

1 Transient intestinal colonization by a live-attenuated oral cholera vaccine induces  
2 protective immune responses in streptomycin-treated mice

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15 **Abstract**

16 Current mouse models for evaluating the efficacy of live oral cholera vaccines  
17 (OCVs) have important limitations. Conventionally raised adult mice are resistant to  
18 intestinal colonization by *Vibrio cholerae*, but germ-free mice can be colonized and have  
19 been used to study OCV immunogenicity. However, germ free animals have impaired  
20 immune systems and intestinal physiology; also, live OCVs colonize germ free mice for  
21 many months, which does not mimic the clearance kinetics of live OCVs in humans.  
22 Here, we leverage antibiotic-treated, conventionally raised adult mice to study the  
23 effects of transient intestinal colonization by a live OCV *V. cholerae* strain. In a single  
24 dose vaccination regimen, we found that HaitiV, a live-attenuated OCV candidate, was  
25 cleared by streptomycin treated adult mice within a week after oral inoculation. This  
26 transient colonization elicited far stronger adaptive immune correlates of protection  
27 against cholera than did inactivated whole-cell HaitiV. Infant mice from HaitiV  
28 vaccinated dams were also significantly protected from choleric disease than pups from  
29 inactivated-HaitiV dams. Our findings establish the benefits of antibiotic treated mice for  
30 live OCV studies as well as its limitations and underscore the immunogenicity of HaitiV.

31

32 **Importance**

33 Oral cholera vaccines (OCVs) are being deployed to combat cholera but current  
34 killed OCVs require multiple doses and show little efficacy in young children. Live OCVs  
35 have the potential to overcome these limitations but small animal models for testing  
36 OCVs have shortcomings. We used an antibiotic treatment protocol for conventional  
37 adult mice to study the effects of short-term colonization by a single dose of HaitiV, a  
38 live OCV candidate. Vaccinated mice developed vibriocidal antibodies against *V.*  
39 *cholerae* and delivered pups that were resistant to cholera, whereas mice vaccinated  
40 with inactivated HaitiV did not. These findings demonstrate HaitiV's immunogenicity and  
41 suggest that this antibiotic treatment protocol will be useful for evaluating the efficacy of  
42 live OCVs.

43 **Introduction**

44 *Vibrio cholerae* is the cause of cholera and, following ingestion of water or food  
45 contaminated with this Gram-negative rod, humans can develop the severe and  
46 sometimes lethal dehydrating diarrhea that characterizes cholera. Cholera remains a  
47 major threat to global public health, with approximately 2.9 million cases and 95,000  
48 deaths reported annually (1). Serologic classification of *V. cholerae* is based on the  
49 structure and chemistry of the abundant LPS O-antigen and the O1 serogroup of *V.*  
50 *cholerae* has given rise to all cholera pandemics. The O1 serogroup is further  
51 subdivided into Ogawa and Inaba serotypes that differ by the presence or absence of  
52 methylation of the terminal perosamine on their respective O-antigens (2). Current  
53 pandemic cholera is predominantly caused by an O1 'variant' El Tor biotype strain, such  
54 as the strain responsible for the Haitian epidemic in 2010 (3).

55 Prior exposure to *V. cholerae* can elicit long-lived protective O-antigen-specific  
56 responses against *V. cholerae* (4, 5), suggesting that vaccination has the potential to  
57 elicit protective immunity if vaccines can safely mimic elements of natural infection. As  
58 such, several vaccination strategies for cholera have been developed over the decades.  
59 Among these, oral cholera vaccines (OCV) are attractive options as they stimulate  
60 immunity at the intestinal mucosal surface, the site of infection, and because of their  
61 ease of administration. Killed whole-cell OCVs, such as Shancol, are being increasingly  
62 adopted as frontline public health tools both in endemic regions (6) as well as to limit  
63 outbreaks (7).Live-attenuated OCVs have also been developed and have theoretical  
64 advantages over killed OCVs including *in vivo* replication which enables continuous  
65 presentation of *in vivo*-induced antigens at the intestinal mucosal surface (8). In contrast

66 to killed OCVs, live vaccines will likely offer single dose efficacy, a particularly important  
67 feature for reactive vaccination campaigns during epidemics. Furthermore, live OCVs  
68 appear to be more effective in children less than 5 years of age (9, 10), a group that is  
69 highly susceptible to death from cholera and that is not adequately protected by killed  
70 OCVs. In volunteer studies, these vaccines have shown great promise (11, 12), but  
71 none are approved for use in cholera endemic regions.

