ANALYSIS OF THE STABILITY OF A TYPE III SECRETION SYSTEM CONTAINING PATHOGENICITY ISLAND (VPI-3) IN THE HUMAN PATHOGEN VIBRIO CHOLERAE

by

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A thesis submitted to the Faculty of the University of Delaware in partial fulfillment of the requirements for the Honors Bachelor of Science Degree in Biological Sciences with a concentration in Cell and Molecular Biology & Genetics with Distinction

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by

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ABSTRACT

*Vibrio cholerae* is an enteric pathogen that is the causative agent of the secretory diarrhea, cholera, that affects millions of people each year. While the *V. cholerae* O1 serogroup pathovar strains are well studied due to their propensity to cause epidemic and pandemic cholera, a second pathovar has been identified that causes inflammatory diarrhea. Strains of this pathovar encode a Type III Secretion system (T3SS) that is present on a Pathogenicity Island (PAI), a mobile genetic element integrated into the chromosome of some non-O1 serogroup strains. Our hypothesis is that this PAI was horizontally acquired and was an essential acquisition in the emergence of this pathovar. The aims of this study were to investigate the genetic structure of T3SS island regions, examine the excision behavior of the region in strain NRT36S and reconstruct the evolutionary history of the region. To accomplish this, we first performed a bioinformatics analysis among a group of strains that contain a T3SS. We constructed a genetic deletion of cognate T3SS island integrase, *intV2*, and determined the excision phenotype of the island using a two stage nested PCR assay. We showed that *intV2* is necessary for the excision of the region. Lastly, we performed a phylogenetic analysis of the conserved T3SS ATPase protein and associated integrases to show unique clustering patterns, indicating that this island is mosaic in structure. Additionally, phylogenetic analysis revealed the presence of two different variants of T3SSs within *V. cholerae*, T3SS and T3SS. The grouping pattern on the tree showed a close relationship between the T3SS ATPase of *V. cholerae* NRT36S, *V. parahaemolyticus*, and *V. mimicus* suggesting that this virulence
system may have been passed horizontally among these different species of bacteria in the past.
Chapter 1

INTRODUCTION

1.1 Introduction to Vibrios

The genus *Vibrio* is a diverse group of aquatic bacteria in the Class Gamma-Proteobacteria. These bacteria thrive in marine, estuarine, or riverine environments and can be free swimming or survive due to commensal or pathogenic relationships with aquatic organisms from fish to phytoplankton (Romalde et al., 2013, Paillard et al., 2004). While most *Vibrio* species thrive in the mesophilic temperature range, some *Vibrio* species have also been found in extreme environments such as hydrothermal deep-sea vents (Colwell 2006). Vibrios have fermentative metabolic processes and it is thought that some environmental bacteria in this genus play an important role in the nutrient cycling and nitrogen fixation of the ecosystems in which they live (Colwell 2006). Although the importance of environmental vibrios is clear, the scientific community has typically focused on pathogenic species that pose a threat to human health. In 2012, incidence of infection for reported cases of *Vibrio*-related illness increased by 43% (Centers for Disease Control and Prevention, 2013). Comparatively, incidence rates for other foodborne pathogens such as *Listeria*, *Salmonella*, and *Shigella* did not increase during the same two-year time period, as shown in Figures 1 and 2 (Centers for Disease Control and Prevention, 2013). *Vibrio parahaemolyticus*, *Vibrio vulnificus*, and *Vibrio cholerae* have been extensively studied because they cause human illness.
Figure 1 Relative rates of laboratory-confirmed infections with *Campylobacter*, Shiga toxin-producing *Escherichia coli* (STEC) O157, *Listeria*, *Salmonella*, and *Vibrio* in the United States during 1996-2012, compared with 1996-1998 rates, by year. In comparison with 1996-1998, incidence of infection was significantly lower for *Campylobacter*, *Listeria*, *Shigella*, STEC O157, and *Yersinia*, whereas incidence of *Vibrio* infection was higher (Centers for Disease Control and Prevention, 2013).
Figure 2 Estimated percentage change in incidence of laboratory-confirmed bacterial and parasitic infections in the United States in 2012, compared with average annual incidence during 2006-2008, by pathogen. The estimated incidences of infection were higher in 2012 compared with 2006-2008 for Campylobacter (14% increase; 95% confidence interval: 7%-21%) and Vibrio (43% increase; 95% confidence interval: 16%-76%) and unchanged for other pathogens (Centers for Disease Control and Prevention, 2013).

1.2 Vibrio cholerae: Background and history

Sanskrit records dating from around 500 BC describe one of the first recorded cases of cholera. The origin of the name of the disease is thought to come from the ancient Greek words “chole” meaning bile and “rein” meaning flow (Colwell 2006). To follow the historical epidemiology of cholera is to follow the course of human history. It has been speculated that cholera, or a disease like it, affected the outcome of historically important battles such as Gallipoli, El Alamein, and the Oman war (Colwell 1996). The second cholera pandemic, which began in Russia in 1829, spread across the globe to New York City. The arrival of this pandemic to London in 1849
sparked the birth of the public health field of epidemiology (Colwell 1996). Modern sanitation technologies have reduced the number of global instances of cholera, but the seventh pandemic, which began in 1961 in Indonesia and spread to South America in 1991, is still occurring today (Colwell 1996). Developing areas with poor sanitation, crowded living conditions, and low hygiene standards, or countries experiencing natural disasters and turmoil, are more likely to report outbreaks or epidemics of cholera. In today’s global world, an epidemic can more easily become a pandemic. In 1996, Colwell et al. showed a correlation between the sea surface temperatures surrounding Bangladesh and the number of cases of cholera in the region. Thus, as the world’s oceans continue to warm due to climate change, cholera may become an even greater public health threat.

1.3 Vibrio cholerae: A Public Health Threat

Over the past 20 years, the incidence of most food and water borne bacterial infections has either decreased or remained constant. This fact does not hold true for Vibrio infections, which have steadily increased in incidence since 2000 (WHO, 2014). In 2013, 47 countries across the globe reported 129,064 cases of cholera to the WHO. 2,102 of these cases were fatal, equaling a case fatality rate of 1.63% (WHO, 2014). V. cholerae is often misreported or underreported; therefore, the global burden of cholera is likely much greater than is represented by the data available. An estimated 1.4-4.3 million cases of cholera occur each year, leading to up to 142,000 un-reported deaths (WHO, 2014).

Researchers are frequently isolating newly evolved strains of V. cholerae. V. cholerae is naturally competent (able to take up DNA) in the presence of chitin, a substance that constitutes the shells of many sea-dwelling crustaceans and bivalves
(Meibom et al., 2005). Thus, *V. cholerae* in their natural environment are able to take up foreign DNA and potentially integrate this DNA into their chromosomes (Meibom et al., 2005). Newly evolved strains pose a greater public health threat, especially in areas where cholera is endemic, such as India and Bangladesh, because the acquired immune response does not protect against novel strains. Some strains isolated from these regions also demonstrate multi-drug antibiotic resistance (Faruque et al. 2007). While antibiotics are usually used only in the most severe cases of cholera infection, resistance to drugs commonly employed in rural or impoverished communities could potentially increase the case fatality rate in these areas where cholera outbreaks are common (WHO, 2014).

As of 2013, there are two vaccinations used to mitigate the effects of a cholera outbreak. Dukoral and Shanchol are approved by the World Health Organization and are Oral Cholera Vaccines (OCVs). Unlike vaccines for the measles or chickenpox that confer lifetime immunity to the infection, these two OCVs only provide limited immunity. Dukoral protects against O1 serogroup strains of *V. cholerae*. 85% of recipients gain immunity to infection for 4-6 months and 60% gain immunity for up to 2 years. Additionally, Dukoral must be given in a minimum of two doses, a problem for public health professionals working in areas where patient follow-up is extremely difficult (“Weekly Epidemiological Record: Cholera 2013,” 2014). Shanchol is another OCV option that protects against both O1 and O139 serogroups of *V. cholerae*. 67% of patients administered 2 doses of Shanchol gain immunity for 2 years with no booster doses, and 65% are protected for 5 years (CDC, 2014). Due to the fact that recent outbreaks and the use of vaccines in areas with endemic cholera have depleted the global vaccine stockpile, the WHO notes that OCVs are primarily used as
a form of outbreak control rather than as a form of primary prevention (“Weekly Epidemiological Record: Cholera 2013,” 2014).

