CDC Laboratory Guidance for Processing Environmental Samples

Introduction

Illness caused by the gram-negative bacteria in the genus *Legionella* is referred to as legionellosis. Legionellosis consists of two distinct clinical syndromes, Legionnaires' disease and Pontiac fever. Legionnaires' disease is characterized by pneumonia whereas Pontiac fever is a self-limiting, nonpneumonic, influenza-like illness. Inhalation of aerosols containing the bacterium is presumed to be the primary means of acquiring legionellosis. Aerosolized waters from cooling towers, evaporative...
condensers, showers, and humidifiers have been identified as sources of infection. *Legionella* species have been recovered from a wide variety of domestic water systems and are ubiquitous in freshwater environments. Although once considered transient contaminants of natural and domestic waters, *legionellae* are now known to be free-living organisms surviving as natural components of freshwater ecosystems. Domestic systems are complex environments in which concentrations of *legionellae* can fluctuate considerably depending upon water temperature, biocide levels, and presence of natural hosts (i.e. protozoa) for *legionellae* to parasitize. The choice of procedure used to recover *legionellae* from water samples is dependant upon the expected degree of bacterial contamination in a particular water source. Potable waters generally have low bacterial densities and are either cultured directly or concentrated to detect *legionellae*. Nonpotable waters, such as those from cooling towers, generally do not require concentration because of their high bacterial concentrations.

![Image](https://www.cdc.gov/legionella/labs/procedures-manual.htm)

**Figure 1. Nucleopore filter funnel assembly for concentrating water samples.**

This manual describes the procedures currently employed by the Centers for Disease Control and Prevention to process environmental samples obtained during investigations of legionellosis outbreaks. It includes information of the collection and concentration of water samples, preparation of samples for bacteriologic examination, formulas for media, sources of reagents, and air sampling techniques.

### Collection of Specimens

An environmental sampling protocol addressing selection of the appropriate sites to sample has been previously published (Barbaree et. al, 1987). Whenever possible, a collection of 1 (one) liter of water is preferred. Larger volumes of water (1 to 10 liters) are occasionally needed to detect *legionellae* in water that has very low concentrations of these bacteria such as municipal water supplies. If a liter cannot be collected from a sample source, a smaller volume is acceptable. Water samples should be collected in a sterile 1-liter wide-mouth screw cap polypropylene plastic bottle. If the water source has been recently treated with chlorine, add 0.5 ml of 0.1N sodium thiosulfate to each 1 liter sample to neutralize the disinfectant.
Swabs of faucet aerators and shower heads should be taken before water samples from these sites. The sample should be taken with the aerator or shower head removed if possible. Polyester swabs with wooden shafts work well for this purpose. The swabs should be submerged in 3-5 ml of water taken at the same time to prevent drying during transport.

All samples should be transported to the laboratory in insulated coolers as protection against extreme heat or cold. Samples that will not reach the laboratory within 72 hrs should be refrigerated before shipping. Samples that reach the laboratory but cannot be processed within 72 hrs of collection should be refrigerated.

Preparation of Specimens for Bacteriologic Examination

a) Filtration (potable water)

Samples are filter concentrated in a biological safety cabinet by pouring the samples into a sterile 47-mm filter funnel assembly containing a 0.2um polycarbonate filter (Fisher Scientific, 3970 Johns Creek Ct., Suite 500, Suwanee, GA 30024). A vacuum source and side-arm flask are necessary to operate the apparatus (Figure 1). Care should be taken to avoid drying of the filters. When the sample has passed through the filter, the filter is removed aseptically from the holder with sterile filter forceps, folded to the outside, and placed into a sterile, 50-ml centrifuge tube containing 5ml of sterile water. The centrifuge tube is then vortexed for one minute to free bacteria and organic material from the filter. If more than one filter is required to concentrate a sample, the additional filters are added to the same sample tube.

b) Direct Plating (nonpotable water)

Nonpotable water rarely requires concentration and can be processed directly. Samples that are found to have high concentrations of bacteria upon direct plating can be treated with a low pH buffer or serially diluted with sterile deionized water, an appropriate buffer, or sterile chorine-free tap water to improve primary isolation of legionellae.
Plating of Specimens

Buffered charcoal yeast extract (BCYE) agar containing 0.1% alpha-ketoglutarate is the base medium used for the recovery of Legionella from environmental and clinical specimens (Appendix 1). Two types of selective BCYE agar supplemented with antimicrobial agents are included in the standard regimen of media used in the processing of environmental samples. The first, designated PCV, contains polymyxin B, cycloheximide, and vancomycin. The second, GPCV, is identical to PCV except for the addition of the amino acid glycine to the medium. PCV without L-cysteine [PCV(-)] may be used as a negative control medium. Several variations in the antimicrobial supplements have been described and appear to be adequate for the recovery of legionellae. Figure 2 shows a water sample cultured on the four media described above.