72 The development of both inactivated and live OCVs has been hampered by the  
73 lack of a small animal model that closely recapitulates cholera pathogenesis in the  
74 setting of normal immune reactivity. *V. cholerae* readily colonizes the intestines of infant  
75 mice and infant rabbits, reviewed in (13), where cholera-like disease can be observed,  
76 but these models lack mature immune systems. Conventionally raised adult mice  
77 cannot be orally colonized by *V. cholerae*, likely due to their resident gut microbiota  
78 (14). Germ free (GF) adult mice, which lack a microbiota, can be colonized by *V.*  
79 *cholerae* and have been used to profile OCV immunogenicity, but immune and intestinal  
80 physiological development is aberrant in these animals (15). Nonetheless, vaccinated  
81 GF mice develop immune correlates of clinical protection against toxigenic *V. cholerae*,  
82 including circulating vibriocidal antibodies and antigen specific antibody responses (14,  
83 16, 17). Another limitation of the GF adult mouse model is that following oral  
84 administration of a live OCV, they remain consistently colonized by *V. cholerae* for  
85 periods exceeding 3 months (14, 17), making single-dose live OCV regimens difficult to  
86 interpret; moreover, the prolonged colonization in this model precludes challenging  
87 vaccinated animals with virulent *V. cholerae*. The consequence and significance of long-  
88 term monocolonization with a live OCV in GF mice also remains unknown.

89        Many studies have shown that oral administration of broad-spectrum antibiotics  
90    to mice depletes the gut microbiota and enables intestinal colonization by diverse  
91    bacteria (18). This is beneficial as antibiotic treated mice are conventionally raised and  
92    do not display the immunological and developmental defects that characterize GF mice.  
93    Antibiotic treatments can enable wild-type *V. cholerae* intestinal colonization for similar  
94    durations, typically 5-7 days, that humans are colonized by live OCVs (19, 20). We  
95    sought to leverage this model to profile HaitiV, a live-attenuated OCV derived from  
96    HaitiWT, a virulent *V. cholerae* O1 Ogawa clinical strain isolated during the Haitian  
97    cholera outbreak (21). HaitiV is non-toxigenic and highly engineered for biosafety and in  
98    infant rabbits HaitiV provides unprecedented rapid protection against virulent *V.*  
99    *cholerae* within 24 hours of administration (21); furthermore, we showed that this  
100   vaccine is immunogenic in GF mice (17).

101        Here, we adopted a streptomycin treated adult mouse model of *V. cholerae* (22)  
102   to profile HaitiV's immunogenicity. Furthermore, we modified this model to assess the  
103   protective efficacy of immune responses in immunized female mice, by challenging their  
104   pups with HaitiWT. Our findings demonstrate the ease and utility of this approach for  
105   studies of live-attenuated OCVs.

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111 **Results**

112 **Vaccine inoculation protocol and colonization kinetics.**

113 We modified the protocol presented in Bueno et al (22) and orally treated 4-  
114 week-old C57BL/6 female mice with streptomycin (Sm) to deplete their intestinal  
115 microbiota to enable a longitudinal study of HaitiV intestinal colonization and  
116 immunogenicity (Fig. 1A). Three days after initiating Sm treatment, mice were orally  
117 gavaged with either a single dose of  $10^9$  CFU of HaitiV, an equivalent dose of formalin-  
118 inactivated HaitiV (FI-HaitiV) to mimic oral vaccination with an inactivated OCV, or  
119 sodium bicarbonate as a vehicle control. There were no apparent untoward effects of  
120 any of these regimens and over the course of the experiment, all mice gained weight  
121 (Fig. 1B).

122 Plating of fresh fecal pellets (FP) from all animals inoculated with HaitiV revealed  
123  $\sim 10^8$  CFU/g FP for 5-7 days post inoculation (dpi), suggesting that initially the vaccine  
124 robustly colonized the intestines of Sm treated adult mice. However, HaitiV was no  
125 longer detectable in FPs by 8-12 dpi (Fig. 1C, Fig. S1B), indicating clearance of the  
126 vaccine strain. No CFUs of HaitiV were recovered from Sm treated mice that had been  
127 orally inoculated with FI-HaitiV or buffer control. The clearance kinetics of HaitiV from  
128 these mice resembles that previously charted by Nygren et al (19) for wild type cholera  
129 toxin-producing *V. cholerae* isolates, suggesting that cholera toxin does not play a  
130 substantial role in intestinal colonization in this model.