1.4 Non-O1/Non-O139 *Vibrio cholerae*

Each pandemic of cholera to date has been caused by O1 or O139 serogroup strains of *V. cholerae*. Due to the obvious health threat that they pose to the public, O1/O139 strains have also been the predominant strains focused on by researchers studying *V. cholerae*. However, non-O1/O139 toxigenic serogroups of *V. cholerae* have emerged that cause disease symptoms such as inflammatory diarrhea, gastroenteritis, wound infections, and septicemia (Ceccarelli et al., 2015). Non-O1/O139 strains have been associated with individual infections or localized outbreaks, especially in locations where cholera is endemic (Chatterjee et al., 2009). These genetically diverse strains do not carry the cholera toxin (CT) or toxin co-regulated pilus (TCP) genes, but can encode a variety of genes that contribute to alternative methods of virulence. Many non-O1/O139 strains encode genes for hemolysin, protease, mobility, Type VI Secretion systems (T6SS), Type III Secretion systems (T3SS), and biofilm formation proteins (Chatterjee et al., 2009). While very few non-O1/O139 isolates secrete the cholera toxin, some isolates encode genes for various other toxins including the MARTX toxin, a multifunctional autoprocessing repeats-in-toxin, *rtxA*, heat-stable enterotoxin (Nag-St), and the Cholix toxin (Schirmeister et al., 2014, Ceccarelli et al., 2015). A 2013 study conducted in the Chesapeake Bay area discovered that between 5-10% of all *Vibrio* infections are due to *V. cholerae* non-O1/O139 strains (Jones et al., 2013). Furthermore, it has been observed that non-O1/O139 *V. cholerae* isolates are more likely to be resistant to
penicillin and ampicillin, as well as sporadically resistant to other commonly used antibiotics such as erythromycin (Ceccarelli et al., 2015).

*V. cholerae* NRT36S is a serogroup O31 strain that is cholera toxin (CT) negative and heat-stable enterotoxin (Nag-ST) positive (Chen et al., 2007). This strain causes severe inflammatory diarrhea in patients, which is different from the secretory diarrheal symptomatic of infection with O1/O139 strains that encode CT, but can be just as severe (Dziejman et al., 2005, Chen et al, 2007). *V. cholerae* NRT36S, relies on a T3SS to cause disease in its host due to its lack of CT and Toxin coregulated pilus (TCP) genes (Shin et al., 2011).

### 1.5 Type III Secretion System (T3SS) in *Vibrio* species

Bacterial T3SSs are contact-dependent protein delivery organelles. A T3SS is a needle-like structure anchored in the bacterial cell membranes and spans the gap between the bacterial cell and its host (Galan et al., 2014). After making contact with a eukaryotic host cell, the T3SS can actively pump proteins, called effector proteins, from the bacterial cell cytosol through the host’s plasma membrane into the host cytosol. Once in the host cytosol, effector proteins can disrupt multiple cellular processes such as actin remodeling or signal transduction, causing disease (Galan et al., 2014). The presence of a T3SS can also cause damage in the absence of effector proteins. It has been shown that the translocation of the needle tip of the *Pseudomonas aeruginosa* T3SS into the host cell membrane is enough to cause host lung cell damage (Audia et al., 2013). This machinery is thought to be evolutionarily derived from the structure of the bacterial flagella due to the fact that the architecture of the membrane components resembles those of the bacterial flagella (Figure 3) (Galan et al., 2014). Additionally, *Mycoccus xanthus* bacteria display an intermediate non-
functional flagella-like structure that is capable of secreting proteins, providing experimental evidence for this theory (Konovalova et al., 2010, Galan et al., 2014). Of the proteins included in the T3SS apparatus, it has been found that both the inner membrane export apparatus proteins and the cytosolic component proteins are highly conserved, even amongst T3SSs of different species and genera (Galan et al., 2014). Of the conserved cytosolic proteins, the T3SS ATPase protein is often used for evolutionary and phylogenetic analysis due to its highly conserved nature.
T3SS play a crucial role in the pathogenesis of non-O1/O139 *V. cholerae* species. In the absence of the CT and TCP genes, these strains must utilize other means of virulence. T3SS genes have been found in clinical strains isolated from patients with diarrhea that lack CT and TCP virulence factors (Dziejman et al., 2005).
In addition to causing disease through the translocation of effector proteins, it has also been shown that the T3SS contributes to effective colonization in the absence of TCP genes in the infant mouse model (Dziejman et al., 2005). In fact, in *V. cholerae* AM-19226, a T3SS positive strain, both colonization and the secretory diarrheal response were found to be just as severe as infections with the cholera causing strain N16961 in the infant rabbit colonization model (Dziejman et al., 2005).

In this study we use *V. cholerae* NRT36, which encodes a T3SS that shares 99% sequence similarity to that of strain AM-19226 (Chen et al., 2007). A functional T3SS is required for the virulence of *V. cholerae* AM-19226 and presumably NRT36S (Shin et al., 2011). In NRT36S, the T3SS genes are encoded on a pathogenicity island that bears significant homology to VPI-2 of *V. cholerae* N16961 (Chen et al., 2007, Murphy and Boyd, 2008).

### 1.6 Pathogenicity Islands

Pathogenicity islands (PAIs) are large regions of DNA that contain virulence factors in the chromosome of pathogenic isolates and are absent from non-pathogenic isolates. (Hacker & Kaper, 2000). Pathogenicity islands have the following main characteristics that distinguish them from core chromosomal DNA. PAIs have a codon usage and a GC content that is different from that of the host core chromosome, and they integrate into the host chromosome via site specific integration at a tRNA gene. This process is mediated by an integrase and results in direct repeat sequences flanking the integrated PAI (Hacker & Kaper, 2000). These features indicate that PAIs have been acquired by horizontal gene transfer. PAIs can include genes that encode alternate metabolic proteins, confer antibiotic resistance, or increase bacterial virulence such as the T3SS. PAIs can also encode genes for self-regulation such as
toxT and tcpPH present in the Vibrio pathogenicity island-1 (VPI-1) (Hacker & Kaper, 2000). Although not self-mobilizable like a phage, PAIs are able to excise out of the core chromosome of the bacteria under certain conditions such as stress, ultra-violet light, or antibiotic exposure (Almagro-Moreno, Napolitano, & Boyd, 2010). Excision occurs when two direct repeat sites on the left and right sides of the island, attL and attR respectively, recombine and the island exists as a circular intermediate outside of the core genome of the bacteria (Rajanna et al., 2003). When these islands occur in a circular intermediate state, they can be transferred to other bacteria via conjugation or transduction (Hacker & Kaper, 2000). Throughout the course of evolution, PAIs can lose mobility genes such as integrase proteins and recombination directionality factors (RDFs) and become a permanently integrated part of the core chromosome of the bacteria, as can be seen in some Salmonella and E. coli spp (Marcus, Brumell, Pfeifer, & Finlay, 2000). This occurs when evolution favors the selection of bacteria containing the PAI due to the increased fitness it confers to the bacteria, either in terms of metabolism or other functions (Hacker & Kaper, 2000).

Pathogenicity Islands have been characterized in both E. coli and Salmonella species. In Salmonella, T3SS machinery has been found to be associated with PAIs (Galan, 1999). Additionally, PAIs present in uropathogenic and EPEC strains of E. coli were some of the first PAIs studied (Hacker et al., 1997). Consistent with what was later found in the PAIs of V. cholerae, E. coli PAIs also integrate site-specifically near tRNA synthase genes (Hacker et al., 1997). In EPEC strains of E. coli, the PAI present is responsible for the enterohaemorrhagic effects of the bacteria due to the presence of a T3SS on the island (Hacker et al., 1997, Marcus et al., 2000).
In choleraegenic *V. cholerae* serogroup O1 and O139 isolates, the *Vibrio* pathogenicity island-2 (VPI-2) region is integrated at a tRNA-ser gene and contains genes for an integrase, a type I restriction modification system (RM), a sialic acid catabolism (SAC) region, a phage-like region, and two RDFs named *vefA* and *vefB* (Figure 4) (Jermyn & Boyd, 2002). Although it was originally thought that this region was only present in choleraegenic strains of *V. cholerae*, homologous integrase, RDF, and SAC proteins have been found in AM-19226 and NRT36S, among other strains (Chen et al., 2007, Dziejman et al., 2005, Murphy & Boyd 2008, Jermyn & Boyd 2005). It is theorized that the two versions of VPI-2, one encoding a RM system and a phage, and one encoding a T3SS, have a common ancestor and have diverged throughout the course of evolution.

