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# Review



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# Vibrio bacteria in raw oysters: managing risks to human health

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The human-pathogenic marine bacteria *Vibrio vulnificus* and *V. parahaemolyticus* are strongly correlated with water temperature, with concentrations increasing as waters warm seasonally. Both of these bacteria can be concentrated in filter-feeding shellfish, especially oysters. Because oysters are often consumed raw, this exposes people to large doses of potentially harmful bacteria. Various models are used to predict the abundance of these bacteria in oysters, which guide shellfish harvest policy meant to reduce human health risk. *Vibrio* abundance and behaviour varies from site to site, suggesting that location-specific studies are needed to establish targeted risk reduction strategies. Moreover, virulence potential, rather than simple abundance, should be also be included in future modeling efforts.

### 1. Introduction

All coastal waters and estuaries contain Vibrio bacteria. Although many Vibrio species are harmless, several can cause serious disease in humans or animals. Vibrio vulnificus and V. parahaemolyticus are the most common types of sometimes deadly foodborne and wound Vibrio infections. Recognized infections from Vibrio species are on the rise, and although there is some uncertainty, most researchers predict that climate change will increase cases [1–5]. Research on microbial ecology, food safety and ecology of shellfish reservoirs has revealed insights into Vibrio ecology and led to tools for predicting risky periods or locations. Nonetheless, regional and strain variation complicate predictions. Vibrio vulnificus and V. parahaemolyticus are detectable in coastal waters and estuaries with salinities greater than 5% [6]. These bacteria are 'particlelovers' and will attach to particulates and other organisms in the water column. This attachment facilitates their uptake by filter-feeding molluscs, such as oysters, that share similar habitats. Through this substrate filtration, oysters can concentrate these bacteria, including the pathogenic forms of these bacteria, by up to 100 times the concentration found in the surrounding waters [7]. Because oysters are often eaten raw, this can pose an infection risk for people consuming them [8]. Most infections are unreported, and it is estimated that in the USA approximately 84 000 people contract a food-borne infection from Vibrio spp. every year, the highest rate since nationwide reporting began [9]. V. parahaemolyticus is, by far, the most common infective agent of the two species. Infections with V. parahaemolyticus produce symptoms that include diarrhoea accompanied by abdominal cramps, nausea, vomiting, headache, chills and low-grade fever [10]. Some strains with increased virulence have been implicated in recent outbreaks, with serotype O4:K12 garnering lots of recent attention [11-13]. Foodborne V. vulnificus infections are less common than V. parahaemolyticus, but the morbidity associated with these infections is far more severe. With this species, the initial symptoms can be the same as V. parahaemolyticus, but the infection can progress rapidly (within hours) to primary septicaemia [14,15]. Here the symptoms are grave, including hypotension and secondary lesions that develop on the extremities. When this occurs, the incubation period is often very short, and death can occur 24-72 h after eating a single oyster [16]. V. vulnificus is the single most fatal foodborne pathogen in the USA, where it comprises 95% of all seafood-related deaths and boasts a fatality rate nearing 50%, even with aggressive medical treatment. Rarely seen in most bacteria, these bacteria also have a second route of infection. They can enter the body via wounds, either preexisting or obtained through coast-related activities such as fishing, swimming, oyster harvesting or handling seafood. Wound infections from these bacteria progress rapidly, and lead to necrotizing fasciitis, also known as 'flesh-eating disease', at the site of infection. Jones & Oliver [16] and Letchumanan et al. [17] describe details on pathogenesis and diseases caused by V. vulnificus and V. parahaemolyticus, respectively.

Almost all (90%) V. vulnificus and V. parahaemolyticus infections stem from eating undercooked or raw oysters, and there are in place several regulations and tools that reduce risk. The Interstate Shellfish Sanitation Conference (ISSC) was formed in 1982 with the goal of promoting a safe shellfish product by addressing sanitation and cooperating with state and federal agencies, industry and academia. The ISSC issues a National Shellfish Sanitation Program guide for the control of molluscan shellfish (NSSP) recognized by the US Food and Drug Administration (FDA). Included in the guide is a Model Ordinance with provisions that ensure shellfishproducing states are in compliance with safety guidelines [18]. Contained within the model ordinance are outlines for shellfish dealer certification, shellfish handling and processing and plant inspection procedures. Furthermore, there are sections that specifically consider how to control *V. parahaemolyticus* and *V. vulnificus*.

