Differential Production and Secretion of Potentially Toxigenic Extracellular Proteins from Hypervirulent Aeromonas hydrophila Under Biofilm and Planktonic Culture

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Abstract

Background: Hypervirulent *Aeromonas hydrophila* (vAh) is an emerging pathogen in freshwater aquaculture that results in the loss of over 3 million pounds of marketable channel catfish, *Ictalurus punctatus*, and channel catfish hybrids (*I. punctatus, I. furcatus*) each year from freshwater catfish production systems in Alabama, U.S.A. vAh isolates are clonal in nature and are genetically unique from, and significantly more virulent than, traditional *A. hydrophila* isolates from fish. Even with the increased virulence, natural infections cannot be reproduced in aquaria challenges making it difficult to determine modes of infection and the pathophysiology behind the devastating mortalities that are commonly observed. Despite the intimate connection between environmental adaptation and plastic response, the role of environmental adaption on vAh pathogenicity and virulence has not been previously explored. In this study, secreted proteins of vAh cultured as free-living planktonic cells and within a biofilm were compared to elucidate the role of biofilm growth on virulence.

Results: Functional proteolytic assays found significantly increased degradative activity in biofilm secretomes; in contrast, planktonic secretomes had significantly increased hemolytic activity, suggesting higher toxigenic potential. Intramuscular injection challenges in a channel catfish model showed that *in vitro* degradative activity translated into *in vivo* tissue destruction. Identification of secreted proteins by HPLC-MS/MS revealed the presence of many putative virulence proteins under both growth conditions. Biofilm grown vAh produced higher levels of proteolytic enzymes and adhesins, whereas planktonically grown cells secreted higher levels of toxins, porins, and fimbrial proteins.

Conclusions: This study is the first comparison of the secreted proteomes of vAh when grown in two distinct ecological niches. These data on the adaptive physiological response of vAh based on growth condition increase our understanding of how environmental niche partitioning could affect vAh pathogenicity and virulence. Increased secretion of colonization factors and degradative enzymes during biofilm growth and residency may increase bacterial attachment and host invasiveness, while increased secretion of hemolysins, porins, and other potential toxins under planktonic growth (or after host invasion) could result in increased host mortality. The results of this research underscore the need to use culture methods that more closely mimic natural ecological habitat growth to improve our understanding of vAh pathogenesis.

Background

*Aeromonas hydrophila* is a wide-spread and diverse species of Gram-negative bacterium ubiquitous in freshwater aquatic ecosystems. As a rapidly growing and metabolically diverse generalist (1-5), *A. hydrophila* is capable of exploiting a variety of ecological habitats and a broad range of hosts. *A. hydrophila* has been isolated from almost every freshwater aquatic environment and from diseased mammals, reptiles, amphibians, insects, and fish (1, 6-8). *A. hydrophila* has been found in association with processed poultry, meats, fish, and even bottled water. It is capable of withstanding chlorination and is resistant to multiple antibiotics (1, 9).
In aquaculture, *A. hydrophila* is an important cause of disease in most freshwater production systems. Historically, *A. hydrophila* has been an important secondary pathogen in catfish production systems, commonly responsible for cutaneous ulceration and muscle necrosis. Occasionally following fish stress (low oxygen, poor water quality, etc.) the bacterium can cause a septicemia (motile aeromonad septicemia [MAS]), resulting in high mortalities (10-14). In 2009, a new, highly virulent strain of *A. hydrophila* was isolated from a diseased channel catfish, *Ictalurus punctatus*, within a production pond in West Alabama. This strain, referred to as hypervirulent *Aeromonas hydrophila*, or vAh, was responsible for outbreaks of peracute motile aeromonad septicemia of epidemic proportions (11, 15-19). vAh apparently acts as a primary pathogen, and may not be preceded by immune insult (11). To date, vAh has been responsible for the loss of 30 million pounds of marketable channel catfish from production farms in West Alabama. In 2017, *A. hydrophila* infections were responsible for the loss of 3.4 million pounds of farm-raised catfish in Alabama alone, more than twice as much as the second leading cause of loss, *Flavobacterium columnare*. vAh has been the primary or secondary cause of catfish loss in Alabama since the primary outbreak in 2009 (Hemstreet, AL Fish Farming Center).

While much current research is focused on MAS disease prevention, there are many important unanswered areas of research to understanding the mechanisms of vAh pathogenesis, bacterial-host interactions, and bacterial adaptive responses under different environmental conditions. *A. hydrophila* are known to secrete a multitude of degradative and cytotoxic extracellular proteins which are widely accepted as virulence determinants (20-22), and which likely contribute to the environmental adaptability and broad species host range.

