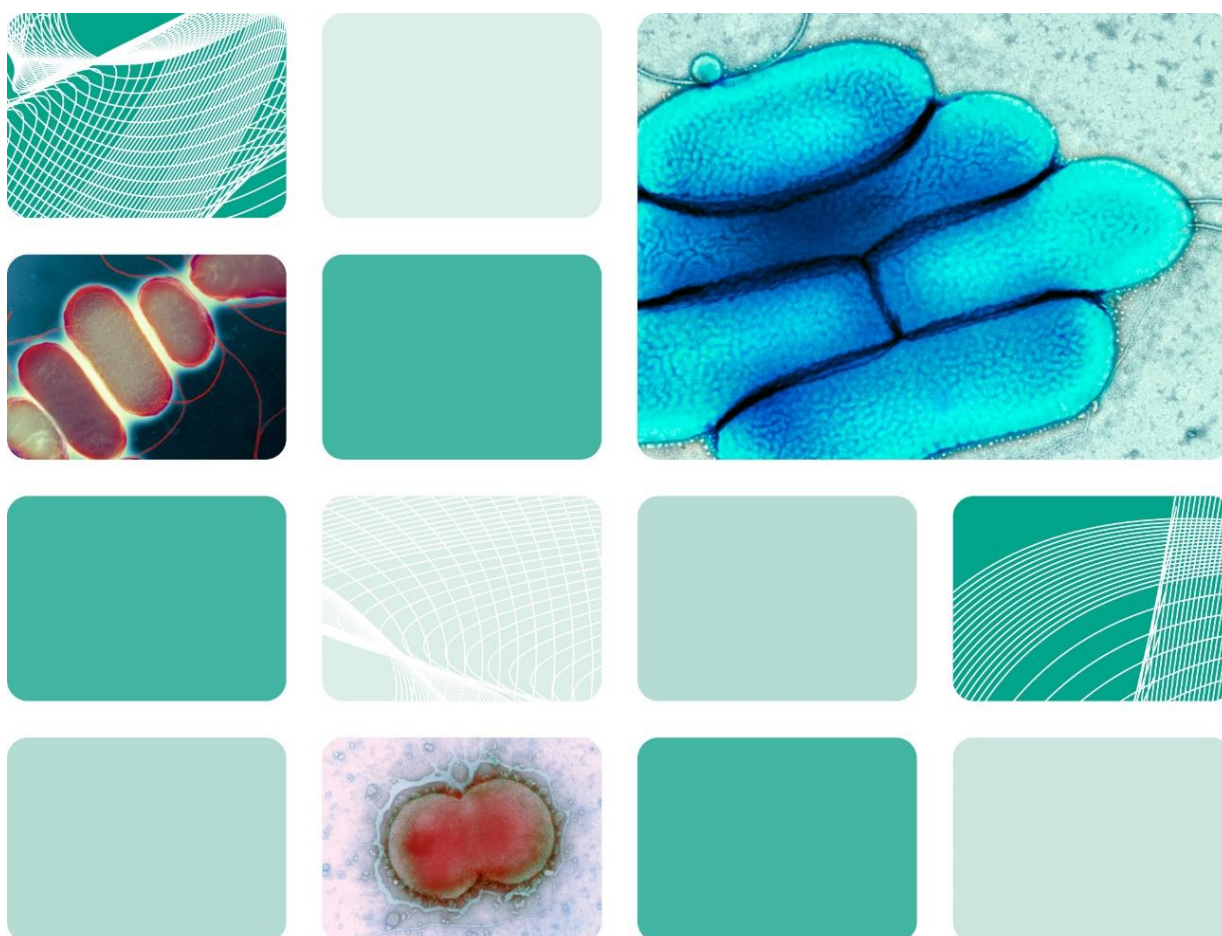




UK Health
Security
Agency

UK Standards for Microbiology Investigations

Identification of *Clostridium* species



Acknowledgments

UK Standards for Microbiology Investigations (UK SMIs) are developed under the auspices of UKHSA working in partnership with the partner organisations whose logos are displayed below and listed on [the UK SMI website](#). UK SMIs are developed, reviewed and revised by various working groups which are overseen by a [steering committee](#).