When a threshold of laboratory-confirmed V. parahaemolyticus cases is exceeded, the shellfish harvest areas that are implicated are closed for at least 7-21 days. After the minimum time has passed and prior to re-opening, shellfish samples must be collected from the harvesting area to ensure fewer than 10 pathogenic bacteria are counted per gram of oyster meat. Although the specific requirement to quantify pathogenic V. parahaemolyticus via tdh or trh (see below for detail on these genes) is commendable, the correlation between these markers and infections is not straightforward.

To control for *V. vulnificus*, the NSSP requires that oysters for raw consumption must meet minimum time-to-temperature requirements during harvest periods, where average monthly water temperatures exceed 26.6°C (80°F). Neither of these, nor any of the other regulations set forth by the NSSP, are arbitrary. Rather, they were carefully conceived using an extensive riskper-serving model for *V. parahaemolyticus* and the assessment that less than 30 V. vulnificus cells per gram of oyster is of little risk [19,20]. Additionally, the NSSP contains a model for V. parahaemolyticus growth rate inside oysters, which is used to determine control measures. States are required to institute control plans for *V. vulnificus* if there have been two infections within the last 10 years. V. parahaemolyticus control plans are initiated based on two infections in a 3 year period, an outbreak having occurred within 5 years, or if average water temperatures during harvest exceed 27.2°C (81°F) [18]. Although these regulations reduce infection risk, the number of infections is still increasing.

To further reduce infections during autumn 2009, the FDA announced that Gulf Coast oysters harvested during warm water months could not be served raw without first undergoing post-harvest processing (PHP) [21]. Through PHP, oysters are treated using such techniques as high pressure, irradiation, pasteurization-like heating or flash freezing to reduce the number of harmful bacteria. PHP leaves the oyster raw, but no longer alive. Treatment could have lowered the infection rate from uncooked or raw shellfish, and such

regulations have already existed in California since 2003. Nevertheless, the response to this ban was overwhelmingly negative. State officials, local legislators, the shellfishing industry and the public raised such an outcry over increased production costs, altered taste of oysters, and even the 'right' to eat whatever one chooses that the FDA suspended the ban a month later [22]. A similar attempt at a warmweather Gulf Coast oyster ban failed in 1995, when the ISSC rejected the FDA-proposed action, instead enacting strict time-to-temperature rules [23].

Harvest restrictions, while effective at reducing infections, do not eliminate the problem. Furthermore, outright bans on marketing raw shellfish severely affect the industry and are unwelcome by some consumers. There is a need for tiered regulation that simultaneously reduces infection risk without ceasing to provide raw shellfish altogether. A further complication is that overstating risk can cause desensitization to risk-based statements. Desensitization can be reduced by educating shellfish growers and the shellfish consuming public. For example, California was the first state, in 1991, to require that those locations serving Gulf Coast oysters warn customers about infection risk [23]. The warning has since been modified from the original wording and now reads, in both English and Spanish, 'This facility offers raw oysters from the Gulf of Mexico. Eating these oysters may cause severe illness and even death in persons who have liver disease (for example alcoholic cirrhosis), cancer or other chronic illnesses that weaken the immune system. If you eat raw oysters and become ill, you should seek immediate medical attention. If you are unsure if you are at risk, you should consult your physician' [24, p. 2]. Although many states now have similar warnings, Mouzin et al. [23] concluded that a warning fails to reduce infections.

Even though the Vibrio models referenced by the NSSP are high quality and robust, they have limitations. For V. vulnificus, most studies find that temperature drives bacterial concentrations [25-34], V. vulnificus most easily being isolated in oyster tissues with a temperature range from 15°C to 17°C [26,27,29,32]. Exceptions include Parvathi et al. [35], Jones et al. [36] and Givens et al. [37], though a common limitation to these studies is a lack of variation in temperature over the sampling period. Models assume that salinity does not play a factor in V. vulnificus abundance, but reported effect of salinity on V. vulnificus varies from location to location, including insignificant [25,30,32,37], positive [29], inhibitory [26,28,31,38] or nonlinear [27] associations. Finally, both regulations are 'reactive' rather than 'proactive', requiring cases to be noted, or concentrations of bacteria in warmer waters to be high prior to action. Thus, there is vast room for improvement in monitoring and modelling pathogenic forms of vibrios, and data gaps that can be filled. Furthermore, the FDA V. parahaemolyticus model predicts bacterial abundance based on water temperature [20], but it is becoming obvious that models should consider other environmental factors. For example, in the Gulf Coast region, Zimmerman et al. [39] found significant, but site-dependent, relationships between V. parahaemolyticus and turbidity and salinity, but not chlorophyll a.