**Figure 4** VPI-1 and VPI-2 of *V. cholerae* N16961. The genes shown in orange represent the cognate integrase of the islands, *intV1* and *intV2* respectively. Shown in green is the tRNA synthase locus of integration.
1.7 Integrase Background

The excision of PAIs is mediated by the function of a protein called an integrase. Integrases are site-specific recombinase proteins that are necessary for the integration and excision of the PAI from the bacterial chromosome (Rajanna et al., 2003, Murphy & Boyd, 2008, Almagro-Moreno, Napolitano, & Boyd, 2010). Phage integrase proteins have three conserved domains, a domain of unknown function (DUF) that is commonly found on the N-terminus of phage integrase proteins, a phage integrase sterile alpha-like motif (SAM) that is also related to the n-terminus of the protein, and the phage integrase family domain (Figure 5) (Marchler-Bauer et al., 2015). The phage integrase family domain is located at the C-terminus of the protein, and houses the active site where double stranded DNA is cut during integration and excision (Marchler-Bauer et al., 2015). PAI excision has been found to be dependent on integrase proteins in both *E. coli* and *Yersinia pseudotuberculosis* strains as well as in *V. cholerae* (Lesic et al., 2004, Hochhut et al., 2006). In *V. cholerae* VPI-1, Rajanna et al. found that the island could excise without the cognate integrase, *intV1*, present (Rajanna et al., 2003). This could be due to the presence of *vpiT*, a transposase gene encoded on the island, or possibly crosstalk between VPI-1 and *intV2*-- the integrase protein of the second PAI present in *V. cholerae*, VPI-2 (Rajanna et al., 2003, Carpenter and Boyd, unpublished data) (Figure 4). It was also found that while inactivation of *intV1* did not abolish excision of VPI-1, the complementation of the gene on a plasmid in the mutant increased the rate of excision of the island (Rajanna et al., 2003). This result was not consistent with what is found in VPI-2. When the cognate integrase of VPI-2, or *intV2*, is deleted, excision of the island no longer occurs (Murphy & Boyd, 2008, Almagro-Moreno, Napolitano, & Boyd, 2010). The lack of a transposase gene on VPI-2 could explain this key difference in excision behavior.
between two PAIs in *V. cholerae*. Two of the aims of this study focus on the cognate integrase of VPI-3, the VPI-2 variant containing a T3SS in strain NRT36S. The first aim is to show that, like VPI-2, the integrase encoded on VPI-3 is necessary for excision. The second aim of this research is to study the evolution of T3SS containing PAIs using the sequence of the cognate integrase encoded on the island. Integrase sequence has been previously shown to be a reliable target of analysis in *E. coli* PAI evolution (Napolitano et al., 2011).

Figure 5 Conserved domains of phage integrase proteins. Figure adapted from the NCBI conserved domains database (Marchler-Bauer et al., 2015).
Chapter 2
MATERIALS AND METHODS

2.1 Bacterial Strains, Plasmids, & Growth Conditions

Bacterial strains were grown in 3mL lysogeny broth (LB) (Fisher Scientific) overnight (12 hours), at 37° C with shaking at 225 rpm in aerobic conditions unless otherwise noted. In the case of *E. coli* β2155 Δpir, the LB media was supplemented with Diaminopimelic acid (DAP) to a final concentration of 0.3 mM. When needed, the media was also supplemented with antibiotics in the following concentrations: 200 µg/ml streptomycin (Sm) (Fisher Scientific), 25 µg/ml chloramphenicol (Cm) (Fisher Scientific) and 100 µg/ml ampicillin (Amp) (Fisher Scientific). LB agar was supplemented with 10% sucrose for double cross screening.

Table 1 List of Plasmids and bacterial strains used in this study.

<table>
<thead>
<tr>
<th>Strain Name</th>
<th>Background</th>
<th>Source</th>
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<tr>
<td><em>V. cholerae</em> Strains:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NRT36S</td>
<td>O31, VPI-2, partial VSP-II, Sm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Chen et al. 2007</td>
</tr>
<tr>
<td>NRT36S ΔintV2</td>
<td>NRT36S, ΔVC1758, Sm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>This study</td>
</tr>
<tr>
<td>N16961</td>
<td>O1, El Tor strain, VPI-1, VPI-2, VSP-I, VSP-II, Sm&lt;sup&gt;R&lt;/sup&gt;</td>
<td>Heidelberg et al., 2000</td>
</tr>
<tr>
<td><em>E. coli</em> strains:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>β2155Δpir</td>
<td>Donor bacteria in conjugation experiment, DAP auxotroph, produces the π protein needed for plasmid replication outside of the bacterial chromosome</td>
<td></td>
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<tr>
<td>DH5αΔpir</td>
<td>Laboratory designed strain</td>
<td></td>
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of *E. coli* for cloning, high transformation frequency, produces the π protein needed for plasmid replication outside of the bacterial chromosome

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<thead>
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<th>Plasmid strains:</th>
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<th>Fermentas</th>
<th>This study</th>
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<tr>
<td>pJET1.2</td>
<td>cloning vector, Am&lt;sup&gt;r&lt;/sup&gt;</td>
<td></td>
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<tr>
<td>pJET1.2ΔintV2</td>
<td>Cloning vector, Am&lt;sup&gt;r&lt;/sup&gt;, contains truncated ΔVC1758 product</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pDS132</td>
<td>Suicide Vector, SacB, Cm&lt;sup&gt;r&lt;/sup&gt;</td>
<td>Philippe, Alcaraz et al. 2004</td>
<td></td>
</tr>
<tr>
<td>pDS132ΔintV2</td>
<td>pDS132 containing truncated ΔVC1758 product, Cm&lt;sup&gt;r&lt;/sup&gt;</td>
<td>This study</td>
<td></td>
</tr>
</tbody>
</table>

**2.2 Mutant Construction**

**2.2.1 PCR amplification and cloning**

The truncated gene was created using the Splicing by Overlap-Extension (SOE) method of PCR (Horton et al., 1989). The restriction enzyme cut sites used in this primer design were SacI (GAGCTC) and XbaI (TCTAGA). All primers were designed to create an in-frame gene deletion and were ordered from Integrated DNA Technologies. 2.0 μl of 1x dNTPs, 5.0 μl of 10x PCR Buffer, 2.0 μl of each primer, and 2.38 μl of Taq polymerase was used in each PCR reaction (Fisher), 1 μl of 10 ng/μl template DNA was also added to all reactions besides the control. In the SOE method, the AB and CD products of the gene are amplified individually in a 50 μL PCR reaction. The PCR conditions used to amplify the AB product were: 95°C for 5 mins, 29 cycles of 94°C for 30 sec, 48.3°C for 30 sec, and 72°C for 30 sec, 72°C for 7 min. Conditions used to amplify the CD product were: 95°C for 5 mins, 29 cycles of 94°C for 30 sec, 58.3°C for 30 sec, and 72°C for 41 sec, 72°C for 7 min. The AB
primers (VC1758A & NRTintv2B) amplified a region of 523 base pairs, and the CD primers (NRTintv2C & NRTintv2D) amplified a region of 678 base pairs. These products were used as DNA template in a third 50 μL PCR reaction, where the reverse complement of primer B, which is included in the sequence of primer C, anneals the AB and CD products together. The conditions used for this third PCR were: 95˚C for 5 mins, 29 cycles of 94˚ C for 30 sec, 55.4˚ C for 30 sec, and 72˚ C for 1:15, 72˚ C for 7 min resulting in formation of a truncated intV2 gene product. This truncated 1,161 base pair product is then cloned into the cloning vector pJET1.2 using the blunt end protocol (ThermoScientific). The blunt end protocol was used so that the DNA fragment can ligate directly into pJET1.2. A 1:3 vector to insert reaction was set up using 10 μL reaction buffer, 1 μL of T4 DNA ligase enzyme, and 19 μL of water. This reaction was kept at 16˚C for 14 hrs to ensure optimal ligation of the truncated ΔintV2 gene into pJET1.2

pJET1.2 is a plasmid designed to facilitate the cloning of DNA fragments and the positive selection of bacteria containing the truncated gene product. The plasmid contains the rep gene that is responsible for the replication of the plasmid, an origin of replication, the bla gene that confers ampicillin resistance to bacteria containing the plasmid, and the eco47IR gene that is lethal to the plasmid unless disrupted. The multiple cloning site of the plasmid is located within the eco47IR gene; therefore, when the truncated ΔintV2 gene is ligated into pJET1.2, this lethal gene is disrupted and the plasmid survives.

