open ocean environment which has limited nutrient availability (Pregnall, 1983). A surprising result was the difference in abundance of Sphingobacteria seen in the Ulva thallus population compared to that of the rocky shore population. This difference may be due to Sphingobacteria being better adapted to local environmental conditions present on the rocks, or more likely, due to unaccounted differences in environmental conditions present at the location and time of sampling such as local wheather activity and/or nutrient levels in the seawater. One such condition, that could affect Bacteroidetes populations, is the occurrence of a phytoplankton bloom as it has been reported that Bacteroidetes populations are increased at the time of such blooms (Pinhassi et al., 2004).

Microbial communities are highly influenced by the seasonal changes that take place in the marine environment. This has been comprehensively demonstrated by phylogenetic, meta-genomic and metabolomic studies conducted over broad time periods at the L4 sampling site in the Western English Channel which show significant changes in bacterial populations depending on the time of sampling (Gilbert et al., 2010; Gilbert et al., 2009). The bacterial community on the surface of Ulva has also been shown to have a high degree of temporal variability (Tujula et al., 2010). Density Gradient Gel Electrophoresis (DGGE) studies analysing the U. australis epiphytic population over one year identified up to 40% seasonal variability, this study also puts forward the case for Ulva spp. possessing a core bacterial population which remains unchanging throughout the year. This hypothesis was however discredited by 16S rRNA phylogenetic analysis of the U. australis community (Burke et al., 2011; Tujula et al., 2010). In addition, Ulva itself undergoes drastic growth changes throughout the

year with biomass and production being at the lowest during the winter period (Maggs and Callow, 2001). Therefore, the bacterial population on Ulva spp. will alter in response to these conditions. Seasonal differences in the bacterial community in response to the changing growth cycle of algae has been demonstrated in the brown algal species Laminaria hyperborean and Laminaria saccharina (Bengtsson et al., 2010; Staufenberger et al., 2008). The bacterial population identified in this study can only be attributed to the time of year when sampling was carried out. In order to gain a more comprehensive understanding of the Ulva bacterial population, sampling and phylogenetic analysis must be conducted throughout the course of Ulva's annual growth cycle.

3.3.2 Signal Molecule Production by Marine Bacteria Representative of the Ulva Bacterial Population

Through various detection methods this study has shown that many of the bacteria isolated from the Ulva thallus and/or representing the Ulva bacterial population are actively producing a range of AHL signal molecules, (see section 3.2.4 and 3.2.5). AHL production by bacterial species isolated from the marine environment and specifically from Ulva has been reported previously (Tait et al., 2009; Mohamed et al., 2008; Gram et al., 2002). Initially AHL detection assays were conducted via both T streak and plate based methods using bio-reporters to detect the presence of AHLs (Winson et al., 1998; McClean et al., 1997). In general the T streak method proved inconclusive; potentially due to abiotic disruption of the homoserine lactone ring via hydrolysis under the increased pH of marine broth, abolishing the biological activity of the AHL (Hmelo and Van Mooy, 2009; Yates et al., 2002). The original T streak

method did however produce one clear result. Isolate RUBI03, identified as a strain belonging to the genus Paracoccus was shown to induce bioluminescence in the bioreporter E. coli pSB401 suggesting that its cognate AHLs possess fatty acid side chains 6 to 8 carbons in length, later confirmed via LCMS analysis. These data are intriguing given that Steindler and Venturi (2007) showed that Paracoccus spp. produced long chain AHLs (Steindler and Venturi, 2007). Assaying dichloromethane extractions from acidified culture supernatants identified AHL production by a wider range of bacteria isolated from Ulva. Performing signal molecule analysis via LCMS to characterise AHLs being produced by different strains isolated from Ulva and strains representative of the Ulva population proved far more beneficial than both T streak and plate based detection assays as it unequivocally revealed which AHLs were being produced by each strain assayed. LCMS analysis showed a wide range of signal molecules being produced by the marine bacteria isolated from the Ulva thallus. It was interesting to note that the AHLs being produced by the strains isolated from Ulva were, in general, found to be producing short to mid range sized fatty acid side chains and not the longer side chains which are more stable in sea water and predominantly show increased Ulva zoospore settlement (Tait et al., 2005). The predominant AHLs identified to be present in rocky shore biofilms were C8-HSL and C10-HSL: these molecules, along with the longer chain C12-HSL and C14-HSL were also seen to be produced by bacteria isolated from the rocky shore (Tait et al., 2009). Although both bacteria isolated from the rocky shore and bacteria isolated from the Ulva thallus surface show a wide range of AHL production, the Ulva thallus isolates tend to produce AHLs with shorter fatty acid side chains. The production of AHLs with short to medium length fatty acid side chains differs from other studies that have investigated AHL production in bacterial strains, isolated from a variety of marine environments, which show a predominant bias to long chain AHL production (Wagner-Döbler et al., 2005; Taylor et al., 2004; Gram et al., 2002).

3.3.3 Effect of AHLs on Ulva Zoospore Germination

A relationship between Ulva and bacterial signalling molecules has been established with regards to Ulva zoospore settlement, with Ulva zoospores being shown to preferentially settle on bacterial biofilms which produce AHLs and on agarose slides permeated with synthetic AHLs (Tait et al., 2005; Joint et al., 2002). Preferential settlement of zoospores was also shown to be abolished either by the hydrolysis of AHLs due to the alkaline pH of sea water, the actions of AHL-inactivating enzymes such as the lactonase aiiA and by AHL inactivation from other bacteria such as Shewanella spp. (Tait et al., 2009; Tait et al., 2005). The method by which Ulva zoospores sense AHLs remains uncertain although it has been shown that in response to an AHL source Ulva zoospore motility is reduced, therefore allowing an accumulation of zoospores at the AHL source promoting increased settlement (Wheeler et al., 2006).

In addition to affecting Ulva zoospore settlement bacteria have also been shown to affect the growth of Ulva spp. and other marine algae (Matsuo et al., 2003). Axenic cultures of the green alga M. oxyspermun and U. linza lack the ability to differentiate naturally. Adding cognate bacterial isolates back to axenic U. linza cultures restores wild type morphology; additionally five isolates were shown to increase the growth rate of the plant (Marshall et al., 2006; Matsuo et al., 2003). Based on the reported effects bacteria have on the growth and morphology of Ulva and on the attraction effects AHLs have on Ulva zoospore settlement the prediction that synthetic AHLs, and

bacteria activity producing AHLs, would increase Ulva zoospore germination was a reasonable assumption. However, this study has shown that Ulva zoospore germination and the early growth of the Ulva germling is reduced when AHLs are present. The effect of AHLs on Ulva zoospore germination was initially observed when exposing germinating Ulva zoospores to varying concentrations of exogenously added synthetic AHLs. High concentrations of C6-HSL, C10-HSL and C12-HSL significantly reduced both the percentage zoospore germination and the length of Ulva zoospore germlings in comparison to when no exogenous AHL was added. Increased consistency in observations of the effect of AHLs on both percentage zoospore germination and average Ulva germling length, was observed when exposing zoospores to high concentrations of C12-HSL. In addition, exposing zoospores to concentrations ranging between 5 µM and 50 µM of C12-HSL also affected percentage zoospore germination and germling length, with germling growth and percentage germination both in general reduced in the presence of C12-HSL in comparison to when no exogenous C12-HSL was present. The increased consistency seen when assaying zoospore germination in response to long chain AHLs can be explained by the fact that AHLs are unstable in seawater and therefore have a short time period of being biologically active. The longer the AHLs fatty acid side chain the more resistant the AHL is to being deactivated by the increased pH of seawater (Hmelo and Van Mooy, 2009). Interestingly, the presence of high concentrations of C8-HSL appeared to have the opposite effect and increase Ulva germling length. This contradictory result may suggest that Ulva zoospores exhibit a differential response to various AHLs, further study using C8-HSL and the 3oxo and 3-hydroxyl AHL derivatives would be required to confirm this speculation. It is also of interest to note that shorter chain AHLs, shown by this study to be produced by Ulva's indigenous bacterial population, had a reduced effect on Ulva zoospore germling length when added exogenously, however the longer chain AHLs which are generally reported to be produced by many marine bacteria had a clear effect, reducing Ulva zoospore germling length. This difference in the effect on Ulva germination between long chain and short chain AHLs may however be due to the high rate of turnover of C4-HSL attributed to relatively high pH of seawater during the experiment as observed in previous studies (Tait et al., 2005). To address this, experiments using RhII-expressing E. coli showed a significant decrease in Ulva germling length. In this experiment the biofilm would have been constantly producing C4-HSL which would replace the hydrolysed signal in contrast with the experiment using exogenously added C4-HSL where replacement of hydrolysed signal did not occur.

Exposing Ulva zoospores to biofilms composed of bacteria which were producing AHLs had the effect of reducing both the percentage of zoospores which germinated and the average length of Ulva germlings, suggesting a reduced growth rate when compared to bacterial biofilms lacking the ability to produce AHLs. Reduced Ulva germling growth and percentage germination in the presence of AHL-producing biofilms was observed with V. anguillarium, E. coli producing AHLs from recombinant AHL synthases and bacteria indigenous to Ulva's population. As previously discussed, biofilms composed of signalling bacteria are producing a continuous source of AHLs therefore maintaining an in situ AHL concentration gradient increasing the reliability of experimental observations of zoospore germination over a 48 h period. Inhibition of algal growth and spore germination by bacteria has been previously reported (Dobretsov et al., 2006). A primary example of marine bacteria which effects spore germination is the gammaproteobacterial species Pseudoalteromonas tunicate, which produces an extracellular component that prevented spore germination in both U.

lactuca and the red alga Polysiphonia spp. (Egan et al., 2001). P. tunicate has been identified as coloniser of Ulva and this study also isolated Pseudoalteromonas from the surface of Ulva (Rao et al., 2006). Additionally a number of bacteria isolated from the marine environment, including a Vibrio isolate have been shown to disrupt Ulva zoospore germination (Bernbom et al., 2011).

Experimental evidence showing AHLs have a reducing effect on Ulva zoospore germination and the early growth of the Ulva germling presents a misnomer; why do AHLs promote zoospore settlement but reduce zoospore germination and growth? This study and studies by Tait et al. have demonstrated that Ulva zoospores and germlings are exposed to a variety of AHL-producing bacteria in the natural environment (Tait et al., 2009; Tait et al., 2005). Therefore, this study speculates that the observation of reduced zoospore germination and germling growth when Ulva is exposed to AHLs and AHL-producing bacteria reflects what takes place in the natural environment as the plant is always going to be exposed to AHL-producing bacteria. The observation that Ulva zoospore germination and growth is greater in the absence of AHLs is most likely an artefact of the experiments conducted by this study as Ulva is unlikely to experience such conditions in the intertidal rocky shore environment. However, in the natural environment fluctuations in the local concentrations of AHLs due to the production of AHL inactivating enzymes by marine bacteria may also have an effect on the germination of the Ulva zoospore and early germling growth. Previous studies have shown Ulva requires a healthy epiphytic bacterial community to form wild type growth morphology (Marshall et al., 2006; Matsuo et al., 2003). This study therefore hypothesises that Ulva uses the AHL signalling by marine bacteria as a cue to reduce early growth in order to become sufficiently colonised by bacteria present in rocky

shore biofilms. The observation by this study that Ulva appears to obtain its epiphytic bacterial community from the biofilms present on the rocky shore lends further credence to this theory. Although this study has clearly shown that AHLs affect not only Ulva zoospore settlement but also Ulva germination, further research is needed to prove the hypothesis that reduced Ulva growth due AHL cues promotes successful colonisation of Ulva. The positive effects with regards to the AHL-producing bacteria of increased Ulva zoospore settlement and reduced zoospore growth are unknown, therefore the relationship between Ulva and bacteria cannot currently be considered as mutualistic. The negative effect of AHLs on Ulva germination could potentially be exploited by industry as an antifouling treatment, as Ulva related biofouling presents a significant burden to the marine economy.

3.3.4 Summary Conclusions and Future Directions

This study has identified the epiphytic bacterial community and species richness present on the thallus of a mature Ulva plant and used these data to select strains for further research. AHL profiling of these selected strains showed bacteria associated with Ulva produce a range of AHL signalling molecules. Additionally, the AHL-mediated interaction between Ulva and bacteria has been shown not just to effect Ulva zoospore settlement but also Ulva germination and early growth of the Ulva germling. In conclusion, it is likely from the phylogenetic study of the Ulva bacterial community that horizontal transmission of bacteria takes place between the rocky shore biofilms and the Ulva plants colonising those rocks. The finding that AHLs reduce zoospore germination and growth is significant as it adds further complexity to the relationship Ulva has with the bacteria present in the marine environment. Further experimentation

is required to investigate why AHLs have such a negative effect on zoospore germination and growth if we are to truly understand this relationship.

Research in this area should progress in the following directions:

- Identification of the mechanism by which AHLs effect Ulva zoospore settlement, germination and germling growth. Previous studies have implicated AHL mediation of calcium efflux to control settlement (Joint et al., 2007). This work could act as a starting point to identify AHL-mediated alterations in Ulva zoospores or Ulva germlings.
- Carrying out comparative protein/RNA expression profiles of Ulva zoospores/germlings exposed to and deprived of AHLs, may also identify where AHL signalling molecules are acting, mediating the behavioural alterations observed by this study and previous work by Tait et al.
- Another area of research would be to identify the levels of AQ production in the
 environment colonised by Ulva. Initial work by this study failed to identify AQ
 production in marine bacteria associated with Ulva, however a targeted LCMS
 approach may reveal AQ production omitted by this study.
- If AQ signalling molecules are present in the environment colonised by Ulva, investigation of their effects on Ulva zoospore settlement, germination and germling growth could be conducted. However, preliminary data not presented in this thesis exposing Ulva zoospores to AQ producing and non-producing P. aeruginosa biofilms did not show any significant differences in germination or germling growth. This work would need repeating to rule out any effect of AQs on the Ulva growth cycle.

Chapter 4

AHL Production and Inactivation in Marine Bacteria Indigenous to the Ulva Epiphytic Community

4.1 Introduction

The green seaweed Ulva possesses an indigenous bacterial population growing in biofilms on the surface of the thallus and the holdfast. This community appears to originate from the rocks colonised by Ulva spp. and the seawater it is exposed to during the tidal cycle. Bacteria from the Altermonadaceae family, to which Shewanella belongs and from the phylum Bacteroidetes, are found in high abundance within this community. These groups of bacteria are also abundant in the biofilms found on rocks in the intertidal zone and in the general marine environment.

4.1.1 The Genus Shewanella

Shewanella spp. were first isolated from spoiled butter in 1931 and identified as Achromobacter putrefaciens, (Derby and Hammer, 1931). However, after much reclassification the genus Shewanella was created in 1985, owing to phylogenetic work from 5S rRNA sequence data. The genus was named after the microbiologist who initially isolated the bacterium, Dr James Shewan (MacDonell and Colwell, 1985). Shewanella bacteria are Gram-negative, rod shaped and lack the ability to form spores. Shewanella cells are in general 2-3 µm in length, 0.4-0.7 µm in diameter and possess a single polar flagellum conferring swimming motility (Venkateswaran et al., 1999). On agar, Shewanella colonies appear smooth and circular with regular edges. Many Shewanella species appear to have a pinkish colouration when grown on agar or in liquid broth. Two species of Shewanella have be shown to be bioluminescent; Shewanella hanedai and an isolate from the Alboran Sea which was designated Shewanella woodyi (Makemson et al., 1997). These species were shown to possess lux

genes homologous to the V. fischeri lux operon, potentially acquired via horizontal transfer (Urbanczyk et al., 2008; Makemson et al., 1997).

To date there are currently 56 typed species of Shewanella (NCBI Taxonomy). The phylogenetic relationship between 39 different species based on 16S rDNA sequencing is shown in Figure 4.1. Shewanella bacteria have been isolated from a variety of different environments and are widely distributed in both marine and fresh water (reviewed by Hau and Gralnick, 2007). Shewanella spp. can grow at a range of different temperatures, however most Shewanella strains display optimal growth at 16-30°C, as a result they have a wide ranging environmental distribution (reviewed by Hau and Gralnick, 2007). A number of species such as Shewanella frigidimarina and Shewanella livingstonensis have been proven to be either psychrotolerant or psychrophillic, growing at temperatures below 5°C (Bozal et al., 2002; Brown and Bowman, 2001). In addition to growth at low temperature, Shewanella spp. such as Shewanella benthica, Shewanella violacea and Shewanella piezotolerans have been proven to be barophillic, growing at high pressures (Wang et al., 2008; Kato and Nogi, 2001).

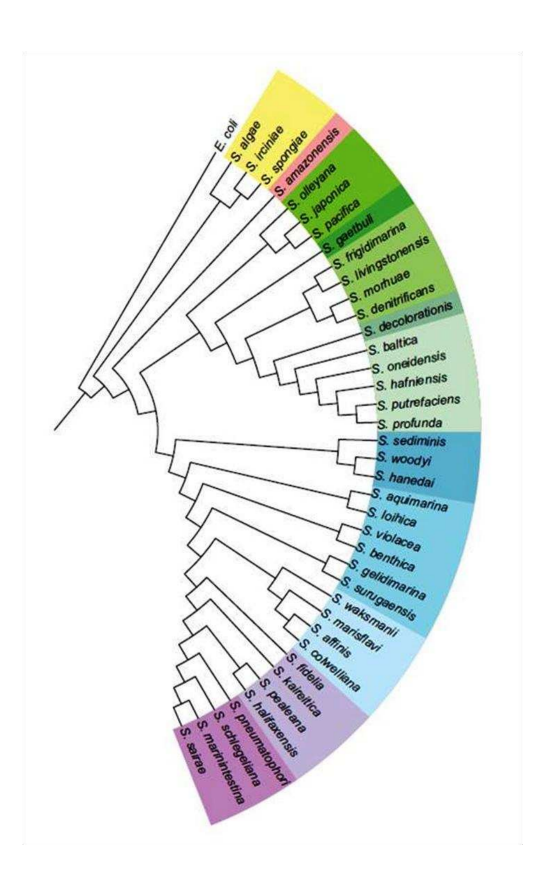


Figure 4.1. Phylogenetic analysis of 16S rDNA from 39 type-strain species of Shewanella. Shewanella abyssi was not considered in this analysis as obtaining 16S rDNA from this species is experimentally difficult. The different colour blocks represent Shewanella species isolated from similar environments (figure sourced from Hau and Gralnick, 2007).

Shewanella spp. are facultative anaerobes and it is their anaerobic metabolism which makes this genus of bacteria of interest to science: Shewanella spp. show a greater diversity in their ability to utilise compounds and elements as electron acceptors than any other group of bacteria (reviewed by Nealson and Scott, 2006). Shewanella spp. do not however have the ability to utilise complex organic carbon sources during metabolism and show a preference for lactate, pyruvate and simple amino acids (reviewed by Venkateswaran et al., 1999; Scott and Nealson, 1994). When growing on lactate in an anaerobic environment, Shewanella spp. release CO₂ and acetate which can be utilised by other anaerobic bacteria such as methanogens. In turn, Shewanella spp. can themselves utilise many metabolic products produced as a result of fermentation by other anaerobic bacteria (Kato and Nogi, 2001).

Shewanella spp. have been shown to reduce more than 20 different electron acceptors and as such sequenced Shewanella genomes have been found to contain a large variety of cytochrome genes in comparison with other bacteria with similar genomes sizes (reviewed by Nealson and Scott, 2006). Shewanella spp. can use Oxygen, Nitrate, Sulphate and elemental sulphur as electron acceptors along with elemental metal oxides including Fe (III), Mn (IV), U (VI), Cr (VI), Tc (VII) and arsenate (reviewed by Nealson and Scott, 2006). Many enzymes involved in the reduction of metal oxides are found on the outer membrane of Shewanella bacteria (Beliaev et al., 2001). Metal reduction by bacteria accounts for the majority of the carbon cycling in a number of different environmental niches, including the marine environment (Lovley and Phillips, 1988). In the central Baltic Sea the majority of the organic carbon cycling at the oxic/anoxic interface is due to the metabolic actions of Shewanella baltica (Ziemke et al., 1997). Shewanella spp. are unique amongst aerobic bacteria as they

have the ability to use elemental sulphur as an electron acceptor producing hydrogen sulphide, as such they can grow at environmental interfaces with sulphate-reducing bacteria such as those found in the Black Sea (Moser and Nealson, 1996).

There have been a number of proposals to exploit the metabolic diversity of Shewanella bacteria for the purposes of bioremediation, (the biotic decontamination of environments contaminated with organic and inorganic pollutants). Using Shewanella spp. to decontaminate land and ground water contaminated with uranium, technetium, cobalt and chromium have been experimented within a laboratory setting (Hau and Gralnick, 2007). In addition to using Shewanella spp. for the bioremediation of environments contaminated with inorganic metal pollutants, it has been proposed to use Shewanella spp., in concert with other bacteria, to decontaminate land contaminated with halogenated solvents, as Shewanella spp. possess the ability to utilise halogenated organic compounds as electron acceptors (Becker et al., 2005; Picardal et al., 1995).

A number of species of Shewanella have been shown to either be pathogenic or cause the spoilage of food. Human disease caused by Shewanella spp. are highly rare opportunistic infections, occurring mainly in geographic regions having warm climates and usually caused by exposure to contaminated seawater (Holt et al., 2005). There have only been two species of Shewanella shown to be associated with human infection; Shewanella algae and Shewanella putrifaciens (Khashe and Janda, 1998). These species have been associated with acute ear infection, skin and soft tissue infection following trauma, acute eye infection and more seriously bacteraemia (Pagani et al., 2003; Butt et al., 1997; Holt et al., 1997; Domínguez et al., 1996). In addition to causing human infection Shewanella bacteria are often implicated in food spoilage.

Shewanella spp. are the dominant organisms found on stored fish products and account for the rotting fish smell due to the production of trimethylamine and hydrogen sulphide (Gram and Huss, 1996). Many of the species that cause food spoilage (i.e. S. putrefaciens) are able to utilise a wider number of carbon sources in comparison to other Shewanella species (Bowman et al., 2000).

4.1.2 AHL Production in Shewanella

AHL production was first characterised in a Shewanella spp. by Tait et al., 2009, in which it was demonstrated that a rock-pool isolate Shewanella sp. B21 produced 3-oxo-C4-HSL, 3-oxo-C10-HSL and 3-oxo-C12-HSL. AHL production was shown to take place during the late exponential growth phase; an unusual result, as AHL production is normally seen in stationary phase bacterial cultures. Shewanella biofilms were also shown to produce AHLs with long chain AHLs being produced during early biofilm growth and short chain AHL production taking place during the later stages of biofilm growth (Figure 4.2) (Tait et al., 2009). AHL production has also been observed in a Shewanella strain (MIB015) isolated from the intestinal tracts of fish (Morohoshi et al., 2008). AHL inhibition by Shewanella spp. during stationary phase has also been observed. Cell free supernatant extracts from cultures over 18 h old failed to activate AHL bio-reporters and disrupted AHL signalling by the Alphaproteobacterial strain Sulfitobacter sp. BR1 (Tait et al., 2009). The turnover of AHL signalling molecules by Shewanella spp. was additionally shown to detrimentally affect Ulva zoospore settlement (Tait et al., 2009).

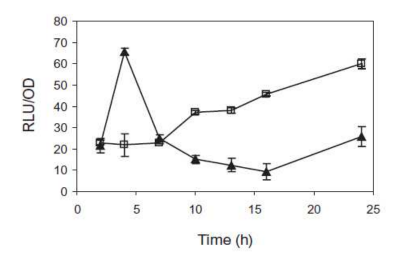


Figure 4.2. AHL production in Shewanella spp. biofilms. Luminescence as a function of growth (RLU/OD) was observed in the lux-based short chain AHL bio-reporter E. coli pSB536 upon exposure to extract from mature Shewanella spp. biofilms (□). In contrast, extract from early growth stage Shewanella spp. biofilms induced luminescence in the lux-based long chain AHL E. coli bio-reporter pSB1075 (▲) (figure sourced from Tait et al., 2009).

4.1.3 The Phylum Bacteroidetes

Bacteria belonging to the phylum Bacteroidetes were first described by Winogradsky as non-spore forming Gram-negative mainly rod shaped bacteria with gliding motility (Kirchman, 2002). After a considerable amount of re-classification the current taxonomic position of the Bacteroidetes phylum was created via the consolidation of the Cytophaga-Flavobacteria/ Flexibacter-Bacteroidetes (CFB) division in 2002 (Garrity et al., 2002; Paster et al., 1985). Following the consolidation of these divisions the newly created Bacteroidetes phylum was taxonomically divided into four separate classes of bacteria; Bacteroidia, Flavobacteria, Sphingobacteria and Cytophagia (Krieg et al., 2010). The foremost characterised class of Bacteroidetes are the Bacteroidia which consist mainly of strict anaerobic bacteria. Members of this class are highly dominant in bacterial populations colonising the human gastrointestinal tract

(Eckburg et al., 2005). The Bacterodia species Bacteroides fragilis is known to be an opportunistic human pathogen causing anaerobic bacteraemia and acute diarrheal disease (Lassmann et al., 2007; Myers et al., 1984). Other Bacteroidia species such as Prevotella intermeddia are commonly found colonising the oral cavity and are associated with oral infections such as dental cares and periodontal abscesses (Tanaka et al., 2008).

Of more interest to environmental microbiology are the Bacteroidetes classes Flavobacteria, Cytophagia and Sphingobacteria. These classes are broad groups of heterotrophic bacteria found colonising fresh water, salt water, soil and sediment environments (Martinez-Alonso et al., 2010; Kormas et al., 2006; Lydell et al., 2004; Eilers et al., 2001). Flavobacteria and Sphingobacteria have also been shown to be associated with marine phytoplankton and marine animals such as sponges (Grossart et al., 2005; Webster et al., 2001). Several Fluorescence in situ Hybridisation (FISH) studies of various open ocean water environments and costal bacterial plankton populations have shown that Bacteroidetes bacteria are the most abundant group of bacteria present (Kirchman, 2002; Cottrell and Kirchman, 2000). In freshwater environments FISH studies have also shown that Bacteroidetes bacteria are highly abundant in biofilm populations (Manz et al., 1999). In addition to studies using FISH to characterise marine bacterial populations, phylogenetic clone libraries of marine environmental niches have also shown high abundances of Bacteroidetes species (Tait et al., 2009; Alonso et al., 2007). Bacteroidetes populations do however tend to be underrepresented in 16S rDNA clone libraries in comparison with FISH based studies, this may be due to the limited copy of the 16S rDNA gene in the genomes of Bacteroidetes species (Alonso et al., 2007).

It is not just the sheer abundance of the Bacteroidetes in the marine environment that makes this phylum of bacteria of interest to science. The Bacteroidetes are metabolically specialised to degrade complex organic matter such as polysaccharides and proteins; an example being cellulose that is insoluble in seawater (Thomas et al., 2011; Edwards et al., 2010). Fibres of cellulose immersed in the Irish Sea were colonised by biofilms composed of Bacteroidetes species that within one month were able to breakdown the cellulose, liberating a source of carbon that could be utilised by other marine organisms higher up the trophic levels (Edwards et al., 2010; Kirchman, 2002). In the ocean, dissolved organic matter (DOM) accumulates into flocculent particles colloquially referred to as marine snow (Alldredge and Silver, 1988). Bacteria readily colonise marine snow particles and can reach densities of between 10⁶-10⁸ cells per aggregate, a figure much greater than that of planktonic bacteria in the marine environment (Munn, 2004). Bacteroidetes have been shown to be highly abundant in these marine snow communities (DeLong et al., 1993), possibly metabolising the DOM in marine snow and hence significantly contributing to the cycling of organic carbon in the marine environment (Kirchman, 2002).

4.1.4 AHL Production in Non Proteobacterial Gram-negative Bacteria

Since the initial discovery of bacterial QS in the marine bacterium V. fischeri the predominant focus of QS research has been centred in the Proteobacteria phylum, with a bigger emphasis on bacterial species that act as human pathogens (Williams et al., 2007; Hastings and Nealson, 1977). Aside from these pathogens, the number of marine species reported to be engaging in AHL-mediated QS, since the initial discovery in V. fischeri, has increased and includes bacteria such as V. harveyi, V. anguillarium,

Aeromonas hydrophyla and Roseobacter sp., (Milton, 2006; Gram et al., 2002; Swift et al., 1999; Hastings and Nealson, 1977). However, in spite of this, little research has been carried out into AHL-mediated QS in other Gram-negative phyla outside the Proteobacteria (reviewed by Williams, 2007). In recent years, the range of bacteria producing AHLs has been extended beyond the Proteobacteria phylum with AHLproducing bacteria being identified in the Cyanobacteria and Bacteroidetes phyla (Romero et al., 2010; Sharif et al., 2008). The initial study to show AHL production by species grouping to the Bacteroidetes phylum was a study in which an strain of Flavobacteria isolated from the North Sea activated the gfp-based short chain AHL bio-reporter E. coli pJBA132. However the specific AHL activating this bio-reporter was not identified (Wagner-Döbler et al., 2005). In a similar study, Huang et al., (2009) showed AHL production in marine biofilms over a number of days using both C. violacium and Agrobacterium tumefaciens AHL bio-reporters. In the same paper the biofilms which activated both reporters were shown to contain bacteria whose 16S rDNA clustered to the Bacteroidetes phylum (Huang et al., 2009). The first AHL identified being produced by Bacteroidetes bacteria was C4-HSL which was being synthesised by a number of strains of Tenacibaculum maritimun (Romero et al., 2010). Studies have also showed that T. maritimum had the ability to quench the activity of C10-HSL (Romero et al., 2011; Romero et al., 2010).

4.1.5 The Relevance of Shewanella and the Bacteroidetes within the Ulva Epiphytic Population

The reason for selecting the Bacteroidetes phylum for further studies in this thesis was primarily due to this phyla's abundance within the Ulva spp. epiphytic

population as seen in the 16S rDNA clone library. In particular, 36.49% of clones in the library grouped to the Bacteroidetes phylum (see Section 3.2.1 and Figure 3.3). Additionally a Bacteroidetes strain (Cytophaga sp. MBIC 04683) was shown to produce the plant growth effector thallusin, implicated to affect to the growth morphology of Ulva spp. (Matsuo et al. 2005).

The Altermonadaceae, the family to which Shewanella belongs to are also abundant within the Ulva spp. epiphytic population with 9.21 % of clones from this library grouping to the Altermonadaceae, this figure represents 41.18 % of the clones grouping to the Gammaproteobacteria phylum within the Ulva spp. 16S clone library (see Section 3.2.1 and Figure 3.3). These two groups of bacteria were also found in abundance within the biofilms present on the rocks where Ulva spp. can be found colonising (Tait et al., 2009). The Shewanella are of particular interest as they can be isolated from both rocky shore biofilms and from the Ulva thallus (Tait et al., 2009). Shewanella isolates from both these niches have been shown to produce AHLs and interestingly turn over their cognate signal and inhibit the signalling of other AHL-producing bacteria found in the rocky shore biofilms, affecting Ulva zoospore settlement (Tait et al., 2009). Additionally AHL synthase and response regulator genes have not, as yet, been identified in the genus Shewanella. All these above factors make Shewanella bacteria of interest for further study and relevant to this thesis.

4.1.6 Experimental Aims

Bacterial belonging to the Bacteroidetes phylum and the genus Shewanella have been shown to be abundant within the epiphytic bacterial population of Ulva and interact with the seeweed affecting the Ulva growth process. The aims of this study were to further investigate the extent of AHL production and inhibition by members of these two groups of bacteria. The presence of AHL synthases, AHL response regulators and AHL inhibiting enzymes within both Shewanella spp. and the Bacteroidetes phylum was investigated using in silico analysis of genomic sequences. Owing to previous research that showed AHL inhibition by Shewanlla spp. to directly effect Ulva zoospore settlement, AHL inhibition by members of the Shewanella genus was therefore investigated using a number of molecular biological techniques.

4.2 Results

4.2.1 AHL Production in Bacteroidetes and Shewanella

4.2.1.1 AHL Production in Shewanella

AHL production has already been reported in Shewanella isolates from the rocks colonised by Ulva spp. and from fish intestines (Tait et al., 2009; Morohoshi et al., 2008). This study utilised a number of Shewanella strains provided by Dr K Tait isolated from Ulva spp. and decaying fish. These strains were identified by 16S rDNA typing and screened for AHL production using the lux-based bio-reporters E. coli pSB536, pSB401 and pSB1142. AHL bio-reporter assays showed that these strains were producing both short and long chain AHLs as cell free supernatant extracts activated both the bio-reporters pSB536 and pSB1142 (Table 4.1).

Table 4.1 Shewanella strains screened for AHL production. A number of Shewanella strains isolated from decaying fish were identified by 16S rDNA typing and screened for AHL production using AHL bio-reporters.

					Bio-reporter Activation *		
Strain	Phylogenetic Identity	Accession Number	% Identity	Source	pSB536	pSB401	pSB1142
A1	Shewanella baltica gene for 16S rRNA, partial sequence, strain: X1410	AB205580	99	Fish	+	-	+
A2	Shewanella sp. CsQ2 16S Ribosomal RNA Gene	EU075118	98	Fish	+	-	+
A4	Shewanella baltica strain KB30 16S Ribosomal RNA Gene	JF327458	98	Fish	+	-	+
A7	Shewanella sp. MPU12 gene for 16S ribosomal RNA, partial sequence	AB334772	98	Fish	+	-	+
A8	Shewanella sp. MPU12 gene for 16S ribosomal RNA, partial sequence	AB334772	97	Fish	+	-	+
B1	Shewanella putrefaciens SS6 16S ribosomal RNA gene, partial sequence	JX032786	97	Fish	+	+	+
B8	Shewanella baltica 16S rDNA X1410	AB902206	98	Fish	-	-	+
P3 †	Shewanella sp. 16S ribosomal RNA P3	AY902206	100	Rock	+	-	+

^{*} Strains marked with a + were shown to induce bioluminessence in the bio-reporter, strains marked with a - failed to induce bioluminessence in the bio-reporter.

