2015

The Characterization of a Putative Protease Expressed by Sneathia amnii

Rana Mehr
Virginia Commonwealth University

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CHARACTERIZATION OF A PUTATIVE PROTEASE EXPRESSED BY *SNEATHIA AMNII*

A thesis submitted in partial fulfillment of the requirements for the degree of Master of Science at Virginia Commonwealth University

by

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July, 2015
Acknowledgements

I would first like to express my deepest gratitude to my mentor Dr. Kimberly Jefferson. Her continuous mentorship, trust, and support in academic, scientific, and personal experiences have empowered me to successfully complete my graduate career both academically and scientifically. She has aided my development as an independent scientist which would have not been possible without guidance.

I would also like to thank the members of my graduate advisory committee: Dr. Dennis Ohman and Dr. Darrell Peterson. Their advice and direction have allowed me to better understand my project and their invaluable knowledge has made me a better scientist. Furthermore, I would like to thank Dr. Gail Christie and Dr. Cynthia Cornelissen for allowing me access to their laboratory resources and equipment.

Finally, I would like to thank all of my lab-mates, both past and present: Dr. Shreni Mistry, Dr. Zhumei Cronk, Dr. Michael Harwich, Melissa Prestosa, Abdallah Abdelmaksoud, and Matthew Allen-Daniels. They have provided invaluable input, guidance, and patience throughout my graduate career. I would like to specially thank my two lab-mates Dr. Jamie Brooks and Amy Sanford for their never ending laughter, friendship, advice, and support that has abetted in my successful conclusion to my graduate career. My closing acknowledgement is extended Dr. Kristin Lane and Laura Klenow, who both always have advice whenever I have needed it.
Dedication

I would like to dedicate this work to my parents for their unyielding love, support, dedication, and guidance which have led me to my successes. I would also like to dedicate this to my brother who has taught me to be a strong independent person. Finally, I must thank my boyfriend, Eric Hall, who has supported me and encouraged me throughout my scientific career. My successes are due to the endless love and support my loved ones have given me.
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<tr>
<td>°C</td>
<td>degrees centigrade</td>
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<tr>
<td>μg</td>
<td>microgram</td>
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<tr>
<td>μl</td>
<td>microliter</td>
</tr>
<tr>
<td>AA</td>
<td>amino acids</td>
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<tr>
<td>Amp</td>
<td>ampicillin</td>
</tr>
<tr>
<td>Amp’-</td>
<td>ampicillin resistance</td>
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<tr>
<td>BHI</td>
<td>brain heart infusion</td>
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<tr>
<td>B-ME</td>
<td>beta-mercaptoethanol</td>
</tr>
<tr>
<td>bp</td>
<td>base pair</td>
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<tr>
<td>BSA</td>
<td>bovine serum albumin</td>
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<tr>
<td>BV</td>
<td>bacterial vaginosis</td>
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<tr>
<td>C</td>
<td>cytosine</td>
</tr>
<tr>
<td>C-section</td>
<td>Cesarean section</td>
</tr>
<tr>
<td>CaCl₂</td>
<td>calcium chloride</td>
</tr>
<tr>
<td>CFU</td>
<td>colony forming unit</td>
</tr>
<tr>
<td>Cm</td>
<td>chloramphenicol</td>
</tr>
<tr>
<td>Cm’</td>
<td>chloramphenicol resistance</td>
</tr>
<tr>
<td>CRH-R1</td>
<td>corticotropin releasing hormone receptor</td>
</tr>
<tr>
<td>DMEM</td>
<td>Dulbecco’s modified eagle media</td>
</tr>
<tr>
<td>DNA</td>
<td>deoxyribonucleic acid</td>
</tr>
<tr>
<td>dNTP</td>
<td>deoxynucleotide triphosphate</td>
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<tr>
<td>ECM</td>
<td>extracellular matrix</td>
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<tr>
<td>E. coli</td>
<td>Escherichia coli</td>
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<tr>
<td>FIRS</td>
<td>fetal inflammatory response syndrome</td>
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<tr>
<td>g</td>
<td>gram</td>
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<tr>
<td>G</td>
<td>Glycine</td>
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<tr>
<td>IgA</td>
<td>Immunoglobulin A</td>
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<td>IL-1a</td>
<td>interleukin-1a</td>
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<tr>
<td>IL-1b</td>
<td>interleukin-1b</td>
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<tr>
<td>IL-6</td>
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<tr>
<td>IL-8</td>
<td>interleukin-8</td>
</tr>
<tr>
<td>IL-10</td>
<td>interleukin-10</td>
</tr>
<tr>
<td>IPTG</td>
<td>isopropyl β-D-1-thiogalactopyranoside</td>
</tr>
<tr>
<td>IUD</td>
<td>intrauterine device</td>
</tr>
<tr>
<td>K</td>
<td>Lysine</td>
</tr>
<tr>
<td>KCl</td>
<td>potassium chloride</td>
</tr>
<tr>
<td>kDa</td>
<td>kilodalton</td>
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<tr>
<td>L</td>
<td>Leucine</td>
</tr>
<tr>
<td>L</td>
<td>liter</td>
</tr>
<tr>
<td>L. crispatus</td>
<td>Lactobacillus crispatus</td>
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Preterm birth, birth prior to 37 weeks gestation, is the leading cause of neonatal mortality and morbidity worldwide. While the uterine cavity and amniotic fluid largely remain sterile throughout gestation, bacterial infections can occur and are associated with preterm birth and/or preterm premature rupture of the fetal membranes (PPROM). *Sneathia amnii* can be detected as a component of the vaginal flora in healthy women; however, it’s also associated with bacterial vaginosis and preterm birth. Sn35, an isolate of *S.amnii*, was identified and sequenced through the Vaginal Human Microbiome Project at VCU. Our objective was to classify potential
virulence determinants in Sn35 and we successfully identified a putative zinc endopeptidase. The zinc endopeptidase appeared to cleave itself in a site-specific manner under calcium-depleted conditions, resulting in a truncated protein. The truncated protein did have collagenase activity and bacteriolytic activity as well.
Clinical Significance of Preterm Birth

Preterm birth, defined as birth prior to 37 weeks gestation, is the leading cause of neonatal morbidity and mortality worldwide (Jefferson, 2012; Wiktin, 2014). The clinical definition of preterm labor is regular contractions that are accompanied with cervical change in less than 37 weeks gestation and is now believed to be a syndrome that is initiated by multiple routes (Goldenberg et al., 2008). Preterm birth can occur where labor is induced or infant is delivered through C-section, spontaneous preterm labor while fetal membranes are intact, or preterm premature rupture of the membranes, PPROM (Goldenberg et al., 2008). It is associated with 70% of neonatal death, 75% of neonatal morbidity and 50% of long-term neurological sequelae (Lacovidou et al., 2010; Whidbey et al., 2013). Classifications of preterm birth are as follows; extremely preterm is defined as birth after 20 weeks gestation but before 28 weeks of completed gestation, very preterm is defined as birth after 28 weeks but before 32 weeks of completed gestation, and moderate preterm is defined as birth after 32 weeks but prior to 37 weeks (Lawn et al., 2010). There are multiple physical risk factors associated with preterm birth including: tobacco and drug use, carrying multiples, uterine abnormalities, extremes in maternal age, systemic and/or intrauterine infections, bacterial vaginosis, and cervical length (Ganu et al.,
The greatest risk factor for preterm birth is having had a previous preterm birth (Jefferson, 2012). Non-physical factors that are associated with preterm birth are low socioeconomic status and race (Fettweis et al., 2014; Goldenberg et al., 2008).

The annual cost of health care associated with premature birth in the United States was at minimum $26 billion in 2005, and in 2004 12.5% of births in the U.S. were preterm (Behrman et al., 2007; Hyman et al. 2013). While the U.S. is a developed country with modern medicinal technology, it currently holds a higher rate of preterm birth than the average rate, 7.5%, for developed countries and has an equal rate of preterm births with countries that are least developed. Additionally, within the U.S., African American women have an increased risk of delivering preterm; 18.4% of African American women delivered preterm and 11.7% of non-Hispanic white women delivered preterm.

**Intrauterine and Amniotic Cavity Invasion by Vaginal Bacteria**

While preterm birth is caused by a multitude of factors, between 25-40% of preterm births are caused by intrauterine infection and inflammation (Wen et al., 2013; Rubens et al., 2010; Gravett et al., 2010). There is strong evidence supporting that infection induces preterm birth; experimental infection or LPS injection in mice has been shown to induce preterm birth, intrauterine infections among women correlates with preterm birth, systemic infections among women is associated with preterm birth and chorioamnionitic tissue isolated from women who had spontaneous labor at less than 30 weeks were infected with bacteria (Hirsch et al., 1995; Schantz-Dunn et al., 2009; Goldenberg et al., 2000; Hillier et al., 1988). Pregnant women with systemic malaria infections were shown to have inflammatory responses that correlate with
preterm birth; however, it has been shown that the majority of preterm birth is due to bacterial intrauterine infections (Jefferson, 2012; Romero et al., 2001). The accepted theory is that vaginal bacteria ascend through the cervix to cause infection and inflammation (Jefferson, 2012).

Chorioamnionic tissue sampled from women undergoing C-section for spontaneous labor indicated that 75% of tissue was culture positive (Goldenberg et al., 2008; Jefferson, 2012). Bacteria isolated from preterm tissue and amniotic fluid has been found in the vagina as well (Romero et al., 1989). While oral flora can be found in amniotic fluid, the general notion is that the oral species first enters the vagina through oro-genital contact and then begins ascension in the uterine cavity to invade the amniotic cavity rather than hematogenous spread (Figure 1.1) (Jefferson, 2012; Kim et al., 2009). The association of bacterial vaginosis, otherwise known as BV, and preterm birth therefore is not likely due to vaginal pathology, but ascension of BV-associated bacteria from the vagina into the uterine cavity (Jefferson, 2012).