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Amendment table

Each UK SMI document has an individual record of amendments. The amendments are listed on this page. The amendment history is available from standards@ukhsa.gov.uk.

Any alterations to this document should be controlled in accordance with the local document control process.

Amendment number/date	8/11.07.25
Issue number discarded	4.1
Insert issue number	4.2
Section(s) involved	Amendment
Whole document.	<p>This is an administrative point change.</p> <p>The content of this UK SMI document has not changed.</p> <p>The last scientific and clinical review was conducted on 12/01/15.</p> <p>Hyperlinks throughout document updated to Royal College of Pathologists website.</p> <p>Public Health England replaced with UK Health Security Agency throughout the document, including the updated Royal Coat of Arms</p> <p>Partner organisation logos updated.</p> <p>Broken links to devolved administrations replaced.</p> <p>References to NICE accreditation removed.</p> <p>Scope and Purpose replaced with General and Scientific information to align with current UK SMI template.</p>

Amendment No/Date.	7/01.03.16
Issue no. discarded.	4
Insert Issue no.	4.1
Section(s) involved	Amendment
Section 4.	Error in flowchart corrected.

Amendment No/Date.	6/12.01.15
Issue no. discarded.	3.2
Insert Issue no.	4
Section(s) involved	Amendment
Whole document.	Hyperlinks updated to gov.uk.
Page 2.	Updated logos added.
Whole document.	Document presented in a new format. Reorganisation of some text. Edited for clarity. Information regarding <i>Clostridium difficile</i> updated. Test procedures updated. Removal of Reference Laboratory contact details.
Scope of document.	The scope has been updated to include webpage link for B 10 document.
Introduction.	The taxonomy of <i>Clostridium</i> species has been updated. More information has been added to the Characteristics section. The medically important species are mentioned and their characteristics described. Use of up-to-date references. Section on Principles of Identification has been updated to inform the users of the appropriate reference laboratory to send <i>Clostridium</i> species for further identification.
Technical Information/Limitations.	Addition of information regarding Gram stain, antibiotic susceptibility, sporulation and commercial identification systems has been described and referenced.
Safety considerations.	Update on Laboratory-acquired infections with references.
Target Organisms.	The section on the Target organisms has been updated and presented clearly. References have been updated.

Identification of *Clostridium* species

Identification.	Amendments have been done on 3.1, 3.2, 3.3 and 3.4 have been updated to reflect standards in practice. Subsection 3.5 has been updated to include the Rapid Molecular Methods. 3.6 has been rephrased and informs users to refer to appropriate laboratory user manual for referrals.
Identification Flowchart.	Modification of flowchart for identification of species has been made for easy guidance.
Reporting.	Subsection 5.2 and 5.6 has been updated to reflect reporting practice.
Referral.	The address of the reference laboratories has been updated.
References.	Some references updated.

*Reviews can be extended up to 5 years where appropriate

1 General information

[View general information](#) related to UK SMIs.

2 Scientific information

[View scientific information](#) related to UK SMIs.

3 Scope of document

This UK SMI describes the identification of *Clostridium* species.

There are many species of clostridia which may be found naturally in the environment and animal faeces. Only species associated with human infection will be discussed in this UK SMI.

For more information on *Clostridium difficile*, refer to [UK SMI B 10 - Investigation of Faecal Specimens for *Clostridioides difficile*](#).

This UK SMI should be used in conjunction with other UK SMIs.

4 Introduction

4.1 Taxonomy

The genus *Clostridium* belongs to the family *Clostridiaceae* and it currently contains 203 species and 5 subspecies, with only a few species being pathogenic to humans. Of these species, 21 have been reclassified to other genera, 5 have been reclassified within the genus and 1 has been de-accessioned¹.

