MODELING COASTAL PATHOGEN POPULATIONS: ADVANCEMENTS IN PREDICTIVE MODELING OF *VIBRIO VULNIFICUS* IN A TROPICAL URBAN ESTUARY IN HONOLULU, HAWAIʻI

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This thesis is dedicated to my mother and grandmother who lit a path for women in the fields of biology and chemistry.

I am forever inspired by their strength and passion.
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ABSTRACT

Vibrio vulnificus is a deadly pathogenic bacterium found worldwide in estuarine habitats. Predicting the presence and abundance of V. vulnificus is becoming more critical as sea level rises into densely populated urban areas and increasing water temperatures accelerate growth rates and expand habitat range. I explored two methods of modeling microbial populations, (1) statistical correlation with environmental covariates and (2) numerical modeling of growth and mortality rates, in order to improve predictive modeling capabilities of V. vulnificus in the Ala Wai Canal in Honolulu, Hawai‘i.

Chapter one of this thesis details research testing the hypothesis that incorporating measurements of dissolved organic and inorganic nutrients will improve the predictive power of statistical and forecasting models of V. vulnificus. In this field-based study, I parameterized linear regression models of V. vulnificus abundance using physical, optical, and chemical water properties in high spatial and temporal resolution to develop a real-time spatially-explicit model of V. vulnificus pathogen risk. A model including fine scale organic matter characterization and a small set of easily measured, highly significant environmental predictors could explain nearly 70% of variation in V. vulnificus abundance over an annual cycle. Using an oceanographic circulation model, I show how these models can be applied to predict plumes of V. vulnificus from the canal into the coastal environment. I also describe a framework for how continuous monitoring technology can be employed to nowcast and forecast the abundance of an aquatic pathogen. Long-term forecasts based on climate change projections estimate an overall increase in V. vulnificus density of up to 40% by the end of the century due to increasing temperature. These novel findings will have significant ramifications to coastal pathogen monitoring programs and ongoing efforts to predict how climate change will alter the dynamics of estuarine microbial ecosystems.

Chapter two compares experimental approaches to calculate mortality rates of natural populations of V. vulnificus in order to improve an existing numerical population model of V. vulnificus. This model uses the environmental constraints of growth rates coupled to a physical circulation model, however, model predictions are inconsistent with field observations. Characterizing the environmental constraints of mortality rates would likely improve the model’s
accuracy and usefulness as a tool to communicate exposure risk. I tested the eukaryotic inhibition and dilution experiment techniques in order to quantify the contributions of grazing and viral lysis to natural mortality of the microbial community as a whole and *V. vulnificus*. My results show that eukaryotic inhibition may be a more effective technique to calculate mortality from predation than the dilution experiment technique. I outline several recommendations to improve the experimental design for interpretation and application. Ultimately, improving these predictive models of microbial populations is critical for management of waterborne pathogen risk exposure, particularly given the uncertainty of a changing global climate. This research represents a significant step forward to developing robust predictive modeling frameworks for coastal pathogens.
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Incorporating the spatiotemporal dynamics of dissolved nutrients and organic matter improves predictive models of *Vibrio vulnificus* concentrations in a tropical urban estuary

Keywords: *Vibrio vulnificus*, Pathogen, Estuary, Hawai‘i, Predictive Modeling, Climate Change, Nutrients, Dissolved Organic Matter, Groundwater

1.1 ABSTRACT

The south shore of O‘ahu, Hawai‘i is one of the most visited coastal tourism areas in the United States with some of the highest instances of recreational waterborne disease. The pathogenic bacterium, *Vibrio vulnificus*, lives in the warm temperature and intermediate salinity waters of the estuarine Ala Wai Canal in Honolulu which surrounds the heavily populated tourism center of Waikīkī. Heavy rainfall has been shown to transport water from the canal into the nearshore coastal system threatening human and ecological health as well as the state’s tourism-based economy. I developed a biological-physical statistical model to predict *V. vulnificus* dynamics using environmental measurements that can be captured with moored oceanographic sensors in real time. I parameterized and validated the model using a novel approach that integrates data from three sources: 1) monthly profile measurements at each of 18 sites spanning an annual cycle which included *in situ* discrete sampling at 3 depths (surface, pycnocline, and bottom water) and continuous sensor profiling, 2) moored instrument continuous time series from the Pacific Islands Ocean Observing System (PacIOOS) and 3) the Waikīkī Regional Ocean Modeling System (ROMS). Discrete samples (n = 213) were analyzed for 36 biogeochemical variables and *V. vulnificus* density using quantitative PCR of the hemolysin A toxicity gene (vvhA). I found significant monthly variation in *V. vulnificus* density ranging over 4
orders of magnitude largely predictable by seasonality in biogeochemistry. The best multiple regression model of *V. vulnificus* concentration, explaining 70% of variation, included salinity, five-day average rainfall, daily maximum air temperature, silicate, dissolved oxygen and two metrics of dissolved organic matter fluorescence (visible humic-like fluorescent dissolved organic matter and the Humification Index). This result suggests a consistent response of *V. vulnificus* to freshwater inputs, particularly groundwater, and changes in organic matter resources. These linear models, paired with the PacIOOS time series and ROMS products, were then implemented to predict *V. vulnificus* density in the canal and coastal waters in real time across the nearshore Waikīkī region and were validated against measured values. Long-term climate model projections of local rainfall and air temperature were used to predict an overall increase in *V. vulnificus* density of approximately 20 to 40 % by 2100. Improving these predictive models of microbial populations is critical for management of waterborne pathogen risk exposure, particularly in the wake of a changing global climate.
1.2 INTRODUCTION

**Vibrio vulnificus Biology and Vibriosis Infection**

*Vibrio* is a genus of Gram-negative heterotrophic bacteria with diverse life history strategies as free-living, host-associated, or particle-attached members of marine and estuarine microbial communities worldwide (Baker-Austin et al., 2018; Heng et al., 2017; Huang et al., 2016). Several *Vibrio* species are known human pathogens, including *V. cholerae*, *V. parahaemolyticus*, and *V. vulnificus*. Vibriosis infections occur through foodborne or recreational exposure with an annual healthcare cost to the U.S. nearing $300 million with approximately 85% of those costs attributed to *V. vulnificus* alone (Ralston et al., 2011). *V. vulnificus* is found in both free-living in brackish water and associated with sediment, algae, fish, molluscs, crustaceans, and zooplankton in warm estuarine environments globally (DePaola et al., 1994; Givens et al., 2014; Johnson et al., 2010; Maugeri et al., 2006; Turner et al., 2009). Although uncommon (approximately 100 cases are reported in the U.S. annually), the majority of *V. vulnificus* infections are from wound exposure to contaminated water, resulting in cellulitis and necrotizing fasciitis with a 25% case-fatality rate (Horseman & Surani, 2011). There is minimal monitoring for *V. vulnificus* in coastal waters worldwide (Baker-Austin et al., 2018), particularly at ecologically relevant scales that would allow for reliable forecasting of pathogen risk of exposure in contaminated waters. Prediction of the presence and abundance of *V. vulnificus* will become even more critical as sea level rises into populated areas and increasing water temperatures accelerate the pathogen’s growth rate. Unlike many other aquatic human pathogens, *V. vulnificus* concentrations cannot be predicted effectively with fecal indicator bacteria because the pathogen lives naturally in estuarine habitats (Strom & Paranjpye, 2000).

*The Estuarine Habitat of Vibrio vulnificus*

Estuaries are heterogenous and idiosyncratic with unique biogeochemical features such as sharp gradients over short geographic areas and small depth ranges, high biological diversity and activity, and anoxic sediments (Cai, 2011). They are known to be effective traps for nutrients, sediments, and pollutants from inflowing rivers and runoff (Shiller, 1996). By nature of their proximity to populous areas, estuaries are at high risk of anthropogenic impact from
development and pollution. Due to complex and location-specific hydrology, oceanography, climatology, and human development, it is unknown how climate change will modify these environments, particularly estuarine microbial communities and the abundance of human pathogens (Ghosh & Bhadury, 2019; Kan et al., 2007).

**Vibrio vulnificus Response to Climate Change**

There is empirical evidence for an increased abundance of *V. vulnificus* in bacterioplankton communities and infection rates over the past 50 years (Newton et al., 2012) which has been linked to increased sea surface temperature (Vezzulli et al., 2012; 2013; 2016). Additional climatological changes, such as increased frequency of hurricanes, storm surges, sea level rise, and coastal flooding, have been hypothesized to increase the geographic distribution of *V. vulnificus* and the incidence of *V. vulnificus* infections (Martinez-Urtaza et al., 2010; Muhling et al., 2017; Paz et al., 2007). Muhling et al. (2017) found significant increases in season length and spatial habitat for *V. vulnificus* in the Chesapeake Bay based on statistical downscaling of climate change projections. There is evidence for habitat expansion into higher latitudes (Baker-Austin et al., 2013; Paz et al., 2007) and a response of *V. vulnificus* to drought conditions (Wetz et al., 2013). Baker-Austin et al. (2017) even argue that *V. vulnificus*, given its response to temperature, be used as a barometer of climate change in marine environments. Despite these concerns, routine monitoring of natural *V. vulnificus* populations is infrequent in the U.S. Predictive models, based on the environmental conditions that govern *V. vulnificus* distribution and growth, may be a viable alternative to survey-based monitoring in order to provide an early warning of the risk of infection. Previous work by Jacobs et al. (2014) has shown promise at constructing such predictive spatially-explicit models. However, these efforts have been limited to probability of occurrence of *V. vulnificus* rather than absolute abundance which is more directly related to the risk of exposure.

**Environmental Controls of Vibrio vulnificus**

The environmental controls of *V. vulnificus* populations are poorly constrained aside from well-established effects of salinity and temperature on growth rates in laboratory cultures (Kaspar & Tamplin, 1993; Motes et al., 1998). Surveys and laboratory experiments have
documented that *V. vulnificus* is most prolific in water temperatures above 18 °C and salinity between 15 and 25 (Huehn et al., 2014; Jones & Oliver, 2009; Randa et al., 2004). There are several studies that assess the effects of additional parameters, such as turbidity, chlorophyll, nutrients and pH, on *V. vulnificus* growth and distribution (Lipp et al., 2001; O’Neill et al., 1992; Pfeffer et al., 2003; Tamplin et al., 1982; Wright et al., 1996). Fewer address the *in situ* environment with multiple interacting factors affecting growth and mortality. Environmental measurements of *V. vulnificus* density show strong seasonal dynamics where *V. vulnificus* density is higher in warm summer months in temperate and subtropical areas (Jacobs et al., 2014; Pfeffer et al., 2003; Tamplin et al., 1982). However, there is minimal data from tropical areas where water temperatures remain above 18 °C year-round and other environmental controls may be more important determinants of *V. vulnificus* density.