2.2.2 Bacterial Transformation

Once ΔintV2 is ligated into pJET1.2, it is then necessary to transform the plasmid pJET1.2 ΔintV2 into E. coli DH5α, an E. coli strain with a high
transformation frequency. pJET1.2ΔintV2 is transformed into *E. coli* DH5α using a CaCl₂ and heat shock reaction. The pDNA enters the *E. coli* cell when the plasma membrane is disrupted at a high temperature (42˚) in the presence of CaCl₂. Consequently, all *E. coli* bacteria that obtain the pJET1.2ΔintV2 plasmid through transformation also acquire ampicillin resistance from the plasmid. This resistance allows the *E. coli* containing the plasmid to be isolated on selective media. Once enough copies of the plasmid containing the truncated gene have been obtained through bacterial replication, pJET1.2 is purified from *E. coli* DH5α (QIAGEN Plasmid Mini Kit (2005)), and digested at specific cut sites with the SacI and XbaI restriction enzymes (ThermoScientific). The digested plasmid is run out on a 0.8% agarose gel. When viewed under a UV light, there are two bands present on the gel. The larger band is the linear plasmid, and the smaller band is the truncated ΔintV2 product that has been cut out by restriction enzymes. This product is purified from the agarose using gel cut purification. The purified product is then ligated into another plasmid, pDS132 overnight at 16˚ C (Philippe, Alcaraz et al. 2004).

pDS132 is a suicide vector that cannot extrachromosomally replicate in strains of bacteria that do not encode the *pir* gene. In the case that pDS132 is transformed into a strain of bacteria that does not encode the *pir* gene, such as *V. cholerae* NRT36S, the plasmid can only replicate if incorporated within the chromosome during DNA replication. Additionally, pDS132 contains the *cat* gene that confers Cm resistance. *Cat* encodes the enzyme chloramphenicol acetyltransferase, which detoxifies Cm to the bacteria by attaching an acetyl group to the antibiotic. This prevents the antibiotic from attaching to and disabling the bacterial ribosomes. Once the truncated gene product is ligated into pDS132, pDS132ΔintV2 is transformed into *E. coli* β2155 using
the same previously mentioned CaCl₂ transformation protocol. *E. coli* β2155 serves as the donor strain in bacterial conjugation with *V. cholerae* NRT36S.

### 2.2.3 Conjugation Protocol

Conjugation was used in order to transfer pDS132Δ*intV2* from *E. coli* β2155 to *V. cholerae* NRT36S. Following overnight growth, *E. coli* β2155 was plated on LB agar with 0.3mM of DAP and incubated at 37°C for five hours. *E. coli* β2155 is a DAP auxotroph, therefore adding DAP to the media was necessary for growth. However, no antibiotics were spread onto this plate because unlike *E. coli* β2155 + pDS132, *V. cholerae* NRT36S is not resistant to Cm. Following this incubation, 150 μL of exponential phase *V. cholerae* NRT36S was plated onto the *E. coli* β2155 plate and incubated at 37°C for 12 hours. The lawn of bacterial overnight growth was then resuspended in 3 mLs of LB, and 150 μL was spread on an LB agar plate supplemented with Sm and Cm. The absence of DAP and the presence of Sm on this plate ensured that NRT36S grew, but not *E. coli* β2155. The presence of Cm did not allow any *V. cholerae* NRT36S to grow that had not received the pDS132 plasmid. Thus, any bacteria grown on this plate should be *V. cholerae* NRT36S + pDS132Δ*intV2*.

Several conjugated colonies were selected and passaged on LB agar + Cm + Sm plates until robust, single colonies were obtained and confirmed by PCR (Figure 6). The double cross was then screened for. The selective media used to identify the double crossover event is LB agar + 10% sucrose + Str. pDS132 contains a sucrose sensitivity gene, *sacB*, which encodes the enzyme levansucrase. This enzyme catalyzes the synthesis of levans, which are fructose polymers toxic to *V. cholerae*. If the plasmid is incorporated into the genome of NRT36S, as it is after the single cross
has occurred, the bacterial colonies will appear lysed and mucoid in the presence of sucrose due to the presence of a functional *sacB* gene. In the event of the double crossover, the plasmid is removed from the core genome of the bacteria leaving only the truncated AD product. If the plasmid is no longer integrated, the bacteria are no longer sensitive to sucrose and their growth should appear phenotypically normal on the selective LB+ 10% sucrose media. Both PCR analysis and genomic sequencing (Genewiz) confirmed the expected deletion of the 1,108 base pairs between primer B and primer C in the IntV2 region of NRT36S (Figure 7).

![Agarose gel confirming the occurrence of the single crossover event. Because the plasmid becomes incorporated into the core chromosome of NRT36S after single crossover occurs, the PCR product using the flanking primers is too large to amplify using PCR.](image-url)
Figure 7 PCR amplification using AD and Flanking primer sets confirmed the deletion of the intV2 gene in the mutant strain. Amplification of a correctly sized product by the flanking primers indicates that the double crossover event has occurred and the plasmid is no longer incorporated into the bacterial chromosome.

2.3 Excision Assays

In order to detect whether the excision of VPI-2 has occurred in the absence of IntV2, a two-stage nested PCR was done to amplify the attB2 site. After one round of PCR, no attB2 product can be seen on the gel; however, after two rounds a strong band can be visualized if excision has occurred (Figure 8). The two stages of this assay are necessary due to the fact that the excision of the PAI is rare, and a more sensitive assay is needed to detect the genomic attB2 site that is left behind after the island excises. The first round PCR conditions to detect attB2 are 95°C for 5 min, then 30 circuits of 94°C for 30 sec, 60°C for 30 sec, 72°C for 1 min 30 sec, concluded by 72°C for 5 min. The primers used in this round were VPI-2fk1 and VPI-2fk2 (Table 2). A 1:10 dilution of the genomic DNA used in this assay was made and heated at
60°C for 20 min. After heating, the concentration of the 1:10 diluted gDNA was taken with the Nanodrop 2000. A subsequent dilution was made after the concentration was taken in order to bring the final concentration of the gDNA to 10 ng/μl. 1 μl of gDNA is used as template in the PCR reaction. The PCR conditions for the second round are as follows: 95°C for 5 min, then 30 circuits of 94°C for 30 sec, 60°C for 30 sec, and 72°C for 30 sec, concluded by 72°C for 5 min. The primers used in this second round were VPI-2attF and VPI-2attR (Table 2). 1 μl of PCR product from the first round of PCR was used as template for the second round of the excision assay. The sequences for all primers used in these excision assays can be found in Table 2.