Although the majority of Legionella spp. grow readily on BCYE, some require supplementation with bovine serum albumin to enhance growth. Legionella micdadei and several strains of Legionella bozemanii show a preference for BCYE with 1.0% albumin (ABCYE) over standard BCYE. The inclusion of ABCYE in the regimen of plating media may be beneficial when clinical evidence indicates that a species other than Legionella pneumophila may be the causative agent.

Plating of Filtered Samples

1. After vortexing of filter (see Preparation of Specimens), inoculate 1 BCYE, 2 PCV, and 2 GPCV and 1 PCV (-) plate each with 0.1 ml of the suspension and spread with a sterile glass or sterile disposable plastic rod.
2. Incubate plates at 35°C in a candle jar or in a humidified incubator with an atmosphere of 2.5% CO₂ in air.
3. Store the remainder of the sample at 4°C.

Direct Plating
1. Place 0.1 ml of the sample onto BCYE and modified BCYE plates as described above and spread with a sterile glass rod or sterile disposable plastic spreader.

2. Incubate plates as described above.

**Acid Treatment**

For some water samples with high concentrations of bacteria, it is necessary to use a selective procedure to reduce the number of non-**Legionella** bacteria before culture. Acid treatment and heating of samples are routinely used for this purpose. **Legionellae** are more resistant to lower pH and brief exposure to higher temperatures than many other freshwater bacteria. The following procedure is used by CDC laboratories to prepare these samples for culture.

1. Place 1.0 ml of the vortexed suspension into a sterile 15 ml centrifuge tube containing 1.0 ml of acid buffer and mix (Appendix 2).

2. Incubate the acidified suspension for 15 minutes at room temperature. (Acid treatment can be extended to 30 minutes if plates are overgrown after initial 15-minutes treatment).

3. Place 0.1 ml of the suspension onto BCYE and modified BCYE plates and spread with a sterile glass rod or sterile disposable plastic spreader.

4. Incubate plates at 35°C in a candle jar or in a humidified incubator with an atmosphere of 2.5% CO₂ in air.

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**Examination of Cultures for Legionellae**

1. Examine all cultures after 72 to 96 hours incubation. Macroscopic examination should be done with a dissecting microscope and oblique lighting to detect bacterial colonies resembling **Legionella**. These colonies are convex and round with entire edges (Figure 3). The center of the colony is usually bright white in color with a textured appearance that has been described as “cut-glass like” or speckled. The white center of the colony is often bordered with blue, purple, green or red
autofluorescence. Further examination of primary isolation plates with long-wave UV light can easily detect these fluorescent colonies.

2. Aseptically pick each suspect colony onto (a) BCYE agar plate without L-cysteine [BCYE(-)] or a BCYE/BCYE(-) biplate. If a L-cysteine (-) biplate is not available, a less desirable alternative is another L-cysteine media such a blood agar plate. Pick 2-4 representative colonies per water sample

3. Streak the inoculated portion of each plate with a sterile loop to provide areas of heavy growth, and incubate the plates for 24 hrs.

4. If no growth is detected after 24 hrs, continue incubation for an additional 24 hrs. Colony picks that grow on BCYE agar, but not BCYE(-) agar, are presumptive Legionella species and must be examined further for definite identification using the direct fluorescent antibody (DFA) or slide agglutination test (SAT).

5. Since Legionella is a relatively slow-growing bacterium, negative plates should be reincubated until 7 days postinoculation and then re-examined for Legionella colonies. Discard plates after the seventh day.

Heat Enrichment

Heat enrichment (or “coculture”) of specimens that are negative for Legionella upon primary culture can improve recovery of this organism significantly. This procedure is based on the finding that many water samples contain protozoa that can serve as host for the intracellular multiplication of legionellae. Incubating the samples at 35ºC allows the legionellae to multiply in these protozoa and achieve a culturable concentration. Results from several epidemic investigations indicate that subsequent replating of specimens results in the isolation of Legionella spp. from approximately 30% of all samples negative upon primary isolation. Twenty-five milliliter aliquots of samples are incubated at 35ºC in 50 ml plastic screw cap centrifuge tubes. With swab specimens, the entire swab sample is incubated. The tube caps should be tightened to minimize evaporation of the accompanying 3-5 ml of water sample. Incubated samples are cultured at 2-week intervals for as long as 6 weeks. Samples are plated on the same regimen of plates as in primary isolation. If overgrowth of media occurs, samples can be acid treated as previously described.

