Safety and Immunogenicity of Live Oral Cholera Vaccine CVD 103-HgR in Children Aged 2–5 Years in the United States

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Abstract. In a phase 4, randomized, placebo-controlled, double-blind, multicenter study, to assess the safety and immunogenicity of live, attenuated cholera vaccine PXVX0200 in children aged 2–5 years in the United States, 172 volunteers were randomized 6:1 to receive a single dose of 1 × 10⁹ colony forming units (CFU) of PXVX0200 or placebo. Immunogenicity endpoints included serum vibriocidal antibody (SVA) levels on days 1, 11, and 29. Safety was assessed by comparing solicited signs and symptoms on days 1–8, unsolicited adverse events through day 29, and serious adverse events (SAEs) through day 181. The SVA seroconversion rates 10 days after immunization were 98.1% and 0% in vaccine and placebo recipients, respectively, and the vaccine seroconversion rate was non-inferior to the 93.5% rate seen in the bridging population of adults aged 18–45 years from a lot consistency study. Most reactogenicity was mild to moderate, and there were no study-related SAEs. PXVX0200 appears safe and immunogenic in children aged 2–5 years.

Vibrio cholerae remains a major cause of diarrhea in much of the world and in its most severe form (cholera gravis) can lead to severe dehydration and death, an outcome seen most commonly in children younger than 5 years.1 It is estimated that 1.3–4 million cases of cholera occur each year and result in up to 21,000–143,000 deaths worldwide.2 Cholera represents an ongoing risk to travelers to countries with both endemic and epidemic disease.

PXVX0200 is the research name for Center for Vaccine Development (CVD) 103-HgR, a live attenuated strain of V. cholerae O1, for the prevention of cholera diarrhea. It was licensed in the United States in 2016 under the trade name Vaxchora® (Emergent Travel Health Inc., Redwood City, CA) for use in individuals aged 18–64 years traveling to cholera-affected areas.3 The safety, efficacy, and immunogenicity of PXVX0200 have been established in adults in four randomized, double-blind, placebo-controlled, multicenter trials.4–7 These included a phase 3 cholera challenge study in which PXVX0200 demonstrated 90% efficacy at 10 days and 80% efficacy at 3 months versus placebo following ingestion of 1 × 10⁶ CFU of wild-type V. cholerae O1 El Tor Inaba strain N16961.3 Because previous studies in children in developing countries of CVD 103-HgR suggested that the immune response, as measured by serum vibriocidal antibody (SVA) seroconversion rates and geometric mean titers (GMTs), was lower than the response in adults, a pediatric trial of PXVX0200 in children was recently performed in the United States.8–10

In this study, CVD 103-HgR was shown to be safe, immunogenic, and well tolerated in children and adolescents aged 6–17 years.10 Here, we report for the first time the safety and immunogenicity results from cohort 3 of that study, that is, children aged 2–5 years.

This was a multicenter phase 4 randomized, double-blind, placebo-controlled trial designed to assess the immunogenicity, safety, palatability, and acceptability of PXVX0200 in children aged 2–17 years. Study methods have been previously described and were similar for cohort 3, except that palatability was assessed by the caregiver instead of the subject.10 The study was performed in children aged 2–5 years at eight U.S. sites from July 2017 through September 2019. For serum antibody assays, blood was collected from all participants on days 1, 11, and 29. For cohort 3, vaccine was prepared as previously described, except that 50 mL of buffer solution was used with each dose, rather than 100 mL. Placebo consisted of 50 mL of physiological (0.9%) saline. The primary endpoint was the proportion of participants achieving seroconversion, defined as a 4-fold or greater rise over baseline SVA, at day 11 after one dose of PXVX0200 in the immunogenicity evaluable population (IEP), which included the set of participants who received vaccination, consumed at least 80% of the study treatment, had evaluable results from both day 1 and day 11, and had no major protocol violations that would affect immunogenicity. The co-primary immunogenicity objectives were to demonstrate a minimum seroconversion rate of 70% and non-inferiority to the seroconversion rate in the bridging population of adults aged 18–45 years from the lot consistency study.6