[†] Data provided by Dr. Karen Tait

4.2.1.2 AHL Production in the Bacteroidetes

The 16S rDNA phylogenetic typing showed that a number of strains isolated from Ulva spp. were Bacteroidetes bacteria. Strains UI08 and UI13, isolated from the Ulva thallus were identified as being Cellulophaga sp. members of the Flavobacteria class of Bacteroidetes. Phenotypically both UI08 and UI13 possessed similar colony morphology of bright yellow-pigmented rough circular colonies. After several days incubation on the bench at $\approx 22^{\circ}$ C both strains spread across plates and UI08 showed agarlytic activity. Acidified cell free supernatant extracts from strains UI08 and UI13 showed activation of the E. coli pSB536 AHL bio-reporter suggesting the presence of short chain AHLs (see Section 3.2.4 and Table 3.3).

In order to positively identify the specific AHLs being produced by these Bacteroidetes strains, solvent extracts from cell free stationary phase cultures of both UI08 and UI13 were analysed by LCMS. In addition to these two Bacteroidetes strains isolated from the Ulva thallus the stationary phase cell free supernatant extract from Bacteroidetes strain P13, isolated from biofilms found on rocks colonised by Ulva sp. were also assayed for short chain AHL production using LCMS. Extraction ion chromatograms and their corresponding compound mass spectra from the LCMS analysis showed all three strains produce C4-HSL (Figure 4.3). LCMS analysis also showed the presence of C4-HSL with the lactone ring opened, indicating that a proportion of C4-HSL was being inactivated by either the alkaline pH of marine broth media actions AHL lactonase. or by the of an

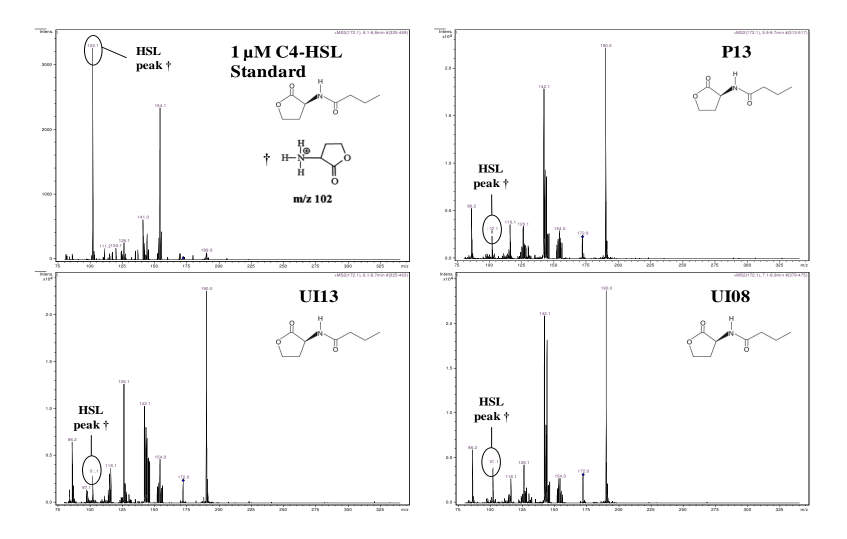


Figure 4.3. LCMS analysis of Bacteroidetes strains isolated from Ulva spp. and rocky shore biofilms. Compound mass spectra from the breakdown of precursor ion m/z 172 (C4-HSL) of C4-HSL standard P13, UI13 and UI08. Daughter ion of m/z 102.1 (HSL) labelled in all spectra.

The Bacteroidetes strains isolated from Ulva spp. which were shown to produce C4-HSL all grouped within the Flavobacteria class. As the Bacteroidetes population present on Ulva spp. also contains bacteria from the Sphingobacteria class, it was decided to look for AHL production in strains of Bacteroidetes from a wider range of phylogenetic backgrounds. A number of Bacteroidetes strains from a range of phylogenetic backgrounds were selected from the PML culture collection and the National Collection of Industrial, food and Marine Bacteria (NCIMB) to be screened for AHL production, (Table 4.2).

Table 4.2. List of Bacteroidetes strains from varying phylogenetic backgrounds obtained from the PML and NCIMB culture collections.

Strain	Phylogenetic Identity	Phylogeny	Accession Number	% Identity	Source
1363	Saprospira grandis 16S rRNA partial sequence	Sphingobacteria	AB071781	99	Rock pool
476	Flexibacter aggregans gene for 16S rDNA IFO15975	Cytophagia	AB078039	95	Seawater
14225	Algoriphagus chordae Strain LMG 21970 16S ribosomal RNA	Cytophagia	NR025603	98	Algae
397	Cytophaga marinoflava 16S ribosomal RNA gene	Cytophagia	AF203475	98	Seawater
14312	Lewinella marina MKG38 16S ribosomal RNA partial sequence	Sphingobacteria	NR041594	99	Marine Sediment
699	Balneola alkaliphila CM41_14b 16S ribosomal RNA	Sphingobacteria	NR044367	92	Seawater

All strains listed in table 4.2 were screened for AHL production using LCMS with the exception of strain 699. Strain 699 was discounted from this screening as it displayed exceedingly slow growth, only reaching an OD_{600} of 0.3 in 14 days. In all

Bacteroidetes strains screened only C4-HSL was detected in the supernatant extracts, (Figure 4.4).

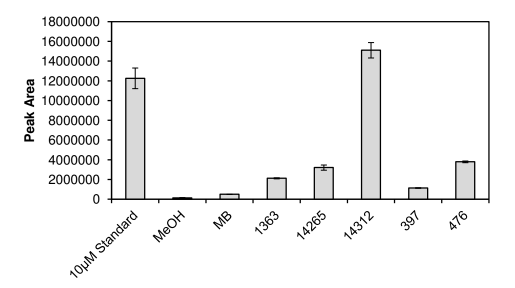


Figure 4.4. Screening of Bacteroidetes strains of varying phylogeny for AHL production via LCMS. Graph showing the areas of the C4-HSL extraction ion chromatogram peaks from the LCMS analysis of Bacteroidetes strains of differing phylogeny. As controls synthetic C4-HSL, extract from marine broth (MB) and Methanol were used. Error bar represent standard deviation from the mean peak area.

4.2.2 In silico search for Shewanella spp. AHL Synthase and AHL Response Regulator Sequences

Annotated and semi-annotated Shewanella genome sequences were screened for AHL synthase and AHL response regulator protein sequences. To accomplish this, AHL synthase and AHL response regulator consensus sequences were compiled from the majority consensus of ClustalW alignments of 24 published LuxI and 31 published LuxR homologues (Appendices 4 and 5). Consensus sequence were aditionally compiled from the majority consensus of ClustalW alignments of LuxM homologues from V. fischeri, V. anguillarium and V. harveyi (Milton et al., 2001, Hanzelka et al.,

1999, Bassler et al., 1993), and 10 published LuxN protein sequences (Appendices 6 and 7). This enabled the screening for the LuxM/N-type two component AHL synthese and response regulator systems. All 4 consensus sequences, (Figure 4.5), were used to screen protein sequences from 19 separate Shewanella genomes using the Genomic BLASTp program (NCBI).

LuxI Consensus Sequence

MLXIFDXXXVSYDXLSEEKXXXKELFRLRKKXFKDRLGWDVNCXNGMXXXEFDXXQYDNXNTRYLLGIXD XXXGQXVGCVRLIETTXXXPNMLTGTFFXLLXDDGALPEXXGXYIXESSRFFXXXDKARARLLXGNXXXY PLSLVXLFLSMINYARANGYTGIVTVVSRAMERILKRSGWPIERIGQXXGHXXEKGEXIYLXLHLPIDXX XQERLARRINQPLQGPSSXLLTWPLSLP

LuxR Consensus Sequence

LuxM Consensus Sequence

MLSLLSXXXXXXPVQDSCPTLVASALIQNWSVRDTWLSFTYAPQXXNYCFPSYGYSEFTRLQLFTPSSL SKCYXXEFDNEFKXQLSDTQAVCEVFTLRLTVXXXXYFLYLAQKELMSVLHQAGYKXXXXXXXIIEQPFM LNFYRAIDAKAYFHSFTGYCDLNDDGKQTYRGFWNFEMMVKAFSNIDFRGYKRXRASRKRGSLERDEHV

LuxN Consensus Sequence

Figure 4.5. AHL synthase and AHL response regulator consensus sequences. The consensuses were generated from the majority consensus of ClustalW sequence alignments of multiple LuxI, LuxR, LuxM and LuxN homologues, X represents regions where no majority consensus was present in the alignment (see Appendices 4-7).

BLASTp screening using the LuxI-type AHL synthase consensus sequence yielded no hits to any sequences within the 19 Shewanella genomes searched. BLAST screening using the LuxR type AHL response regulator consensus sequence yielded 164 hits to sequences within Shewanella genomes. Most of these hits possessed very low similarity scores (< 25%) to the consensus sequence and any similarity was due to the presence of helix-turn-hexix DNA binding domains. The top hit (percentage Identity 34%), corresponded to SVI_2459, which was annotated as a LuxR family response regulator in Shewanella violacea DSS12. This sequence had regions of similarity to the LuxR consensus sequence in both the C-terminal and N-terminal regions. Structural analysis of this sequence using the Pfam database (Punta et al., 2012), showed both an AHL-binding domain and a LuxR type DNA-binding domain to be present, characteristic of the LuxR family of proteins (Kolibachuk and Greenberg, 1993). Analysis of the upstream and downstream ORFs of SVI_2459 using Artemis genome viewer showed no sequences with any similarity to the LuxI consensus.

BLASTp screening using the LuxM-type AHL synthase consensus sequence produced three hits in the Shewanella genomes. These hits possessed very low sequence identity and amino acid coverage values, (28% and 27% respectively), so are therefore unlikely to be true LuxM homologues (data not shown). BLAST screening using the LuxN type AHL response regulator consensus sequence yielded a number of hits within the Shewanella genomes. These hits possessed sequence identity values of up to 46% and amino acid coverage values of up to 52%. However, the genomes of these top 5 BLAST hits lacked any sequences resembling LuxU and LuxO which are key elements in the phospho relay cascade integral to the function of the two

component AHL response regulator system (Freeman and Bassler, 1999a; Freeman and Bassler, 1999b). It is therefore most likely that these hits are not true LuxN homologues.

4.2.3 In silico Search for Shewanella spp. AHL Inactivating Enzyme Sequences

In order to screen for potential AHL-inactivating enzymes, consensus sequences were constructed from protein sequences of published AHL lactonases and AHL acylases, (Appendix 8), using the same approach as with the construction of the LuxI/R consensus sequences (see Section 4.2.2). The resultant consensus sequences (Figure 4.6) were used to screen 19 separate Shewanella genomes. No potential AHL lactonase sequences were identified within these Shewanella genomes. Using the AHL acylase consensus sequence 26 hits were identified within the Shewanella genomes including S. putrefaciens, S. baltica and Shewanella amazonensis. With the exception of one, all the identified sequences possessed between 29-40% identity with the consensus and 90-94% amino acid sequence coverage. The top 5 hits are displayed in table 4.3.

AHL Lactonase Consensus Sequence

MXXXXXKLXXLQSGXXRCDXXHISMNXGXXXXGKXYXXXXXXXEIPVPAXLIXHXDGFTLIDTGLXXEG AXDPSGRWXXGSXXXXQXXPXXXXSXEQGCVXEQLKXXXGLXPXDIXYVVLSHLHLDHAGAIGRFPNAXH IVQRXEYEYAFAXXXWFXXGAXXXXYIRKDFXXXXXXLPXLNWQFXNGIEDDRYXVXPGVXXTLLXXFTP GHXPGHQSXLVRLPKXGPFLLTIDAAYTLDXXXXXXALAGXLXXTIDXXQSVQKLRQXAERYXATVIFGH DPEQWARFXKXAPXFYX

AHL Acylase Consensus Sequence

Figure 4.6. AHL inactivating enzyme consensus sequences. The consensuses were generated from the majority consensus of ClustalW sequence alignments of multiple AHL lactonase and AHL acylase homologues, X represents regions where no majority consensus was present in the alignment (Appendix 8).

Table 4.3 BLAST hits resulting from screening Shewanella genomes for AHL inactivating acylases. Table listing NCBI accession number, sequence annotation, percentage identity, and percentage amino acid coverage values of the top 5 AHL acylase BLAST hits from the screening of Shewanella genomes.

AHL Acylase					
Accession Number	Sequence Annotation	%	%		
		Identity	Coverage		
YP_964683	Peptidase S45, penicillin amidase,	39	93		
	Shewanella sp. W3-18-1				
YP_006008798	Acyl-homoserine-lactone acylase, S.	39	94		
	putrefaciens 200				
YP_001182388	Peptidase S45, penicillin amidase, S.	39	94		
	putrefaciens 200				
NP_716547	aac gene product, S. oneidensis MR1	40	93		
YP_001367743	Peptidase S45, penicillin amidase, S.	40	93		
	baltica OS185				

4.2.4 In silico Search for Bacteroidetes AHL Synthases and AHL Response Regulator Sequences.

Having identified AHL production in a number of Bacteroidetes strains, annotated and semi-annotated Bacteroidetes genome sequences were screened for AHL synthase and AHL response regulator protein sequences using the LuxI, LuxR, LuxM and LuxN consensus sequences compiled for the in silico analysis of Shewanella genomes (Figures 4.5). These consensus sequences were used to screen protein sequences from 180 separate Bacteroidetes genomes using the Genomic BLAST program (NCBI).

BLAST screening using the LuxI-type AHL synthase consensus sequence yielded a total of 5 hits all of which had a low % similarity and % coverage score (< 38% and < 36% respectively). Similarity to the consensus was only apparent to the C-terminal ends of the identified hits. Aligning the BLAST hits to published LuxI homologues showed that no sequence similarity was present in the reported active site region located between residues 25 and 70 (Hanzelka et al., 1997). It is therefore unlikely that these hits are true LuxI homologues.

BLAST screening using the LuxR-type AHL response regulator consensus sequence yielded over 100 hits all of which had a slightly higher but still low % similarity and % coverage score in comparison to the LuxI hits (< 50% and 24% respectively). Again, these hits only possessed regions of similarity to the C-terminal end of the consensus. Structural analysis using the Pfam database (Punta et al., 2012) of the top 5 BLAST hits showed LuxR-type DNA-binding domains to be present within the protein sequences however all 5 hits lacked AHL-binding domains, characteristic of

the LuxR family of proteins (Kolibachuk and Greenberg, 1993). Additionally, there were no luxI-type predicted gene sequences near these BLAST hits suggesting the absence of any LuxI/R system within the genomes searched.

BLAST screening using the LuxM-type AHL synthase consensus produced one hit in the Bacteroidetes genomes. This hit possessed very low sequence identity and amino acid sequence coverage (36% and 21% respectively). It is therefore unlikely that these hits are true LuxM homologues. BLAST screening using the LuxN-type AHL response regulator consensus sequence produced over 135 hits in Bacteroidetes genomes with many having high levels of similarity over the entirety of the sequences identified (48% and 49% respectively). Structural analysis using the Pfam data base, (Punta et al., 2012), of the BLAST hits showed them to be transmembrane proteins containing phosphokinase domains. The top 5 BLAST hits were added to an alignment of published LuxN homologues compiled by Swem et al. 2008. The BLAST hits lacked many of the conserved residues that are thought to be required for the activity of LuxN (Swem et al., 2008). As with the Shewanella LuxN BLAST hits the genomes of these Bacteroidetes top 5 BLAST hits also lacked any sequences resembling LuxU and LuxO and therefore again it is most likely that these hits are not true LuxN homologues.

4.2.5 In Silico Search for Bacteroidetes AHL-Inactivating Enzymes

AHL inactivation by Bacteroidetes bacteria has been reported (Romero et al., 2011, Romero et al., 2010). Therefore, in addition to carrying out in silico screens for putative AHL synthase and response regulator sequences, a screen for AHL-inactivating enzymes was conducted. The AHL lactonase and AHL acylase consensus

sequences used to screen Shewanella genomes (Figure 4.6) were used to screen 180 separate Bacteroidetes genomes. The AHL lactonase consensus sequence produced 53 BLAST hits in Bacteroidetes genomes. The top 5 hits are displayed in Table 4.4 and have a percentage identity score with a 14% - 59% range and a percentage amino acid coverage score with a 46% to 24% range.

Table 4.4. BLAST hits resulting from screening Bacteroidetes genomes for AHL inactivating lactonases. Table lists NCBI accession number, sequence annotation, % Identity and % amino acid coverage values of the top 5 AHL lactonase BLAST hits from the screening of Bacteroidetes genomes.

AHL Lactonase Hits						
Hit Number	Accession Number	Sequence Annotation	% Identity	% Coverage		
BLAST Hit 1	ZP_01051078.1	Metallo-beta-lactamase superfamily protein [Dokdonia donghaensis MED134]	59	25		
BLAST Hit 2	YP_004430745.1	AttM/AiiB family protein [Krokinobacter sp. 4H-3-7-5]	63	27		
BLAST Hit3	YP_003715653.1	AttM/AiiB family protein [Croceibacter atlanticus HTCC2559]	60	24		
BLAST Hit 4	ZP_01692265.1	Zn-dependent hydrolase GumP [Microscilla marina ATCC23134]	47	25		
BLAST Hit 5	ZP_03967553.1	possible metal-dependent hydrolase [Sphingobacterium spiritivorum ATCC 33300]	14	46		

The top 5 BLAST hits were added to a protein sequence alignment containing the sequences of 3 published AHL lactonases; AiiA from Bacillus sp. 240B1. (Dong et al., 2000), AiiB from Agrobacterium tumefaciens and AttM from A tumefaciens (Carlier et al., 2003). The sequence alignment (Figure 4.7) showed that all five hits contained high levels of sequence similarity and that all 5 hits contained the conserved motif HXHXDH~H~D~H, a structural motif encompassing the Zn-binding domain

previously shown to be integral for the function of an AHL lactonase (Liu et al., 2007; Dong et al., 2000).

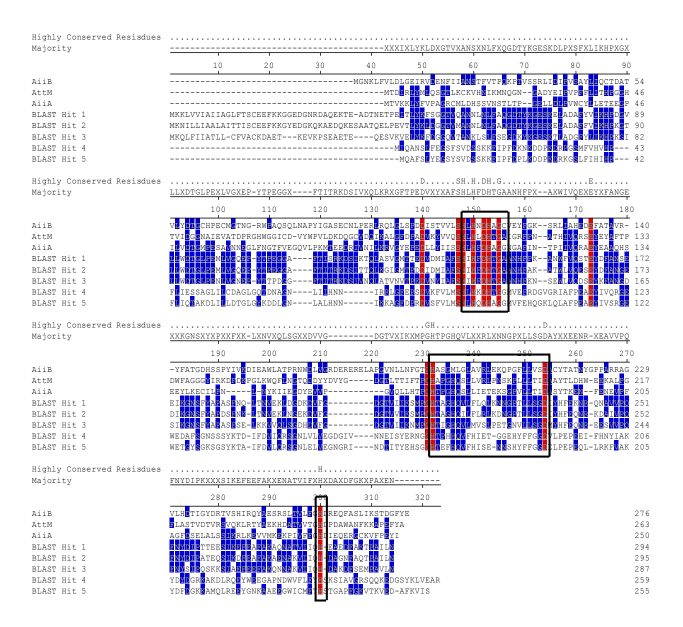


Figure 4.7. Protein sequence alignment of AHL lactonase BLAST hits with the AHL lactonases; AiiB, AttM and AiiA. BLAST hits 1-5 (see Table 4.4) identified by screening Bacteroides genomes using the AHL lactonase consensus sequence were aligned against published AHL lactonase sequences. Areas of sequence similarity are highlighted in blue. Residues highlighted in red are where sequence similarity is conserved throughout the alignment. The residues enclosed by black boxes indicate conserved residues that form the Zn-binding domain present in AHL lactonases.

Screening Bacteroidetes genomes using the AHL acylase consensus sequence produced 59 BLAST hits. The top 5 hits are displayed in Table 4.5 and have a percentage identity score with a 29%-47% range and a percentage coverage score with a 24% to 29% range.

Table 4.5. BLAST hits resulting from screening Bacteroidetes genomes for AHL inactivating acylases. Table listing NCBI accession number, sequence annotation, % identity and % amino acid coverage values of the top 5 AHL acylase BLAST hits from the screening of Bacteroidetes genomes.

AHL Acylase Hits							
Hit Number	Accession	Sequence Annotation	%	%			
	Number		Identity	Coverage			
BLAST Hit 1	YP_003194092.1	Penicillin amidase superfamily protein	30	24			
		[Robiginitalea biformata HTCC2501]					
BLAST Hit 2	ZP_01692595.1	Glutaryl 7-aminocephalosporanic acid acylase [Microscilla marina ATCC 23134]	29	24			
BLAST Hit3	ZP_03701247.1	Peptidase S45 penicillin amidase [Flavobacteria bacterium MS024-3C]	47	29			
BLAST Hit 4	YP_445269.1	Penicillin amidase superfamily protein [Salinibacter ruber DSM 13855]	43	29			
BLAST Hit 5	YP_003571196.1	Penicillin amidase [Salinibacter ruber M8]	43	29			

The BLAST hits from the AHL acylase screen were aligned with the AHL acylase sequences of AiiD from Ralstonia sp. XJ12B (Lin et al., 2003), PvdQ from P. aeruginosa PAO1 (Huang et al., 2003) and AhlM from Streptomyces sp. M664 (Park et al., 2005). The alignment (Figure 4.8) shows areas of sequence similarity between the BLAST hits and the known AHL acylase sequences. Additionally there is a highly conserved glycine-serine paring present in all the BLAST hits that is present in AHL

acylases and which has been previously shown to be integral to the post translational modification of the acylase propertide (Lin et al., 2003).



Figure 4.8. Protein sequence alignment of AHL Acylase BLAST hits and known AHL acylases; AiiD, PvdQ and AhlM. BLAST hits 1-5 (see Table 4.5) identified by screening Bacteroides genomes using the AHL acylase consensus sequence were aligned against published AHL acylase sequences. Areas of sequence identity are highlighted in blue. Residues highlighted in red are regions of sequence identity conserved throughout the alignment. The residues enclosed in black boxes indicate conserved glycine-serine pairing common in acylase sequences.

4.2.6 AHL Inactivation by Shewanella sp.

Shewanella spp. have been previously shown to produce AHLs in mid log phase and then to inactivate them during stationary growth phase (Tait et al., 2009). To identify the gene responsible for AHL inactivation, a genomic library of Shewanella sp. P3 chromosomal DNA, a strain shown previously to have good AHL inactivating activity (Tait, un-published data), was constructed via restriction digest, transformed into an E. coli host and screened using the Quorum Sensing Inhibitor Selector (QSIS) system present on plasmid pMH655. The plasmid pMH655 constitutively expresses the lasR AHL response regulator gene and also possesses the levansucrase gene sacB, which is under the control of the LasR regulated promoter, PlasB (Rasmussen et al., 2005). The Shewanella sp. P3 library was screened by growing transformant colonies on media containing sucrose, and seeded with exogenous 3-oxo-C12-HSL. Bacteria possessing the QSIS system and no AHL inhibitor sequence would not survive, as the exogenous 3-oxo-C12-HSL will be transduced by LasR, activating P_{lasB} allowing the expression of sacB resulting in cell death due to the presence of the sucrose in the media. If an AHL inactivating enzyme sequences was present, exogenously added AHLs will be inactivated and the sacB gene will not be expressed, resulting in cell survival in the presence of sucrose. After screening approximately 6000 transformant clones, 6 clones appeared to have increased growth after an 18h period and as such were selected as potentially having AHL inactivating activity.

The six clones putatively possessing a gene encoding an AHL inactivating enzyme were grown in the presence of synthetic 3-oxo-C12-AHL. This AHL was selected as a previous study showed that Shewanella spp inactivate 3-oxo-C12-HSL during stationary phase (Tait et al., 2009). The lux-based AHL bio-reporter E. coli

pSB1142 was used to detect any inactivation of the signal molecule in the culture supernatants of these clones using a microtitre plate reader over a 24h period. Inactivation of AHLs would present itself as a loss or absence of luminosity by the bioreporter over time. The six clones were compared to a positive control, an E. coli strain carrying the QSIS system and expressing the AHL inactivating lactonase gene aiiA, (E. coli JM109 pSU18aiiA pMH655). Of the six clones assayed which had been incubated with 3-oxo-C12-HSL, supernatants from clones 1 and 4 were shown not to cause any activation of the bio-reporter. The AHL inactivating positive control strain showed an initial activation of the bio-reporter followed by a fall in luminescence indicating AHL inactivation. Clones 2, 3, 5 and 6 showed activation of the bio-reporter, as did the negative control E. coli DH5α pMH655 which had been seeded with 3-oxo-C12-HSL (Figure 4.9).

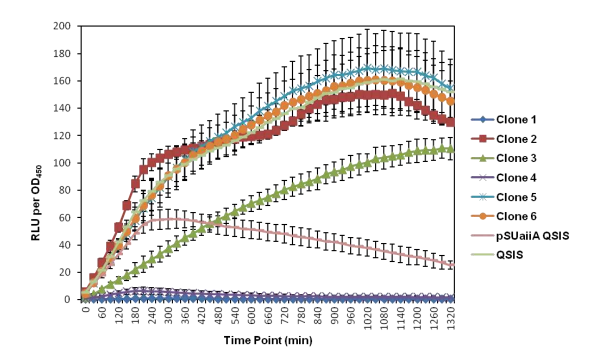


Figure 4.9. AHL degradation assay of Shewanella sp. P3 genomic library clones. Clones possessing suspected AHL inactivating gene sequences as identified through QSIS screening were assayed for AHL inactivation in the presence of 3-oxo-C12-HSL over a 24 h period at 37°C.

Clones 1 and 4 which appeared to show AHL inactivation were subjected to miniprep plasmid extraction in order to detect insert DNA within the pME6000 shuttle vector. Although the clones appeared to show AHL inhibiting activity using the bioreporter neither of the clones possessed insert DNA so were therefore false positive results.

4.2.7 Putative AHL Acylase AacS

4.2.7.1 Screening Shewanella Strains for the aac Gene

Screening a Shewanella genomic library for AHL inactivating enzymes produced a number of false positive results and was therefore abandoned. Previous work by Tait et al. (2009) showed that AHL inactivation in Shewanella takes place via the actions of an acylase enzyme which cleaves the acyl side chain from the homoserine lactone ring rendering the AHL biologically inactive (Tait et al., 2009). A putative AHL acylase has been identified in a Shewanella strain isolated from a fish intestinal tract in Japan (Shewanella sp. MIB015). This acylase showed high levels of similarity to the Aculeacin A acylase gene aac in S. oneidensis strain MR1 (Morohoshi et al., 2008). Using degenerate primers designed from sequence data of the upstream and downstream regions of putative AHL acylases found within genomic sequences of S. baltica, S. putrefaciens and S. oneidensis the isolates listed in Table 4.6 were screened by PCR for potential AHL inactivating enzymes with similarity to aac. The majority of strains screened produced PCR fragments of approximately 2.5 Kb; the expected size of the Shewanella aac gene. These fragments were cloned into the vector pGEM T easy, transformed into E. coli DH5a and subsequently sequenced using the universal primers M13F and M13R. The resultant sequences were compared to the

NCBI nucleotide database using a BLAST search and found to be homologous to either aac from S. oneidensis MR1, or to putative AHL acylase genes identified during genomic sequencing of various Shewanella species (Table 4.6).

Table 4.6. Shewanella strains screened for the presence of the aac gene. Table listing the Shewanella strains screened via PCR for the aac gene, their phylogenetic identity as identified by 16S rDNA typing, where they were sourced from and whether they possessed aac homologues.

Strain	Phylogenetic Identity	Accession Number	% Identity	Source	aac Gene †
A1	Shewanella baltica gene for 16S rRNA, partial sequence, strain: X1410	AB205580	99	Fish	+
A2	Shewanella sp. CsQ2 16S Ribosomal RNA Gene	EU075118	98	Fish	+
A4	Shewanella baltica strain KB30 16S Ribosomal RNA Gene	JF327458	98	Fish	-
A7	Shewanella sp. MPU12 gene for 16S ribosomal RNA, partial sequence	AB334772	98	Fish	+
A8	Shewanella sp. MPU12 gene for 16S ribosomal RNA, partial sequence	AB334772	97	Fish	-
B1	Shewanella putrefaciens SS6 16S ribosomal RNA gene, partial sequence	JX032786	97	Fish	+
B8	Shewanella baltica 16S rDNA X1410	AB902206	98	Fish	-
E1	Shewanella sp.		99	Fish	+
NCIMB 2157	Shewanella hanaidai	N/A	N/A	NCIMB	+
NCIMB 13526	Shewanella woodyi	N/A	N/A	NCIMB	+
MR1	Shewanella onediensis	N/A	N/A	PML	+

[†] Strains possessing an aac homologues are marked with +, strains not found to possess an aac homologue are marked with -

4.2.7.2 AHL Inactivation by the Shewanella putrafaciens B1 aac Clone

The putative AHL inactivating gene clone pGEMB1aac from strain B1 (identified as S. putrefaciens), harboured by E. coli DH5α was selected for further study. The DNA sequence and translated protein sequence of this putative AHL acylase from S. putrefaciens B1 (named aacS and AacS respectively) can be seen in Figure 4.10.

aacS DNA Sequence 2,550bp

ATGAAATTCAACAAACTCGTGATTGCTATGGGAATGGCCTGCGGTGTAATACTGACCGGCTGTAACGATA GCGAAGATAGCACTACGCCTACCGAACCTGAAACTCAACTGCAAGCTTTTGCCCCCAATGGTTTACTCAA AGCCAGTATTCGCCGTACTACCTTTGGCGTGCCGCATATCCAAGCAGACAACTTAGAAAGTTTAGGTTTT GGTAGTGGTTATGCGCAGGCACAGGACAACTTATGTGTATTAGCCGATGGTTTTATCAAGGCGAACTCAC AGCGTTCTATGTACTTTGGTCCCCATGCGTCGATTGACTTCACTACGGGTCAACCTACGGCGGAAGATAA CGGTAACCTTATCTCAGATTTTGCCTATAAAGCGTTAAAGATCAGAGCGCAAGCCGAAGAAAAATGGCCG AAATTTAGTGAAAACTCTCGGGCACTTATCCAAGGTTTTACCTCTGGTTATAACCAATATCTTGCTGATG TCGAAGCGGGCACACAAACGGCAGAACCTTTCTGTGGCGGTCAGCCTTGGGTGAAACCCATAGTGCCAGA GGATGTGGTGACTTATTTGTTTTCTATCGCCTTATTACCGGGCGCAGCTAACTTTCTCGATCTGATTTTT TACGCTAACCCAGGAGATGCACAGGAATACATGCCGCGTATCGTTGGGCCCGCGCCTAGCCAAGACCAAA CTAGGTAATCCGCATTTCCCGCATACGGGTAACCTGCGTTTTTGGCAATCCCATATTACGATCCCAGGGC ATTTAGATATGATGGGCGGCTCGTTAGTGGGTATGCCTGGACCGATTAATATTGGTTTCaATAAAGATCT $\tt CGCTTGGACTCATACCTTCTACCGCTGAGCATTTTGTGATGTATAACCTTGAGTTAGTCTCGGGTGAT$ CGGATGCAGTATTTGTTTGATGGTAAACCTATGCCGATCACCAAAGAGACAGTATCGGTTCTAGTGAATG CAGGTCCTGCGGGCATGCTGGTCGCCGAGAAAGATATTTATACTACGGCCAAAGGCCCTATGGTCGAAGC TCCTCCTGCTTTAGCGCCTTTTGGTTGGGATGATGGCAGTGCCTTTATGATCCAAGATGCCAATATGGGG ACTATGGACCCTGTTGACCATTGGTTAGCCATGAACATGGCGACGAACAAGAAGAGTTCCAACAGGCAT TTAAGGACTACGATGGTGTCATCTTCAATAACACTATGTACGCCGATAAAGAGGGCAATGCTTTTTATAT CGACGACTCAACTGTGCCAGGATTATCTGAATCGGCAGTTGTGTTGTTAAAAAACCTCACCGGATATTAAA GCGGCTAAGGCAAAAGCCGGATTTACGATTTTACCCGGTAACACTTCGTTGTTTAGCTTCGATGGTCCTA CGCCCTATGAGCGTGCGCCCAAGCTTGAGCGCAGCGACTTTGTGCAAAACTCCAATGACTCCTTTTGGTC GACTAACTTGAATGAACCGCTGACTTACTACTCGCCTATTTTACGGCGCAGAAGGCTGGACAGCTATCAC TGCGGACACGGATGGGCTATGCCGCGGGGGGCTGATGGCAAGTTCAGCTGGAAGAACTCGAAGCGGGCTG TGCTGTCGAATCGCAGTTATCTCGCCGAGTTAGTGCTGCCTGACTTGATTGCCCAGTGTGATGCCCAAGG CAGTACACCTGTGGTGGTGTCGGCAAGTTTATCTAAGGATGTCTCTTTTGGCTTGTGCGGCATTAAAGGCG TGGAACGGTAAGCAAGATAACGACAGTAAAGGTGGTGCTTTACTACGTGAATTTGCCCATCTATTCAGCC AAAAGACCATGTTGACCCAAGGATTCGATCCCGCTAATGCGGCAACGACACCTAAAACCTTGACTACGGA TGGCAGTGCCTTAAAGGCCTTGGCGCACAGTGCGCTTAACCTTGAGGCCGCAGGCTTTGCATTAGATGCG CCATTGGGGGATGTGCAATTTGTGGAGAAATCGCTGCCAGATGGAACGCCAAGCGGGTCGCGTTTACCTT GGCCGGGTAGTCATAATGCTGAAGGTGGATTCAACGTGTTTTCAACCAGTTTTGTCGGGTGATGACACTTT GATCCCGCAGCACAAATACGCCGCTGTGATGGATGTGTCACAGGCAAGGCGATGAGTTCTGGCTTAACG GCGAAAGGATACCAAGTGCGTTACGGTTCGAGTTGGATGATGGCGGTGAACTTTACCGACGAGGGGCCTG TGGCGCGGGGGATTTTAACTTACTCTGAGTCGAGTAATATCTTAACGCCTGCGTTTGCTGATCAAAGTAT CTTGTACTCGAGTGCTAAAAGCTTCCGTCCTTTGTTGTTCAAAGAGGCTGACATAGCACCTGCTGTAGTG TCGACGACTGAGCTGACACTACAAAAACCA

AacS Protein Sequence 850 amino acid residues, predicted size 91.82 KDa

MKFNKLVIAMGMACGVILTGCNDSEDSTTPTEPETQLQAFAPNGLLKASIRRTTFGVPHIQADNLESLGF GSGYAQAQDNLCVLADGFIKANSQRSMYFGPHASIDFTTGQPTAEDNGNLISDFAYKALKIRAQAEEKWP KFSENSRALIQGFTSGYNQYLADVEAGTQTAEPFCGGQPWVKPIVPEDVVTYLFSIALLPGAANFLDLIF YANPGDAQEYMPRIVGPAPSQDQTAFVADMQSKLIARAARITTPETNPRDLGSNGWGLGKDKTENGKGMV LGNPHFPHTGNLRFWQSHITIPGHLDMMGGSLVGMPGPINIGFNKDLAWTHTFSTAEHFVMYNLELVSGD RMQYLFDGKPMPITKETVSVLVNAGPAGMLVAEKDIYTTAKGPMVEAPPALAPFGWDDGSAFMIQDANMG TMDPVDHWLAMNMATNKEEFQQAFKDYDGVIFNNTMYADKEGNAFYIDDSTVPGLSESAVVLLKTSPDIK AAKAKAGFTILPGNTSLFSFDGPTPYERAPKLERSDFVQNSNDSFWSTNLNEPLTYYSPILRRRLDSYH CGHGWAMPRGADGKFSWKNSKRAVLSNRSYLAELVLPDLIAQCDAQGSTPVVVSASLSKDVSLACAALKA WNGKQDNDSKGGALLREFAHLFSQKTMLTQGFDPANAATTPKTLTTDGSALKALAHSALNLEAAGFALDA PLGDVQFVEKSLPDGTPSGSRLPWPGSHNAEGGFNVFSTSLSGDDTLIPQHKYAAVMDVVTGKAMSSGLT AKGYQVRYGSSWMMAVNFTDEGPVARGILTYSESSNILTPAFADQSILYSSAKSFRPLLFKEADIAPAVV STTELTLQKP

Figure 4.10. aacS DNA sequence and translated protein sequence.