*Previous work on Vibrio vulnificus in Hawai‘i*

In Hawai‘i, recreational exposure to *Vibrio* contaminated water is likely high, and rates of *Vibrio* infections are reportedly five times higher than any other U.S. state per capita (National Center for Emerging & Infectious Diseases). *V. vulnificus* is found ubiquitously in the Ala Wai Canal, a prominent waterway in Honolulu, Hawai‘i (Nigro, 2012). Previous surveys of the canal have shown that variability in *V. vulnificus* abundance scales with temporal resolution (Nigro, 2012). The range and variability of *V. vulnificus* abundance is smaller for shorter time scales (weekly, daily, hourly) than for monthly or seasonal patterns in abundance. Therefore, I focused my research in this study on capturing aspects of seasonal variation as important determinants of *V. vulnificus* variation. Preliminary work from 2008 – 2009 found significantly higher abundances of *V. vulnificus* in the canal during the rainy season from October – March (Nigro, 2012). In addition, Nigro (2012) found evidence to suggest a varied response of *V. vulnificus* to different freshwater sources with a salinity optimum of 12. There also appeared to be different dynamics between the infectious C-type strain and noninfectious E-type strain where the proportion of C-type increased with higher salinity values. However, the Nigro (2012) study was limited to surface sampling and a limited number of environmental measurements (salinity, temperature, chlorophyll-a, particulate carbon, phosphate, nitrate, nitrite, and ammonium) from bottle samples. These previous results hint at a response of *V. vulnificus* to freshwater flow and
nutrient resources that are useful for characterizing environmental controls but require further exploration for statistical and predictive modeling. Previous modeling attempts using a coupled physical circulation and numerical model of *V. vulnificus* growth rate in response to salinity and temperature by Nuss (2016) resulted in significant discrepancies between modeled and measured *V. vulnificus* abundance and patterns of variation in the canal. The addition of nutrient and organic matter resources into predictive models of *V. vulnificus* abundance may therefore improve predictive capabilities.

**Project Objectives**

In order to parameterize the distribution and abundance of this pathogen in the Ala Wai Canal and adjacent recreational coastal waters, I measured temporal and spatial variation in *V. vulnificus* over an annual cycle in concert with an array of biogeochemical parameters. Using these dynamic relationships, I built a spatially-explicit predictive model to assist mitigation efforts and limit the threat to human health. Past work has demonstrated that salinity and temperature variation alone do not sufficiently explain *V. vulnificus* population dynamics for predictive modeling, especially for this tropical location where water temperature is always in an optimal range. I hypothesized that model parameterization with measurements of nutrient and organic matter concentrations would improve model strength and capture significantly more seasonal and geographical variation in *V. vulnificus* density. I present a robust predictive model of *V. vulnificus* density based on comprehensive measurements of environmental variables at high-resolution ecologically-relevant spatial and temporal scales. Leveraging an existing coastal oceanographic time series and turbidity plume forecast model (Johnson et al., 2013) maintained by the Pacific Islands Ocean Observing System (PacIOOS), I applied the statistical model to construct a real time model of *V. vulnificus* density throughout nearshore Waikīkī in order to provide more timely information on pathogen risk assessment. In addition, I report a long-term forecast projection of *V. vulnificus* densities with climate change.
1.3 MATERIALS AND METHODS

The Ala Wai Canal Study Site

The Ala Wai Canal is a prominent waterway in urban Honolulu, Hawai‘i on the island of O‘ahu. The canal was constructed during the 1920s by the U.S. Army Corps of Engineers in order to drain the Waikīkī wetlands for coastal development. The canal is channelized with cement and extends for roughly 3 km parallel to the shoreline varying from 51 to 83 m wide with one major bend towards the ocean. Currently, the canal operates as a tidally-influenced estuary with freshwater input from the Makiki stream, Mānoa-Pālolo drainage, and several small urban runoff drains. The mouth of the canal is located at the Ala Wai Boat Harbor centered between the popular Waikīkī beaches to the east and Ala Moana Beach Park to the west. In this potential outflow area, there is high recreational activity, including surfing, paddling, sailing, swimming, and fishing. Therefore, there is a concerning risk of exposure to *V. vulnificus* in this area. The Clean Water Branch (CWB) of the Hawai‘i State Department of Health is responsible for the protection and monitoring of the coastal ecosystem in the state. The CWB tests coastal waters for fecal indicator bacteria concentrations (*Enterococcus* spp. and *Clostridium perfringens*), as well as water chemistry (pH, salinity, DO, turbidity, and temperature) (http://cwb.doh.hawaii.gov/CleanWaterBranch/WaterQualityData/default.aspx). However, the CWB sampling in the Ala Wai Canal has been suspended indefinitely due to budgetary constraints and *V. vulnificus* is not routinely monitored by the CWB anywhere in Hawai‘i. Previous studies investigating rain-driven effluent plumes from the canal suggest that the canal could be a source of pathogenic bacteria, including *V. vulnificus*, and pathogen indicator bacteria, specifically *Enterococcus* sp. (Connolly et al., 1999; Johnson et al., 2013) into coastal waters. *V. vulnificus* is found and regularly cultured from the canal and has been shown to vary over a seasonal cycle with little variation attributed to water temperature (Nigro, 2012). The variation in response to low salinity has been attributed to a strong correlation with groundwater flow (reduced nitrogen and silicate) rather than surface runoff (Nigro, 2012).

Field Surveys and Sample Collection

Biogeochemical survey data were collected over 9 sampling dates between October 2018
and October 2019 (21 October 2018; 29 November 2018; 20 January 2019; 18 February 2019; 22 March 2019; 20 April 2019; 4 June 2019; 31 August 2019; 29 September 2019). Depth profiles were collected from 18 sites along the Ala Wai canal and offshore (Figure 1.1). Continuous sensor profiles were conducted at each site on each date for salinity, temperature, turbidity, chlorophyll fluorescence, and dissolved oxygen. Discrete bottle samples were collected at 8 of these sites roughly 0.5 km apart along the transect. For discrete sampling, water was pumped into 1 L polycarbonate bottles at 3 depths per site: surface (25 cm below the air-water interface), middle (targeting the pycnocline), and bottom (25 cm above the sediment-water interface). Sampling was conducted sequentially along the sites and against the tidal flow over a period of ~3 h. I targeted sampling from the back of the canal at the lowest low tide for consistency and maximal gradient in salinity. Bottle samples were processed or preserved within 8 h of collection.

**Hydrological and Climatological Data Sources**


**Water Chemistry Data Collection**

**In Situ Sensor Data Collection**

Temperature and conductivity were measured using an Aanderaa 4319A conductivity sensor (1.4 Hz). Dissolved oxygen was measured using an Aanderaa 4330F optode (~ 1 Hz). Turbidity (approximated from optical backscatter) and chlorophyll-a fluorescence were
measured using a Sea-Bird ECO FLNTU sensor (~ 6 Hz). Depth was measured using a custom-built water pressure probe (~ 1 Hz). Precise latitude and longitude were recorded using an onboard Simrad GPS.

*Nutrient Measurements*

All nutrient samples were analyzed at the University of Hawai‘i at Mānoa School of Ocean and Earth Science and Technology (SOEST) Laboratory for Analytical Biochemistry (http://www.soest.hawaii.edu/S-LAB/) using a Seal Analytical AA3 HR Nutrient Autoanalyzer. Dissolved nitrate plus nitrite, ammonium, phosphate, and silicate were analyzed according to Armstrong et al. (1967) and Grasshoff et al. (1983), Kérouel & Aminot (1997), Murphy & Riley, (1962), and Grasshoff et al. (1983), respectively, following filtration through a 0.22 μm pore size polyethersulfone filter capsule (Sterivex, Millipore) using a peristaltic pump. Total nitrogen and total phosphorus were measured from non-filtered samples following the modified Autoanalyzer procedure developed by the University of Hamburg.

*Particulate and Dissolved Organics Measurements*

Particulate organic carbon (POC) and particulate organic nitrogen (PON) samples were collected on combusted GF/F filters (Whatman) and analyzed using an Exeter Analytical model CE 440 elemental analyzer according to methods by Gordon (1969) and Sharp (1974). Filtrate was collected and acidified to pH 2 to measure dissolved organic carbon (DOC) and total dissolved nitrogen (TDN) using a Shimadzu High-Temperature TOC-L Combustion Analyzer. Samples for fluorescent dissolved organic matter (fDOM) were analyzed using a Horiba Aqualog scanning fluorometer according to Nelson et al. (2015). Relative fluorescence of known spectral peaks in excitation emission matrices was determined based on Coble (1996).

*pH Measurements*

Measurements of pH were made from whole water bottle samples in the lab using a Hach sensor probe calibrated for each sampling event.
**Microbial Data Collection**

**Chlorophyll-a Measurements**

Chlorophyll-a was measured from bottle samples using acetone extraction from material collected on 0.45 µm HAWP filters (Millipore) and fluorescence spectroscopy on a modified Turner 10AU fluorometer according to Smith et al. (1981) with a detection limit of 0.025 µg L⁻¹ using the EPA Method 445.0.

**Total Prokaryotic Abundance Measurements**

Total prokaryotic abundance was determined using an Attune Acoustic Focusing Cytometer according to Nelson et al. (2015) from whole water samples fixed with paraformaldehyde (0.5% final concentration) and stained with 1X SYBR Green I.

**DNA Extraction Protocol**

Samples for DNA extraction were collected by passing water through 0.22 µm polyethersulfone filters (Sterivex, Millipore); the exact filtrate volume was recorded (250-500 mL). Filters were added to MP Biomedicals Lysing Matrix A (No. 116910100) tubes with MC 1 lysis buffer and homogenized using a MP Biomedicals FastPrep-96 bead beater. DNA extractions were completed using the Macherey-Nagel NucleoMag Plant Extraction Kit (No. 744400.4) with KingFisher accessory kit (No. 744951). Samples were eluted to a final volume of 110 µL.

**Quantifying Vibrio vulnificus**

There is a consistent log correlation between CHROMagar *Vibrio* blue colony forming units and *V. vulnificus* hemolysin A (vvhA) gene copy concentrations (Nigro & Steward, 2015). Therefore, I used a 16S rRNA 5’ nuclease vvhA quantitative PCR assay (Campbell & Wright, 2003; Holland et al., 1991; forward primer: 5’-TGTATGTGAGAAACCGGTGACA-3’; reverse primer: 5’-TTCTGGTAGCCCCAACCTTG-3’; 5’-nuclease probe: 5’-/56-FAM/CCGTTAACC/ZEN/GAACCACCCGCAA/31ABkFQ/-3’) to determine total *V. vulnificus* abundance. A 25 µL reaction mixture was prepared with final concentrations of 1X Kapa Probe Force Master Mix (Kapa Biosystems), 0.9 µM of each primer, 0.5 µM of the labeled probe, 0.56
mg/ml bovine serum albumin (Thermofisher), and 5 μL of DNA template (11.4 – 22.7 mL sample water). All qPCR reactions were performed in triplicate with the final replicate diluted 10-fold to check for inhibition of amplification as suggested by Bustin et al. (2009). The cycling protocol consisted of an initial denaturation step at 95 °C for 10 minutes, followed by 40 cycles of denaturation at 95 °C for 15 s and a combined annealing/extension at 60 °C for 30 minutes. I created an eight-point standard curve (10 – 50,000 copies per reaction well) of genomic DNA from *V. vulnificus* (strain YJ016; Chen et al., 2003) with known gene copy numbers per reaction run in triplicate along with the environmental samples. Assay efficiency was calculated from multiple standard curves using the formula \( E = -1 + 10^{(-1/slope)} \) (Pfaffl, 2001).