A total of 187 participants aged 2–5 years were screened, 176 were randomized across the sites, and 155 completed the study through the day 181 visit (Figure 1). The demographic and baseline characteristics of the two groups were similar and, other than age, were also similar to cohorts 1 and 2. The day 11 SVA seroconversion rate in the IEP of cohort 3 was 98.1%, which was non-inferior to the 93.5% rate in the adult bridging population and was also greater than 70% (Table 1). Thus, both primary immunogenicity objectives were met. No additional seroconversions occurred between day 11 and day 29. The SVA GMTs of cohort 3 peaked at 4,852 on day 11 and decreased to 1,014 by day 29, whereas the geometric mean fold increase (GMFI) of serum vibriocidal antibodies peaked at 182 on day 11 and decreased to 38 by day 29. The incidence of vomiting was higher in the placebo group; otherwise there were no significant differences in the frequency and severity of solicited reactogenicity signs and symptoms between PXVX0200 and placebo recipients (Table 2). There was one serious adverse event (SAE), a hospitalization for an asthma exacerbation and pneumonia, in a placebo subject, and one “life-threatening” fever (> 40°C, per protocol definition) in a PXVX0200 subject, which were both considered not related to the study product.

Of the 172 cohort 3 subjects dosed, ≥ 80% of the dose was consumed by 82.7% and 84.6% of Vaxchora and placebo...
recipients, respectively. This was less than the 99.4% and 93.7% of vaccine recipients in the 12- to 17- and 6- to 11-year-old cohorts, respectively, who consumed ≥ 80% of the dose. 

Taste was ranked as very good, good, or neutral by the caregivers of 91 (62.3%) and 18 (69.2%) of Vaxchora and placebo recipients, respectively. Optional Pure Via Stevia sweetener was added to all, but one, of the doses; thus, assessment of its effect on palatability was not possible.

Cholera is a threat to children and adults who travel to endemic areas and can cause severe and fatal disease. A single dose vaccine that can provide rapid and effective immunity would be ideal. Here, we document the safety, immunogenicity, and tolerability of single-dose, live oral cholera vaccine PXVX0200 in children aged 2–5 years in a developed country. Serum vibriocidal antibody seroconversion was measured as the primary endpoint in this bridging study comparing rates in children with those in adults aged 18–45 years. In cohort 3, seroconversion occurred in 98.1% of vaccine recipients aged 2–5 years versus 0% in placebo recipients (P < 0.0001). These rates were non-inferior to the 93.5% seroconversion rate in the adult bridging population of the phase 3 lot consistency study, and because the lower limit of the 98.3% CI for seroconversion was greater than 70%, both primary objectives were met. Because SVA seroconversion was shown to be a strong correlate of protection against moderate to severe cholera diarrhea in the adult challenge study, it is possible that children aged 2–5 years vaccinated with PXVX0200 will be protected against cholera-induced diarrhea. In this study, GMT and GMFI at day 11 were lower than those seen in cohorts 1 and 2 and the adult bridging study, possibly reflecting immunologic or intestinal immaturity in the youngest cohort. However, because seroconversion rates were similar and SVA seroconversion is the best correlate of protection against moderate to severe diarrhea, it is hoped that because challenge studies cannot be performed in children, the efficacy of PXVX0200 in the youngest cohort would be similar to the efficacy in adults.

Previous studies of single-dose CVD 103-HgR oral cholera vaccine in developing countries demonstrated that SVA seroconversion rates were lower in children than those seen in adults, and higher dose formulations with $5 \times 10^9$ CFU were developed, with seroconversion seen in only 22–78% of
vaccine recipients at this dose.\textsuperscript{8,9,12,17} A large efficacy trial of CVD 103-HgR in Indonesia failed to demonstrate protection in children, although herd immunity may have contributed to the unexpected reduction of cholera cases in placebo recipients in that study.\textsuperscript{2,16} A mass vaccination campaign using single-dose CVD 103-HgR during a cholera outbreak in Micronesia, which included children aged 2–5 years, demonstrated a vaccine efficacy of 79.2%.\textsuperscript{21} The reduced immune responses in developing countries may be due to multiple factors, including malnutrition, small bowel overgrowth with intestinal mucosal damage (chronic environmental enteropathy), and preexisting immunity due to natural exposure to cholera infection.\textsuperscript{18,19} None of these conditions are common in the United States where children mount a more robust immune response to live oral cholera vaccine.

The study vaccine was well accepted by 82.7% of cohort 3 subjects versus 99.4% and 93.7%, respectively, of subjects in cohorts 1 and 2, and by 84.6% of placebo recipients in cohort 3, further demonstrating the impact of age on the willingness and ability to consume the oral solution.\textsuperscript{11} As noted in cohorts 1 and 2, there was no significant difference in palatability assessments between vaccine and placebo recipients in cohort 3. Previously in a trial of CVD 103-HgR in infants and toddlers in Chile, similar seroconversion rates occurred in participants who consumed partial doses of vaccine versus full doses.\textsuperscript{15} However, these results may not be applicable to the current study.