Figure 8 PCR amplification of the A. attB and B. attP sites of wild type V. cholerae N16961 and NRT36S strains. Due to the low excision rate of this island, no product from PCR1 was detected; therefore, product from PCR 1 was used as template for PCR 2 to amplify attB2 and attP2. In B., V. cholerae SG7 was included as a negative control, as SG7 does not contain VPI-2. These results demonstrated that VPI-3 of NRT36S can excise from the chromosome and can be detected through our attB2 and attP2 assays.
Table 2 Oligonucleotide primers used for PCR in this study. Primers were designed for the study and prepared by Integrated DNA Technologies

<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence</th>
<th>Tm (˚)</th>
<th>GC(%)</th>
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</thead>
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<td>50</td>
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<tr>
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<td>GGATTTCTGCTACTACCGT</td>
<td>54</td>
<td>50</td>
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<tr>
<td>VC1785A*</td>
<td>TCTAGAGATTCGGTGAGTTGTCGGAG</td>
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<td>55</td>
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<tr>
<td>NRTintV2B</td>
<td>CATGAGCGAGAATTACTTGG</td>
<td>52</td>
<td>45</td>
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<tr>
<td>NRTintV2C**</td>
<td>CCAAGTAATTCTGCTCATGAAGCTACAGTGCCTGGTG</td>
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<tr>
<td>NRTintV2D***</td>
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<td>AGAGTGAAAGTCGCCAAAG</td>
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<td>47</td>
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<tr>
<td>VPI-2attR</td>
<td>GGTGCAATTTCCAGATTTG</td>
<td>59</td>
<td>52</td>
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</tbody>
</table>

* Bases in bold represent XbaI restriction enzyme cut sites
** Bases in italics represent the reverse complement of primer NRTintV2B
*** Bases in bold represent SacI restriction enzyme cut sites

2.4 Bioinformatics

2.4.1 Annotation of VPI-3 in ARTEMIS

The Artemis program is a bioinformatics tool developed by the Sanger Institute to analyze genome sequence data. Genebank files can be imported from outside databases into Artemis, which allows the user to edit these sequences or perform other functions such as analysis of RNA sequencing data, calculation of GC percentage, and also calculation of certain statistics such as the percent of pseudogenes, gene density, and protein coding bases (Rutherford et al., 2000).
In this study Artemis was used to annotate VPI-3 in *V. cholerae* NRT36S from raw whole genome sequence data. The FASTA sequence of the island was assembled from unordered contig level data obtained from Colin Stine’s lab using the homologous region of *V. cholerae* AM-19226 as a guideline to determine gene order. This was particularly useful when genes were located sequentially on AM-19226, but on separate contigs in NRT36S. This FASTA file was then imported into Artemis, which detected the open reading frames (ORFs) in the sequence. Each ORF was then run through NCBI’s protein BLAST algorithm and annotated in one of three ways. If the ORF was located in the T3SS region, it was annotated using the locus tag of the most homologous gene in *V. cholerae* AM-19226. If the ORF was located in the Sialic Acid Catabolism region of the island, it was annotated using the locus tag of the most homologous gene in *V. cholerae* N16961. If the ORF did not blast to a protein from either of these strains, the locus tag of the protein from the BLAST result with the highest homology and query cover was chosen for annotation. Recombination directionality factors *vefA* and *vefB* were identified by searching the sequence manually due to the fact that the ORFs were too small for the default settings of Artemis to find in the initial ORF search. The annotation was then saved in Genebank file format for further use.

### 2.4.2 ARTEMIS Comparison Tool (ACT) Alignments

Artemis is designed primarily for the viewing, analysis, and annotation of a single sequence, in comparison Artemis Comparison Tool is used for comparing and viewing homology between multiple large genomic sequences. Generating an ACT comparison, such as the one shown in Fig. 3, requires 2 different software programs. ACT allows the user to view a comparison and an external program is used to generate
a blastout or crunch file that can then be imported into ACT for viewing. Sanger offers a web based platform called WebACT that accomplishes this task and allows the user to download the blastout files to save for use with ACT (www.webact.org). On this web-based platform, FASTA files for up to 5 sequences can be uploaded for analysis and the program will generate 4 sequential blastout files. This means that a specific sequence is only compared to those sequences directly before and after it. When the sequences and the comparison files are loaded into ACT, the program generates the visual comparison. In ACT a red band connecting two sequences indicates a region of homology between the two sequences. The “intensity” of the band will vary depending on the percent of homology between the two regions (Carver et al. 2005). A blue band between the two sequences indicates a reverse match. Each ORF in NRT36S was given a number from 1 (intV2) to 70 (vefB), which corresponds to the numbers labeling the ORFs in Table 3. Annotations for V. cholerae N16961 and AM-19226 were taken in Genebank file format from the NCBI database and loaded into ACT. Nine additional strains with T3SS containing PAIs were selected for analysis with ACT (Table 4). The FASTA sequence of VPI-3 for each of these strains was downloaded from either the NCBI or IMG database. Background information for each of these strains including serogroup, location of isolation, year of isolation, and accession number for the genome can be found in Table 4.

2.4.3 Construction of Phylogenetic Trees

All Phylogenetic trees were constructed using the software program MEGA. Protein amino acid sequences were aligned by the ClustalW algorithm using the default parameters. A tree was then constructed from the aligned sequences using the neighbor-joining statistical method. Evolutionary distances were computed using the
Poisson correction method (Saitou & Nei, 1987, Zuckerkandl & Pauling, 1965, Tamura et al., 2011). Positions lacking data or containing gaps were removed from the analysis. All V. cholerae amino acid sequences were collected from NCBI and IMG genome databases. Analysis of T3SS ATPase proteins located near PAI integrase proteins was crucial, as we only wanted to examine T3SSs present on PAIs. Cognate integrase sequences on PAIs lacking a T3SS were also excluded from the analysis. The “Gene Neighborhood” tool in the IMG database was used to find associated integrase and T3SS ATPase proteins. 11 V. cholerae strains with T3SS containing PAIs, in addition to strain NRT36s, were selected for phylogenetic analysis due to the availability and assembly status of their genomes (Table 4).

The phylogenetic tree shown in Figure 17 was generated from MEGA using the same method and parameters as previously described (Saitou & Nei, 1987, Zuckerkandl & Pauling, 1965, Tamura et al., 2011). However, T3SS ATPases located outside of PAIs were not excluded from this analysis. A total of 27 T3SS ATPase sequences were chosen for the final analysis. The main criterion when selecting sequences was genus and species level diversity in the final data set.
Chapter 3

RESULTS

3.1 Structure of the Type III Secretion System Containing PAI in NRT36S

Construction of NRT36S VPI-3 from raw sequence data was the crucial first step in understanding the excision behavior and evolutionary history of the island. The presence of a VPI-2 like island in pathogenic V. cholerae non-O1/O139 was previously shown in a number of studies (Jermyn and Boyd, 2005, Chen et al., 2007, Murphy and Boyd, 2008). In this study we show that the island in V. cholerae NRT36S is made up of 70 Open Reading Frames (ORFs), which were sequentially numbered and will be referred to as ORF1-ORF70 (Table 3). The island shows extremely high homology with a VPI-2-like region of V. cholerae strain AM-19226 (Figure 10). We have named this island region in NRT36S as VPI-3. In NRT36S, VPI-3 ORFs 1-43 encode the various components of a Type III secretion system, including effector proteins. Homologous T3SS proteins are present in V. cholerae strains AM-19226, HE-25, TMA-21, VC12129, EM1676-A, VC35, CP1115, P-18785 and NHCC-008D (Table 4). ORFs 54-65 encode genes necessary for the metabolism of sialic acids. This sialic acid metabolism region is not only homologous in all of the strains listed above; it is also homologous to a similar region found in VPI-2 of O1 strains such as V. cholerae N16961. Gene maps of VPI-3 of NRT36S and VPI-2 of N16961 can be found in Figure 9. Figure 10 shows an ACT alignment of VPI-2 of N16961, VPI-3 of NRT36S, and VPI-3 of AM-19226. This figure clearly depicts
regions of significant genetic difference in the structures of VPI-2 in an O1/O139 strain of *V. cholerae* and VPI-3 of two pathogenic non-O1/O139 strains.

