The Microbiology of Drinking Water (2007) – Part 6 – Methods for the isolation and enumeration of sulphite-reducing clostridia and *Clostridium perfringens* by membrane filtration

*Methods for the Examination of Waters and Associated Materials*
The Microbiology of Drinking Water (2007) - Part 6 - Methods for the isolation and enumeration of sulphite-reducing clostridia and *Clostridium perfringens* by membrane filtration

**Methods for the Examination of Waters and Associated Materials**

This booklet contains two methods for the isolation and enumeration of sulphite-reducing clostridia and *Clostridium perfringens* by membrane filtration, and supersedes “The Microbiology of Drinking Water (2004) - Part 6 - Methods for the isolation and enumeration of sulphite-reducing clostridia and *Clostridium perfringens* by membrane filtration”.

Within this series there are separate booklets, each dealing with different topics concerning the microbiology of drinking water. Other booklets include

The Microbiology of Drinking Water (2002)
Part 1 - Water quality and public health
Part 2 - Practices and procedures for sampling
Part 3 - Practices and procedures for laboratories
Part 4 - Methods for the isolation and enumeration of coliform bacteria and *Escherichia coli* (including *E. coli* O157:H7)
Part 5 - The isolation and enumeration of enterococci by membrane filtration
Part 8 - Methods for the isolation and enumeration of *Aeromonas* and *Pseudomonas aeruginosa* by membrane filtration
Part 10 - Methods for the isolation and enumeration of *Yersinia*, *Vibrio* and *Campylobacter* by selective enrichment.

The Microbiology of Drinking Water (2004)
Part 11 - Taste, odour and related aesthetic problems
Part 12 - Methods for micro-organisms associated with taste, odour and related aesthetic problems.

The Microbiology of Drinking Water (2006)
Part 5 - The isolation and enumeration of enterococci by membrane filtration
Part 9 - The isolation and enumeration of *Salmonella* and *Shigella* by selective enrichment, membrane filtration and multiple tube-most probable number techniques

The Microbiology of Drinking Water (2007)
Part 7 - Methods for the enumeration of heterotrophic bacteria
Part 13 - The isolation and enumeration of aerobic spore-forming bacteria by membrane filtration

Whilst specific commercial products may be referred to in this document, this does not constitute an endorsement of these products but serves only as an illustrative example of the type of products available. Equivalent products may be available and it should be understood that the performance of the method might differ when other materials are used and all should be confirmed by validation of the method.
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About this series

Introduction

This booklet is part of a series intended to provide authoritative guidance on recommended methods of sampling and analysis for determining the quality of drinking water, ground water, river water and sea water, waste water and effluents as well as sewage sludges, sediments, soils (including contaminated land) and biota. In addition, short reviews of the most important analytical techniques of interest to the water and sewage industries are included.

Performance of methods

Ideally, all methods should be fully evaluated with results from performance tests. These methods should be capable of establishing, within specified or pre-determined and acceptable limits of deviation and detection, whether or not any sample contains concentrations of parameters above those of interest.

For a method to be considered fully evaluated, individual results from at least three laboratories should be reported. The specifications of performance generally relate to maximum tolerable values for total error (random and systematic errors) systematic error (bias) total standard deviation and limit of detection. Often, full evaluation is not possible and only limited performance data may be available.

In addition, good laboratory practice and analytical quality control are essential if satisfactory results are to be achieved.

Standing Committee of Analysts

The preparation of booklets within the series “Methods for the Examination of Waters and Associated Materials” and their continuing revision is the responsibility of the Standing Committee of Analysts. This committee was established in 1972 by the Department of the Environment and is now managed by the Environment Agency. At present, there are nine working groups, each responsible for one section or aspect of water quality analysis. They are

1 General principles of sampling and accuracy of results
2 Microbiological methods
3 Empirical and physical methods
4 Metals and metalloids
5 General non-metallic substances
6 Organic impurities
7 Biological methods
8 Biodegradability and inhibition methods
9 Radiochemical methods

The actual methods and reviews are produced by smaller panels of experts in the appropriate field, in co-operation with the working group and main committee. The names of those members principally associated with these methods are listed at the back of this booklet.

Publication of new or revised methods will be notified to the technical press. If users wish to receive copies or advanced notice of forthcoming publications or obtain details of the index of methods then contact the Secretary on the Agency’s web-page (www.environment-agency.gov.uk/nls) or by post.

Every effort is made to avoid errors appearing in the published text. If, however, any are found, please notify the Secretary.

Dr D Westwood
Secretary
July 2006

Warning to users

The analytical procedures described in this booklet should only be carried out under the proper supervision of competent, trained analysts in properly equipped laboratories.

All possible safety precautions should be followed and appropriate regulatory requirements complied with. This should include compliance with the Health and Safety at Work etc Act 1974 and all regulations made under the Act, and the Control of Substances Hazardous to Health Regulations 2002 (SI 2002/2677). Where particular or exceptional hazards exist in carrying out the procedures described in this booklet, then specific attention is noted.

Numerous publications are available giving practical details on first aid and laboratory safety. These should be consulted and be readily accessible to all analysts. Amongst such publications are: “Safe Practices in Chemical Laboratories” and “Hazards in the Chemical Laboratory”, 1992, produced by the Royal Society of Chemistry; “Guidelines for Microbiological Safety”, 1986, Portland Press, Colchester, produced by Member Societies of the Microbiological Consultative Committee; and “Safety Precautions, Notes for Guidance” produced by the Public Health Laboratory Service. Another useful publication is “Good Laboratory Practice” produced by the Department of Health.
A Enumeration of sulphite-reducing clostridia by membrane filtration

A1 Introduction

Tests for sulphite-reducing clostridia play only a subsidiary role in water examination. The organisms form spores which are environmentally resistant and their presence may indicate soil contamination, although some species may grow in deposits, and be associated with corrosion of distribution pipes. *Clostridium perfringens* is a sulphite-reducing species and is associated with faecal contamination. The significance of sulphite-reducing clostridia and *Clostridium perfringens* in water treatment and supply are described elsewhere\(^\text{(1)}\) in this series.

A2 Scope

The method is suitable for the examination of drinking waters, including samples from all stages of treatment and distribution, and those source waters of moderate turbidity.