In 1994 the heterogeneity of this species was confirmed by 16S rRNA gene sequencing². This has been reaffirmed by the work of Yutin *et al* that 16S rRNA and ribosomal protein sequences are better indicators of evolutionary proximity than phenotypic traits. This genus like several others has undergone a number of revisions with the increasing availability of genomic data. An analysis of proteins from a number of members of this genus suggested another revision³. The main findings from the proposal suggested that:

- The *Selenomonas-Megasphaera-Sporomusa* group are still members of the genus *Clostridium*
- *Clostridium difficile* and its close relatives are placed within the family *Peptostreptococcaceae*. Under this proposal, the species *Clostridium difficile* would become *Peptoclostridium difficile*
- Members of the family *Ruminococcaceae* belong to the genus *Clostridium*

- It was also proposed to create six new genera to accommodate the 78 validly described species that fell outside the family *Clostridiaceae*. These genera are: *Erysipelatoclostridium*, *Gottschalkia*, *Lachnoclostridium*, *Peptoclostridium*, *Ruminiclostridium* and *Tyzzzeria*

The type species is *Clostridium butyricum*.

4.2 Characteristics

Clostridium are phylogenetically heterogeneous and are Gram positive but can decolourise easily and appear Gram negative or Gram variable, spore formers and non-spore formers, rods and cocci and anaerobic and non-anaerobic bacteria⁴.

Medically significant *Clostridium* strains tend to be Gram positive rods (some are Gram variable), 0.3 – 2.0 x 1.5 – 20.0µm which are often arranged in pairs or short chains, with rounded or sometimes pointed or square ends. They are commonly pleomorphic and vary considerably in their oxygen tolerance. Some species such as *Clostridium novyi* type A and *Clostridium haemolyticum* may require extended incubation on pre-reduced or freshly prepared plates and total handling in an anaerobic chamber. Conversely, *Clostridium tertium*, *Clostridium histolyticum* and *Clostridium carnis* are aerotolerant and will form colonies on blood agar plates incubated in an atmosphere of air with 5-10% added CO₂⁵.

Virtually all of the members of the genus, except *Clostridium perfringens*, are motile with peritrichous flagellae and form oval or spherical endospores that may distend the cell. They may be saccharolytic or proteolytic and are usually catalase negative. Many species produce potent exotoxins.

Toxins of *Clostridium* species

Clinically significant *Clostridium* species produce a variety of toxins. It is the production of these toxins which leads to the distinctive clinical features of the diseases they cause, eg tetanus and botulism result from the production of neurotoxins that are amongst the most lethal substances known to man⁶. Clostridial toxins are biologically active proteins that are antigenic in nature and can therefore be neutralised with specific antisera.

Detection of particular toxins directly from some clinical samples may render the isolation of the organism unnecessary for primary investigation eg *C. difficile* (refer to [UK SMI B 10 - Investigation of Faecal Specimens for *Clostridioides difficile*](#)). Culture is required for typing (outbreaks and incidents) and susceptibility testing.

Clostridium perfringens is the most commonly isolated *Clostridium* species. Five types (A-E) may be distinguished by the combinations of major lethal toxins they produce⁵.

Clostridium tetani produces two exotoxins, tetanolysin and tetanospasmin – Tetanolysin causes lysis of RBCs and serves no known benefit to *C. tetani* infections while tetanospasmin is a neurotoxin that causes the clinical manifestations of tetanus⁶.

Clostridium botulinum also produces neurotoxins (which are the most potent natural poisons known) that cause botulism, a disease characterized by a symmetrical, descending paralysis⁷. There are seven toxin types (A-G), man is susceptible to type A, B, E, F and G toxins; types A, B, C and D cause intoxication in animals. Although less common, bivalent strains that express two different toxin types exist and are

designated by the predominant toxin produced. Strains of *C. baratii* and *C. butyricum* have been implicated as causative agents of botulism as they also produce the types F and E respectively^{6,8}. *C. argentinense* (formerly *C. botulinum* type G) produce botulinum neurotoxin.

Clostridium difficile is the most common toxigenic *Clostridium* species. They produce two potent exotoxins namely – Toxin A (enterotoxin) and toxin B (cytotoxic activity)⁶.

Clostridium novyi comes in three types, labelled A, B, and a non-pathogenic type C distinguished by the range of toxins they produce. The toxins are designated by Greek letters⁶.

Clostridium sordellii produces three toxins in common with non-pathogenic *C. bifermentans* namely; a lecithinase, an oxygen-labile haemolysin and a fibrinolysin. It also has a major lethal toxin referred to as beta-toxin that distinguishes it from *C. bifermentans*. This beta-toxin actually contains two toxins: lethal toxin (LT) and haemorrhagic toxin (HT)⁶.