**NRT36S VPI-3 (67.9 kb):**

![Gene map of NRT36S VPI-3]

**N16961 VPI-2 (57.3 kb):**

![Gene map of N16961 VPI-2]

Figure 9 Gene maps of VPI-3 and VPI-2 regions of NRT36S and N16961 respectively. The conserved integrase of the islands, IntV2, is shown in orange. Green ORFs represent tRNA-ser, the locus of island integration. The T3SS is depicted as yellow ORFs. The purple region shown in VPI-3 is a variable region that can be divergent among multiple species containing this island. The SAC region (blue) contains the amino sugar sialic acid catabolism region. RM (grey) contains a restriction modification system. Red ORFs represent recombination directionality factors *vefA* and *vefB*.
Figure 10 The Artemis comparison tool (ACT) was used to examine sequence differences between VPI-2 and VPI-3 in *V. cholerae* strains N16961 (top), NRT36S (middle), and AM-19226 (bottom). Red blocks represent forward regions of homology between the three island regions and blue lines indicate reverse matches (Carver et al. 2005). In NRT36S, the T3SS is found inserted between the conserved integrase and SAC region rather than the RM system found in N16961. NRT36S also lacks the Mu phage region present in N16961. Two major differences are presented between the otherwise homologous T3SS region of NRT36S and AM-19226: 1) a 1,051 bp region at the 13 kb position (*) which comprises ORF17 and the intergenic region between ORF17 and ORF18 of NRT36S and 2) repeated sequence region (**) indicative of several 201 bp duplications between ORF 26 and ORF 27. All of the default settings were used in ACT, and all comparison files were generated using the webACT tool available at www.webact.org.

### 3.2 Excision of VPI-3

It has been previously shown that VPI-2 of *V. cholerae* El Tor biovar N16961 has the capability to excise out of the core chromosome of the bacteria, and that this excision is dependent upon the presence of the cognate integrase encoded on the island (Murphy and Boyd, 2007, Almagro-Moreno, Napolitano, & Boyd, 2010). The ability of VPI-3 to excise out of the core chromosome of NRT36S was tested using a two-stage nested PCR reaction. Two sets of primers (VPI2 fk1, VPI2 fk2 & VPI-2 attF,
VPI-2 attR) were designed in NRT36S to amplify a 524 bp area, including the 22 bp attB site, which could only be amplified by PCR if the island was not present in the core genome of the bacteria. When wild-type strains were tested (Figure 8a) the expected band was observed for both NRT36S and N16961, a positive control strain. This indicates that the rare excision event is occurring, allowing the primers to bind and the PCR to amplify the 524 bp site that includes the attB site. The same result is found when the wild type strains are tested for amplification of a region containing the 497 bp attP site (Figure 8b).

In order to test whether the cognate integrase of NRT36S VPI-3 (intV2) is also necessary for the excision of VPI-3, the same assay described above was performed on NRT36SΔintV2 (Figure 10). No expected band was observed following our two round PCR reaction for NRT36SΔintV2 (Figure 11). The primers were not able to amplify the 524 bp site due to the fact that the island was unable to excise; therefore, the area between the primers was too large for amplification via PCR. The absence of the cognate integrase from the island causes this result, suggesting that this protein is crucial for the excision of VPI-3 as well as in the strains where it has been previously shown in V. cholerae (Rajanna et al., 2003; Murphy and Boyd, 2008, Almagro-Moreno, Napolitano, & Boyd, 2010). All PCR reactions described above were repeated a minimum of three times. When examining the amino acid sequence of V. cholerae strain CP1115, a premature stop codon was found that prevented the translation of the first 60 amino acids of the cognate integrase protein of VPI-3. As a result of this, the DUF domain of this integrase was truncated (Figure 12).
Figure 11 The two-stage *attB* PCR amplification assay was used to detect excision of VPI-3 in Δ*intV2* and wild-type NRT36S strains. As expected, no PCR product was detected in the first round. A wild-type *attB* excision product of 524 bp was detected in the second round, however no product was detected for Δ*intV2* indicating excision had been abolished in this strain.

Figure 12 Conserved domains of the integrase protein present on VPI-3 of *V. cholerae* strain CP1115. The truncated DUF domain at the N-terminus of the protein due to the disruption of translation is shown in green. Figure adapted from the NCBI conserved domain database (Marchler-Bauer et al., 2015).
3.3 ORFs within the NRT36S VPI-3 Region

The 62.9 kb region that makes up the VPI-3 region of NRT36S is made up of 70 Open Reading Frames (ORFS) (Table 3). Most of these ORFs fall within the two distinct regions of the island, the Type III Secretion system region (ORFs 3-45) and the Sialic Acid catabolism region (ORFs 54-65). ORFs 1-14 are 99% homologous to A33_1660 to A33_1671 in AM-19226. ORF16 has 97% identity with its homolog in AM-19926 (A33_1673), and lies in a region that is 96% identical to homologous sequence in AM-19226. ORF 17 has 78% identity with its homolog in AM-19226, and lies in a region of sequence with significant non-homology to AM-19226 (13117-14005). ORF 18 has 97% homology to its homolog A33_1675, and lies within a region of sequence that is 97% homologous to corresponding sequence in AM-19226. ORF 19 is 99% homologous to its AM homolog, but lies within a region of sequence that is 97% identical to corresponding AM sequence. ORF 20 is 87% identical to its AM-19226 homolog, A33_1677 and lies within a region of non-homology between the two sequences (15185-15356). The sequence containing ORFs 21-27 is 99% homologous to the corresponding sequence in AM-19226 (A33_1678-A33_1683). The sequence containing ORFs 29-40 is 99% homologous to the corresponding sequence in AM-19226 except for a 22 bp gap at 29.6 KB (29652-29674). ORFs 41-52 are encoded in a region of sequence that is 99% homologous to between NRT36S and AM-19226. The sequence containing ORFs 53-71 is 99% homologous to the sequence containing A33_1707-AM_1723. Although the nucleotide sequence is identical between the two strains in this region, no annotated protein in AM_19226 is homologous to ORF 53 found by ARTEMIS (hypothetical protein). ORFs 66-70 are
all 100% homologous to their AM-19926 homologs, except for ORF 72, which contains 2 gaps between in homology between it and A33_1724.

Table 3 VPI-3 Open Reading Frames, gene name, amino acid length, and % homology

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<th>ORF</th>
<th>Gene Name</th>
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<th>N16961 Homolog (% ID)</th>
<th>Other Homolog (% ID)</th>
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### 3.4 Presence of VPI-3 in other *V. cholerae* strains

Using the ARTEMIS comparison tool, the T3SS containing island of NRT36S was compared to nine other T3SS containing strains of *V. cholerae* (Figure 13). While the main components of the T3SS and the sialic acid metabolism region were generally conserved amongst the ten strains, there were regions of non-homology present, suggesting divergence throughout the evolution of VPI-3 in *V. cholerae*. The first major gap in homology is at 13.2 kb in NRT36S. The gap extends through the region between ORF16 and ORF18 (ToxR2), including the absence of most of ORF17 (hypothetical protein). These ORFs lie within the T3SS of NRT36S. This gap in homology is present in each strain selected for comparison except for CP115. At 33 kb, the region of non-homology between *V. cholerae* AM-19226 and TMA-21 spans the region between A33_1695 (hypothetical protein) and A33_1697 (Zinc finger domain protein). In AM-19226 this region encodes effector protein VopF.

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Interestingly, when we examine the region at 33 kb in NRT36S an ORF with homology to VopF was not identified. The absence of an effector protein in two otherwise highly homologous strains of *V. cholerae* is noteworthy, especially if VopF is translocated into the host cell via the T3SS. Another significant gap in homology exists between NRT36S and *V. cholerae* strains TMA-21 and V51 at around 29 kb, a region that encodes a hypothetical protein. These gaps in homology could indicate hot spots for recombination events that allow virulence elements to enter and exit the island during horizontal gene transfer or the evolution of novel effector proteins. For example, in *V. cholerae* El Tor biovar N16961 the Mu-phage region is found inserted between *vefA* and *vefB*, whereas this region is absent in NRT36S. Gaps within the T3SS itself could lend insight into the evolution of this virulence system.
Figure 13 Artemis comparisons of the T3SS containing PAIs of 10 *V. cholerae* strains. In each comparison, *V. cholerae* NRT36S is sequence one. Strains compared to NRT36S in A-I are AM-19226, TMA-21, VC12129, EM-1676A, NHCC008D, VC35, HE-25, P-18785, and CP1115 respectively. Two prominent and conserved sites of non-homology are the gaps at the 13 kb and 29 kb positions.

