A Direct Fluorescent-Monoclonal Antibody Staining Kit for Detection and Enumeration of *Vibrio cholerae* O139

For Laboratory Use Only. Not for Human Use.

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INTENDED USE
The Bengal DFA Cholerae Test Kit, by New Horizons Diagnostics, is intended for the direct detection of *Vibrio cholerae* O139 in clinical, food, and water samples.

INTRODUCTION
Until recently, only toxigenic *V. cholerae* serogroup O1 was believed to be capable of causing epidemic cholera. The other 137 serogroups, collectively called the “non-O1” serogroups (O2-O138) were not thought to have pandemic potential, and to include only organisms that cause sporadic diarrheal illness or occasional limited outbreaks. However, in the late part of 1992, large epidemics of cholera broke out in southern and eastern India and southern Bangladesh, and has since spread to the entire Indian subcontinent and several neighboring countries. The epidemic strain is a new serogroup of *V. cholerae* non-O1 and assigned to serogroup O139 with the synonym Bengal, to refer to its first isolation from areas surrounding the Bag of Bengal. By January, 1994, the organism had officially been reported from Bangladesh, India, Malaysia, Nepal, Pakistan, China, Thailand, the United Kingdom, and the United States.

New epidemics of cholera due to O139 strains are affecting persons of all ages in an area where most of the population except for young children has some level of acquired immunity to *V. cholerae* O1. This suggests that prior to immunity to O1 does not protect against O139 infection. It also suggests that existing and experimental O1 cholera vaccines will not induce immunity to this strain. Widespread transmission of *V. cholerae* O139 with outbreaks similar to those caused by the O1 can, therefore, be expected to occur in Latin America once O139 is introduced into the region.

The vibrios are aquatic bacteria found in a wide variety of environmental water sources. Cholera is spread primarily by ingestion of contaminated water or raw, poorly cooked, or recontaminated seafood. A rapid and reliable test for *V. cholerae* O139 is of great value to public health officials in monitoring and in controlling spread of the bacterium.

The Bengal DFA Test consists of a monoclonal antibody, specific for the surface antigen of O139 lipopolysaccharide in the outer membrane of *V. cholerae* O139 that is directly labeled with fluorescein isothio cyanate (FITC) for the rapid, simple detection and enumeration of *V. cholerae* O139 in clinical food and water samples. New Horizons also manufactures a similar kit,
Cholera DFA, for direct detection and enumeration of *V. cholerae* serogroup O15.

**PRINCIPLE**

The test kit is comprised of the Bengal DFA reagent and two control reagents. Water samples are concentrated and a sample is fixed onto a microscope slide. The test sample and control samples are then incubated with the DFA reagent. If the sample contains *V. cholerae* O139, the FITC-labeled monoclonal antibody will bind *V. cholerae* O139. After washing, the slide is examined under the fluorescent microscope.

**MATERIALS PROVIDED**

For 100 determinations

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>QUANTITY</th>
<th>DESCRIPTION</th>
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<tbody>
<tr>
<td>1. Cholera DFA Reagent</td>
<td>1 amber bottle (lyophilized)</td>
<td>Mouse monoclonal antibody to <em>V. cholerae</em> O139 - labeled with fluorescein. Contains 0.01% sodium azide, and protein in phosphate buffer.</td>
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<tr>
<td>2. Cholera Positive Control</td>
<td>1 bottle (1 mL)</td>
<td>Formalin fixed inactivated <em>V. cholerae</em> O1. Contains 0.05% sodium azide as a preservative.</td>
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<tr>
<td>3. Cholera Negative Control</td>
<td>1 bottle (1 mL)</td>
<td>Formalin fixed inactivated <em>V. cholerae</em> non-O1, non-O139. Contains 0.05% sodium azide as a preservative.</td>
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<tr>
<td>4. Glass Slide</td>
<td>10 slides</td>
<td>Reusable glass slide with 10 wells for samples.</td>
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<tr>
<td>5. Fluorescent Mounting Medium</td>
<td>1 bottle</td>
<td>Vial at working dilution. Contains an inhibitor that retards photobleaching of fluorescein.</td>
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MATERIALS REQUIRED BUT NOT SUPPLIED