Other *Clostridium* species produce similar toxins to that produced by *C. perfringens*.

The medically important species are:

Clostridium perfringens

They are non-motile straight-sided gram-variable rods with blunt ends that occur singly or in pairs, 0.6 – 2.4µm wide by 1.3 – 19.0µm long, and rarely produce spores. They grow vigorously at temperatures between 20 and 50°C, with an optimum of 45°C for most strains. On blood agar, large discrete colonies are produced after overnight incubation. They may be flat and rough-edged or smooth and domed, and either non-haemolytic or with a narrow zone of complete haemolysis inside a larger zone of partial haemolysis. Haemolysis is more pronounced on sheep blood agar than on horse blood agar⁹. They are positive for lecithinase, nitrate, and fermentation of sugars but negative for lipase, indole and urease tests.

Clostridium tetani

Cells are 0.5 – 1.7 by 2.1 – 18.1µm and often possess terminal endospores that give a “drumstick” appearance. Cells in culture older than 24hr may appear Gram negative. They are also motile by peritrichous flagella. The optimal growth temperature is 37°C and little or no growth takes place at 25 or 42°C.

Growth may appear as a film rather than discrete colonies because of swarming due to the vigorous motility after 48hr incubation. On blood agar, the colonies are flat, translucent, and grey with a matte surface, showing a zone of β- haemolysis and are 4 to 6mm in diameter. Colonies have irregular and rhizoid margins. They are negative for fermentation of sugars, lecithinase, lipase, urease, nitrate reduction tests but give variable results for indole test.

Clostridium botulinum

Cells are gram variable bacilli that show profuse sub-terminal and free spores. The proteolytic types A, B and F initially produce discrete rhizoidal colonies that spread and coalesce. Haemolysis is variable, but the odour is strong and redolent of rotten eggs due to production of H₂S. They are positive for lipase but negative for indole and urease tests. They give variable test results for lecithinase reaction.

They have been isolated from clinical samples such as – faeces, wounds, tissue, and pus as well as from foods.

Clostridium difficile

Cells are motile rods, with dimensions of 0.5 -1.9 by 3.0 – 16.9µm, which forms oval sub-terminal spores and show optimum growth on blood agar at human body temperatures in the absence of oxygen. Colonies of *C. difficile* are 4 - 6mm in diameter, irregular, raised, opaque, and grey-white after 48hr incubation. They may be isolated from faecal specimens using cycloserine cefoxitin fructose agar (CCFA) or cycloserine cefoxitin egg yolk agar (CCEY). They ferment sugars but are negative for lecithinase, lipase and indole tests.

Refer to [UK SMI B 10 - Investigation of Faecal Specimens for *Clostridioides difficile*](#) for further information.

Clostridium novyi

Cells are motile, gram variable rods with occasional sub-terminal spores. Cell dimensions are 0.5 – 1.6 by 1.6 – 18µm except for *C. novyi* type B, which are larger, 1.1 – 2.5 by 3.3 – 22.5µm. Isolating and identifying *C. novyi* is difficult due to its extreme anaerobic nature. Because of their fastidious nature and difficulty in culturing, they require the presence of thiols for growth¹⁰. Growth is stimulated by fermentable carbohydrates, serum or peptic digest of blood. On blood agar after overnight incubation anaerobically, colonies appear as small, flat, rough or rhizoidal, translucent, haemolytic colonies with a spreading edge and 1 - 5mm in diameter and after incubation for 48–72hr, colonies will often coalesce to give a fine spreading growth that may cover the entire plate, often with a marked β-haemolysis so as to make the blood agar plate completely transparent. There is poor growth in nutrient broth or cooked meat broth. They ferment glucose and liquefy gelatin. Proteolytic activity is variable. They are positive for lecithinase and lipase reactions but give variable results for indole test.

C. novyi type A is usually unreactive in commercial anaerobe identification kits and commonly is not identified by this approach. *C. novyi* type B has different phenotypic characteristics and can be distinguished by its biochemical reactions⁹.

Clostridium sordellii

Colonies are large, grey-white and irregular, sometimes with a “fern-leaf” edge. They produce indole and lecithinase as well as ferment sugars. They are also urease positive, which differentiates them from *C. bifermentans*, generally regarded as a non-pathogen.