Users wishing to employ this method should verify its performance under their own laboratory conditions\(^\text{(2)}\).

A3 Definitions

Sulphite-reducing clostridia are Gram-positive anaerobic spore-forming rod-shaped bacteria, which in the context of this method reduce sulphite to sulphide at 37 °C within 24 hours.

A4 Principle

A volume of sample is filtered and the membrane filter placed on the surface of an agar medium containing sulphite, iron(III) and D-cycloserine (which inhibits other bacteria and reduces the size of colonies that develop). The agar medium is then incubated under anaerobic conditions at 37 °C. Sulphite-reducing clostridia usually produce black colonies as a result of the reduction of sulphite to sulphide which reacts with the iron(III) salt. If only a spore count is required then the sample is heat-treated at 60 °C prior to filtration in order to kill vegetative bacteria.

A5 Limitations

The method is suitable for most types of aqueous samples except those with high turbidities which tend to block the membrane filter. This will limit the volume of sample that can be filtered. Accumulated deposit on the membrane filter may mask or inhibit the growth of indicator organisms. The maximum number of colonies that should be counted from a single membrane filter is approximately 100. Some clostridia may produce spreading colonies which may reduce the potential maximum count.

A6 Health and safety

Media, reagents and bacteria used in this method are covered by the Control of Substances Hazardous to Health Regulations\(^\text{(3)}\) and appropriate risk assessments should be made before adopting this method. Standard laboratory microbiology safety procedures should be followed and guidance is given elsewhere\(^\text{(2)}\) in this series.
A7 Apparatus

Standard laboratory equipment should be used which conforms to the performance criteria outlined elsewhere\(^{(2)}\) in this series. Principally, appropriate membrane filtration apparatus and incubators (fan assisted, static temperature) are required. Other items include:

A7.1 Sterile sample bottles of appropriate volume, made of suitable material, containing sufficient sodium thiosulphate pentahydrate to give a final concentration in the sample of not less than 18 mg/l (for example, 0.1 ml of a 1.8 % m/v solution of Na\(_2\)S\(_2\)O\(_3\).5H\(_2\)O per 100 ml of sample, or equivalent).

A7.2 Incubator capable of maintaining a temperature of 37.0 ± 1.0 °C.

A7.3 Anaerobic jars, or similar equipment, and anaerobic gas-generating system (for generating atmospheres of approximately 9 - 13 % carbon dioxide).

A7.4 Filtration apparatus, sterile filter funnels, and source of vacuum.

A7.5 Sterile gridded membrane filters, for example, white, 47 mm diameter cellulose-based, 0.45 µm nominal pore size.

A7.6 Smooth-tipped forceps.

A8 Media and reagents

Commercial formulations of these media and reagents may be available, but may possess minor variations to their formulation. The performance of all media and reagents should be verified prior to their use in the method\(^{(2)}\). Variations in the preparation and storage of media should also be verified. Water should be distilled, deionised or of similar quality. Unless otherwise stated chemical constituents should be added as anhydrous salts. If the pH of the medium is not within its stated range, then, before heating, it should be adjusted accordingly. Where media are stored in a refrigerator they should be allowed to reach room temperature before use.

A8.1 \textit{Tryptose sulphite cycloserine agar without egg yolk} \(^{(4,5)}\)

\begin{align*}
\text{Yeast extract} & \quad 5 \text{ g} \\
\text{Tryptose} & \quad 15 \text{ g} \\
\text{Soya peptone} & \quad 5 \text{ g} \\
\text{Sodium metabisulphite} & \quad 1 \text{ g} \\
\text{Iron(III) ammonium citrate} & \quad 1 \text{ g} \\
\text{Agar} & \quad 14 \text{ g} \\
\text{Water} & \quad 1 \text{ litre}
\end{align*}

Suspend the ingredients in the water and dissolve by heating and stirring the mixture. Sterilise the solution by autoclaving at 121 °C for 15 minutes. Allow the medium to cool to 46 ± 2 °C. Add 4 ml of a filter-sterilised solution of D-cycloserine in water at a concentration of 100 mg/ml. Mix the solution thoroughly, and dispense into Petri dishes. The final pH of the medium should be 7.6 ± 0.2. The Petri dishes should be the vented type to ensure anaerobic conditions for the medium during storage and incubation.

Performance of the medium deteriorates during storage due to exposure to oxygen. Prepared media may be stored in a refrigerator under anaerobic conditions at a temperature
of 5 ± 3 °C for up to one week. However, some anaerobic generating systems may not work satisfactory at this temperature). When fresh medium is used, the colony characteristics that are observed tend to be more defined. The medium, once removed from the refrigerator, should be discarded if not used.

A8.2 Other media

Standard and commercial formulations of other media and reagents used in this method include quarter-strength Ringer’s solution and maximum recovery diluent.

A9 Analytical procedure

A9.1 Sample preparation

The volumes, and dilutions, of samples should be chosen so that the number of colonies to be counted on the membrane filter lies, if possible, between 20 and 80. With some waters, it may be advantageous to filter a selection of different volumes of sample so that the number of colonies on one of the membrane filters is likely to fall within this range. For treated waters, filter 100 ml of the sample. For polluted waters, either filter smaller volumes, or dilutions of the sample made with quarter-strength Ringer’s solution or maximum recovery diluent.

If it is the intention to count only the spores of sulphite-reducing clostridia then the volume of sample should be heated to 60 ± 2 °C (for example, in a water bath) and the whole volume maintained at this temperature for 15 ± 1 minutes. The temperature may be monitored by placing an appropriate thermometer in a similar bottle containing the same volume of water as the sample volume being treated.

A9.2 Sample processing

Place the sterile filtration apparatus in position and connect to a source of vacuum, with the stopcock turned off. Remove the funnel and, holding the edge of the membrane filter with sterile smooth-tipped forceps, place a sterile membrane filter, grid-side upwards, onto the porous disc of the filter base. Replace the sterile funnel securely on the filter base. Pour or pipette the required volume of sample into the funnel. When the volume of sample to be filtered is less than 10 ml, add 10 - 20 ml of sterile diluent (for example, quarter-strength Ringer’s solution or maximum recovery diluent) to the funnel before addition of the sample. This aids the dispersion of the bacteria over the entire surface of the membrane filter during the filtration. Open the stopcock and apply a vacuum not exceeding 65 kPa (500 mm of mercury) and filter the sample slowly through the membrane filter. Close the stopcock as soon as the sample has been filtered.