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133 **Transient HaitiV colonization elicits antibodies targeting multiple *V. cholerae***  
134 **serotypes.**

135

136 Sera from all mice were individually assayed to quantify the circulating vibriocidal  
137 antibody titers (VATs) targeting Ogawa and Inaba *V. cholerae* strains. By 7dpi most  
138 mice did not have detectable VATs but by 14dpi when HaitiV was no longer detectable  
139 in FPs, most (3/5 mice) seroconverted and developed high circulating VATs against  
140 both serotyped matched (Ogawa) and serotype mismatched (Inaba) *V. cholerae* (Fig.  
141 2A, Fig. S1C, Fig. S1D), though highest geometric mean VATs were directed against  
142 serotype matched (Ogawa) isolates (Fig. 2A, Fig. S1C). Only one mouse in the FI-  
143 HaitiV group developed circulating VATs against Ogawa serotype *V. cholerae*, and no  
144 mice in this group developed vibriocidal antibodies against Inaba *V. cholerae* in this  
145 group (Fig. 2B). None of the Sm treated mice that received sodium bicarbonate  
146 developed detectable vibriocidal antibodies. Thus, transient colonization by HaitiV is  
147 sufficient to elicit the generation of *V. cholerae* specific circulating markers of immunity.

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149 **Streptomycin treated adult mice resist recolonization by *V. cholerae*.**

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151 Administration of live-OCVs to GF mice leads to long term intestinal colonization  
152 (14, 17), precluding the possibility of challenge studies with wild-type *V. cholerae*. Since  
153 the Sm-treated mice cleared HaitiV, we investigated whether animals immunized with  
154 this OCV could be successfully challenged with HaitiWT. At 28dpi, when all mice had  
155 stopped shedding HaitiV, they (live HaitiV, FI-HaitiV, and vehicle control) were orally

156 treated with sulfamethoxazole and trimethoprim (SXT) to reduce the intestinal  
157 microbiota (Fig. 3A); unlike HaitiV, HaitiWT is resistant to SXT. All three groups of mice  
158 from above and another group, specific pathogen free (SPF) mice that had not been  
159 Sm-treated or exposed to HaitiV was also included in this experiment, to test whether  
160 SXT treatments modify susceptibility to HaitiWT colonization. All 4 groups of mice were  
161 then orally gavaged with  $10^9$  CFU of HaitiWT, and their FPs monitored for HaitiWT  
162 colonization. Unexpectedly, the three groups of mice (HaitiV, FI-HaitiV, and sodium  
163 bicarbonate) that had been previously treated with Sm failed to be colonized by HaitiWT  
164 (Fig. 3B). In contrast, the SXT-treated SPF mice were robustly colonized by HaitiWT  
165 ( $\sim 10^8$  CFU/g FP) (Fig. 3B), indicating that although SXT treatment does susceptibilize  
166 mice to *V. cholerae* colonization, the prior Sm treatment of the vaccinated animals  
167 rendered them resistant to recolonization with *V. cholerae*.

168         Although the FPs of the previously Sm treated and HaitiWT challenged mice  
169 lacked detectable HaitiWT on the Sm agar plates used to detect *V. cholerae*, these  
170 plates contained high numbers of colonies of non-*V. cholerae* bacteria. 16s rRNA  
171 sequencing of Sm resistant small colonies showed that these colonies corresponded to  
172 either *Escherichia coli* or SXT resistant *Lactobacillus murinus*, which together were  
173 present at  $\sim 10^8$  CFU/g FP. These observations suggest that a bloom of Sm-resistant  
174 organisms and other changes in the gut microbiota associated with prior oral Sm  
175 treatment render mice resistant to *V. cholerae* colonization.

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179 **Offspring of vaccinated dams are protected from virulent *V. cholerae*.**

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181 As challenge studies could not be performed in Sm treated adult mice, we turned  
182 to the suckling mouse model of cholera to assess the protective efficacy of the immune  
183 response that was elicited by the transient HaitiV intestinal colonization. We recently  
184 found that the survival of suckling mice in this lethal challenge model can be used to  
185 gauge OCV efficacy (17). As such, animals from HaitiV, FI-HaitiV, and sodium  
186 bicarbonate groups were mated with SPF males and their neonatal pups infected with a  
187 lethal dose of HaitiWT (Fig. 4A).

188 Pups from HaitiV immunized dams survived significantly longer (median 30hpi)  
189 than pups from dams treated with FI-HaitiV or sodium bicarbonate control (median  
190 ~25hpi) (Fig. 4B). Despite the increased survival in the offspring of HaitiV immunized  
191 dams, at time of death, there were no differences in the burden of HaitiWT CFU  
192 recovered from the small intestines of the 3 groups (Fig. 4C). Thus, oral immunization  
193 and transient colonization of Sm-treated mice with HaitiV induces immune responses  
194 that significantly protect against choleric disease in mice.