Clostridium septicum

Cells are gram variable rods with numerous sub-terminal spores. On blood agar, they grow rapidly and usually produce a thick haemolytic swarming growth. In culture, it has no characteristic odour. They are negative for lecithinase, lipase, indole and urease tests.

They are easily recognised by use of commercially available identification kits. The most common source of *C. septicum* isolates seen in recent years has been from blood cultures from patients with malignancies of the colon or caecum⁹.

4.3 Principles of Identification

Clues to the identity of certain pathogenic species may be obtained by observing characteristics such as colonial appearance, Gram stain appearance and the presence or absence of β -haemolysis. Other phenotypic tests may also be applied to obtain a presumptive identification¹¹. It is important to ensure the culture is pure, as the fine spreading growth of some *Clostridium* species may mask contaminating organisms.

If confirmation of *Clostridium* species is required, isolates should be referred to the Anaerobe Reference Unit, Public Health Wales, Cardiff.

If *C. difficile* confirmation is required, refer to [UK SMI B 10 - Investigation of Faecal Specimens for *Clostridioides difficile*](#).

If *C. botulinum* is suspected, samples of patient's serum, faeces and implicated foodstuff should be referred directly to the Foodborne Pathogens Reference Section, Colindale.

5 Technical Information/Limitations

Antibiotic susceptibility

Reduced susceptibility of *C. difficile* to metronidazole has been demonstrated¹².

Sporulation

Several species of *Clostridium*, including *C. carnis*, *C. histolyticum* and *C. tertium* can grow, but not sporulate, in air⁵.

Gram stain

It is important to ensure the culture is pure, as the fine spreading growth of some *Clostridium* species may mask contaminating organisms.

There can be failure to determine the Gram reaction correctly (many anaerobes over decolourise and appear Gram negative). For example, Gram negatively staining *Clostridium* species, especially *C. clostridioforme*, can be misidentified as *Bacteroides*¹¹.

Commercial identification Systems

The use of commercially available anaerobe identification kits alone may not give an accurate identification eg *C. novyi* type A is usually unreactive in commercial anaerobe identification kits and commonly is not identified by this approach. *C. novyi* type B has different phenotypic characteristics and can be distinguished by its biochemical reactions⁹.

6 Safety Considerations¹³⁻²⁹

Hazard Group 2 organisms

Laboratory acquired infections have been reported^{30,31}.

Refer to current guidance on the safe handling of all Hazard Group 2 organisms documented in this UK SMI.

Laboratory procedures that give rise to infectious aerosols must be conducted in a microbiological safety cabinet²¹.

The above guidance should be supplemented with local COSHH and risk assessments.

Compliance with postal and transport regulations is essential.

7 Target Organisms

Clostridium species Reported to have Caused Human Disease^{5,11}

C. perfringens, *C. septicum*, *C. novyii* type A, *C. sordellii*, *C. tetani*, *C. difficile*, *C. botulinum*, *C. butyricum*, *C. baratii*, *C. tertium*, *C. histolyticum*

Non-pathogenic Clostridium species Commonly Isolated that may have Caused Human Infections^{5,32,33}

C. sporogenes, *C. ramosum*, *C. innocuum*, *C. paraputrificum*, *C. cadaveris*, *C. bifermentans*, *C. fallax*, *C. clostridioforme*

8 Identification

8.1 Microscopic Appearance

([UK SMI TP 39 - Staining Procedures](#))

Gram stain

Clostridium species are Gram positive rods, which may possess a single endospore. Some species may be Gram variable.

Spore stain

This is used to determine the shape and position of the spore (phase contrast microscopy is an alternative option).

C. perfringens Oval, subterminal

Note: *C. perfringens* spores are rarely seen in vivo or usual in vitro conditions. They do not sporulate on normal agar media. *C. perfringens* also have non-spore forming strains.

C. botulinum Oval, subterminal

<i>C. difficile</i>	Oval, subterminal
<i>C. novyi</i>	Oval, central or subterminal
<i>C. sordellii</i>	Oval, subterminal
<i>C. septicum</i>	Oval, subterminal
<i>C. tetani</i>	Round, terminal (giving a drumstick appearance)

8.2 Primary Isolation Media

Agar containing blood incubated anaerobically at 35-37°C for 40–48hr.