In order to assay the ability of the aacS clone from S. putrefaciens B1 to inactivate AHLs, the E. coli DH5α pGEMaacS strain was grown to stationary phase in media seeded with 1.25 µM C4-HSL, C6-HSL and C12-HSL. An AHL concentration of 1.25 µM was found via reapeated AHL inactivation assays exposing a range of exogenous AHL concentrations to the AHL lactonase AiiA to be sufficient to activate the bio-reporters but low enough for AHL inactivation to be detected (data not shown). AHLs were extracted from the resultant culture supernatant using acidified ethyl acetate and detected using appropriate E. coli bio-reporters in a 96 well plate assay (Figure 4.11). After 4 h incubation E. coli DH5α pGEMaacS failed to activate the AHL bioreporters pSB536, pSB401 and pSB1142 indicating that the AHLs in seeded media had been degraded. Extracts from culture supernatants from E. coli DH5a pGEMaacS, which had been acidified to pH 2.0 prior to extraction, also failed to activate the AHL bio-reporters. Based on previous work characterising enzymic methods of AHL inactivation this result indicated that the AHLs were inactivated through the actions of an acylase as acidification would counter act AHL inactivation via a lactonase enzyme (Yates et al., 2002) (Figure 4.11).

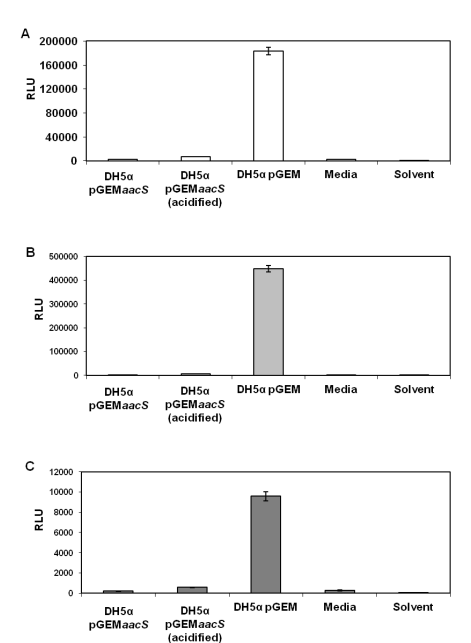


Figure 4.11. Initial AHL Inactivation Assays. Luminescence (RLU) resulting from the activation of AHL bio-reporters (**A**) E. coli pSB536, (**B**) E. coli pSB401 and (**C**) E. coli pSB1142 in the presence of cell free supernatant extracts from E.coli DH5α pGEMaacS, E.coli DH5α pGEMaacS (acidified) and E.coli DH5α pGEM cultured in media seeded with 1.25 μM C4-HSL, C6-HSL and 3-oxo-C12-HSL respectively. Extracted LB media and solvent are plotted as negative controls. Error bars represent standard deviation from the mean.

4.2.7.3 AacS Protein Expression

The putative AHL acylase from S. putrefaciens B1, AacS, was expressed in order to purify the resultant protein for further studies. Expression was carried out using the pCOLD expression system (Hayashi and Kojima, 2008). The pCOLD vector regulates expression using the E. coli cold shock protein (cspA) promoter, only allowing expression at 15°C. This vector also adds an N-terminal his-tag onto the expressed protein allowing easy purification. The aacS ORF was amplified via PCR and cloned into the pCOLD vector creating the expression construct pCOLDaacS. This expression construct was transformed into E. coli BL21 and cultured for 24 h at 15°C. The successful expression of the acylase gene from E. coli pCOLDaacS resulted in an approximately two-fold decrease in growth, as measured by OD₆₀₀, in comparison to E. coli pCOLD cultured under the same conditions.

The recombinant AacS protein from E. coli BL21 pCOLDaacS was analysed on a 12% polyacrylamide gel (Figure 4.12). The expressed AacS protein was present in the insoluble fraction of the protein extraction. The size of this expressed protein was between 50 KDa and 60 KDa, much smaller than the predicted 91.82 KDa, indicating that there may have been post-translational modification. This assumption of post-translational modification in given further credence by the findings of Lin et al. who state that AHL acylases possess conserved glycine-serine parings which are integral to post-translational modification (Lin et al. 2003).

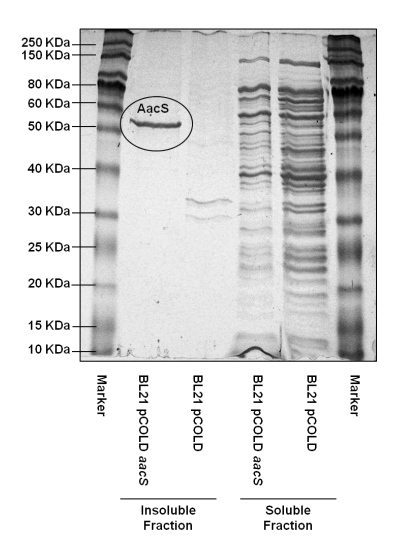


Figure 4.12. Expression of the S. putrefaciens **B1 recombinant AHL inactivating enzyme AacS from E. coli BL21 pCOLDaacS**. Polyacrylamide gel showing soluble and insoluble proteins extracted from both E. coli BL21 pCOLDaacS and E. coli BL21 pCOLD. The expressed AacS protein is marked in the E. coli BL21 pCOLDaacS insoluble fraction. The protein size marker used was the ColourPlus Pre-stained Protein Ladder, Broard Range 10-230 KDa (NEB).

4.2.7.4 AacS AHL Inactivation Specificity Assays

The specificity of AacS was tested by assaying its ability to inactivate a range of different AHLs. E. coli DH5 α pGEMaacS and E. coli DH5 α pGEM were grown separately to stationary phase in LB media supplemented with different AHLs at a

concentration of 1.25 μ M. Cell free supernatants from these cultures were extracted using DCM and assayed for AHL inactivation as described in section 2.9.3. Bioluminescence and absorbance was measured over a 12 h period every 30 min and the total amount of bioluminescence as a function of absorbance was plotted against time (Figure 4.13). Extracts from E. coli DH5 α pGEM grown in media seeded with AHL activated the appropriate bio-reporters as expected. In contrast a complete absence of bio-reporter activation was observed with extracts from E. coli DH5 α pGEMaacS grown in media seeded with C4-HSL, C10-HSL, 3-oxo-C10-HSL, 3-OH-C10-HSL, C12-HSL, a reduction in bio-reporter activation was observed with extracts from E. coli DH5 α pGEMaacS grown in media seeded with the remaining AHLs in comparison to extract from E. coli DH5 α pGEM (Figure 4.13 B). Extracts from the LB culture media and solvent had no effect on the AHL bio-reporters (data not shown).

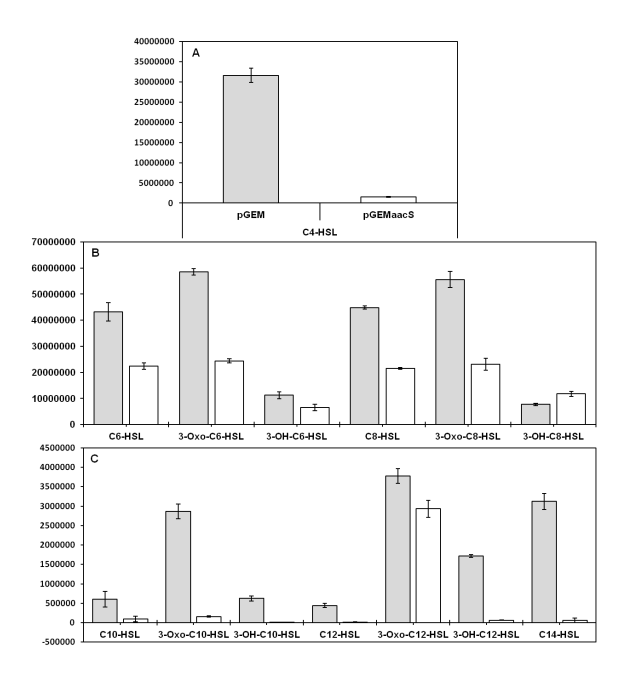


Figure 4.13. AacS AHL Inactivation Specificity Assay. AHL bio-reporter E. coli pSB536 (**A**), pSB401 (**B**), and pSB1142 (**C**) activation in the presence of cell free supernatant extracts of DH5α pGEMaacS (white bars), DH5α pGEM (grey bars) grown in LB seeded with C4-HSL, C6-HSL, 3-oxo-C6-HSL, 3-OH-C6-HSL, C8-HSL, 3-oxo-C8-HSL, 3-OH-C8-HSL, C10-HSL, 3-oxo-C10-HSL, 3-OH-C10-HSL, C12-HSL, 3-oxo-C12-HSL, 3-OH-C12-HSL and C14-HSL. Error bars represent standard deviation from the mean.

In addition to assaying the specificity of AacS to inactivate synthetic AHLs exogenously added to culture media, AacS was also assayed for its ability to inactivate the AHLs produced by the human pathogen Yersinia pseudotuberculosis. Y. pseudotuberculosis has the ability to produce 24 separate AHL's. Of these 24 only eight were found to be present at levels above 17% of the total AHL output and were therefore produced at abundancies of biological significance (Ortori et al., 2007). These eight are 3-oxo-C6-HSL, 3-oxo-C7-HSL, C6-HSL, 3-OH-C8-HSL, 3-oxo-C8-HSL, C8-HSL, 3-oxo-C10-HSL and C7-HSL (Ortori et al., 2007). The pGEMaacS vector was transformed into Y. pseudotuberculosis YPIII and the resultant strain YPIII pGEMaacS subsequently grown to stationary phase. AHLs from cell free supernatants of YPIII pGEMaacS, the parental YPIII (without plasmid) and, to act as a positive control for AHL inactivation in Yersinia, YPIII harbouring the AHL lactonase aiiA on the pGEM vector were extracted using acidified ethyl acetate. These extracts were assayed for their ability to activate the AHL bio-reporters E. coli pSB401 and pSB1142. Extract from YPIII pGEMaacS cultures activated the bio-reporter E. coli pSB401 to a comparable level as extract from YPIII wt cultures (Figure 4.14). This result indicates that AacS failed to inactivate the shorter chain AHLs produced by Y. pseudotuberculosis. Extract from YPIII pGEMaacS cultures did not activate the bioreporter pSB1142, showing comparable levels of bio-reporter activation to extract from the AHL-inactivating YPIII pGEMaiiA cultures (Figure 4.14). This indicated that AacS preferentially inactivates the longer chain AHLs produced by Y. pseudotuberculosis; a result consistent with AHL inactivation assays using synthetic AHLs.

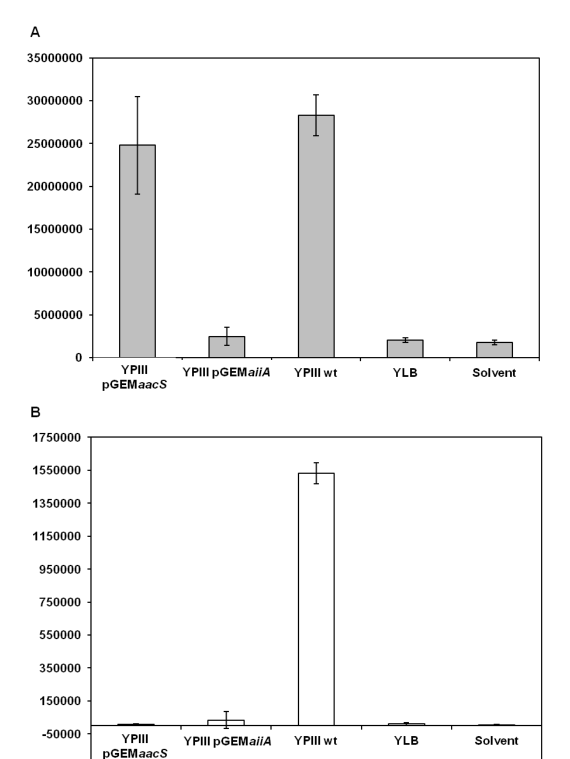


Figure 4.14. AacS AHL inactivation assay within a Y. pseudotuberculosis host. E. coli pSB401 (**A**) and pSB1142 (**B**) total luminescence as a function of growth in the presence of cell free supernatant extracts of YPIII pGEMaacS, YPIII wt and YPIII pGEMaiiA. Solvent and media controls are also plotted. Error bars represent standard deviation from the mean.

-250000

4.3. Discussion

4.3.1 AHL Production in Shewanella and Bacteroidetes Strains Abundant in the Ulva Epiphytic Bacterial Population.

This chapter reports AHL signal molecule production and inactivation in bacteria belonging to the genus Shewanella and the bacterial phylum Bacteroidetes. Both either can be isolated from the Ulva thallus surface or from other marine environmental niches colonised by Ulva spp. In an attempt to identify AHL synthases, response regulators and AHL inactivating protein sequences, this study screened both Shewanella and Bacteroidetes genomes via in silico analysis. Owing to the greater amount of previous work conducted on AHL signalling and inactivation in Shewanella spp. and the effects of both these phenotypes on the biology of both Ulva spp. and the bacteria present in rocky shore biofilm populations, this study also went on to investigate AHL inactivation within a number of different Shewanella stains.

4.3.1.1 AHL Production by the Bacteroidetes

The majority of research into AHL-mediated QS has focused on the proteobacterial phylum, specifically in species which are involved in human pathogenesis (reveiwed by Williams et al., 2007). However, in the last few years an increasing number of marine bacterial strains belonging to Cyanobacteria and Bacteroidetes phyla have been reported to produce AHL signalling molecules (Huang et al., 2009; Sharif et al., 2008). Initial studies reporting AHL signalling in biofilms containing Bacteroidetes bacteria and strains of the Bacteroidetes genus Flavobacterium presented conflicting results and were limited in their detection

methods; relying on the activation of AHL bio-reporters to prove the existence of AHL signalling mechanisms (Huang et al., 2009; Wagner-Döbler et al., 2005). This study and others has found that the sole use of bio-reporters to confirm AHL signalling is unreliable when analysing marine strains. This is due to the conditions of the marine culture media such as the increased pH affecting bio-reporter activation due to the hydrolysis of AHLs (Romero et al., 2010; Tait et al., 2005). True confirmation of AHL signal molecule production requires chemical analysis of cell free supernatant extracts using high performance liquid or gas chromatography coupled with mass spectroscopy (Ortori et al., 2007). Presently, AHL production by Bacteroidetes has only been proved unequivocally using LCMS in T. maritimum, which has been confirmed to be a producer of C4-HSL (Romero et al., 2010). Using LCMS as a detection method for AHL production, this study has extended the range of Bacteroidetes species proven to be producing AHLs to include Cellulophaga strains isolated from the Ulva thallus and other Bacteroidetes strains from varying taxonomic backgrounds (Table 3.3 and Figure 4.4). Having shown AHL production in the Sphingobacteria strain Lewinella marina and the Cytophaga strains Cytophaga marinoflava, Algoriphagus chordate, Flexibacter aggregans and Saprospira grandis this is additionally the first study to demonstrate that Bacteroidetes strains outside of the Flavobacteria class produce AHLs. This study has shown that the Bacteroidetes make up a substantial percentage of the epiphytic bacterial community of the green seaweed Ulva (Figure 3.5). Previous work has also shown the Bacteroidetes to be prevalent in biofilm communities on the rocks where Ulva spp. are found colonising (Tait et al., 2009). Ulva zoospore settlement, Ulva zoospore germination and the early growth of the Ulva germling is affected by the production of AHLs by the bacteria in the plants proximal environment, (Tait et al., 2005; Joint et al., 2002). Owing to the prevalence of Bacteroidetes in this proximal environment and the evidence of AHL production by a wide range of Bacteroidetes strains from varying phylogenetic backgrounds, it is therefore likely that the Bacteroidetes play a significant role in Ulva spp. growth and development.

Interestingly, the only AHL being found to be produced by Bacteroidetes strains assayed in this study was the short chain C4-HSL. C4-HSL production appears to be ubiquitous within the Bacteroidetes phylum as T. maritimum was also shown to be only producing C4-HSL, (Romero et al., 2010). The sole production of C4-HSL is at odds with other marine bacterial strains which tend to produce AHLs with longer acyl side chains (Tait et al., 2009). AHLs with acyl short side chain, such as C4-HSL, have been shown to be rapidly hydrolysed by the alkaline pH of seawater rendering them biologically inactive (Hmelo and Van Mooy, 2009; Yates et al., 2002). As such the production of C4-HSL by Bacteroidetes may be to facilitate communication between cells within close proximity to each other potentially limiting the possibility of 'eavesdropping' by competitive bacterial species. The production of a single type of signal molecule makes the Bacteroidetes differ from proteobacterial species such as marine Vibrio spp., which in general produce two specific types of AHL, and human pathogens such as Y. pseudotuberculosis which has been shown to produce a range of different AHLs (Atkinson et al., 2006; Milton, 2006).

4.3.1.2 Speculative Role of AHL Signalling in the Bacteroidetes and Shewanella sp.

The primary purpose of bacterial AHL production is to regulate gene expression and therefore various phenotypes in a cell density-dependent manner (Williams et al., 2007). Many proteobacterial phenotypes have been connected to AHL-mediated QS and in the cyanobacterial phyla AHLs have been shown to regulate alkaline phosphatases, affecting phosphorus acquisition in Trichodesmium consortia (Van Mooy et al., 2012; Williams et al., 2007). Currently no phenotype within Shewanella spp. or the Bacteroidetes phylum has been shown to be regulated by AHLs; however a number of potential candidate phenotypes can be proposed. Flavobacterial species such as Flavobacterium johnsoniae and Flavobacterium psychrophilum exhibit rapid gliding motility linked to a novel protein secretion mechanism (Sato et al., 2010; Jarrell and McBride, 2008). Shewanella is also a highly motile bacterial species with swimming motility being conferred by the presence of a single polar flagella (Paulick et al., 2009). Motility is a phenotype widely regulated by AHL-mediated quorum sensing in a number of proteobacterial species including Y. pseudotuberculosis, P. aeruginosa and Serratia liquefaciens (Atkinson et al., 1999; Glessner et al., 1999; Lindum et al., 1998). Swimming motility, the type of motility exhibited by Shewanella has been conclusively shown to be regulated by QS in Y. pseudotuberculosis, where the AHL-mediated QS system YpsR/I controls the expression of FlhDC, the flagella master regulator (Atkinson et al., 2008). Similar regulatory mechanisms with regard to swimming motility could account for AHL production in Shewanella spp. In addition motility similar to the type reported to be exhibited by Flavobacteria has also been reported to be regulated by AHL-mediated quorum sensing systems in Serratia marcescens (Morohoshi et al., 2007). Virulence factor production may also be a candidate as to why these two groups of bacteria are producing AHLs. A number of Bacteroidetes strains such as F. psychrophilum are implicated in fish disease and have been shown to produce virulence factors such as secreted proteases and cytolytic toxins (Duchaud et al., 2007; Nematollahi et al., 2003). The regulation of similar virulence factors by AHL-mediated quorum sensing systems in a range of bacterial pathogens has been widely reported (Williams, 2007). If Bacteroidetes virulence is regulated by AHLmediated QS it could be of importance not just in marine ecosystems but also in the field of human disease due to the prevalence of Bacteroidetes bacteria in intestinal and dental infections, (Tanaka et al., 2008; Lassmann et al., 2007). As little is known about Shewanella pathogenesis this study is unable to connect AHL production to virulence factor expression within this group of bacteria. Another Bacteroidetes phenotype that could potentially be regulated by AHL-mediated quorum sensing is pigment production. In general Bacteroidetes species are reported to be highly pigmented, (Kirchman, 2002), a characteristic also observed in the strains assayed for AHL production by this study. Pigment production has been reported to be connected to AHL-mediated quorum sensing in a number of bacteria including violacein production in C. violaceum and pyocyanin production in P. aeruginosa, of which both pigments additionally act as antimicrobials (Dietrich et al., 2006; McClean et al., 1997). Secondary metabolite and exoenzyme production and export are regulated by AHL-mediated QS in a number of different bacteria (reviewed by Williams et al., 2007). Shewanella species such as S. putrefaciens produce and export siderophores, which are utilised in iron accession (Gram, 1994). AHL-mediated QS has been implicated in the regulation of siderophore production in Burkholderia pseudomallei and P. aeruginosa (Song et al., 2005; Stintzi et al., 1998). Bacteroidetes bacteria have also been shown to produce exoenzymes that facilitate the degradation of complex organic matter such as chitin and cellulose (Teeling et al., 2012; Cottrell and Kirchman, 2000). As the production of exoproteases and other exoenzymes has been reported to be regulated by AHL-mediated QS in marine bacteria such as Aeromonas hydrophyla and Aeromonas salmonicida, potential QS regulation of this phenotype in the Bacteroidetes is not unfounded (Swift et al.,

1999). AHL production in Shewanella spp. may also be used in the regulation of organism's novel metabolic processes such as the use of nitrate and nitrites. Nitrogen metabolism has been shown to be regulated by QS in P. aeruginosa (Juhas et al., 2004). It is however important to stress that currently these candidate phenotypes are only speculations as to why both Shewanella and Bacteroidetes strains are producing AHLs; further phenotypic analysis of these bacteria would be required to link any of these behaviours to AHL production.

4.3.2 AHL Synthase, AHL Response Regulator and AHL Inactivating Enzymes in Shewanella and the Bacteroidetes

Although this study and previous work have established AHL production in both Shewanella spp. and the Bacteroidetes phylum, as yet no AHL synthase or response regulator has been identified in these two groups of bacteria (Romero et al., 2010; Huang et al., 2009; Tait et al., 2009). One putative AHL synthase sequence has been identified in the flanking DNA upstream of the lux operon in S. hanedai, however AHL production by the product of this gene has not been investigated and the sequence of this gene does not appear in other Shewanella genomes (Kasai et al., 2007). This study screened 19 separate Shewanella and 180 separate Bacteroidetes genomic sequences in an attempt to identify putative AHL synthases and response regulators but failed to identify any sequences with homology to LuxI in either Shewanella or the Bacteriodetes. Screening for AHL response regulator homologues produced a number of results, however in both the Shewanella and the Bacteroidetes genomes the resultant hits did not possess AHL binding domains so therefore cannot be classed as AHL transducing proteins. Only one identified LuxR hit, found in genome of S. violacea,

was shown to possess both an AHL binding domain and a LuxR type DNA binding domain. However, this S. violacea genome lacked any ORF with homology to luxI. This putative S. violacea LuxR sequence therefore maybe an orphan LuxR homologue, which has the ability to detect and transduce exogenously produced AHL signals, similar to those reported in P. aeruginosa, E. caratova and E. coli (Barnard and Salmond, 2007; Chugani et al., 2001). Further investigation, in vivo, using S. violacea would need to be conducted to confirm this theory. Similar false positive results were seen when screening Shewanella and Bacteroidetes genomes for two component AHL response regulators similar to the LuxN system in V. harveyi (Bassler et al., 1993). Bacteroidetes genomes that possessed hits to the LuxN consensus lacked the constitutive LuxM type AHL synthases. A speculative theory that these hits may be 'orphan' LuxN homologues is not plausible as these genomes also did not contain LuxO and LuxU homologues, the components essential for the function of LuxN as an AHL receptor/response regulator (Freeman and Bassler, 1999a; Freeman and Bassler, 1999b). It is therefore credible to assume that both Shewanella and the Bacteroidetes possess novel and yet to be identified AHL synthase/response regulator systems.

In general, AHLs are biologically inactivated by two classes of enzyme; AHL lactonases which catalyse the hydrolysis of the homoserine lactone ring and AHL acylases which facilitate the cleavage of the fatty acid side chain from the homoserine lactone ring (Dong et al., 2000; Leadbetter and Greenberg, 2000). AHL inactivation has been identified in a number of marine bacteria including two species of Bacteroidetes; T. maritimum and T. discolor (Romero et al., 2011; Romero et al., 2010). This study screened Bacteroidetes genomic sequences for putative AHL lactonase and AHL acylase sequences. Genomic screening revealed a number of hits to both AHL lactonase

and AHL acylase consensus sequences. Further bioinformatic analysis of these sequences revealed conserved amino acid residues to be present that are integral for AHL lactonase and acylase activity (Lin et al., 2003; Dong et al., 2002). The presence of these integral conserved residues within the identified hits supports the idea that these sequences are putative novel AHL inactivating enzymes. In order to confirm these hits as true AHL lactonase or acylase enzymes, AHL inactivation assays would need to be carried out. The putative AHL inactivating enzyme sequences identified in Bacteroidetes genomes by this study were however, not found in the genomes of strains known to be either producing or degrading AHLs. As this study failed to identify any AHL lactonase or AHL acylase sequences within the genomes of Bacteroidetes strains currently reported to be either actively producing or inactivating AHLs, it is credible to assume that Bacteroidetes bacteria possess novel AHL inactivating enzymes yet to be alluded.

4.3.3 AHL Inactivation by Shewanella Aculeacin A Acylase.

AHL inactivation has additionally been reported in Shewanella spp., where long chain AHLs are rapidly inactivated during the stationary growth phase (Tait et al., 2009). The study by Tait et al. also identified that AHL inactivation by Shewanella spp. takes place through the actions of an acylase enzyme. This method of AHL inactivation was first identified using AHL bio-reporters and then confirmed in HPLC experiments which showed a reduction in 3-oxo-C12-HSL coupled to a rise in homoserine lactone levels after incubation with Shewanella sp. BL21 (Tait et al., 2009). There has also been one study to identify an AHL acylase in a Shewanella strain isolated from fish intestinal tracts. Morohoshi et al. (2008) reported that the aculeacin A acylase gene

(aac) cloned from Shewanella sp. MIB015 possessed the ability to inactivate AHLs and shared 26.8% identity with the AHL acylase AiiD from Ralstonia sp. XJ12B (Morohoshi et al., 2008; Lin et al., 2003). In silico analysis carried out in this study revealed a number of putative AHL acylase sequences in genomes from various different Shewanella species including S. ondeiensis, S. putrefaciens, S. baltica and S. amonzoniensis. Many of these sequences also possessed high levels of identity to Aac from Shewanella sp. MIB015. Owing to these high levels of sequence identity, this study screened a number environmental isolates for homologues of the aac gene using PCR primers derived from the sequences identified during in silico screening. PCR screening showed that the majority of the strains assayed did possess aac homologues and, as such, it can be inferred that the aac gene is highly conserved within the Shewanella genus. The aac gene from a Shewanella strain, B1, (identified as most closely related to S. putrefaciens), aacS, was selected for further study. AacS was shown to possess 33% identity with the AHL acylase PvdQ found in multiple Pseudomonas species (Huang et al., 2003). Purification of AacS with the intention to carry out AHL inactivation assays was attempted. However, expression of AacS resulted in the protein being present within the insoluble fraction of the protein extraction and, due to time constraints, it was not possible to carry out the purification of this protein. Interestingly, the purified AacS protein expressed in this study was smaller than the size predicted by analysis of the AacS sequence. This may be due to post-translational modification of the polypeptide. Previous studies have indicated that AHL acylase proteins are modified and that a serine-glycine paring within the acylase is integral to this modification (Lin et al., 2003). To confirm that the AacS protein expressed using the pCOLD system by this study is an active AHL acylase, it may be possible to run the protein through a non-denaturing gel and overlay the gel with a bioreporter that has the ability to detect AHL inactivation.

The AHL inactivating activity of AacS was however assayed using the recombinant clones, harbouring aacS within the pGEM vector. AacS was shown to inactivate a wide variety of AHLs with the exception of 3-OH-C8-HSL. Inactivation was found to be most pronounced in AHLs with long acyl side chains. In addition to the inactivation of synthetic long chain AHLs, AacS also inactivated the long chain AHLs produced by Y. pseudotuberculosis, however had little to no effect on the other AHLs this bacterium produces. Inactivation of long chain AHLs is consistent with previous results that showed long chain AHL inactivation in Shewanella biofilms and with the activity of AHL acylases PvdQ from P. aeruginosa and AiiD from Ralstonia sp. XJ12B (Tait et al., 2009; Huang et al., 2003; Lin et al., 2003). AHL inactivation by Aac homologues does not however appear to be consistent in all Shewanella strains as previous work demonstrated that the Aac from S. oneidensis MR1 did not show AHL inactivating activity, as such it is likely that Shewanella spp. may possess another AHL acylase (Tait et al., 2009).

In general, bacteria inactivate AHLs in order to gain a competitive advantage over other bacteria growing within a particular environmental niche (Dong et al., 2004). For example Bacillus thuringiensis produces an AHL lactonase that has the ability to inhibit AHL signalling in E. carotovora. The inactivation of this signal results in the 'disarming' of E. carotovora as it inhibits antibiotic and virulence factor expression preventing the spread of E. cartovora and therefore conferring B. thuringiensis with an advantage (Dong et al., 2004). AHL acylase production by Shewanella spp. has been

shown to inactivate the AHL signals produced by bacteria found growing in the same environmental niche (Tait et al., 2009). It is therefore possible that Shewanella produces AHL inactivating enzymes in order to gain a competitive advantage over these bacteria. This still however poses the question of why does Shewanella inactivate its own AHL signal molecules? As Shewanella appears to inactivate its cognate signal during stationary phase it may be that inactivation of signal is to prevent autoinduction of AHLs which are no longer required during this growth stage, therefore preventing the metabolic burden associated with AHL production. An example of a bacterium using an AHL inactivating enzyme to turn over its cognate AHL signal molecules for this purpose is not unprecedented. Agrobacterium tumefaciens inactivates cognate 3oxo-C8-HSL production using an AHL lactonase, AttM (Carlier et al., 2003). As with Shewanella AHL inactivation, AHL inactivation by AttM also takes place during stationary phase and in this case is triggered by a stress alarmone (p)ppGpp (Zhang et al., 2004). This inactivation of cognate AHLs terminates energy consuming processes regulated by AHL-mediated QS and allows the A. tumefaciens to timely adapt to starvation stress (Zhang et al., 2004). Another explanation of Shewanella cognate AHL inactivation may also be that Shewanella inactivates its long chain AHLs at stationary phase in order to switch to the regulation of gene expression by short chain AHLs, as it is these short chain AHLs that are found in mature Shewanella biofilms (Tait et al., 2009). As with potential roles of AHL production in Shewanella presented in section 4.3.1.2, the purposes for AHL inactivation by Shewanella are only speculations and require proving via experimental investigation.

4.3.4 Summary Conclusions and Future Directions

In summary, this study has further extended the range of Bacteroidetes bacteria known to be producing AHLs. This study has also further investigated AHL inactivation by Shewanella bacteria and shown that the AHL acylase Aac is highly conserved in Shewanella spp. and is responsible for long chain AHL inactivation. In spite of this, our knowledge of AHL signalling and QS in Shewanella and the Bacteroidetes phylum is still in its infancy as no AHL synthase and response regulator systems have yet been identified within these two groups of bacteria. Further research into AHL production and QS in both Shewanella and the Bacteroidetes would be of benefit to our understanding of marine ecosystems due to the abundance of these groups of bacteria within the marine environment. In addition to furthering our understanding of the marine environment researching AHL production and QS in the Bacteroidetes may provide benefit to medical science, as these bacteria are responsible for a large number of common infections.

Research in this area should progress in the following directions:

• Carry out mutation of AacS in S. putrafaciens B1. Construction of an AacS knock out mutant has been initiated by this study, however is not discussed in this thesis. An AacS mutation construct has been constructed via PCR amplification of an 80 bp region in the centre of aacS which was ligated into the suicide vector pKNOCK, (Alexeyev, 1999), in order to perform homologous recombination, an approach previously shown to successfully construct knock out mutants in Shewanella spp. (Bodor et al., 2011). A potential S. putrafaciens B1 ΔaacS strain has been constructed but not tested.