Observed regional variation in Vibrio ecology suggests a need for site-specific models. A study based outside of the Gulf Coast region, in the Chesapeake area, by Parveen et al. [39] found that unlike along the Gulf Coast, in the Chesapeake area neither salinity nor chlorophyll a were significant factors, but rather that V. parahaemolyticus abundance was driven by water temperature, turbidity and dissolved oxygen [40]. An ambitious project by Johnson et al. [25] incorporated data from multiple regions, including the Gulf Coast, Northwest and Northeast regions of the USA. They noted that total suspended solids, dissolved organic carbon and salinity, in addition to water temperature, were significant controlling factors of V. parahaemolyticus in oysters, while salinity and chlorophyll a were not [25]. They also concluded that stateby-state modelling should be considered, based on the variation that they observed at each site they conducted experimentation. When multiple sites in South Carolina were sampled, the combinatory impacts of salinity and temperature appeared to be complicated, with some sites showing that both salinity and temperature correlated with V. parahaemolyticus abundance, some sites neither, or some with only temperature [41]. In a recent North Carolina-specific study, no correlation with salinity was observed [26]. This study also found 10fold fewer V. parahaemolyticus cells per gram of shellfish than has been reported on average for similar water temperatures in oysters from the Gulf Coast, further highlighting the importance of region-specific studies. Outside of the USA, a study on shellfish in Mexico found that turbidity actually had a negative effect on V. parahaemolyticus concentrations in oysters [42]. While vibrios are often particle attached, Lopez-Hernandez et al. [42] hypothesized that non-digestible particles led to downshifts in filter-feeding in oysters. In Brazil, V. parahaemolyticus was found to be negatively correlated with salinity, and studies in New Zealand have demonstrated that salinity is unrelated or negatively correlated to V. parahaemolyticus shellfish abundances, depending on the study [43,44].

For *V. vulnificus*, the story is less convoluted but still variable. Nearly all studies that examine the number of *V. vulnificus* cells in oysters find that temperature is one of the major driving forces in determining bacterial concentrations [25-34]. Exceptions are by Parvathi et al. [35], Jones et al. [36] and Givens et al. [37], and a common limitation to these studies was the lack of variation in temperature over the sampling period. Of those that have shown positive correlations to temperature, there was tight agreement on a temperature threshold for when V. vulnificus is most easily isolated in oyster tissues. This temperature range is from 15°C to 17°C [26,27,29,32]. Conversely, reports on salinity for V. vulnificus are more variable from location to location. The effect of salinity on V. vulnificus oyster populations has been calculated to be insignificant by some [25,30,32,37], positive by others [29], inhibitory by yet others [26,28,31,38] or in the case of the study by Motes et al. [27], nonlinear.

One important note for correlation assessments for both V. vulnificus and V. parahaemolyticus is that even when environmental parameters have been important driving factors of abundance, there are still 'missing factors'. Regional differences in both the bacterial communities and their reactions to environmental stimuli make clear the need for multiple, focused studies of the Vibrio species. When predictive models or risk assessments are generated, these tools appear to only be relevant to the local area at best. In fact, even when a single region is examined, there are often siteby-site differences in prediction capabilities making even regional management of risk a challenge. It is not possible to manage Vibrio-associated infection risk under a single modelling system, and any attempt to do this would be difficult and inaccurate. Each oyster-producing state would need to develop their own dataset to increase confidence in our ability to forecast, now-cast or hindcast these bacteria

in oysters. But these spatial aberrations are only the first of several obstacles.

Strains of V. vulnificus and V. parahaemolyticus vary in virulence, with some being pathogenic and others seemingly harmless [10,16]. The FDA and ISSC acknowledge that strain virulence should be considered, but at the time the NSSP was created there was not enough data to implement such fine detail. It seems evident that simple quantification of V. vulnificus and V. parahaemolyticus does have merit, and has been successfully used in regulation and in other microbial risk assessment tools [18-20,45,46]. Still, evidence continues to mount that the quantity of a particular Vibrio species inside an oyster may not be as important as the number of patho-

For *V. parahaemolyticus*, potential pathogenicity is indicated by the presence of genes for thermostable direct haemolysin (tdh) and tdh-related haemolysin, trh. Briefly, the strains that possess at least one of these genes are often associated with cytotoxicity within the host [47,48]. However, there are other virulence-related factors, and there are also tdh- or trh-positive strains found in environmental samples and clinical isolates that contain neither gene [17]. Most studies that attempt to correlate tdh- and/or trh-positive V. parahaemolyticus strains in oysters to environmental parameters have had little success. Often, there are not enough of these strains above the limit of detection to make statistically significant comparisons [7,37,40,44,49,50]. Occasionally, when there are sufficient data on pathogenic strains, correlations with other measured data are still non-significant [51]. Though admittedly variable, Johnson et al. [32] and Zimmerman et al. [39] found that turbidity was correlated with trh- and tdh-positive strains and not with concentrations of total *V. parahaemolyticus*. Intriguingly, temperature was not significantly related to concentrations of pathogenic strains [32,39]. By contrast, Jones et al. [36] observed that when temperature did not affect total concentrations of V. parahaemolyticus, it did have an effect on tdh- and trh-positive strains.