Table 4 Background information of *V. cholerae* strains used in bioinformatic comparisons of the VPI-3 region

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### 3.5 Discovery of a repeated region of homology in T3SS+ strains of V. cholerae

During the ACT comparison of the highly homologous VPI-3 regions of NRT36S and AM-19226 a series of 3 repeated sequences was discovered beginning at approximately 22.4 kb. The three repeats are 201 base pairs long and are 95% identical between the sequence of NRT36S and AM-19226 (Figure 14a). In AM-19226 each of these repeats is separated by 234 base pairs; the regions separating the repeats are 96% identical, and the repeats themselves are between 95-100% identical. The repeats fall in an intergenic region between NRT36S ORFS 26, a VsaC lipoprotein, and ORF 27, a hypothetical protein. In contrast, two separate ORFs are not found in the annotation of AM-19226. In that region a 1,328 amino acid hypothetical protein, A33_1684, has been identified and annotated (Alam et al., 2011). In order to confirm that A33_1684 is the only ORF present in that region of AM-
19226, the sequence was entered into NCBI’s ORFinder program (Figure 14b). This program confirmed the presence of a single large ORF in the region containing the repeated sequences. Further analysis revealed the presence of a 46 bp smaller repeated sequence directly downstream of the 201 bp repeated sequence in NRT36S. This sequence is also repeated three times in AM-19226 and is 82% identical to the sequence that originates in NRT36S. In AM-19226 two of the repeats are identical and one has a one bp T to C substitution at the 19 bp site, a G to A substitution at the 103 bp site, an A to G substitution at the 113 bp site, a T to C substitution at the 118 bp site, and a C to G substitution at the 145 bp site (Figure 18).

Variations of this repeated region were found in nine strains of *V. cholerae* containing a T3SS, including AM-19226. Background information on these strains can be found in Table 3. When the sequence of the island region of these strains was compared to that of NRT36S, it was found that each of these strains has a unique number of repeats, and the sequence that is repeated between the comparison strain and NRT36S can vary (Figure 15). In seven of the nine strains, the repeats disrupt the transcription of the ORFs in the region, leading to the transcription of a large ORF similar to what was discovered in AM-19226 (Figure 15).
Figure 14 Further examination of the repeated region at 22.4 kb between *V. cholerae* strains NRT36S and AM-19226. **A.** Enlarged Artemis comparison of NRT36S and AM-19226 reveals the difference in annotation of the ORFs in this region. In NRT36S, 2 ORFS separated by a 265 bp intergenic region were detected. However, in AM-19226, one large ORF (A33_1684) encompassing 1328 amino acids is detected in the corresponding region. A33_1648 encodes a probable effector protein (Alam et al. 2011). **B.** NCBI ORFinder display confirming that no alternative ORFs could be found in the region encoding A33_1684 in AM-19226.
Figure 15 Comparison of the repeated region of NRT36S to the corresponding region of 9 different *V. cholerae* strains. The number of repeats varies in all cases; however, the repeat always originates in the intergenic region of NRT36S VPI-3 between ORF 26 and ORF 27.
3.6 Phylogenetic Analysis of the T3SS ATPase of VPI-3

The evolution of a PAI can be speculated by examining the evolution of its component biological function genes. The phylogenetic trees of integrase proteins and T3SS ATPase proteins of various *V. cholerae* PAIs were compared (Figure 16). The T3SS ATPase amino acid sequence was chosen for analysis as it is conserved not only across T3SSs of different species and genera of bacteria, but across different types of T3SS as well, such as T3SS2α and T3SS2β. Only T3SS containing islands with an associated integrase were included in the phylogenetic analysis. Figure 16a shows that the T3SS ATPase proteins found on VPI-3 of *V. cholerae* strains NRT36S, AM-19226, CP1110, EM-1676A, V51, TMA21, 12129, VC35, HE-25 and P-18785 are all highly homologous. These strains encode a T3SS2α. *V. cholerae* strains 1587 and 623-39 cluster together separately because they encode the variant T3SS2-β. A phylogenetic analysis of the cognate integrase of VPI-3 was also performed (Figure 16b). Amino acid sequences of integrase proteins found on PAIs containing T3SS were included. The 11 *V. cholerae* strains that all encode highly homologous T3SS ATPase proteins display a different clustering pattern when analyzing the cognate integrase of VPI-3. It is important to note the difference in scale between the two phylogenetic trees. On the T3SS ATPase tree, one scale bar equals 0.01 amino acid substitutions per site. One the integrase tree this number is much lower; one scale bar equals 0.002 amino acid substitutions per site. Hence, the different clades *V. cholerae* integrase proteins are more highly related than the two distinct clades of T3SS ATPase proteins.

The T3SS ATPase from 27 different *Vibrionaceae* and *Salmonella* bacteria was also chosen for phylogenetic analysis. The tree shown in Figure 17 illustrates the relationships between the two varieties of T3SSs found in different *Vibrios* and
*Salmonella*. The *Salmonella* strains examined contain a T3SS1, whereas the *V. cholerae* strains encode a T3SS2. *V. parahaemolyticus* strains studied have either T3SS1 or T3SS2, and strains such as RIMD2210633 acquired both systems on separate chromosomes or other mobile genetic elements. There are two types of T3SS2, T3SS2α and T3SS2β. Various *Vibrio* species contain either the T3SS2α or T3SS2β. The clustering pattern of the phylogenetic tree shown in Figure 17 indicates that the T3SS ATPase of strain NRT36S is most closely related to strain AM-19226. The T3SS of NRT36S is also closely related to certain strains of *V. parahaemolyticus*, *V. mimicus* and *G. hollisae* (Figure 17). The clustering of T3SS containing strains of bacteria of different genera and species on the tree suggests that the T3SS was transferred amongst these species horizontally. The exact direction of this transfer is unknown.
Figure 16 A. Phylogenetic tree of T3SS ATPases in *V. cholerae* strains with an associated integrase protein encoded on a PAI. B. Phylogenetic tree of integrase proteins encoded on PAIs containing T3SS. Sequences were obtained from either NCBI or IMG databases and uploaded into MEGA where they were aligned using the ClustalW algorithm (Tamura et al., 2007). After alignment, MEGA was used to generate a neighbor-joining tree, using the bootstrap method (Saitou & Nei, 1987; Felsenstein, 1985). 1,000 bootstraps were used. In all phylogenetic trees the entry after the strain name indicates the locus tag of the protein on the tree, and the second entry is the IMG database gene ID number. Note the difference in the scales of the two trees.
Figure 17. Phylogenetic tree of the T3SS ATPase protein in multiple species of Gram-negative bacteria. Tree was constructed using the neighbor-joining method described in Figure 13.
Chapter 4
DISCUSSION

4.1 Excision behavior of T3SS containing PAI

It has been demonstrated that the cognate integrase of VPI-2 is necessary for the excision of VPI-2 in *V. cholerae* (Rajanna et al., 2003, Murphy & Boyd, 2008, Almagro-Moreno, Napolitano, & Boyd, 2010). However, excision in a variant of VPI-2 containing a T3SS, which we named VPI-3, was not characterized before this study. It was confirmed that the cognate integrase of VPI-3 is necessary for the excision of the island. This observation is relevant not only because the biological function genes on the islands are drastically different, but also because VPI-3 is approximately 10 kb larger than VPI-2 in *V. cholerae* N16961. It is possible that the truncated DUF domain could affect the folding or functioning of VPI-3’s cognate integrase in strain CP1115; however, this result is unlikely due to the fact that the truncated domain is not present in the phage integrase domain nor is it near the active site of the protein. Further investigation into the effects of this mutation is required.

4.2 Evidence for the mosaic evolution of *V. cholerae* PAIs

It is known that PAIs are involved in the evolution of novel and virulent strains of bacteria; however, less is understood about the evolution of the PAI regions themselves. It is possible that PAIs evolve as one defined unit over time, or more fluidly as a mosaic-like structure where biological function genes could evolve within the island. In this model, the evolution of the island does not follow a sequential path where mutations occur in one region while the rest of the island stays consistent. Instead, the island evolves in a manner where the components of the island are not a
discrete unit and have evolved independently over the course of time. If the PAI followed the sequential path of evolution, the biological function genes such as the T3SS proteins, would have evolved in a similar manner to the structural components of the island, such as the integrase protein. In strains of *V. cholerae* containing the T3SS2, the ATPase proteins are separated into two clades representing T3SS proteins present in T3SS2α and T3SS2β, but are otherwise identical (Figure 15a). However, the integrase protein is not genetically identical in the majority of these strains, as shown in the phylogenetic tree above (Figure 15b). The incongruent structure of the two trees suggests that the integrase and T3SS proteins did not co-evolve as would be expected if the island was a discrete entity. There are two evolutionary distinct clades present on the T3SS ATPase tree representing the ATPase present in the T3SS2α or T3SS2β. In contrast, the integrase proteins cluster into four separate clades that are very closely related, even between strains with T3SS2α and T3SS2β present on the island. If these two proteins evolved together over time, the structures of the trees presented in Figure 15 would be the same and the same strains would consistently cluster together. This divergence implies that the T3SS was acquired at a different point in the course of the evolution of VPI-3 for each of these strains, providing evidence to support the theory that PAIs can evolve in a mosaic-like manner.