- Assay Shewanella sp. ΔaacS mutant for AHL production at all stages of growth using the method outlined in section 2.9.2 and 2.9.3 to confirm AacS as the AHL inactivating enzyme in Shewanella and to identify if AacS is the only AHL inactivating enzyme present.
- Use Shewanella sp. ΔaacS mutant to identify Shewanella spp. AHL synthase and response regulators using a transposon library or genomic library approach as previously used in enteric bacteria (Hao et al., 2010; Swift et al., 1993).
- Screen an AHL-producing Bacteroidetes strain for AHL synthase and response regulators using a transposon library or genomic library approach as previously used in enteric bacteria (Hao et al., 2010; Swift et al., 1993).

Chapter 5

Quorum-quenching Activity in

Microalgae

5.1. Introduction

The inhibition of bacterial QS has been proven to affect an array of phenotypes such as virulence factor expression (Dong et al., 2000). The targeting and subsequent inhibition of key components in QS systems is often referred to as quorum-quenching (reveiwed by Zhang, 2003). Quorum-quenching has been shown to be achieved either by the production of AHL inhibiting enzymes such as AHL lactonases and AHL acylases discussed in Section 1.3.2 or chemically by AHL inhibiting compounds (reveiws by Dong and Zhang, 2005, Zhang, 2003). Such compounds have been identified to be produced by a number of different organisms including other bacteria, terrestrial plants and marine eukaryotes (reveiwed by Zhang, 2003). The production of quorum-quenching compounds by marine microalgae and its potential effects on both human and marine pathogens is the focus of the data presented in this chapter.

5.1.1. Algal Compounds have the Ability to Inhibit Bacterial Quorum Sensing

Bacterial/algal associations have been studied for a long period of time and as such interactions between these two kingdoms have been well documented. A number of these interactions are discussed previously (Sections 1.2.2, 3.1.1, 3.1.3 and 3.3.3). Recent studies however have focused on bacterial/algal interactions involving bacterial signalling. Algae have been shown to interfere with bacterial QS in order to inhibit colonisation, which may lead to infection (Teplitski et al., 2004, Hentzer et al., 2002). Algae accomplish this interference using two methodologies; either by producing molecules that disrupt bacterial QS by quorum-quenching activity or by producing molecules that mimic bacterial QS signals (Teplitski et al., 2004, Manefield et al., 2002). Algal quorum-quenching behaviour was first observed in a species of

macroalgae, the red seaweed Delisea pulchra which was shown to produce halogenated furanones which are chemically similar to AHLs (Manefield et al., 1999). The furanone compounds were proven to affect QS regulation of the motility master operon flhDC within Serratia liquefaciens, reducing swarming motility (Rasmussen et al., 2000). Halogenated furanone disruption of QS was further confirmed by observations that furanones produced by D. pulchra reduced V. fischeri bioluminescence by 50 to 100 fold, a clear example of quorum-quenching activity (Rasmussen et al., 2000). Halogenated furanone compounds are produced by D. pulchra within vesicles in the algas thallus and are then exported out onto the surface of the thallus and are in greatest concentration at the apical tips of the plant (Dworjanyn et al., 1999). Manefield et al. (2002) proposed a biochemical mechanism by which furanone compounds produced by D. pulchra disrupt QS regulated gene expression. Manefield suggests that furanones interact with the signal response regulator LuxR forming a LuxR-Furanone complex. This complex causes a conformational change within LuxR leading to its increased susceptibility to proteolytic degradation within the bacterial cytosol, thus abolishing or severely disrupting QS gene regulation (Manefield et al., 2002). Halogenated furanones have also been shown to effect virulence factor expression in P. aeruginosa and reduce the expression of genes thought to be regulated by AI-2 in both E. coli and Salmonella enterica (Janssens et al., 2008, Ren et al., 2004, Hentzer et al., 2002). It is not only D. pulchra which has the ability to disrupt QS, the bryozoan Flustra foliacea has also been shown to produce brominated alkaloid compounds. The brominated alkaloid compounds have been proven to reduce the growth of bacteria isolated from F. foliacea. Using a P. aeruginosa model the brominated alkaloids were shown to block AHLdependent QS and have a phenotypic effect on the bacteria by reducing extracellular protease production (Peters et al., 2003). Oxidised halogen compounds produced by

many marine macroalgae have been shown to affect bacterial biofilms (Wever et al., 1991). Borchardt et al. (2001) showed that such compounds also react with 3-oxo substituted AHLs forming halogenated homoserine lactone and fatty acids (Borchardt et al., 2001). In the same study oxidised bromide compounds produced by the brown alga Laminaria digitata were shown to react and therefore biologically inactivate AHLs produced by P. aeruginosa biofilms (Borchardt et al., 2001). QS inhibitory compounds effecting C8-HSL and the A. tumefaciens TraR AHL response regulator have also been shown to be produced by the red algae Ahnfeltiopsis flabelliformis (Liu et al., 2008). A broad ranging study was undertaken by Skindersoe et al. (2008) investigating QS inhibition using extracts from 284 marine organisms in conjunction with the Quorum Sensing Inhibitor Selector (QSIS) systems described in section 4.2.7 (Skindersoe et al., 2008, Rasmussen et al., 2005). This study showed that 23% of the 284 extracts tested showed some QS inhibitory activity including 5 out of 35 algal extracts (Skindersoe et al., 2008). The study also went on to characterise QS inhibition by three C₂₅ sesterterperne metabolites produced by the marine sponge Luffariella variabilis (Skindersoe et al., 2008).

Less research has been carried out on the effect of molecules produced by microalgal species, in spite of studies showing complex bacterial communities to be present on these algae (Fukami et al., 1997). One example of such behaviour has been demonstrated in the freshwater microalgae Chlamydomonas reinhardtii, which has been shown to produce a number of molecules that stimulated the AHL response regulators LasR (P. aeruginosa) and CepR (Pseudomonas putida). These compounds did not show any effect on other AHL response regulators including LuxR (V. fischeri), AhyR (A. tumefaciens) and CivR (C. violaceum), indicating that the mimic activity was

receptor specific (Teplitski et al., 2004). Owing to their abundance in both the marine and freshwater environments and the increasing economical importance of microalgae, further research into their interactions with bacteria is warranted.

5.1.2. Microalgae, Ecology and Biotechnology

5.1.2.1 Microalgal Ecology

Microalgae are eukaryotic, unicellular algal species (reveiwed by Satyanarayana et al., 2011). In the marine environment microalgae are found existing either individually, in chains, or in groups and are therefore classed as part of the marine phytoplankton (reveiws Satyanarayana et al., 2011; Munn, 2004). Microalgal metabolism is driven by photosynthesis; as such these organisms are limited to the photic zone of the marine water column (Munn, 2004). The photosynthetic activity of marine microalgae, (including cyanobacterial species), accounts for the majority of carbon fixation in the marine environment and approximately 50% of global carbon fixation (Munn, 2004). When living, microalgae can be prey for zooplankton and other higher marine organisms and in death microalgal organisms provide carbon sources for heterotrophic marine microbes in the form of dissolved organic matter (Munn, 2004). These key roles in carbon fixation and in the biological marine carbon cycling makes microalgae a significant group of organisms in the study of marine ecosystems and ocean processes. In addition to being found in the marine environment microalgae can additionally be found in fresh water environments (Harris, 2001).

5.1.2.2 Usage of Microalgae in Aquaculture and Industry

Microalgae or the metabolic products of microalgae are already used in a number of biotechnological applications including use in the cosmetics industry, food industry, aquaculture and use as biofuels. In the cosmetics industry frozen microalgal bio-matter is utilised in the production of anti wrinkle creams and the high proportions of carbohydrates, proteins, omega-3 and omega-6 oils produced by microalgae are utilised in the food industry (reveiwed by Guil-Guerrero et al., 2004). Microalgae can be used in the production of biofuels, where lipids produced by microalgae are utilised as replacements for fossil fuels in the production of diesel or utilised in the production of bio-ethanol; a substitute for petroleum in the production of motor fuel (John et al., 2011, Li et al., 2008). The main economic purpose of growing microalgal species is however for usage in aquaculture (reveiwed by Borowitzka, 1997). Aquaculture can be defined as the farming of aquatic organisms such as fish, molluscs, crustaceans and aquatic plants. Statistics provided by the 2012 United Nations Food and Agriculture Organisation (UNFAO) report into world fisheries and aquaculture show inland and marine aquaculture production to have risen from 47.3 million tonnes in 2006 to 63.6 million tonnes in 2011. Total aquacultural algal production has also seen a distinct rise from 3.8 million tonnes in 1990 to 19 million tonnes in 2010 providing an economic value of \$5.7 billion in 2010 to the global economy (UNFAO, 2012). The majority of this production is related to the farming of macroalgal species, which are farmed for hydrocolloids and in both animal and human food production (Abbott, 1996, Bixler, 1996). However the production of marine microalgae in aquaculture has also seen a rise in the last 20 years with a total of 3.1 million tonnes produced in 2010, (UNFAO, 2012). Aquacultural production of microalgae has multiple uses including; food sources or feed additives in the commercial rearing of aquatic animals for direct human

consumption; food sources for commercial fish species at the larval stage; food sources for rearing rotifers which are intern used has a food in aquacultural fisheries and for the quality enhancement of aquacultural produce by providing pigments such as astaxanthin (Chien and Shiau, 2005; Borowitzka, 1997). Microalgal cultures can be reared from two sources either from natural populations or from unialgal cultures (Austin and Day, 1990; New, 1990). A great deal of research has been focused on the usage of optimal strains of microalgae in order to gain the highest levels of aquacultural production at reduced economic cost to the producer. This has lead to the discoveries of microalgal strains with specific biological properties that are of benefit to aquaculture such as high lipid containing strains of Tetraselmis spp. which have been shown to enhance the growth of oyster larvae (Wikfors et al., 1996).

5.1.3. Experimental Aims

This study investigated three microalgal species indigenous to the marine environment: Nannochloropsis oculata, Tetraselmis suecica and Isochrysis galbana. These species were selected for study due their importance in aquaculture in the rearing of rotifers, crustaceans and molluscs respectively (Borowitzka, 1997). By further understanding the biology of these commercially significant species, particularly focusing on how they may interact with bacterial populations, it may be possible to exploit such interactions for commercial benefit. Based on research discussed previously, this study investigated the possibility that these three microalgal species interact with bacteria by modulating bacterial QS (Teplitski et al., 2004, Manefield et al., 2002). As such, extracts from axenic cultures of all three species were assayed for

their ability	to	produce	QS	mimic	molecules	or	actively	quench	bacterial	AHL-
mediated signalling.										

5.2. Results

5.2.1. Activation of AHL Bio-reporters by Microalgal Extracts

Axenic N. oculata, T. suecica and I. galbana cultures were subjected to solvent extraction as described in Section 2.10.1. The resultant extracts were concentrated via rotary evaporation and assayed for their ability to produce bioluminescence in the luxbased AHL bio-reporters E. coli DH5α pSB536, JM109 pSB401 and JM109 pSB1142. If the extracts contained compounds that interacted with and activated the various AHL response regulator genes contained in these reporters, bioluminescence would occur due to the transcription of a lux cassette also contained in these reporters (Winson et al., 1998, Swift et al., 1997). The bio-reporters produced approximately the same levels of bioluminescence after co-incubation with extracts from all three microalgal species as co-incubation with extracted F/2 media and solvent negative controls (Figure 5.1). A minor increase in the bioluminessence produced in pSB536 was observed after exposure to the microalgal extracts in comparison to the solvent and media controls (Figure 5.1). This rise is however thought to be due to pSB536 possessing a 'leaky' promoter and not due actual activation of the bio-reporter. The AHL bio-reporters did however produce bioluminescence when exposed to appropriate synthetic AHLs: C4-HSL, C6-HSL and 3-oxo-C12-HSL, respectivly (Figure 5.1).

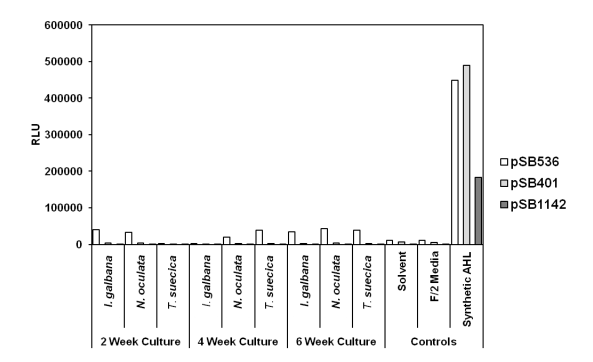


Figure 5.1. Activation of AHL bio-reporters by microalgal extracts. AHL bio-reporters exposed to extracts from N. oculata, T. suecica and I. galbana cultures grown for 2, 4 and 6 weeks produced levels of bioluminessence comparative to both solvent and F/2 media controls, indicating a lack of any compound within the extract that could activate these reporter stains. Synthetic AHL controls (C4-HSL, C6-HSL and 3-oxo-C12-HSL) activated all three bio-reporters, indicating that the reporters were acting as expected.

5.2.2. Quorum-quenching Activity by Marine Microalgal Cultures

N. oculata, T. suecica and I. galbana extracts were also assayed for the ability to disrupt bioluminescence in the E.coli lux-based AHL bio-reporters in the presence of exogenously added AHLs. This was achieved by co-incubating E.coli pSB536, pSB401 and pSB1142 with C4-HSL, C6-HSL and 3-oxo-C12-HSL respectively in the presence of extracts from the three microalgal cultures grown for 2, 4 and 6 weeks.

In the presence of synthetic AHLs, the F/2 media control extracts had no effect on bioluminescence (Figures 5.2-5.4), whereas a number of the microalgal extracts repeatedly had an effect on the levels of bioluminescence produced by the AHL bioreporters in the presence of synthetic AHLs. Significantly reduced levels of bioluminescence were observed using the bio-reporter pSB536 in the presence extracts from 2 wk old cultures of I. galbana and extracts from 4 wk old cultures of T. suecica when compared to the F/2 media control (2-Tailed T Test P < 0.01) (Figure 5.2). When exposing the bio-reporter pSB401 to extracts from 2 and 4 wk old cultures of T. suecica significantly reduced bioluminescence was observed in comparison to F/2 media extracts (2-Tailed T Test P < 0.01) (Figure 5.3). Allthough not found to be significant, reductions in pSB401 bioluminessence after exposure to extracts from 6 wk old cultures of I. galbana and extracts from 2 and 6 wk old cultures of N. oculata in comparison to F/2 media extracts was also apparent (Figure 5.3). Finally, significantly reduced levels of bioluminescence were observed in the bio-reporter pSB1142 after exposure to extracts from 2, 4 and 6 wk old cultures of I. galbana; extracts from 2 and 6 wk old cultures of N. oculata and extracts from 2, 4 wk old cultures of T. suecica, in comparison with F/2 media extracts (2-Tailed T Test P < 0.01) (Figure 5.4). Extracts from both F/2 media and the microalgal cultures did not affect the growth of any of the AHL bio-reporters used in this assay and the solvent used to reconstitute these extracts did not affect the AHL bio-reporters (Figure 5.2-5.4).

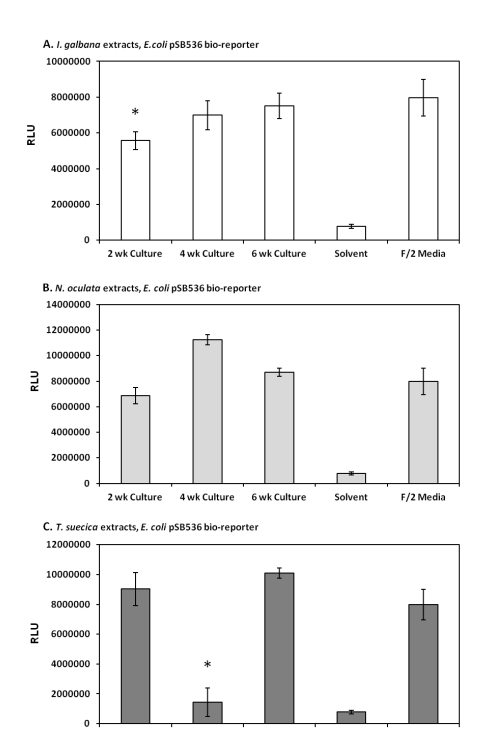


Figure 5.2. Microalgae quorum-quenching assay using AHL bio-reporter pSB536. Total luminescence produced from the bio-reporter JM109 pSB536 cultured with microalgal extracts and extracted F/2 media in the presence of C4-HSL. Significantly reduced luminescence was observed after exposure to extracts from I. galbana and T. suecica cultures grown for 2 and 4 weeks respectively. A solvent control was shown to have no effect on the bio-reporter. Error bars represent standard deviation from the mean. Asterix represent significantly significant difference from the control (2-Tailed T Test P < 0.01).

6 wk Culture

Solvent

F/2 Media

4 wk Culture

2 wk Culture

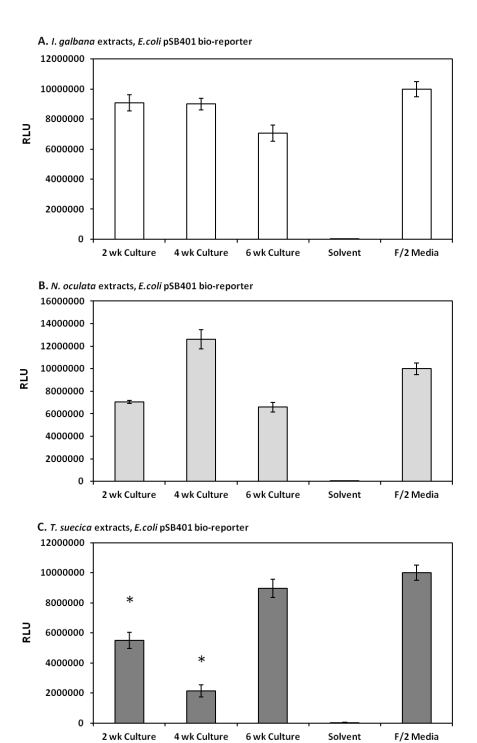


Figure 5.3. Microalgae quorum-quenching assay using AHL bio-reporter pSB401. Total luminescence produced from the bio-reporter JM109 pSB401 cultured with microalgal extracts and extracted F/2 media in the presence of C6-HSL. Significantly reduced luminescence was seen after exposure to extracts from T. suecica cultures grown for 2 and 4 weeks. A solvent control was shown to have no effect on the bio-reporter. Error bars represent standard deviation from the mean. Asterix represent significantly significant difference from the control (2-Tailed T Test P < 0.01).

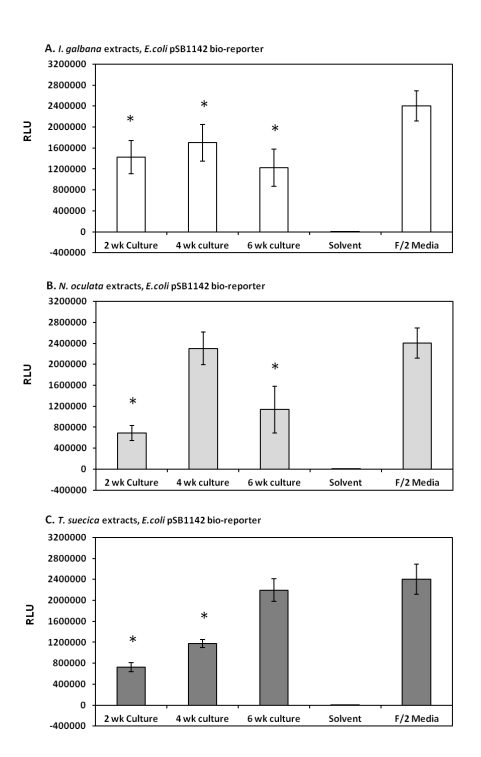


Figure 5.4. Microalgae quorum-quenching assay using AHL bio-reporter pSB1142. Total luminescence produced from the bio-reporter JM109 pSB1142 cultured with microalgal extracts and extracted F/2 media in the presence of 3-oxo-C12-HSL. Significantly reduced luminescence was seen after exposure to extracts from I. galbana cultures grown for 2, 4 and 6 weeks, extracts from N. oculata cultures grown for 2 and 6 weeks and extracts from T. suecica cultures grown for 2 and 4 weeks. A solvent control was shown to have no effect on the bio-reporter. Error bars represent standard deviation from the mean. Asterix represent significantly significant difference from the control (2-Tailed T Test P < 0.01).

5.2.3. Microalgal Quorum-quenching Activity Effects P. aeruginosa Virulence

As extracts from microalgal cultures affected the activation of AHL bioreporters in the presence of their cognate signal molecules. Extracts micro algal cultures
were also assayed for their ability to affect the expression of P. aeruginosa genes
regulated by QS. P. aeruginosa PAO1 lux-based transcriptional fusions to rhII, lasI,
rhlA and lasB (Unpublished, provided by Dr. James Lazenby) were cultured in the
presence of pooled extracts from a number of stationary phase axenic cultures of N.
oculata, T. suecica and I. galbana. Pooled extracts were used inorder to scale up the
number of experiments which could be carried out and to overcome incosistanceies
observed in previous assays using extracts from separate stationary phase microalgal
cultures.

Several of the extracts from microalgal cultures did produce lower amounts of bioluminescence in the P. aeruginosa transcriptional bio-reporters in comparison to extract from F/2 culture media (Figure 5.5). Significantly reduced levels of bioluminescence were seen in the rhlI, lasI and rhlA bio-reporters upon exposure to extracts from the three microalgal cultures (2-Tailed T Test P < 0.01) (Figures 5.5). Significantly reduced of bioluminescence were also seen in the lasB bio-reporter when exposed to extracts from I. galbana and N. oculata cultures (2-Tailed T Test P < 0.01) (Figure 5.5).

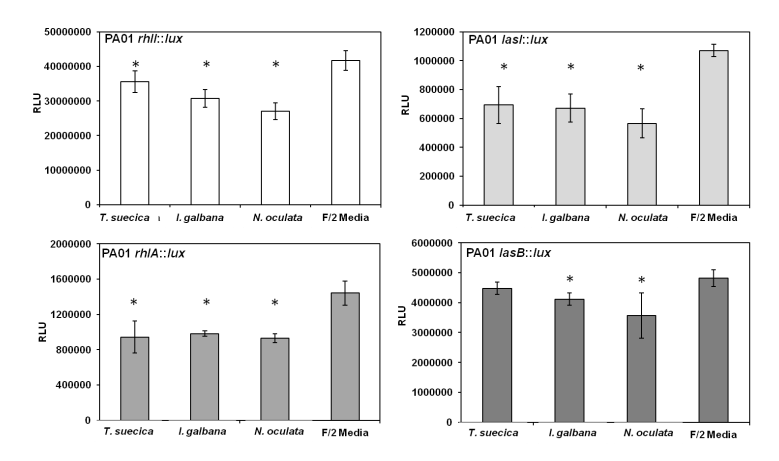


Figure 5.5. Microalgal quorum-quenching assay using P. aeruginosa transcriptional bio-reporters. Total bioluminescence obtained from bio-reporters P. aeruginosa PAO1 rhlI::lux, PAO1 lasI::lux, PAO1 rhlA::lux and PAO1 lasB::lux co-incubated with pooled extracts from stationary phase microalgal cultures and F/2 media over a 12 h period. Asterix mark extracts, which caused significant reductions in reporter bioluminescence in comparison to the F/2 media control (2-Tailed T Test P < 0.01). Error bars represent standard deviation from the mean.

The microalgal extracts were checked to ensure that they did not have any effect on the activity of the lux genes. This was achieved by repeating the assay using a P. aeruginosa PAO1 strain harbouring the mini CTX::lux construct constituently expressing the lux gene under the Km constitutive promoter (Becher and Schweizer, 2000). After carrying out 2-tailed Tests no significant reduction in bioluminescence was observed when the strain was cultured with extracts from the three microalgal cultures in comparison to culturing the strain with an extract from F/2 media (2-Tailed T Test P > 0.01) (Figure 5.6).

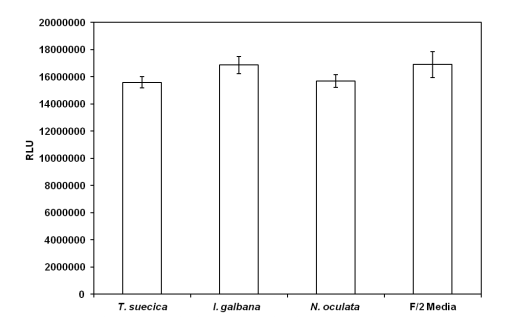


Figure 5.6. PAO1 mini CTX::lux bioluminescence upon exposure to microalgal extracts. Extracts from the three micro algal species were assayed with PAO1 harbouring the mini CTX::lux construct expressing under the constitutive P_{km} promoter to ascertain if they affected the bioluminescence process itself. Comparing bioluminescence after exposure to each microalgal extract to bioluminescence after exposure to F/2 media showed no significant differences (2-Tailed T Test P > 0.01). Error bars represent standard deviation from the mean.

5.2.4. Fractionation of Extracts Microalgal Cultures

In an attempt to identify the compounds produced by the microalgae which were showing activity against the bio-reporter and transcriptional fusions, extracts from stationary phase cultures of the three microalgal species were fractionated using solid phase extraction columns as described in section 2.10.5.

5.2.4.1 P. aeruginosa Virulence Assay

The re-constituted fractions were assayed for their ability to affect the P. aeruginosa lux-based transcriptional fusions described in section 5.2.4. Small but significant reductions in bioluminessence were observed in the PAO1 lasI::lux and PAO1 lasB::lux reporters when exposed to the 60% Methanol I. galbana fraction (2-Tailed T Test P < 0.05) (Figure 5.7). The 70% Methanol N. oculata fraction also showed small but significantly reduced bioluminessence in the PAO1 rhlI::lux in comparrsion to the solvent control but not the F/2 media control (2-Tailed T Test P < 0.05). However, the 70% Methanol N. oculata fraction did significantly reduce bioluminessence in the PAO1 rhlA::lux reporter in comparrsion to both the solvent and F/2 media controls (2-Tailed T Test P < 0.05) (Figure 5.8). Other fractions had no observable effect on the P. aeruginosa bio-reporters (Data not shown).

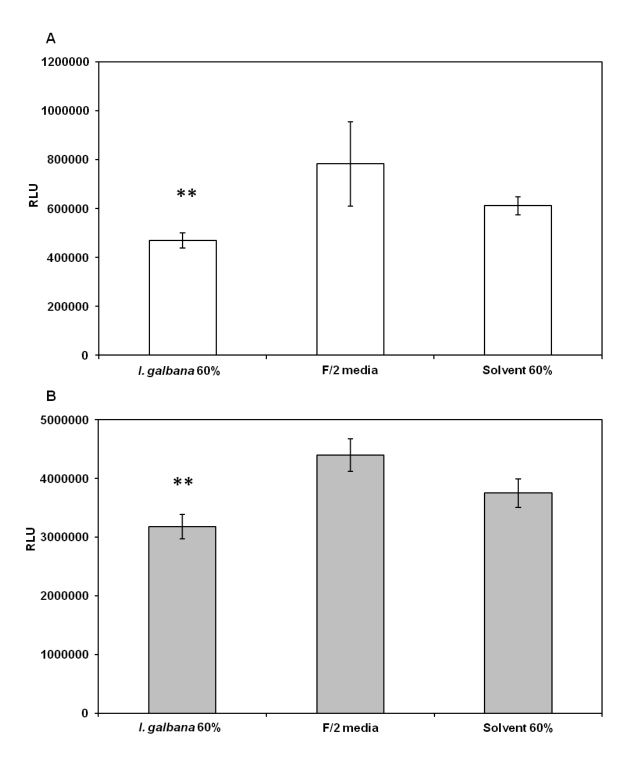
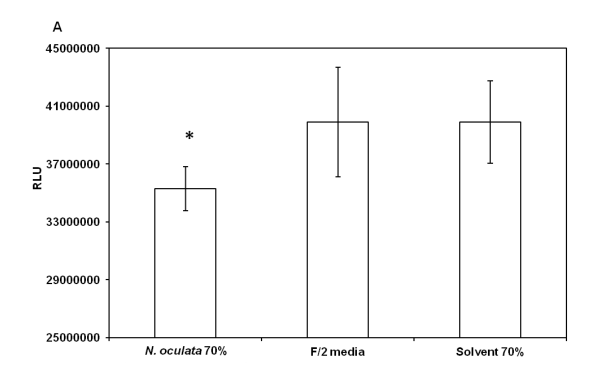
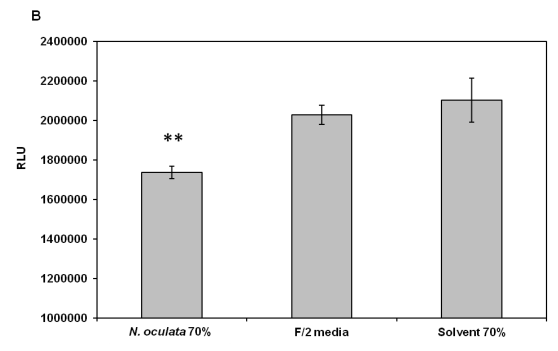


Figure 5.7. P. aeruginosa transcriptional bio-reporter assay using fractionated I. galbana culture. Total bio-reporter luminescence produced by (A) PAO1 lasI::lux and (B) PAO1 lasB::lux after 12 h exposure to 60% I. galbana fraction. ** Marks significant reductions in bioluminescence is seen in both reporters exposed to the 60% methanol I. galbana fraction in comparison to the F/2 media and solvent controls (2-Tailed T Test P < 0.05). Error bars represent standard deviation from the mean.





5.8. P. aeruginosa transcriptional bio-reporter assay using fractionated N. oculata culture.

Total bio-reporter luminescence produced by (A) PAO1 rhlI::lux and (B) PAO1 rhlA::lux after 12 h exposure to 70% N. oculata fraction. * Marks significant reduction in bioluminescence was seen the rhlI reporter in comparison to the solvent control (2-Tailed T Test P < 0.05). ** Marks significant reduction in bioluminescence seen in the rhlA reporter in comparison to both F/2 media and solvent controls (2-Tailed T Test P < 0.05). Error bars represent standard deviation from the mean.

5.2.4.2 Marine Bacteria Protease Assays

The fractionated extracts from the three microalgal cultures were also used to investigate any effects on protease production in 4 species of marine bacteria, V. anguillarium, Vibrio tubiashii, Vibrio corrallilyticus and Aeromonas hydrophila. These species are commonly found to cause infection in fish species reared for aquacultural purposes via the AHL regulated production of protease enzymes that act as virulence factors; (Tait et al., 2010, Croxatto et al., 2002, Swift et al., 1999). As such, inhibition of AHL signalling may affect protease production. The four bacterial strains were cocultured with the fractionated extracts from T. suecica, I. galbana and N. oculata and resulting protease production in each bacterium was compared between the fractions. Protease activity was determined using the methodology described in section 2.10.5. Initial assays comparing the different fractions from each microalgal culture indicated reductions in protease activity when cultures of A. hydrophila were exposed to the 60% fraction from I. glabana and when cultures of V. tubiashii were exposed to the 60% fraction from I. glabana and the 70% fractions from both T. suecica and N. oculata extracts (Appendix 9). As such, these fractions were used to repeat the assay in all four bacterial strains comparing protease activity to cultures exposed to a solvent control.

The 60% fraction of I. galbana extract and the 70% fraction of N. oculata extract caused significant reductions in A. hydrophila protease activity in comparison to the solvent control (2-Tailed T Test P < 0.05) (Figure 5.9). However, only the 70% fraction of N. oculata extract significantly affected protease activity in V. anguillarium and in V. tubiashii (2-Tailed T Test P < 0.05) (Figure 5.9). No significant reductions in protease activity were observed in V. corrallilyticus upon exposure to any of the solvent fractions assayed in comparison to the solvent control (2-Tailed T Test P > 0.05).

Culture OD_{600} measurments showed that in comparison to the solvent control the fractions from all three microalgal culture extracts did not affect bacterial growth (data not shown).

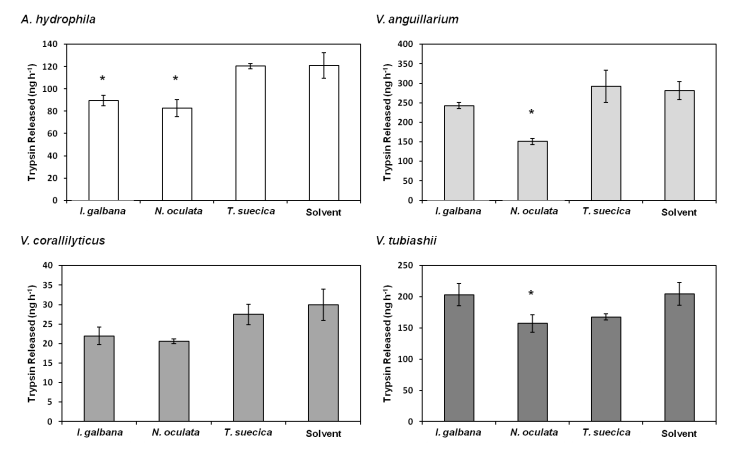


Figure 5.9. Protease assay of fractionated extracts from microalgal cultures. Protease activity in marine bacteria as a result of co-incubation with the 60% fraction from I. glabana extract and from the 70% fractions from N. oculata and T. suecica extracts in pathogenic marine bacteria. The 60% fraction of I. glabana extract significantly reduced protease activity in A. hydrophila. The 70% fraction N. oculata extract significantly reduced protease activity in A. hydrophila, V. anguillarium and V. tubiashii. The Error bars represent standard deviation from the mean. Asterix represent significantly significant difference from the control (2-Tailed T Test P < 0.05).