**Data Analysis**

**Data Processing**

Data collected from sensors (salinity, temperature, dissolved oxygen, chlorophyll, and turbidity) were resampled to 1 Hz frequency with linear interpolation using Python 3.7.4 (Python Core Team, 2015) pandas package (McKinney, 2011) in order to coordinate the datasets based on timestamp. These data were then averaged over the time the bottle sample was collected to combine with discrete bottle measurements (nutrients, organics, pH, chlorophyll-a, total prokaryotes, *vvhA*). When necessary, data were transformed to a Gaussian distribution, typically via log10, to satisfy residual error distribution assumptions for linear regression statistical analysis. Replicate qPCR measurements of *vvhA* gene copy number were averaged and those below the assay limit of detection (10 copies per reaction) were set to the lowest detectable measurement of 0.1 copies mL\(^{-1}\) sample volume. *V. vulnificus* density values (*vvhA* gene copies mL\(^{-1}\)) were log10 transformed.

**Statistical Analysis and Model Selection**

I hypothesized that spatial and temporal variation in environmental parameters could explain significant variation in *V. vulnificus* density. In particular, I hypothesized that including additional measurements, such as nutrients and organics, in coordination with salinity and temperature could improve statistical models of *V. vulnificus* density in the Ala Wai Canal. To avoid collinearity between model predictors, I assessed pairwise correlation of all predictors to
generate a correlation matrix. A hierarchical cluster based on the correlation matrix was generated and one or two predictors were chosen from each major cluster (Figure 1.2) to be included in the full model for further analysis. To maximize the usefulness of the final model, I prioritized parameters that are widely measured, lower cost, and available on continuous monitoring platforms.

To test my hypothesis, I log$_{10}$-transformed the response variable (V. vulnificus density) and fit a linear model with an error distribution assumption of normal. The model included main effects of predictors only with no interaction terms in order to maximize reproducibility in other systems. Model assumptions were assessed by inspection of residuals for normality and uniformity. Linear regression analyses were performed using R version 3.1.6 (R Core Team, 2019). The best combination of predictors was determined by fitting all possible predictor combinations using the dredge function in the MuMIn package (Bartoń, 2019) and comparing the Akaike Information Criterion value (Akaike, 1973) corrected for small samples sizes (AICc). The importance of predictors was evaluated using the importance function from the MuMIn package to get an importance measure of each predictor. The importance value is the sum of the model Akaike weights for all models that contain the particular parameter. Leave-one-out cross-validation was used to evaluate the prediction error (PE) of different models based on the root mean squared error (RMSE) of predictions versus observations, where PE is calculated as the log$_{10}$-corrected RMSE.

**Short-term Forecasting**

Short-term forecasting, or nowcasting, of V. vulnificus density was developed using the PacIOOS nearshore sensor NS02 (http://www.pacioos.hawaii.edu/water/sensor-hawaiiyachtclub/) and the PacIOOS turbidity plume forecast (http://www.pacioos.hawaii.edu/water/model-plume-alawai/) based on the Regional Ocean Modeling System for the Waikīkī area (Waikīkī ROMS). The PacIOOS NS02 is moored at 1 m depth at (21.286407°N, -157.84276°E) and records continuous measurements of temperature and conductivity (Sea-Bird Electronics, SBE16plus V2 SEACAT), and fluorescence (470/695 nm) from which chlorophyll-a is estimated, and optical backscatter (700 nm) as an approximation of turbidity (WET Labs, ECO FLNTUS) at 0.0042 Hz frequency (McManus, 2008). An additional
sensor for continuous measurements of fDOM (WET Labs, WETStar DOM Fluorometer, 370 nm excitation/460 nm emission, 0.0014 Hz; Belzile et al., 2006) was deployed at this location from September 2018 – May 2019. These measurements were validated based on depth profile measurements in this study at site 2. *V. vulnificus* density was predicted using NS02 salinity, National Weather Service five-day average rainfall and daily maximum air temperature and compared to measured *V. vulnificus* density. Air temperature and water temperature are highly correlated, so air temperature was used since it is easier to measure remotely. A second model of *V. vulnificus* density which included fDOM measurements was compared to the previous model to assess the effect of adding DOM as a model parameter. Goodness of fit and model prediction error were evaluated by RMSE.

The PacIOOS turbidity plume forecast based on the Waikīkī ROMS predicts three-dimensional circulation as well as salinity, temperature, and turbidity based on the PacIOOS NS02 measurements for the canal and adjacent coastal region. The model is spatially resolved within the canal and nearshore environment to 40 m with 14 depth layers (0.25, 1, 2, 5, 10, 20, 30, 50, 75, 100, 125, 150, 200, 250 m) at 1 h time intervals. *V. vulnificus* density was predicted for the surface 0.25 m depth using the forecasted salinity and turbidity, and NWS five-day average rainfall, and daily maximum air temperature.

**Long-term Forecasting**

End-of-century projections of air temperature and precipitation were used to estimate overall change in *V. vulnificus* density due to climate change. The dynamical downscaling projections that were used (Zhang et al., 2012, 2016a, 2016b) are based on the IPCC AR5 CMIP5 global model for representative concentration pathways (RCP) 4.5 and 8.5 (IPCC, 2014). The forecasting assumes a linear response of *V. vulnificus* with increasing temperature and no significant changes to other aspects of environmental variation.
1.4 RESULTS

Spatial and Temporal Variation of *V. vulnificus*

*V. vulnificus* density (*vvhA* gene copies mL\(^{-1}\)) ranged from 0.1 – 266 with a geometric mean of 4.0 for the warm, rainy season (October – March) and ranged from 0.1 – 529 with a geometric mean of 17.6 for the hot, dry season (April – September). There were significant differences between month (F = 18.24, p < 0.001), site (F = 8.88, p < 0.001), and depth (F = 26.80, p < 0.001) in *V. vulnificus* density (Figure 1.3). *V. vulnificus* density was significantly elevated in the surface water, within the canal (sites 1 – 6), and during the summer months.

Correlation with Biogeochemical Predictors

*V. vulnificus* density was most strongly correlated with dissolved inorganic nutrient concentrations and metrics of fluorescent DOM characteristics (Figure 1.2; r > 0.6). The strongest additional univariate correlations (r > 0.4) were rainfall, total prokaryotes, turbidity (Figure 1.4), and bulk measurements of dissolved and particulate organic carbon and nitrogen. *V. vulnificus* concentrations were strongly negatively correlated (r < - 0.4) with pH and salinity. I found no significant correlation between *V. vulnificus* (p < 0.05) and stream discharge and *V. vulnificus* and chlorophyll-a measured *in situ*.

Predictive Model Selection

To construct predictive models, highly correlating biogeochemical parameters were removed to mitigate overfitting, unstable coefficient estimates, and high prediction error. I prioritized parameters that are widely measured, lower cost and available on continuous monitoring platforms. Out of 36 biogeochemical measurements, 12 were chosen for model selection analysis, with 1-2 parameters each representing one of 9 clusters of covarying parameters (Figure 1.2). Five-day average rainfall, daily maximum air temperature, salinity, turbidity, dissolved oxygen, particulate organic carbon, silicate, phosphate, visible humic-like fDOM, the fDOM humification index (HIX), and total prokaryote density had reasonably low pairwise collinearity and were used as terms to construct a linear model of *V. vulnificus* density in log\(_{10}(vvhA\) gene copies mL\(^{-1}\)) (Table 1.1). Model residuals were well-behaved (uniformly
distributed relative to predicted values) and normally distributed, following model assumptions. For the full model with all terms, the residual standard error was 0.54 on 179 degrees of freedom with an $R^2$ value of 0.70.

Comparing models representing all combinations of predictors, 24 models were within $\Delta$AICc < 4 of the best-specified model and rainfall, air temperature, salinity, silicate, visible humic-like fDOM, and HIX were included in all 24 of those low AICc models. The best-specified model based on AICc included rainfall, air temperature, salinity, dissolved oxygen, silicate, visible humic-like fDOM, and HIX had an $R^2$ of 0.70 (Model 6 in Table 1.2). The second lowest AICc model ($\Delta$AICc = 1.03, $R^2 = 0.69$) does not include dissolved oxygen. The importance value for each predictor, calculated using the MuMIn package, is reported in Table 1.1.

**Operational Model Comparison and Cross Validation**

Based on the important predictors from the best-specified model (see Predictive Model Selection) and model applications (see Model Forecasting Applications), six linear regression models were compared to assess predictive power of *V. vulnificus* density (Table 1.2, Figure 1.5). The models were compared using AICc and RMSE as well as the average RMSE from LOOCV to assess model fit and prediction error. $\Delta$AICc is relative to model 6, the best-specified model. As RMSE is the average error between the predicted and observed value, the “prediction error” is the log-corrected RMSE. Model 1 included rainfall and air temperature only as predictors of *V. vulnificus* density and performed the worst ($R^2 = 0.325$, $\Delta$AICc = 140, RMSE = 0.79, LOOCV RMSE = 0.80, Prediction Error = 6.3). Model 2 included water chemistry parameters: salinity and water temperature. Model 2 performed better than model 1 and explains more variation in *V. vulnificus* density ($R^2 = 0.46$, $\Delta$AICc = 96, RMSE = 0.70, LOOCV RMSE = 0.71, Prediction Error = 5.1). Model 3 combined climate and water chemistry parameters: rainfall, air temperature, and salinity. Water temperature was removed to avoid collinearity with air temperature and air temperature explained slightly more variation in *V. vulnificus* density as well as being easier to measure remotely. Model 3 performed better than model 2 and explains more variation in *V. vulnificus* density ($R^2 = 0.53$, $\Delta$AICc = 71, RMSE = 0.67, LOOCV RMSE = 0.67, Prediction Error = 4.7). Model 4 builds from model 3 to include turbidity which is modeled
in the Waikīkī ROMS forecast. Again, model 4 performed better than model 3 and explains more variation in *V. vulnificus* density ($R^2 = 0.58$, $\Delta$AICc = 55, RMSE = 0.63, LOOCV RMSE = 0.64, Prediction Error = 4.4). Model 5 replaces turbidity with visible humic-like fDOM which is measured at the NS02 time series. Model 5 performed better than model 4 and explains more variation in *V. vulnificus* density ($R^2 = 0.62$, $\Delta$AICc = 34, RMSE = 0.59, LOOCV RMSE = 0.61, Prediction Error = 4.1). Model 6, is the best-specified model from model selection analysis and performed as such ($R^2 = 0.69$, $\Delta$AICc = 0, RMSE = 0.53, LOOCV RMSE = 0.55, Prediction Error = 3.6).