During gestation, the cervical mucus plug becomes more viscous due to the presence of progesterone to form a mechanical barrier. Additionally, alpha and beta defensins, antimicrobial compounds like lactoferrin, secretory IgA, and host immune system factors like complement are present within the cervical mucus plug (Jefferson, 2012). Once bacteria have crossed the cervical mucus plug, they can remain in the choriodecidual space or invade the chorion, cause chorioamnionitis, traverse the fetal membranes or enter the amniotic cavity and are capable of eliciting a chronic infection that results in an inflammatory host response which can lead to preterm birth (Goldenberg et al., 2000; Jefferson, 2012). Triggering the inflammatory response mediates the release of tumor necrosis factor alpha (TNF-α), IL-6, IL-8, IL-1a, IL-1b and granulocyte colony-stimulate factor (Estrada-Guiterrez et al 2010; Goldenberg et al., 2000; Jefferson, 2012). The release of the cytokines stimulates secretion of other cytokines,
Figure 1.1: Schematic representation of bacterial ascension into the amniotic cavity.

Bacterial ascension occurs at the vagina where select bacteria are able to traverse the fetal membranes and enter into the amniotic cavity to cause infection. Adapted from (Kim et al., 2009) and reproduced with permission of publishing journal.
prostaglandins, matrix metalloproteinases, which weaken the fetal membranes and eventually lead to PPROM.

Therefore it is hypothesized that bacteria that contain certain virulence factors can contribute to preterm birth and/or PPROM. The most frequently isolated BV-associated bacteria accompanying infection and preterm birth are *Mycoplasma hominis*, *Ureaplasma urealyticum*, *Gardnerella vaginalis*, *Sneathia amnii*, and *Bacteriodes* (Goldenberg et al., 2000; Goldenberg et al., 2008; Jefferson, 2012). Since numerous bacteria associated with microbial invasion of the amniotic cavity (MIAC) are difficult to culture, the utilization of molecular techniques, like PCR, has aided research of bacterial species that are present in amniotic fluid and fetal membranes (Goldenberg et al., 2008).

**Bacterial Vaginosis**

Bacterial vaginosis, otherwise known as BV, is a complex polymicrobial disorder that is caused by a chemical and biological imbalance in the normal flora of the vagina. The virulence factors associated with BV are poorly understood and it is frequently misdiagnosed as trichomoniasis or a yeast infection. BV is associated with an overgrowth of commensal anaerobic bacteria where none is the clear pathogen. It is the most common vaginal infection worldwide and affects 29% of women in the United States (Allsworth et al., 2007; Fluery, 1981). It is associated with preterm birth, low birth weight, increased risk of acquiring HIV-1 and other sexually transmitted disease, pelvic inflammatory disease, and post-operative infections (Allsworth et al., 2011; Lambert et al., 2013; Lamont et al., 2011; Madden et al., 2012; Schellenberg et al., 2012). While the etiology is unknown, risk factors for BV include; race, douching, IUDs, and sexual activity (Madden et al., 2012).
BV is characterized by a decrease in *Lactobacillus* species, particularly those that are able to produce hydrogen peroxide; the depletion of *Lactobacillus* allows for an increase in anaerobic bacteria (Muzny et al., 2013). *Lactobacillus spp* are gram positive rod shaped bacterium that are facultative anaerobes and are the primary colonizer of the vagina. In addition to producing hydrogen peroxide, which inhibits the growth of other bacterial species, *Lactobacillus* produce lactic acid by fermentation lowering the overall pH (< 4.5) of the vagina to favor the vaginal microbiota composition towards acidophiles (Fettweis et al., 2012). A shift in *Lactobacillus* composition allows for anaerobic bacteria like *Gardnerella vaginalis*, *Prevotella bivia*, *Mycoplasma hominis*, *Mobiluncus spp.*, *Atopobium vaginae*, *Fusobacterium spp.*, and *Sneathia amnii* to flourish (Harwich et al., 2012; Munzy et al., 20113).

Research has indicated that race plays an important role for BV and consequently preterm birth. While 10-20% of Caucasian women experience BV, African American women have a 30-50% rate. This racial disparity is also seen in preterm birth; African American women have a twofold increased risk of experiencing preterm birth and a threefold risk of extremely preterm birth (Sobel, 2005). While both races are colonized by *Lactobacillus*, there is a difference in colonization that attributes to BV and preterm birth. There are six species of *Lactobacillus* that colonize the vagina; *Lactobacillus crispatus*, *Lactobacillus iners*, *Lactobacillus jensenii*, *Lactobacillus gasseri*, *Lactobacillus vaginalis* and *Lactobacillus johnsonii* (Fettweis et al., 2014; Jefferson, 2012). These species of *Lactobacillus* vary in their stability and their ability to protect the vagina from BV associated bacterial colonization (Tamrakar et al., 2007). More stable species are not easily displaced from changes in pH, hormonal changes, nutrient sources and semen deposition. Women of European decent are predominantly colonized with *L. crispatus*, which is more stable and protective against BV. Women colonized with *L.crispatus* have a
fivefold decreased risk of developing BV. African American women however, tend to be colonized with *L. iners* which is less stable and less protective than *L. crispatus* and therefore women colonized with *L. iners* have a tenfold increased risk of developing BV (Fettweis et al., 2014; Verstraelen et al., 2009). Furthermore, African American women have a higher diversity of microflora that is associated with microbial invasion of the amniotic cavity and preterm birth (Fettweis et al., 2014).

Another associated risk factor of BV is douching, since it mechanically causes an imbalance in normal flora through physical and chemical irritants; douching solutions often contain surfactants that lyse the normal flora and allow for other bacterial growth. Additionally, the use of intrauterine devices is a risk factor for BV. It is thought that the presence of the IUD string in the vagina allows for colonization of anaerobic bacteria associated with BV (Madden et al., 2012). Finally, for reasons not completely understood, having multiple sex partners increases the risk for BV especially women who have sex with women, WSW, have a higher BV prevalence than women who do not (Koumans et al., 2009; Munzy et al., 2013).

Clinical diagnosis of BV is done either by the Amsel criteria or by the Nugent scoring system. For the Amsel criteria, three of the four criteria characterizes a positive diagnosis of BV: a thin, homogenous grey-white discharge; pH greater than 4.5; presence of clue cells on wet mount; positive whiff amine test where a fishy smell is present with 10% KOH is added to sample of discharge (Amsel et al., 1983; Jefferson, 2012; Srinivasan et al., 2012). The Nugent system utilizes the gram stain, which is the gold standard of BV diagnosis (Srinivasan et al., 2012). The scoring system is based on the presence of lactobacilli, curved rods and small gram negative rods (Nugent et al., 1991). In the majority of cases, women who test positive for BV are asymptomatic, so it is unclear why or what factors promote symptoms (Srinivasan et al., 2012).
Treatment for BV includes either oral or topical administration of clindamycin or metronidazole; however, relapse among women is very high (Hay, 1989; Srinivasan et al., 2012). This complex polymicrobial syndrome includes many species of bacteria, like *Sneathia amnii*, that play a role in both vaginal and gestational health.

*Sneathia amnii* strain Sn35

*Sneathia amnii*, formerly classified under *Leptotrichia*, is an anaerobic gram negative, non-motile bacterium that is part of the normal urogenital flora of women (Harwich et al., 2012). This species, however, is associated with serious medical complications such as; preeclampsia, spontaneous abortion, bacterial vaginosis, preterm birth, preterm premature rupture of the fetal membranes and post-partum bacteremia in both mother and neonate (Bachy et al., 2011; Harwich et al., 2012). The uterine cavity and amniotic fluid are thought to remain sterile throughout gestation; however bacterial infections can occur and *S.amnii* is frequently isolated from the amniotic cavity. (Harwich et al., 2012) *S.amnii* infiltration of the fetal membranes and amniotic cavity can lead to chorioamnionitis and/or amnionitis (Han et al., 2009). Other studies have shown that *Sneathia* colonization of the male urogenital tract can occur, suggesting that this organism can be sexually transmitted (Nelson et al., 2010). Additionally, through the Vaginal Microbiome Project at VCU, mid-vaginal isolates of 736 women were sequenced and *Sneathia* was found to be present in 43.4% of women with an abundance threshold of 0.1% of total reads (Harwich et al., 2012). *S.amnii* is a fastidious bacterium that requires blood or human serum for growth. Growth can occur on chocolate agar and BHI agar with supplemented blood; however it cannot survive on Brucella Sheep’s blood agar, making this organism human specific (Harwich et al., 2012).
Strain Sn35, a mid-vaginal isolate, was from an African American woman in her early 20’s enrolled in the Vaginal Microbiome Project at VCU who had extremely preterm birth at 26 weeks (Harwich et al., 2012; Fettweis et al., 2011). Sequencing and identification was done using the V1-V3 regions of the 16S rRNA gene and was shown that *S. amnii* has high homology to another vaginal bacterium, *Sneathia sanguinegens*; which often co-occurs in the vagina (Harwich et al., 2012). *Sneathia amnii* has the smallest genome of the Fusobacteriaceae family with a genome size of approximately 1.34 Mbp and has 28% GC content. *S. amnii* has 1,282 protein encoding genes, making up 92% of the genome and has at least 110 overlapping genes (Harwich et al., 2012).

With scanning electron microscopy, *S. amnii* was found to have a variety of morphologies; it has been seen as a long rod, short rod, and both amorphous rods and cocci. The shorter morphologies have been identified in other bacterial species such as, *Streptococcus moniliformis* and are referred to as “L forms” where the cell wall is deficient (Dienes, 1947). “L forms” of *S. amnii* were typically found in older cultures and active cultures were dominated by bacillus (Harwich et al., 2012). Occasionally, the rod shaped bacterium exhibits bulbous protrusions at the ends of the rods (Harwich et al., 2012).

Preliminary research on *Sneathia amnii* identified that it has weak alpha hemolysin activity, and has the ability to ferment glycogen, maltose and glucose while is unable to ferment starch, mucin, galactose, sucrose or fructose (Harwich et al., 2012). Similar to other pathogenic bacteria, *S. amnii* does not have enzymes necessary for synthesis of most amino acids; however, it does have enzymes required to convert L-amino acids to D-amino acids and the enzymes necessary for the conversion of L-aspartate to fumarate, L-asparagine and oxaloacetate. Additionally, genes required for de novo amino acid biosynthesis have not been identified in the
species; however, salvage pathways for purine and pyrimidine synthesis are present (Harwich et al., 2012). A putative DNA restriction modification system was identified; however, it lacks a complete system of genetic competence (Harwich et al., 2012). Furthermore, the sequence of *S. amnii* identified a putative Type II secretion system, and has genes related to ion transport and multi-drug/lipid/protein pumps (Harwich et al., 2012).