5.3. Discussion

5.3.1 Quorum-quenching Activity if Marine Microalgae

Previous studies investigating algal/bacterial interactions in the marine environment have shown that large scale surface water bioluminescence, commonly referred to as milky sea events, result from AHL-mediated QS by V. fischeri bacteria associated with algal plankton blooms (Nealson and Hastings, 2006). As the bacterial densities measured during milky sea events was below that previously reported to initiate AHL-mediated regulation of bioluminescence in V. fischeri, it has been theorised that the bioluminescence apparent during such events may be stimulated by QS mimic compounds produced by microalgae (Nealson and Hastings, 2006, Fuqua et al., 1994). This hypothesis was given credence due to the evidence that secondary metabolite production by freshwater microalgal species C. reinhardtii mimicked AHL bacterial signalling molecules (Teplitski et al., 2004). As such, this study investigated potential AHL mimic production and quorum-quenching activity in three microalgal species found in the marine environment. The species were I. glabana, N. oculata and T. suecica, species which are commonly grown for usage in aquaculture, acting as feed stocks for bivalve molluscs, crustaceans and rotifers, which are in turn used to feed reared fish species (reviewed by Borowitzka, 1997). In further investigating if these algae can modulate bacterial behaviour via the stimulation or inhibition of QS, it may be possible to exploit such behaviour for aquacultural or medical benefit.

This study failed to identify any secondary metabolite being produced by these three microalgal species that had the ability to activate AHL response regulators AhyR (A. hydrophyla), LuxR (V. fischeri) and LasR (P. aeruginosa), as extracts from these microalgae did not produce bioluminescence in reporters pSB536, pSB401 and

pSB1142, respectively (Winson et al., 1998; Swift et al., 1997). This does not entirely rule out the possibility that these species produce compound(s) which mimic AHLs, as such compounds may affect AHL response regulators other than the three present in theses bio-reporters. It may also be the case that any mimic compound(s) interact with these AHL response regulators inhibiting AHL-mediated QS and there for would obviously not produce bioluminescence in these reporters.

This study went onto investigate if these species were producing compound(s) that inhibited the activation of these AHL response regulators in the presence of exogenous AHLs. When culturing the lux-based AHL bio-reporter pSB536 with extract from I. glabana and T. suecica; pSB401 with extract from T. suecica and pSB1142 with extracts from all three cultures in the presence of their cognate AHLs bioluminescence was significantly decreased in comparison to culturing with extracted sterile F/2 microalgal media. The microalgae used in this study were obtained as axenic cultures, grown in conditions that would prevent bacterial contamination and checked for bacterial contamination via microscopy throughout the study supporting the notion that inhibition of AHL bio-reporter luminescence is due to the microalgal species and not epiphytic bacteria. In addition, extracts from all three microalgal cultures were shown to have no effect on the transcription of the genes in the lux operon that encode the bioluminescence producing enzyme luciferase. As such, it is likely that compounds(s) produced by these microalgae are inhibiting AHL-mediated QS in the bio-reporters accounting for the reductions in bioluminescence observed in the presence of exogenous AHLs. Natrah et al. (2011) published results that support this study's finding that axenic strains of marine microalgae produce secondary metabolite(s) that inhibit the activation of AHL bio-reporters in the presence of their cognate signal

molecules (Natrah et al., 2011). The study published by Natrah et al. (2011) only shows inhibition of AHL bio-reporters by extracts from microalgal cultures and does not identify any compound(s) that may be causing such inhibitory activity. The study does however included the freshwater microalgae Chlorella vulgaris and Chlorella saccharophila to the list of species known to be producing potential QS inhibitors (Natrah et al., 2011). With the exception of bioluminescence in the fish pathogen V. harveyi, in which they show a reduction in luminescence due to the presence of microalgal extracts, using a V. harveyi AHL bio-reporter; the Natrah et al. (2011) study does not investigate if this quorum-quenching activity effects any genes regulated by AHL-mediated QS (Natrah et al., 2011).

This current study did investigate if extracts from microalgal species effected the expression of genes known to be, in part regulated by AHL-mediated QS using the opportunistic human pathogen P. aeruginosa. P. aeruginosa possesses two AHL-mediated QS systems, the LasR/I system which directs the production and transduction of 3-oxo-C12-HSL and the RhlR/I system which directs the production and transduction of C4-HSL (Latifi et al., 1995). Further research has also shown that these two QS systems function in a hierarchical fashion, with the LasR/I systems regulating the RhlR/I system (reviewed by Latifi et al., 1996). These two AHL-mediated QS systems, alongside other regulatory systems are responsible for the regulation of a large amount of genes in P. aeruginosa, including many responsible for the production of extracellular virulence factors, biofilm formation, siderophore production and autoregulation of their own AHL synthase genes (reviewed by Whiteley et al., 1999). Extracts from all microalgal cultures were shown to significantly affect the luminescence produced by lux-based transcriptional fusions reporting the transcription

of both lasI and rhlI, with reduced luminescence observed in the presence of extracts from all three microalgal cultures. Additionally a number of the extracts effected luxbased transcriptional fusions reporting the transcription of two P. aeruginosa genes, which are in part regulated by AHL-mediated QS, lasB and rhlA. Both these gene provide important roles in Pseudomonas pathogenicity (Reviewed by Whitelely et al., 1999). All three extracts produced significant reduction in biolumminesnce in the rhlA bio-reporter and extracts from I. galbana and N. oculata significant reduced biolumminesnce in the lasB bio-reporter. The lasB gene encodes a metalloprotease, elastase, which acts as a major virulence factor in P. aeruginosa infections (Fukushima et al., 1997, Gambello and Iglewski, 1991). The rhlA gene belongs to a gene cluster along with rhlB. These two genes encode rhamnosyltransferase, which catalyses the production of rhamnolipids; glycolipids produced by P. aeruginosa and used as a surfactant promoting motility (Soberón-Chávez et al., 2005, Lang and Wullbrandt, 1999). P. aeruginosa rhlI mutants lack the ability to produce rhamnolipids and are therefore rhlI is implicated in the regulation of the rhlAB gene cluster (Ochsner and Reiser, 1995). As the microalgal extracts were shown to reduce bioluminescence in these transcriptional bio-reporters and in the lux-based E. coli AHL bio-reporters it is possible that these extracts contain compound(s) which may inhibit AHL-mediated QS in P. aeruginosa.

In addition to studying potential quorum-quenching activity by marine microalgae on QS regulated P. aeruginosa genes, this study also initiated the identification of potential quorum-quenching compound(s) by fractionating the extracts obtained from the three microalgal cultures. The separate solvent fractions were tested for their ability to inhibit bioluminescence in lux-based transcriptional bio-reporters of

genes regulated by QS in P. aeruginosa and also to inhibit protease activity in 4 marine bacteria shown to be pathogenic to fish species reared in aquaculture, A hydrophila, V. anguillarium, V. coralliyticus and V. tubiashii. Results showing inhibition of bioluminescence in P. aeruginosa virulence factor transcriptional fusion bio-reporters by the fractionated extracts were not as clear as when the whole extract was assayed using the same reporters. The lack of clear bio-reporter inhibition using the fractionated extracts may indicate that the compound(s) causing inhibition when assaying the whole extract did not bind to the solid phase extraction columns used by this study in the fractionation process. In spite of this, small by significant reductions in the bioluminescent resulting from the activation of the lasI and lasB bio-reporters were observed after exposure to the I. galbana 60% methanol fraction and in the bioluminescent resulting from the activation of the rhII and rhIA bio-reporters when exposed to the N. oculata 70% fraction.

The fractionated microalgal extracts were also show to decrease protease activity in all four marine bacterial pathogens assayed. Previous studies have shown that all four of these species produce AHLs and have also shown them to possess AHL-mediated QS systems (Santos Ede et al., 2011, Temperton et al., 2011, Tait et al., 2010, Milton et al., 2001, Swift et al., 1997). Previous studies have also shown that in A hydrophila and in various Vibrio species, the production of protease virulence factors is regulated by both AHL and AI-2 mediated QS systems (Milton, 2006, Swift et al., 1999). As such, the inhibition of protease activity in marine pathogens observed by this study, has a result of exposure to extracts from microalgal cultures, may indicate quorum-quenching activity by compound(s) produced by these marine microalgae. Interestingly, the fractions from microalgal extracts effecting the transcription of AHL

regulated genes in P. aeruginosa were, in the most part, consistent with those that affected protease activity in the marine pathogens. As such, it is therefore likely that any compound(s) inhibiting AHL-mediated QS are present in these fractions.

The evidence that different solvent fractions from the two species affected different P. aeruginosa bio-reporters and protease activity in marine pathogens may indicate the production of a range of secondary metabolites that elute into different fractions, which in combination have the ability to effect AHL regulated gene expression by potential quorum-quenching abilities. Previous studies have shown eukaryotic species including C. reinhardtii, D. pulchra and Medicago truncatula to produce up to 30 separately identifiable compounds that inhibit bacterial QS (Teplitski et al., 2004, Gao et al., 2003, Manefield et al., 2002). Further fractionation combined with chemical analysis techniques such as LCMS or nuclear mass spectroscopy would be required to positively identify the compound(s).

The biological aim of producing compound(s), which inhibit AHL regulated QS may by to disrupt bacterial signalling in order to prevent infection. This is clearly demonstrated to be the case for the production of halogenated furanones by the marine macroalgae D. pulchra as these compounds were shown to prevent swarming motility in S. liquefaciens reducing the bacteria's ability to colonise. Previous studies have shown marine microalgae have the ability to modulate the bacterial population to which they are associated with (Sapp et al., 2007, Salvesen et al., 2000). Inhibition of bacterial QS by producing quorum-quenching compounds may be one method by which microalgae can control the bacterial population to their advantage. As microalgal cultures are heavily used in aquaculture it may be possible to utilise these antimicrobial

V. anguillarium by Roseobacter spp. due to co-culturing with N. oculata has been recently reported (Sharifah and Eguchi, 2011). It is of note that both V. anguillarium and Roserbacter clade bacteria have been reported to regulate gene expression using AHL-mediated QS. It maybe that quorum-quenching activity by N. oculata is promoting Roseobacter growth at the expense of V. anguillarium. Results presented by this study showing inhibition of protease production in marine pathogens noted to effect aquacultural stocks by three microalgal species currently used as feed sources in aquaculture further expands these findings, lending credence to the potential of exploiting microalgal quorum-quenching to control infection in aquaculture. In addition, this study has shown that compound(s) produced by marine microalgae may affect virulence gene expression in human pathogens. This makes microalgae and the quorum-quenching compounds they produce attractive candidates for novel antimicrobials as they can reduce the virulence of a pathogen without inducing a selective pressure, causing antimicrobial resistance (Müh et al., 2006, Zhang, 2003).

5.3.2. Summary Conclusions and Future Directions

This study has identified quorum-quenching activity in three marine microalgal species. This activity has been shown to effect AHL bio-reporters bioluminescence in the presence of various exogenous AHLs, the expression of a number of P. aeruginosa virulence factors previously shown to be in part regulated by AHL-mediated QS and protease expressing in a number of marine bacterial pathogens. Although advanced by this study, research into QS inhibition by marine microalgae is clearly still in its infancy. As yet, no compound(s) have been identified as the cause of the quorum-quenching

activity seen in this study. The identification of these compounds is clearly the next step in this research.

Research in this area should progress in the following directions:

- Continue fractionation of extracts from stationary phase microalgal cultures to purify and therefore identify quorum-quenching compound(s) produced by these species. This can be conducted by continuing with the method laid out in this thesis (Section 2.10.5) or by the use of reverse-phase C18 analytical HPLC (see Teplitski et al., 2004), assaying each fraction for quorum-quenching activity using AHL bioreporters (see Section 2.10.4).
- Once quorum-quenching compound(s) have been sufficiently purified, initiate compound(s) identification using chemical analysis techniques such as LCMS, or for more precise identification, nuclear magnetic resonance spectroscopy.

Chapter 6

General Discussion

6.1 General Discussion

The first aim of this study was to examine the relationship the green seaweed Ulva has with its epiphytic bacterial community, concentrating on the impacts of AHL signal molecule production by these bacteria. As such, this thesis has characterised Ulva's epiphytic bacterial population in terms of both taxonomic composition and the AHL signal molecules being produced by members of this population. As previous work has shown that AHL signal molecule production affects the settlement of Ulva zoospores and that bacteria affect the growth morphology of Ulva spp. This thesis presented data examining the effects AHLs, and bacteria producing AHLs have on Ulva zoospore germination and the early growth of the Ulva germling. This thesis then progressed to investigate AHL production and degradation in two groups of bacteria that can be readily isolated from biofilms present on both the surface of the Ulva thallus and on the rocks colonised by Ulva spp., Shewanella and the Bacteroidetes. Finally, this thesis diverged into a study of quorum-quenching behaviour in three species of marine microalgae prolifically used in aquaculture.

Marine algae have been shown to possess diverse populations of epiphytic bacteria; these bacteria in general exist in aggregated biofilm communities and as such will behave in a different manner to planktonic bacteria in the water column. These epiphytic bacterial populations have been shown to interact with their algal host organisms in a number of different ways, many of which have been discussed in this thesis (Tait et al., 2009; Rao et al., 2006; Miller et al., 2005; Matsuo et al., 2003,). In some cases, these interactions may be symbiotic. Examples of potential symbiotic interactions include the utilisation of bacterially produced cyanocobalamin (Vitamin

B₁₂) by algae, many of which are cyanocobalamin auxotrophs (Croft et al., 2006). In return the bacteria benefit from carbon sources produced by algal metabolism which provide growth substrates (Croft et al., 2006; Croft et al., 2005). However, the true symbiotic nature of this interaction has been questioned with suggestions that algae merely scavenge vitamins (Droop, 2007). These interactions have also been proven to be dynamic and can be either beneficial or detrimental to the host. The prominent microalgal species Emiliania huxleyi and bacteria from the Roseobacter clade show both a mutualistic and pathogenic interaction depending on the senescence of the algal host and is mediated by small molecule production by both host and symbiont (Seyedsayamdost et al., 2011). In healthy E. huxeyli cultures the algae provide a substrata and nutrients for bacterial growth and the bacteria produce antibiotics preventing algal infection. As the algae matures it releases a signal (pCA) which prompts Roseobacter species to produce algaecides killing the host and allowing the bacteria to access the subsequent nutrient released proliferate to a healthy host (Sevedsayamdost et al., 2011).

6.2 Ulva Zoospore Settlement, Germination and Germling Growth

Small molecule mediated algal/bacterial interactions have also been proven to take place utilising the signalling molecules that many species, including those from the marine environment, produce for bacterial QS. A prime example is the interaction between bacteria and Ulva spp., where AHL signalling molecules attract Ulva zoospores to microcolonies within bacterial biofilms via the mediation of zoospore swim speed (Wheeler et al., 2006; Tait et al., 2005; Joint et al., 2002). A key finding

presented in this thesis has shown that the interaction between Ulva spp. and their indigenous signal molecule producing bacterial population does not just affect zoospore settlement but also the germination process and the early growth of the Ulva zoospore. In the presence of AHLs Ulva zoospore germination and the growth of the resultant Ulva germlings were both shown to be retarded. This reduction in the rate of early growth may allow the Ulva germling to obtain an indigenous epiphytic bacterial population which is of benefit as bacteria have been shown to modulate the healthy differentiation of Ulva spp. into mature adult plants. This hypothesis is supported by data presented in sections 3.2.1 and 3.2.2 that show Ulva possess an epiphytic bacterial population similar to the previously characterised population found in rocky shore biofilms (Tait et al., 2009).

The interactions between Ulva spp. and bacteria is often cited as an example of 'cross kingdom communication' owing to AHL bacterial signalling molecules, which are normally involved in population density dependent communication, mediating these interactions (Joint et al., 2007). However, the data presented in this study and the previous studies into AHL-mediated Ulva/bacterial interactions only reveals the algal response to AHL signal molecules. As yet, no data has been presented showing Ulva spp. to produce any form of signal molecule which mediates a phenotypic change in bacterial populations. As true communication requires both communicating partners to understand and participate in the communicative process, labelling this interaction as an example of cross kingdom communication may be a somewhat broad ranging statement. Additionally, very little is understood of the mechanism behind reduced swimming speed of Ulva zoospores by AHLs and this study has not eluded any mechanism for how AHLs modulate zoospore germination and germling growth. As such, it is most

likely that Ulva spp. utilise the AHLs being produced by members of its indigenous population as cues for various growth processes, be this zoospore settlement, zoospore germination or the early growth of Ulva germlings. Since the interaction between AHLproducing bacteria and Ulva has been established, the mechanisms which drive this interaction should be investigated. Joint et al. (2007) propose that an influx of calcium into the cytosol caused by the presence of AHLs is responsible for the modulation of Ulva zoospore swim speed, which in turn leads to increased zoospore settlement in the presence of AHL-producing biofilms (Joint et al., 2007; Wheeler et al., 2006). This hypothesis was supported by manganese quenching experiments in which the fluorescence of a calcium indicator dye was reduced as a result of manganese entering the cell, a direct result of calcium influx channels opening when zoospores were exposed to high concentrations of 3-oxo-C12-HSL (Joint et al., 2007; Taylor et al., 1996). Calcium has also been shown to affect flagella swim speed in other marine eukaryotes such as Chlamydomonas spp. (Kamiya and Witman, 1984). Comparative genomic and proteomic analysis of Ulva to both bacterial and micro-eukaryotic species sequence information may identify potential receptors which have the ability to detect AHLs. Such analysis may also identify subsequent signalling pathways regulating these previously reported alterations in calcium influx which appear to modulate swim speed. However, to accomplish this, it will be necessary to first sequence the genomes of key Ulva species. Additionally, comparisons of both RNA and protein expression profiles between Ulva zoospores or germlings exposed to AHLs with those not exposed to AHLs may also provide information on the transcription of proteins involved in modulating this interaction and answer the question of whether AHLs produced by bacteria elicit a change in Ulva's gene expression.

There is currently no direct evidence that epiphytic bacteria benefit from being associated with Ulva spp. However, it is possible to hypothesise that epiphytes may obtain nutrients because of the association with Ulva spp. A number of studies have shown that bacteria belonging to the Rhodobacteraceae have the ability to degrade dimethylsulphoniopropionate (DMSP) to dimethylsulphide (DMS) (Wagner-Döbler and Biebl, 2006). DMSP is a sulphonium compound produced by a range of marine algae including Ulva, this compound is released into the water and its degradation can provide a source of sulphur for marine bacteria (Wagner-Döbler and Biebl, 2006; Yoch, 2002; Van Alstyne et al., 2001). Species belonging to the Rhodobacteraceae have been shown to be able to degrade DMSP via two biochemical pathways and are also closely associated with marine algae, including, as this study has shown, Ulva (Miller and Belas, 2004). It is therefore possible that bacteria benefit by being provided with a source of sulphur from the DMSP being produced by Ulva spp. Similar relationships have been observed between Rhodobacteraceae bacteria and the microalgal species Pfiesteria spp. and DMSP has also been shown to act as a chemo-attractant to these bacteria (Miller and Belas, 2004). As this is only a hypothesis as to how bacteria benefit from association with Ulva spp., the Ulva/bacterial interaction cannot yet be truly considered as symbiotic. It would therefore be interesting to observe if the bacterial population associated with Ulva spp. changes during the growth of the plant in response to the different nutrients available as a result of the different stages in plant growth. Currently the only studies looking at Ulva bacterial populations either have taken snap shots from one point in the Ulva life cycle as this study did, or have tracked seasonal change in the Ulva bacteria/population (Burke et al., 2011; Tujula et al., 2010).

In the presence of a variety of synthetic AHLs Ulva zoospore germination and germling growth was assessed. There was no clear pattern of specificity seen with regards to which AHLs affected germination or germling growth, although assays were more consistent using AHL with long acyl side chains. This increased consistency was attributed to long chain AHLs being more stable in the alkaline pH of seawater (Hmelo and Van Mooy, 2009; Tait et al., 2009). As bacterial biofilms producing AHLs with short and mid-length acyl side chains showed significant effects on Ulva zoospore germination and germling length in this study, and to Ulva zoospore settlement in previous studies (Tait et al., 2005), it is possible that Ulva does not respond to a specific AHL as a cue during zoospore settlement, germination and early growth. This would be consistent with data presented by this study and previous studies suggesting that the bacteria associated with Ulva spp. and the surfaces colonised by Ulva spp. produce a wide variety of different AHL signalling molecules (Tait et al., 2009). As other signalling molecules such as alkyl quinolones have been shown to be present in the marine environment being produced by marine bacteria such as Alteromonas spp., (Long et al., 2003), it would be interesting to investigate if these molecules affect Ulva zoospore settlement and germination. As such, similar assays to those used by this study could be carried out using synthetic alkyl quinolone molecules and model organisms such as P. aeruginosa, which produce these quinolone molecules.

This study has increased our understanding of the complex relationship that exists between the algae and bacteria and in to the future the commercial benefits of these observations may be far reaching. Ulva is a prominent species involved in marine macrofouling which incurs massive costs to the marine industry (Callow and Callow, 2002). Currently antifouling methods involve treating submerged surfaces with

chemical paints, which can be detrimental to the marine environment (Thomas et al., 2001; Tolosa et al., 1996). A greater understanding of Ulva growth and development in conjunction with associated bacterial species may lead to new antifouling treatments that are more effective with reduced environmental implications. This study and previous work suggest that AHL signalling is beneficial to the growth processes of Ulva spp. therefore one such avenue of research for producing novel antifouling treatments may be to develop treatments that inhibit AHL signalling and therefore reduce Ulva settlement and affect the wild type growth process of the Ulva germling. Alkyl quinolone molecules have additionally been shown to be algaecidal, inhibiting the growth of phytoplankton at concentrations above $10~\mu M$ (Long et al., 2003). As such, investigating the effects quinolone producing bacteria have on Ulva growth may be another avenue for novel antifouling treatments.

6.3 AHL-mediated Quorum Sensing in Shewanella and the Bacteroidetes

Section 4.2.2 described AHL signal molecule production in two groups of marine bacteria, Shewanella and the Bacteroidetes. These two bacterial groups were deemed of relevance to this study for separate reasons. Shewanella spp. can be readily isolated from Ulva spp. and has previously been shown to modulate Ulva zoospore settlement via the production and subsequent inactivation of AHLs. The Bacteroidetes are of relevance as this study has shown that they are highly abundant in the Ulva epiphytic community and previous work has shown similar abundance in rocky shore biofilms colonised by Ulva spp. (Tait et al., 2009). This study has shown AHL production by a number of different strains of both Shewanella and Bacteroidetes

bacteria. Concerning the Bacteroidetes, this is of particular interest as AHL production in this phylum is currently a novel topic for research and expands our understanding of AHL signal molecule production to bacteria outside of the proteobacterial phylum. As previous work has only proven AHL production in one strain of Bacteroidetes, the discovery that a number of Bacteroidetes strains from differing taxonomic backgrounds produce AHLs, specifically C4-HSL, points towards conserved signal molecule production by this bacterial phylum. The study of Bacteroidetes AHL production also underpinned the importance of using the latest techniques to identify AHL production. In this study the use of LCMS to characterise AHL production in both the Bacteroidetes and in other marine isolates produced an accurate AHL profile which would have been virtually impossible using traditional bio-reporters because of the alkaline nature of marine media hydrolysing AHLs (Yates et al., 2002).

This study has increased the range of Shewanella bacteria known to be producing AHLs to include strains of S. putrafaciens and S. baltica. As it is clear that both the Bacteroidetes and Shewanella spp. are actively producing AHLs this study attempted to identify AHL synthase and response regulators in both groups. In silico analysis failed to identify any sequences homologous to LuxI/LuxR or LuxM/LuxN. This study therefore concludes that both these groups of bacteria possess novel, yet to be identified, AHL synthases and response regulators. Given the abundance of these two groups and other bacteria in the marine environment and the expanding list of marine bacteria known to be producing AHLs, it is possible that there is a wide variety of novel systems used by marine bacteria to produce and detect AHLs. Further genomic sequencing of the ocean's bacterial populations and subsequent bioinformatic analysis currently being carried out by studies such as Global Ocean Sampling Expedition may

reveal such novel systems (Venter et al., 2004). Alternatively, the identity of the AHL synthase(s) may be revealed using a transposon library constructed in Bacteroidetes strains which are actively producing AHLs. AHL bio-reporters which detect C4-HSL production would then be used to identify mutants which were unable to produce AHLs, this technique does however present drawbacks owing to the high levels of background light associated with C4-HSL bio-reporters which could lead to a number of false positive results. A Bacteroidetes genomic library may also be used to identify novel AHL synthases using bio-reporters which respond to C4-HSL. These strategies have been used for the detection of AHL-mediated QS systems in a number of enteric bacteria and in environmental metagenomic libraries (Hao et al., 2010; Swift et al., 1993). In Shewanella, the screening of genomic libraries being expressed in a number of different AHL bio-reporters to identify AHL synthases and transducers has been exhaustively attempted and lead a number of false positive results (Tait et al., 2009). The turnover of Shewanella AHLs by the expression of AHL acylases complicates this process and as such mutating the cognate Shewanella AHL inactivating enzyme would be required before attempting this screening procedure. As this study has identified a conserved AHL acylase in a number of Shewanella strains, its mutation would be the next logical step to determining if Shewanella spp. express more than one AHL inactivating enzyme. If this proved successful, then putative AHL synthase and response regulator screening genes could be screened in a transposon library generated from the newly constructed mutant.

As well as investigating AHL production in Shewanella spp. this study also concentrated on the phenomenon of AHL inactivation by Shewanella. Enzymic AHL inactivation by bacteria is generally carried out either by AHL lactonase enzymes or

AHL acylase enzymes, both of which inactivate AHLs via hydrolysis but at different points in the AHL molecule (Lin et al., 2003; Dong et al., 2000). AHL inactivation in Shewanella spp. has been observed in previous studies and one study has identified an acylase enzyme in an environmental Shewanella isolate with the ability to cleave the homoserine lactone ring from the acyl side chain of AHLs known to be produced by Shewanella (Tait et al., 2009; Morohoshi et al., 2008). Via in silico screening this study found putative AHL acylase sequences to be present in a number of different Shewanella genomes. Using this genomic information, this study confirmed the presence of sequences with strong identities to AHL acylase genes in a number of Shewanella strains isolated from the marine environment. The identified acylase, named AacS, cloned from one such strain inactivated synthetic AHLs and the long chain AHLs produced the human pathogen Y. pseudotuberculosis. Inactivation of long chain AHLs was consistent with previous studies investigating AHL turn over by Shewanella spp. (Tait et al., 2009). AHL inactivation by Shewanella is not a unique phenotype in the marine environment as AHL inactivating activity in other marine bacteria such as Vibrio spp. and in the Bacteroidetes has been reported (Romero et al., 2011; Romero et al., 2010; Tait et al., 2010). AHL inactivation would therefore affect algal bacterial interactions mediated by QS signal molecules. Tait et al. (2009) have already shown that the enzymatic inactivation of AHLs in poly-microbial biofilms abolishes the attraction of Ulva zoospores during settlement (Tait et al., 2009). Data presented in this thesis showed both Ulva zoospore germination to be repressed by AHLs. This study also revealed poly-microbial communities composed of Vibrio spp., Bacteroidetes and Shewanella spp. to be present on Ulva spp. It is therefore conceivable that enzymic inactivation of AHLs will have stimulatory effects on germination as oppsed to the negative effect reported by Tait et al. (2009) on zoospore settlement (Tait et al., 2009).

6.4 Quorum-quenching by Marine Microalgae

Finally, this study investigated the effects marine microalgae have on bacterial QS. The microalgae selected for this study are species commonly utilised in aquaculture (Muller-Feuga, 2000; Borowitzka, 1997). Many bacteria shown to be pathogenic to fish populations reared in aquaculture have been shown to possess AHL-mediated QS systems, such bacteria include A hydrophyla, A salmonicida, V. anguillarium, V. tubiash, V. vulnificans and Yersinia ruckeri (Temperton et al., 2011; Fernández et al., 2007; Milton, 2006; Swift et al., 1997). Infection in aquacultural stocks is routinely prevented by supplementing feed with antibiotics; this has the potential to increase the prevalence of antibiotic resistant bacteria which is currently a significant cause for concern for both the UK Health Protection Agency and the World Health Organisation (Cabello, 2006). As such, alternative treatments that do not exert a selective pressure on pathogenic bacteria are currently being investigated.

One such alternative is to inhibit QS in these pathogens as QS often regulates virulence factor expression (Fernández et al., 2007; Swift et al., 1999). Inhibition of QS can result from the production of signal molecule inactivating enzymes, such as AHL lactonases and acylases, (Section 1.3.2), or by the production of compounds which have the ability to quench QS (Lin et al., 2003; Dong et al., 2000; Manefield et al., 1999). Compounds that inhibit QS by quenching the effects of AHL type signalling molecules

have been shown to be produced by a number of different marine organisms, including the freshwater microalgae C. reinhardtii (Skindersoe et al., 2008; Teplitski et al., 2004). This study has identified that the marine microalgal species N. oculata, I. galbana and T. suecia also produce and excrete a compound(s) that appears to have the ability to inhibit AHL-mediated QS. This was shown using AHL bio-reporters (Section 5.2.2). In assays using marine bacterial fish pathogens and in the opportunistic human pathogen P. aeruginosa, the expression of a number of genes previously linked with AHL-mediated QS were also shown to be inhibited by microalgal extracts. As such this is the first study to demonstrate a phenotypic effect of putative microalgal quorum-quenching apart from in an AHL bio-reporter. This study did not identify specifically what compound(s) were causing this apparent quorum-quenching activity; however, purification and characterisation of this compound(s) may reveal novel antibiotic compounds capable of attenuating virulence. Such compounds may be of benefit to the bio-control of marine pathogens that can affect aquacultural productivity and potentially human health and in the control of human pathogens in a broader medical setting.

Exploiting microalgal quorum-quenching compound production to attenuate virulence in marine fish pathogens detrimental to aquaculture benefits productivity without resorting to supplementing feed with antibiotics routinely used to treat human infections. This could both reduce the likelihood of spreading antibiotic resistance and reduce aquacultural costs as we are already using these microalgal species as food stocks. Additionally, exploiting quorum-quenching compound production by microalgal feed stocks in the bio-control of marine pathogens that effect human health would also be beneficial to aquaculture. The main route of transmission of the

pathogenic marine Vibrio species such as Vibrio parahaemolyticus is through the ingestion of raw oysters (Daniels et al., 2000). V. parahaemolyticus has been shown to regulate its virulence factors via QS and the V. harveyi like AHL synthase LuxM has been shown to be present in the V. parahaemolyticus genomes (Henke and Bassler, 2004; Makino et al., 2003). By inhibiting QS, virulence may be reduced to levels sufficient to prevent disease in oysters and subsequently prevent human disease. This is compounded by data presented in section 5.2.4.2 that showed extracts from these microalgae effected QS mediated protease production in a number of marine pathogens both detrimental to aquacultural production and similar to marine pathogens that effect human health such as V. parahaemolyticus and V. vulnificus.

Sections 5.2.3 and 5.2.4.1 revealed that putative quorum-quenching compounds produced by the marine microalgae affected the expression of genes associated with virulence in the opportunistic human pathogen P. aeruginosa. This is of importance as the development of novel antimicrobial compounds, specifically those that do not induce a selective pressure on bacterial populations is a prominent area of current medical research. As such, it would be of interest to investigate how the production of quorum-quenching compounds in microalgae is controlled. Understanding how these compounds are produced, it may be possible to culture microalgae under conditions that select for quorum-quenching compound production. The initial step in further understanding quorum-quenching in microalgae would be to advance the work in this study by identifying these compounds(s). This can be achieved by continuing the fractionation of microalgal extracts initiated by this study and screening the resultant fractions with AHL bio-reporters. This study used solid phase extraction cartridges to carry out fractionation, however a technique such as reverse-phase HPLC could also be

used and has proven to be successful in previous studies in identifying quorum-quenching compound(s) (Teplitski et al., 2004). Identification of the quorum-quenching compound(s) would be best carried out using chemical analysis technique such as mass spectroscopy as has been used previously in the identification of similar acting compounds produced by freshwater microalgal species which inhibit bacterial QS, (Teplitski et al., 2004), or full characterisation using techniques such as nuclear magnetic resonance spectroscopy. These compounds could then be easily purified and investigated for use in medicine to tackle both acute and systemic bacterial infections, where virulence is highly regulated by QS, such as in P. aeruginosa.

From an ecological perspective, it would also be interesting to investigate if microalgal quorum-quenching activity could affect macroalgal/bacterial interactions such as those observed in Ulva zoospore settlement, Ulva zoospore germination and the early growth of Ulva spp. germlings in a similar way to enzymic AHL inactivation by bacteria such as Shewanella. If so, it would add a deeper layer of complexity to these bacterial/algal interactions taking place in the marine environment. Additionally if such microalgal quorum-quenching compounds do affect these interactions, they may also have potential usages in the production of novel antifouling treatments. Overall, this thesis has furthered our understanding of how bacterial signalling molecules and their inactivation either by enzymes or by quorum-quenching compounds facilitate bacterial/algal interactions in the marine environment. It is also likely that as interactions are taking place between epiphytic bacteria and macroalgae like Ulva, similar bacterial/algal interactions may be facilitated by the production and turnover of QS signalling molecules in other marine species. As such, our current understanding of the importance of QS signalling in the marine environment and its potential biological,

environmental and biotechnological implications is limited and therefore merits further
research.

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Appendices

Appendix 1: Bacterial QS Systems

Table listing a number of bacteria known to regulate gene expression via QS, the type of signal molecule they produce, the synthase and response regulator genes and the phenotypes regulated.