195

196 **Discussion**

197 The wide availability of diverse mutant mice and reagents for their study make  
198 mice a preferred model mammal for studies of human disease and therapeutics.  
199 However, adult mice are not susceptible to intestinal colonization with *V. cholerae*,  
200 confounding evaluation of OCVs. Here, we found that Sm-treated adult mice can be  
201 used to investigate the immunogenicity as well as the protective efficacy of live-

202 attenuated OCVs. Sm-treated adult mice orally inoculated with HaitiV were colonized by  
203 this live-attenuated vaccine for 5-7days and this transient colonization was sufficient to  
204 elicit vibriocidal antibodies against both *V. cholerae* serotypes. Furthermore, pups born  
205 to HaitiV immunized Sm-treated mice exhibited prolonged survival following lethal  
206 challenge with HaitiWT compared to pups born to dams immunized with FI-HaitiV or  
207 dams treated with vehicle control. Together, these findings suggest that this model  
208 should be valuable for further studies of the efficacy and protective mechanisms of live  
209 OCVs.

210 Antibiotic treatment of adult mice limits the roles that the intestinal microbiota  
211 plays in inhibiting live OCV colonization. Unexpectedly, we found that oral Sm treatment  
212 enables the expansion of Sm and SXT-resistant microbes that also inhibit *V. cholerae*  
213 colonization (Fig. 3). Detailed analyses of the composition of the microbiota that are  
214 initially killed by oral Sm administration, and those that bloom after the antibiotic is  
215 withdrawn would offer valuable clues into which bacteria inhibit *V. cholerae* colonization.  
216 Though SXT treatment did not enable HaitiWT colonization of previously Sm treated  
217 animals (Fig. 3), it is possible that different antibiotic cocktails would facilitate re-  
218 challenge studies; for example, clindamycin was recently found to enable *V. cholerae*  
219 intestinal colonization of adult mice (23). Establishing conditions for re-challenge studies  
220 would simplify the model, bypassing the requirement for challenge studies in the  
221 offspring of immunized mice; however, only intestinal colonization resistance could be  
222 assayed in adult re-challenge studies since unlike suckling mice, adult mice are  
223 resistant to choleric diarrhea.

224            Although pups from HaitiV immunized Sm-treated dams survived longer than  
225    control animals, there was no difference in the CFU burden of HaitiWT in pups from  
226    immunized or control dams at the time they became moribund (Fig. 4). It is possible that  
227    vaccination partially delayed the replication of HaitiWT or, in addition, vaccination may  
228    elicit antibodies that antagonize toxic factors, such as cholera toxin, that promote  
229    disease, but do not directly modulate bacterial colonization. HaitiV ectopically expresses  
230    the GM1-binding B subunit of cholera toxin and immune responses to this non-toxic  
231    component of cholera toxin are linked to short term protection against cholera and  
232    enterotoxigenic *E. coli* (24).

233            In contrast to GF mice, which were colonized HaitiV for many months in our  
234    previous study (17), Sm-treated mice were only colonized by HaitiV for several days  
235    (Fig. 1). This is similar to the duration that live OCVs colonize the human intestine (20,  
236    25, 26) meaning that Sm treated mice may provide a more physiologically relevant  
237    model to gauge OCV immunogenicity. However, it is important to note that although  
238    Sm-treated mice display HaitiV clearance kinetics that mimic human OCV clearance,  
239    mice are not natural hosts for *V. cholerae*. *V. cholerae* intestinal colonization in  
240    susceptible adult mice occurs primarily in the colon and does not rely on TCP, a critical  
241    factor for colonization of the small intestine in humans as well in infant mice and rabbits  
242    (13, 19). Nonetheless, since adaptive protective immune responses are stimulated by  
243    HaitiV in both Sm treated and GF mice, these models have value for testing vaccine  
244    immunogenicity.