**Model Forecasting Applications**

**Short-term Forecasting**

The PacIOOS time series data from NS02 were validated against measured observations in this survey from site 2 (Figure 1.6). Overall, the NS02 data were roughly equivalent to the survey data with high $R^2$ values and a linear fit with intercepts near zero and slopes near one. Salinity and turbidity are exceptions with salinity $R^2$ of 0.41, attributable to fine-scale variation in sampling location in a stratified estuarine habitat, and turbidity having an intercept of 2.1 potentially caused by an instrument offset. Using the NS02 data, *V. vulnificus* density was hindcast for the sampling period (Figure 1.7) and validated against measured observations (Figure 1.8). Two model frameworks were compared to assess the improvement of adding fDOM measurements (model 3 vs. model 5, see *Operational Model Comparison*). Model 3 (“–fDOM”) uses salinity, five-day average rainfall, and daily maximum air temperature to predict *V. vulnificus* density ($R^2 = 0.53$) with RMSE of 0.33 and $R^2$ of 0.53 compared to measured values. Model 5 (“+ fDOM”) uses salinity, five-day average rainfall, daily maximum air temperature, and visible humic-like fDOM to predict *V. vulnificus* density ($R^2 = 0.62$) with RMSE of 0.12 and $R^2$ of 0.98 compared to measured values. The addition of fDOM measurements in Model 5 improves predictions of *V. vulnificus* density (Figure 1.8).

A three-day spatial forecast of *V. vulnificus* density was constructed using the PacIOOS turbidity plume forecast and Waikīkī ROMS model framework which includes 4 m resolution projections of salinity, temperature, and turbidity in addition to water velocity at 1 h intervals. Model 4 (see *Operational Model Comparison*) was used to predict *V. vulnificus* density based on
the forecast output. For an example rain event in February 2019, this model demonstrated *V. vulnificus* plume dynamics exiting the Ala Wai Canal (Figure 1.9).

**Long-term Forecasting**

Long-term climate predictions were based on Model 1, rainfall and air temperature, using projections of monthly average rainfall and air temperature. Model 1 was first validated against survey measurements of *V. vulnificus* density (Figure 1.10). The climate model does not accurately predict the spatial variation in *V. vulnificus* density (\( R^2 = 0.42, \text{RMSE} = 0.65, \text{PE} = 4.47 \)), but does well to predict the overall average *V. vulnificus* density within the canal (\( R^2 = 0.90, \text{RMSE} = 0.04, \text{PE} = 1.1 \)). Model coefficients were used to predict the long-term effects of changing rainfall and air temperature on overall average *V. vulnificus* density for the canal. Three climate projections were considered based on dynamical downscaling (250 m resolution) of recorded data (1990-2009) and the AR5 RCP 4.5 and RCP 8.5 modeled scenarios (2080-2099). RCP 4.5 is a 580-720 ppm CO2-eq by 2100 and 1.7-3.2 °C increase relative to 1850-1900 and RCP 8.5 is a >1000 ppm CO2-eq by 2100 and 3.2-5.4 °C increase relative to 1850-1900. There was no significant difference (\( p = 0.45 \)) in annual average rainfall between the three projections (1990-2009, RCP 4.5, and RCP 8.5) (Figure 1.11). There was a significant difference (\( p < 0.0001 \)) in annual average air temperature between the three projections. Mean comparisons using Tukey-Kramer HSD show that RCP 8.5 (mean = 27.6 °C) is significantly higher (\( p < 0.0001 \)) than RCP 4.5 (mean = 26.0 °C) and RCP 4.5 is significantly higher (\( p < 0.0001 \)) than 1990-2009 (mean = 24.2 °C). There is also a significant difference (\( p < 0.0001 \)) in predicted annual average *V. vulnificus* density between the three projections (Figures 1.11 and 1.12). Mean comparisons using Tukey-Kramer HSD show that RCP 8.5 (mean = 61.0 vvhA gene copies mL\(^{-1} \)) is significantly higher (\( p < 0.0001 \)) than RCP 4.5 (mean = 36.5 vvhA gene copies mL\(^{-1} \)) and RCP 4.5 is significantly higher (\( p < 0.0001 \)) than 1990-2009 (mean = 19.3 vvhA gene copies mL\(^{-1} \)). The RCP 4.5 and 8.5 values in *V. vulnificus* density correspond to a 21.5 % and 38.9 % increase respectively relative to 1990-2009 (Figure 1.12).
1.5 DISCUSSION

**General Patterns in Vibrio vulnificus Seasonality**

Hawai‘i has consistently higher *Vibrio* infection rates per capita than any other state in the U.S. This study evaluates statistical models of *V. vulnificus* concentrations in the Ala Wai Canal, a tropical estuarine environment in a densely populated area of Honolulu, Hawai‘i. The results demonstrate how accounting for spatiotemporal dynamics of dissolved nutrient resources improves the predictive capabilities of these statistical models and model applications. Despite the high degree of both temporal and spatial variation, climate and biogeochemistry can explain up to 70 % of the variation in *V. vulnificus* density using the best-specified model (Model 6 in Table 1.2). There is a significant increase in *V. vulnificus* density in the summer months which corresponds with higher water temperatures and an enhanced chlorophyll density likely generating the observed increase in dissolved oxygen and organic matter (Figure 1.4). These results contrast previous findings in the canal from a decade prior (Nigro, 2012) which show overall higher abundances during the rainy season and may reflect changes to the environment such as increasing water temperature in the summer months over 10 years or differences in the rain events captured between the studies. Rain events, which transport nutrients and dissolved organics as well as low salinity water into the canal through the streams and runoff are also positively correlated with *V. vulnificus* density in this study (see the February 2019 rain event in Figures 1.3 and 1.9). These findings are consistent with a previous survey that show elevated concentrations of *V. vulnificus* in the harbor region after a large rain event (Nigro, 2012). These results suggest that conditions for *V. vulnificus* growth are improved in this outflow region during rain events in addition to water movement transporting *V. vulnificus* out of the canal. Since the harbor and outflow regions are sites of high and concentrated recreational activity, summer rain storms may be particularly concerning for human exposure to *V. vulnificus*. Interestingly, Fiedler et al. (2014) also reported a significant increase in *V. vulnificus* density near the PacIOOS NS02 following the Japan tsunami. The correlation and importance of SiO$_2$ (De Carlo et al., 2007) and HIX (Nelson et al., 2015) in the modeling exercises suggests a positive response of *V. vulnificus* to groundwater flow which is in keeping with the findings of Nigro (2012).
The pattern and order-of-magnitude seasonal variation in *V. vulnificus* density recorded in this study is comparable to previous studies in other areas (Jacobs et al., 2014; Pfeffer et al., 2003; Tamplin et al., 1982), but significantly lower than previous reports in the Ala Wai Canal where canal-wide average concentrations ranged from 23.1 $vvhA$ gene copies mL$^{-1}$ to 3640 $vvhA$ gene copies mL$^{-1}$ (Nigro, 2012). This variation is still notable since the water temperature remains well above the 18 °C minimum for optimal growth (ranging between 22 – 31 °C). For this environment, salinity and temperature alone explain only 20 – 30 % of the variation in *V. vulnificus* density.

**Improvements with Dissolved Nutrients and Organics**

The quality of nutrient resources available may be an important and overlooked determinant in predicting the population dynamics of *V. vulnificus*. My results indicate that the addition of fluorescent dissolved organic matter (fDOM) measurements to resolve humic components (often targeting a range of wavelengths referred to as “cDOM” or visible humic-like components; Nelson & Siegel, 2013) could resolve 10 % more variation in *V. vulnificus* density than salinity and climate parameters alone (Model 3 vs. 5 in Table 1.2). In application, the addition of a real time deployed fDOM sensor at NS02 reduced prediction error by 36 % (Figure 1.8). Although dissolved inorganic nutrients, particularly silicate, are also important in the models and strongly correlated with *V. vulnificus* concentrations, nutrient analysis currently requires water sample collection and laboratory analysis. Fluorescent DOM can be measured with a deployed real time sensor and fDOM parameters were almost equally correlated with *V. vulnificus* as nutrients were with *V. vulnificus* (Figure 1.2). However, advances are being made in real-time sensors for inorganic nutrients such as nitrate (Sea-Bird Scientific, SUNA V2 Nitrate Sensor based on Johnson & Coletti, 2002) and orthophosphate (Sea-Bird Scientific, HydroCycle-Phosphate Analyzer). These real-time sensors may become valuable tools for future coastal pathogen monitoring programs.

There are only a few studies on natural populations of *V. vulnificus* in tropical estuaries. In the case of a tropical environment where temperature is always facilitating rather than limiting growth, other factors, such as macronutrient resource abundance and quality, may be more important controls on the population density of *V. vulnificus*. Vibrios are copiotrophic
bacterioplankton with broad capacity to degrade organic compounds and extract macronutrients from DOM (Baker-Austin et al., 2018). As such, the character and quantity of DOM is a reasonable parameter modulating growth dynamics of this pathogen and other naturally-occurring estuarine organisms. Considering nutrient and DOM resources in predictive modeling seems to be a promising route forward. Measurements of fluorescent dissolved organic matter, which can be recorded easily with deployed sensors (Belzile et al., 2006; Carstea et al., 2020; Ruhala & Zarnetske, 2017), may be the key to generating robust predictive models of *V. vulnificus* population dynamics. There are even promising efforts at forecast modeling DOM in estuarine environments (Bowers & Brett, 2008) which could be integrated into coastal pathogen modeling.

**Conclusions and Climate Change Implications**

This study parameterizes *V. vulnificus* absolute abundance with comprehensive physical, optical, and chemical water properties in high spatial and temporal resolution to develop a real-time spatially-explicit model of *V. vulnificus* pathogen risk. It is critical that we improve our predictive capabilities of the *V. vulnificus* human pathogen in the face of increased water temperature, storm surge, sea level, hurricanes, and flooding due to climate change which may be favorable to *V. vulnificus*. The habitat range of *V. vulnificus* may expand due the sea level rise and coastal flooding given its preference for mixed salinity water. In addition, *V. vulnificus* population growth will likely increase due to increasing water temperature. There is evidence for significantly increased season length and spatial habitat from work in the Chesapeake Bay (Muhling et al., 2017). Based on local dynamical downscaling of the CMIP5 global model projections for rainfall and air temperature (Zhang et al., 2012, 2016a, 2016b), my models suggest a 21.5 – 38.9 % increase in average *V. vulnificus* density by 2100 if atmospheric CO2 concentrations reach between RCP 4.5 and 8.5. If climate change proves more favorable for *V. vulnificus* and expands areas of coastal flooding, we will likely see increased exposure of the general public to *V. vulnificus* and increased infections. Therefore, improving predictive models of microbial pathogen populations is critical to public safety and incorporating real time measurements of nutrient resources may be a crucial step forward.
1.6 TABLES

**Table 1.1** Model Selection: Full-model likelihood ratio test based on linear least squares regression analysis of *V. vulnificus* density in log_{10}(vvhA gene copies mL^{-1}) and AIC-weighted importance of predictors based on model comparison of all possible predictor combinations.