Bacteria	Signal Molecule	Synthase/ response	Phenotypes Regulated	References
		regulator		
Acidithiobacillus ferrooxidans	AHLs	AfeR/I	Biofilm formation	(Rivas et al., 2005)
			Nutrient acquisition	(Gonzalez et al., 2012)
Aeromonas hydrophila	AHLs	AhyR/I	Biofilm formation	(Swift et al., 1997)
			Exoproteases	(Swift et al., 1999)
			Virulence	(Lynch et al., 2002)
Aeromonas salmonicida	AHLs	AsaR/I	Exoprotease	(Swift et al., 1997)
				(Swift et al., 1999)
Agrobacterium tumefaciens	AHLs	TraR/I	Plasmid Conjugation	(Fuqua and Winans, 1994)
				(Piper et al., 1993)
Agrobacterium vistis	AHLs	AvsR/I	Virulence	(Wang et al., 2008)
				(Hao et al., 2005)
Berkholderia cenocepacia	AHLs	CepR/I	Exoenzymes	(Lewenza et al., 1999)
	AQs		Biofilm formation	(Sokol et al., 2003)
			Moltility	(Huber et al., 2001)
			Siderophore production	

Burkholderia pseudomallei	AHLs	BpsR/I	Virulence	(Song et al., 2005)
				(Ulrich et al., 2004)
Chromobacterium violaceum	AHLs	CviR/I	Virulence	(McClean et al., 1997)
			Pigmentation	
Enterobacter agglomerans	AHLs	EagR/I	?	(Swift et al., 1993)
Enterococcus faecalis	Autoinducing	Cyl operon	Cytolysin production	(Reading and Sperandio, 2006)
	Peptides			
Erwinia carotova	AHLs	CarR/I	Carbapenem production	(Bainton et al., 1992)
			Virulence	(Barnard and Salmond, 2007)
Obesumbacterium proteus	AHLs	OprR/I	?	(Swift et al., 1999)
Pantoea stewartii	AHLs	EsaR/I	Exopolysaccharide production	(Beck von Bodman and Farrand, 1995)
Pseudomonas aeruginosa	AHLs	LasR/I, RhlR/I	Exoenzyme	(Latifi et al., 1996)
	AQs	PQS operon	Biofilm formation	(Glessner et al., 1999)
			Motility	(Latifi et al., 1995)
			Virulence	(De Kievit et al., 2001)
Pseudomonas fluorescens	AHLs	PhzR/I	Mupirocin production	(Shaw et al., 1997)
				(El-Sayed et al., 2001)
Pseudomonas putida	AHLs	PpuR/I	Biofilm formation	(Dubern et al., 2006)
Pseudomonas syringae	AHLs	PsyR/I	Exopolysaccharide production	(Swift et al., 1999)
			Motility	(Quinones et al., 2005)
			Virulence	
Rhizobium leguminosarum	AHLs	RhiR/I	Root nodulation	(Rodelas et al., 1999)
			Plasmid transfer	(Wisniewski-Dye and Downie, 2002)
Rhodobacter sphaeroides	AHLs	CerR/I	Cellular aggregation	(Puskas et al., 1997)

Serratia liquifacines	AHLs	SwrR/I	Motility	(Eberl et al., 1996)
			Biofilm formation	(Horng et al., 2002)
			Biosurfactant production	(Van Houdt et al., 2007b)
Serratia marcescens	AHLs	SpnR/I	Motility	(Horng et al., 2002)
			Biofilm formation	(Van Houdt et al., 2007a)
			Biosurfactant production	(Eberl et al., 1996)
Sinorhizobium meliloti	AHLs	SinR/I	Root nodulation	(Marketon and Gonzalez, 2002)
			Symbiosis	(Marketon et al., 2003)
Staphylococcus aureus	Autoinducing	Arg operon	Virulence	(Novick, 2003)
	Peptides		Biofilm formation	(Ji et al., 1995)
Streptococcus mutans	Autoinducing	Com operon	Biofilm formation	(Reading and Sperandio, 2006)
	Peptides			(Li et al., 2002)
Vibrio anguillarum	AHLs	VanR/I, VanM/N	Virulence	(Milton et al., 1997)
				(Defoirdt et al., 2005)
Vibrio cholera	CAI-1	CqsA	Biofilm formation	(Hammer and Bassler, 2003)
	AI-2	LuxS		(Waters et al., 2008)
Vibrio fisheri	AHLs	LuxR/I, AinS	Bioluminescence	(Engebrecht and Silverman, 1984)
				(Hastings and Nealson, 1977)
Vibrio harvyi	AHLs	LuxM/N	Bioluminescence	(Bassler et al., 1993)
	AI-2	LuxS		
Yersinia enterocolylica	AHLs	YenR/I	Motility	(Throup et al., 1995)
				(Atkinson et al., 2008)
Yersinia pestis	AHLs	YpeR/I	Biofilm formation	(Throup et al., 1995)
				(Bobrov et al., 2007)

Yersinia pseudotuberculosis	AHLs	YpsR/I, YtbR/I	Biofilm Formation	(Atkinson et al., 1999)
			Motility	(Atkinson et al., 2011)

Appendix 2: F/2 Media Supplements

F/2 Trance Metal Solution

In 11 d.H₂O (Autoclave)

 Na_2EDTA 4.36 g FeCl₃ 6H₂O (Ferric Chloride) 3.15 g

Primary trace metals 1ml of each stock solution

Primary Trace Metal Solutions

	In 10 ml d.H ₂ O (filter sterilise)
CuSO ₄ 5H ₂ O	0.100 g
ZnSO ₄ 7H ₂ O	0.220 g
CoCl ₂ 6H ₂ O	0.100 g
MnCl ₂ 4H ₂ O	0.180 g
NaMoO ₄ 2H ₂ O	0.063 g

F/2 Vitamin Stock solution

In 100 ml d.H₂O (Filter

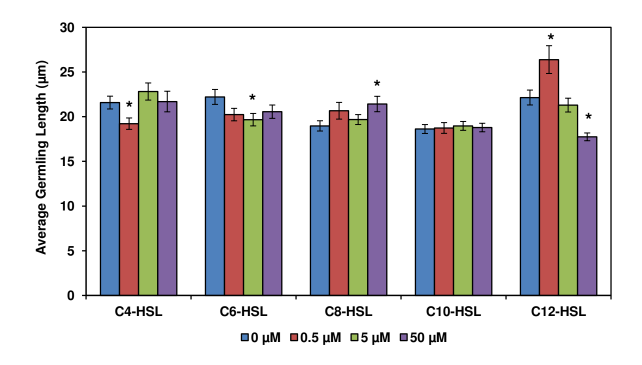
sterilise)

 $\begin{array}{lll} \text{Biotin} & 0.10 \text{ mgml}^{-1} \text{ solution} & 1.00 \text{ ml} \\ \text{Vitamin B}_{12} & 1.00 \text{ mgml}^{-1} \text{ solution} & 0.10 \text{ ml} \\ \text{Thiamine HCl} & 0.02 \text{ g} \end{array}$

Appendix 3: Ulva Zoospore Germination in the Presence of Synthetic AHLs

Graph displaying the average length of Ulva germlings exposed to various concentrations of synthetic AHLs after 24 h incubation. Error bars represent 95% confidence limits.

* Represents where average zoospore length is significantly different from that where no exogenous AHL was added (0 μ M) (one-way ANOVA p < 0.05).



Appendix 4: Construction of LuxI-Type AHL Synthase Consensus Sequences

LuxI-Type AHL Synthase Consensus Sequence

LuxI Consensus Sequence

MLXIFDXXXVSYDXLSEEKXXXKELFRLRKKXFKDRLGWDVNCXNGMXXXEFDXXQYDNXNTRYLLGIXD XXXGQXVGCVRLIETTXXXPNMLTGTFFXLLXDDGALPEXXGXYIXESSRFFXXXDKARARLLXGNXXXY PLSLVXLFLSMINYARANGYTGIVTVVSRAMERILKRSGWPIERIGQXXGHXXEKGEXIYLXLHLPIDXX XQERLARRINQPLQGPSSXLLTWPLSLP

Protein Sequences of Published LuxI-Type AHL Synthases

AhyI (Aeromonas hydrophyla)

MLVFKGKLKEHPRWEVENELYRFRNRVFSDRLGWDVESHRGLEQDSFDTPDTHWVLIEDEEGLCGCIRLL SCAQDYMLPSIFPTALAGEAPPRSNDVWELTRLAIDAERAPRLGNGISELTCVIFREVYAFAKAQGIREL VAVVSLPVERIFRRLGLDIERLGHRQAVDLGAVRGVGIRFHLDERFARAVGQPLQGAYDEARELVTE

AsaI (Aeromonas salmonicida)

MLVFKGKLKEHPRWEVENELYRFRNRVFSDRLGWDVESHRGLEQDSFDTPDTHWVLIEDEEGLCGCIRLL SCAQDYMLPSIFPTALAGEAPPRSSDVWELTRLAIDANRAPRMGNGVSELTCVIFREVYAFARAKGIREL VAVVSLPVERIFRRLGLPIERLGHRQAVDLGAVRGVGIRFHLDERFARAVGHPMQGEYADARELVTE

Carl (Erwinia carotovora)

MLEIFDVNHTLLSETKSGELFTLRKETFKDRLNWAVQCTDGMEFDFTQYDNNNTTYLFGIKDNTVICSLR FIETKYPNMITGTFFPYFKEINIPEGNYLESSRFFVFTDKSRAKDILGNEYPISSMLFLSMINYSKDKGY DGIYTIVSHPMLTILKRSGWGIRVVEQFTGLSEKEERVYLVFLPVDDENQEALARRINRSGTFMSNELKQ WPLRVPAAIAQA

CerI (Rhodobacter sphaeroides)

MIFIIDSLNLREHADIVKDMFRLRKRVFADRLGWDVQISQGMERDFTRFDDLDPAHVVSVDDEGRVVGCM RLLQTTGPHMLSDVFSSILDGEPPLRSATLWEATRFFTCVDTDRLVSGRARNSIAYVTSEVMIGAFEFAM SAGVTDAVAVIDPVMDRVLKRSGNAPQFTGYVGTPKPMGKVTALAALMDCSEERVKRIRDFAGIYHDVTQ PQTVIA

EagI (Pantoea agglomerans)

MLEIFDVSYNDLTERRSEDLYKLRKITFKDRLDWAVNCSNDMEFDEFDNSGTRYMLGIYDNQLVCSVRFI DLRLPNMITHTFQHLFGDVKLPEGDYIESSRFFVDKNRAKALLGSRYPISYVLFLSMINYARHHGHTGIY TIVSRAMLTIAKRSGWEIEVIKEGFVSENEPIYLLRLPIDCHNQHLLAKRIRDQSESNIAALCQWPMSLT VTPEQV

EchI (Erwinia chrysanthemi)

MLEIFDVSFSLMSNNKLDEVFTLRKDTFKDRLDWAVNCINGMEFDEYDNEHTTYLLGVKEGKVICSVRFI EIKYPNMITGTFYSYFDNLKIPEGNYIESSRFFVDRDRVRNLIGTRNPACVTLFLAMINYARKYHYDGIL TIVSHPMLTLLKRSGWRISIIQQGLSEKQERIYLLHLPTDDDSRHALIERITQMTQAESEQLKTLPLLVP LA

EsaI (Pantoea stewartii)

MLELFDVSYEELQTTRSEELYKLRKKTFSDRLGWEVICSQGMESDEFDGPGTRYILGICEGQLVCSVRFT SLDRPNMITHTFQHCFSDVTLPAYGTESSRFFVDKARARALLGEHYPISQVLFLAMVNWAQNNAYGNIYT IVSRAMLKILTRSGWQIKVIKEAFLTEKERIYLLTLPAGQDDKQQLGGDVVSRTGCPPVAVTTWPLTLPV

ExpI (Pectobacterium carotovorum)

MLEIFDVSYTLLSEKKSEELFTLRKETFKDRLNWAVKCINGMEFDQYDDDNATYLFGVEGDQVICSSRLI ETKYPNMITGTFFPYFEKIDIPEGKYIESSRFFVDKARSKTILGNSYPVSTMFFLATVNYSKSKGYDGVY TIVSHPMLTILKRSGWKISIVEQGMSEKHERVYLLFLPVDNESQDVLVRRINHNQEFVESKLREWPLSFE PMTEPVG

LasI (Pseudomonas aeruginosa)

MIVQIGRREEFDKKLLGEMHKLRAQVFKERKGWDVSVIDEMEIDGYDALSPYYMLIQEDTPEAQVFGCWR ILDTTGPYMLKNTFPELLHGKEAPCSPHIWELSRFAINSGQKGSLGFSDCTLEAMRALARYSLQNDIQTL VTVTTVGVEKMMIRAGLDVSRFGPHLKIGIERAVALRIELNAKTQIALYGGVLVEQRLAVS

LuxI (Vibrio fischeri)

MTIMIKKSDFLAIPSEEYKGILSLRYQVFKQRLEWDLVVENNLESDEYDNSNAEYIYACDDTENVSGCWR LXPTTGDYMLKSVFPELLGQQSAPKDPNIVELSRFAVGKNSSKINNSASEITMKLFEAIYKHAVSQGITE YVTVTSTAIERFLKRIKVPCHRIGDKEIHVLGDTKSVVLSMPINEQFKKAVLN

PhzI (Pseudomonas chlororaphis)

MHMEEHTLNQMSDELKLMLGRFRHEQFVEKLGWRLPAHPSQAGCEFTWDQYDTEHARYLLAFNEDRAIVG CARLIPTTFPNLLEGVFGHTCAGAPPKHPAIWEMTRFTFTTREPQLAMPLFWRSLKTASLAGADAIVGIV NSTMERYYKINGVHYERLGPVTVHQNEFTKILAIKLSAHREHHRSAVAPSAFMSDTLLRETA

PsyI (Pseudomonas syringae)

MSSGFEFQLASYTTMPVTLLETLYSMRKKIFSDRLEWKVRVSHAFEFDEYDNAATTYLVGSWNGVPLAGL RLINTCDPYMLEGPFRSFFDCPAPKNAAMAESSRFFVDTARARSLGILHAPLTEMLLFSMHNHAALSGLQ SIITVVSKAMARIVRKSGWEHHVLSTGEASPGETVLLLEMPVTADNHQRLLGNIALRQPVTDDLLRWPIA LGVSGSAPQACMHSAA

RhlI (Pseudomonas aeruginosa)

MIELLSESLEGLSAAMIAELGRYRHQVFIEKLGWDVVSTSRVRDQEFDQFDHPQTRYIVAMGRQGICGCA RLLPTTDAYLLKEVFAYLCSETPPSDPSVWELSRYAASAADDPQLAMKIFWSSLQCAWYLGASSVVAVTT TAMERYFVRNGVILQRLGPPQKVKGETLVAISFPAYQERGLEMLLRYHPEWLQGVPLSMAV

Soli (Rhodobacter sphaeroides)

MRTFVHGGGRLPEGIDAALAHYRHQVFVGRLGWQLPMADGTFERDFTQYDRDDTVYVVARDEGGTICGCA RLLPTTRPYLLKDVFASLLMHGMPPPESPEVWELSRFTFAARSGAPCPRSGRADWAVRPMLASVVQCAAQ RGARRLIGATFVSMVRLFRRIGVRAHRFTAGPVRCIGGRPVVACWIDIDASTCAALGIPSASAAPGPVLQ

SwrI (Serratia liquefaciens)

MIELFDVDYNLLPDNRSKELFSLRKKTFKDRLDWLVNCENNMEFDEYDNRHATYIFGTYQNHVICSLRFI ETKYPNMISDGVFDTYFNDIKLPDGNYVEASRLFIDKARIQALQLHQAPISAMLFLSMINYARNCGYEGI YAIISHPMRIIFQRSGWHISVVKTGCSEKNKNIYLIYMPIDDANRNRLLARINQHATKMG

Tral (Agrobacterium tumefaciens)

VanI (Vibrio anguillarum)

MTISIYSHTFQSVPQADYVSLLKLRYKVFSQRLQWELKTNRGMETDEYDVPEAHYLYAKEEQGHLVGCWR ILPTTSRYMLKDTFSELLGVQQAPKAKEIYELSRFAVDKDHSAQLGGVSNVTLQMFQSLYHHAQQYHINA YVTVTSASVEKLIKRMGIPCERLGDKKVHLLGSTRSVALHIPMNEAYRASVNA

YenI (Yersinia enterocolytica)

MLKLFNVNFNNMPERKLDEIFSLRKITFKDRLDWKVTCIDGKESDQYDDENTNYILGTIDDTIVCSVRFI DMKYPTMITGPFAPYFSDVSLPIDGFIESSRFFVEKALARDMVGNNSSLSTILFLAMVNYARDRGHKGIL TVVSRGMFILLKRSGWNITVLNQGESEKNEVIYLLHLGIDNDSQQQLINKILRVHQVEPKTLETWPIIVP GIIK

YepI (Yersinia pestis)

MLEIFDVRXDEXTDIRSEDLYKLRKKTFKDRLNWEVNCSNGMEFDEYDNSDTRYLLGIYQGQLICSVRFI ELHLPNMITHTXNALFDXXALPKRGYIESSRFFVDKTRAKLLFGNHYPISYLFFLSIINYSRHNGYTGIY TIVSRAMLTILKRSGWQVEVIKXAHITEKERIYLLHLPIDRDNQARLXLQVNQRLQDPCSVLSTWPISLP VMPESA

YpeI (Yersinia pestis)

MLKVFNVNFDRMSENKLDEIFTLRKITFKDRLDWKVTCIDGKESDQYDDENTNYLLGTIDDTLVCSVRFV EMQYPTMITGPFAPYFRDLDLPIDGFIESSRFFVEKALARDKLGNNGSLSAILFLSMVNYARNRGYKGIL TVVSRGMYTILKRSGWGITVINQGESEKNEVIYLLHLSIDSNSQQQLIRKIQRVHNIDTHTLASWPLVVP SMTK

YpsI (Yersinia pseudotuberculosis)

MLKVFNVNFDRMSENKLDEIFTLRKITFKDRLDWKVTCIDGKESDQYDDENTNYLLGTIDDTLVCSVRFV EMQYPTMITGPFAPYFRDLDLPIDGFIESSRFFVEKALARDKLGNNGSLSAILFLSMVNYARNCGYKGIL TVVSRGMYTILKRSGWGITVINQGESEKNEVIYLLHLSIDSNSQQQLIRKIQRVHNIDTHTLESWPLVVP SMTK

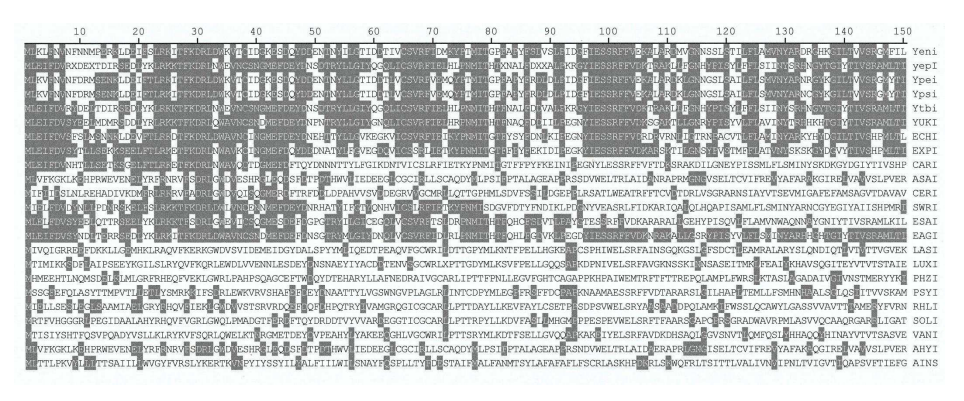
YtbI (Yersinia pseudotuberculosis)

MLEIFDVRYDELTDIRSEDLYKLRKKTFKDRLNWEVNCSNGMEFDEYDNSDTRYLLGIYQGQLICSVRFI ELHLPNMITHTFNALFDDVALPKRGYIESSRFFVDKTRAKLLFGNHYPISYLFFLSIINYSRHNGYTGIY TIVSRAMLTILKRSGWQVEVIKEAHITEKERIYLLHLPIDRDNQARLLLQVNQRLQDPCSVLSTWPISLP VMPESA

YukI (Yersinia ruckeri)

MLEIFDVSYEELMDMRSDDLYRLRKKTFKDRLQWAVNCSNDMEFDEYDNPNTRYLLGIYGNQLICSVRFI ELHRPNMITHTFNAQFDDIILPEGNYIESSRFFVDKSGAKTLLGNRYPISYVLFLAVINYTRHHKHTGIY TIVSRAMLTILKRSGWQFDVIKEAFVSEKERIYLLRLPVDKHNQALLASQVNQVLQGSDSALLAWPISLP VIPELV

Lux I-Type AHL Synthase Sequence Alignment



160 170 180 190 200 210 220	
LKRSGWNITV <mark>ln</mark> ggesekneviyllhigidnosogolinkilrvhovepktletweiivpgiik.	Yeni
LKRSGW <mark>QVE</mark> VIK <mark>XAHITEK</mark> ERIYLLHLPID <mark>RUNGARUXL</mark> OVNGRLODPCSVUSTWEISLPVMPESA.	yepI
LKRSGMGUTVI <mark>n</mark> ggesb <mark>knev</mark> iyllhisidsnsocqlirkiqrvhnidthtl <mark>as</mark> me <mark>lvvpsmtk.</mark>	Ypei
LKRSGAG <mark>etvin</mark> ogesekneviyllhlsidsnsoscier <mark>kior</mark> vh <mark>nidtht</mark> leswelvvpsmtk.	Ypsi
LKRSGM <mark>QVE</mark> VIK <mark>EAHITEK</mark> ERIYLIHLPIDR <mark>INGARULLGV</mark> NG <mark>RLODPC</mark> SVUSTWEISLPVMPESA.	Ytbi
LKRSGW <mark>QFD</mark> VIK <mark>EAFVSEK</mark> ERIYLIR <mark>IJEVOKHN</mark> QALI <mark>ASOVNQVL</mark> OGSDSAILAWEISLPVIPELV.	YUKI
LKRSGWRUSIIQQQLSEKQERIYELHIPTODOSRHAFIERITQMTQAESEQFKTIELLVPLA	ECHI
LKRSGMKU <mark>SIVE</mark> OGMSDKHERVYLLIFILEVONESODVIVKRINHNQEFVESKUREWELSFEPMTEPVG	EXPI
MLTILKRSGWGIRVVEQFTGLSEKEERVYLVFLPVDDEN EALARRINR GTFMSNELKQWPLRVPAAIAQA.	CARI
IF RLC PIERLGHRQAVDLGAVRGVGIRFHLDERFARAVGHPMQGEYADARELVTE.	ASAI
IDPVMDRVLKRSGNAPQFTGYVGTPKPMGKVTALAA MDCSEERVKRIRDFAGIYHDVTQPQTVIA.	CERI
IFQRSGWHISVVKTGCSEKNKNIY IYMPIDDANRNRLLARINQHATKMG	SWRI
TRSGWQIKVIKEAFLTEKERIYL TLPAGQDEKSCLGGDVVSRTGCPPVAVTTWPLTLPV	ESAI
AKRSGWETEVIKEGEVSENEPIYLIRIPIDCHNCHLIAKRIRDQSESNIAAICQWEMSLTVTPEQV	EAGI
MMIRAC DVSRFGPHLKIGIERAVALRIELNAKT LALYGGVL EURLAVS	LASI
RFLKRIKVPCHRIEDKEIHVLGDTKSVVLSMPINEQFKKAVLN.	LUXI
NGVHYERLGPVTVHQN FTKILAIKLSAHREHHRSAVAPSAFMSDTLLRETA.	PHZI
ARIVRKSGWEHHVLSTGEASPGETVL LEMPVTADNHQRLLGNIALRQPVTDDLLRWPIALGVSGSAPQACMHSAA	PSYI
GVILQR GPPQKVKGETLVAISFPAYQERGLEMLLRYHPEWL GVPLSMAV	RHLI
FVSMVRIFRRIGVRAHRFTAGPVRCIGGRPVVACWIDIDASTCAALGIPSASAAPGPVLQ.	SOLI
KLIKRMG PCERICDKKVHLLGSTRSVALHIPMNEAYRASVNA	VANI
IFRIC DIERLGHRQAVDLGAVRGVGIRFHLDERFARAVGQPLQGAYDEARELVTE.	AHYI
PFAPLFFLNAFLFVILTSINFFK	AINS

Appendix 5: Construction of LuxR-Type AHL Response Regulator Consensus Sequences

LuxR-Type AHL Response Regulator Consensus Sequence

LuxR Consensus Sequence

Protein Sequences of Published LuxR-Type AHL Response Regulators

AcuR (Aeromonas veronii)

MKQEQLFEYLEHFTSVTDGDRLAGLIGRFTVDMGYDYYGFTLIIPMSMQRPKVVLFNQCPISWVQTYTDN NMLACDPVIQLARKQTLPIYWNRLDERARFLQEGSMDVMGLAAEFGLRNGISFPLHGAAGENGILSFITS ERASSDLLLESSPILSWMANYIFEAAIRVVRLRDSDQQAALTDRETECLFWASEGKTSGEIACILGITER TVNYHLNQVTRKTGSMNRYQAIAKGISSGILLPNLEQVVVTNFPKLMQ

AhyR (Aeromonas hydrophyla)

 $\label{thm:local} $$ MKQDQLLEYLEHFTSVTDGDRLAELIGRFTLGMGYDYYRFALIIPMSMQRPKVVLFNQCPDSWVQAYTAN$$ HMLACDPIIQLARKQTLPIYWNRLDERARFLQEGSLDVMGLAAEFGLRNGISFPLHGAAGENGILSFITA$$ ERASSDLLLESSPILSWMSNYIFEAAIRIVRVSLREDDPQEALTDRETECLFWASEGKTSGEIACILGIT$$ ERTVNYHLNQVTRKTGSMNRYQAIAKGVSSGILLPNLEQVVVTNFPKLMQ$$$

AsaR (Aeromonas salmonicida)

MKQDQLLEYLEHFTSVTDGDRLAELIGRFTLGMGYDYYRFALIIPMSMQRPKVVLFNQCPDSWVQAYTAN HMLACDPIIQLARKQTLPIYWNRLDERARFLQEGSLDVMGLAAEFGLRNGISFPLHGAAGENGILSFITA ERASSDLLLESSPILSWMSNYIFEAAIRIVRVSLREDDPQEALTDRETECLFWASEGKTSGEIACILGIT ERTVNYHLNQVTRKTGSMNRYQAIAKGVSSGILLPNLEQVVVTNFPKLMQ

BsmR (Burkholderia pseudomallei)

MRAAMGNWAEDLLAGLDSARSEEEAFRSVETAAAALDFEYCAYGLRVPWPLSRPRIETRSNFPEQWKRRY VEAGFLDVDPILAHGRRSQQPVVLAETLFASAHQMWVEAQSFGLRFGWAQSSFDAYGGMGMLALVRSCEP VTAAELDAKEYRMRWLVRTAHAALGRMMLPKLMADPERGLTEREVEVLKWAADGKTSGEISKILAISVDT VNFHVKNAILKLRTANKTAAVVRAAMLGLLS

BviR (Burkholderia cepacia)

MQAWREKYLNGFATAKSEADVFLEFSADVRALGFDHCSFGLRIPLPISKPQFMLQSNYPQTWVERYVSQN YFAVDPTVRHGLSRMSPLIWRADSQTQCVQFWEEADQHGLRHGWCMPSVSRTGAIGLITMVRSGEPIEER

ELAEKGYQMSWLANTANYAMSMHLLQRLVPEYTVELTVREREALQWSAAGKTYAEIGKIMHVDDRTVKFH LVNAMRKLNAANKTEAAVKATMLGLLF

CarR (Erwinia carotovora)

MDHEIHSFIKRKLKGVGDVWFSYFMMSKNSTSQPYIISNYPEAWMKEYIKKEMFLSDPIIVASLARITPF SWDDNDIVTLRAKNQDVFISSVQHDISSGYTFVLHDHDNNVATLSIANHLEDANFEKCMKNHENDLQMLL VNVHEKVMAYQRAINDQDNPPDNSRNALLSPRETEVLFLVSSGRTYKEVSRILGISEVTVKFHINNSVRK LDVINSRHAITKALELNLFHSPCEPVVMKHMDAR

CerR (Rhodobacter sphaeroides)

MDIIDLSTVATDDASFLDYIDQLCQKLGFDYASYATTSPMTGAVQFTGYANYPDSWKMHYMRRNLHRVDP TIHKSALSIAPVDWSRFERDERFRAVFFAAEDFGITFTPQGLTVPVRGPYGDRGLLSVTRNCARPEWEKH KRAVIGELQVAAVHLHDAVMRSDVISRFTALRQPRLSTREIEVLQWAAAGKSQTDIGDILGISHRTVEVH LRSAREKLGTLSTVQAVGFTRAIGLGLVYPR

CinR (Rhizobium leguminosarum)

MQENEHSTTYPDAFSAMKNAATVAAALDEFQSHYPIDFVTFHLARTIVDNVDAPFVRTTYPDSWVSRYLL NDYINVDPIIREGFSRQLPFDWREIDITETAQEFMVDAELHGIGTNGVSVPIVDKSRRSLLSINSQKSDE EWTLIERQFLPEWLELGFLLHRKAVFELHGENDPVPALGSREIECLHWASRGKDSKDIGKILGLSEHTTR GYLKSARYKLGCPTLSAAIAHAVHLNLITPHVGTPS

EchR (Erwinia chrysanthemi)

MSISFSNFDFINSTIQNYLNRKLKSYGDLKYAYLIMNKKKPTDVVIISNYPSEWVEIYRSNNYQHIDPVI LTAINKISPFSWDDDLVISSKLKFSRIFNLSKEYDIVNGYTFVLHDPGNNLATLSFMFEENRSGELEEIV QNNKEKLQMLLISAHEKLTSLYREMSKNKNNSKSQEPNIFSQRENEILYWASMGKTYQEIALILGITTST VKFHIGNVVKKLGVLNAKHAIRLGVEMNLIKPVEPVKARS

EsaR (Pantoea stewartii)

MFSFFLENQTITDTLQTYIQRKLSPLGSPDYAYTVVSKKNPSNVLIISSYPDEWIRLYRANNFQLTDPVI LTAFKRTSPFAWDENITLMSDLRFTKIFSLSKQYNIVNGFTYVLHDHMNNLALLSVIIKGNDQTALEQRL AAEQGTMQMLLIDFNEQMYRLAGTEGERAPALNQSADKTIFSSRENEVLYWASMGKTYAEIAAITGISVS TVKFHIKNVVVKLGVSNARQAIRLGVELDLIRPAASAAR

ExpR (Pectobacterium carotovorum)

MSQLFYNNETISRIIKSQFDMALSHYGDIKYAYMVLNKKKPTEILIISNHHDEWREIYQANNYQHIDPVV IAALNKITPFPWDEDLLVSTQLKMSKIFNLSREHNITNGYTFVLHDHSNNLVMLSIMIDESNVSNIDDVI ESNKDKLQMTLMTIHAETISLYREMIRNKEDERSNDKDIFSQRENEILYWASMGKTYQEIALILDIKTGT VKFHIGNVVKKLGVLNAKHAIRLGIELQLIRPVQS

Lask (Pseudomonas aeriginosa)

MALVDGFLELERSSGLEWSAILQKMASDLGFSKILFGLLPKDSQDYENAFIVGNYPAAWREHYDRAGYAR VDPTVSHCTQSVLPIFWEPSIYQTRKQHEFFEEASAAGLVYGLTMPLHGARGELGALSLSVEAENRAEAN RFMESVLPTLWMLKDYALQSGAGLAFEHPVSKPVVLTSREKEVLQWCAIGKTSWEISVICNCSEANVNFH MGNIRRKFGVTSRRVAAIMAVNLGLITL

LuxR (Vibrio fischeri)

MKNINADDTYRIINKIKACRSNNDINQCLSDMTKMVHCEYYLLAIIYPHSMVKSDISILDNYPKKWRQYY DDANLIKYDPIVDYSNSNHSPINWNIFENNAVNKKSPNVIKEAKSSGLITGFSFPIHTANNGFGMLSFAH SEKDNYIDSLFLHACMNIPLIVPSLVDNYRKINIANNKSNNDLTKREKECLAWACEGKSSWDISKILGCS KRTVTFHLTNAQMKLNTTNRCQSISKAILTGAIDCPYFKS

MalR (Burkholderia pseudomallei)

MHDFLQFWLNEFSRSENPQHVISVLTRAAATLGYEYAAYGMRRPFPISNPPILMVSNYPARWQERYIEAR FANIDGAVKAALGSDRPVTWSAPANASKSAFWAEALSFGIAHGWSSASRGADGAIGVLTLSRTQDPIDTA EKFRNESIVHWLANVAHASMAPFLPAADEFDPDLTRRETDVLKWTADGKTAYEIALILSISESTVNFHVK NIVSKLGSTNKIQAVAKAALMGML

OccR (Agrobacterium tumefaciens)

MNLRQVEAFRAVMLTGQMTAAAELMLVTQPAISRLIKDFEQATKLQLFERRGNHIIPTQEAKTLWKEVDR AFVGLNHIGNLAADIGRQAAGTLRIAAMPALANGLLPRFLAQFIRDRPNLQVSLMGLPSSMVMEAVASGR ADIGYADGPQERQGFLIETRSLPAVVAVPMGHRLAGLDRVTPQDLAGERIIKQETGTLFAMRVEVAIGGI QRRPSIEVSLSHTALSLVREGAGIAIIDPAAAIEFTDRIVLRPFSIFIDAGFLEVRSAIGAPSTIVDRFT TEFWRFHDDLMKQNGLME

PhzR (Pseudomonas chlororaphis)

MELGQQLGWDAYFYSIFARTMDMQEFTAVALRALRELRFDFFRYGFTMCSVTPFMRPRTYMYGNYPEDWV QRYQAANYAVIDPTVKHSKVSSSPILASNELFRGCPFTDLWSEANDSNLRHGLAQPSFNTQGRVGVLSLA RKDNPISLQEFEALKVVTKAFAAAVHEFTKISELESDVRVFNTDVEFSGRECDVLRWTADGKTSEEIGVI MGVCTDTVNYHHRNIQRKFTIGASNRVQASRYAVAMGYI

ProR (Serratia proteamaculans)