245            We adopted the strategy that Sit et al (17) used for studying the protective  
246    efficacy of OCVs in immunized GF mice and coupled the suckling mouse model of

247 cholera with the Sm-treated adult model of OCV immunogenicity. However, in contrast  
248 to the GF model, when the offspring of HaitiV immunized Sm-treated mice were  
249 inoculated with HaitiWT, their dams had undetectable HaitiV CFUs in their feces and  
250 VATs in their sera (Fig. 1, Fig. 4); these conditions more closely mimic those that will be  
251 present when immunized humans are exposed to *V. cholerae*. Despite the absence of  
252 circulating VATs, which are known to be relatively short-lived (27) , the offspring of  
253 HaitiV-immunized dams exhibited significantly delayed death due to cholera-like  
254 disease, but all succumbed (Fig. 4). In contrast, the offspring of HaitiV-immunized GF  
255 dams were completely protected from challenge with HaitiWT (17), demonstrating the  
256 increased potency of HaitiV immunization in GF mice. The greater effectiveness of  
257 HaitiV vaccination in GF mice is likely attributable to the constant stimulation of the  
258 intestinal mucosa by HaitiV in the persistently mono-colonized GF animals and could be  
259 consistent with multiple-dose live OCV regimens. Thus, even though GF mice have  
260 immune defects, they may overestimate the potency of HaitiV and other live OCVs.

261           Killed whole cell vaccines like Shancol have ~50% efficacy in single dose trials  
262 (29). Notably, in marked contrast to HaitiV, we found in the Sm-treated mice model that  
263 a single dose of FI-HaitiV, which is similar to current killed whole cell vaccines, did not  
264 elicit protective immunity, consistent with the idea that live-attenuated HaitiV is far more  
265 immunogenic than killed vaccines, at least when administered as a single dose.

266           In summary, we have described an adult mouse model for the investigation of the  
267 protective efficacy of live OCVs. Using this model, we found that a single oral dose of  
268 HaitiV and transient colonization by the vaccine can elicit vibriocidal antibody titers and  
269 protective immune responses. Future work should enable refinement of this model to

270 allow re-challenge of immunized animals and further characterization of these protective  
271 immune responses. Furthermore, given the availability of genetically defined mutant  
272 C57BL/6 mice, the model described here should be a valuable approach for unraveling  
273 the molecular determinants of live vaccine-mediated protection against cholera and  
274 other mucosal pathogens.

275

## 276 **Materials and Methods**

277

### 278 **Bacterial Strains and culture conditions**

279 *Vibrio cholerae* strains were grown in Luria-Bertani (LB) broth supplemented with  
280 relevant antibiotics: streptomycin (Sm) at 200 $\mu$ g/ml and sulfamethoxazole trimethoprim  
281 (SXT) at 80 $\mu$ g/mL and 16 $\mu$ g/mL respectively at 37°C with continuous shaking at  
282 220rpm. For LB agar plates 1.5% agar was used and was supplemented with 5-bromo-  
283 4-chloro-3-indolyl- $\beta$ -d-galactopyranoside (X-Gal) at 60 $\mu$ g/mL. Bacteria were kept as -  
284 80°C stocks in LB with 40% glycerol.

### 285 **Oral immunization scheme**

286 4-week-old C57BL/6 female mice were purchased (Charles River) and housed in a BL-2  
287 facility under conventional rearing conditions for the duration of the studies. On day 0,  
288 all mice were briefly anesthetized with isoflurane and orally gavaged with 20mg of  
289 streptomycin in 100 $\mu$ L of sterilized water. Mice were then provided drinking water  
290 supplemented with 5mg/mL Sm for 72 hours (22). After this, mice were then gavaged  
291 with 10<sup>9</sup> CFU of an overnight culture of either HaitiV, or HaitiV inactivated by 10%  
292 formalin (FI-HaitiV) for 15 minutes, and then resuspended in 2.5% Na<sub>2</sub>CO<sub>3</sub>. Control

293 mice were also gavaged with 100uL of 2.5% Na<sub>2</sub>CO<sub>3</sub> alone. Mice were then provided  
294 drinking water supplemented with 200 $\mu$ g/ml Sm for 14 days before being returned to  
295 unsupplemented water. All mice were weighed weekly and blood samples were  
296 retrieved from each mouse by tail vein incision at each weighing. Blood samples were  
297 clotted for 45 minutes at room temperature, centrifuged at 20000xg for 10 minutes, and  
298 the serum stored at -20°C for future analysis.

299 Fresh fecal pellets were collected daily from each mouse, weighed, and plated in serial  
300 dilutions on LB+Sm+X-Gal agar to determine the colonization of each mouse by HaitiV.  
301 HaitiV colonizes are white due to a disrupted *lacZ*. The limit of detection of this assay  
302 represents the lowest CFU count that could be detected for a fecal pellet of that weight.