While most gram negative bacteria are inherently vancomycin resistant, *Sneathia amnii* is vancomycin sensitive. It is also sensitive to metronidazole, a common drug used to treat bacterial vaginosis; however it is resistant to nafcillin, ciprofloxacin, and tetracycline. Women who are treated for BV with metronidazole during gestation still have a twofold increased risk of preterm birth. During BV, biofilm formation on the vaginal epithelium can occur which can lead to antibiotic resistance; *S. amnii* can become resistant or tolerant of antibiotic measures if associated with a biofilm (Harwich et al., 2012; Swidsinski et al., 2008). While this species is part of the normal urogenital microflora of women little is known about this organism; however, *Sneathia amnii* seems to be a significant opportunistic pathogen that plays a role in vaginal health, MIAC, and preterm birth.

**Research Objectives**

*Sneathia amnii* is part of the vaginal normal flora; however, it has serious clinical implications such as bacterial vaginosis and preterm birth. Since bacterial vaginosis is a risk factor for preterm birth and the greatest risk factor for preterm birth is having previously had preterm birth, it is essential to understand the ability of species like *S. amnii* to play a part in both clinical pathologies. Additionally, most extreme preterm births arise from intrauterine infections and inflammation and pose a more serious threat to the neonate. While *Sneathia amnii* can be
found in 25% of amniotic fluid samples of patients presenting with preterm birth, the ability of *S. amnii* to traverse the cervical mucus plug and enter into the amniotic cavity remains unknown. Recent studies in our lab indicating that *S. amnii* is cytotoxic to ME-180 cervical epithelial cells show a small glance of the virulence potential of *S. amnii*. The goal of this project is to utilize a *Sneathia amnii* mid-vaginal strain isolated from a woman who gave birth at 26 weeks to better elucidate details of *S. amnii*’s virulence determinants and its ability to traverse the fetal membranes. First, to characterize protease interactions with fetal membrane substrates and secondly, to explore bacterial ascension into the uterine cavity. These objectives have resulted in the characterization of a novel protease expressed by *S. amnii*, a potential virulence factor.
CHAPTER 2
Materials and Methods

Bacterial Strains and Growth Conditions

The *S. amnii* and *E.coli* strains used throughout this study are listed in Table 1. *Sneathia amnii* was maintained under anaerobic conditions at 37°C on SBHI blood plates. SBHI blood plates were composed of brain heart infusion (BHI) agar (Oxoid, Cheshire, England) + 10% gelatin + 0.5% starch + 0.5% dextrose + 5% yeast and 10% female pooled human blood. Liquid media was composed of SBHI broth and pooled human serum (MP Biomedicals, Santa Ana, CA). SBHI broth was composed of brain heart infusion broth + 10% gelatin + 0.5 % starch + .5% dextrose + 0.5% yeast. After autoclaving, 10% human serum was added to the SBHI broth and then filter sterilized prior to *S.amnii* inoculation. Anaerobic conditions were met through the Shell Lab BacBasic Anaerobic Chamber with anaerobic gas mixture; 5% carbon dioxide, 5% hydrogen and 5% nitrogen. *Escherichia coli* strains were grown aerobically at 37°C on Luria-Bertani broth (LB) agar plates containing the appropriate antibiotic. *E.coli* liquid cultures were grown in LB, incubated aerobically at 37°C, 200 rpm. Appropriate antibiotics were incorporated at the following concentrations 100µg ampicillin (Amp)/ml for *E. coli*, 35µg chloramphenicol (Cm)/ml for *E. coli*. 

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Table 1. Bacterial Strains

<table>
<thead>
<tr>
<th>Strain</th>
<th>Description</th>
<th>Source</th>
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<tbody>
<tr>
<td>CH3 Blue</td>
<td>Chemically competent cells derived from <em>E. coli</em> K12 cloning strain (Amp(^r) when transformed with plasmid)</td>
<td>Bioline (Tauton, MA)</td>
</tr>
<tr>
<td>BL21-CodonPlus (DE3)-RIL</td>
<td>Chemically competent cells derived from <em>E. coli</em> BL21 (Amp(^r) when transformed with plasmid, Cm(^r))</td>
<td>Bioline (Tauton, MA)</td>
</tr>
<tr>
<td>BL21+pET32xT+peg1</td>
<td>BL21-CodonPlus(DE3)RIL cells containing peg1 from Sn35 <em>in trans</em> using pET32xT (Amp(^r), Cm(^r))</td>
<td>This study</td>
</tr>
<tr>
<td>Sn35</td>
<td>Mid-vaginal isolate from woman in her early 20’s presenting with extreme preterm birth at 26 weeks</td>
<td>Vaginal Microbiome Consortium, VCU</td>
</tr>
<tr>
<td><em>Lactobacillus crispatus</em></td>
<td>Mid-vaginal Isolate</td>
<td>Vaginal Microbiome Consortium, VCU</td>
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Polymerase chain reaction (PCR) and product purification

For all PCR amplifications of DNA, general conditions were used with some modifications that were dependent upon the DNA template in use, the melting temperature ($T_m$) of the primer, and the size of the product being amplified. Reactions were composed of a 50µL total volume, 1 µL 10mM dNTPS, 1 µL template, 1µL of both forward and reverse primer (10pmol dilutions in sterile water), 1µL Phire II polymerase (Thermo Scientific, Waltham, MA), 10 µL 5X Phire buffer and 25 µL of sterile water. Primers utilized in this study are listed in Table 2 and primer design was based off S.amnii isolate Sn35. Thermocycler reaction conditions were programmed as the following: heated lid at 105 ºC, initial denaturation 99ºC for 2 minutes, followed by 31 cycles of denaturation at 94ºC for 30 seconds, annealing at 60ºC for 30 seconds, and extension at 68ºC for 2 minutes (+2 minutes, depending upon size of amplified region), and an additional cycle of final extension at 68ºC for 10 minutes. PCR products were then held at 4ºC in thermocycler or placed in -20ºC freezer until ran on DNA gel electrophoresis and subsequently purified using QIAquick Gel Extraction kit (Qiagen, Valencia, CA) using manufacturer’s instructions.

DNA Agarose Gel

1% DNA agarose gels were made with 1X Tris-acetate- EDTA (TAE) buffer (Life Technologies, Waltham, MA), agarose, and 0.15µg/mL of ethidium bromide. 5X Loading DNA Buffer (Bioline, Taunton, MA) was added to DNA samples for a final concentration of 1X. Agarose gels were electrophoresed at 124 volts in 1X TAE buffer and examined under UV light. The size of the product was determined against Bioline Hyperladder 1kb weight markers (Bioline, Taunton, MA).
<table>
<thead>
<tr>
<th>Primer Name</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>PCR4TOPO T3</td>
<td>5′-ATTAACCCTCACTAAAGGGA-3′</td>
</tr>
<tr>
<td>PCT4TOPO T7</td>
<td>5′-TAATACGACTCACTATAGGG-3′</td>
</tr>
<tr>
<td>Sn35Coll2FWD</td>
<td>5′-GGCCGGGATCCTTTTCAGATGCTCGTATTTGATAAGAATGAAAGGA-3′</td>
</tr>
<tr>
<td>Sn35Coll2REV</td>
<td>5′GGCCGGCTCGAGGTTATAGATACCTCGAGATTCCACAATG-3′</td>
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</tbody>
</table>
Plasmid DNA purification and modification

Plasmid purifications were done using the QIAprep Spin Miniprep Kit (Qiagen, Valencia, CA) using the manufacturer’s instructions. Single colony transformants of *E. coli* were grown overnight in 3 ml LB with 100µg ampicillin/ml. Plasmid modification/ DNA manipulation restriction enzymes, buffers, and BSA were purchased from New England Biolabs (Ipswich, MA) and were used according to the manufacturer’s protocols. Plasmid restriction digests were done using BamHI and XhoI, and the 5’ phosphoryl groups were removed using Antarctic Phosphatase (New England Biolabs, Ipswich, MA) to prevent re-ligation of plasmid.

Plasmid Construction

DNA fragments were subcloned into PCR4-TOPO (Life Technologies, Waltham, MA) using manufacturer’s recommended procedure and were transformed into CH3 Blue chemically competent *E.coli* (Bioline, Taunton, MA). After incubation overnight on LB agar with ampicillin, single colonies were selected and grown overnight in LB broth with ampicillin. Plasmids were purified using QIAprep Spin Miniprep Kit (Qiagen, Valencia, CA) and digested with New England Biolabs restriction enzymes and buffers according to the manufacturer’s recommendations. Digests were electrophoresed on DNA agarose gel. Bands that were the appropriate insertion size, based off Hyperladder 1kb (Bioline, Taunton, MA) were gel extracted and purified. Ligation of purified DNA fragment insert into digested pET vectors was done using Ready-To-Go T4 DNA Ligase and Ligase Buffer (Amersham Biosciences, Piscataway, NJ). Ratio used for ligation was a 3µL insert to 1µL plasmid, at 16ºC for 1 hour.
Transformation of chemically competent *E. coli*, CH3 Blues

A vial of chemically competent *E. coli* cells was thawed on ice, 3µL of pET32xT plasmid ligation reaction was added to the cells and reaction was incubated on ice for 30 minutes. The tube was placed in a 42°C water bath for 30 seconds, immediately afterward 200µL of warmed SOC media (85.5mM NaCl, 2.5mM KCl, 10mM MgCl₂, 2% tryptone, 0.5% yeast extract, and 20mM glucose) was added to the reaction. The cells recovered on a shaker at 37°C for 1 hour. Cells were plated on LB plates containing 100µg/mL of ampicillin.

Transformation of chemically competent *E. coli*, BL21 DE3 RIL

A vial of electrocompetent *E. coli* cells was thawed on ice; 0.7µL of β-mercaptoethanol was added to the cells. The reaction was incubated on ice for 10 minutes and gently swirled every 2 minutes before adding 3µL of ligated plasmid and incubating on ice for 10 minutes. SOC media (85.5mM NaCl, 2.5mM KCl, 10mM MgCl₂, 2% tryptone, 0.5% yeast extract, and 20mM glucose) was pre-warmed in a 42°C water bath. The reaction was heat pulse in a 42°C water bath for 45 seconds and 250µL of warmed SOC media was immediately added to the reaction. The reaction was incubated for 1 hour at 37°C, shaking at 200 rpm.