Remove the funnel and transfer the membrane filter carefully to a Petri dish of well-dried tryptose sulphite cycloserine agar. Ensure that no air bubbles are trapped between the membrane filter and the medium. ‘Rolling’ the membrane filter onto the medium minimises the likelihood of air bubbles becoming trapped.

As the spores of sulphite-reducing clostridia are very resilient, funnels that have been used once should be sterilised by autoclaving before being used again. Placing funnels in a water bath at this stage may not be sufficient to kill spores. If different volumes of the same sample are to be examined, the funnel may be re-used without sterilising the funnel provided that the smallest volume, or highest dilution of the sample, is filtered first. For different samples, take a fresh pre-sterilised funnel and repeat the filtration process. During
the filtration of a series of samples, the filter base need not be sterilised unless it becomes, or is suspected of being, contaminated or a membrane filter becomes damaged. When funnels are not in use they should be covered with a sterile lid or a sterile Petri dish lid.

The time between the end of the filtration step and the beginning of the incubation stage should be as short as possible and no longer than 2 hours.

Incubate the Petri dishes at 37 °C in an anaerobic jar or similar system containing an indicator of anaerobiosis and an atmosphere containing 9 - 13 % carbon dioxide. Examine the dishes after 21 ± 3 hours incubation.

A9.3 Reading of results

After incubation, count all black or grey colonies (see Figure A1).

![Figure A1 Typical colonies of sulphite-reducing clostridia on tryptose sulphite cycloserine agar](image)

A9.4 Confirmation tests

The specificity of tryptose sulphite cycloserine agar is such that confirmation of isolates is not usually required.

A10 Calculations

A10.1 Confirmed sulphite-reducing clostridia

The number of confirmed sulphite-reducing clostridia colonies is generally quoted as the number of colonies per 100 ml. Calculate the confirmed count as follows:

\[
\text{Confirmed count/100 ml} = \frac{\text{Number of colonies} \times 100 \times \text{DF}}{\text{Volume of sample filtered (ml)}}
\]

Where DF is the appropriate dilution factor.
A11 Expression of results

Counts for sulphite-reducing clostridia are expressed in colony forming units per volume of sample. For drinking water, the volume is typically 100 ml.

A12 Quality assurance

New batches of media and reagents should be tested with appropriate reference strains of target bacteria (for example Clostridium perfringens) and non-target bacteria (for example Bacillus species). Petri dishes should be incubated for 21 ± 3 hours at 37 °C. Further details of media and analytical quality control are given elsewhere\(^{2}\) in this series.

A13 References


**B** Enumeration of *Clostridium perfringens* by membrane filtration

**B1** Introduction

Tests for *Clostridium perfringens* play only a subsidiary role in water examination. The organisms form spores which are resistant to environmental stress and can persist in the environment for some time. *Clostridium perfringens* is associated with faecal contamination. If found at a time when other faecal indicator organisms are no longer detectable, the organism may indicate remote or intermittent pollution. The significance of *Clostridium perfringens* in water treatment and supply are described elsewhere\(^{(1)}\) in this series.

**B2** Scope

The method is suitable for the examination of drinking waters, including samples from all stages of treatment and distribution, and those source waters of moderate turbidity.

Users wishing to employ this method should verify its performance under their own laboratory conditions\(^{(2)}\).

**B3** Definitions

*Clostridium perfringens* is a Gram-positive anaerobic spore-forming rod-shaped bacterium, which in the context of this method reduces sulphite to sulphide at 44 °C within 24 hours. *Clostridium perfringens* is non-motile, reduces nitrate, ferments lactose and liquefies gelatin. *Clostridium perfringens* also produces the enzyme acid phosphatase, which is a diagnostic characteristic for this species amongst the clostridia.

**B4** Principle

A volume of sample is filtered and the membrane filter placed on the surface of an agar medium containing sulphite, iron(III) and D-cycloserine (which inhibits other bacteria and reduces the size of colonies that develop). The agar medium is incubated under anaerobic conditions at 44 °C. *Clostridium perfringens* usually produces black colonies as a result of the reduction of sulphite to sulphide which reacts with the iron(III) salt. If only a spore count is required, then the sample is heat-treated at 60 °C prior to filtration in order to kill vegetative bacteria.

**B5** Limitations

The method is suitable for most types of aqueous samples except those with high turbidities which tend to block the membrane filter. This will limit the volume of sample that can be filtered. Accumulated deposit on the membrane filter may mask or inhibit the growth of indicator organisms. The maximum number of colonies that should be counted from a single membrane is approximately 100.

**B6** Health and safety

Media, reagents and bacteria used in this method are covered by the Control of Substances Hazardous to Health Regulations\(^{(3)}\) and appropriate risk assessments should be made before adopting this method. Standard laboratory microbiology safety procedures should be followed and guidance is given elsewhere\(^{(2)}\) in this series.
B7 Apparatus

Standard laboratory equipment should be used which conforms to the performance criteria outlined elsewhere\(^{(2)}\) in this series. Principally, appropriate membrane filtration apparatus and incubators (fan assisted, static temperature) are required. Others items include:

B7.1 Sterile sample bottles of appropriate volume, made of suitable material, containing sufficient sodium thiosulphate pentahydrate to give a final concentration in the sample of not less than 18 mg/l (for example, 0.1 ml of a 1.8 % m/v solution of Na\(_2\)S\(_2\)O\(_3\).5H\(_2\)O per 100 ml of sample, or equivalent).

B7.2 Incubators capable of maintaining temperatures of 37.0 ± 1.0 °C and 44.0 ± 0.5 °C.

B7.3 Anaerobic jars, or similar equipment, and anaerobic gas-generating system (for generating atmospheres of approximately 9 - 13 % carbon dioxide).

B7.4 Filtration apparatus, sterile filter funnels, and source of vacuum.

B7.5 Sterile gridded membrane filters, for example, white, 47 mm diameter, cellulose-based, 0.45 \(\mu\)m nominal pore size.