303 **Quantification of vibriocidal antibody titers**

304 Circulating titers of vibriocidal antibodies were quantified by determining the minimal  
305 serum dilution required to lyse PIC158 (Ogawa) or PIC018 (Inaba) *V. cholerae* as  
306 described previously (30, 31) with minor modifications. Briefly, serial dilutions of serum  
307 were incubated with guinea-pig complement (Sigma) and the target strain, and then  
308 allowed to grow in BHI media. Reported titers are the dilution of serum that caused  
309 more than 50% reduction in target strain optical density when compared to normal  
310 saline control wells. A mouse monoclonal antibody 432A.1G8.G1.H12 targeting *V.*  
311 *cholerae* O1 OSP was a positive control for the assay. The limit of detection of this  
312 assay represents the lowest serum dilution at which no inhibition of growth could be  
313 detected.

314

315 **Colonization of immunized mice with toxigenic *V. cholerae***

316 Mice from all three groups (HaitiV, FI-HaitiV, and vehicle control) as well as 6 week old  
317 female SPF mice (Charles River) were anesthetized with isoflurane and gavaged with  
318 16mg of sulfamethoxazole and 3.2mg trimethoprim (SXT) in 100uL of water and  
319 provided drinking water supplemented with 4mg/mL SX and 0.8mg/mL T for 72 hours.  
320 All mice were then gavaged with 10<sup>9</sup> CFU of an overnight culture of HaitiWT and  
321 switched to 0.16mg/mL SX and 0.032mg/mL T in their drinking water. Fecal pellets were  
322 collected daily and plated on both SXT agar and Sm agar. Non-*V. cholerae* colonies  
323 were isolated by streaking on fresh Sm plates, and 16s V1/V2 DNA sequences were  
324 amplified by colony PCR using primers F341 (TCG TCG GCA GCG TCA GAT GTG  
325 TAT AAG AGA CAG CCT ACG GGN GGC WGC AG) and R805 (GTC TCG TGG GCT  
326 CGG AGA TGT GTA TAA GAG ACA GGA CTA CHV GGG TAT CTA ATC C) and  
327 sequenced. Identification of the bacteria was performed using BLAST (NCBI)

328 **Infant mouse survival assay**

329 The infant mouse survival assay was adapted from a previous report (17). Female mice  
330 were mated with age matched SPF male mice (Charles River) and singly housed at E18  
331 for delivery. At the third day of life (P3) pups were orally gavaged with 10<sup>7</sup> CFU of  
332 HaitiWT and returned to their dams. Pups were monitored every 6 hours until the first  
333 signs, typically including diarrhea and dehydration, were evident. At this point,  
334 monitoring was performed every 30 minutes until pups were moribund. Pups were then  
335 removed from the nest, euthanized with isoflurane, and decapitated and dissected. The  
336 small intestines of each pup were excised, homogenized, and plated on LB agar with  
337 Sm and X-Gal.

338 **Statistical analysis**

339 All statistical analyses were performed with Prism 8 (Graphpad). Infant mouse survival  
340 curves were analyzed with a log rank (Mantel Cox) test and CFU data were analyzed  
341 with a Mann Whitney U test.

342 **Animal use statement**

343 All experiments in this study were performed was approved by the Brigham and  
344 Women's Hospital IACUC (protocol 2016N000416) and in compliance with the NIH  
345 Guide for Use and Care of Laboratory animals.

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490 **Figure legends:**

491 **Figure 1. Transient intestinal colonization of HaitiV in Sm-treated adult mice**  
492 **following treatment with streptomycin.** A) Schematic of streptomycin treatment and  
493 single oral inoculation with HaitiV. Black arrowheads indicate peroral treatment with  
494 either streptomycin in sodium bicarbonate, or oral gavage with bacteria. B) Bodyweight  
495 of all mice over the course of this study. C) Fecal shedding of HaitiV from mice  
496 inoculated with HaitiV; open symbols depict CFU levels below the limit of detection.

497

498 **Figure 2. Serum vibriocidal antibody titers in Sm-treated mice immunized with**  
499 **HaitiV.** Circles indicate the lowest dilutions at which specific vibriocidal activity was  
500 detected, and the height of the bars represent geometric mean titers in each group.  
501 Ogawa *V. cholerae* PIC158 was used to measure Ogawa serotype-specific antibodies  
502 (black) and Inaba *V. cholerae* PIC018 was used to measure Inaba serotype-specific  
503 antibodies (red). Titers below the limit of detection are indicated by open symbols.

504

505 **Figure 3. SXT treatment and re-colonization of Sm-treated adult mice by *V.***  
506 ***cholerae.*** A) All mice were treated with sulfamethoxazole and trimethoprim (SXT) and  
507 dosed with  $10^9$  CFU of HaitiWT on day 0. B) Fecal shedding of HaitiWT from all mice;  
508 open symbols represent fecal pellets from which no CFUs of HaitiWT could be detected.  
509 An asterisk denotes differences with a P value of  $<0.05$  as determined by a Mann  
510 Whitney U test.