1. Pipettes and micropettes and safety pipetting devices.
2. Coverslips, 22x50mm, No. 1 thickness.
3. Filters, 10.0 µm pore size and 0.45 µm pore sizes (available at NHD).
4. Absolute ethanol.
5. Filtering device with hand pump (available at NHD).
6. Distilled or deionized water.
7. Phosphate buffered saline (PBS).
8. Moist chamber - could be a petri dish with wet paper towel in bottom.
9. Incubator (35°C).
10. Epifluorescent Microscope. Refer to manufacturer instructions manual for the filter system which gives optimum results for FITC. (Maximum excitation wavelength = 490 nm and maximum emission wavelength = 520 nm.)

PRECAUTIONS

1. FOR USE WITH WATER SAMPLES ONLY. NOT FOR USE IN DIAGNOSIS OF HUMANS OR ANIMAL DISEASE.
2. Positive control and negative control materials have been fixed with formalin (2%). Good laboratory procedures dictate that these materials, as well as the samples, be handled and disposed of as potentially hazardous material.
3. Some of the reagents contain sodium azide. Sodium azide may react with lead and copper plumbing to form a highly explosive metal azide. On disposal, flush liberally with water.
REAGENT PREPARATION

1. Bengal DFA Reagent. Reconstitute in 1.0 mL of distilled or deionized water. Keep at 4-8°C following reconstitution. Store reagent away from bright light.

SAMPLE PREPARATION

1. Water samples of 100 to 500 mL should be collected in a clean container. If the water sample is turbid, it should be initially filtered through a 10.0 µm filter.

2. Using negative pressure from pump or other vacuum source, concentrate the 100 to 500 mL water sample. Place the filter on a clean petri dish and add 1 mL of sterile PBS onto it in order to obtain a thick suspension of organisms.

3. Stool samples need to be filtered through specimen filtering devices (available upon request from NHD).

SUGGESTED PROCEDURE

1. Prepare Bengal DFA Reagent and samples. All materials should be at room temperature.

2. Make a thin smear of resuspended sample by adding 5 µl on a well, then spreading the contents to cover the well.

3. Controls should be run at least once a day. Make a thin smear of the positive control by adding a small drop of the control on a well, then spreading the drop to cover the well. Make a similar thin smear of the negative control. Make a thin smear of the sample by adding approximately 5 µl of the sample to the slide and spreading.

4. Air dry.

5. Add one drop of absolute ethanol to each control or sample well to fix the smear, then air dry.

6. Add 10 µl of reconstituted Bengal DFA Reagent to each well.
7. Place the slides in a covered, moist chamber, and incubate at 35°C for 30 ± 5 minutes. Protect from light.

8. Rinse the slides thoroughly with PBS. Protect from light.

9. Absorb off excess moisture using a blotter paper.

10. Add a small volume (5-7 µl) of Fluorescent Mounting Medium to each well and cover with a 22 x50mm, No. 1 coverslip.

11. For best results, the slides should be read immediately at a magnification of 1000 X with oil immersion. Equivalent readings may be obtained if the slides are read within 24 hours. The slides must, however, be kept cool, in the dark, and sealed, or kept humid to prevent drying.

INTERPRETATION

1. Quality Control: For the test samples to be valid, the Positive Control should exhibit bright, apple-green, fluorescence and there should be no fluorescence on the Negative Control well.

2. The sample is considered Positive if apple-green fluorescence is observed that appears in a ring shaped pattern.

LIMITATIONS

The DFA procedure is capable of detecting as little as 10,000 organisms/mL. Given a concentration of the sample of at least 100-fold, the test should be capable of detecting the presence of any significant amount of V. cholerae O139 contamination in a sample. The sensitivity of the procedure will be affected by a number of factors, including the number of V. cholerae O139 in a sample, the quality of the sample, the sample age, etc. The presence of a heavy load of other organisms may mask a positive reaction or contribute to non-specific binding. Questionable samples should be repeated and additional samples from the same source should be obtained. Detection can also be enhanced by enriching the sample in alkaline peptone water (APW, pH 8.4 ± 0.2) for 6-8 hours. Call the Technical Service Department at New Horizons Diagnostics (NHD) at 443/543-5746 if you require assistance.
REFERENCES


Reorder No.: 89-116111

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