The data from this study demonstrate that PXVX0200 is well tolerated in children aged 2–5 years, with a safety profile similar to subjects aged 6–17 years and adults aged 18–45 years in the lot consistency study. Other than vomiting, there were no significant differences in the incidence of solicited reactogenicity, including diarrhea or fever, between vaccine and placebo recipients. One vaccine recipient developed a fever which was classified as "life-threatening" by predefined criteria but was not considered related to study product. There were no vaccine-related SAEs in any pediatric cohort.

Two other cholera vaccines are available outside of the United States, whole cell recombinant B subunit (Dukoral\textsuperscript{6}, SBL Vaccin AB, Stockholm, Sweden) and bivalent, killed whole cell vaccine (Shanchol\textsuperscript{6}, Shantha Biotechnics Limited, Andhra Pradesh, India, Euvichol\textsuperscript{6}, Eubiologics, Seoul, South Korea) administered as two or three doses depending on criteria but was not considered related to study product. There were no vaccine-related SAEs in any pediatric cohort.

### Table 1

Seroconversion, GMT, and fold increase for children and adults

| Adults (18–45 years) | Cohort 1 (12–17 years) | Cohort 2 (6–11 years) | Cohort 3 (2–5 years) |
|----------------------|------------------------|-----------------------|----------------------|
| PXVX0200             | Placebo                | PXVX0200              | Placebo              |
| (n = 2,688)          | (N = 334)              | (n = 157)             | (n = 23)             |
|                      |                        | (n = 139)             | (n = 24)             |
|                      |                        | (n = 103)             | (n = 25)             |
| N (analyzable)       | 2,687                  | 157                   | 139                  | 103                  |
| N (%) seroconverted  | 2,513 (93.5)           | 156 (99.4%)           | 136 (97.8%)          | 101 (98.1%)          |
| 98.3% CI             | [92.3%, 94.6%]         | [95.4%, 99.9%]        | [92.5%, 99.4%]       | [91.5%, 99.6%]       |
| Difference (cohort minus adults) | – | 5.8% | – | 4.3% | – | 4.5% | – | 4.5% | – | 4.5% | – | 4.5% | – | 4.5% |
| 96.7% CI on difference | – | [2.4%, 7.1%] | – | [–0.3%, 6.2%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] |
| GMT                  | –                      | –                     | –                     | –                     |
| –                    | 0.0095                 | 0.0455                | 0.0628                | 0.0628                |

#### Seroconversion

**Day 11 visit**

- N (analyzable): 2,687
- N (~) seroconverted: 2,513 (93.5%)
- 98.3% CI: [92.3%, 94.6%]
- Difference (cohort minus adults): –
- 96.7% CI on difference: –
- GMT: 2.687
- P-value: 0.0095

**Day 1 visit**

- N (analyzable): 2,688
- N (~) seroconverted: 2,688
- 95% CI: [90.67, 95.31]
- Minimum, maximum: 20,10,240

**Day 29 visit**

- N (analyzable): 2,680
- GMT: 6,170
- Minimum, maximum: 640,10,960

**Geometric mean fold increase**

**Day 11 visit**

- N (analyzable): 2,687
- Mean fold increase: 2.72
- Maximum: 6,170

**Day 29 visit**

- N (analyzable): 2,680
- Mean fold increase: 4.3
- Minimum, maximum: 640,10,960

\* GMT is geometric mean titer.

| P-value | 0.0095

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**TABLE 1**

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| 98.3% CI             | [92.3%, 94.6%]         | [95.4%, 99.9%]        | [92.5%, 99.4%]       | [91.5%, 99.6%]       |
| Difference (cohort minus adults) | – | 5.8% | – | 4.3% | – | 4.5% | – | 4.5% | – | 4.5% | – | 4.5% |
| 96.7% CI on difference | – | [2.4%, 7.1%] | – | [–0.3%, 6.2%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] | – | [–1.1%, 6.4%] |
| GMT                  | –                      | –                     | –                     | –                     |
| –                    | 0.0095                 | 0.0455                | 0.0628                | 0.0628                |

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- GMT: 2.687
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- N (analyzable): 2,680
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\* GMT is geometric mean titer.

\* P < 0.0001; P-values are based on t-statistics assuming normal distribution of the log titer.

\* Data for adults for PXVX0200 and placebo from the immune substudy population.
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