

doses usually separated by 14 days. They provide ~60–85% protection for the first few months. Booster immunizations of WC-rBS are recommended after 2 years for individuals  $\geq 6$  years of age and after 6 months for children 2–5 years of age. For BivWC, which was developed more recently, no formal recommendation regarding booster immunizations exists. However, BivWC was associated with ~60% protection over 5 years among recipients of all ages in a study in Kolkata, India; the rate of protection among children  $\leq 5$  years of age approximated 40%. Models predict significant herd immunity when vaccination coverage rates exceed 50%. The killed vaccines have been safely administered among populations with high rates of HIV.

Oral live attenuated vaccines for *V. cholerae* are also in development. These strains have in common the fact that they lack the genes encoding cholera toxin. One such vaccine, CVD 103-HgR, was safe and immunogenic in phase 1 and 2 studies but afforded minimal protection in a large field trial in Indonesia. Other live attenuated vaccine candidate strains have been prepared from El Tor and O139 *V. cholerae* and have been tested in studies of volunteers. A possible advantage of live attenuated cholera vaccines is that they may induce protection after a single oral dose. Conjugate and subunit cholera vaccines are also being developed. Recognizing that it may be decades before safe water and adequate sanitation become a reality for those most at risk of cholera, the WHO has now recommended incorporation of cholera vaccination into comprehensive control strategies and has established an international stockpile of oral killed cholera vaccine to assist in outbreak responses. No cholera vaccine is commercially available in the United States.

### OTHER VIBRIO SPECIES



The genus *Vibrio* includes several human pathogens that do not cause cholera. Abundant in coastal waters throughout the world, noncholera vibrios can reach high concentrations in the tissues of filter-feeding mollusks. As a result, human infection commonly follows the ingestion of seawater or of raw or undercooked shellfish (Table 193-5). Most noncholera vibrios can be cultured on blood or MacConkey agar, which contains enough salt to support the growth of these halophilic species. In the microbiology laboratory, the species of noncholera vibrios are distinguished by standard biochemical tests. The most important of these organisms are *Vibrio parahaemolyticus* and *Vibrio vulnificus*.

The two major types of syndromes for which these species are responsible are gastrointestinal illness (due to *V. parahaemolyticus*, non-O1/O139 *V. cholerae*, *Vibrio mimicus*, *Vibrio fluvialis*, *Vibrio hollisae*, and *Vibrio furnissii*) and soft tissue infections (due to *V. vulnificus*, *Vibrio alginolyticus*, and *Vibrio damsela*). *V. vulnificus* is also a cause of primary sepsis in some compromised individuals.

### SPECIES ASSOCIATED PRIMARILY WITH GASTROINTESTINAL ILLNESS



***V. parahaemolyticus*** Widespread in marine environments, the halophilic *V. parahaemolyticus* causes food-borne enteritis worldwide. This species was originally implicated in enteritis

in Japan in 1953, accounting for 24% of reported cases in one study—a rate that presumably was due to the common practice of eating raw seafood in that country. In the United States, common-source outbreaks of diarrhea caused by this organism have been linked to the consumption of undercooked or improperly handled seafood or of other foods contaminated by seawater. Since the mid-1990s, the incidence of *V. parahaemolyticus* infections has increased in several countries, including the United States. Serotypes O3:K6, O4:K68, and O1:K-untypable, which are genetically related to one another, account in part for this increase. Serotypes O4:K12 and O4:KUT were initially unique to the Pacific Northwest but caused recent outbreaks in the eastern United States and Spain. The enteropathogenicity of *V. parahaemolyticus* is linked to its ability to cause hemolysis on Wagatsuma agar (i.e., the *Kanagawa phenomenon*). Although the mechanisms by which the organism causes diarrhea are not fully defined, the genome sequence of *V. parahaemolyticus* contains two type III secretion systems, which directly inject toxic bacterial proteins into host cells. The activity of one of these secretion systems is required for intestinal colonization and virulence in animal models. *V. parahaemolyticus* should be considered a possible etiologic agent in all cases of diarrhea that can be linked epidemiologically to seafood consumption or to the sea itself.

Infections with *V. parahaemolyticus* can result in two distinct gastrointestinal presentations. The more common of the two presentations (including nearly all cases in North America) is characterized by watery diarrhea, usually occurring in conjunction with abdominal cramps, nausea, and vomiting and accompanied in ~25% of cases by fever and chills. After an incubation period of 4 h to 4 days, symptoms develop and persist for a median of 3 days. Dysentery, the less common presentation, is characterized by severe abdominal cramps, nausea, vomiting, and bloody or mucoid stools. *V. parahaemolyticus* also causes rare cases of wound infection and otitis and very rare cases of sepsis.

Most cases of *V. parahaemolyticus*-associated gastrointestinal illness, regardless of the presentation, are self-limited. Fluid replacement should be stressed. The role of antimicrobials is uncertain, but they may be of benefit in moderate or severe disease. Doxycycline, fluoroquinolones, or macrolides are usually used. Deaths are extremely rare among immunocompetent individuals. Severe infections are associated with underlying diseases, including diabetes, preexisting liver disease, iron-overload states, or immunosuppression.

**Non-O1/O139 (Noncholera) *V. cholerae*** The heterogeneous non-O1/O139 *V. cholerae* organisms cannot be distinguished from *V. cholerae* O1 or O139 by routine biochemical tests but do not agglutinate in O1 or O139 antiserum. Non-O1/O139 strains have caused several well-studied food-borne outbreaks of gastroenteritis and have also been responsible for sporadic cases of otitis media, wound infection, and bacteremia; although gastroenteritis outbreaks can occur, non-O1/O139 *V. cholerae* strains do not cause epidemics of cholera. Like other vibrios, non-O1/O139 *V. cholerae* organisms are widely distributed in marine environments. In most instances, recognized cases in the United States have been associated with the consumption of raw oysters or with recent travel. The broad clinical spectrum of diarrheal illness caused by these

**TABLE 193-5** FEATURES OF SELECTED NONCHOLERA VIBRIOSES

Organism	Vehicle or Activity	Host at Risk	Syndrome
<i>Vibrio parahaemolyticus</i>	Shellfish, seawater	Normal	Gastroenteritis
	Seawater	Normal	Wound infection
Non-O1/O139 <i>Vibrio cholerae</i>	Shellfish, travel	Normal	Gastroenteritis
	Seawater	Normal	Wound infection, otitis media
<i>Vibrio vulnificus</i>	Shellfish	Immunosuppressed <sup>a</sup>	Sepsis, secondary cellulitis
	Seawater	Normal, immunosuppressed <sup>a</sup>	Wound infection, cellulitis
<i>Vibrio alginolyticus</i>	Seawater	Normal	Wound infection, cellulitis, otitis
	Seawater	Burned, other immunosuppressed	Sepsis

<sup>a</sup>Especially with liver disease or hemochromatosis.

Source: Table 161-3 in *Harrison's Principles of Internal Medicine*, 14th edition.