B7.6 Smooth-tipped forceps.

B8 Media and reagents

Commercial formulations of these media and reagents may be available, but may possess minor variations to their formulation. The performance of all media and reagents should be verified prior to their use in the method\(^{(2)}\). Variations in the preparation and storage of media should also be verified. Water should be distilled, deionised or of similar quality. Unless otherwise stated chemical constituents should be added as anhydrous salts. If the pH of the medium is not within its stated range, then, before heating, it should be adjusted accordingly. Where media are stored in a refrigerator they should be allowed to reach room temperature before use.

B8.1 *Tryptose sulphite cycloserine agar without egg yolk*\(^{(4, 5)}\)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Yeast extract</td>
<td>5 g</td>
</tr>
<tr>
<td>Tryptose</td>
<td>15 g</td>
</tr>
<tr>
<td>Soya peptone</td>
<td>5 g</td>
</tr>
<tr>
<td>Sodium metabisulphite</td>
<td>1 g</td>
</tr>
<tr>
<td>Iron(III) ammonium citrate</td>
<td>1 g</td>
</tr>
<tr>
<td>Agar</td>
<td>14 g</td>
</tr>
<tr>
<td>Water</td>
<td>1 litre</td>
</tr>
</tbody>
</table>

Suspend the ingredients in the water and dissolve by heating and stirring the mixture. Sterilise the solution by autoclaving at 121 °C for 15 minutes. Allow the medium to cool to 46 ± 2 °C. Add 4 ml of a filter-sterilised solution of D-cycloserine in water at a concentration of 100 mg/ml. Mix the solution thoroughly, and dispense into Petri dishes. The final pH of the medium should be 7.6 ± 0.2. The Petri dishes should be the vented type to ensure anaerobic conditions for the medium during storage and incubation.

Performance of the medium deteriorates during storage due to exposure to oxygen. Prepared media may be stored in a refrigerator under anaerobic conditions at a temperature
of 5 ± 3 °C for up to one week. (However, some anaerobic generating systems may not work satisfactorily at this temperature). When fresh medium is used, the colony characteristics that are observed tend to be more defined. Medium, once removed from the refrigerator, should be discarded if not used.

B8.2  **Buffered nitrate-motility medium**\(^6\)

<table>
<thead>
<tr>
<th>Component</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beef extract</td>
<td>3 g</td>
</tr>
<tr>
<td>Peptone</td>
<td>5 g</td>
</tr>
<tr>
<td>Potassium nitrate</td>
<td>5 g</td>
</tr>
<tr>
<td>D-Galactose</td>
<td>5 g</td>
</tr>
<tr>
<td>Glycerol</td>
<td>5 g</td>
</tr>
<tr>
<td>Disodium hydrogen phosphate</td>
<td>2.5 g</td>
</tr>
<tr>
<td>Agar</td>
<td>3 g</td>
</tr>
<tr>
<td>Water</td>
<td>1 litre</td>
</tr>
</tbody>
</table>

Dissolve the solid ingredients in 950 ml of water by heating to boiling point whilst stirring the mixture continuously. Dissolve the glycerol in 50 ml of water in a separate container and add this solution to the base medium and mix thoroughly. Dispense the resulting solution, typically in 10 ml aliquots, in appropriately sized tubes. Cap the tubes. Sterilise the medium by autoclaving at 121 °C for 15 minutes. The final pH of the medium should be 7.3 ± 0.2. Prepared tubes may be stored at a temperature of 5 ± 3 °C for up to one month if protected against dehydration.

Before use, stored media should be heated for 10 - 15 minutes in a boiling water bath, to ensure that the contents have melted and to eliminate any absorbed oxygen. The tubes should then be allowed to cool and the media to solidify ready for use.

B8.3  **Nitrate reduction test reagents**\(^7\)

**Reagent A**

- Sulphanilic acid 0.8 g
- Acetic acid 30 ml
- Water 100 ml

Warm gently to aid dissolution.

**Reagent B**

- N, N-dimethyl-1-naphthylamine 0.6 ml
- Acetic acid 30 ml
- Water 100 ml

Dissolve the amine in the acetic acid solution. To aid dissolution, warm gently (for example, by placing in a water bath at 40 - 60 °C).

The reagents may be stored at a temperature of 5 ± 3 °C for up to several months.

For the combined reagent, mix equal volumes of reagents A and B immediately prior to use. Prepare in small volumes sufficient for the tests to be performed. The combined reagent may be stored at a temperature of 5 ± 3 °C, protected from direct light, and should be used within 24 hours.
B8.4 Lactose-gelatin medium\(^{(4)}\)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tryptose</td>
<td>15 g</td>
</tr>
<tr>
<td>Yeast extract</td>
<td>10 g</td>
</tr>
<tr>
<td>Disodium hydrogen phosphate</td>
<td>5 g</td>
</tr>
<tr>
<td>Gelatin</td>
<td>120 g</td>
</tr>
<tr>
<td>Lactose</td>
<td>10 g</td>
</tr>
<tr>
<td>Phenol red (0.4 % m/v solution)</td>
<td>12.5 ml</td>
</tr>
<tr>
<td>Water</td>
<td>1 litre</td>
</tr>
</tbody>
</table>

Dissolve the ingredients, except the gelatin, lactose and phenol red, in the water. Add the gelatin gradually whilst stirring the mixture continuously and warming gently to aid dissolution. Adjust the pH to 7.5 ± 0.2. Add the lactose and phenol red and mix thoroughly to dissolve. Dispense the resulting solution, typically in 10 ml aliquots, in appropriately sized tubes. Cap the tubes. Sterilise the medium at 121 °C for 15 minutes. The final pH should be 7.5 ± 0.2. Prepared media may be stored at a temperature of 5 ± 3 °C for up to one month if protected against dehydration.

Before use stored media should be heated for 10 - 15 minutes in a boiling water bath, to ensure that the contents have melted and to eliminate any absorbed oxygen. The tubes should then be allowed to cool and the media to solidify ready for use.