Several different methods are available for detecting potentially pathogenic V. vulnificus strains. Most oyster studies use genetic differences found in either the virulence-correlated gene (vcg) or in the 16S rDNA gene [52-56]. Clinically isolated V. vulnificus strains are associated with the 'C' genotype of vcg and the 'B' type of the 16S rRNA gene, whereas strains isolated from the environment correlate with the 'E' and 'A' types of vcg and 16S, respectively. For simplicity, the potentially pathogenic strains, determined by either method, will be termed 'P' while those less likely to cause septicaemia will be called 'NP' (nonpathogenic). There are only a handful of studies that examine the prevalence of virulent V. vulnificus in oysters. As with V. parahaemolyticus, the relatively low occurrence of these strains, especially outside of the Gulf Coast region, makes even targeted analyses difficult [26,37]. Several authors agree that the P strains of V. vulnificus are more temperature-sensitive than NP strains, where the P strains in oysters increase in relative abundance to NP strains as water temperatures increase [31,57,58]. A similar reaction has been observed in V. vulnificus as has been noted in V. parahaemolyticus, where the normal drivers of abundance, temperature and salinity in the case of V. vulnificus, are not the factors that appear to influence the concentration of pathogenic strains. Factors such as dissolved oxygen and pH correlate with P strains while not seeming to affect total V. vulnificus (B. A. Froelich and R. T. Noble 2016, unpublished data). Thus, it

appears that the pathogenic strains of both V. vulnificus and V. parahaemolyticus have a niche different from that of the species as a whole, hence deserving further attention.

Neither total V. parahaemolyticus nor total V. vulnificus concentration appears to be related to the observed concentration of pathogenic strains in oysters [26,31,32,37,58]. One notable exception is presented by Han et al. [51], who found significant correlations between total and pathogenic V. parahaemolyticus in oysters, despite a ratio between the two that varied widely. However, most pathogenic strains in that study were obtained from Chinese markets, rather than directly from the environment, and the authors remark that handling practices could alter pathogenic V. parahaemolyticus levels [51].

Although it seems obvious to link water quality with oyster quality, most studies that examine environmental effects on V. vulnificus or V. parahaemolyticus find less variation in the water column than in oysters. In fact, it is not uncommon for bacterial concentrations inside oysters to be quite different from the water that they were harvested from, even when sampling is tightly controlled temporally and spatially [26,31]. Furthermore, variation in Vibrio concentrations from oyster to oyster can be orders of magnitude, even when oysters are collected from the exact same clutch. It seems that while Vibrio spp. in the water column react quickly to changing conditions, once they colonize the interior of the oysters they remain relatively stable. It is hypothesized that the Vibrio colonization of oysters occurs at the larval settlement stage, and exogenous bacteria in adult oysters are merely transient [59]. Major ecological disruptions, such as large upshifts in salinity, can disrupt these resident populations, but nominal conditions may not have a large intra-oyster effect [38,59-61]. This independence from the surrounding water makes it difficult to predict pathogenic Vibrio concentrations within oysters from environmental parameters.

In the face of likely changing Vibrio concentrations with impending climate change and warming estuarine and coastal waters, we are left with hurdles to improving regulations that aim to reduce food-borne infections. Current US regulations are based on obsolete data, are relevant mostly to a specific region of the USA, and make simplifying assumptions. Moreover, current risk assessment does not take into account the differences in growth rates of different strains among different oyster species. Simultaneously, outbreaks of illness are expanding latitudinally to locations never suspected of Vibrio-associated risk, while oyster growers and commercial markets are increasing with the burgeoning boutique oyster business [62,63]. The current rules are protective and reduce risk, yet infections are still increasing. Current guidelines might not be appropriate for every state or region, as differences in Vibrio population structure and abundance vary from location to location. Underprotection leads to health risks, while overprotection increases costs. Increasing the number of location-specific studies would help strengthen the understanding of these dynamic naturally occurring pathogens. Future studies should include specific quantification of pathogenic forms of each species, as it is clear that concentrations at the species level do not necessarily indicate health risks. For V. vulnificus, this can include either C/E genotyping, pilF polymorphism to determine serum resistance potential, or AB ribotyping [54,55,64]. For *V. parahaemolyticus*, at the minimum, research should include detection and quantification of the tdh and trh genes. Newer types of equipment, such as digital PCR, and advanced techniques, including nextgeneration sequencing, are becoming cheaper and more available. Sequencing environmental samples and using digital PCR to detect rare sequences can help overcome the problem of pathogen rarity and improve pathogen-specific modelling. Region-specific, hydrodynamic models that take into account environmental and biogeochemical factors might also improve understanding. Including environmental factors, such as salinity, temperature, pH, dissolved oxygen and chlorophyll a, along with novel factors, such as oyster ploidy, oyster condition, particulates, chitin or strain pathogenicity might further the understanding of this complex and important problem.