Induction of BL21 DE3 RIL- Coll2 bacterial cells

Once plasmid was sequenced to confirm proper insertion of template into pET32xT, plasmid containing bacterial cells were inoculated in 6 mL overnight cultures containing LB, 100µg/ml ampicillin (Amp) and 35µg/ml chloramphenicol (Cm) at 37°C on shaker. The overnight cultures were diluted 1:200 in 1L of fresh LB containing 100µg/ml ampicillin (Amp) and incubated at 37°C for 4 hours. The culture was induced with 1mM IPTG and incubated for
an additional 2 hours at 37°C. Prior to induction, 500µL of the culture was collected, the bacteria were obtained through centrifugation. After 2 hour induction, 500µL of culture was collected, the bacteria were obtained through centrifugation. Uninduced and induced controls were electrophoresed on a NuPAGE SDS-PAGE Gel system (Invitrogen, Grand Island, NY) at 200V for 30 minutes in 1X NuPAGE Running Buffer. Lanes were stained with Imperial Protein Stain (Thermo Scientific, Waltham, MA) for 1 hour and then destained with water until bands were clearly visible. Bacterial cells were collected from cultures by centrifugation at 8,000 rpm at 4°C for 15 minutes. Pellets were suspended in 25 mL of 1X sterile PBS (Invitrogen, Grand Island, NY) and were subsequently harvested by centrifugation at 4,000 rpm at 4°C for 15 minutes. The pellet was frozen at -80°C.

Polyacrylamide gel electrophoresis

All protein gels used the NuPAGE SDS-PAGE Gel system (Invitrogen, Grand Island, NY). Samples were mixed using a 30:10:4 ratio, 30 µL of sample, 10 µL of Novex NuPAGE LDS Sample Buffer 4X (Life Technologies, Waltham, MA) and 4 µL of NuPAGE Sample Reducing Buffer (Invitrogen, Grand Island, NY). Samples were boiled for 10 minutes and loaded into the wells of a NuPAGE 4-12% Bis-Tris 1.0mm Gel (Invitrogen, Grand Island, NY). Gel electrophoresis was for 30 minutes at 200Vs in 1X NuPAGE Running Buffer. Gels were then stained with Imperial Protein Stain for one hour and destained in water until bands were visible. Peptide size was determined using Precision Plus Protein Kaleidoscope Standard (BioRad, Hercules, CA).
Protein Purification

The bacterial pellet from 1 L induction was thawed on ice and was resuspended in a solution of 25 mL 50mM NaP, 300mM NaCl buffer and one Complete, EDTA Free Protease Inhibitor Cocktail Tablet (Roche, Indianapolis, IN). The resuspended pellet was then lysed by passing through a French Pressure Cell Press (Thermo Scientific, Waltham, MA) 4 times. The solution was then lysed by sonication on ice 6 times with 30 second sonication periods and 1 minute rests between each pulse, bubbles were not introduced. The lysed solution was then centrifuged at 12,000 rpm at 4°C for 20 minutes. The lysate was filtered with 0.45µm filter and immediately loaded onto a 3 mL HisPur Cobalt Resin (Thermo Scientific, Waltham, MA) column. After flow through collection, the column was subjected to 6 washes consisting of 10 mL of 50mM NaP + 300mM NaCl and 10mM imidazole. Washes were collected and incubated on ice, wash 7 consisted of 10 mL 50mM NaP + 300mM NaCl and 20mM imidazole. Final elution was consisted of 3 mL of 50mM NaP + 300mM NaCl and 250mM imidazole. The Final elution was performed in 3, 1 mL washes where 1 mL of elution buffer was rocked gently in resin for 5 minutes before eluate was collected. A280 reading was collected on elute. The Eluate was dialyzed overnight at 4°C against 3L of 20mM Tris-HCl pH 8.0 with 150mM NaCl using a 3,500 MWCO Slide-A-Lyzer Dialysis Cassette G2 (Thermo Scientific, Waltham, MA).

Thrombin Cleavage

The protein was removed from overnight dialysis cassette, A280 reading was recorded. Thrombin CleanCleave Kit (Sigma Aldrich, St. Louis, MO) was used for removal of the purified protein’s Thio-HIS tag. Thrombin cleavage was performed according to the manufacturer’s instructions except the following deviations; 150 mM of CaCl₂ was added to purified protein,
thrombin beads, and cleavage buffer. Cleavage reaction was rocked at 25°C for 2 hours. The reaction was centrifuged at 4,000 rpm for 2 minutes; supernatant was collected and ran through HisPur Cobalt Resin (Thermo Scientific, Waltham, MA) column. Flow thru was collected and centrifuged at 4,000 rpm for 2 minutes, elute was collected. The 3 mL final collection of purified protein was then diluted with 12 mL sterile water and was concentrated using Amicon Ultra-4 Centrifugal Filters, 30k MWCO (Millipore, Billerica, MA) at 4,000 rpm until 500µL of elute remained. Thrombin cleaved purified elute was either stored at -80°C or immediately used.

Casein/Mucin/ Gelatin Zone of Activity Plates

General conditions were used for all zone of activity plates, with the exception of the substrate used; casein, gelatin from cold water fish skin (Sigma, St. Louis, MO), and mucin (Sigma, St. Louis, MO). 1% agarose, 1% substrate plates were made by dissolving agarose in 1X Tris-acetate-EDTA (TAE) buffer (Life Technologies, Waltham, MA). The agarose was heated until boiled; substrate was added and poured in 35x10mm plates (Fisher Scientific, Pittsburg, PA). After plates were solidified, 10 µL of sample were pipetted 1 inch apart. Samples included; 1X sterile PBS (Life Technologies, Waltham, MA), sterile water, uncleaved purified protein, cleaved purified protein, proteinase K (1:10 dilution, in sterile water) (Qiagen, Valencia, CA). Plates were placed in 37°C incubator overnight. The plates were observed to identify zones of activity.

Biotinylation of Collagen Type I and Collagen Type IV

Both collagen type IV (Sigma Aldrich, St. Louis, MO) and collagen type I (Sigma Aldrich, St. Louis, MO) were from human placental tissues. Collagen types were dissolved in 0.5
M acetic acid to achieve 2mg/ml. Solutions were gently vortexed until completely dissolved, 242.5 µL of 5M NaOH was added to neutralize the solutions. Appropriate calculations were done to identify the amount of biotin necessary for each collagen type. EZ-link Sulfo-NHS Biotin kit (Thermo Scientific, Waltham, MA) was used to biotinylate collagen type I and IV. 10mM Biotin was prepared with ultrapur water; 71.9 µL of biotin solution was added to each collagen solution. Reaction was then incubated on ice for 2 hours. Reaction clean-up was done with Zeba Spin Desalting Columns (Thermo Scientific, Waltham, MA) according to the manufacturer’s instructions. Biotinylated collagen type I and IV were aliquoted in 100 µL in Siliconized Polypropylene Extended Capacity Tubes (Sigma Aldrich, St. Louis, MO) and were stored at -80ºC. Dot blot was performed on biotinylated collagen type I and IV to verify Biotinylation. Collagen type I and IV were spotted onto Pure Nitrocellulose Blotting Membrane (Pall, Pensacola, FL) in 2 µL aliquots in various dilutions. Membrane was left to dry and blocked in 5% non-fat milk in PBS for 1 hour. Afterward 1.1 µL of Streptavidin- HRP (Thermo Scientific, Waltham, MA) was added to the membrane and shook for 1 hour at 25ºC. Membrane was washed 3 times in 0.005% PBST and subsequently incubated with ECL Prime Western Blotting Detection Reagent (GE Healthcare, Buckinghamshire, UK) and developed.

**Collagenase Assay**

Samples used in collagenase assay included; cleaved purified protein, 1X sterile PBS (Life Technologies, Waltham, MA), 1:10 dilution Proteinase K solution (Qiagen, Valencia, CA) and overnight cultures of *Sneathia amnii* isolate, Sn35. Biotinylated collagen type I and IV were diluted to a 1:25 working concentration in 1X sterile PBS (Life Technologies, Waltham, MA), 10 µL of sample was incubated with collagen in Siliconized Polypropylene Extended Capacity
Tubes (Sigma Aldrich, St. Louis, MO) overnight at 37°C. Samples were boiled for 10 minutes and were loaded NuPAGE 4-12% Bis-Tris 1.0mm Gel (Invitrogen, Grand Island, NY). Gel electrophoresis ran for 30 minutes at 200Vs in 1X NuPAGE Running Buffer. After electrophoresis, gel was rinsed with deionized water. A 0.45µm PVDF membrane (Invitrogen, Grand Island, NY) was soaked in 100% methanol while sponges and Criterion Blotter Filter Paper (BioRad, Hercules, CA) were presoaked in 1X NuPAGE Transfer Buffer (Invitrogen, Grand Island, NY) with 10% methanol. Sponges, filter paper, gel and membrane were stacked upon one another in the Invitrogen transfer system so the arrangement was; 4-5 sponges, 1filter paper, gel, PVDF membrane, 1 filter paper, and finally 4-5 sponges, more sponges allowed for tight contact between the PVDF membrane and the gel. The transfer was run at 45V for 90 minutes on ice in 1X NuPAGE transfer buffer with 10% methanol. Membrane was blocked overnight at 4ºC in 5% nonfat milk in 1X PBS and subsequently probed with 1:15,000-diluted Streptavidin-HRP (Thermo Scientific, Waltham, MA) and 1X PBST (PBS + 0.05% Tween 20) for 1 hour, rocking at 25ºC. The membrane was washed 3 times for 4 minutes with 1X PBST and developed using ECL Prime Western Blotting Detection Reagent (GE Healthcare, Buckinghamshire, UK), where membrane was immersed in solution for 1 minute, covered in plastic wrap and exposed to Blue Devil Interleaved Autoradiography Film (Genesee Scientific) and developed.