B8.5 Columbia agar base

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Special peptone</td>
<td>23.0 g</td>
</tr>
<tr>
<td>Starch</td>
<td>1.0 g</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>5.0 g</td>
</tr>
<tr>
<td>Agar</td>
<td>10.0 g</td>
</tr>
<tr>
<td>Water</td>
<td>1 litre</td>
</tr>
</tbody>
</table>

Suspend the ingredients in the water and dissolve by heating and stirring the mixture. Sterilise the solution by autoclaving at 121 °C for 15 minutes. Cool and dispense into Petri dishes. The Petri dishes should be the vented type to ensure anaerobic conditions for the medium during incubation. The final pH of the medium should be 7.3 ± 0.2. Sterile media may be stored at a temperature of 5 ± 3 °C for up to one month, if protected against dehydration.

B8.6 Acid phosphatase reagent\(^{(8)}\)

<table>
<thead>
<tr>
<th>Ingredient</th>
<th>Quantity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetate buffer</td>
<td></td>
</tr>
<tr>
<td>Acetic acid</td>
<td>1.2 ml</td>
</tr>
<tr>
<td>Sodium acetate</td>
<td>1.6 g</td>
</tr>
<tr>
<td>Water</td>
<td>to 100 ml</td>
</tr>
</tbody>
</table>

Thoroughly mix the ingredients. The final pH value should be 4.6 ± 0.2.
Complete reagent

1-naphthyl phosphate disodium salt 0.4 g
o-dianisidine tetrazotized zinc chloride complex 0.8 g
(Fast Blue B)
Acetate buffer 20 ml

Add the ingredients to the acetate buffer and shake well to dissolve. Store the reagent at 5 ± 3 °C for one hour. Filter the solution to remove any precipitate. The reagent may be stored at 5 ± 3 °C for up to two weeks.

B8.7 Other media

Standard and commercial formulations of other media and reagents used in this method include zinc powder, quarter-strength Ringer’s solution and maximum recovery diluent.

B9 Analytical procedure

B9.1 Sample preparation

The volumes, and dilutions, of samples should be chosen so that the number of colonies to be counted on the membrane filter lies, if possible, between 20 and 80. With some waters, it may be advantageous to filter a selection of different volumes of sample so that the number of colonies on one of the membrane filters is likely to fall within this range. For treated waters, filter 100 ml of the sample. For polluted waters, either filter smaller volumes, or dilutions of the sample made with quarter-strength Ringer’s solution or maximum recovery diluent.

If it is the intention to count only the spores of Clostridium perfringens then the volume of sample should be heated to 60 ± 2 °C (for example in a water bath) and the whole volume maintained at this temperature for 15 ± 1 minutes. The temperature may be monitored by placing an appropriate thermometer in a similar bottle containing the same volume of water as the sample being treated.

B9.2 Sample processing

Place the sterile filtration apparatus in position and connect to a source of vacuum, with the stopcock turned off. Remove the funnel and, holding the edge of the membrane filter with sterile smooth-tipped forceps, place a sterile membrane filter, grid-side upwards, on the porous disc of the filter base. Replace the sterile funnel securely on the filter base. Pour or pipette the required volume of sample into the funnel. When the volume of sample to be filtered is less than 10 ml, add 10 - 20 ml of sterile diluent (for example, quarter-strength Ringer’s solution or maximum recovery diluent) to the funnel before addition of the sample. This aids dispersion of the bacteria over the entire surface of the membrane filter during filtration. Open the stopcock and apply a vacuum not exceeding 65 kPa (500 mm of mercury) and filter the sample slowly through the membrane filter. Close the stopcock as soon as the sample has been filtered.

Remove the funnel and transfer the membrane filter carefully to a Petri dish of well-dried tryptose sulphite cycloserine agar. Ensure that no air bubbles are trapped between the membrane filter and the medium. ‘Rolling’ the membrane filter onto the medium minimises the likelihood of air bubbles becoming trapped.
As the spores of *Clostridium perfringens* are very resilient, funnels that have been used once should be sterilised by autoclaving before being used again. Placing funnels in a water bath at this stage may not be sufficient to kill spores. If different volumes of the same sample are to be examined, the funnel may be re-used without sterilising the funnel provided that the smallest volume, or highest dilution of sample, is filtered first. For different samples, take a fresh pre-sterilised funnel and repeat the filtration process. During the filtration of a series of samples, the filter base need not be sterilised unless it becomes, or is suspected of being, contaminated or a membrane filter becomes damaged. When funnels are not in use they should be covered with a sterile lid or a sterile Petri dish lid.

The time between the end of the filtration step and the beginning of the incubation stage should be as short as possible and no longer than 2 hours.

Incubate the Petri dishes at 44 °C in an anaerobic jar or similar system containing an indicator of anaerobiosis and an atmosphere containing 9 - 13 % carbon dioxide. Examine the dishes after 21 ± 3 hours incubation.

**B9.3 Reading of results**

Under anaerobic conditions at 44 °C colonies of clostridia are typically black or grey in colour (see Figure B1). However, on occasion colourless colonies may be encountered. All colonies growing on tryptose sulphite cycloserine agar at 44 °C should, therefore, be counted as presumptive *Clostridium perfringens*.

![Figure B1 Typical colonies of *Clostridium perfringens* on tryptose sulphite cycloserine agar](image)

**B9.4 Confirmation tests**

Depending on the intended purpose of the analysis and the required accuracy, sub-culture a suitable number of colonies. If the aim is to estimate the number of organisms present, then for the greatest accuracy, all colonies should be sub-cultured if fewer than ten are present or, at least ten colonies should be sub-cultured if more than ten are present.
Clostridium perfringens can be confirmed by testing for reduction of nitrate, motility, fermentation of lactose and liquefaction of gelatin (i.e. the NMLG tests). Alternatively, Clostridium perfringens can be confirmed by testing for the production of acid phosphatase.

B9.4.1 Confirmation by the NMLG tests

For each isolate, inoculate a tube of buffered nitrate-motility medium by stabbing the medium with a straight wire or inoculator to just above the bottom of the tube and incubate anaerobically at 37 °C for 21 ± 3 hours. Growth of motile clostridia will be seen as cloudy growth throughout the medium. Growth of non-motile clostridia will be restricted to along the length of the stab (see Figure B2).