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# References

- 1. Burge CA et al. 2014 Climate change influences on marine infectious diseases: implications for management and society. Annu. Rev. Mar. Sci. 6, 249-277. (doi:10.1146/annurev-marine-010213-135029)
- 2. Levy S. 2015 Warming trend: how climate shapes Vibrio ecology. Environ. Health Perspect. 123, A82 – A89. (doi:10.1289/ehp.123-A82)
- Martinez-Urtaza J, Bowers JC, Trinanes J, DePaola A. 2010 Climate anomalies and the increasing risk of Vibrio parahaemolyticus and Vibrio vulnificus illnesses. Food Res. Int. 43, 1780-1790. (doi:10. 1016/j.foodres.2010.04.001)
- Urquhart EA, Zaitchik BF, Waugh DW, Guikema SD, Del Castillo CE. 2014 Uncertainty in model predictions of Vibrio vulnificus response to climate variability and change: a Chesapeake Bay case study. PLoS ONE 9, e98256. (doi:10.1371/journal. pone.0098256)
- Foodborne Diseases Active Surveillance Network (FoodNet) & Centers for Disease Control and Prevention. 2011 Vital signs: incidence and trends of infection with pathogens transmitted commonly through food. MMWR Morbid. Mortal. Wkly Rep. 60, 749 – 755.
- Hidetoshi U, Irma NGR. 2006 Aquatic environment. In The biology of Vibrios (eds FL Thompson, B Austin, J Swings), pp. 175 – 189. Washington, DC: ASM Press.
- DePaola A, Nordstrom JL, Bowers JC, Wells JG, Cook DW. 2003 Seasonal abundance of total and pathogenic Vibrio parahaemolyticus in alabama oysters. Appl. Environ. Microbiol. 69, 1521-1526. (doi:10.1128/AEM.69.3.1521-1526.2003)
- Colwell RR. 2006 A global perspective. In The biology of Vibrios (eds FL Thompson, B Austin,

- J Swings), pp. 3-11. Washington, DC: American Society of Microbiology.
- Centers for Disease Control and Prevention. 2015 Foodborne Diseases Active Surveillance Network (FoodNet): FoodNet Surveillance Report for 2013 (Final Report). Atlanta, GA: US Department of Health and Human Services. See http://www.cdc. gov/foodnet/reports/annual-reports-2013.html.
- 10. Yeung PSM, Boor KJ. 2004 Epidemiology, pathogenesis, and prevention of foodborne Vibrio parahaemolyticus infections. Foodborne Pathoq. Dis. **1**, 74-88. (doi:10.1089/153531404323143594)
- 11. Abbott SL, Powers C, Kaysner CA, Takeda Y, Ishibashi M, Joseph SW, Janda JM. 1989 Emergence of a restricted bioserovar of Vibrio parahaemolyticus as the predominant cause of Vibrio associated gastroenteritis on the West Coast of the United States and Mexico. J. Clin. Microbiol. 27, 2891 - 2893. (doi:10.1128/IAI.01046-08)
- 12. Martinez-Urtaza J, Baker-Austin C, Jones JL, Newton AE, Gonzalez-Aviles GD, DePaola A. 2013 Spread of Pacific Northwest Vibrio parahaemolyticus strain. N. Engl. J. Med. **369**, 1573 – 1574. (doi:10.1056/ NEJMc1305535)
- 13. Turner JW, Paranjpye RN, Landis ED, Biryukov SV, González-Escalona N, Nilsson WB, Strom MS. 2013 Population structure of clinical and environmental Vibrio parahaemolyticus from the Pacific Northwest Coast of the United States. PLoS ONE 8, e55726. (doi:10.1371/journal.pone.0055726)
- 14. James DO. 2006 Vibrio vulnificus. In The Biology of Vibrios (eds FL Thompson, B Austin, J Swings), pp. 349-366. Washington, DC: American Society of Microbiology.
- 15. Centers for Disease Control and Prevention. Vibrio illness (vibriosis). US Department of Health and Human Services. See http://www.cdc.gov/vibrio/ vibriov html
- 16. Jones MK, Oliver JD. 2009 Vibrio vulnificus: disease and pathogenesis. Infect Immun. 77, 1723-1733. (doi:10.1128/IAI.01046-08)
- 17. Letchumanan V, Chan K-G, Lee L-H. 2014 Vibrio parahaemolyticus: a review on the pathogenesis, prevalence, and advance molecular identification techniques. Front. Microbiol. 5, 705. (doi:10.3389/ fmicb.2014.00705)
- 18. NSSP. 2013 Guide for the Control of Molluscan Shellfish 2013 Revision, Interstate Shellfish Sanitation Conference. See http://www.issc.org/ nssp/default.aspx.
- World Health Organization. 2005 Risk assessment of Vibrio vulnificus in raw oysters. Geneva, Switzerland: WHO.
- 20. Center for Food Safety and Applied Nutrition. 2005 Quantitative risk assessment on the public health impact of pathogenic Vibrio parahaemolyticus in raw oysters.
- 21. Taylor M. 2009 Address at the Interstate Shellfish Sanitation Conference Biennial Meeting.
- 22. US Food and Drug Administration. 2009 FDA Statement on Vibrio vulnificus in raw oysters.
- 23. Mouzin E, Mascola L, Tormey M, Dassey D. 1997 Prevention of Vibrio vulnificus infections: assessment