511

512 **Figure 4. Transient colonization by HaitiV is protective in an infant mouse model**

513 **of cholera. A)** Survival curves of pups born to dams inoculated with HaitiV (n=15),

514 formalin inactivated-HaitiV (n=24) or vehicle buffer (n=16) after HaitiWT challenge.

515 Differences in the survival curves were determined by a log rank (Mantel Cox) test

516 (p=0.0013 for HaitiV vs FI-HaitiV). **B)** HaitiWT CFUs recovered from the small intestines

517 of the neonatal mice at the time they were moribund. Differences in CFU burdens were

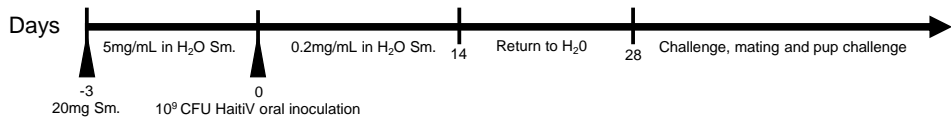
518 determined by a Mann Whitney U test.

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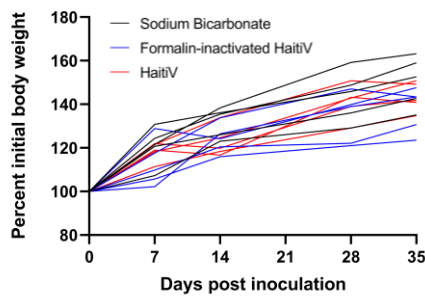
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Figure 1

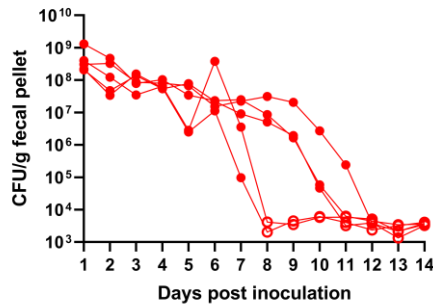
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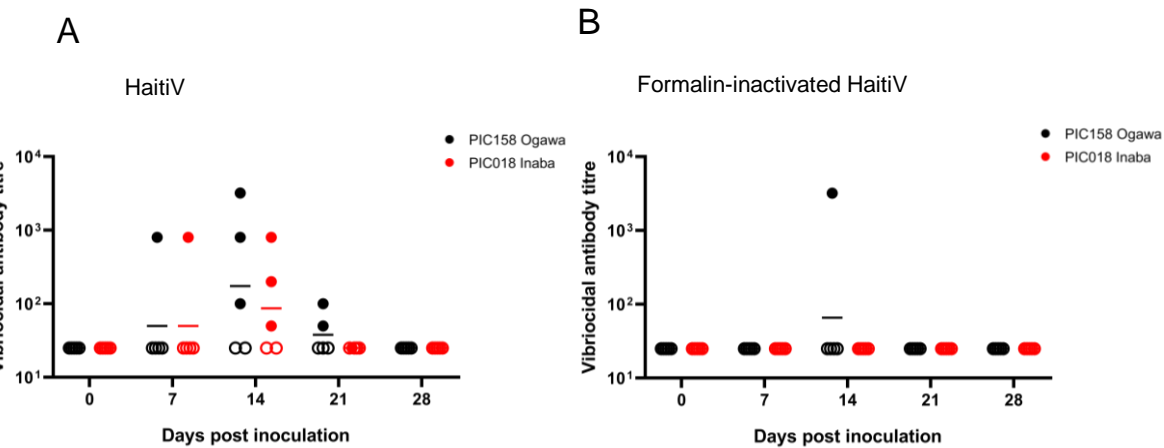


C



**Figure 1. Transient intestinal colonization of HaitiV in Sm-treated adult mice following treatment with streptomycin.** A) Schematic of streptomycin treatment and single oral inoculation with HaitiV. Black arrowheads indicate peroral treatment with either streptomycin in sodium bicarbonate, or oral gavage with bacteria. B) Bodyweight of all mice over the course of this study. C) Fecal shedding of HaitiV from mice inoculated with HaitiV; open symbols depict CFU levels below the limit of detection.