**4.3 Evolution of new effector proteins due to multiple repeated sequences**

This study identified a repeated region in multiple strains of *V. cholerae* encoding T3SS PAIs (Figure 15). The ORFs shown by blue arrows in the comparison strains were found by NCBI’s ORFinder program (Figure 15). Although the exact reason and mechanism for these duplications is unknown, we hypothesize that they do affect the transcription of nearby genes by causing frameshift mutations, disrupting
start and stop codons, and potentially contributing to the formation of novel effector proteins. Figure 14 shows one such disruption when comparing the DNA sequence between NRT36S and AM-19226. It appears as if the duplications that occur in the sequence of AM-19226 disrupt the stop codon of ORF 27, leading to the transcription of one large ORF in that region as opposed to the two smaller ORFs seen in NRT36S. Alam et al. found that this ORF in AM-19226 (A33_1684) was translocated into the host cell, providing experimental evidence that this protein may be an effector (Alam et al., 2011). This fusion has occurred in all strains examined except for *V. cholerae* EMA-1676A and CP1115. It has been previously shown that the fusion of ORFs is a contributing factor to the evolution of effector proteins (Alam et al., 2011). This region of repeats may be evidence of this evolutionary phenomenon.

4.4 Relevance to the scientific community

Understanding the evolution of non-O1/O139 strains of *V. cholerae* is an extremely important aspect of *V. cholerae* research. Because these strains do not present classical choleraegenic symptoms, they can often go misdiagnosed as “acute watery diarrhea” in treatment centers (WHO, 2014). Thus, while the mortality rate from *V. cholerae* strains lacking CT and TCP may be low, it is estimated that the morbidity levels from non-O1/O139 strains causing localized and patient level outbreaks is much higher than reported. Furthermore, as discussed previously, these non-O1/O139 strains evolve rapidly and may have acquired antibiotic resistance. They are not recognized by the immune response in areas where cholera is endemic and native populations have acquired immune responses to O1/O139 strains. No vaccines are available to provide protection against non-O1/O139 strains of *V. cholerae* if a major outbreak were to occur, and in strains that encode a T3SS, disease symptoms
can be just as severe as infection with classical or El Tor strains. Expanding the existing knowledge base to include the evolution of the molecular mechanisms of pathogenesis in these non-O1/O139 strains is crucial if the medical and public health fields are to effectively prepare for future outbreaks and possible epidemics.
REFERENCES


Marchler-Bauer A et al. (2015), "CDD: NCBI's conserved domain database.", *Nucleic Acids Res.* 43(D)222


Marchler-Bauer A et al. (2009), "CDD: specific functional annotation with the Conserved Domain Database.", *Nucleic Acids Res.* 37(D)205-10.


55


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Appendix A

ALIGNMENTS OF REPEATED REGIONS IN AM-19226

A.

<table>
<thead>
<tr>
<th>Score</th>
<th>Expect</th>
<th>Identities</th>
<th>Gaps</th>
<th>Strand</th>
</tr>
</thead>
<tbody>
<tr>
<td>366 bits(198)</td>
<td>1e-106</td>
<td>198/198(100%)</td>
<td>0/198(0%)</td>
<td>Plus/Plus</td>
</tr>
</tbody>
</table>

Query 1: TCTGCACTGCACTGTCACCCGCCCGCTCGGCAACGCTGTAAGGT
Sbjct 1: TCTGCACTGCACTGTCACCCGCCCGCTCGGCAACGCTGTAAGGT

Query 61: ATGGTCGCGCTTCTCACTGGTAAGTTCCGGCTTCTCGGCAACGCTGTAAGGT
Sbjct 61: ATGGTCGCGCTTCTCACTGGTAAGTTCCGGCTTCTCGGCAACGCTGTAAGGT

Query 121: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT
Sbjct 121: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT

Query 181: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT
Sbjct 181: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT

B.

<table>
<thead>
<tr>
<th>Score</th>
<th>Expect</th>
<th>Identities</th>
<th>Gaps</th>
<th>Strand</th>
</tr>
</thead>
<tbody>
<tr>
<td>324 bits(175)</td>
<td>9e-94</td>
<td>189/196(95%)</td>
<td>0/198(0%)</td>
<td>Plus/Plus</td>
</tr>
</tbody>
</table>

Query 1: TGGTCACTGCACTGTCACCCGCCCGCTCGGCAACGCTGTAAGGT
Sbjct 1: TGGTCACTGCACTGTCACCCGCCCGCTCGGCAACGCTGTAAGGT

Query 61: ATGGTCGCGCTTCTCACTGGTAAGTTCCGGCTTCTCGGCAACGCTGTAAGGT
Sbjct 61: ATGGTCGCGCTTCTCACTGGTAAGTTCCGGCTTCTCGGCAACGCTGTAAGGT

Query 121: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT
Sbjct 121: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT

Query 181: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT
Sbjct 181: TCTGACGCAGACGTGGGCTGCGGCTCGGCAACGCTGTAAGGT

Figure 18 BLAST alignments of the 201 base pair repeated region found in *V. cholerae* AM-19226. A. The subject of the alignment is the first repeat in the sequence and the query is the third repeat. B. The subject of the alignment is the first repeat in the sequence and the query is the second repeat. Note that repeats one and three are 100% identical, while repeats one and two are only 95% identical.
Appendix B

EXAMINING PAI EXCISION IN RECA, HIMA, AND FIS MUTANTS IN V. CHOLERAE

The genes himA and fis have been found to affect excision rates of the λ bacteriophage and SXT, which are mobile genetic elements that behave similarly to PAIs. Another gene, recA is known to influence homologous recombination, whereas site-specific recombination, the mechanism by which PAIs excise, can occur independently of recA. We used three mutant strains of V. cholerae, Bah-3 (ΔrecA), Sm463 (ΔhimA), and Sm465 (Δfis), to determine if the absence of these genes, which are located outside of the PAI, would affect excision of VPI-1 and VPI-2 in V. cholerae. We found that himA, fis, and recA do not play a significant role in VPI-1 and VPI-2 excision.

Table 5 Mutants examined, size of selected gene in wild-type, and size of selected gene

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Size of Mutant (bp)</th>
<th>Size of WT (bp)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sm465 (Δfis)</td>
<td>192</td>
<td>399</td>
</tr>
<tr>
<td>Sm463 (ΔhimA)</td>
<td>75</td>
<td>297</td>
</tr>
<tr>
<td>Bah-3 (ΔrecA)</td>
<td>~540</td>
<td>1656</td>
</tr>
</tbody>
</table>
Figure 19 Diagram of excision related proteins examined in this study. A. Excision is found to be most efficient in the λ bacteriophage when the phage encoded excisionase (Xis) protein is bound to one of two binding sites (shown in teal) and the Fis protein is bound to a separate site (shown in grey). B. HimA is the alpha subunit of the protein Integration Host Factor (IHF). The binding of IHF to specific sites on the MGE chromosome (shown in orange) is required in order for the attR and attL sites on the bacterial chromosome to recombine and reform the attB and attP sites on the chromosome and the PAI (i.e. for the excision event to occur).

Figure 20 Growth curve analysis of mutants included in this study. No significant growth defects resulted from the deletion of the recA, himA, or fis genes.
Excision occurs in all deletion mutants, indicating that the presence of the examined genes is not necessary for the excision of *V. cholerae* pathogenicity island VPI-1.

Excision occurs in all deletion mutants, indicating that the presence of the examined genes is also not necessary for the excision of *V. cholerae* PAI VPI-2.