Appendix 1

Preparation of Media

Buffered charcoal yeast extract (BCYE) agar

A. Composition

<table>
<thead>
<tr>
<th>Component</th>
<th>Amount</th>
</tr>
</thead>
<tbody>
<tr>
<td>Norit SG charcoal</td>
<td>2.0 g</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>10.0 g</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------</td>
<td>------------</td>
</tr>
<tr>
<td>ACES buffer</td>
<td>10.0 g</td>
</tr>
<tr>
<td>Ferric pyrophosphate (soluble)</td>
<td>0.25 g</td>
</tr>
<tr>
<td>L-cysteine HCl-H2O</td>
<td>0.40 g</td>
</tr>
<tr>
<td>Agar, bacto (Oxoid)</td>
<td>17.0 g</td>
</tr>
<tr>
<td>Potassium alpha-ketoglutarate</td>
<td>1.0 g</td>
</tr>
<tr>
<td>Distilled water</td>
<td>1,000 ml</td>
</tr>
</tbody>
</table>

Antibiotic components for media

1. **GPCV medium**
   - Glycine: 0.3%
   - Polymyxin B: 100 units/ml
   - Vancomycin: 5 ug/ml
   - Cycloheximide: 80 ug/ml

2. **PCV medium – GPCV medium without glycine**

**ABCYE medium**

- Bovine serum albumin (fraction V): 10g/L

Dehydrated BCYE agar base is available from several commercial sources.

B. Directions

1. Add ACES buffer to 940 ml of distilled water and dissolve in a 50°C water bath.
2. Add enough 1.0 N KOH (about 40 ml) to the buffer solution to bring the pH up to 6.9 and mix. (Do not use NaOH because it has been found to be inhibitory to *Legionella pneumophila*).
3. Into a second flask, add charcoal, yeast extract, alpha-keto-glutarate, and agar. Mix the dry powders.
4. Pour the buffer solution into the second flask containing the dry powders and mix.
5. Carefully heat to dissolve the agar, then sterilize by autoclaving at 121°C for 15 minutes.
6. Immediately place the medium in 50°C water bath.
7. For complete medium, add membrane-filtered solution of L-cysteine to the medium and mix thoroughly.
8. Add membrane-filtered soluble ferric pyrophosphate, and mix thoroughly. Do not mix iron and cysteine before adding to medium.
9. Adjust the pH of the medium to 6.9 at room temperature.

Since reagents may vary, each laboratory must determine the amount of KOH required. Hold the completed medium at 50°C, pour a 10 ml sample, and check the pH at room temperature. When necessary, adjust the completed medium with either 1.0 N KOH or 1.0 HCl. Note that the pKa of
ACES buffer is 6.9 at 20ºC and 6.8 at 25ºC. Its pKa is affected by temperature (0.02 pH unit/o).
However, once the agar has solidified, the pH does not appear to change with temperature but
remains at 6.9.
10. For preparation of BCYE + antibiotics, add membrane-filtered antibiotics and mix.
11. For BCYE + albumin agar, dissolve the albumin in distilled water and filter sterilize before addition to
the medium.
12. Dispense 20 ml per 15 X 100-mm Petri dish.

The medium must be mixed frequently during the pouring to keep the charcoal particles suspended.
After the medium has solidified, the plates should be stored in plastic bags in the refrigerator in the dark.
They should be good for approximately 4 months, provided they pass quality control.

C. Precautions

1. The soluble ferric pyrophosphate must be kept dry and in the dark.
2. Fresh solutions of L-cysteine and ferric pyrophosphate must be prepared each time. Since L-cysteine
   is a chelating agent, it must be added to the medium first. Both solutions must be added slowly. This
   is also true for adding the 1.0 N KOH.
3. Do not use ferric pyrophosphate if its green color becomes yellow or brown.

Quality Control of Media

Each batch of medium should be tested for pH and the ability to support the growth of \textit{L. pneumophila}.
Ideally, all media should be tested with a standardized tissue inoculum. However, since this material is
not generally available, stock culture inoculum is recommended.

1. The pH of the medium must be 6.90 +/- 0.05 Measure with a pH meter. To check solid media, use a
   surface electrode (if available), or emulsify the agar from one plate in distilled water, and measure
   the pH of the emulsion. The pH is critical. If given lots of ingredients do not conform to the
   acceptable pH range, adjust the pH of the completed liquid medium by adding 1.0 N KOH or 1.0 N
   HCl.
2. Check for support of growth in the following manner:
   a. Prepare a standard stock culture inoculum by suspending \textit{L. pneumophila} (log phase) in sterile
distilled water to a concentration of 10⁶ cells/ml.
   b. Inoculate the medium with 0.05 ml of the standard inoculum. Streak the plates to obtain isolated
      colonies, and incubate at 35ºC in air plus 2.5% CO₂.
   c. Examine the media daily. On agar plates, growth should be present in the heavily inoculated area
      after one day. Isolated colonies should be macroscopically visible in 2 to 3 days. If cells in
      standard inoculum have been refrigerated or frozen, growth will be slower.
3. If possible, selective media should be checked using an environmental water specimen containing a
   known concentration of legionellae and other bacteria.
Appendix 2

Reagents for Acid Treatment Procedure

KCl-HCl Acid

A. 0.2 M KCl (14.9 g/L in distilled water)
B. 0.2 M HCl (16.7 g/L in distilled water)
   1. Mix 18 parts of (A) with 1 part of (B)
   2. Dispense into screw-capped tubes in 1.0 ml volumes and sterilze by autoclaving

References

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