Figure 2

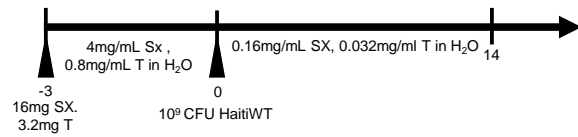


**Figure 2. Serum vibriocidal antibody titers in Sm-treated mice immunized with HaitiV.**

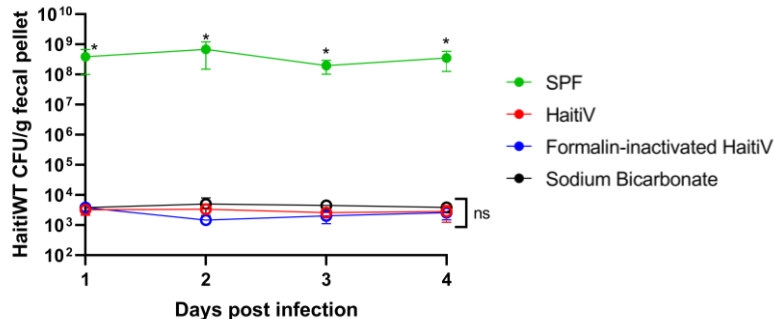
Circles indicate the lowest dilutions at which specific vibriocidal activity was detected, and the height of the bars represent geometric mean titers in each group. Ogawa *V. cholerae* PIC158 was used to measure Ogawa serotype-specific antibodies (black) and Inaba *V. cholerae* PIC018 was used to measure Inaba serotype-specific antibodies (red). Titers below the limit of detection are indicated by open symbols.

Figure 3

A

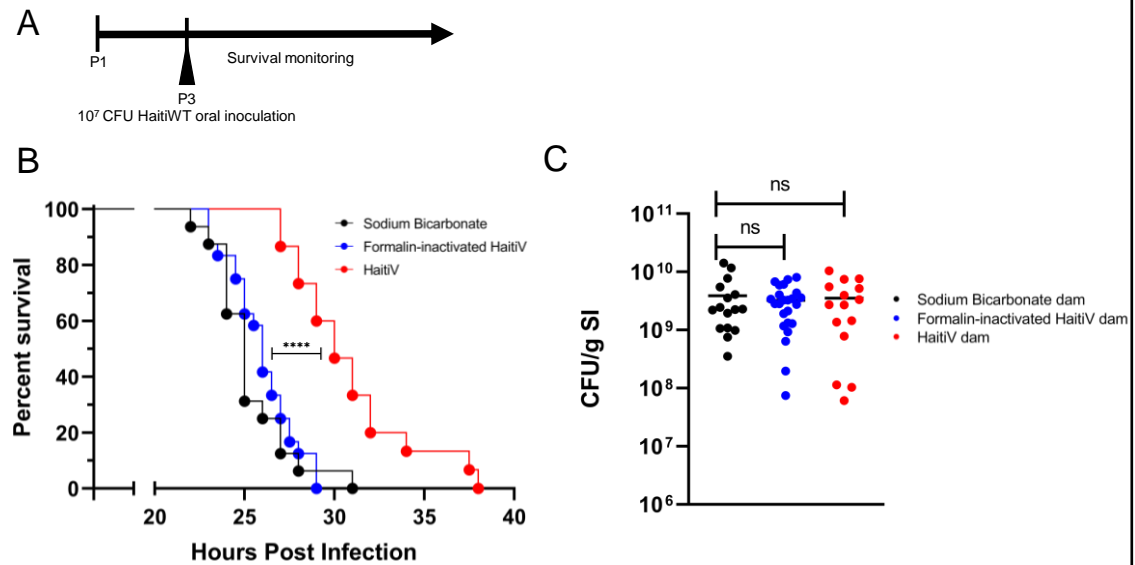


B



**Figure 3. SXT treatment and re-colonization of Sm-treated adult mice by *V. cholerae*.** A) All mice were treated with sulfamethoxazole and trimethoprim (SXT) and dosed with 10<sup>9</sup> CFU of HaitiWT on day 0. B) Fecal shedding of HaitiWT from all mice; open symbols represent fecal pellets from which no CFUs of HaitiWT could be detected. An asterisk denotes differences with a P value of <0.05 as determined by a Mann Whitney U test.

Figure 4



**Figure 4. Transient colonization by HaitiV is protective in an infant mouse model of cholera.** A) Survival curves of pups born to dams inoculated with HaitiV (n=15), formalin inactivated-HaitiV (n=24) or vehicle buffer (n=16) after HaitiWT challenge. Differences in the survival curves were determined by a log rank (Mantel Cox) test ( $p = 0.0013$  for HaitiV vs FI-HaitiV). B) HaitiWT CFUs recovered from the small intestines of the neonatal mice at the time they were moribund. Differences in CFU burdens were determined by a Mann Whitney U test.