Fibronectin assay

Samples were incubated with fibronectin from Human Plasma (Sigma Aldrich, St. Louis, MO) at 37°C overnight. Fibronectin was diluted 1:10 with sterile 1X PBS (Invitrogen, Grand Island, NY) and each incubated reaction had 10µL of fibronectin. Fibronectin was incubated in a
variety of salt solutions; 10mM CaCl$_2$ + 15mM ZnCl$_2$, 15mM ZnCl$_2$ +10 µL of cleaved purified protein, 2mM CaCl$_2$ + 5mM ZnCl$_2$, 2mM CaCl$_2$ + 15mM ZnCl$_2$ + 10 µL of cleaved purified protein, 5mM ZnCl$_2$ +10µL of cleaved purified protein diluted 1:5, 5mM ZnCl$_2$ +10 µL of cleaved purified protein diluted 1:25, 5mM ZnCl$_2$ +10 µL of cleaved purified protein diluted 1:125. Samples were mixed using a 30:10:4 ratio, 30 µL of sample, 10 µL of Novex NuPAGE LDS Sample Buffer 4X (Life Technologies, Waltham, MA) and 4 µL of NuPAGE Sample Reducing Buffer (Invitrogen, Grand Island, NY). Reactions were boiled for ten minutes and then loaded onto a NuPAGE 4-12%Bis-Tris 1.0mm Gel (Invitrogen, Grand Island, NY). Gel electrophoresis ran for 30 minutes at 200Vs in 1X NuPAGE Running Buffer where band size was determined using Precision Plus Protein Kaleidoscope Standard (BioRad, Hercules, CA). Gels were then stained with Imperial Protein Stain (Thermo Scientific, Waltham, MA) for one hour and destained in water until bands were visible. Degradation of fibronectin was evaluated based on complete disappearance of fibronectin peptide band, 220kDa.

**Laminin Assay**

Samples were incubated with laminin from Human Placental Liquid (Sigma Aldrich, St. Louis, MO) at 37ºC overnight. Laminin was diluted 1:10 with sterile 1X PBS (Invitrogen, Grand Island, NY) and each incubated reaction had 20µL of laminin. Laminin was incubated in a variety of salt solutions; 10mM CaCl$_2$ + 15mM ZnCl$_2$, 15mM ZnCl$_2$ +10 µL of cleaved purified protein, 2mM CaCl$_2$ + 5mM ZnCl$_2$, 2mM CaCl$_2$ + 15mM ZnCl$_2$ + 10 µL of cleaved purified protein, 5mM ZnCl$_2$ +10 µL of cleaved purified protein diluted 1:5, 5mM ZnCl$_2$ +10 µL of cleaved purified protein diluted 1:25, 5mM ZnCl$_2$ +10 µL of cleaved purified protein diluted 1:125. The samples were mixed using a 30:10:4 ratio, 30 µL of sample, 10 µL of Novex
NuPAGE LDS Sample Buffer 4X (Life Technologies, Waltham, MA) and 4 µL of NuPAGE Sample Reducing Buffer (Invitrogen, Grand Island, NY). The reactions were boiled for ten minutes and then loaded onto a NuPAGE 4-12% Bis-Tris 1.0 mm Gel (Invitrogen, Grand Island, NY). Gel electrophoresis ran for 30 minutes at 200 Vs in 1X NuPAGE Running Buffer where band size was determined using Precision Plus Protein Kaleidoscope Standard (BioRad, Hercules, CA). Gels were then stained with Imperial Protein Stain (Thermo Scientific, Waltham, MA) for one hour and destained in water until bands were visible. The degradation of laminin was evaluated based off complete disappearance of laminin peptide bands at 50 kDa and 130-160 kDa.

**Coagulase Assay**

BBL Coagulase Plasma Rabbit (BD, Franklin Lakes, NJ) vials were rehydrated in sterile purified water; one vial was aliquoted into six separate tests. Reactions with coagulase test included; 1X Sterile PBS (Invitrogen, Grand Island, NY), live overnight culture of *Sneathia amnii* isolate SN35, cleaved purified protease + 5 mM ZnCl₂, cleaved purified protease, 5 mM ZnCl₂, 2 mM CaCl₂. The reaction vials were gently mixed and incubated in 37°C water bath. The tubes were examined by tipping the tube to identify clotting after 4 hours and then again at 24 hours to identify any weak enzyme production.

**Turbidity Assay**

A 2 mL overnight culture of SN35 was grown and subcultured into 10 mL of fresh media. Culture grew for 6 hours to reach exponential stage and was removed from the anaerobic chamber. Culture was divided into two 5 mL cultures. One 5 mL culture was centrifuged at
4,000rpm for 4 minutes. Supernatant was removed and pellet was washed with sterile water and centrifuged. The pellet was resuspended in 4 mL of sterile water and aliquoted into two 2 mL tubes. Non-centrifuged culture was separated and aliquoted into two 2 mL tubes. The initial sample optical density readings, OD$_{600nm}$, were determined using a spectrophotometer; sterile SBHIs and sterile water were used as controls. Afterward, 100µL of cleaved purified protease was added to 1 tube of Sn35 in SBHI media and 1 tube of Sn35 in sterile water; 100µL of dialysis buffer was added to other Sn35 tubes. The optical density was determined immediately, then after 30 minutes, 1 hour, 2 hours, and 24 hours.

Collagen Type IV Biofilm Assay

    Cell Adhesion Human Collagen Type IV Strips (Millipore, Temecula, CA) were rehydrated in 200 µL of 1X sterile PBS (Invitrogen, Grand Island, NY) for 15 minutes at room temperature; PBS was removed from the strips. Overnight culture of Sn35 was grown and diluted until OD$_{600nm}$ was 1. The strip wells were inoculated as such; SBHI alone, 100 µL of 1:10 dilution Sn35 OD=1, 100 µL of 1:100 dilution Sn35 OD=1, 100 µL of 1:10 dilution Sn35 OD=1 boiled for 5 minutes, 100 µL of cleaved purified protease, 100 µL of heat killed Sn35 + cleaved purified protease, SBHI alone, and last lane was left as BSA. The collagen strips were kept under anaerobic conditions at 37ºC overnight and suspensions were discarded and well strips were washed gently 2-3 times with 200 µL of PBS. The wells were stained with 100 µL of safranin; adherence was determined by visualization of the degree of safranin staining.
Edman Degradation

The protein was purified and cleaved using methods mentioned above. Thrombin treated protein was run on a 4-12% Bis-Tris 0.1mm NuPAGE gel for 30 minutes at 200V. Gel was electroblotted onto PVDF membrane using methods mentioned previously. PVDF membrane was then stained using Imperial Protein Stain (Thermo Scientific, Waltham, MA) for 1 hour. The membrane was destained for 15 minutes using 15% methanol/10% acetic acid and was rinsed under deionized water for 4 hours. The band of interest was excised and placed in microcentrifuge tube with deionized water. Edman degradation was performed by Iowa State University Protein Facility; membrane was washed 6 times in deionized water and sample was analyzed by 494 Procise Protein Sequencer/140C Analyzer from Applied Biosystems, Inc.

Bacteriolyssin Assay

A 10% SDS PAGE separating gel containing 0.003g of dry Micrococcus luteus was poured, allowed to solidify and a 4% SDS PAGE stacking gel was prepared and poured on top of the separating gel. After the gels polymerized, cleaved purified protein was mixed with 5X Loading buffer and was electrophoresed at room temperature at 175V for 40 minutes in 1X SDS PAGE running buffer. Gels were rinsed for 1 hour under a slow stream of sterile water for one hour and then immersed in 100mM phosphate buffer pH 7.0 + 10mM MgCl2. Gels remained in buffer, rocked overnight at 37ºC. Gels were rinsed with sterile water and subsequently stained with Imperial Protein Stain (Life Technologies, Waltham, MA) for one hour. Gels were destained in sterile water until banding patterns were visible.
Antiserum Production

Cleaved purified protein (> 1.5mg/ml) was produced using methods mentioned previously and shipped to New England Peptides. Prior to immunization, 5 mL of pre-immune sera was collected from one rabbit (New Zealand White- SPF), the rabbit was immunized with protein 3 times. Boosts occurred at day 14 and 28, bleeds occurred on days 35 and 40. Upon arrival, sera were aliquoted into cryovials and stored at -80°C.

Antisera Detection and Subsequent Westerns with antisera

All western analysis performed with antisera had a positive control sample consisting of frozen cleaved purified protein. Samples were electrophoresed with SDS NuPAGE gel as previously mentioned in methods. After electrophoresis, gels were rinsed with deionized water.

0.45µm PVDF membranes (Invitrogen, Grand Island, NY) were soaked in 100% methanol while sponges and Criterion Blotter Filter Paper (BioRad, Hercules, CA) were presoaked in 1X NuPAGE Transfer Buffer (Invitrogen, Grand Island, NY) with 10% methanol. Sponges, filter paper, gel and membrane were stacked upon one another in the Invitrogen transfer system so the arrangement was; 4-5 sponges, 1filter paper, gel, PVDF membrane, 1 filter paper, and finally 4-5 sponges, more sponges allowed for tight contact between the PVDF membrane and the gel. Transfers were run at 45V for 90 minutes on ice in 1X NuPAGE transfer buffer with 10% methanol. Membranes were blocked overnight at 4°C in 5% nonfat milk in 1X PBS and subsequently probed with anti-sera, 1:2,500 in 1X PBST( PBS + 0.05% Tween 20) for one hour; rocking at room temperature. Membranes were washed 4 times for 15 minutes with 1X PBST and then probed with 1:15,000 diluted Goat-Anti-Rabbit IgG ( H+L) Horseradish Peroxidase Conjugate ( Life Technologies, Waltham, MA) for 45 minutes; rocking at room temperature.
Membranes were washed 4 times for 15 minutes in 1X PBST. The membrane blots were incubated in ECL Prime Western Blotting Detection Reagent (GE Healthcare, Buckinghamshire, UK) for 1 minute, covered in plastic wrap and developed with autoradiography film.

**Sn35 Growth Curve Assay**

A 250 mL culture of SBHI + 10% Human Serum was inoculated with a 5 mL overnight culture of Sn35 under anaerobic conditions at 37°C. At 4 hours, 100 mL of growing culture was collected and centrifuged at 4,000 rpm for 20 minutes. The supernatant was discarded and pellet was stored at -80 °C. Additional time points of 6, 8, 12, 16, 24, and 48 hours were collected, centrifuged, and stored at -80°C as well. For each time point the culture collected was halved so that at 48 hours 1.56 mL was collected. Pellets were resuspended in NuPAGE SDS Loading Buffer and boiled for 10 minutes, afterward A280 reading was performed to confirm protein concentration. Samples were loaded into wells so that each contained equivalent amounts of protein. SDS page gels were electrophoresed, transferred onto PVDF membrane and subsequently probed with antisera.