- of regulatory educational strategies. JAMA 278, 576 - 578. (doi:10.1001/jama.1997.03550070 068040)
- 24. State Department of Health Services. In press. Raw Gulf oysters: labeling, written warnings and additional requirements. Chapter 5 Subchapter 2. Group 1.
- 25. Johnson CN et al. 2012 Ecology of Vibrio parahaemolyticus and Vibrio vulnificus in the coastal and estuarine waters of Louisiana, Maryland, Mississippi, and Washington (United States). Appl. Environ. Microbiol. 78, 7249-7257. (doi:10.1128/ AEM.01296-12)
- 26. Froelich BA, Ayrapetyan M, Fowler P, Oliver JD, Noble RT. 2015 Development of a matrix tool for the prediction of Vibrio species in oysters harvested from North Carolina. Appl. Environ. Microbiol. 81, 1111 – 1119. (doi:10.1128/AEM.03206-14)
- 27. Motes ML, DePaola A, Cook DW, Veazey JE, Hunsucker JC, Garthright WE, Blodgett RJ, Chirtel SJ. 1998 Influence of water temperature and salinity on Vibrio vulnificus in Northern Gulf and Atlantic Coast oysters (Crassostrea virginica). Appl. Environ. Microbiol. 64, 1459-1465.
- 28. Staley C, Chase E, Harwood VJ. 2013 Detection and differentiation of Vibrio vulnificus and V. sinaloensis in water and oysters of a Gulf of Mexico estuary. *Environ. Microbiol.* **15**, 623 – 633. (doi:10.1111/ 1462-2920.12045)
- 29. Tamplin M, Rodrick GE, Blake NJ, Cuba T. 1982 Isolation and characterization of Vibrio vulnificus from two Florida estuaries. Appl. Environ. Microbiol. **44**, 1466 – 1470.
- 30. Lin M, Payne DA, Schwarz JR. 2003 Intraspecific diversity of Vibrio vulnificus in Galveston Bay water and oysters as determined by randomly amplified polymorphic DNA PCR. Appl. Environ. Microbiol. 69, 3170-3175. (doi:10.1128/AEM.69.6. 3170-3175.2003)
- 31. Warner EB, Oliver JD. 2007 Population structures of two genotypes of Vibrio vulnificus in oysters (Crassostrea virginica) and seawater. Appl. Environ. *Microbiol.* **74**, 80 – 85. (doi:10.1128/AEM.01434-07)
- 32. Johnson CN, Flowers AR, Noriea NF, Zimmerman AM, Bowers JC, DePaola A, Grimes DJ. 2010 Relationships between environmental factors and pathogenic vibrios in the Northern Gulf of Mexico. Appl. Environ. Microbiol. 76, 7076-7084. (doi:10. 1128/AEM.00697-10)
- 33. Chase E, Young S, Harwood VJ. 2015 Sediment and vegetation as reservoirs of Vibrio vulnificus in the Tampa Bay Estuary and Gulf of Mexico. Appl. Environ. Microbiol. 81, 2489-2494. (doi:10.1128/ AEM.03243-14)
- 34. Wright AC, Hill RT, Johnson JA, Roghman MC, Colwell RR, Morris JG. 1996 Distribution of Vibrio vulnificus in the Chesapeake Bay. Appl. Environ. Microbiol. **62**, 717-724.
- 35. Parvathi A, Kumar HS, Karunasagar I, Karunasagar I. 2004 Detection and enumeration of Vibrio vulnificus in oysters from two estuaries along the Southwest Coast of India, using molecular methods. Appl. Environ. Microbiol. **70**, 6909 – 6913. (doi:10.1128/ AEM.70.11.6909-6913.2004)