MDTHLQPLMDALLTSQPDRKVFLSQLAPCAQALGFEYFSYTVFSCYPASRPKMLIEGNFPECYLEDYRKL RVYLQDPVIEQAHHSTLQFYWDEHFYQDKPELWWRMAQFGIREGWSQSVKDCYGRLGILTFAGKSIPVQS PQASARNETFFLWLAQMVHKTLREALISVNDEAIKDVLTLREKDILRWCSEGKTSEETALLMGLSERTVN FHIGNSIKKLSVANKTAATAKAVYLQLI

PsyR (Pseudomonas syringae)

MEVRTVKAQLDCPPLKINGAPAPLRQLIEDFENDLHHIGDFTYAYFSTPKTRNVKPVILSNYPDSWLKSY VASNYHLIDPIIKHAWHSITPFFWREAECCSGRRTDDFLKRSAKYQLSSGATFTLHDASWLFAALSLCNA RQQNDFDQRIREKAADIQMSLIRFHDRLIKTRAPHELFPQPAQCKLSTRETGVLKWVAMGKSYSEIAEIF SISERTVKFHMSNVSSKLKVRTAKQAVYKAINMGMV

RaiR (Rhizobium etli)

MSPSHAEQFSFFLLSGPDLRIADIAGSGNDAGRSRPHLCDIAYGSPCDLAGATDSNPLLMLTYPPEWVKQ YRDRDYFSIDPVVRLGRRGFLPVEWSASGWDSGRAYGFFKEAMAFGVGRQGVTLPVRGPQGERSLFTVTS NHPDAYWRQFRMDSMRDLQFLAHHLHDRAMVLSGMRKVADLPRLSRRELQCLEMTANGLLAKQICARLSI SVSAVQLYLASARRKLTVATTSEQLLGPRRSN

RhiR (Rhizobium leguminosarum)

MKEESSAVSNLVFDFLSESASAKSKDDVLLLFGKISQYFGFSYFAISGIPSPIERIDSYFVLGNWSVGWF DRYRENNYVHADPIVHLSKTCDHAFVWSEALRDQKLDRQSRRVMDEAREFKLIDGFSVPLHTAAGFQSIV SFGAEKVELSTCDRSALYLMAAYAHSLLRAQIGNDASRKIQALPMITTREREIIHWCAAGKTAIEIATIL GRSHRTIQNVILNIQRKLNVVNTPQMIAESFRLRIIR

RhlR (Pseudomonas aeriginosa)

MRNDGGFLLWWDGLRSEMQPIHDSQGVFAVLEKEVRRLGFDYYAYGVRHTIPFTRPKTEVHGTYPKAWLE RYQMQNYGAVDPAILNGLRSSEMVVWSDSLFDQSRMLWNEARDWGLCVGATLPIRAPNNLLSVLSVARDQ QNISSFEREEIRLRLRCMIELLTQKLTDLEHPMLMSNPVCLSHREREILQWTADGKSSGEIAIILSISES TVNFHHKNIQKKFDAPNKTLAAAYAAALGLI

SmaR (Erwinia billingiae)

MSNSFFNNTSINISIKNYLEKNLKVFNNIKYAYAIMNKKNPNDFAIISNRMEWFDFYTKNNLQFIDPVLI TASCCFTPFLWDENIMISSGLKMPKIFNMAKNYDVINGYTFVLHDHNHNLVVLSIIMDKSCDDDIEKIIV DKKNDLQMLLLTTHEKLITLYQEINDTHQFNKKNQKEILSKRENEILYWASMGKSYQEIALILGIKLTTV KYHVGNAVKKLGVTNAKHAIRLGVELKLIRPILPDAE

SolR (Rhodobacter sphaeroides)

MEPDFQDAYHAFRTAEDEHQLFREIAAIARQLGFDYCCYGARMPLFTPVSKPAVAIFDTYPAGWMQHYQA SGFLDIDPTVRAGASSSDLIVWPVSIRDDAARLWSDFTARDAGLNIGVARSSWTAHGAFGLLTLARHADP LTAAELGQLSIATHWLANLAHTLMSPFFTLVPQLVPESNAVLTTREREVLCWTGEGKTAYEIGQILRISE RTVNFHVNNVLLKLAATNFTKVQAVVKAIATGLI

VanR (Vibrio angullarium)

MYKILRLIQENQQITSHDDLENVLNGLNNLIGHEFFLFGLSFQPTLKTSETLVTDNYPNSWRQQYDESGF MHIDPIVKYSITNFLPIRWDDAKRVNNDGRVIFEEARCNGLKAGFSIPIHGLRGEFGMISFATSDTKSYD LNQQSIHTSQLIVPLLAHNIGNITRYHKDAKPRAVLTAREVQCLAWAAEGKSAWEIATIINTSERTVKFH FSNACKKLGATNRYQAITKAILGGYINPYL

VsmR (Pseudomonas aeruginosa)

MRNDGGFLLWWHGLRCEMQPIHDSQGVFAVLEKEVRRLGFDYYAYGVRHTIPFTRPKTEVHGTYPKAWLE RYQMQNYGAVDPAILNGLRSSEMVVWSDRLFDQSRMLWNEARDWGLCVGATLPIRAPNNLLSVLSVARDQ QNISSFEREEIRLRLRCMIELLTQKLTDLEHPMLMSNPVCLSHREREILQWTADGKSSGEIAIILSISES TVNFHHKNIQKKFDAPNKTLAAAYAAALGLI

YenR (Yersinia enterocolytica)

MIIDYFDNESINEDIKNYIQRRIKTYGDLCYSYLVMNKKTPLHPTIISNYPLDWVKKYKKNSYHLIDPVI LTAKDKVAPFAWDDNSVINKKSTDSAVFKLAREYNIVNGYTFVLHDNSNNMATLNISNGSDDSISFDERI EINKEKIQMLLIITHEKMLGLYQSNSDKNENRNTQIERDIFSPRENEILYWASVGKTYAEISIILGIKRS TVKFHIGNVVRKLGVLNAKHAIRLGIELKLIKPI

YpeR (Yersinia pestis)

MIINFFDNESINEDIKNYIQRRIKAYGNIRYSYLLMNKKVPLHPAIISNYPLDWVKKYKKNSYHLIDPVI LTAKGKVAPFAWDDNSVINIKSTDSAVFNLAREYNIVNGYTFVLHDNNNNMATLNVSSGDDDSIFFDESI EVNKEKIQMLLIFIHDKMLGLYNKSHHENNTLNKKENKREIFSPRENEILYWASVGKTYSEIAIILGIKK STVKFHIGNIVRKLGVLNAKHAIRLGIELQLIKPI

YpsR (Yersinia pseudotuberculosis)

MIINFFDNESINEDIKNYIQRRIKAYGNIRYSYLLMNKKVPLHPAIISNYPLDWVKKYKKNSYHLIDPVI LTAKGKVAPFAWDDNSVINIKSTDSAVFNLAREYNIVNGYTFVLHDNNNNMATLNVSSGDDDSIFFDESI EVNKEKIQMLLIFIHDKMLGLYNKSHHENNTLNKKENKREIFSPRENEILYWASVGKTYSEIAIILGIKK STVKFHIGNIVRKLGVLNAKHAIRLGIELQLIKPI

YspR (Yersinia pestis)

MHSVFNRSNEVIETLRDYIDRKLTIYDSPKYTYMVINKKNPGDIFIVTSYPNEWAELYTNNNYQNIDPVV LIAFRRFSPFSWDENITVLSELKLSKIFTLSKKYNIVNGFTFVLHDTMNNLAMLSLIMDDSALNGVESRV LNDRDRLQMLLIETHEKMLTLSQRNMNIQERQGKGMPGKAILSPRENEVLYWASMGKTYQEIAIITNITP RTVKYHIGNVVKKLGVINAKQAIGLGVELEIIKPILA

YtbR (Yersinia pseudotuberculosis)

MHSVFNRSNEVIETLRDYIDRKLTIYDSPKYTYMVINKKNPGDIFIVTSYPNEWAELYTNNNYQNIDPVV LIAFRRFSPFSWDENITVLSELKLSKIFTLSKKYNIVNGFTFVLHDTMNNLAMLSLIMDDSALNGVESRV LNDRDRLQMLLIETHEKMLTLSQRNMNIQERQGKGMPGKAILSPRENEVLYWASMGKTYQEIAIITNITP RTVKYHIGNVVKKLGVINAKQAIGLGVELEIIKPILA

YukR (Yersinia ruckeri)

RKLERYDSPRYTYMVIDKKNPVDVFIVTSYPDEWADIYTSQNYQHIDPIVLTAFKRISPFAWDENITILS DLKSSKIFALSKKYNIVNGFTFVLHDHMNNLAMLSLIMDNNADKGLNSRIESDKDRLQMNLIKIHEKMLM LEQNKLGVSNGKNTDTSGKGILSPRENEVLHWASMGKTYPEIALIAGITTRTVKHHMGNVVKKLGVINAR QAIRLGVELELIKPVLV

Lux R-Type AHL Response Regulator Sequence Alignment

	10	20	30	40	50	60	70	80	90	100	110	120	130	140	150
									VI MA <mark>KGKVA</mark> PE						- m
									VI. MAKGKVA						
	HSV	IRSNEVT TIR		DSBK TVMVTI	AREAN P	CDTERVTS	A BILDWAKKSK	MNNIVONIEDE	V <mark>i</mark> lta <mark>kdkva</mark> pe VVl <mark>i</mark> a <mark>f</mark> rrespe	APPUNSVINK SEPERATUVI S	FI ISA	TETTI SKKVNITA	NEGYTEVUM Neg y men	NSN-NMATER	NISNGS YenR
									VVLIA F RRESPE						
									IVLWA <mark>fk</mark> rises						
									IIVASLARIT						
									V <mark>i</mark> lta <mark>inki</mark> see						
									VV <mark>IAALNKIT</mark>						
									VLITA <mark>SCOFT</mark> EF TV <mark>SHCTQSVL</mark> EI						
MRNDGG	FLLWWHGLF	CEMOPIH SO	GVFAVI EKEV	/RR GFD AY	GVRHTIPI	TREKTEVHGT	KAWLE O	MONYGAVIDE	AI NGL SSEMV	Vos RL	OSRM	LW E DW LC	V- A LPIR	APNIN-LI SVIIS	Vario VsmR
	MQHWLDKL1	LAATEG EC	ILKTG ADIA	ADHFGFTG AY	LHIQHR-	HITAVT	HRQ QSTYF	DKKFEALDE	WKRARSKHIE	TASGEHERPT	LSKD-ERA	TYDHASDFO R	S-CITIPIK	ran <mark>g-fm</mark> smft	MASOK TraR
									VVRLGREGELEV						
									IIREGFSRQLPH						
MYKI	LRL	QENQQITSHD	DLENVINGLI	NNLIGHEFFLF	GLSFQPTI	KTSETLVTD	NS RQQ D	ESGFMH D	IV <mark>KYSITN</mark> I.FI IIQL∕RKQTI :I	R AKRV	NNDGRV	EEA CNOLK	A- FSIPI (GLRG-EFG I	FTS VanR
MKNTN-	DQLLEII	WKIKACESNN TEULISAI GD	DINOCUSDM	PKMVHCEVVI.I.	ALTINHSN	WKS TS T.D.	CEDSEA ÓUEL	A HMLAC	I DYSNSNH I	NENT FENNAN	NK SDN	MG ABECLK Treeksselt	T- IS PINC	PAG-ENGIDO	FITAL ASAR
									LNPENFSQGHL						
MRAA	AMGNWAEDLI	AGLDSARSEE	EAFRSVET A	AAAD DFEYCAYO	GLRVWPI	SRERIETRSN	F EQUKR YV	EAGFLDVD	ILAHGR SQQ V	VLAETL A	SAHQI	WVE-QSFCLR	F-CWAQSSF	YG-GMGMTA	VRSC BsmR
-MELGÇ	QLGWDAY 1	SIFARTM MQ	EFTAVA R	LRE RFDFFRY	GFTMCSVTPI	MR RTYMYG	Y FEDWQR Q	AA WAV UD	TVKHSKVSS	LASNELDF	GCPFTD	LWSE NDSNLR	H-LAQPSF1	NTQG-RVGV	RKU PhzR

	160	170	180	190	200	210	220	230	240	250	260	270	280	
131 D	DSIFFDESIEVXK	EKIOMBLILF D	KMLGLY <mark>NKS</mark>	HHENNTLNK	ENKRETFOR	ENEILYWAS	/GKTY <mark>S</mark> ELALL	LGI <mark>KKS</mark> TVK	FHION <mark>IVR</mark> KL	VLNAK <mark>H</mark> AI	RIGIEIQIII	BI		YpsR
131 D	DS <mark>IFFDES</mark> IE <mark>V</mark> NR	ek <mark>i</mark> cvelefied	KMLGLY <mark>NKS</mark>	HHENNTLAK	ENKRE F. G	ENEILYWAS	/GKTY <mark>S</mark> EIAII	LGI <mark>KKS</mark> TVK	FHIGN <mark>I</mark> V <mark>R</mark> KL	e <mark></mark> vinak <mark>h</mark> ai	rugi niqini	PI.		YpeR
	DS <mark>ISE</mark> DER JEINK													YenR
131 8	ALNGV SRVLNDR	DR <mark>LQMLLIETHE</mark> I	KWI TI SCRN	MNIQERQG	<mark>mpg</mark> kailspr	ene <mark>v</mark> lywasi	/GKTYQETATI	INTPRIVE	<mark>Y</mark> HIGNVVKKL	VINAKQAI	GLGWELEIN	ILA.		YtbR
	ALNGV SRVLNDR													YepR
111 N	ADKGLNSRIESDA	DRLOM <mark>NUJKI</mark> HE	KMIMUEONK	LGVSNGK	TSGK <mark>G</mark> LLSPR	ene <mark>v</mark> lhwasi	GKTYPETALI	ACHTTRIVK	<mark>H</mark> :M=NVVKKL	VINA <mark>RQ</mark> AI	RIGVELELII	ALA		YukR
121 E	AN- KCMKNHE	NDEOMET VNV HE	VMAYQRAI	NDQDNPPI	NSRNAL SER	TOV FLVS	S RIYK VSRI	LGISE <mark>V</mark> TVK	FIINNSVRELI	DVINSRHAI	TKAL I N FI	ISPCE VVMKH	IMDAR	CarR
131 N	RSGELSEIVQNAK	EKLOMILI <mark>SA</mark> NE	KLTS LYREM	ISKNKN S	SQEPNIFSQR	ENEILYWASI	igktyqeta <mark>l</mark> t	LCITTSTVK	FHIGNVVKKL/	3vlnak <mark>h</mark> ai	PALGWEMNIAU	VEPVK	KARS .	EchR
131 S	NVSNIDDVIESNK	DKLOMTLMTI A	ETIS REM	IIRNKEDEI	rsndkdifsór	ENEILYWASI	4GK <u>T</u> YQETA <mark>L</mark> I	LDI <mark>KTG</mark> TVK	EHIGNVVKKL	:vlnak <mark>h</mark> ai	RIGIELQUII	R VQS		ExpR
130	CDDDI KI VDKK	ndlowli <mark>lt</mark> ime	KLITLYÇEI	NDTHQFN	KNOKETTSKE	ENEILYWASI	/GK <mark>S</mark> YQETA <mark>L</mark> I	LGI <mark>KLT</mark> IVK	YH <mark>V</mark> GN <mark>A</mark> VKKI.	3––v <mark>t</mark> nakhai	RIGVELKIJI	R ILPDA	Œ	SmaR
	NRAEANRFM::SVL											CL		LasR
141 Q	NISSE REEIRLR	LRCMUFLLT	QKI TDLEH-	PMLI	IS-NPVCLSHR	PRE LOWTAI	osk <mark>ssg</mark> etatt	LSLSESTVN	HKNIQKKFI	DAPNKTL <i>AF</i>	AYAAAUGUI			VsmR
135 P	VID-LDRE DAVA	AAATIGO HARI	SFURTTP		AEDA CVD K	ATY RWIA	/SKIME IIIADV	E VKYNS R	VKLRERMARFI	DVRSKAHLI	'ALAIRRKLI			TraR
143 P	AYWRQFRMDSMR	-Du∢FuAHHL#DI	RAMV SG	MR	VADLPRESRE	LQC EMTAN	Nellako car	ISISVSAVQ	LYLASARR	ΓVATTSEQI	LGPRRSN			RaiR
	EE-WT IERQFL												S	CinR
	KSYDINQQ <mark>S</mark> IH													VanR
142 R	ASSDLL ESSPIL	SWMSNYIFDAAI	RIVRVSLR-	EI	DD-QEA TOR	TDCLFWASE	ECKT <mark>SC</mark> ETA <mark>C</mark> I	LGII <mark>TE</mark> RIIVN	Y LNQVTRKT	SMNRYQ	AKG SSCIL	LPNLEQVVVTN	IFPKLMQ.	AsaR
143 K	NYIDS FLHACM	NIPHIVPSLV	DNYRKIN	IA1	INKSNNDETKE	KECLAWACE	ECKSSWDISKI	Leckrivi	LTNAQMKUI	NTTNRCQS	SKAILTCAU	C YFKS		LuxR
141 R	REIPILSDELQL	MQ VRESLI	MA MRLND-	EIVI	IT-SEMNFSKR	K R TA	eskt <mark>sa</mark> ela <mark>m</mark> i	LSIBENTVN	D QKNMQKKII	NAPNKTQVA	CYAAAT			SdiA
139 E	PVTAA DAKEYR	MRWTVRTAH	AAL RMML-	PKLI	1AD ERGITER	V V KWAI	XEKTSGETSKI	ALSVDIVN	VKNAILKU	RTANKTA/N	VRAAM OLL	3		BsmR
144 N	PISLO FEALKVV	TAFAAAVHEFT	ISE ESD-	VRVI	N-TDVEFSGR	CDV R TAI	OGKT SEFT GV	MCVCTD	Y HR IQRKF	rigas rvo s	RYA AMCY II	į.		PhzR

Appendix 6: Construction of LuxM-Type AHL Synthase Consensus Sequences

LuxM-Type AHL Synthase Consensus Sequence

LuxM Consensus Sequence

MLSLLSXXXXXXPVQDSCPTLVASALIQNWSVRDTWLSFTYAPQXXNYCFPSYGYSEFTRLQLFTPSSLSKCYX XEFDNEFKXQLSDTQAVCEVFTLRLTVXXXXYFLYLAQKELMSVLHQAGYKXXXXXXXIIEQPFMLNFYRAIDAK AYFHSFTGYCDLNDDGKQTYRGFWNFEMMVKAFSNIDFRGYKRXRASRKRGSLERDEHV

Protein Sequences of Published LuxM-Type AHL Synthases

LuxM (V. harveyi)

 ${\tt MLSLLSLSQVGKHFIVLKHPVQDSCPTLVASALIQNWSVRDTWLSFTYAPQFSNEQWNYCFPSYGYSEFTRLQLFTPSSLSKCYSLPEFDNEFKLQLSDTQAVCEVFTLRLTVSGNAQQKLYFLYLAQKELMSVLHQAGYKIGFTIIEQPFMLNFYRAIDAKAYFHSFTGYCDLNDDGKQTYRGFWNFEMMVKAFSNIDFRGYKRAVRASRKRGSLERDEHV$

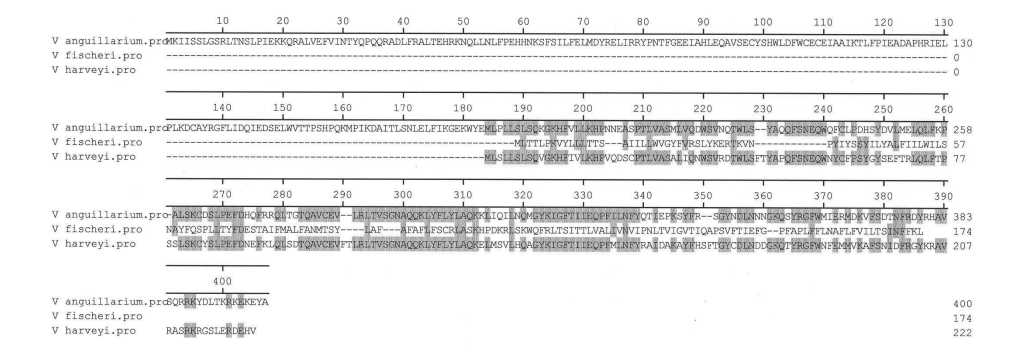
VanM (V. anguillarium)

MKIISSLGSRLTNSLPIEKKQRALVEFVINTYQPQQRADLFRALTEHRKNQLLNLFPEHHNKSFSILFELMDYRE LIRRYPNTFGEEIAHLEQAVSECYSHWLDFWCECEIAAIKTLFPIEADAPHRIELPLKDCAYRGFLIDQIEDSEL WVTTPSHPQKMPIKDAITLSNLELFIKGEKWYEMLPLLSLSQKGKHFVLLKHPNNEASPTLVASMLVQDWSVNQT WLSYAQQFSNEQWQFCLPDHSYDVLMELQLFKPALSKCDSLPEFDHQFRRQLTGTQAVCEVLRLTVSGNAQQKLY FLYLAQKKLIQILNQMGYKIGFTIIEQPFILNFYQTIEPKSYFRSGYNDLNNNGKQSYRGFWMIERMDKVFSDTN FRDYRHAVSQRRKYDLTKRKEKEYA

AinS (V.fischeri)

MLTTLPKVYLLLTTSAIILLWVGYFVRSLYKERTKVNPYIYSSYILYALFIILWILSNAYFQSPLLTYFDESTAI FMALFANMTSYLAFAFAFLFSCRLASKHPDKRLSKWQFRLTSITTLVALIVNVIPNLTVIGVTIQAPSVFTIEFG PFAPLFFLNAFLFVILTSINFFKL

LuxM-Type AHL Synthase Sequence Alignment



Appendix 7: Construction of LuxN-Type AHL Response Regulator Consensus Sequences

LuxN-Type AHL Response Regulator Consensus Sequence

Protein Sequences of Published LuxN-Type AHL Response Regulators

LuxN (P. phosphorem)

MPDLPLLIFSEPRGALLFFAAGIILAWLGYFSFTLFTSRPGANRNVYYPYLAYSVSIFLWILSNAYFQSP
LLTYYSESTAVTMALFANLVSFCAFISAYSFSCRLISTQPDSNLSLYQKLFISIISLYALIINSSPGLTV
KHVDIVAPGDFVIIFGPQTSWFFLCLMSAVFLTFHNFLIYKKAGSPLIQKKSQYMILGVIIFMLSTLIVH
LIIPFMLDDFSLTWVPPALAIFETLLIGYALLFNRFYSPRYIISQFISHLVNVTLYLSPYLLIIAIGYED
NPLLIGLWIALIGLGWKSSLIQIKRGTNRLLYGKNGSPSENIQRVIGHFQYSTEYGLGKLNELLNTRSGQ
ILNINTHSDLAALKIYFEGKHSVLVKDELEFQIQYETHTELSNISWLKKNMDANNSALVLPIVSKNGDIS
HLFMVSKKDRDGLFSSEEIDALQVLFEQANQYIRSEEQVRKSQVLAGSIAHEIRNPLSKIQYHFERIDAD
LFDVNNNSAHPFLSEQMKGLYKELTESKKAVQLGTRFIDIIIDEIKGNSINSQTFSSHSAGRLTEQALSE
YGFVGNTYQARIIANTQNDFQFWGNETLFSFVMFNLVKNALHYFSQYPQSTLSIHLERGESENCIIVTDT
GPGIADNVIPHIFDEFYTLGKSDGSGLGLAYCRRVINAFGGNIHCQSKYGSYTRFTLTFPIINEERIPNN
LFNELKEALTGKQVLVIGHKENTTLISSLLSGFNIIVSTVDNGKSAAKYIGNNNVDFAFYDLSLSPTQFE
ALKKIRSGDFGANAQKIPLIALSNENTRSTRFDTNVFQGEFRISDSLPLFAQSLKLLIDSGSLKPLGHLI
GKRVLVVDDMQINRMLVQSYLAQEGITVLQAHNGSVALCIAEQERPDLILMDIHMPEMDGLEVSRILRQR
GYNIPIIALSGECCNEVTKEISQYMNAYLMKPITRQQLIQKLQYWIPESEADKVISKQDIHIVHSI

LuxN (P. profundem)

MHDFIQSTLANMVAIFLVAIALVVVIWATYFARILAKHLPGSSRQVYFPYTLYSVFISAWILSNAYFQSD
LLVYFGADTAIIMALLANIFSGLAFAYAFLFSCRLVSERTSFQLKTWQWILFSLTCIIILVTNCVPGLNV
KSVDIEGIGSFVIHFGPTIGVFFGNLLLLILTLGNFILSSRSQLKLKQIKANYMIFGMMAFIISTFFAH
FLIPIFLNDFSKAWLPPALSIIEVIIVGYALLHHRFYSIRYIGLITLSFVINAAIYIIPIASVGFVGTQD
STLLLVIWTLITGICWYKSLAIIRRSVNRLLYKEKGDPVENICNLIGEFSYSTDQAVIKLNQVLNAKSGR
IQKVSGNTENNIFVSYFHGNRSVLIKEEIEYQLKHEKPEGTKELSNVTREMVNMGVSLVLPITNERNEVT
QLYMVSKEKENVLFSSEEIMGLQLLFDKANCFIVTEDKIRKSQVLVGTIAHEIRNPLTKIKYHFERIDAD
MFGIENTSLSPFASKEMKKIYQELSEGQKAVQLGSRFIDAILDELRGESIGTTLFDNYSVAKLTHQALND
FCFNSEEHKLRINIDTQSDFFFHGSDTLYSFVLFNLIKNAVYYFDTYPNSQIRIYFQKERNYNKVHVVDT
GPGISPDHQKHILEEFYTNGKVQGNGLGLSYCKRVIESFGGTITCQSELGEYTEFILSFPSIDEKIHSEM
SKEKIKSYLTGMSGLVLGSVEVGNWLSSEFKSLGVELCTAPDVKTGLHHLSQQAVDFIIMDHMLLNREMG
SIKMLRAGTHGHQAQTTPMFLYGYTENSEHLNSIELSPFFQGQIDGINDHQAFLHSLESLIDNDLFAKLG
SLIGKTVLVVDDMQVNRMLVQAYLASEGITVVQASSGDEAIEKVKKEPFNLVLMDIQMPGMSGIEATHQI
RHLFDAIPIVALSGEYNEEITRAISETMNDHLVKPINKQQLLOTLTKWMT

LuxN (V. algiolyticus)

MLDVHLHGLFYPKAMALYATVLIVFAWLLYYCYRLKQKSESILGSHHAPYIAYSSCIIVWISSNAYFHTD LLPELGSVGGIFMAKLANLASFFAFAFAFYFSCQLTADVKKTAVKVWQKVVFVALATYSLYINLVPNLTV ENVTISGPSQFVIEFGPHTSYFFISLLAFVVLTLLNLIAMRANSSKLTLAKSNYMIAGILVFMLSTAVIH LGMTYFLGDFSLTWLPPALSISEMLFVGYALLTSRFYSAKYLTYLTVSALLVCAIFVLPLGAIFIPISED NQWLVAVPLCALIGITWHLLFKRVSRYASYFIYGKRHTPVQQILGLEEEFKRSIDDAMRQLASLLNIPNN KLQLVTSNYTETFYEEYLPSSKSVLVLDELSEEIDYASSSKGSMRKLYERMRSSNTALVMPLFGRGKSVT HLLISSHKIDNKLFSNEEISALQTLLVRIQSTIEADRKVRQSRALANSIAHEMRNPLAQVQLQFEALKQH IESNASLDTLKREIDKGEAAIQRGRQLIDIILREVSDSSPEHEPLALTSIHKAIDQAVSRYGFENDQIIE RINLPQAHDFVAKLNETLFNFVIFNLIRNAIYYFDSYPDSQIEIRTQTGAYENILIFRDSGPGIDSSILH KIFDEFFSYQKSGGSGLGLGYCQRVMRSFGGRIECQSELNEFTEFYLYFPVVPNAPKPETLRAPDFDSWK ATASHSENHSAQHVQVCTDAPTVLIVDDKEVQRTLVQMYLKRLGVNSLQANNGASAVELFHSHKIDLVLM DVQMPVMNGFDASQRIKQITSSVPIIALSGESGARELELISKLMDDRLEKPTTLNALQVVIQRWLQNENF APSNTF

Van N (V. anguillarium)

MLNLNLDPILYPKAITLIAAVAMVLVWLTYYCYRLKQKNEVIFGTHHASYIAYSSCIIAWIGSNAYFHTD WLVELGVNRAIFMAEIANISASLAFVFAYYFSCQLSAEQRKGKVHLWQRLIFITIAAYSVLINLQSNLTV KHVDIVGPSEFVIEFGPHTPYFFNAMLCSVILTLFNLVVMRTNSSKLALAKTNYIIAGILVYMLSTLVIH IGITFFFQDFSLSWLPPALSISEMMFVGYALITSRFYSVKYLAYLCLNTALVCGVLFIPLGAIFIPLTDS NQWLIAIPLCALIGITWNPLYKRLSRYASLLIYGNQQTPVEQILALEDDFKRSIDDAMRRLGQLLYIADD KLQFVNSNYNETVYERYLSSKQTALVFDELFEKLDNKTAAKNSIKALYDKMSSNNTALVMPLFGHSKLVT HLLISPHKINNQMFSNEEIAALQTLLTRIQSIIEADRRVCQSRALANSIAHEMRNPLAQVQLHFEILKQH IDSQAPAQQIKQDIENGQAAIQRGRQLIDIILREVSDSSPEHEPITMTSIHKAVDQAVSQYGFENEKVIE RIHLPQQDDFVAKLNETLFNFVIFNLIRNAIYYFDSYPNSQIEITTQIGTYENILIFRDTGPGIDDAISY KIFDDFFSYQKSGGSGLGLGYCQRVMRSFGGRVECKSKLGEFTEFHLYFPMVPNAPQADSLRTPDFKSWQ QPKPNTEQRTVDNIQPIDKPFLINNKAPTVLIVDDKEVQRSLVQMYLNQLGVNNLQANNGENAVEIFKAN SIDLILMDIQMPVMNGFEASQIIKAHSPQVPIIALSGESGERELEMISKLMDGRLEKPTSLNALQQVISH WLNKDIVPNAHTAKSGTVI

LuxN (V. angustum)

MADLYQAVTTNVIAIFLIAISAVIAVWTGYFARFLHSKPSLSHDKRIYFPYIIYTSFISLWILSNAYFQS
SLLIERSDIVAVNIALAANIFSGLAFIFAYLFSCRITSKKDNFSLTFTQKFLLYTSIIITLLTNIIPRIN
ITSIDIKAIGVFYINFGELSFIFFGMLIIILLSTIINLLILHKNNTCINRVKAKYMITGIIAFISSTFLI
HFIAAVIFHDFTAAWLPPALSVIEVFLIGYALFNSRFYSLKYIIFITSSTFINIIFYTAPVILLELYHIK
ETPFFLVLWTLITGFFWHRTLRLVRLFANKIIYHKKGNPVENITKIISEFKISTDLGISKLNTVIHSNNG
IIVQVSNKNQLLRDYFKTGRNILLKQDLDVLLNDNVLADNHLHLVSEQLHKMGVTLVVPILDESKKITHF
YIASKEMSNVLFSCEEIMGLQRLFERANRFIDTEEKVRKSQVLAGSIAHEIRNPLSKIKYHFEKIDSDFL
SVHKESINSLATLEIEKIHQELTEGKKALQLGTKFSDVILDELRGSSISTSFFQHYSAASLTSQALNDFS
LYSEEHKKRIHLEATNNFYFYGSDTLFSFVLFNLLKNAVYYFDTFPESHISIQFEKGLKHNKIHVRDTGP
GITEEQLENLFDEFYSFGKVSGNGLGLAYCKKVMESFSGSISCHSILGEFTEFTLTFPAINIQSNGELTN
PRIKQHLSGQSCLILSASSLSKKLTESFNGLNMNIECSNDPSIGFTRIKDCPFNFIVIDHRLYITHYDQI
SMLREGKYGYLAQITPIFIFNSTSINLNNDRINVPKYTQGYIDTLNGALAFECSLEAIINDTKFAPLGSL
NDKTVLVVDDMHANRLLVKAYLSKEGINVIQAASGYEAIEQVKKNNIDLIFMDIHMPGMNGIETAKQLKE
LDSTKPIIAISGEYGEKIVSDIHKVMDDYIVKPIEKSTLVSLTSKWLIINKVKD

LuxN (V. fischeri)

MLTTLSKVYLLLTTSAIILLWVGYFVRSLYKERTKVNPYIYSSYIFYALFIILWILSNAYFQSPLLTYFD ESAAIFMALFANMTSYLAFAFAFLFSCRLASKHPDKRLSKWQFGLTSITTFAALIVNVIPNLTVIGVTIQ APSVFTIEFGPFAPLFFLNAFLFVILTSINFFKLRKSNIKLNKEKSIYLMVGIFIYMISTIASQIIIPVI WADFSYTWVPPALSVTEALLIGYTLLYHRLYSFKYLLFWSLSYSINLILYLIPIIIIYDLTTPSDLLYIC IIEIIFTGLFWDKTLKKTKKIASIIIYKDKQTPVEKIYKIAEEFKYSSSNAIIKLASILNTPKEELLLIG KNTNYNIFIPHLNQSHSALVKDELDYQIHYSPKTANAELHQVQEKMSESKTALILPIFGENKLISHFLIS ANKHDNTTFSNEEISAIQWVLTKVQGYIESERKVRQSQALANSIAHEMRNPLSQLQYHFEKIKHHYQKNT EHEKQEQLIKNELNQGCLAIQKGAQLIDIILSEAKNTAISDDLFHHHSISLLTQQIIDEYVFDSEEMKQK ITLDLEDDFIVNINDTLYGFILFNLLRNATYYFDEYNSSISIRLVKGFATNKLIFRDTGPGIDSHILPNI FDDFFTHNKEGGSGLGLSYCLRVMHAFEGNIACYSTKGEFTEFVLSFPHIEGDINALNSHKSNTPPLINK KDNSLKTVLIVDDKKVQRMLIHTFINKDNLTLLQAENGEEAVEIATNNKLDLIFMDSRMPVMNGIDAAKK IKIIYPNLPIIALTGESSHEEISAITQVMDGYLTKPVSKAQLQQVVDKWL