While vAh has established itself as a primary pathogen in natural settings (11), laboratory-cultured vAh appears to mimic its opportunistic relatives during immersion challenges. Planktonically-cultured vAh is extremely virulent, causing death in a matter of hours in intraperitoneal injection challenges. However, models meant to mimic more natural infections including submersion and gavage have been unreliable, even when challenged with artificially high colony forming units (CFUs) of the bacterium (2, 23). Current studies of vAh pathogenesis and virulence are performed almost exclusively with planktonically-cultured bacteria despite the fact that most free-living generalist bacteria in aquatic systems reside primarily within biofilm (24-29). One study using only planktonically-cultured vAh reported the presence of 228 extracellular proteins (ECPs) in the supernatant of vAh broth cultures, at least 23 of which were putative virulence factors (18), and a recent study comparing the secretomes of wild-type vAh with that of a group four capsule (gfc) - deficient vAh mutant reported the presence of multiple degradative and hemolytic proteins under planktonic culture (30). A comparative proteomics study by Wang et al (31) found differential expression of 33 proteins, many of which were involved in proteolysis, in response to iron starvation, underscoring the role of secreted proteins in environmental adaptation of vAh. Though niche adaptation clearly plays an important role in protein secretion, no studies have evaluated the secretome of biofilm grown vAh. A recent study by Cai et al. (2018) found no vAh present in the water column through the survey period, (July- October), while vAh resident in biofilm and pond sediment was detected at an increasing rate in the same sampling period, suggesting that biofilms serve as a stable reservoir for vAh survival when planktonic conditions are less favorable. Biofilm-associated bacteria generally have
increased adhesive properties (24, 25, 32-34) and may have increased production of proteolytic enzymes, both of which could increase virulence (12, 35-37). Redfield (2002) suggested that extracellular proteases are expressed when diffusion and/or mixing is reduced. vAh residing within a biofilm may have an advantage in attaching to and invading fish tissues due to increased secretion of proteolytic enzymes and adhesins. Given the data supporting the presence of vAh within pond biofilms, it is important to identify virulence factors secreted during biofilm residence that could impact host attachment and invasion. In this study, we compared the secreted protein profiles (secretomes) of biofilm- or planktonically cultured vAh strain ML09-119 to determine if niche occupation could influence vAh pathogenicity in natural environments.

**Results**

**Biofilm grown vAh express higher protease activity but less hemolytic activity than planktonic cultures.** Protease activity in biofilm samples was observed to be more than 2 times higher than in planktonic samples, and 1.2 times higher than the trypsin positive control ($p < 0.05$; Figure 1). Similarly, elastase activity was significantly higher for biofilm grown vAh, which expressed 2.7 greater elastase activity than observed for planktonically grown cultures ($p < 0.05$; Figure 2). In contrast, hemolytic activity was greatly increased in planktonic cultures, with more than 6 times higher hemolytic activity compared to biofilm grown vAh ($p < 0.05$; Figure 3).

**Severe fish tissue necrosis was induced by the secretome of biofilm-grown vAh.** To determine if the increased proteinase and elastase activities observed from vAh when grown as an *in vitro* biofilm would result in tissue damage indicative of MAS disease, 10 µg of secreted proteins from each growth condition was injected intra-muscularly into channel catfish. Two hours post-injection, loss of dermal pigment was noted at the injection site in biofilm-injected fish, but no changes were observed for the injected planktonic secretome. After 24 hours, substantial tissue necrosis was observed grossly in all biofilm-injected fish (Figure 4). Fish injected with planktonic-associated ECPs developed no gross lesions even after 7 days. No control fish developed any gross lesion at the injection site after 7 days.

Histopathology was performed on skin and subcutaneous tissues collected from injection sites of channel catfish. Biofilm-injected fish tissue was edematous, hemorrhagic, and there was extensive tissue necrosis at the site of injection (Figure 4). Despite substantial tissue damage, few inflammatory cells were present. In contrast, fish injected with planktonic ECPs were identical to the control fish, with no perceptible damage to skin, subcutaneous adipose tissue, or muscle.

**Biofilm and planktonically grown vAh have distinct secretomes.** The differences in enzyme activities and tissue damage observed for biofilm versus planktonically-cultured vAh supported the hypothesis that niche occupancy has a significant influence on vAh exoprotein expression. To further test this hypothesis, a secretome analysis was performed to identify differentially secreted proteins present under the two culture conditions. A total of 272 proteins were identified in the secretomes of biofilm and planktonically-cultured vAh. Eighty-two proteins were identified that were present in both secretomes, while 98 were
identified only in biofilm secretomes and 92 were unique to planktonic secretomes. ROTS and T-test analyses identified 53 proteins that significantly (FDR < 0.05, \( p = 0.01 \)) varied in abundance. The protein abundance ratios of 52 ROTS-identified proteins were above the significant fold change threshold of \( \geq 1.5 \) (Table 1). Thirty-five proteins were significantly increased in the biofilm secretomes, 20 of which were uniquely present in samples from biofilm grown vAh; for planktonic secretomes, 15 proteins were significantly increased in their abundance relative to biofilm samples, and these included nine proteins that were only observed from planktonic cultures. Of the proteins that varied significantly in their abundance, at least 15 from planktonic secretomes and 30 from biofilm secretomes have been indicated in virulence (Table 1, in bold font). However, not all secreted putative virulence proteins were differentially secreted, with many putative virulence factors identified in secretomes under both conditions.

Functional group comparisons based on gene ontology (Table 1, Figure 5) revealed extensive secretion of degradative enzymes and toxins in both biofilm and planktonic secretomes, with degradative enzymes, such as elastase, metalloprotease, chitinase, and endochitinase, dominating biofilm secretomes and cytotoxic and cytotoxic toxins, such as \( ahh1 \)-type hemolysin and extracellular lipase enriched in planktonic secretomes. In both planktonic and biofilm secretomes, degradative enzymes and toxins made up the majority of significant proteins, representing 79.8% of planktonic proteins and 55.7% of biofilm proteins. Proteins involved in transport (16.5%), carbohydrate metabolism (8.5%), and pilus and flagellin (3.6%) contributed significantly to the biofilm secretome, while pilus and flagellin proteins (5.8%), outer membrane proteins (4.0%), and proteins involved in transcriptional regulation and electron transport (3.5%) were other significant contributors to planktonic secretomes (Figure 5). Of particular interest were the presence of polar flagellar proteins (\( AHML\_09345 \) and \_09350\)) present in higher quantities in the biofilm secretome and type I pili proteins (\( AHML\_2665 \) and \_2690\)) that were present in planktonic secretomes, but absent from biofilm secretomes. Polar flagella, typically considered motility flagella, are important in adhesion and invasion in \( A. \ hydrophila \) that lack lateral flagella, such as vAh (38), while type I pili are thought to contribute to host colonization, but not host invasion (39).