**Adhesion Binding Assay**

A 200 mL culture of SBHI 5% human serum was inoculated with SN35 and grown overnight. Bacterial cells were harvested by centrifugation at 3,000g for 20 minutes at 4°C. Supernatants were discarded and pellets were gently resuspended in 7 mL of TSE buffer (200mM Tris-HCL pH 8.0, 500mM sucrose, 1mM EDTA, protease inhibitor cocktail). Cells were incubated on ice for 30 minutes and then centrifuged at 16,000g for 30 minutes at 4°C. Supernatants were transferred to an ultracentrifuge tube and centrifuged at 60,000g for 1 hour at
4°C. The final spin pellet contained outer membrane proteins while the supernatant contained periplasmic proteins. Pellet was resuspended in 200 µL of TSE buffer, if not immediately used pellet was stored at -80°C. Eight CellStar T75 Flasks (Greiner Bio-One, Monroe, NC) of VK2/E6E7 (ATCC, Manassas, VA), human epithelial vaginal mucosal cells, in 1X Keratinocyte Serum Free Media (Life Technologies, Waltham, MA) accompanied with Supplements for Keratinocyte Serum Free Media (Life Technologies, Waltham, MA) were prepared. Once cells were grown to 80-90% confluency, outer membrane cell surface proteins were biotinylated and extracted using Pierce Cell Surface Protein Isolation Kit (Thermo Scientific, Waltham, MA). Manufacturer’s instructions were followed for Biotinylation, Cell Lysis, and Isolation of Labeled Proteins. After the streptavidin beads were washed in final wash buffer, they were washed with 1X sterile PBS (Life Technologies, Waltham, MA) and centrifuged at 1000g, this was repeated 3 more times. Outer membrane surface proteins isolated from Sn35 were added to streptavidin beads and rocked for 30 minutes at room temperature with end-over-end rotator. Flow thru was collected via centrifugation at 1000g for 1 minute. The streptavidin beads were subject to a series of stringent washes where each wash was collected by centrifugation at 1000g for 1 minute. All washes were done twice and consisted of 500µL. Washes went in the following order; 10mM Tris-HCl pH 8.0 + 10mM NaCl, 10mM Tris-HCl pH 8.0 + 25mM NaCl + 10mM KCl, 10mM Tris-HCl pH8.0 + 100mM NaCl + 25mM KCl, 10mM Tris-HCl pH 8.0 + 150mM NaCl + 50mM KCl, 10mM Tris-HCl pH 8.0 + 150mM NaCl + 200mM KCl. The final wash was Novex NuPAGE LDS Sample Buffer 4X (Life Technologies, Waltham, MA). Samples, including beads, were boiled and run on a NuPAGE SDS gel. The gel was rinsed with sterile water, stained with Imperial Protein Stain for one hour and destained until band patterns were visible.
Placental Invasion Assay

Full term healthy caesarean section, c-sectioned, placentas were provided by consenting women who gave birth at VCU Medical Center through IRB approval #HM10906_CR1 through Dr. Scott Walsh. Placental tissues were immediately placed in a sterile hood using sterile gloves and 4cm x 4cm sections of placental tissues were cut and submerged in a solution of 1X DMEM (Life Technologies, Waltham, MA) and 1mg/mL Penicillin Streptomycin (Life Technologies, Waltham, MA) for 2 hours in Series II Water Jacket CO$_2$ incubator (Thermo Scientific) at 37°C. Sectioned tissues were washed 5 times with 1X sterile PBS (Life Technologies, Waltham, MA). Washed tissue sections were then placed on the outside of a 0.4µm Sterile cell Strainer (Fisher Scientific, Pittsburg, PA) that did not have a bottom strainer. The tissue was kept in placed with a sterile rubber band. Cell strainer + tissue were then placed in a 6 Well Tissue Culture Plate (Grenier, Nurtingen, DE). Each well had 7 mL of antibiotic free DMEM (Life Technologies, Waltham, MA) and 10% pooled human serum. The 6-well plate was incubated overnight in BacBasic Anaerobic Chamber overnight. Live cultures of SN35 and *Lactobacillus crispatus* were inoculated into separate strainers and were left under anaerobic condition overnight. Placental tissues were removed from the cell strainer and were gently washed in 1X sterile PBS 4 times. Tissues were then submerged in 4% paraformaldehyde + 2% glutaraldehyde in 0.1M Cacodylate Buffer. Tissues were embedded in paraffin, sectioned and mounted on microscope slides, and observed under light microscopy. Media from the top portion of cell strainer and the media at the bottom of the well were inoculated onto SBHI Blood agar plates and grown under anaerobic conditions, CFUs were noted.
ME180 Cytotoxicity Assay

A 6-well plate (CellStar) of human epithelial cervical carcinoma ME-180 cells (ATCC, Manassas, VA) were grown to 99% confluency in McCoy’s 5A Media Modified (Cassion Labs, Logan, UT). Cells were incubated in CO₂ incubator at 37ºC during growth. Wells were washed 3 times with 1X sterile PBS (Life Technologies, Waltham, MA) and subsequently incubated in 2 mL of 1X sterile PBS. Wells were then either inoculated with cleaved purified protease, cleaved purified protease + 50mM ZnCl₂, 50mM ZnCl₂, or 2mM CaCl₂. The plates were incubated overnight in CO₂ incubator at 37ºC. Monolayers were examined via light microscopy.
CHAPTER 3
Analyses of Pathogenicity Potential of Sneathia amnii isolate Sn35

Introduction

Once bacteria have invaded the amniotic cavity; the bacterial infection is able to elicit an inflammatory response. Pathogen associated molecular patterns, PAMPs, are recognized by toll-like receptors and activate the innate immune response and the production of cytokines like IL-1α, IL-1β- IL-6, and TNF-α are stimulated (Chiesa et al., 2015; Goldenberg et al., 2008). An increase in IL-6 is a hallmark of preterm birth and the fetal inflammatory response syndrome; while IL-6 levels are modest during term labor, they are excessively higher in preterm births (Chiesa et al., 2015; Romero et al., 2007). IL-6 is produced by a variety of cells and functions to regulate the immune response, inflammation and the acute phase response (Chiesa et al., 2015). The concentration of IL-6 serves as an indicator of the severity of inflammation and if highly elevated levels of IL-6 exist, the fetal inflammatory response syndrome (FIRS) is activated (Chiesa et al., 2015; Jefferson, 2012). Furthermore, IL-6 recruits neutrophils and other leukocytes to the fetal membranes to secrete pro-inflammatory cytokines.

The activation of pro-inflammatory cytokines is a normal process during parturition and leads to the increased activity of matrix metalloproteinases and production of prostaglandins (Goldenberg et al., 2008). Throughout gestation, IL-10, an anti-inflammatory cytokine, is expressed to prevent premature responses; however the production of IL-10 decreases towards the end of gestation and allows for increased activity of other cytokines to induce labor (Hanna et
al., 2000). In order for labor to occur, the cervix must soften, shorten and dilate, and the fetal membranes have to become weakened in order to rupture for birth. During labor estrogen, prostaglandins, and oxytocin hormones work together to cause the uterus to contract forcefully; birthing the fetus (Jefferson, 2012). The decrease in IL-10 allows for corticotropin releasing hormone receptor (CRH-R1) to be induced and thereby increases prostaglandin and placental estrogen release (Jefferson, 2012). In addition to cytokines, women with MIAC will have elevated amniotic fluid levels of matrix metalloproteinase-8 (Combs et al., 2014). Matrix metalloproteinases, MMPs, like MMP-8 and MMP-9 can degrade collagen which allows for the fetal membranes to weaken and aides in the onset of labor (Estrada-Gutierrez et al., 2010).

The fetal membranes are composed of collagens type I, III, IV and V and allow for high tensile strength. Collagen is the most abundant protein found in mammalian bodies and is typed based upon the structure it forms. Collagen type IV forms the basal lamina found in the basement membranes, collagen type V is found in the placenta, collagen type III is typically found with collagen type I which is found in organs and ligaments. The collagen in fetal membranes allows for resilience; however, at the end of term labor MMPs degrade collagen to induce labor.

PPROM and preterm birth have been correlated with elevated activity of collagensases (MMP-8) as well as other MMPs (Maymon et al., 2001). The increase in MMPs can be due to inflammation, but also bacteria. MMP-9 has elevated expression in the fetal membranes in the presence of bacteria. Additionally, select bacterial species are capable of producing their own matrix metalloproteinases in order to degrade collagen (Jefferson, 2012). The work done on this study was to analyze the Sneathia amnii genome to identify MMPs that could be responsible for fetal membrane destruction and PPROM.
Results

Invasion of human fetal membranes by *Sneathia amnii*

It has been determined that MIAC is a cause of preterm birth. Invasion of the amniotic cavity depends upon the bacteria’s ability to produce virulence factors to promote entry into the fetal membranes and cause infection. Previous data showed that *Sneathia amnii* is detectable in the amniotic cavity and can cause chorioamnionitis and amnionitis leading to preterm birth (Harwich et al., 2012). Therefore it is thought that *Sneathia amnii* is ascending from the vagina and ultimately flourishing in the amniotic cavity. Placental tissues from full term healthy c-sectioned patients were incubated with isolate Sn35 in anaerobic conditions at 37ºC to mimic the *in vivo* environment. We found that placental tissues incubated with Sn35 had extensive damage to the chorion layer causing the cells to round up (Figure 3.1). Additionally, tissues incubated anaerobically with no bacteria saw no damage to the chorion layer; however, tissue incubated with Sn35 no longer had tight cell to cell contact. This observation supported previous hypothesis of bacterial ascension and additionally these results determined that Sn35 can invade the placental tissues over a short amount of time. However, media plated from the bottom portion of the well did not have any microbial growth from either *Lactobacillus crispatus* or Sn35, suggesting that the bacteria did not traverse the fetal membranes within 24 hours. While media plated from the top of the cell strainer did have colony growth suggesting that the bacteria survived the experimental conditions. Previous results suggest that *L. crispatus* does not traverse the fetal membranes. The media plated from the top was to ensure that bacterial survival could occur in DMEM 10% Human serum. The lack of Sn35 in the bottom portion of the well indicated that while Sn35 has the ability to traverse the first layer of the fetal membranes, it may need additional time to fully traverse the fetal membranes.
Figure 3.1: Fetal membranes incubated with or without Sn35. Placental tissue not incubated with bacteria (top image) remained intact and did not have cell or tissue damage. Placental tissue incubated with Sn35 (bottom image) had extensive tissue and cell damage. Highlighted in the box, the chorion layer no longer possessed tight cell:cell contact and cells appeared to be rounding; indicating that Sn35 was able to traverse the chorion layer.
**Sn35 adhesion capability with VK2 cell line**

Bacterial adhesion to a eukaryotic cell always precedes invasion. In order to identify ligand: ligand receptor interaction, outer membrane proteins from both Sn35 and VK2 cells were isolated. VK2 outer membrane proteins were biotinylated and bound to streptavidin beads while outer membrane proteins from SN35 were added to the VK2 beads. A series of stringent salt solutions was used to wash the beads in hopes of identifying a bacterial peptide with high affinity for a eukaryotic protein. However, we did not detect a peptide that remained on the streptavidin beads following the initial washes. Therefore we did not isolate a Sn35 adhesin.