<table>
<thead>
<tr>
<th>Model Term</th>
<th>Coefficient Estimate</th>
<th>Standard Error</th>
<th>t-value</th>
<th>p-value</th>
<th>AIC-weighted Importance</th>
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<tbody>
<tr>
<td>(Intercept)</td>
<td>5.78E-02</td>
<td>1.17E+00</td>
<td>0.05</td>
<td>9.61E-01</td>
<td>NA</td>
</tr>
<tr>
<td>1. Five-day Average Rainfall</td>
<td>3.11E-01</td>
<td>5.15E-02</td>
<td>6.03</td>
<td>9.00E-09</td>
<td>*** 1.00</td>
</tr>
<tr>
<td>2. DOM Humification Index</td>
<td>-2.25E-01</td>
<td>5.12E-02</td>
<td>-4.39</td>
<td>1.93E-05</td>
<td>*** 1.00</td>
</tr>
<tr>
<td>3. Air Temperature</td>
<td>8.02E-02</td>
<td>2.05E-02</td>
<td>3.91</td>
<td>1.31E-04</td>
<td>*** 1.00</td>
</tr>
<tr>
<td>4. Visible Humic-like DOM</td>
<td>1.13E+00</td>
<td>3.90E-01</td>
<td>2.89</td>
<td>4.31E-03</td>
<td>** 1.00</td>
</tr>
<tr>
<td>5. Silicate</td>
<td>5.11E-01</td>
<td>2.00E-01</td>
<td>2.56</td>
<td>1.13E-02</td>
<td>* 1.00</td>
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<tr>
<td>6. Salinity</td>
<td>-1.48E-05</td>
<td>6.34E-06</td>
<td>-2.34</td>
<td>2.04E-02</td>
<td>* 1.00</td>
</tr>
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<td>7. Dissolved Oxygen</td>
<td>-2.52E-02</td>
<td>1.66E-02</td>
<td>-1.51</td>
<td>1.31E-01</td>
<td>0.61</td>
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<tr>
<td>8. Particulate Organic Carbon</td>
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<td>1.67E-01</td>
<td>-0.68</td>
<td>5.00E-01</td>
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<tr>
<td>9. Phosphate</td>
<td>-1.87E-01</td>
<td>2.06E-01</td>
<td>-0.91</td>
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<td>10. Total Prokaryotes</td>
<td>-7.61E-02</td>
<td>1.92E-01</td>
<td>-0.40</td>
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<td>11. Nitrate + Nitrite</td>
<td>3.52E-02</td>
<td>1.51E-01</td>
<td>0.23</td>
<td>8.16E-01</td>
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<td>12. Turbidity</td>
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<td>1.36E-01</td>
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<td>9.94E-01</td>
<td>0.14</td>
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Significance codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’
**Table 1.2** Operational Model Comparison: Linear least squares regression analysis of *V. vulnificus* density in log$_{10}$(vhA gene copies mL$^{-1}$) comparing six models using AIC model selection analysis and root-mean-square-error (RMSE) for both the full dataset and leave-one-out-cross-validation (LOOCV).

<table>
<thead>
<tr>
<th>Model</th>
<th>&quot;Climate-only&quot;</th>
<th>R$^2$ = 0.325</th>
<th>AICc = 464</th>
<th>RMSE = 0.79</th>
<th>LOOCV</th>
<th>RMSE = 0.80</th>
<th>PE = 6.31</th>
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<tr>
<td></td>
<td>Estimate</td>
<td>SE</td>
<td>t value</td>
<td>p-value</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(Intercept)</td>
<td>-4.00E+00</td>
<td>7.42E-01</td>
<td>-5.35</td>
<td>2.50E-07</td>
<td>***</td>
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<tr>
<td>Five-day Average Rainfall</td>
<td>4.41E-01</td>
<td>5.85E-02</td>
<td>7.55</td>
<td>1.80E-12</td>
<td>***</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Air Temperature</td>
<td>1.42E-01</td>
<td>2.45E-02</td>
<td>5.85</td>
<td>2.10E-08</td>
<td>***</td>
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</table>

<table>
<thead>
<tr>
<th>Model</th>
<th>&quot;Water Chemistry&quot;</th>
<th>R$^2$ = 0.462</th>
<th>AICc = 420</th>
<th>RMSE = 0.70</th>
<th>LOOCV</th>
<th>RMSE = 0.71</th>
<th>PE = 5.13</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Estimate</td>
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AICc is the Akaike Information Criterion value corrected for small sample sizes. RMSE is the average error between the predicted and observed value. LOOCV is the leave-one-out-cross-validation technique to test for prediction error. PE, or prediction error, is the log-corrected RMSE. SE is the standard error value from the likelihood ratio test. Significance codes: 0 '***', 0.001 '**', 0.01 '*', 0.05 '.}'
Figure 1.1 Map of the Ala Wai Canal in Honolulu, Hawai’i depicting monthly survey sampling locations from October 2018 - September 2019. Full depth profiles for salinity, temperature, dissolved oxygen, turbidity, and chlorophyll were collected from all sites (n = 18) marked with a pin. Yellow numbered sites (1-8) are locations where discrete bottle samples were collected at three depths (surface, pycnocline, and bottom water) for additional nutrient, organic matter, and microbial measurements. The PacIOOS nearshore sensor NS02 at the Ala Wai Harbor is marked with a grey box.
**Figure 1.2** Heat map of the correlation matrix showing pairwise correlation of all variables. The dendrogram (right) depicts the hierarchical cluster structure based on the correlation matrix values. Light grey rectangles overlaying the dendrogram mark the nine major clusters of highly collinear variables.
Figure 1.3 Spatial and temporal variation of *V. vulnificus* density in the Ala Wai Canal in Honolulu, Hawai‘i from October 2018 - September 2019 divided by site (horizontal axis) and sample depth (panels). *V. vulnificus* density varies significantly between months, sites, and depths (p < 0.001). *V. vulnificus* density significantly increases from offshore (site 1) to the back end of the canal (site 8) and in the surface waters (0.25 m from the surface) relative to the pycnocline or bottom waters (0.25 m from the bottom). Seasonally, canal-averaged *V. vulnificus* density rises through the year beginning in October 2018 and peaks in September 2019 with anomalously elevated levels during the February 2019 rain event.
Figure 1.4 Continuous depth profiles of biogeochemistry in the Ala Wai Canal from October 2018 – September 2019. Data are separated by the rainy season (October through March; top set of panels) and the dry season (April through September; bottom set of panels).
Figure 1.5 Operational Model Comparison. Model predictions, based on leave-one-out-cross-validation (LOOCV), are compared to observations of *V. vulnificus* density in $\log_{10}(vvhA \text{ gene copies mL}^{-1})$ (see Table 1.2). The 1:1 line is in red. RMSE is the root mean squared error.
Figure 1.6 Validation of NS02 measurements with survey data from site 2 surface values. Linear fit equations are shown.
Figure 1.7 Time series of data from PacIOOS NS02, USGS, and NOAA National Weather Service. *V. vulnificus* density is hindcast using a linear model of five-day average rainfall, air temperature, salinity, and visible humic-like DOM (when data were available). Grey vertical dashed lines mark dates when field surveys were conducted.
Figure 1.8 Predicted *V. vulnificus* density based on two models (see Model 3 and Model 5, Table 1.2) of NS02 data compared to measured *V. vulnificus* density from the survey site 2 surface values. Model 3 (– fDOM) uses salinity, five-day average rainfall, and daily maximum air temperature to predict *V. vulnificus* density ($R^2 = 0.53$) with RMSE of 0.33 and $R^2$ of 0.53 compared to measured values. Model 5 (+ fDOM) uses salinity, five-day average rainfall, daily maximum air temperature, and visible humic-like fDOM to predict *V. vulnificus* density ($R^2 = 0.62$) with RMSE of 0.12 and $R^2$ of 0.98 compared to measured values.
Figure 1.9 Applied model predictions (Table 1.2, Model 4) using the PacIOOS turbidity plume forecast showing plume dynamics of *V. vulnificus* density into recreational areas during a rain event (16-18 of February 2019).
Figure 1.10 Climate-modeled *V. vulnificus* density based on rainfall and air temperature versus measured *V. vulnificus* density within the canal (sites 2-8). Measured *V. vulnificus* density is shown as the geometric mean with standard error around the mean value. The linear fit ($y = 1x - 3.55e-15$) is practically 1:1 and the 95% confidence interval is shown in purple. The climate model does not predict the spatial variation in *V. vulnificus* density ($R^2 = 0.42, RMSE = 0.65, *PE = 4.47$), but expertly predicts the overall monthly average of *V. vulnificus* density within the canal ($R^2 = 0.90, RMSE = 0.04, *PE = 1.1$). Model coefficients were used to predict the long-term effects of changing rainfall and air temperature on overall average *V. vulnificus* density for the canal (see Figure 1.10). Model: $\log_{10}(vvhA) = 0.4414 \log_{10}(\text{five-day average rainfall}) + 0.1405 \log_{10}(\text{air temperature}) - 3.7750$
Figure 1.1 Monthly average climate projections of rainfall and air temperature based on dynamical downscaling (250 m resolution) of recorded data (1990-2009) and the AR5 RCP 4.5 and RCP 8.5 modeled scenarios* (2080-2099). Monthly average *V. vulnificus* density for the canal is predicted from rainfall and air temperature. Geometric mean values are shown with standard error around the mean value. There is no significant difference (p = 0.45) in annual average rainfall between the three projections (1990-2009, RCP 4.5, and RCP 8.5). There is a significant difference (p < 0.0001) in annual average air temperature and in *V. vulnificus* density between the three projections. Mean comparisons using Tukey-Kramer HSD show that RCP 8.5 (mean = 27.6 °C, 61.0 *vvhA* gene copies mL\(^{-1}\)) is significantly higher (p < 0.0001) than RCP 4.5 (mean = 26.0 °C, 36.5 *vvhA* gene copies mL\(^{-1}\)) and RCP 4.5 is significantly higher (p < 0.0001) than 1990-2009 (mean = 24.2 °C, 19.3 *vvhA* gene copies mL\(^{-1}\)).