![Figure B2: Motility test reactions for non-motile Clostridium perfringens and motile Clostridium bifermentans in buffered nitrate-motility medium](image)

To test for nitrate reduction, add a few drops, approximately 0.2 - 0.5 ml, of the combined nitrate reduction test reagent to each tube. A red colour forming within 15 minutes indicates nitrate reduction to nitrite and the test is regarded as being positive (See Figure B3).

![Figure B3: Nitrate reduction test reactions for Clostridium perfringens (positive) and Clostridium bifermentans (negative) in buffered nitrate-motility medium](image)
If a red colour does not develop within 15 minutes, add a small amount of zinc powder and leave to stand for 10 minutes. If, after this time, there is still no red colour, this indicates that nitrate has been reduced to nitrite, which has been further reduced to nitrogen. The test is regarded as being positive. However, if a red colour subsequently develops after the addition of zinc powder, this indicates that nitrate has not been reduced and the test is regarded as being negative.

In addition, inoculate a tube of lactose-gelatin medium by stabbing the medium with a straight wire and incubate anaerobically at 37 °C for 44 ± 4 hours. After incubation, the medium will be liquid, irrespective of whether gelatin liquefaction has occurred or not. In order to establish whether gelatin liquefaction has occurred, the tube should be placed in a refrigerator for at least one hour. Gelatin liquefaction will have occurred in tubes where the medium remains liquid after refrigeration. If necessary, the tubes may be examined after incubating at 37 °C for 21 ± 3 hours and refrigerated (for example, for about one hour) and if gelatin liquefaction occurs, i.e. the test is regarded as positive, the result is recorded. If negative, i.e. the medium remains solid after refrigeration, the tubes should be returned to the incubator. Incubation should be continued until the total incubation period of 44 ± 4 hours has been achieved. The tubes are then re-examined.

A set of control tubes inoculated with appropriate positive and negative strains should be incubated and tested in parallel.

*Clostridium perfringens* is confirmed by the following reactions:

(i) Non-motile - growth along the line of the stab and not spread through buffered nitrate-motility medium.

(ii) Nitrate reduction - red colour after addition of combined nitrate reduction test reagent to buffered nitrate-motility medium, or remaining colourless after addition of zinc powder.

(iii) Lactose fermentation - orange/yellow colouration of lactose-gelatin medium.

(iv) Gelatin liquefaction - contents of the lactose-gelatin medium tube become liquefied.

Further identification may be carried out by means of appropriate biochemical and other tests. Suitable commercial identification kits may be used following appropriate performance verification at the laboratory.

B9.4.2 *Confirmation by the acid phosphatase test*

*Clostridium perfringens* can be confirmed by demonstration of production of acid phosphatase. Data on the verification of the performance of the acid phosphatase confirmation procedure are given in appendix B1.

Sub-culture presumptive colonies onto Columbia agar base and incubate anaerobically at 37 °C for 21 ± 3 hours. Place two or three drops of acid phosphatase reagent onto the growth. Development of a purplish or dark brown colour within three minutes is considered positive (see Figure B4).
Figure B4  Positive (*Clostridium perfringens*) and negative (*Clostridium bifermentans*) acid phosphatase reactions by dropping acid phosphatase reagent on colonies on Columbia agar base

Alternatively, soak a filter paper with the acid phosphatase reagent, transfer some of the colonies from the Columbia agar base and smear them onto the pre-soaked filter paper. The development of a purplish colour within three minutes is considered positive (see Figure B5).

Figure B5  Positive (*Clostridium perfringens*) and negative (*Clostridium bifermentans*) acid phosphatase reactions by streaking colonies from Columbia agar base onto filter paper soaked in acid phosphatase reagent

At the same time test positive (for example *Clostridium perfringens*) and negative (for example *Clostridium bifermentans*) controls. Isolates producing acid phosphatase are confirmed as *Clostridium perfringens*.

Further identification may be carried out by means of appropriate biochemical and other tests. Suitable commercial identification kits may be used following appropriate performance verification at the laboratory.
B10 Calculations

B10.1 Presumptive Clostridium perfringens

The number of presumptive Clostridium perfringens colonies is generally quoted as the number of colonies per 100 ml. Calculate the presumptive count as follows:

\[
\text{Presumptive count/100 ml} = \frac{\text{Number of colonies} \times 100 \times \text{DF}}{\text{Volume of sample filtered (ml)}}
\]

where DF is the appropriate dilution factor.

B10.2 Confirmed Clostridium perfringens

The number of confirmed Clostridium perfringens colonies is calculated by multiplying the number of presumptive Clostridium perfringens by the proportion of the isolates that are either non-motile, reduce nitrate, ferment lactose and liquefy gelatin, or produce acid phosphatase.

B11 Expression of results

The number of presumptive and confirmed Clostridium perfringens is expressed in colony forming units per volume of sample. For drinking waters, the volume is typically 100 ml.

B12 Quality assurance

New batches of isolation medium should be tested with appropriate reference strains of target bacteria (for example Clostridium perfringens) and non-target bacteria (for example Bacillus species). Petri dishes should be incubated for 21 ± 3 hours at 37 °C or 44 °C as appropriate. New batches of confirmatory media and reagents should be tested with appropriate reference strains of bacteria chosen to verify positive and negative reactions in each case. Further details of media and analytical quality control are given elsewhere\(^{(2)}\) in this series.

B13 References


Appendix 1 Verification of the acid phosphatase test for the confirmation of *Clostridium perfringens* isolated from various waters

1 Introduction

In a previous document\(^1\) in this series procedures are described for confirming presumptive *Clostridium perfringens* from membrane filters incubated on tryptone sulphite cycloserine (TSC) agar involving sub-culture to buffered nitrate-motility medium and lactose-gelatin medium (i.e. the NMLG tests) to test for nitrate reduction, motility, lactose fermentation and gelatin liquefaction. Isolates that reduce nitrate, are non-motile, ferment lactose and liquefy gelatin are considered to be confirmed as *Clostridium perfringens*.

An alternative method for confirming *Clostridium perfringens* based upon the demonstration of the production of acid phosphatase has been reported\(^2\) where the acid phosphatase test was reportedly more specific for *Clostridium perfringens* than the NMLG tests. However, some strains of other species of *Clostridium* were also found to be acid phosphatase-positive. Of 114 environmental isolates of *Clostridium perfringens*, 108 (i.e. 94.7 %) were acid phosphatase-positive compared to 104 (i.e. 91.2 %) that produced typical reactions in the NMLG tests\(^2\). Failure to reduce nitrate was the most common atypical result from the NMLG tests, which is in accordance with reports\(^3, 4\) that 10 % or more of strains of *Clostridium perfringens* are nitrate-negative.