**Selection of virulence determinant expressed by Sn35**

Bioinformatic analysis revealed a gene of interest in Sn35 labeled as a putative zinc endopeptidase with homology to the M23 peptidase family comprised of glycine glycine endopeptidases and matrix metalloproteinases. Analysis using SignalP4.1 suggested that the putative zinc endopeptidase had a signal peptide for secretion via the sec pathway. The isoelectric point of the putative zinc endopeptidase is 10.63 pH(I). With the signal peptide, the gene is 1311 bp and 436 amino acids and is 48.95 kDa. Additionally, it also has an active site characteristic of the M23 peptidase family, HXXXD, where HXXXD is HAILD, and is found AA site 165-169 (Figure 3.2) This putative gene was of interest to our research since it could be a potential virulence determinant that allows Sn35 to traverse the fetal membranes. The zinc endopeptidase gene, without its signal sequence, was then amplified by PCR with specific primers, and was named protein encoding gene 1 (Peg1). Serial cloner indicated that the gene is 1233 bp (Figure 3.3).
**Figure 3.2 Gene Map of Putative Zinc Metallopeptase.** The nucleotide sequence is shown first and the amino acid sequence is directly below it. The number, seen at the end of every line, correlates to the position of the last amino acid on the line. The signal peptide is highlighted in green, the active site HAILD is highlighted in blue, and the M23 peptidase is highlighted in orange.
**Figure 3.3: DNA gel of Peg1 PCR amplification.** PCR amplification of Peg1 was done using Peg1 specific primers. Lane 1 (left most lane) contains Hyperladder 1kb, Lane 2 contains PCR product Peg1, 1123kb which is highlighted in the box. Molecular weight markers indicated.

DNA sequence of Peg1 is noted below DNA gel.
Expression and purification of recombinant Peg1

Peg1 was cloned into pET32xT (generously donated by Dr. David Williams, VCU, Richmond, VA) in BL21 DE3 RIL E.coli cells, clones were restriction digested (Figure 3.4) and sequenced to ensure successful cloning. The vector, pET32xT, is an IPTG inducible vector which attached a C-terminal Thio-HIS tag to the protein. Expression of the Thio-His-tagged Peg1 product was confirmed through SDS PAGE gel with the presence of the protein at approximately 64.1kDa upon induction with 1M IPTG (Figure 3.5). Peg1 is 46.1kDa while the His-tag is 18kDa. The expressed recombinant protein was isolated through column purification using cobalt resin (Figure 3.6) and the Thio-His tag was removed by thrombin cleavage and the addition of calcium (Figure 3.7). If 150mM of CaCl$_2$ was not added to the thrombin cleavage reaction, Peg1 rapidly degraded in a site specific fashion leaving two peptide bands at 35kDa and 12kDa (Figure 3.8). The degradation appeared to be autocatalytic as no contaminating proteases were detected. While Peg1 does not have an obvious (HEXXH) zinc binding motif, which is common in M23 peptidases, our results determined that CaCl$_2$ does hinder Peg1’s autocatalytic degradation. Additionally, these results determined that the protein degrades itself in a site specific manner to produce a C-terminal peptide and an N-terminal peptide.

Edman Degradation

Site specific cleavage is also seen in a zinc endopeptidase expressed by Bacteriodes fragilis (Freer et al., 1999). The protein is cleaved into a proprotein and a mature protein between an Arginine$^{211}$ and Alanine$^{212}$. We hypothesized that since Sneathia amnii is also a zinc metalloprotease, Peg1 could cleave itself at the same motif. Sn35 has an Arginine Alanine site where if cleaved at this site it would produce a protein around 35 kDa. However, when the
Figure 3.4F: DNA agarose gel analysis of restriction digest of BL21 DE3 RIL *E. coli* cells containing pET32xT+Peg1. Lane 1 contains HyperLadder 1kb, lane 2 contains BL21 DE3 RIL colony miniprep that has been digested with BamHI and XhoI. The fragments 5900 bp (pET32xT) and 1123 bp (Peg1) indicate successfully cloning by restriction digest confirmation.
Figure 3.5: Confirmation of induced Peg1 with 1mM IPTG. Imperial stained SDS-PAGE of uninduced (Lane 2) and induced (Lane 3) cell lysates of E.coli BL21 DE3 RIL cells containing pET32xT+ Peg1. Molecular weight markers are shown above. Peg1 with Thio-HIS tag is 64.1 kDa, indicated in box, is induced under IPTG.
Figure 3.6: Purified protein by cobalt resin column confirmed with SDS PAGE. Lane 5 indicates purified protein was collected after 250mM imidazole wash, peptide band boxed in is Peg1 which is 64.1 kDa. Molecular weights are indicated. Lane 2 is flowthrough, Lane 3 and 4 are 10mM imidazole washes.
Figure 3.7: Thrombin cleavage of Thio-HIS tag with CaCl$_2$. SDS-PAGE gel shows supplementation of CaCl$_2$ in the thrombin reaction allowed for recovery of 46.1 kDa peptide, boxed in Lane 2. Molecular weight markers are noted.
**Figure 3.8: Thrombin cleavage without CaCl$_2$ allows for site specific degradation.** Imperial stained SDS PAGE gel depicts autodegradation of Peg1 into 2 peptides sized 12 kDa and 37 kDa in Lane 3. Lane 2 is thrombin treated Peg1. Molecular weight markers are indicated.
11.98 kDa peptide was sent for N-terminal sequencing, it was found that Peg1 cleaves itself between a Glutamine\(^{(335)}\) (Q) and Glycine\(^{(336)}\) (G) before a Leucine Lysine (LK) motif leaving the N-terminal peptide at 312 AA and the C-terminal peptide at 92 AA (Figure 3.9). The N-terminal peptide is 34.12 kDa while the C-terminal peptide is 11.98 kDa.

**Extracellular Matrix Component Degrading Activity**

We hypothesized that Peg1 would be able to degrade a substrate that was present in the fetal membranes. First, the protease was tested against a universal protease assay where casein acted as the substrate. Zone of activity plate assays indicated that proteinase K, a broad spectrum serine endopeptidase, was able to degrade casein; however, Peg1 was unable to degrade casein. This same method was tested with gelatin and mucin and the results were the same, proteinase K was able to degrade the three substrates while Peg1 was unable to. Mucin was tested as a control since it was already shown that Sn35 was unable to degrade mucin (Harwich et al., 2012).

Since Blastx indicated homology to matrix metalloproteinases, we next hypothesized that the substrate of Peg1 would be a component in the extracellular matrix. The ECM consists of extracellular molecules secreted by cells that allow for structural support for the surrounding cells, tissues repair, and intercellular communication. Matrix metalloproteases, serine proteases and threonine proteases aide in the dynamic changes of the ECM by degrading and remodeling the ECM. The ECM components include collagen, fibronectin and laminin. Fibronectin is a glycoprotein that binds to integrins, fibrin, collagen and proteoglycans and allows for dynamic movement; it is an insoluble matrix where its monomers are linked together by disulfide bonds. While laminin, a major extracellular matrix protein of the basal lamina aides in cell adhesion and
Figure 3.9: Schematic representation of site specific degradation. Peg 1 autodegrades itself at the N terminus of the 12 kDa peptide. Cleavage occurs between Q\text sub{335} and G\text sub{336}. Full amino acid sequence of signal peptide, N-terminus and C-terminus are noted below.
provides structural scaffolding. Laminin can also bind to collagen, integrins and other cellular domains.

In addition to testing ECM components; we hypothesized that since this protease had homology to putative zinc endopeptidases that it must require zinc for activation. Therefore the substrates and Peg1 were incubated with varying amounts of salt concentrations. Fibronectin and laminin assays were run on SDS PAGE to determine if Peg1 was capable of complete degradation. Fibronectin and laminin substrates were collected from human tissue in attempt to recreate the in vivo substrates for Peg1. Fibronectin produced a band at 220kDa on the SDS PAGE and while it appeared that Peg1 was able to degrade fibronectin, complete degradation was never seen. While lanes 3 and 5 on the SDS PAGE gel appear to have less fibronectin (Figure 3.10), this might be due to salt concentrations effecting the migration of proteins into the gel. The fibronectin sample located in lane 4 was incubated with 5 mM ZnCl$_2$ and 2mM CaCl$_2$ while lane 2 was incubated with three times more ZnCl$_2$ and five times more CaCl$_2$ concentrations. The results indicate that less protein migrated into the gel which can be due to the salt concentration in the samples. Laminin produced two major bands at 50kDa and 130-160kDa. Although the results of the SDS PAGE could indicate that the protease degrades the 130-160 kDa peptide, once again it appears that the salt concentrations affected the migration of the protein into the gels (Figure 3.11). Furthermore, some of the samples remained in the wells. Therefore it was concluded that Peg1 cannot degrade fibronectin or laminin.

**Coagulase potential of Peg1**

Several microorganisms are capable of producing coagulase enzymes to convert fibrinogen to fibrin which results in blood clotting, like *S.aureus*. This conversion is viewed as a
Figure 3.10: SDS PAGE fibronectin degradation assay. Imperial stained SDS-PAGE. Lane 2: Fibronectin+15mM ZnCl₂+10mM CaCl₂, Lane 3: Fibronectin+ 15mM ZnCl₂+ Peg1 , Lane 4: Fibronectin+5mM ZnCl₂+2mM CaCl₂, Lane 5: Fibronectin+5mM ZnCl₂+ (1:5) Peg1, Lane 6: Peg1 alone. Degradation of Fibronectin is not seen.
Figure 3.11: SDS PAGE laminin degradation assay. Lane 2: Laminin+15mM ZnCl₂+10mM CaCl₂, Lane 3: Laminin+15mM ZnCl₂+Peg1, Lane 4: Laminin+5mM ZnCl₂+2mM CaCl₂, Lane 5: Laminin+5mM ZnCl₂+Peg1, Lane 6: Laminin + 5mM ZnCl₂+ (1:5) Peg1, Lane 7: Peg1 alone. Peg1 is unable to degrade laminin.
virulence factor since it coats the bacterial surface with fibrin and therefore can protect the bacterial cell from phagocytosis (Tortora et al., 2013). Peg1 was incubated with rabbit fibrinogen; however, clumping never occurred and therefore Peg1 was not identified as a coagulase. This experiment was done to determine if Sn35 utilizes Peg1 to remain undetected from the immune system while traversing the fetal membranes.