- 36. Jones JL, Lüdeke CHM, Bowers JC, DeRosia-Banick K, Carey DH, Hastback W. 2014 Abundance of Vibrio cholerae, V. vulnificus, and V. parahaemolyticus in oysters (Crassostrea virginica) and clams (Mercenaria mercenaria) from Long Island Sound. Appl. Environ. Microbiol. 80, 7667-7672. (doi:10.1128/AEM. 02820-14)
- 37. Givens CE, Bowers JC, DePaola A, Hollibaugh JT, Jones JL. 2014 Occurrence and distribution of Vibrio vulnificus and Vibrio parahaemolyticus—potential roles for fish, oyster, sediment and water. Lett. Appl. *Microbiol.* **58**, 503 – 510. (doi:10.1111/lam.12226)
- 38. Froelich BA, Williams TC, Noble RT, Oliver JD. 2012 Apparent Loss of Vibrio vulnificus from North Carolina oysters coincides with a drought-induced increase in salinity. Appl. Environ. Microbiol. 78, 3885 – 3889. (doi:10.1128/AEM.07855-11)
- 39. Zimmerman AM, DePaola A, Bowers JC, Krantz JA, Nordstrom JL, Johnson CN, Grimes DJ, 2007 Variability of total and pathogenic Vibrio parahaemolyticus densities in Northern Gulf of Mexico water and oysters. Appl. Environ. Microbiol. **73**, 7589 – 7596. (doi:10.1128/AEM.01700-07)
- 40. Parveen S et al. 2008 Seasonal distribution of total and pathogenic Vibrio parahaemolyticus in Chesapeake Bay oysters and waters. Int. J. Food Microbiol. 128, 354-361. (doi:10.1016/j. ijfoodmicro.2008.09.019)
- 41. Moore JG, Ruple A, Ballenger-Bass K, Bell S, Pennington PL, Scott Gl. 2014 Snapshot of Vibrio parahaemolyticus densities in open and closed shellfish beds in Coastal South Carolina and Mississippi. Environ. Monit. Assess. 186, 7949 – 7960. (doi:10.1007/s10661-014-3979-z)
- 42. López-Hernández KM, Pardío-Sedas VT, Lizárraga-Partida L, Williams J, Martínez-Herrera D, Flores-Primo A, Uscanga-Serrano R, Rendón-Castro K. 2015 Environmental parameters influence on the dynamics of total and pathogenic Vibrio parahaemolyticus densities in Crassostrea virginica harvested from Mexico's Gulf coast. Mar. Pollut. *Bull.* **91**, 317–329. (doi:10.1016/j.marpolbul.2014.
- 43. Cruz CD, Hedderley D, Fletcher GC. 2015 Long-term study of Vibrio parahaemolyticus prevalence and distribution in New Zealand shellfish. Appl. Environ. Microbiol. 81, 2320-2327. (doi:10.1128/AEM. 04020-14)
- Kirs M, DePaola A, Fyfe R, Jones JL, Krantz J, Van Laanen A, Cotton D, Castle M. 2011 A survey of oysters (Crassostrea gigas) in New Zealand for Vibrio parahaemolyticus and Vibrio vulnificus. Int. J. Food *Microbiol.* **147**, 149–153. (doi:10.1016/j. ijfoodmicro.2011.03.012)
- 45. Schijven J, Bouwknegt M, de Roda Husman AM, Rutjes S, Sudre B, Suk JE, Semenza JC. 2013 A decision support tool to compare waterborne and foodborne infection and/or illness risks associated with climate change. Risk Anal. 33, 2154-2167. (doi:10.1111/risa.12077)
- 46. Smith BA, Ruthman T, Sparling E, Auld H, Comer N, Young I, Lammerding AM, Fazil A. 2015 A risk modeling framework to evaluate the impacts of