**Discussion**

While vAh are significantly more virulent that traditional \( A. \ hydrophila \) when challenged by intraperitoneal injection (15), Zhang et al. (40) reported consistent vMAS mortality was attainable in channel catfish immersion trials only following scarification and challenge with \( 2\times10^7 \) CFU/ml of planktonically-cultured vAh. This suggests that some environmental stimuli are not present in artificial broth culture, which, in pond systems, could be responsible for inducing bacterial virulence and resulting in large scale MAS epidemic outbreaks. Since most environmental bacteria spend much of their time in biofilm, either attached to a substrate or floating as bacterial flocs (24, 26, 27, 29, 41), biofilm-associated vAh may produce proteins that increase invasiveness and allow initial colonization *in vivo* (42). The ability to form a biofilm is commonly considered a virulence factor, particularly in human disease conditions (43). Likewise, for \( A. \ hydrophila \) biofilm formation and residency may induce global changes in gene expression resulting in increased production and secretion of degradative enzymes and other factors that
increase pathogenicity or invasiveness. *Aeromonas* spp. produce extracellular enzymes that facilitate nutrient acquisition in aquatic environments and produce adhesins that aid in the attachment and colonization of benthic surfaces (5). In aquatic environments, these enzymes provide nutrients by degrading the organic compounds including suspended detritus and benthic substrates. These enzymes may also be important in the pathophysiology of disease by enabling degradation of animal tissues (5, 24, 44).

Previous research reported the presence of multiple potentially toxigenic extracellular proteins in the supernatant of planktonically-cultured vAh (18, 30). Because many opportunistic bacteria like vAh reside largely in biofilms and not as sustained planktonic populations (45), it was important to evaluate the influence of biofilm growth on vAh exoprotein expression. This study found that degradative activities were significantly increased in the supernatant of biofilm-associated vAh (Figures 1 and 2). Furthermore, when biofilm-grown vAh ECPs were injected into the muscle of channel catfish, significant necrosis and cytolysis occurred within 24 hours, while secreted proteins of planktonically-cultured vAh failed to produce necrotic lesions after seven days.

A secretome analysis was conducted to examine in more detail how biofilm growth influenced vAh exoprotein expression, which revealed significant differences in the secretomes of the two cultures, both in complexity and quantity. The biofilm secretome contained 248 proteins, including 183 unique proteins, while planktonic secretomes contained 183 total proteins, including 101 unique proteins. Of the 82 proteins secreted under both culture conditions, at least 36 had previously been identified as putative virulence factors (18, 21, 46). Under both growth conditions, vAh secreted an abundance of potential virulence proteins, the majority of which were not statistically significant in differential secretion analyses. However, secretomes of vAh cultured in biofilm were significantly more varied and, in general, relative protein abundance was increased.

Assays to measure general and specific proteolytic potential of the secreted proteins revealed significant increases in both caseinolytic and elastinolytic activity in biofilm secretomes when compared to planktonic ECPs (Figures 1 and 2). A significant difference in proteolytic potential was also seen upon inspection of the secretome analysis. ROTS analysis revealed at least seven degradative proteins were present in the biofilm secretomes at significantly higher observed abundance relative to planktonic secretomes. There was a 5-fold increase in elastase abundance in biofilm secretomes, with an average quantitative protein value (QPV) of 122, compared to an average QPV of 23 in planktonic secretomes, which agreed with the results obtained from elastase enzyme activity measurements. There was a 3-fold increase in the M66 - family metalloprotease *AHML_05230* in biofilm secretomes, with average QPVs of 103 and 30 in biofilm and planktonic secretomes, respectively. Both elastase and the M66 zinc metalloprotease are considered significant virulence factors of *A. hydrophila* as well as other pathogens, such as *Vibrio cholerae*, and enterohemorrhagic *Escherichia coli* (47, 48). Five other proteolytic enzymes were secreted in statistically significant quantities in biofilm secretomes but were not detected in the planktonic secretomes and likely increase the overall proteolytic potential of biofilm ECPs (Table 1).
While the majority of the differentially secreted degradative enzymes present in the biofilm secretome were proteolytic, two important glycolytic proteins, chitinase and chitin binding protein (CBP) were found in significantly higher amounts in biofilm secretomes. While chitinase and CBP are integral in the breakdown of environmental chitin, these proteins may also play integral roles in virulence. Though vAh can use chitin as a sole carbon source (49), the lack of chitin in the TSB growth medium would make it unlikely that chitinase and CBP production would be energetically favorable. Therefore, it is hypothesized that these chitin-associated proteins play other roles in vAh fitness or pathogenicity. In other pathogens, chitinases and CBPs are considered virulence factors not because they target chitin but because of their interactions with substrates other than chitin. In some virulent *E. coli* and *V. cholerae*, chitinases and CBPs target host glycoproteins and glycolipids that contain N-acetylglucosamine (GlcNAc), the monomer present in mucus (50, 51). Outer membrane-expressed chitinases and CBPs have also been indicated as accessory molecules responsible for initiating host cell adhesion and invasion (50-52). In a murine model, *E. coli* chitin-binding domain interacts with intestinal epithelial cells, increasing invasiveness and pathogenicity (51). In *V. cholerae*, Bhowmick et al. (2008) found chitinases function to break down the GlcNAc of mucus and reported upregulation of chitinases resulting from exposure to exogenous mucin. Furthermore, the *V. cholerae* chitin binding protein GbpA was shown to bind to the protective mucus layer of mammalian intestinal epithelium, resulting in bacterial colonization and disease initiation. Likewise, chitinases and CBPs produced by clinical *Pseudomonas aeruginosa* strains isolated from patients with cystic fibrosis (CF) were also upregulated in response to mucin-containing sputum and likely play an integral role in primary adhesion to lung epithelium in the initiation of CF (50). In fish, the mucosal barrier covering the gills, skin, and intestinal surfaces are considered the first line of defense against invading pathogens (53, 54). The presence of chitinases and CBP may act to degrade not only the catfish slime coat, but also to bind to and degrade the epithelial mucins in the digestive tract, increasing vAh invasiveness. Peatman et al (2018) reported a direct link between feed consumption and vAh-induced MAS, with survival in vAh-challenged catfish decreasing significantly when fish were fed to satiation 4 hours prior to challenge. The mucus coating of the intestinal epithelium may decrease after eating, as ingesta moves through the digestive tract and takes mucus with it. Chitinases and CBPs may then be capable of breaking down the remaining mucus, gaining access to the underlying epithelium and, eventually, the bloodstream (55). The presence of chitinase and CBP could help explain the intestinal epithelial damage found on necropsy in fish naturally infected with vMAS (56). Although significantly higher in biofilm secretomes, chitinase and CBP was prominent in both planktonic and biofilm secretomes, suggesting they play an important role in bacterial fitness regardless of growth condition.