Analysis of cytotoxicity of Peg1 to ME-180s

Previous studies indicated that Sneathia amnii is cytotoxic to ME-180 cells which are cervical epithelial carcinoma cells. It is thought that the cytotoxic activity may help the bacteria ascend into the intrauterine cavity. While Peg1’s peptides were analyzed through Edman degradation, a cytotoxicity assay was performed based upon Bacteroides fragilis’s zinc endopeptidase’s capability to disrupt tight junctions in epithelial cell monolayers. Hypothesizing that Peg1 could have similar enzymatic capabilities; ME-180 cells were incubated with Peg1 and ZnCl₂. Results indicated that there was a significant increase in cell rounding compared to cells just treated with ZnCl₂ (Figure 3.12). While obvious cytotoxicity was not seen, the monolayers incubated with Peg1 and ZnCl₂ sloughed off.

Analysis of biofilm formation capability of Peg1

Previous attempts to uncover Peg1’s protease capabilities indicated that Peg1 did not have true degradation of substrates utilized in this study. Therefore, it was hypothesized that Peg1 could be a ligand involved in adhesion and biofilm formation. Collagen type IV, which is located in the fetal membranes, was utilized to determine whether Peg1 could bind to collagen type IV and allow S.amnii to produce biofilms. Live S.amnii was incubated in collagen type IV
Figure 3.12: Cytotoxic effect of Peg1 on ME-180s. Monolayer exposed to ZnCl$_2$, top image, remained intact and cells had little to no rounding. Monolayer exposed to Peg1 and ZnCl$_2$, bottom image, came apart and sloughed off. An increase in cell rounding was observed when cells were exposed to Peg1.
strip wells with or without the addition of cleaved purified Peg1. After, the wells were stained with safranin to visually evaluate biofilm formation. No biofilm formation occurred in the presence or absence of Peg1.

**Bacteriolytic Activity**

Production of a bacteriolysin is a virulence factor for the bacterial cell against other species. While autolysins are produced during binary fission to break down the bacteria’s own peptidoglycan layer and allow for two cells to form from one cell. We hypothesized that Peg1 could be either a bacteriolysin or an autolysin since proteases, like proteinase K, can break down *Micrococcus luteus*. M23 peptidase family also contains endopeptidases that lyse bacterial cell wall peptidoglycans. The cleaved purified protease was exposed to *Micrococcus luteus* and a zymogram assay was performed. A zone of clearing was produced and therefore it was concluded that Peg1 appeared to have bacteriolytic activity. (Figure 3.1). When the cleaved purified protease was incubated with Sn35 to analyze autolysin potential via OD$_{600nm}$ measurement it was concluded that Peg1 was not capable of autolysin activity since there was not a significant decrease in OD$_{600nm}$ measurement.

**Collagen degradation**

Collagen, an integral protein of connective tissue, is found in the ECM, skin, tendons, ligaments and other tissues. Collagen exists as procollagen before it is cleaved by proteases to allow for assembly into collagen; collagen type IV is the only type of collagen to be found in the basement membrane which is found in the fetal membranes. It had previously appeared that Sn35 had collagenase activity and could partially degrade collagen type V. To analyze if Peg1...
Figure 3.13: Bacteriolytic effect of Peg1 on *Micrococcus luteus*. *M. luteus* embedded SDS was run and imperial stained to indicate whether or not Peg1 was able to lyse other bacterial species. A zone of clearing was observed in lanes 2-4 where Peg1 was loaded. Therefore Peg1 appears to have bacteriolytic activity.
was able to degrade collagen, collagen type IV and I were biotinylated and incubated with cleave purified Peg1. Western analysis indicated that Peg1 appears to have the same effect on collagen type IV as live bacteria do (Figure 3.14). Peg1 appears to have collagenase activity.

Analysis of Peg1 antisera production

Bioinformatic analysis of PSORTb identified Peg1 as a lipoprotein located on the outer membrane of the bacterial cell wall. Additionally, PSORTb identified that without the 12kDa fragment the protein had no signal. Therefore cleaved purified protein was used for polyclonal antisera production to further help characterize Peg1. To confirm PSORTb results, live Sn35 was probed with anti-sera; however, western analysis indicated that Peg1 was not found on the outer membrane (Figure 3.15) and instead antisera detected the protein in the cytoplasm. While we hypothesized that the protease was expressed on the outer membrane and PSORTb indicated the same results, the results from western analysis cemented that this protease is not expressed on the outer membrane. To better understand the protease using antisera, a temporal expression assay was done to identify when the putative protease is expressed. The growth curve assay indicated that Peg1 is expressed throughout exponential phase, late exponential phase and stationary phase; hours 4 through 48(Figure 3.16).

Discussion, Conclusion, and Future Perspectives

Almost nothing is known about the process through which *S. amnii* causes preterm birth, even though this newly discovered bacterium is associated with serious health complications for
Figure 3.14: Role of Peg1 in collagen degradation. Lanes 1 and 2 are live Sn35 incubated with collagen type IV, lanes 3 and 4 are PBS incubated with collagen type IV, lanes 5 and 6 are Peg1 incubated with collagen type IV. Peg1 appears to have collagenase activity, to the same effect that live Sn35 does.
Figure 3.15: Western analysis of Peg1 expression location. Lane 1: molecular weight marker, Lane 2: cytosolic proteins of Sn35, Lane 3: outer membrane protein of Sn35. Peg1 is visualized in the cytoplasm of Sn35 about 75 kDa.
Figure 3.16: Temporal expression of Peg1. Lane 1: ladder, Lane 2: empty, Lanes 3-9 are cell lysates taken at 4, 6, 8, 12, 16, 24, 48 hours respectively. Peg1 expression was found throughout exponential, late exponential and stationary phase. Peg1 is visualized ~75 kDa.
both mother and neonate. We have hypothesized that \textit{S. amnii} has virulence determinants that allow for the bacterium to traverse the fetal membranes and invade the amniotic cavity. The goal of this project was to identify a potential virulence factor that Sn35 utilizes to traverse the fetal membranes and cause preterm birth. We sought to first better understand traversal of \textit{S. amnii} through the fetal membranes and then to explore potential virulence determinants that aided in the degradation of the extracellular matrix. The initial study focused on bacterial adhesion and ascension through the chorion layer to support MIAC. We have developed a method that simulates MIAC \textit{in vitro}, and indicated that Sn35 is able to invade and traverse the chorion layer. These observations are significant because it supports bacterial ascension and indicates that \textit{S. amnii} utilizes virulence determinants to ascend. The lack of detectable bacteria below the placental tissue indicates that further work is needed to develop a MIAC assay that can stay viable longer than 24 hours to observe microbial traversal. While Sn35 did not produce a strong ligand: ligand interaction with VK2 cell line, this indicates that further work is necessary to identify what cell tissue type Sn35 targets for cell adhesion.

The virulence determinants of \textit{S. amnii} remain elusive. Analysis of its genome indicated promising virulence determinants that could enlighten its ability to traverse the fetal membranes and cause preterm birth. We hypothesized that the putative zinc endopeptidase, Peg1, would degrade substrates found in the fetal membranes due to its homology to matrix mellatoproteinases. We attempted to discover Peg1’s degradation properties by incubating the cleaved purified protein with various substrates found in the extracellular matrix, like fibronectin and laminin; however, results from each experiment either did not support the hypothesis of degradation. Peg 1, under the conditions tested, was not capable of degrading casein, mucin, gelatin, or fibrinogen. Furthermore, Peg1 lacked biofilm formation capability using collagen type
IV as the substrate. However, when Peg1 was incubated with collagen type IV, Peg1 appeared to have the same effect on collagen as the live bacteria. It appears that Peg1 has collagenase activity. Peg1 also appeared to have bacteriolytic activity against *Micrococcus luteus* which is characteristic of the M23 peptidase family. Antisera production of Peg1 identified the protease as being cytosolic, disproving PSORTb bioinformatic analysis that placed Peg1 on the outer membrane. Peg1 does appear to have protease activity, however, the conditions in which the protease is secreted were not found. Further work needs to be devoted to understand the conditions in which Peg1 is secreted.

Peg1 appeared to autodegrade and we determined the cleavage site by Edman degradation. Peg1 required the addition of CaCl₂ in order to remain an intact peptide of 46.1 kDa. If CaCl₂ was not supplemented to the thrombin reaction, Peg1, underwent autocatalytic cleavage. This autocatalytic degradation resulted in two distinct peptides; a peptide sized 34.12 kDa while the other 11.98 kDa. Further identification of the autocatalytic event was examined through N-terminal Edman degradation revealing that site specific cleavage occurred between Glutamine (335) and Glycine (336). Additionally, the cleavage at a glycine residue is often seen in M23 peptidase family, since it is the preferred site of cleavage.

Given the clinical complications that *Sneathia amnii* possesses as an opportunistic pathogen responsible for an array of infections in numerous body sites in both mother and neonate, research into the virulence determinants could have a significant impact on our understanding in their role in BV, MIAC, PPROM and preterm birth. The mucus plug and fetal membranes are supposed to remain impermeable until the end of gestation; however, *S. amnii* can be found in the uterine cavity, fetal membranes, and amniotic fluid during gestation. The work in this study revealed the ability of Sn35 to traverse the chorion layer of the fetal
membrane. Additionally it appears that Peg1 has collagenase and bacteriolytic activity that could be responsible for MIAC; however, conditions in which Peg1 is secreted were not identified. Therefore, the contributions of this study to our understanding of Sn35’s ability to cause preterm birth may help understand its other virulence determinants.
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Vita

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Virginia Branch ASM Student Travel Grant. October 2014