*The Intergovernmental Panel on Climate Change (IPCC) 5th Assessment Report 2014. Representative Concentration Pathways. RCP 4.5: 580-720 ppm CO\(_2\)-eq by 2100 and 1.7-3.2 °C increase relative to 1850-1900. RCP 8.5: >1000 ppm CO\(_2\)-eq by 2100 and 3.2-5.4 °C increase relative to 1850-1900.*
Figure 1.12 Predicted monthly average *V. vulnificus* density from climate projections of rainfall and air temperature based on dynamical downscaling (250 m resolution) of recorded data (1990-2009) and the AR5 RCP 4.5 and RCP 8.5 modeled scenarios (2080-2099). There is a significant difference (p < 0.0001) in predicted annual average *V. vulnificus* density between the three projections. Mean comparisons using Tukey-Kramer HSD show that RCP 8.5 (mean = 61.0 *vvhA* gene copies mL\(^{-1}\)) is significantly higher (p < 0.0001) than RCP 4.5 (mean = 36.5 *vvhA* gene copies mL\(^{-1}\)) and RCP 4.5 is significantly higher (p < 0.0001) than 1990-2009 (mean = 19.3 *vvhA* gene copies mL\(^{-1}\)). RCP 4.5 is a 21.5 % increase in *V. vulnificus* density relative to 1990-2009. RCP 8.5 is a 38.9 % increase in *V. vulnificus* density relative to 1990-2009.
1.8 ACKNOWLEDGEMENTS

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CHAPTER 2

A comparison of experimental techniques for capturing the elusive mortality rate dynamics of *Vibrio vulnificus* in the Ala Wai Canal

Keywords: *Vibrio vulnificus*, Pathogen, Estuary, Hawai‘i, Dilution Experiment, Eukaryotic Inhibition, Grazer, Viral Lysis, Mortality Rate

2.1 ABSTRACT

Mortality is an essential aspect of population control with different dynamics than those governing growth. In order to construct mechanistic models of populations over time, a mortality term is required and is often poorly constrained. Direct measurements of mortality, especially for microbial communities, is reasonably difficult. Several techniques have been developed to assess mortality rates of natural microbial populations. With varying degrees of success, these techniques have been applied to phytoplankton and more recently heterotrophic bacteria including the pathogenic *Vibrio cholerae*. Identifying and incorporating constraints on mortality rates of *Vibrio vulnificus* would likely benefit predictive modeling capabilities. I tested two techniques on *V. vulnificus* from the Ala Wai Canal in Honolulu, Hawai‘i. The dilution method systematically reduces grazing pressure in order to measure the impact of predation on mortality rate. I modified the dilution method to assess both grazing and viral lysis. In addition, I tested a eukaryotic inhibitor to compare and validate the grazing dilution method. In most cases, there was not a significant difference between the control and test treatments and there was large variation in responses between replicates. However, overall, the more diluted treatments grew faster suggesting a positive response to the removal of predators and viruses. The gross growth rates of the dilution experiments were ultimately closer than the inhibition experiment to the gross growth rates expected based on the multifactorial growth experiment (≈0.6 h⁻¹ based on a salinity of 11.19 and 24.57 °C). Mortality rates were comparable to gross growth rates (≈0.2 –
0.7 h⁻¹) suggesting that natural population growth is limited by grazing and viral lysis. Natural populations of *V. vulnificus* may remain constant even with abundant resources due to tightly coupled growth with grazing and viral lysis.
2.2 INTRODUCTION

*Numerical Population Modeling of Vibrio vulnificus in the Ala Wai Canal*

In addition to statistical correlation modeling, efforts have been made to construct a mechanistic numerical population model for *Vibrio vulnificus* in the Ala Wai Canal (Nuss, 2016). The importance of salinity and temperature on *V. vulnificus* growth have been well documented (Motes et al., 1998; Jones and Oliver, 2009; Huehn et al., 2014; Kaspar and Tamplin 1993; Randa et al., 2004). Using a *V. vulnificus* isolate from the Ala Wai Canal, growth rates have been constrained over salinity (0 – 36) and temperature (24 – 36 °C) using multifactorial experiments (Figure 2.1; Steward and Nigro, unpublished). Based on these laboratory experiments of *V. vulnificus* growth, Nuss (2016) constructed a coupled physical-microbial numerical model of *V. vulnificus* population dynamics in the Ala Wai Canal. Their microbial model (Nuss, 2016; equation 2.1) incorporates growth based on the constraints of salinity and temperature. However, the mortality term is assumed to be constant and linearly scaled by the ratio of current *V. vulnificus* abundance to lagged abundance to account for the increase in mortality as the population grows, as well as depth to account for the increase in mortality in marine waters. Model results of *V. vulnificus* abundance are significantly different than observations from November 2008 – March 2009 for the three regions evaluated (the back end of the canal, the elbow where the canal bends towards the ocean, and the Ala Wai Harbor). Therefore, dynamic mortality rates may be more appropriate and improve model results.

*Techniques to Assess Mortality Rate*

I tested two methods of capturing *in situ* mortality rates with whole water (unamended) manipulations and incubations. The first method that is a eukaryotic inhibition with cytochalasin B. The second method is the dilution experiment which was developed by Landry and Hassett (1982). Both methods operate by systematically removing predators and/or viruses, releasing predation pressure and thereby promoting growth of the organism of interest. The change in growth rate is interpreted as the mortality rate. The dilution experiment technique is to serially dilute whole water with filtered water to linearly remove predation pressure. The resulting linear increase in growth rate with dilution is interpreted as the mortality rate (Figure 2.2). It was
originally developed for eukaryotic phytoplankton and grazers (Landry and Hassett, 1982) and later modified for prokaryotic phytoplankton (Worden and Binder, 2003) and heterotrophic prokaryotes (Jacquet et al., 2005). Landry et al. (1995) modified the experimental design to include a nutrient enrichment at the start of the experiment in order to remove any potential for nutrient limitation derived from the dilution itself and the removal of predators as a source of nutrient recycling. The gross growth rate, interpreted as the intercept of the dilution-growth curve which corresponds to all predation pressure removed, is then corrected by the difference in growth rate between the unenriched and enriched whole water. Furthermore, Li et al. (2017) attempted to account for nonlinear responses in the dilution-growth curve using a Holling II feeding response which addresses grazing saturation. Grazing saturation would mean that the grazing capacity is exceeded in the whole water, so the initial dilutions (75 % whole water for instance) have no effect on the grazing rate and therefore no effect on the growth rate of the organism of interest. This creates a depression in the dilution-growth curve which can be modeled to extract a mortality rate (Li et al., 2017). A final modification was to include the “predation pressure” of viral lysis (Jacquet et al. 2005; Kimmance et al., 2007; Mojica et al., 2016; Tijdens et al., 2008). In this design, the diluent is ultrafiltered either at 30 kDa or 100 kDa to remove both grazers and viruses. The results are then compared to a parallel experiment with traditional 0.45 μm or 0.2 μm filtered diluent which removes only grazers in order to approximate the effect of viral lysis on mortality rate. Enrichment is necessary with the viral lysis dilution experiment because the 30 kDa or 100 kDa filtration removes nutrients and depresses phytoplankton growth through nutrient limitation (Kimmance et al., 2007).

Although the dilution experiment design has been used successfully for total heterotrophic prokaryotes (Jacquet et al., 2005; Tijdens et al., 2008), there are only a handful of studies which assess the mortality rate of specific bacteria species using the dilution experiment design. For one, Worden et al. (2006) evaluated the trophic regulation of *V. cholerae*. In this case, *V. cholerae* cells from particular strains were inoculated into whole water rather than tracking a natural population. The efficacy of using these techniques to track a single population in a natural community without inoculation has yet to be shown.
Project Objectives

In order to begin to constrain the mortality rates to improve the coupled physical circulation and microbial population model, I tested these methods on natural microbial communities with *V. vulnificus* from the Ala Wai Canal. *V. vulnificus* predators are thought to be mainly ciliates and nanoflagellates and infrequently other bacteria (Chen et al., 2011). Additional mortality of *V. vulnificus* is likely caused by bacteriophages. *V. vulnificus* bacteriophage VvAW1 was isolated from the Ala Wai Canal and described in Nigro et al. (2012). However, the absolute and relative contribution of predation and viral lysis to *V. vulnificus* mortality is unknown. The dynamics of mortality are also unknown. There are likely seasonal and biogeochemical controls on mortality which could be employed to improve the coupled physical-microbial model and assist predictions for early warnings of pathogen exposure risk in coastal waters. Here, I report the results of three experiments run simultaneously and suggestions for improvement in the future.
2.3 MATERIALS AND METHODS

Water Collection

Whole water was hand collected in three 20-L, acid-cleaned polypropylene carboys on February 18, 2019 at 11:50 from the back end of the canal (21.275267° N, 157.8179596°W) at 0.25 m from the surface. *V. vulnificus* is consistently found at this location (see Chapter 1, Figure 1.3). The salinity was 11.19 and the water temperature was 24.57 °C.

Inhibition Experiment Design

Subsamples of whole water (400 mL) were transferred to 1-L, acid-cleaned flasks in triplicate. At 17:49 (six hours after water collection), 40 µL of cytochalasin B was added to each flask. The experiment was run in the lab at 20.5 °C in ambient lab lighting (overhead fluorescent lights). Samples were collected to analyze total prokaryote abundance and *V. vulnificus* abundance every hour for three hours beginning at 18:40 (T0) and ending at 21:41 (T3). Samples were collected using a peristaltic pump with acid-cleaned tubing kept inside the flasks for the entire experiment duration. Before sample collection, the flasks were mixed by hand and 20 mL was pumped from the flasks through the tubing with no filter attached to flush the lines. The cytochalasin B inhibition treatment was compared to the dilution experiment control samples (whole water with no organic enrichment; see Grazing Mortality Dilution Experiment) to calculate mortality rates.

Grazing Mortality Dilution Experiment

Five dilution treatments were applied to whole water incubations: 100 % whole water without tryptic soy broth (TSB) enrichment, 100 % whole water with TSB enrichment, 70 % whole water and 30 % diluent, 40 % whole water and 60 % diluent, and 20 % whole water and 80 % diluent. For the grazing-only experiment, the diluent was generated by filtering whole water into acid-cleaned, 20-L carboys using an in-line filter capsule (dual-layer 0.8 µm/ 0.2 µm polyethersulfone membrane; Whatman) and a peristaltic pump. Filtering began at 15:17 (roughly three hours after water collection) and intake was rotated among the three collection carboys every five minutes to effectively mix the source water. Filtered and whole water were combined
in correct proportions in 10-L polycarbonate carboys creating 7 L of the two whole water and three diluted treatments. At 17:50, 1.75 mL of TSB was added and mixed thoroughly into the 7 L treatment carboys (~100 µM labile carbon), except the whole water without TSB treatment. Two liters from each treatment and control carboy was then transferred into triplicate 2-L polycarbonate bottles and then randomly arranged on a cart to begin the experiment.

In parallel with the inhibition experiment, the grazing dilution experiment was run in the lab at 20.5 °C in ambient lab lighting. Samples were collected to quantify concentrations of total prokaryotes and *V. vulnificus* every hour for three hours beginning at 18:40 (T0) and ending at 21:41 (T3). Samples were collected using a peristaltic pump with acid-cleaned intake tubing kept inside the bottles for the entire experiment duration. Before sample collection, the bottles were gently hand-mixed and 20 mL was pumped from the bottles with no filter attached to flush the tubing lines.

**Grazing Plus Viral Mortality Dilution Experiment**

This dilution experiment was run in tandem with the two previously described experiments. In this case, the diluent was ultrafiltered at 100 kDa to remove both grazers and viruses for the three dilution treatments (70 %, 40 %, and 20 % whole water). The 0.8 µm/ 0.2 µm filtrate was filtered using tangential flow ultrafiltration (100 kDa ultracel, Millipore) at 1.5 L min⁻¹. The effectiveness of the 0.2 µm and 100 kDa filtration was determined using epifluorescence microscopy to count cells and viruses before and after filtration. The 0.2 µm filtration removed 99 % of bacteria and kept 98 % of viruses. The 100 kDa filtration removed 100 % of bacteria and ~100 % of viruses. The remainder of the experiment was carried out as described above in the grazing-only dilution experiment. Whole water treatments were the same bottles as in the grazing only experiment.