Whereas biofilm secretomes were flush with degradative exoenzymes, such as elastase, chitinases, and multiple Zn-dependent and metalloproteases, planktonic secretomes consistently produced more hemolytic and cytotoxic ECPs. Notably, both aerolysin-type and *ahh1*-type hemolysins were detected in much higher quantities in planktonic secretomes, as were two extracellular serine proteases (neither of which were identified in any biofilm sample) and extracellular lipases, all of which exert hemolytic activity against erythrocytes, and have been shown to be cytotoxic to cells (20, 57). Interestingly, the alpha-hemolysin, phospholipid-cholesterol acyltransferase, which was present in planktonic secretomes but
absent in biofilm, has been reported to produce significant lysis of salmon erythrocytes following activation by serine protease (58). The presence of substantial amounts of both proteins in the planktonic secretomes suggests that the production of these proteins could allow a multi-pronged approach to cell death, with each toxin acting independently, but increasing the collective virulence resulting from multiple exoproteins. Aerolysin-type hemolysin has been implicated as the main virulence factor of *A. hydrophila* (20), and was significantly higher in planktonic secretomes, with a three-fold increase compared to biofilm. However, ahh1-type hemolysin was present in planktonic secretomes at greater than three times the amount of aerolysin-type hemolysin. Ahh1 hemolysins are homologous to hlyA hemolysins of *V. cholerae* (59). The activity of this pore-forming hemolysin is not erythrocyte-specific, but targets erythrocytes, leukocytes, lymphocytes, and epithelial and endothelial cells in a multitude of eukaryotes (60) and, as such, are considered cytotoxins. This supports the *in vitro* hemolysis assay results that found 80% hemolysis of channel catfish erythrocytes in one hour when exposed to planktonic supernatants, compared to less than 15% average hemolysis of erythrocytes that were incubated with biofilm supernatants (Figure 3). The presence of these hemolysins and other cytotoxins in planktonically-cultured vAh may also help explain the rapid mortality seen in catfish when challenged by intraperitoneal injection, as these bacteria may be primed to produce vast amounts of toxins *in vivo*.

Biological functions of secreted proteins as analyzed by gene ontology found carbohydrate utilization to be the dominant function of secreted proteins under both conditions. Proteins involved in hemolysis, lipid and nucleotide catabolism, arginine biosynthesis, protein folding and transport were dominant biological functions of planktonic secretomes. Significant biofilm proteins were largely involved in transmembrane transport, amino acid processing, and transport of ions, amino acids, and carbohydrates. Interestingly, flagellar motility was also important in biofilms. This is likely due to *A. hydrophila*’s use of flagella in biofilm construction and not for bacterial motility (38). This increase in polar flagella may also contribute to an increased host colonization in biofilm-associated vAh. While lateral flagella are often considered imperative for biofilm production and adhesion (33, 61), Aeromonads that lack lateral flagella are capable of using polar flagella for biofilm formation as well as cellular adhesion (34, 38, 62, 63). The increased polar flagella required for biofilm formation could act secondarily as adhesins when biofilm-derived bacteria come into contact with catfish mucosal surfaces and could act in concert with other secreted invasins to colonize and destroy host mucosal barriers.

**Conclusions**

Most aquatic bacterial generalists, such as *A. hydrophila*, spend the majority of time resident in biofilms and host-microbe interactions are likely influenced by niche-specific microbial phenotype. Because biofilm-associated bacteria have emergent properties that cannot be elucidated by the study of free-living cells, it is imperative to study organisms within biofilms to understand how niche adaptations may influence overall pathogenicity and virulence. This study is the first comparison of the secreted proteomes of vAh when grown in two distinct ecological niches. These data on the adaptive physiological response of vAh based on growth condition increase our understanding of how environmental niche partitioning could affect vAh pathogenicity and virulence. Increased secretion of colonization factors and
degradative enzymes during biofilm growth and residency may increase bacterial attachment and host invasiveness, while increased secretion of hemolysins, porins, and other potential toxins under planktonic growth (or after host invasion) could result in increased host mortality. These shifts in protein expression and secretion indicate that growth under biofilm and planktonic conditions results in massive changes in gene expression. Future research should explore the global regulatory factors that affect vAh gene expression under these growth conditions. Taken together, these data may help in our understanding of the unique aspects of this emerging pathogen that contribute to the devastating impact of MAS disease outbreaks.