LuxN (V. harveyi)

MFDFSLEAIVYAKAISLLATVAVVMMWLFYYCYRLKQKNEVIFGTHHAAYIAYSVCIIAWISSNAYFHTD LLPELGASAGMFMAKFANLASFFAFAFAYYFSCQLAAEQRKGKVHRWQQGIFVSLTVYSLFINLRPGLTV EHVDIVGPSQFIIEFGPHTSYFFIGLVSFVVLTLVNLVAMRTNSSKLTLAKTNYMIAGILVFMLSTAVIH LGMTYFMGDFSLTWLPPALSISEMLFVGYALLTSRFYSVKYIAYLALSVLLVCAIFVLPLGAIFIPLTES NQWLIAIPICALIGITWQLLYKKTSRYASFLIYGDKKTPVQQILSLEEDFKLSIDDAMRRLGKLLQIPND KLRLVTSNYNETFYEEYLSSNRSVLVFDELSEELEYKVSAKRSMKALYDKMSSNNTALVMPLFGQGKSVT HLLISPHKSNNQMFSNEEISAVQTLLTRVQSTIEADRRIRQSRALANSIAHEMRNPLAQVQLQFEALKQH IENHAPVEQITLDIENGQAAIQRGRQLIDIILREVSDSSPEHEPIAMTSIHKAVDQAVSHYGFENEKIIE RIRLPQHTDFVAKLNETLFNFVIFNLIRNAIYYFDSYPDSQIEISTKTGPYENTLIFRDTGPGIDETISH KIFDDFFSYQKSGGSGLGLGYCQRVMRSFGGRIECKSKLGTFTEFHLYFPVVPNAPKADTLRTPYFNDWK QNKRSNEHKVAPNVQINNQSPTVLIVDDKEVQRALVQMYLNQLGVNSLQANNGENAVEVFKANHVDLILM DVQMPVMNGFDASQRIKELSPQTPIVALSGESGERELDMINKLMDGRLEKPTTLNALRHVLGNWLNKNTA SSACEAERE

LuxN (V. paraheamolyticus)

MLDIGLSGLLYPKAITLFATVAVVLVWLLYYCYRLKQKNEVILGSYHAPYIAYSTCIIIWISSNAYFHTD LLPLLGSEGGIFMAKLANLASFFAFAFAFYFSCQLAAEQKKGKVKLWQQGIFVALTVYSLVINLRPNLTV ENVLIDGPSQFVIEFGPHTSYFFMGLVTFVVMTLTNLISMRANSSKLSIAKNNYMIAGILVFMLSTAVIH LGMTYFLGDFSLTWLPPALSISEMLFVGYALLTSRFYSAKYLAYLTISVLFVCTIFVLPLGAVFIPMSED NQWLISIPICALIGITWHLVYKRVSRVASFFIYGNRQTPVQQILALEEEFKRSIDDAVHQLSTLLNIPND KLQLVTSNYTETFYEDYLHSNDSVLVLDELSERLDEKPSSKGSIKALYERMRSSNTALVMPLFGREKSVS HLLISSHKSDNKLFSNEEISALQTLLIRVQNTIESDRKIRQSRALANSIAHEMRNPLAQVQLQFEALKQH IDSNASDDKIRSDIEKGQAAIQRGRQLIDIILREVSDTSAVHEPLSLTSIHKAVDLAVSRYGFENEHIIE RVKLPTQNDFVAKINETLFNFVIFNLIRNAIYYFDSYPDSQIEIRTLVGPYENTLVFRDTGPGIDDSILH KIFDDFFSFQKSGGSGLGLGYCQRVMRSFGGRIECKSVTNEFTEFYLHFPVVPNAPKVETLRTPNFYNWN QKVKTKPSPEPVVQINKDAPTVLIVDDKEVQRTLVQMYLNRLGVNSLQANNGANAVELFQSHQVDLVLMD VQMPVMNGFDASEKIKQCSPTTPIIALSGESGEKELEMIAKLMDGRLEKPTTLNALRDVLVRWLHFDKIS VTNSYQIANE

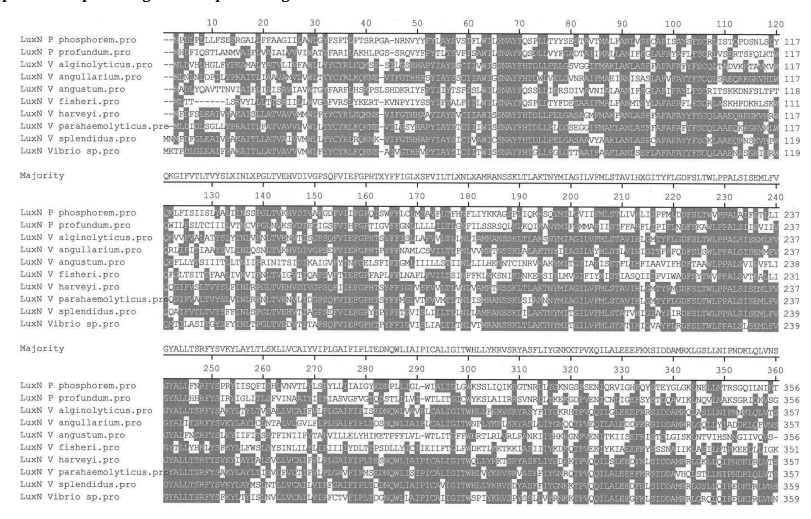
LuxN (V. splendidtus)

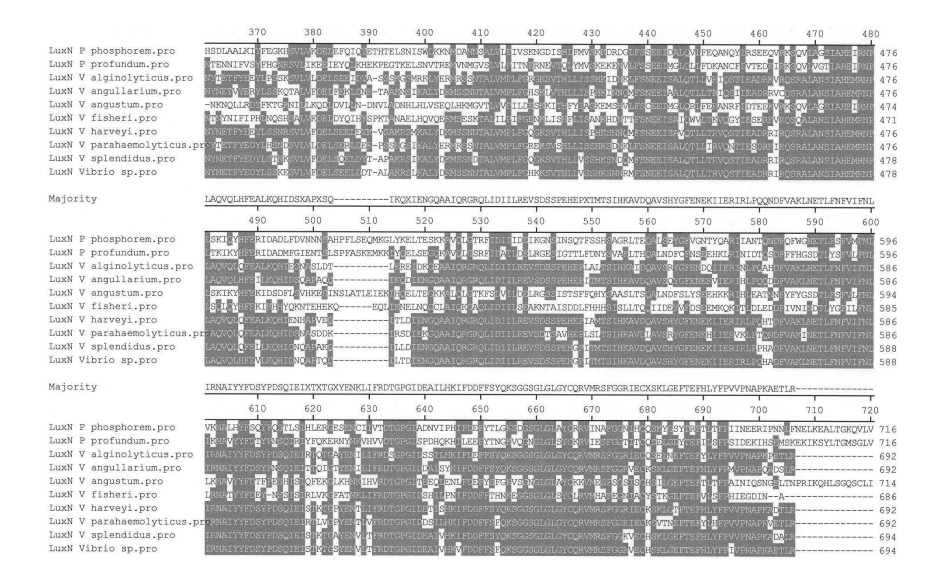
MNMFDFGLEAIVYAKAITLLATVAIVVMWLLYYCYRLRQKNKVIFGTHHAPYIAYSICIVAWICSNAYFH
TDLLPELGASAAVYAAKLANLASFCAFAFAYYFSCKLAAEQRNSKVHPWQQAIFVTLTVYSFFINLSPGL
TVEHVTIAGPSEFVIEFGPYTPYFFTGVISLIILTLLNLLAMRANSSKLILAKTNYMITGILVFMLSTAT
VHIGIAYFIRDFSLTWLPPALSISEMLFVGYALLTSRFYSVKYLAYMSLNTLLVCAILVIPFGAIFIPLT
DDNQWLIAIPICAVIGITWHLLYKRVSDYASFFIYGNKKTPVQQILALEEDFKLSIDDAMRRLGSLLQIP
EDKLRLVNSNYNETFYEDYLSTNKSVLVFDELSQELDYTAPAKRSIKALYDKMSSNDTALVMPLFGQGKS
VTHLLVSSHKSNDQMFSNEEISALQTLLTRVQSTIEADRRIRQSRALANSIAHEMRNPLAQVQLQFELLK
QHIDNQAPAKQILLDIENGQAAIQRGRQLIDIILREVSDSSPEHGPITMTSIHKAVDQAVSHYGFENEKI
IERIRLPPHADFVAKLNETLFNFVIFNLIRNAIYYFDSYPDSQIEISTKTGAYENVLTFRDTGPGIDEAI
VHKIFDDFFSYQKSGGSGLGLGYCQRVMRSFGGKVECHSKLGEFTEFHLYFPVVPNAPKADALRTPYFND
WKSNQAATENKTNVDAKPDNQAATQNSEPTSTLTPGNHLAPTVLIVDDKEVQRTLVQMYLSRLGVNSLQA
KNGENAVELFKTHKVDLILMDVQMPIMNGFDASQIIKARSPQTPIIALSGESGQHELDMISKLMDGRLEK
PTSLKALQHVLDNWLEKGWASNTSKETESEE

LuxN (Vibrio sp.)

MKTFDLGLEAIFYAKAITLLATVAVVVMWLFYYCYRLKQKNEAIVGTHHVPYIAYSICIITWISSNAYFH
TGLLPGLGTTAAIFAAKLANLSSFLAFAFAYYFSCQLAAENRSGKIHRWQKTILASITGYSFYINLTPGL
TVEDVTITAPSQFVIEFGPHTPYFFIGVISLIALTLTNLVTMRANSSKLTLAKTNYMITGILVFMLSTAT
IHIGVAYFLRDFSLTWLPPALSLSEMLFVGYALLTSRFYSFKYLTYISLNVLLVCAILVIPFCTVFIPLT
DGNQWLLAIPICAIIGITWSPIYKRVSPYSSLLVYRNKKTPVQQILALEEGFKLSIDDAMRRLGRQLQIP
EDKLRLVNNNYNETFYEDYLSSKESVLVFDELSEELDDTALAKRSLKALYDKMSSNNTALVMPLFGHKKS
VTHLLVSSHKSNNRMFSNEEISALQTLLTRVQSTIEADRRIRQSRALANSIAHEMRNPLAQVQLHFEVLK
QHIDNQAPTQQILTDIENGQAAIQRGRQLIDIILREVSDSSPEHGPITMTSIHKAVDQAVSHYGFENEKI
IERIRLPQHADFVAKLNETLFNFVIFNLIRNAIYYFDSYPDSQIEISTKTGSYENVLTFRDTGPGIDEAI
VHKVFDDFFSFQKSGGSGLGLGYCQRVMRSFGGRVECHSKLGEFTEFHLYFPIVPNAPKAETLRTPYFNG
WKHNQSTEDKAEADVKPESQTPSGDIEPEPASTLTESKQTERTQAENQPASSHLAPTVLIVDDKEVQRTL
VQMYLSRLGVNSLQAKNGENAVELFRSHKVDLILMDVQMPIMNGFDASQIIKARSPQTPIIALSGESGQR
ELDMIRKLMDGRLEKPTSLNALQHLLDNWLEKGWAPNASKETENE

LuxN-Type AHL Response Regulator Sequence Alignment





	730	740	750	760	770	780	790	800	810	820	830	840
LuxN P phosphorem.pro LuxN P profundum.pro LuxN V alginolyticus.pro LuxN V angullarium.pro		KS <mark>LGVELC AF</mark> DSWK ATASHS E KSW <mark>Q PKPN</mark> IF	PDVKTGLHHLS N Q	SQQAVDFIIMI	OHMLLNREMGS HSAQH RUVDN	IKML AGTH QVCTD IQ IDK	IGHQAQTTPMFL PFL	YGYTENSEH INN	LNSIELSPFFÇ	QGQIDGINDH(AFLHSLESL	IDNDLF 836 719 725
LuxN V angustum.pro LuxN V fisheri.pro LuxN V harveyi.pro LuxN V parahaemolyticus.p LuxN V splendidus.pro	TPYF	NGLNM IECSN NSHKSNTPPLI NDW ONKRSN YNWN KVKUKE NDW SNQAA	N H	KDCPFNFIVII	KKDNS VAPN SPEPV	LK QINNQ QINKD	S					706 720
LuxN Vibrio sp.pro Majority	TPYF	ngwahnos-ive	D	ELQANNGENAN 1 880	KAEAD	VK ES <mark>OTPS</mark>	GDIEPEPASTL MNGFDASQIIK 910	TESK			QTEI	*
LuxN P phosphorem.pro LuxN P profundum.pro LuxN V alginolyticus.pro LuxN V angullarium.pro LuxN V angustum.pro LuxN V fisheri.pro LuxN V harveyi.pro LuxN V parahaemolyticus.p LuxN V splendidus.pro LuxN Vibrio sp.pro	KPLGHLIGKRVIVV AKLGSLIGKIVIVV	DDMQINEMIYC DDKEVORTIYC DDKEVORSIYO DDMANEIIYA DDKKVORMIIH DDKEVORAIYO DDKEVORIIYO DDKEVORIIYO	SYLAGE CITY A LASE CITY MYLK LGYNN TAYLSKE CITY TOTAL	ILOA HAC SVAT IVVOA SSEDEA ILOANNI AS ILOANNI SENA ILOA BOUTE ILOA	CIAEQERP I KVSKEPFN I SANSI I QVKKNNI I ATNNKI I ATNNKI V KANH V KANH V KANH V KANH	LIMPIHMER VIME OF COMPA VIME OF THE OF THE OF THE OF THE OF THE OF THE OF THE OF THE OF VIME OF THE	MOCLEVER LR MSCIEATHOUR MNGFDASCRIK MNGFEASCIK MNGFEASCIK MNGFEASCRIK MNGFEASCRIK MNGFEASCRIK MNGFEASCRIK MNGFEASCRIK	QRGYNI HLFDAI : IV QITSSV : I AHS PQV : I ELDSTK : I IIY : NI : I ELS : CI : IV QCS : I : I I I I ARS : CI : I I I I	ALSGESC <mark>CCNEVT</mark> ALSGESC <mark>A</mark> REI ALSGESCEREI A <mark>I</mark> SGE <mark>Y</mark> SE <mark>KIV</mark> A I GESSHEBI ALSGESCHREI ALSGESCHREI	CKE I SOYMAN CRA I SETMOD EL I SKIMI D PM I SKIMI D SON I HKVMI D SA I TOVMI D DM I NKIMI D D	I MA ITROO II VK INKOO LEKETINA LEKETISINA ITK IVSKAO ITK IVSKAO ITK ITNA	ICKIQ 953 LCTIT 956 VVIQ 832 COVIS 839 VSLTS 954 COVV 817 RHVIG 832 RDVIV 831 CHVII 852
LuxN P phosphorem.pro LuxN P profundum.pro LuxN V alginolyticus.pro LuxN V angullarium.pro LuxN V angustum.pro LuxN V fisheri.pro LuxN V harveyi.pro LuxN V parahaemolyticus.p LuxN V splendidus.pro LuxN V ibrio sp.pro	XWLXKXXASNX 970 Y01PESE DKVISK KMT RUQNENFAPSNTF HUNKDIVPAHTAI KULIINKVKD KM NKNTS-SACE NKNTS-SACE NKNTS-SACE NKNTS-SACE NKNTS-SACE NKNTS-SACE NKNTS-SACE	980 QDIHIVHSI KSGTVI AERE QIANE TESEE										976 960 846 859 964 820 849 850 871

Appendix 8: Protein Sequences of Published AHL Degrading Enzymes

AHL Lactonase Sequences

AhlD (Arthrobacter sp. IBN110)

MEKDQLKVRVLETGVMEADMAWLLLKPGRIIADRNNKERQREWGEIPTHAVLIEHPEGRILWDTGVPRDW SSRWQESGMDNYFPVKTESSSESGFLDSSLAQVGLEPADIDLLILSHLHLDHAGNARLFDNGKTKIVANR KELEGVQEIMGSHLGGHLKADFEGLKIDAIEGDTEIVPGVSVIDTPGHTWGTMSLQVDLPDDGTKIFTSD AVYLRDSFGPPAIGAAVVWNNLLWLESVEKLRRIQERTNAEMIFGHESEQTSQIRWAHQGHYQ

AhlK (Klebsiella pneumoniae 342)

MMPEIKLFMFQSGTQHCRYQHIRMNQGVGEHYEIPVPWFLLTHPDGFTLIDGGLAVEGLKDPSGYWGSTV EQFKPVMSEEQGCVEQLKRIGIAPEDIRYVVLSHLHSDHTGAIGRFPHATHVVQRQEYEYAFAPDWFTSG AYCRRDFDRPQLNWLFLNGLSDDHYDLYGDGTLQCIFTPGHSPGHQSFLIRLPGGTNFTLAIDAAYTLDH YHEKALPGLMTSATDVAQSVRKLRQLTERYHAVFIPGHDPEEWKKNRLAPACYY

AiiA (Bacillus sp. 240B1)

 $\label{thm:constitegklinlpvwcyllettegpilidtgmpesavdnedlfkgtfve} $$ GQILPKMKPDDRIVNILKRVGYAPEDLLCVISSHLHFDHAGGNGSFSHAPIIVQRTEHDAALHRAEYLKE $$ CILPDLNYQMIEGDYEVMPGVQLLYTPGHSPGHQSILVKTEKSGSVLLTIDASYTQENFEQGVPFAGFDS $$ EMASQSINRLKEIVLDEKPIVFFGHDMEQEKRCKTFPEFL $$ $$$

AiiB (Agrobacterium tumefaciens C58)

MGNKLFVLDLGEIRVDENFIIANSTFVTPQKPTVSSRLIDIPVSAYLIQCTDATVLYDTGCHPECMGTNG RWPAQSQLNAPYIGASECNLPERLRQLGLSPDDISTVVLSHLHNDHAGCVEYFGKSRLIAHEDEFATAVR YFATGDHSSPYIVKDIEAWLATPRNWDLVGRDERERELAPGVNLLNFGTGHASGMLGLAVRLEKQPGFLL VSDACYTATNYGPPARRAGVLHDTIGYDRTVSHIRQYAESRSLTVLFGHDREQFASLIKSTDGFYE

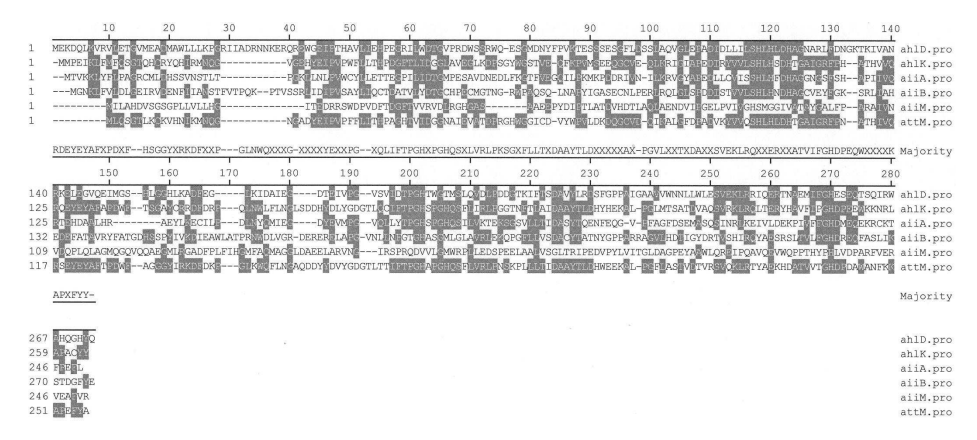
AiiM (Microbacterium testaceum LB037)

MILAHDVSGSGPLLVLLHGITEDRRSWDPVDFTDGFTVVRVDLRGHGASAAEEPYDIPTLATDVHDTLAQ LAENDVIPGELPVIVGHSMGGIVATAYGALFPARAIVNVDQPLQLAGMQGQVQQAEGMLRGADFPLFIHG MFAQMAGGLDAEELARVNGIRSPRQDVVLGMWRPLLEDSPEELAALVSGLTRIPEDVPYLVITGLDAGPE YAAWLQREIPQAVQEVWQPPTHYPHLVDPARFVERVEAFVR

AttM (Agrobacterium tumefaciens C58)

MLQSGTLKCKVHNIKMNQGNGADYEIPVPFFLITHPAGHTVIDGGNAIEVATDPRGHWGGICDVYWPVLD KDQGCVDQIKALGFDPADVKYVVQSHLHLDHTGAIGRFPNATHIVQRSEYEYAFTPDWFAGGGYIRKDFD KPGLKWQFLNGAQDDYYDVYGDGTLTTIFTPGHAPGHQSFLVRLPNSKPLLLTIDAAYTLDHWEEKALPG FLASTVDTVRSVQKLRTYAEKHDATVVTGHDPDAWANFKKAPEFYA

AHL Lactonase Sequence Alignment



AHL Acylase Sequences

Aac (Shewanella sp. MIB015)

MKFNKLMIAMGMACGVMLTGCNDSEDSTLPTEPETQLQTFAPNGLLKANIRRTTYGVPHIQADNLESLGF
GSGYAQAQDNLCVLADGFIKANSQRSMYFGPHASIDFTTGQPTAEDNGNLISDFAYKALKIREQAEAKWP
QFSERSRALIQGFTAGYNQYLADVEVGKQTAEPFCGGQPWVKPIVPEDVVTYLFSIALLPGAANFLDLIF
YANPGDGQEYMPRIVGPAPSQEQTAFVSDMQSKLLARAARITTPETNPRDLGSNGWGLGKDKTENGRGMV
LGNPHFPHTGNLRFWQSHITIPGQLDMMGGSLVGMPGPINIGFNKDLAWTHTFSTAEHFVMYNLELVSGD
RLQYLFDGQPMPINKETVSILVNAGPAGMLVAEKDIYTTAKGPMVEAPPSLAPFGWDDGSAFMIQDANMA
NMDPVDHWLAMNLATNKDEFQQAFKDYDGVIFNNTMFADKEGNAFYIDDSTVPGLSESAVVVLKTSPDIK
AAKQRAGFTILPGNTSLFSFSGPTPYERAPKLERSDFVQNSNDSFWSTNLNEPLTYFSPMYGPEAGQLSL
RTRMGLTLMQDAAGSDGKFNLEELEAAVLSNRSYLAELVLQDLIAQCEAQGSTPVVVSASLSKDLTSACA
ALKAWNGKQDNDSKGGALLREFAHQFSQKTMLTKGFDPANAATTPNTLTTDGSALVALAHAALNLEAAGF
ALDAPLGDVQFVEKSLPYGTASGARLPWPGSHNAEGGFNVFSTSLSGDDTLIPQHKYAPLMDVVTGKAMA
SGMTAKGYQVRYGSSWMMAVSFTDEGPVARGILTYSESSNILSPSFTDQSNLYSSSKSFRPLLFKEADIA
PAVISTTELTLQKAQ

Ahlm (Streptomyces sp. M664)

MRLRNRLRLLGVAGLALFTVSASLPPATASGAGQERHPSGGGLSAVIRYTEYGIPHIVAKDFAQLGFGTG
WAQAADQVCTLADGFLTVRGERSRFFGPDAATDFSLSSAATNLSSDLYFRGVRDSGTVEKLLKVPAPAGP
SRDVKETMRGFAAGYNAWIAQNRITDPACRGASWVRPVTALDVAARGYALAVLGGQGRGIDGITAAQPPT
AAPPAAGVTPEEAASAAKRLLSAQNADMGSNAVAFDGSTTVNGRGLLLGNPHYPWQGGRRFWQSQQTIPG
ELNVSGASLLGATTMSIGHNPDVAWSHTVATGVTLNLHQLTLDPADPTVYLVDGKPERMTKRTVSVPVKG
AADVTRTQWWTRYGPVVTSMGAALPLPWTATTAYALNDPNATNLRMADTGLGFSKARSTKDVERSLRRNQ
GMPWVNTIAADRAGHSFFAQSQVLPRITDDLAERCSTPLGRATYPASGLAVLDGSRKDCALGSDRDAVQP
GIFGPGRMPVLKNQPYVENSNDSAWLTNADRPLTGYERVFGTIATPRSMRTRGAIEDVASMADKGRLRVA
DLQRQQFANRAPAGELAASEVAKWCAALPGGTAVGTGGTPVDVSDACAVLRRWDRSVDSDSRGALLFDRF
WRKTSAVPAAELWKTPFDPADPVRTPRGLNTAAPGVGRALADAVAELRAAGIALDAPLGKHQFVVRNGKR
LPIGGGTESLGIWNKTEPVWNAAGGGYTEVSSGSSYIQAVGWDDSRCPVARTLLTYSQSENPRSRHFSDQ
TRLYAGERWVTSRFCEKDIARSPDLRVVRVHERR

AiiD (Ralstonia sp. XJ12B)

MMQGFALRGTLAMAALAALAGCASSTDGRWGSLSDTGLSAEIRRTGFGIPHIRANDYASLGYGMAYAYAQ DNLCLLADQVVTVNGERSKTFGPEGTVTVSFKPIPNLQSDAFFKGIFDEDGLRAGYAQMSPEARELLRGY IAGFNRYLKDTPPANFPAACRNAAWVRPLTLGDMMRMGEEKAIQASAGAMLAGIVAAQPPGRTPVAEREI PPQAVDTVALDRELQLRDMPIGSNGWAFGADATANRRGVLLGNPHFPWTTTNRFYQVHLTVPGKLDVMGA SIAAFPVVSIGFNKDVAWTHTVSTGRRFTLFELKLAEGDPTTYLVDGTPHKMTTRTVAFDVKLPDGRLER RTHTFYDTIYGPVLSMPSGGMPWTTQKAYALRDANRNNTRSVDSWLHIGQARDVAGIRQAIGNLGIPWVN TIATDRNGRALFADVSTTPDVPAAELQRCAPSPLAGKLFKDAGLVLLDGSRGTCNWQVDPASPVPGLVAP ARMPVLERDDYVANSNDSSWLTNPAQKLTGFSPVMGSVDVPQRLRTRIGLIEIGRRLAGTDGLPGNRIDL PNLQAMIFSNANLAGQLVLGDLLAACKATPAPDADVRDGCAALGQWNRTSNADARAAHLFREFWMRAKDI AQVHAVEFDPADPVHTPRGLRMNDATVRTAVFKALKEAVGAVRKAGFALDAPLGTVQAAHAPDGSIALHG GEEYEGVLNKLQTLPIGPKGLPVYFGTSYIQTVTFDDQGPVADAILTYGESTDHASPHAFDQMRAYSGKH WNRLPFSEAAIAADPALKVMRLSQ

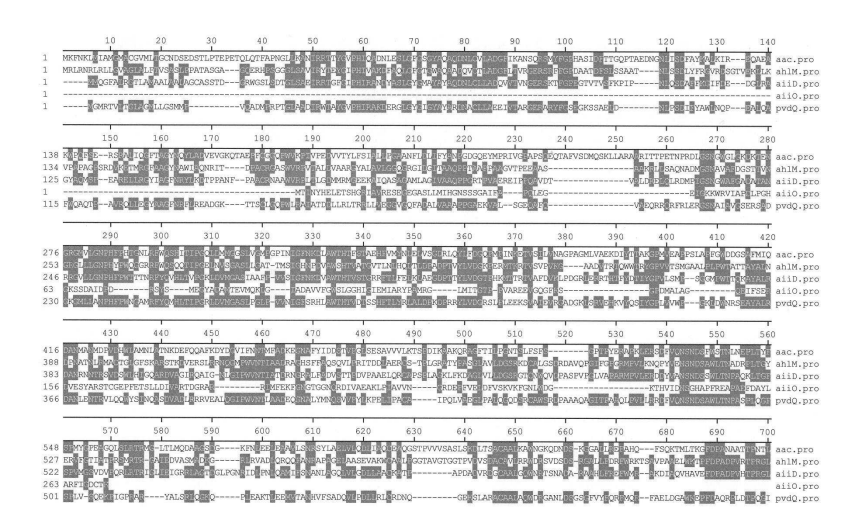
AiiO (Ochrobactrum sp. A44)

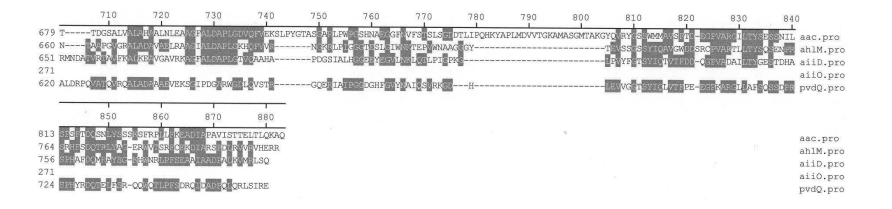
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PvdQ (Pseudomonas aeruginosa PA01)

MGMRTVLTGLAGMLLGSMMPVQADMPRPTGLAADIRWTAYGVPHIRAKDERGLGYGIGYAYARDNACLLA EEIVTARGERARYFGSEGKSSAELDNLPSDIFYAWLNQPEALQAFWQAQTPAVRQLLEGYAAGFNRFLRE ADGKTTSCLGQPWLRAIATDDLLRLTRRLLVEGGVGQFADALVAAAPPGAEKVALSGEQAFQVAEQRRQR FRLERGSNAIAVGSERSADGKGMLLANPHFPWNGAMRFYQMHLTIPGRLDVMGASLPGLPVVNIGFSRHL AWTHTVDTSSHFTLYRLALDPKDPRRYLVDGRSLPLEEKSVAIEVRGADGKLSRVEHKVYQSIYGPLVVW PGKLDWNRSEAYALRDANLENTRVLQQWYSINQASDVADLRRRVEALQGIPWVNTLAADEQGNALYMNQS VVPYLKPELIPACAIPQLVAEGLPALQGQDSRCAWSRDPAAAQAGITPAAQLPVLLRRDFVQNSNDSAWL TNPASPLQGFSPLVSQEKPIGPRARYALSRLQGKQPLEAKTLEEMVTANHVFSADQVLPDLLRLCRDNQG EKSLARACAALAQWDRGANLDSGSGFVYFQRFMQRFAELDGAWKEPFDAQRPLDTPQGIALDRPQVATQV RQALADAAAEVEKSGIPDGARWGDLQVSTRGQERIAIPGGDGHFGVYNAIQSVRKGDHLEVVGGTSYIQL VTFPEEGPKARGLLAFSQSSDPRSPHYRDQTELFSRQQWQTLPFSDRQIDADPQLQRLSIRE

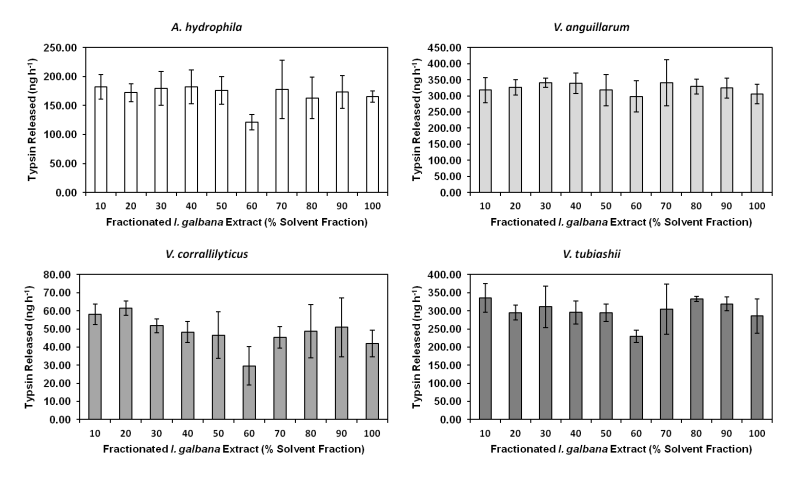
AHL Acylase Sequence Alignment



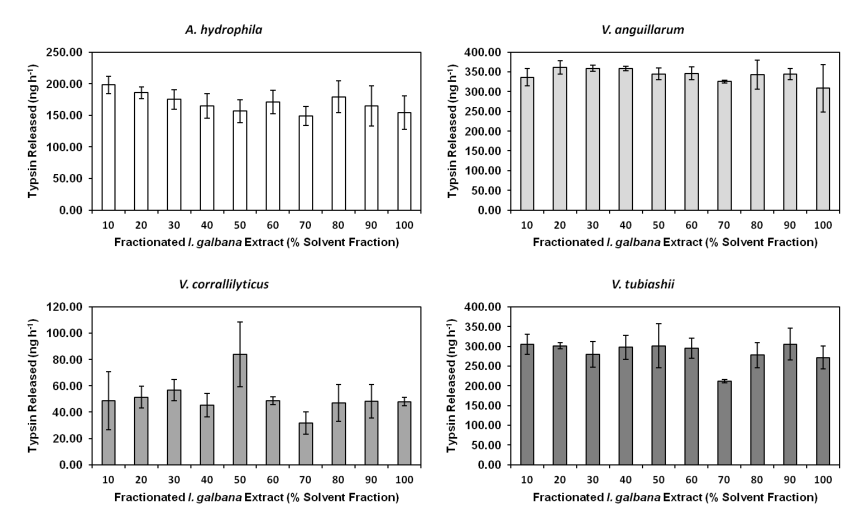


Appendix 9: Initial Microalgal Protease Assays

Protease activity of in four marine bacterial pathogens exposed to fractionated extracts from stationary phase I. glabana culture. Error bars represent standard deviation from the mean.



Protease activity of in four marine bacterial pathogens exposed to fractionated extracts from stationary phase N. oculata culture. Error bars represent standard deviation from the mean.



Protease activity of in four marine bacterial pathogens exposed to fractionated extracts from stationary phase T. suecica culture. Error bars represent standard deviation from the mean.