A multi-laboratory study was therefore organised under the auspices of the Standing Committee of Analysts to assess the acid phosphate test for the confirmation of *Clostridium perfringens* and to demonstrate the equivalency of the method to the NMLG tests procedure\(^1\).

2 Materials and methods

Samples of a range of environmental water types were analysed according to procedures described in “The Microbiology of Drinking Water (2004) - Part 6 - Methods for the isolation and enumeration of Sulphite-Reducing Clostridia and *Clostridium perfringens* by membrane filtration”\(^1\) (using membrane filtration and enumeration on TSC agar). Following incubation, membranes filters exhibiting between 10 - 30 colonies were selected for confirmation, and colonies were counted and presumptive counts recorded.

Colonies of presumptive *Clostridium perfringens* were sub-cultured onto Columbia agar base and incubated anaerobically at 37.0 ± 1.0 °C for 21 ± 3 hours. For each isolate, a tube of buffered nitrate-motility medium was inoculated by stabbing with a straight wire or inoculator to just above the bottom of the tube and incubated anaerobically at 37 °C for 21 ± 3 hours. Testing for nitrate reduction was achieved by adding a few drops, approximately 0.2 - 0.5 ml, of the combined nitrate test reagent to each tube. A red colour forming within 15 minutes indicates nitrate reduction to nitrite and the test was regarded as positive. If a red colour did not develop within this time, a small amount of zinc powder was added and the tube left to stand for 10 minutes. If, after this time there was still no red colour, this indicated that the nitrate has been reduced to nitrite, which had been further reduced to nitrogen, and the test was, therefore, deemed positive. However, if a red colour subsequently developed after the addition of the zinc powder, this indicated that nitrate had not been reduced and the test was regarded as being negative. Motility was assessed as growth along the line of the stab spreading through buffered nitrate-motility medium.

In addition, a tube of lactose-gelatin medium was inoculated by stabbing with a straight wire and incubated anaerobically at 37 °C for a minimum of 21 ± 3 hours and a maximum of
44 ± 4 hours. After incubation the tubes were placed in a refrigerator for at least one hour. A positive gelatin liquefaction reaction was recorded for tubes where the medium remained liquid after refrigeration. Tubes examined after 21 ± 3 hours incubation that were negative, i.e. did not exhibit gelatin liquefaction were returned for further incubation until the total incubation period of 44 ± 4 hours had been achieved. The tubes were then re-examined.

Lactose fermentation was indicated by an orange/yellow colouration of the lactose-gelatin medium.

The remaining growth on the Columbia agar base plate was used for the acid phosphatase test. Two or three drops of acid phosphatase reagent were placed onto growth of each culture. The development of a purplish or brown colour within three minutes was considered as positive. Alternatively, a filter paper was soaked with the acid phosphatase reagent and some growth was smeared onto the filter paper. Development of a purplish colour within three minutes was considered as positive. At the same time positive (Clostridium perfringens) and negative (Clostridium bifermentans) controls were tested.

The participating laboratories were also requested to test a selection of strains giving unusual results by sub-culturing to cooked meat medium (BioMerieux), staining using Gram stain (clostridia being Gram-positive) and identified using API 20A miniaturised identification system (BioMerieux).

The water types examined ranged from surface freshwaters (for example, river, stream, canal and reservoir waters) groundwaters, raw sewage and sewage effluents and saline waters (marine and bathing beach).

3 Results and discussion

Fourteen laboratories participated, of which data from 13 laboratories were suitable for analysis (Table 1). Data were generated for 274 samples of surface freshwaters and similar waters (including one drinking water sample and one unclassified sample) 6 samples of groundwaters, 12 samples of raw sewage, 29 samples of sewage effluent and 4 samples of saline waters, i.e. a total of 325 samples.

Table 1 Numbers and types of waters and isolates of presumptive Clostridium perfringens analysed by 13 participating laboratories

<table>
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<tr>
<th>Laboratory</th>
<th>Fresh waters</th>
<th>Ground waters</th>
<th>Raw sewage</th>
<th>Sewage effluent</th>
<th>Saline waters</th>
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<th>Number of isolates</th>
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<tr>
<td>Total</td>
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<td>12</td>
<td>29</td>
<td>4</td>
<td>325</td>
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</table>
From these samples, 4146 isolates of presumptive *Clostridium perfringens* were tested by the NMLG confirmation tests and the acid phosphatase test, the data for which are summarised in a 2x2 matrix shown in Table 2. The data for each laboratory with respect to all possible combinations of the NMLG profiles and acid phosphatase reactions are summarised in Table 3.

Of the 4146 isolates, 3499 (i.e. 84.4 %) were acid phosphatase-positive, of which 3270 (i.e. 78.9 %) were confirmed as *Clostridium perfringens* according to the NMLG tests (the test profile being + − + + respectively) (see Table 2). Of the 647 (15.6 %) acid phosphatase-negative isolates, 462 (11.1 %) gave NMLG profiles other than that for *Clostridium perfringens*. Of the remaining isolates, 229 (5.5 %) were acid phosphatase-positive but did not confirm as *Clostridium perfringens* by the NMLG tests and 185 (4.5 %) were acid phosphatase-negative but confirmed as *Clostridium perfringens* by the NMLG tests. Thus, there is 90.0 % agreement between the NMLG tests and the acid phosphatase test for confirming *Clostridium perfringens*.

**Table 2** Summary of comparative results from NMLG tests and acid phosphatase test for presumptive *Clostridium perfringens* isolated from various waters

<table>
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<th>Acid phosphatase reaction</th>
<th>NMLG profile</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>+ − + +</td>
<td>3270 (78.9 %)</td>
<td>229 (5.5 %)</td>
</tr>
<tr>
<td></td>
<td>other</td>
<td>185 (4.5 %)</td>
<td>462 (11.1 %)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3455 (83.4 %)</td>
<td>691 (16.6 %)</td>
</tr>
</tbody>
</table>

Three laboratories (laboratories 8, 9 and 10) provided data on the identification of presumptive *Clostridium perfringens* isolates, principally targeting isolates that gave discrepant results between some of the NMLG tests and the acid phosphatase test. These are summarised in Table 4.