**Quantifying Microbial Abundance**

**Total Prokaryote Abundance**

Total prokaryotic abundance was determined using an Attune Acoustic Focusing Cytometer according to Nelson et al. (2015) from whole water samples fixed with paraformaldehyde (0.5% final concentration) and stained with 1X SYBR Green I.
**DNA Extraction Protocol**

Samples for DNA extraction were collected by passing water through 0.22 μm polyethersulfone filters (Sterivex, Millipore); the exact filtrate volume was recorded (250-500 mL). Filters were added to MP Biomedicals Lysing Matrix A (No. 116910100) tubes with MC lysis buffer and homogenized using a MP Biomedicals FastPrep-96 bead beater. DNA extractions were completed using the Macherey-Nagel NucleoMag Plant Extraction Kit (No. 744400.4) with KingFisher accessory kit (No. 744951). Samples were eluted to a final volume of 110 μL.

**Quantifying Vibrio vulnificus**

There is a consistent log correlation between CHROMagar *Vibrio* blue colony forming units and *vvhA* gene copy concentrations (Nigro & Steward, 2015). Therefore, I used a 16S rRNA 5’ nuclease *vvhA* qPCR assay (Campbell & Wright, 2003; Holland et al., 1991; forward primer: 5’-TGTTTATGTTGAGAACGGTGACA-3’; reverse primer: 5’-TTCTTTATCTAGCCCAACTTG-3’; 5’-Nuclease probe: 5’-/56-FAM/CCGTTAACC/ZEN/GAACCACCGCAA/31ABkFQ/-3’) to determine total *V. vulnificus* abundance. A 25 μL reaction mixture was prepared with final concentrations of 1X Kapa Probe Force Master Mix (Kapa Biosystems), 0.9 μM of each primer, 0.5 μM of the labeled probe, 0.56 mg/ml bovine serum albumin (Thermofisher), and 5 μL of DNA template (11.4 – 22.7 mL sample water). All qPCR reactions were performed in triplicate with the final replicate diluted 10-fold to check for inhibition of amplification as suggested by Bustin et al. (2009). The cycling protocol consisted of an initial denaturation step at 95 °C for 10 minutes, followed by 40 cycles of denaturation at 95 °C for 15 s and a combined annealing/extension at 60 °C for 30 minutes. I created an eight-point standard curve (10 – 50,000 copies per reaction well) of genomic DNA from *V. vulnificus* (strain YJ016; Chen et al., 2003) with known gene copy numbers per reaction run in triplicate along with the environmental samples. Assay efficiency was calculated from multiple standard curves using the formula $E = -1 + 10^{(-1/slope)}$ (Pfaffl, 2001).

**Statistical Analysis**

Linear least squares regression analyses in R version 3.1.6 (R Core Team, 2019) were...
performed to calculate growth rates of total prokaryotes and *V. vulnificus* for each bottle replicate ("A, B, C") in the three experiments and across all replicates for each treatment ("R") according to the exponential growth rate formula, net growth rate (h⁻¹) = ln(concentration change) / time (h). To account for a possible lag, or acclimation, in the response of *V. vulnificus*, results were compared including and excluding the T0 time point. Growth rates for total prokaryotes were always lagged until the final time point, so growth calculations were based on changes in prokaryote concentrations between the third and final time point. Total prokaryotes also grew faster than *V. vulnificus* and are reported min⁻¹.

For the inhibition experiment, growth rate in the samples treated with cytochalasin B was interpreted as gross growth rate and growth rate in unamended, undiluted whole water was interpreted as net growth rate. An estimate of mortality rate was calculated as the difference between the two. For the dilution experiments, gross growth rate and mortality rate were calculated according to the following formula.

Net growth rate (h⁻¹) = – mortality rate (h⁻¹) * proportion whole water + gross growth rate (h⁻¹)
2.4 RESULTS

**Total Prokaryote Response**

*Inhibition Experiment*

Total prokaryotes grew at an average of 0.384 cells mL\(^{-1}\) min\(^{-1}\) in the inhibition treatment and 0.634 cells mL\(^{-1}\) min\(^{-1}\) in the control (Figure 2.3). Prokaryote density peaked at the second time point (~1 h) for replicates A and B and at the third time point (~2 h) for replicate B in the inhibition treatments. Prokaryote density declined in the whole water control until the final time point. Net growth rate was calculated individually for the three replicates in each treatment. There was not a significant difference in the growth rates between the whole water control and the inhibition treatment (p = 0.245). The difference growth rates between treatment and control (Figure 2.3) defined a mortality rate of 0.250 cells mL\(^{-1}\) min\(^{-1}\).

*Grazing Only Dilution Experiment*

Total prokaryotes in the grazer only dilution experiment grew only between the third and final time point (Figure 2.4) for all treatments. Individual growth rates per bottle varied from 0.1 to 0.6 cells mL\(^{-1}\) min\(^{-1}\). Average growth rates were similar across all treatments, even the whole water without TSB, resulting in a dilution curve with a weak positive slope (Figure 2.5). Mortality and gross growth rates are interpreted as -0.10 cells mL\(^{-1}\) min\(^{-1}\) and 0.28 cells mL\(^{-1}\) min\(^{-1}\) respectively. However, a negative mortality term implies that the dilution had a negative effect on growth rather than stimulating growth, so the mortality rate is not interpretable.

*Grazing and Viral Lysis Dilution Experiment*

Similarly, in the grazing plus viral lysis dilution experiment, growth in total prokaryotes was only detected between the third and final time points (Figure 2.6). Individual growth rates per bottle varied from 0.1 to 0.8 cells mL\(^{-1}\) min\(^{-1}\). The variance in growth rates of the replicates between treatments was substantial (see 20 % versus 40 % in Figure 2.6). Average growth rates were similar across all treatments, even the whole water without TSB, resulting in a dilution curve with a weak positive slope (Figure 2.7). Mortality and gross growth rates are interpreted as -0.17 cells mL\(^{-1}\) min\(^{-1}\) and 0.32 cells mL\(^{-1}\) min\(^{-1}\), respectively.
Viral Lysis Contribution to Mortality

The contribution of viral lysis to total prokaryote mortality is the grazing plus viral lysis mortality rate minus the grazing only mortality rate. However, both mortality rates were negative which indicates a flaw in the experimental design, so a viral contribution could not be calculated.

Vibrio vulnificus Response

Inhibition Experiment

For the *V. vulnificus* analysis, I compared the growth rates calculated from the whole dataset to growth rates calculated without the initial T0 time point to account for an acclimation lag. For the full dataset, the inhibition treatment growth rate was 0.26 *vvhA* gene copies mL$^{-1}$ h$^{-1}$ ($R^2 = 0.29$) and there was a significant effect of time on concentration ($p < 0.001$; Figure 2.8). The whole water control growth rate was -0.08 gene copies mL$^{-1}$ h$^{-1}$ ($R^2 = 0.02$) with no significant effect of time on density ($p = 0.41$). The inhibition treatment net growth rate (0.26 *vvhA* gene copies mL$^{-1}$ h$^{-1}$) is interpreted as the gross growth rate. The difference in inhibition treatment net growth rate and whole water net growth rate (0.34 *vvhA* gene copies mL$^{-1}$ h$^{-1}$; Figure 2.8) is interpreted as the mortality rate. There was not a significant difference between the treatments ($p = 0.12$).

For the dataset without the initial T0 time point, the inhibition treatment growth rate was 0.21 *vvhA* gene copies mL$^{-1}$ h$^{-1}$ ($R^2 = 0.12$) and there was no significant effect of time on concentration ($p = 0.06$; Figure 2.8). The whole water control growth rate was 0.06 gene copies mL$^{-1}$ h$^{-1}$ ($R^2 = 0.01$) with no significant effect of time on density ($p = 0.68$). The inhibition treatment net growth rate (0.21 *vvhA* gene copies mL$^{-1}$ h$^{-1}$) is interpreted as the gross growth rate. The difference in inhibition treatment net growth rate and whole water net growth rate (0.15 *vvhA* gene copies mL$^{-1}$ h$^{-1}$) is interpreted as the mortality rate. There was not a significant difference between the treatments ($p = 0.28$).

Grazing Only Dilution Experiment

For the full dataset, growth rates ranged from -0.08 to 0.68 *vvhA* gene copies mL$^{-1}$ h$^{-1}$ across the treatments (Figure 2.9). All treatments, except the whole water without TSB, had a significant effect of time on density ($p < 0.05$) suggesting resolvable growth. The highest growth
rate (0.68 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\)) was detected in the 20\% whole water treatment as expected. From the dilution curve (Figure 2.10, top panel), the mortality rate is interpreted as 0.44 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) and the gross growth rate is 0.59 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\).

There was a significant effect of dilution (proportion of whole water) on the net growth rates (p < 0.05) suggesting a resolvable dilution curve.

For the dataset without the initial T0 time point, growth rates ranged from 0.06 to 0.83 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) across the treatments (Figure 2.9). All treatments, except the whole water without TSB, had a significant effect of time on density (p < 0.05) suggesting resolvable growth. The highest growth rate (0.83 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\)) was detected in the 40\% whole water treatment. From the dilution curve (Figure 2.10, bottom panel), the mortality rate is interpreted as 0.37 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) and the gross growth rate is 0.76 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\). There was not a significant effect of dilution (proportion of whole water) on the net growth rates (p = 0.11).

\textit{Grazing and Viral Lysis Dilution Experiment}

For the full dataset, growth rates ranged from -0.08 to 0.57 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) across the treatments (Figure 2.11). All treatments, except the whole water without TSB, had a significant effect of time on density (P < 0.05) suggesting resolvable growth. The highest growth rate (0.57 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\)) was detected in the 40\% whole water treatment. From the dilution curve (Figure 2.12, top panel), the mortality rate is interpreted as 0.26 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) and the gross growth rate is 0.42 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\). There was not a significant effect of dilution (proportion of whole water) on the net growth rates (p = 0.38).