Methods And Materials

Bacterial Strain

vAh strain ML09-119 was isolated from a diseased channel catfish from a MAS outbreak in a West Alabama aquaculture facility in 2009. Molecular characterization and genome sequencing of vAh ML09-119 have been performed (19) and the complete genome sequence deposited in GenBank (Accession CP005966). Aliquots of vAh ML09-119 were cryogenically stored in 10% glycerol freeze medium at -80°C.

Catfish: Specific-pathogen free channel catfish fingerlings maintained under Auburn University IACUC-approved protocol 2018-3251 (Catfish Production and Maintenance) were used for challenges. All challenges were performed adhering to the guidelines of AU-IACUC-approved protocol 2016-2900 (Identification of toxigenic proteins of virulent Aeromonas hydrophila and evaluation of host response).

Culture Media and Culture Conditions

Tryptic soy broth (TSB) (Bacto TSB, BD) prepared according to manufacturer's directions was used as the culture medium for planktonic growth.

Biofilm media was prepared by adding 0.2% agar powder (AlfaAesar) to TSB media prior to sterilization. Approximately 70 ml of molten biofilm agar was poured into deep well petri dishes (Fisher) and allowed to solidify. Bacterial strain vAh ML09-119 was removed from cryogenic storage and inoculated into 25 ml TSB media and grown overnight at 30°C with shaking. A 1 ml aliquot of overnight culture was transferred to 70 ml of TSB and grown at 30°C on an orbital shaker to mid-log phase, approximately 16 hours. Biofilm agar plates were inoculated from overnight culture by stab inoculation. Plates were sealed with parafilm and incubated at 30°C for 72 hours, until an adherent bacterial film covered the agar surface.

Planktonic and biofilm cultures were performed in triplicate.

Secretome Preparation

Planktonic Secretome: vAh ML09-119 was cultured as described above. Cells were pelleted by centrifugation at 20,000 x g for 15 minutes at 4°C and supernatant was decanted and retained. Cells were washed twice with cold, sterile PBS, pelleted as above, and the wash was added to the supernatant.
Remaining cells were removed by passage through a low-binding 0.22 µm vacuum filter (VWR). Cell-free supernatants were used as the starting point for purification of extracellular proteins (ECPs).

**Biofilm Secretome:** vAh ML09-119 cells were gently removed from the biofilm media surface with a sterile cell scraper, transferred to 50 ml conical tube, and washed twice with cold, sterile PBS as described above. The cell wash was decanted and retained. To collect secreted proteins within biofilm media, the plates were disrupted using a sterile disposable probe until the soft agar had formed a slurry. The agar slurry was transferred to a sterile 50 ml conical tube and centrifuged at 20,000 x $g$ for 15 min at 4°C to pellet the agar. Following centrifugation, the liquid media was decanted from the agar plug and retained. The agar plug was then resuspended in 20 ml cold sterile PBS, centrifuged as above, and the wash solution decanted and retained. All wash solutions and liquid media were combined and filtered, first through a low-binding 0.45 µm vacuum filter (VWR), then through a low-binding 0.22 µm vacuum filter to remove any residual agar and bacterial cells. This cell-free supernatant was used at the starting point for biofilm ECP purification.

**Ammonium Sulfate Precipitation:** ECPs were precipitated from cell-free supernatants by the addition of ammonium sulfate crystals (Fisher Scientific) to achieve 65% saturation, followed by incubation at 4°C on a rotary platform shaker with gently mixing for 24 hours. Precipitated proteins were collected by centrifugation at 30,000 x $g$ for 45 min at 4°C, then dissolved in 10 ml cold Tris buffer (20mM Tris-Hcl, pH 7.6) + protease inhibitor (Complete tablets, mini, EDTA-free (Roche)). Resuspended proteins were dialyzed twice, for 18 hours and 12 hours, respectively, against the same buffer in 10 Kda dialysis cassettes (Slide-A-Lyzer (Thermo Fisher)). After dialysis, the total volume was adjusted to 20 ml by the addition of cold Tris buffer. The protein concentration of each sample was determined by the Bradford assay (Pierce Coomassie Plus Protein Assay, Thermo Fisher). These concentrated proteins were used for all assays.

**Enzymatic Activity**

The *in vitro* activity of secreted proteins was measured using specific substrates to determine the degradative and toxigenic potential of planktonic and biofilm secretomes, as described below:

**Hemolysis:** Hemolytic potential was measured using the method of Peatman et al. (2018) with some modifications. In brief, heparinized blood from three channel catfish was pooled and diluted 1:10 in sterile phosphate buffered saline (PBS). A suitable dilution of protein in 150 µl PBS buffer was added to 25 µl diluted blood in sterile microcentrifuge tubes. Tubes were incubated at 30°C in an orbital shaker for 2 h. Positive control tubes representing 100% hemolysis contained 150 µl sterile distilled water in place of protein samples. Negative control tubes contained 150 µl sterile PBS in place of protein samples. Controls were incubated with 25 µl diluted blood as above. Following incubation, tubes were centrifuged at 1,000 x $g$ to pellet un-lysed cells and 150 µl of supernatant was transferred to 96-well flat bottom plates. Erythrocyte lysis was quantified by measuring absorbance of released hemoglobin at 415 nm in multi-mode plate reader (Synergy HTX, Bio-Tek) and hemolysis was reported as percent of positive control.
Universal Protease Activity: Non-specific proteolytic activity was measured using HiLyteFluor 488-labeled casein as the substrate, following manufacturer's protocol with minor modifications (Sensolyte Green Fluorimetric Protease Assay Kit, AnaSpec, Inc.). Briefly, a suitable concentration of protein in 50 µl deionized water was added to triplicate wells of black, flat-bottom 96-well plates with non-binding surface (Greiner Bio-One). Trypsin, diluted 50-fold in deionized water, acted as a positive control and sterile deionized water served as a substrate control. Following the addition of 50 µl labeled casein substrate, plates were mixed briefly and fluorescent intensity was measured at Ex/Em = 490 nm/520 nm every five minutes for one hour in a multi-mode plate reader (Synergy HTX, Bio-Tek) with 30°C incubation temperature. Data were plotted as relative fluorescence units versus time for each sample.

Elastase Activity: Elastase-specific activity was measured using 5-FAM/QXL™ 520 labelled elastin as the substrate, following the manufacturer's protocol with minor modifications (Sensolyte Green Fluorimetric Elastase Assay Kit, AnaSpec, Inc.). Briefly, a suitable concentration of protein in 50 µl deionized water was added to triplicate wells of black, flat-bottom 96-well plates with non-binding surface. Elastase, diluted 50-fold in assay buffer, acted as a positive control and sterile, deionized water was a substrate control. Following the addition of 50 µl labeled elastase substrate, plates were mixed briefly and fluorescent intensity was measured continuously at Ex/Em = 490 nm/520 nm, and data recorded every five minutes for one hour in a multi-mode plate reader (Synergy HTX, Bio-Tek) with 30°C incubation temperature. Data were plotted as relative fluorescence units versus time for each sample.

In vivo Proteolysis

Extracellular protein activity was measured in vivo using channel catfish fingerlings to determine potential proteolytic and cytotoxic tissue effects.

Protein Preparation: Ten microgram aliquots of secreted planktonic and biofilm-associated proteins, prepared as above, diluted in 100 µl sterile PBS were used for injection challenges.

Challenge Model: Channel catfish fingerlings were transferred to 57-liter glass aquaria containing dechlorinated municipal water and acclimated at 30°C for two days prior to challenge. Triplicate tanks containing five fish each represented planktonic ECP, biofilm-associated ECP, and injection control groups. Prior to injection, fingerlings were transferred to sedation aquaria containing 70 mg/ L tricaine methanesulfonate (MS-222) buffered to neutrality with sodium bicarbonate. Following sedation, characterized by decreased opercular movement and loss of equilibrium, 100 µl of sterile PBS containing 10 µg of total protein was injected intramuscularly just below the dorsal fin using tuberculin syringes fitted with 26 gauge needles. Control fish were injected with 100 µl sterile PBS. Fish were then returned to the appropriate aquarium and monitored until fully recovered. Fish were maintained in aquaria at 30°C for 7 days under flow-through conditions at 1 gallon per hour water replacement. Moribund fish or fish developing severe external lesions were euthanized by prolonged exposure to buffered MS-222, the tissues were collected and fixed in 10% neutral-buffered formalin. After 7 days, remaining fish were humanely euthanized and samples were collected and prepared as above.
Histology: Formalin-fixed tissues were paraffin-embedded and 4 micron sections were prepared and stained with hematoxylin and eosin according to standard methods (64). Slides were evaluated and photographed using an Olympus BX53 microscope fitted with an Olympus DP26 digital camera.

Secretome Analysis.

To determine how vAh niche occupancy might influence protein production, secreted protein profiles of vAh cultured within a biofilm and in broth were compared by liquid chromatography with tandem mass spectrometry (LC MS/MS) analysis at the UAB Mass Spectrometry/Proteomics shared facility to identify and quantify proteins present in each sample, as previously described (30), as follows.

Proteomics analysis: Samples were prepared for analysis as follows: 20µg of protein per samples was diluted to 35µl in NuPAGE LDS sample buffer (Invitrogen), reduced with dithiothreitol, and denatured at 70°C for 10 minutes prior to loading onto Novex NuPage 10% Bis-Tris protein gel (Invitrogen). The gel was separated as a short stack (10 min, 200V constant) and stained overnight with Novex Colloidal Blue Staining kit (Invitrogen). Gels were destained and each lane was cut into single molecular weight fractions and equilibrated in 100mM ammonium bicarbonate. Each plug was then digested overnight with Trypsin Gold (Mass Spectrometry grade (Promega)) following manufacturer's instructions and peptide extracts were reconstituted to 0.1µg/µl in 0.1% formic acid.

Mass Spectrometry: Prepared peptide digests (8µl) were injected onto a 1260 Infinity nHPLC stack (Agilent Technologies) and separated using a 71µ l.d. X 15cm pulled-tip C-18 column (Jupiter C-18 300 Å, 5 micron (Phenomenex)) running in-line with a Thermo Orbitrap Velos Pro hybrid mass spectrometer, equipped with a nano-electrospray source (Thermo Fisher). All data were collected in CID mode. nHPLC was configured with binary mobile phases comprised of 0.1% formic acid (solvent A) and 0.1% formic acid in 85% acetonitrile (solvent B). Following each parent scan (300-1200 m/z at 60K resolution), fragmentation data (MS2) were collected on the top most intense 15 ions. For data-dependent scans, charge-state screening and dynamic exclusion were enabled with a repeat count of 2, repeat duration of 30s, and exclusion duration of 90s.