For 67 isolates that were confirmed as *Clostridium perfringens* by the NMLG tests (test profile being + − + + respectively), 33 (i.e. 49.3 %) were identified as *Clostridium perfringens*, of which 13 (19.4 %) were acid phosphatase-negative and 34 (50.7 %) were identified as species other than *Clostridium perfringens*. These were, principally, *Clostridium beijerinckii* / *Clostridium butyricum*, i.e. 26 (38.8 %) of the 67 isolates.

Additionally, 29 acid phosphatase-positive isolates, with non-*perfringens* NMLG profiles, were identified, of which 6 (20.7 %) were identified as *Clostridium perfringens*. Three of these isolates were nitrate-negative, one was gelatin-negative and one was negative for both tests. Thus, of the 39 isolates (i.e. 33 + 6) identified as *Clostridium perfringens*, 10.3 % were nitrate-negative. This figure agrees with figures reported elsewhere (2, 3, 4). Of the 29 non-*perfringens* acid phosphatase-positive isolates, 14 (48.3 %) were species of *Clostridium beijerinckii* / *Clostridium butyricum* and *Clostridium bifermentans*, i.e. being reported as 7 and 7 respectively.

Some NMLG profiles are at variance with the identification of the isolates. For example, two isolates of *Clostridium perfringens* are recorded as being motile (NMLG profiles of + + + + and − + + −). Additionally, the majority (26) of the non-*perfringens* isolates with the *Clostridium perfringens* NMLG profile (+ − + +) were identified as strains of *Clostridium beijerinckii* or *Clostridium butyricum*, despite these species being defined as being nitrate reduction-negative, motile and gelatin liquefaction-negative (3). Similarly, a further seven
strains were identified as *Clostridium bifermentans* although the species does not reduce nitrate nor ferment lactose\(^3\). This highlights problems with either reading results from the NMLG tests or limitations with identifying environmental isolates of *Clostridium* with the API 20A kit. Thus, the identifications reported in this study should be treated with caution.

Taking ratios from the submitted identification data, false-positive and false-negative rates for the two confirmation methods were calculated. However, caution is needed in the interpretation of the results as the isolates selected for identification were primarily taken from those that produced discrepant confirmation results, particularly with respect to the acid phosphatase test. This will skew any assessment of the data, especially as the targeted isolates represent strains from only about 10 % of all the isolates tested and the data are derived from only three of the thirteen participating laboratories. Taking this into account and applying the identification data to that shown in Table 3, the false-positive and false-negative rates of the NMLG and acid phosphatase test methods for the confirmation of *Clostridium perfringens* are:-

\[
\begin{align*}
\text{False-positive rate for NMLG tests} & = 3.9 \% \\
\text{False-positive rate for acid phosphatase test} & = 4.9 \% \\
\text{False-negative rate for NMLG tests} & = 1.3 \% \\
\text{False-negative rate for acid phosphatase test} & = 1.6 \%
\end{align*}
\]

The false-negative rate for the NMLG tests is lower than expected, particularly as 10.3 % of the strains identified as *Clostridium perfringens* did not reduce nitrate.

### 4 Conclusions

The results of this study indicate that the acid phosphatase test for the confirmation of *Clostridium perfringens* from water is at least as reliable as the current method\(^1\) based upon the demonstration of reduction of nitrate, lack of motility, fermentation of lactose and liquefaction of gelatin. The two procedures show similar false-positive and false-negative rates, at a level expected from application to a large number of a wide range of environmental isolates. There is an agreement rate of 90.0 %. The false-positive rates for both procedures are less than 5 % and appear to be primarily due to species of *Clostridium beijerinckii* or *Clostridium butyricum* and *Clostridium bifermentans*, although these identifications need to be treated with caution. The acid phosphatase test is considerably simpler to perform and is potentially more specific\(^2\).
### Table 3  NMLG reactions of acid phosphatase-positive and acid phosphatase-negative presumptive *Clostridium perfringens* isolates from various waters

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<td>13</td>
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<td>11</td>
<td>28</td>
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+ = Nitrate reduction-positive, motility-positive, lactose fermentation-positive and gelatin liquefaction-positive.

− = Nitrate reduction-negative, non-motile, lactose fermentation-negative and gelatin liquefaction-negative.
Table 4  Identification of isolates of presumptive *Clostridium perfringens* with respect to their NMLG profiles and acid phosphatase reactions

<table>
<thead>
<tr>
<th>NMLG profile</th>
<th><em>Clostridium</em> identification</th>
<th>Acid phosphatase positive</th>
<th>Acid phosphatase negative</th>
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<tr>
<td>+ – + +</td>
<td><em>Cl. perfringens</em></td>
<td>20</td>
<td>13</td>
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<tr>
<td></td>
<td><em>Cl. baratii / Cl. paraputrificum</em></td>
<td>2</td>
<td>24</td>
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<tr>
<td></td>
<td><em>Cl. beijerinckii / Cl. butyricum</em></td>
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<td>7</td>
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<tr>
<td>+ – + –</td>
<td><em>Cl. perfringens</em></td>
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<td>2</td>
</tr>
<tr>
<td></td>
<td><em>Cl. baratii</em></td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td><em>Cl. baratii / Cl. paraputrificum</em></td>
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<td>3</td>
</tr>
<tr>
<td></td>
<td><em>Cl. beijerinckii / Cl. butyricum</em></td>
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<td>1</td>
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<tr>
<td></td>
<td><em>Cl. histolyticum</em></td>
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<td><em>Cl. clostridioforme</em></td>
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<tr>
<td></td>
<td><em>Cl. ramosum</em></td>
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<td><em>Cl. tertium</em></td>
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<tr>
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<td><em>Cl. beijerinckii / Cl. butyricum</em></td>
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<td><em>Cl. histolyticum</em></td>
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<td><em>Cl. beijerinckii / Cl. butyricum</em></td>
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<td>– + + –</td>
<td><em>Cl. septicum</em></td>
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5 References


6 Acknowledgements

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