- climate change and adaptation on food and water safety. Food Res. Int. 68, 78-85. (doi:10.1016/j. foodres.2014.07.006)
- 47. Broberg CA, Calder TJ, Orth K. 2011 Vibrio parahaemolyticus cell biology and pathogenicity determinants. Microbes Infect. 13, 992-1001. (doi:10.1016/j.micinf.2011.06.013)
- 48. Nishibuchi M, Kaper JB. 1995 Thermostable direct hemolysin gene of Vibrio parahaemolyticus: a virulence gene acquired by a marine bacterium. Infect. Immun. 63, 2093-2099.
- 49. DePaola A, Hopkins LH, Peeler JT, Wentz B, McPhearson RM. 1990 Incidence of Vibrio parahaemolyticus in U.S. coastal waters and oysters. Appl. Environ. Microbiol. 56, 2299 - 2302.
- 50. Yu W-T, Jong K-J, Lin Y-R, Tsai S, Tey YH, Wong H. 2013 Prevalence of Vibrio parahaemolyticus in oyster and clam culturing environments in Taiwan. Int. J. Food Microbiol. 160, 185-192. (doi:10.1016/ j.ijfoodmicro.2012.11.002)
- 51. Han H et al. 2015 Temporal and spatial variation in the abundance of total and pathogenic Vibrio parahaemolyticus in shellfish in China. PLoS ONE 10, e0130302. (doi:10.1371/journal.pone.
- 52. Kim MS, Jeong HD. 2001 Development of 16S rRNA targeted PCR methods for the detection and differentiation of Vibrio vulnificus in marine environments. Aquaculture 193, 199-211. (doi:10. 1016/S0044-8486(00)00495-6)

- 53. Nilsson WB, Paranjype RN, DePaola A, Strom MS. 2003 Sequence polymorphism of the 16S rRNA gene of Vibrio vulnificus is a possible indicator of strain virulence. J. Clin. Microbiol. 41, 442-446. (doi:10.1128/JCM.41.1.442-446.2003)
- 54. Aznar R, Ludwig W, Amann RI, Schleifer KH. 1994 Sequence determination of rRNA genes of pathogenic Vibrio species and whole cell identification of Vibrio vulnificus with rRNA targeted oligonucleotide. Int. J. Syst. Bacteriol. 44, 330 – 337. (doi:10.1099/00207713-44-2-330)
- 55. Rosche TM, Yano Y, Oliver JD. 2005 A rapid and simple PCR analysis indicates there are two subgroups of Vibrio vulnificus which correlate with clinical and environmental isolation. Microbiol. Immunol. 49, 381-389. (doi:10.1111/j.1348-0421. 2005.tb03731.x)
- 56. Warner JM, Oliver JD. 1999 Randomly amplified polymorphic DNA analysis of clinical and environmental isolates of Vibrio vulnificus and other Vibrio species. Appl. Environ. Microbiol. 65, 526 - 534.
- 57. Jones JL, Lüdeke CHM, Bowers JC, DePaola A. 2013 Comparison of plating media for recovery of total and virulent genotypes of Vibrio vulnificus in U.S. market oysters. Int. J. Food Microbiol. 167, 322-327. (doi:10.1016/j. ijfoodmicro.2013.09.017)
- Lin M, Schwarz JR. 2003 Seasonal shifts in population structure of Vibrio vulnificus in an

- estuarine environment as revealed by partial 16S ribosomal DNA sequencing. FEMS Microbiol. Ecol. 45, 23-27. (doi:10.1016/s0168-6496(03)
- 59. Froelich BA, Noble RT. 2014 Factors affecting the uptake and retention of Vibrio vulnificus in oysters. Appl. Environ. Microbiol. 80, 7454-7459. (doi:10. 1128/AEM.02042-14)
- 60. Audemard C, Kator HI, Rhodes MW, Gallivan T, Erskine AJ, Leggett AT, Reece KS. 2011 High salinity relay as a postharvest processing strategy to reduce Vibrio vulnificus levels in Chesapeake Bay oysters (Crassostrea virginica). J. Food Prot. 74, 1902 – 1907. (doi:10.4315/0362-028X.JFP-11-152)
- 61. Motes ML, DePaola A. 1996 Offshore suspension relaying to reduce levels of Vibrio vulnificus in oysters (Crassostrea virginica). Appl. Environ. Microbiol. **62**, 3875-3877.
- 62. Baker-Austin C, Trinanes JA, Taylor NGH, Hartnell R, Siitonen A, Martinez-Urtaza J. 2013 Emerging Vibrio risk at high latitudes in response to ocean warming. Nat. Clim. Change 3, 73-77. (doi:10.1038/ nclimate1628)
- 63. Weirich C. 2015 North Carolina oyster aquaculture industry update.
- Baker-Austin C, Lemm E, Hartnell R, Lowther J, Onley R, Amaro C, Oliver JD, Lees D. 2012 pilF polymorphismbased real-time PCR to distinguish Vibrio vulnificus strains of human health relevance. Food Microbiol. 30, 17 – 23. (doi:10.1016/j.fm.2011.09.002)