For the dataset without the initial T0 time point, growth rates ranged from 0.06 to 1.02 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) across the treatments (Figure 2.11). All treatments, except the whole water without TSB, had a significant effect of time on density (P < 0.05) suggesting resolvable growth. The highest growth rate (1.02 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\)) was detected in the 40\% whole water treatment. From the dilution curve (Figure 2.12, bottom panel), the mortality rate is interpreted as 0.73 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\) and the gross growth rate is 0.99 \textit{vvh}A gene copies mL\(^{-1}\) h\(^{-1}\). There was not a significant effect of dilution (proportion of whole water) on the net growth rates (p = 0.18).
Viral Lysis Contribution to Mortality

The contribution of viral lysis to total prokaryote mortality is the grazing plus viral lysis mortality rate minus the grazing only mortality rate. For the full dataset, the viral lysis mortality rate is \(-0.18\) \(vvhA\) gene copies mL\(^{-1}\) h\(^{-1}\) which is not interpretable since the combined mortality rate was smaller than the mortality rate of grazing alone. For the dataset without T0, the viral lysis mortality rate is 0.36 \(vvhA\) gene copies mL\(^{-1}\) h\(^{-1}\).
2.5 DISCUSSION

**General Conclusions**

Gross growth rates and mortality rates for all experiments are reported in Table 2.1. There are clearly different responses of the total prokaryote and *V. vulnificus* populations to the treatment manipulations. In the inhibition experiment, total prokaryote density peaked by the second time point (within 1 h). The total prokaryote density was stable or declining until the final time point in the dilution experiments, whereas the *V. vulnificus* population was reliably growing by the second time point for all treatments, even inhibition, except in the whole water without TSB. This result suggests that the whole water manipulations did have the growth stimulating effect that was expected by the experimental design for *V. vulnificus* density. In some cases, particularly when the T0 time point was removed, the 40 % whole water treatment had the highest growth rate which might suggest a saturation in gross growth rate before the most diluted treatment. However, overall, the more diluted treatments grew faster despite having an equivalent amount of TSB enrichment, suggesting a response to the removal of predators and/or viruses. There was also not a discernable nonlinear response in the dilution curve. The gross growth rates of the dilution experiments were ultimately closer than the inhibition experiment to the gross growth rates expected based on the multifactorial growth experiment (~0.6 h\(^{-1}\) based on a salinity of 11.19 and 24.57 °C, Figure 2.1). The lower rate for the inhibition experiment may have been a result of the lower temperature (20.5 °C) where the experiment was run in the lab.

A clear interpretation of these results is difficult to extract. The dilution may have unaccounted for effects, such as removing competitors which would stimulate growth and lead to an overestimation of mortality rate. In contrast, dilution may remove particles and nutrients not replaced by TSB enrichment which would depress or limit growth and lead to an underestimate of mortality rate.

**Recommendations for Future Tests**

There are several modifications to the experimental design which may improve our confidence in the mortality and gross growth rates. For one, this experiment was rather short (3 h duration) compared to the traditional 24 h for these experiments (Mojica et al., 2016; Worden
and Binder, 2003) or even longer incubations such as in Tijdens et al. (2008) who ran incubations for 40 h. A longer experiment duration might lead to a more distinguishable growth curve where the lag, or acclimation, period can be removed with more confidence. There was a clear lag in response by total prokaryotes. It is possible that the total prokaryote density was not changing for the first few time points because the relative abundance of populations within the community was changing and acclimating to the treatment before overall growth was resolvable. There was also a possible lag in response by *V. vulnificus* which motivated evaluation of the data without the initial time point. There was also a long period (~ 7 h) between water collection and the start of the experiment. It took ~ 4 h to prepare the different treatments which meant that the experiment was running in the light of the lab when it was naturally dark outside. The diel cycle of growth is notably important in the growth of microbes (Kelly et al., 2019; Ottesen et al., 2014). A smaller scale experiment with fewer treatments would accelerate the time to collect water, create the treatments, and begin the experiment. Performing the experiment in the field, possibly submerging the bottles *in situ* or at least running the experiment at *in situ* temperature, would likely reduce the time for acclimation. Finally, there is also a concerning amount of variation between the replicates in each treatment for *V. vulnificus* density despite our best efforts to aliquot the replicates immediately preceding the experiment. As in Worden et al. (2015), inoculation with *V. vulnificus* may be necessary in order to standardize the initial population abundance and inflate the natural low abundance population to reliably detectable levels with qPCR. However, this inoculation could change community interactions and affect results.

Ultimately, the variation in mortality rate across the physical and chemical environment of the canal is more important than the absolute rate numbers for modeling the population dynamics. A reliable sampling design with the fewest treatments possible, such as the inhibition experiment or a 2-point dilution experiment, with the fewest sampling times possible would allow for coverage of more locations and environmental conditions. Mortality rate as a function of temperature and salinity would complement the growth rate as a function of temperature and salinity to improve the existing physical-microbial model for real time predictions of *V. vulnificus* in the Ala Wai Canal.
Table 2.1 Summary of mortality and gross growth rates from all three experiments. *V. vulnificus* data are separated into two datasets: the full dataset and the dataset without T0. The viral contribution to mortality is calculated for the *V. vulnificus* dilution curves without T0. It is not calculated for the total prokaryote or *V. vulnificus* full dataset dilution curves because the values are not interpretable.

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<th>Total prokaryotes</th>
<th>V. vulnificus full dataset</th>
<th>V. vulnificus without T0</th>
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<tr>
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<td>Gross growth rate (cells mL⁻¹ min⁻¹)</td>
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<td>Grazing only dilution experiment</td>
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Figure 2.1 Laboratory growth rates of *Vibrio vulnificus* isolated from the Ala Wai Canal at various combinations of salinity and temperature (Steward and Nigro, unpublished).
Figure 2.2 Idealized dilution experiment curve based on Landry and Hassett (1982) and Landry et al. (1995) using mock data. Population abundance is tracked over time for each dilution treatment (left panel) in triplicate. The resulting growth rates are plotted against the proportion of whole water to generate the dilution curve (right panel). The singular growth rate generated from the left panel is circled in red on the right panel. The mortality rate is interpreted as negative the slope of the curve (0.96 h\(^{-1}\)). The gross growth rate is interpreted as the intercept of the curve (the theoretical full dilution where predation pressure would effectively be zero) corrected by the difference between the growth rates of the enriched and unenriched whole water treatments. Gross growth rate = intercept – correction factor = 1.33 – 0.22 = 1.11 h\(^{-1}\). In this case, enrichment is with tryptic soy broth (TSB).
Figure 2.3 Growth rates of total prokaryotes in whole water untreated and with cytochalasin B eukaryotic inhibitor. Values are shown as points in addition to the mean as a bar and standard error around the mean. Gross growth is interpreted as the growth rate in the inhibited treatment (0.384 min⁻¹). Mortality rate is interpreted as the difference in growth rate between the inhibited and whole water treatment (0.634 – 0.384 = 0.250 min⁻¹). There was no significant difference between the growth rates of the control and treatment (t-test, p = 0.245).
Figure 2.4 Growth curves of total prokaryotes for each treatment in the grazing only dilution experiment (0.2 µm diluent). 100% corresponds to the whole water, undiluted treatment. TSB is the organic matter enrichment.

Figure 2.5 Dilution curve for total prokaryote growth rates from the grazing only dilution experiment (see Figure 2.4). Gross growth rate is 0.28 min\(^{-1}\) and mortality rate is -0.10 min\(^{-1}\).
Figure 2.6 Growth curves of total prokaryotes for each treatment in the grazing plus viral lysis dilution experiment (100 kDa diluent). 100% corresponds to the whole water, undiluted treatment. TSB is the organic matter enrichment.

Figure 2.7 Dilution curve for total prokaryote growth rates from the grazing plus viral lysis dilution experiment (see Figure 2.6). Gross growth rate is 0.32 min\(^{-1}\) and mortality rate is -0.17 min\(^{-1}\).
Figure 2.8 Growth rates of *V. vulnificus* in whole water untreated and with cytochalasin B eukaryotic inhibitor. Linear regression analysis was performed with the initial time point (T0) included (grey line) and without the initial time point to account for a lag in growth (black line). Growth rates (h\(^{-1}\)), R\(^2\), and p-values are included for each growth curve. There was not statistically significant growth in the whole water control (p > 0.05).

The two bottom panels are comparing the average growth rates between the treatments where the bottom left panel includes T0 in calculations and the bottom right panel does not include T0. Gross growth is interpreted as the growth rate in the inhibited treatment (0.25 h\(^{-1}\) including T0; 0.21 h\(^{-1}\) without T0). Mortality rate is interpreted as the difference in growth rate between the inhibited and whole water treatment (0.34 h\(^{-1}\) including T0; 0.15 h\(^{-1}\) without T0). There was no significant difference between the growth rates of the control and treatment (t-test, p = 0.117 including T0; p = 0.281 without T0).
**Figure 2.9** Growth curves of *V. vulnificus* from the grazing only dilution experiment (0.2 μm diluent). 100% corresponds to the whole water, undiluted treatment. Linear regression analysis was performed with the initial time point (T0) included (grey line) and without the initial time point to account for a lag in growth (black line). Growth rates (h⁻¹), R², and p-values are included for each growth curve. As intended, there was statistically significant growth in all treatments (p < 0.05) except for the whole water without enrichment.
Figure 2.10 Dilution curves for *V. vulnificus* growth rates from the grazing only dilution experiment (see Figure 2.9). Linear regression analysis was performed with the initial time point (T0) included in growth rate calculations for the top panel (grey line) and without the initial time point for the bottom panel (black line). Replicate R is the average growth rate across all the replicates (A, B, C) of a particular treatment. The dilution curve regression is evaluated with replicate R. The average of the whole water without TSB replicates (open circles) is the correction factor which is subtracted from the intercept to calculate the gross growth rate. Gross growth rates were 0.59 h\(^{-1}\) including T0 and 0.76 h\(^{-1}\) without T0. Mortality rate rates were 0.44 h\(^{-1}\) including T0 and 0.37 h\(^{-1}\) without T0. There was no significant effect of dilution (p = 0.05 including T0; p = 0.11 without T0).
Figure 2.11 Growth curves of *V. vulnificus* from the grazing plus viral lysis dilution experiment (100 kDa diluent). 100% corresponds to the whole water, undiluted treatment. Linear regression analysis was performed with the initial time point (T0) included (grey line) and without the initial time point (black line) to account for a lag in growth. Growth rates (h⁻¹), $R^2$, and p-values are included for each growth curve. As intended, there was significant growth in all treatments (p < 0.05) except for the whole water without enrichment.
Figure 2.12 Dilution curves for *V. vulnificus* growth rates from the grazing and viral lysis dilution experiment (see Figure 2.11). Linear regression analysis was performed with the initial time point (T0) included in growth rate calculations for the top panel (grey line) and without the initial time point for the bottom panel (black line). Replicate R is the average growth rate across all the replicates (A, B, C) of a particular treatment. The dilution curve regression is evaluated with replicate R. The average of the whole water without TSB replicates (open circles) is the correction factor which is subtracted from the intercept to calculate the gross growth rate. Gross growth rates were 0.42 h\(^{-1}\) including T0 and 0.99 h\(^{-1}\) without T0. Mortality rate rates were 0.26 h\(^{-1}\) including T0 and 0.73 h\(^{-1}\) without T0. There was no significant effect of dilution (p = 0.38 including T0; p = 0.11 without T0).
2.8 ACKNOWLEDGEMENTS

I thank Grieg Steward for his assistance with designing and running the experiments and interpreting the data. I thank Brian Powell for attempting to make meaningful results from the data. I thank the interns and volunteers who assisted with collection of samples from the canal and running the experiments: Shaun Wriston, Han Quach, and Brenna Carroll. This project was funded by NOAA Sea Grant.
2.9 LITERATURE CITED