Mass spectrometry data conversion and searches: Xcalibur RAW files were collected in profile mode, centroided, and converted to mzXML using ReAdW v3.5.1 (IonSource). The mgf files were then created using MzXML2Search (included in Trans-Proteomics Pipeline v3.5) for all scans. The data were searched using SEQUEST (Thermo Fisher, San Jose, CA, USA; version 27), which was set for two maximum missed cleavages, a precursor mass window of 20ppm, trypsin digestion, variable modification C at 57.0293, and M at 15.9949. Searches were performed with a species-specific subset of the UniRef 100 database.

Peptide filtering, grouping, quantification and statistical analyses: Scaffold (v. 4.8.4, Proteome Software Inc., Portland, Oregon) was used to validate MS/MS based peptide and protein identifications. The list of peptide IDs generated based on SEQUEST search results were filtered using Scaffold, which filters and groups all peptides to generate and retain only high confidence IDs while also generating normalized spectral counts across all samples to allow for relative quantification. Filter cut-off values required to
accept peptide identifications were set with minimum peptide length of >5 amino acids, with no MH+ charge states, peptide probabilities of >80% C.I., and with the number of unique peptides per protein ≥2. Peptide probabilities were assigned by the PeptideProphet algorithm (65). Protein identifications were accepted if proteins probabilities could be established at >99% C.I., contained at least 4 identified peptides, and with false discovery rate <1.0. Protein probabilities were assigned by the ProteinProphet algorithm (66). Scaffold incorporates the two most common methods for statistical validation of large proteome data sets, the false discovery rate (FDR) and protein probability. Relative quantification across samples were then performed via spectral counting and, when relevant, spectral count abundances were normalized between samples. Proteins present in at least two experimental replicates were included in analyses. To determine statistical significance, two non-parametric statistical analyses were performed between each pair-wise comparison, including reproducibility-optimized test statistic (ROTS) (bootstrapping value = 1000) combined with single-tail t-test (p < 0.05) (67, 68). These were then sorted according to the highest statistical relevance in each comparison. For protein abundance ratios determined by normalized spectral counts, a fold change threshold ≥1.5 was set for significance. Protein abundance of proteins present in only one experimental group was set as the average of the normalized quantitative value.

**Protein Function:** To define the potential function of secreted proteins, major biological processes of statistically significant proteins were determined from gene ontology annotation in UniProt (Consortium, T.U. 2018) and QuickGO (69). Predicted protein function was assessed by determining major biological processes through gene ontology. Using these data, eight functional groups were established, and proteins were sorted into these groups based on their primary biological function. A further comparison was made by compiling all proteins in each functional group from both biofilm and planktonic secretomes and expressing as parts of a whole, with side-by-side comparisons for each secretome type.

**Statistical Analyses**

Reproducibility-optimized test statistic (ROTS) analysis of differentially secreted proteins was performed in R (70). All other statistical analyses were performed in Prism 8.2.0 (Graphpad). One-way ANOVA followed by Tukey's multiple comparisons post-test were performed on triplicate data with significance set at p < 0.05. Graphical representations of data were produced in Prism 8.2.0.

**Declarations**

**Ethics approval and consent to participate**

All animal challenges were approved by the Auburn University Institutional Animal Care and Use Committee (AU-IACUC) and were performed adhering to the guidelines of AU-IACUC-approved protocol 2016-2900 (Identification of toxigenic proteins of virulent *Aeromonas hydrophila* and evaluation of host response).

**Consent for publication**
Availability of data and materials

The datasets generated and analyzed during the current study are included as supplementary files (Additional File 1) or from the corresponding author on reasonable request.

Competing Interests

The authors declare that they have no competing interests.

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Author Contributions

PB, ML, and JN conceived and designed the experiments. PB and JN performed in vivo challenges and JN performed histology. PB performed other experiments and analyzed data. BB provided equipment and critical review. PB, ML, BB, and JN contributed to the writing and editing of the manuscript.

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Tables

Table 1: Differentially secreted proteins of planktonic and biofilm-cultured ML09-119
| Secreted Protein                        | Locus Tag    | ROTS-Statistic | p value | FDR  | Protein Abundance Fold Change | Significant Experimental Group |
|----------------------------------------|--------------|----------------|---------|------|-------------------------------|--------------------------------|
| **Transcription regulation/ e-Transport** |              |                |         |      |                               |                                |
| DNA gyrase inhibitor                   | AHML_21105   | 2.2            | 0.01    | 0    | 16*                           | BIO                            |
| Ribonuclease activity regulator        | AHML_16315   | 1.55           | 0.02    | 0.02 | 2.5                           | BIO                            |
| FKKP-type peptidyl-prolyl cis-trans isomerase | AHML_05355   | -2.10          | 0.01    | 0    | 22*                           | TSB                            |
| Cytochrome d ubiquinol oxidase         | AHML_19200   | -1.71          | 0.02    | 0.02 | 16                            | TSB                            |
| **Free Radical Scavenging**            |              |                |         |      |                               |                                |
| Superoxide dismutase                   | AHML_07590   | 2.95           | 0.01    | 0    | 6                             | BIO                            |
| Glyoxalase/dioxygenase protein         | AHML_09045   | 1.89           | 0.01    | 0.03 | 13*                           | BIO                            |
| **Amino Acid/Cofactor Metabolism**     |              |                |         |      |                               |                                |
| Redox protein (hypothetical)           | AHML_05250   | 2.33           | 0.01    | 0    | 19*                           | BIO                            |
| Diaminopimelate epimerase              | AHML_02440   | 1.82           | 0.02    | 0.03 | 9*                            | BIO                            |
| Ornithine carbamoyltransferase         | AHML_21555   | -1.35          | 0.03    | 0.05 | 3.5                           | TSB                            |
| Riboflavin-biosynthesis protein        | AHML_17795   | 1.4            | 0.02    | 0.05 | 2.4                           | BIO                            |
| Urocanate hydratase                    | AHML_01870   | 3.71           | 0.00    | 0    | 8                             | BIO                            |
| Dihydridopicolinate synthase           | AHML_04540   | 2.35           | 0.01    | 0    | 22*                           | BIO                            |
| Succinylarginine dihydrolase           | AHML_16715   | 5.49           | 0.00    | 0    | 66*                           | BIO                            |
| **Carbohydrate Metabolism**            |              |                |         |      |                               |                                |
| Maltose operon periplasmic protein     | AHML_06220   | 2.57           | 0.01    | 0    | 15*                           | BIO                            |
| Phosphoglyceromutase                   | AHML_01445   | 1.94           | 0.01    | 0    | 13*                           | BIO                            |
| Transaldolase B                       | AHML_16890   | 8.08           | 0.00    | 0    | 89*                           | BIO                            |
| Pullulanase                            | AHML_04415   | 3.62           | 0.00    | 0    | 10                            | BIO                            |
| Ribose-5-phosphate isomerase A         | AHML_14480   | 2.92           | 0.01    | 0    | 3                             | BIO                            |
| Beta-glucosidase                       | AHML_14270   | -1.68          | 0.02    | 0.0  | 14                            | TSB                            |
| **Outer Membrane Proteins**            |              |                |         |      |                               |                                |
| Outer membrane protein A               | AHML_21905   | -1.99          | 0.01    | 0    | 4                             | TSB                            |
| Outer membrane protein A               | AHML_20145   | 1.46           | 0.02    | 0.04 | 13*                           | BIO                            |
| Outer membrane lipoprotein             | AHML_00700   | -1.58          | 0.02    | 0.02 | 13*                           | TSB                            |
| **Secreted Protein**                   |              |                |         |      |                               |                                |
| Hemolysin (Aerolysin-type)             | AHML_02265   | -2.37          | 0.01    | 0    | 3                             | TSB                            |
| Hemolysin (ahlh1-type)                 | AHML_08400   | -2.69          | 0.01    | 0    | 4                             | TSB                            |
| Elastase                               | AHML_04340   | 4.68           | 0.00    | 0    | 5                             | BIO                            |
| Chitinase                              | AHML_05225   | 4.07           | 0.00    | 0    | 3                             | BIO                            |
| Metalloprotease                        | AHML_05230   | 2.73           | 0.01    | 0    | 3                             | BIO                            |
| Basic endochitinase                    | AHML_05235   | 3.15           | 0.01    | 0    | 3                             | BIO                            |
| Zn-dependent carboxypeptidase           | AHML_05535   | 2.15           | 0.01    | 0    | 13*                           | BIO                            |
| Outer membrane porin protein           | AHML_04355   | 2.91           | 0.01    | 0    | 15*                           | BIO                            |
| Extracellular lipase                   | AHML_00550   | -4.16          | 0.01    | 0    | 3                             | TSB                            |
| Chitin-binding domain 3                | AHML_11110   | -1.50          | 0.02    | 0.02| 12                            | TSB                            |
| Zn-dependent protease with chaperone function | AHML_06635  | 1.51           | 0.02    | 0.02| 11*                           | BIO                            |
### Table 1.

| Secreted Protein                                                                 | Locus Tag | ROTS-Statistic | p value | FDR  | Protein Abundance Fold Change | Significant Experimental Group |
|---------------------------------------------------------------------------------|-----------|----------------|---------|------|------------------------------|--------------------------------|
| **Pilus and Flagellin Proteins**                                                |           |                |         |      |                              |                                 |
| Flagellin                                                                       | AHML_09350 | 5.42           | 0.00    | 0    | 43*                          | BIO                            |
| Flagellin-like protein                                                          | AHML_09345 | 2.05           | 0.01    | 0    | 2.6                          | BIO                            |
| Type I pilus assembly protein FimF                                                | AHML_02690 | -2.29          | 0.01    | 0    | 26*                          | TSB                            |
| Fimbrial Protein                                                                | AHML_02665 | -2.61          | 0.01    | 0    | 36*                          | TSB                            |
| **Transport Proteins**                                                          |           |                |         |      |                              |                                 |
| ABC-type sugar transport                                                         | AHML_20895 | 1.77           | 0.02    | 0.02| 14*                          | BIO                            |
| TonB-dependent copper receptor                                                  | AHML_02545 | 1.79           | 0.02    | 0.02| 9                            | BIO                            |
| Peptide ABC transporter                                                         | AHML_17755 | 3.2            | 0.00    | 0    | 98*                          | BIO                            |
| Arginine ABC transporter                                                        | AHML_03370 | 4.58           | 0.00    | 0    | 7                            | BIO                            |
| Oligopeptide ABC transporter                                                    | AHML_13875 | 6.05           | 0.00    | 0    | 93*                          | BIO                            |
| Leucine binding protein (ABC transport)                                         | AHML_00595 | 3.74           | 0.00    | 0    | 18                           | BIO                            |

Differentially secreted proteins of vAh ML09-119 cultured planktonically (TSB) and within a biofilm (BIO). Proteins are grouped based on their major biological process, determined by gene ontology. Protein abundance fold change marked with * denotes protein identified in only one condition and is reported as the average Quantitative Protein Value. Proteins and locus tags in Bold indicate putative virulence proteins. FDR = False Discovery rate.