Egg Yolk agar incubated anaerobically at 35-37°C for 16-24hr.

If culturing for toxigenic *C. difficile*, Cycloserine Cefoxitin Fructose agar (CCFA) or Cefoxitin Cycloserine Egg Yolk agar (CCEY) should also be inoculated and incubated anaerobically at 35-37°C for 24 - 48hr. The antibiotics cycloserine and cefoxitin inhibit the growth of most bacteria other than *C. difficile*¹¹.

8.3 Colonial Appearance

Colonial appearance varies with species and brief descriptions of the most common species are given here:

Organism	Characteristics of growth on agar containing blood after anaerobic incubation at 35–37°C for 40–48hr
<i>C. botulinum/sporogenes</i>	Large (3mm), irregularly circular, smooth, greyish, translucent with a fibrillar edge that may spread. Most strains are β -haemolytic; produces lipase.
<i>C. difficile</i>	Glossy, grey, circular colonies with a rough edge; fluoresce green-yellow under long wavelength UV light (360 nm \pm 20nm). They are usually non-haemolytic, with a characteristic farmyard smell.
<i>C. novyi</i>	Raised, circular colonies, which become flattened and irregular in old cultures. Colonies tend to fuse forming a spreading growth with a double zone of β -haemolysis. Type A produces lecithinase and lipase.
<i>C. perfringens</i>	Large, smooth, regular convex colonies, but may be rough and flat with an irregular edge. Usually has a double zone of β -haemolysis; produces lecithinase.
<i>C. septicum</i>	Usually produce a thick swarming growth with a narrow zone of β -haemolysis.
<i>C. sordellii/bifermentans</i>	Grey-white, convex, circular colonies with crenated edges, which may spread. They may be β -haemolytic; produce lecithinase; indole positive.
<i>C. tetani</i>	Fine swarming growth (may be difficult to see) which may appear β -haemolytic.
Other <i>Clostridium</i> species	Colonial appearances vary, but may produce a spreading growth which may or may not be β -haemolytic.

8.4 Test Procedures

The following tests can be used to differentiate between *Clostridium* species. If clinically indicated, refer to the appropriate Anaerobe Reference Unit for further identification.

8.4.1 Biochemical tests

Nagler test [UK SMI TP 22 - Nagler Test](#)

The nagler test determines the ability of a microorganism to produce the enzyme lecithinase. Lecithinase producing organisms are identified by a zone of opalescence surrounding individual colonies on egg yolk agar. *C. perfringens* lecithinase is inhibited by the antitoxin *C. perfringens* type A.

Clostridium baratii, *Clostridium absonum*, *Clostridium bifermentans*, *Clostridium sordelli* and *Clostridium novyi* also produce lecithinase. *C. sordelli* and *C. bifermentans* produce enzymes that are also closely related to *C. perfringens* alpha toxin (lecithinase) and can produce a partial cross-reaction³⁴.

A Nagler positive result is indicated by lecithinase production and inhibition due to antitoxin.

Note: In recent years, popularity of the Nagler test has declined because the antitoxin has not been widely available. An alternative to the Nagler test used in some laboratories is the reverse CAMP test.

Reverse CAMP test

Reverse CAMP test can be used for differentiation of *C. perfringens* from other *Clostridium* species. Alpha toxin producing *C. perfringens* and group B, β -haemolytic streptococci grow in a characteristic pattern on blood agar; however care must be taken to ensure pure cultures are used³⁵.

Indole test [UK SMI TP 19 - Indole Test](#)

The indole test determines the ability of an organism to produce indole from the degradation of the amino acid tryptophan.

Anaerobes, particularly *Clostridium* species, form indole but can rapidly break it down as it is produced; therefore, false negative reactions may occur³⁶.

C. novyi A strains give variable indole test results but are usually indole negative.

Lipase test

The lipase test determines the ability of microorganisms to produce the enzyme lipase that catalyses the hydrolysis of triglycerides and diglycerides to fatty acids and glycerol. This is shown by the iridescent sheen on and surrounding colonies on plate medium. This aids in differentiation of *Clostridium* species.

C. botulinum, *C. sporogenes*, *C. novyi* A, *C. ghonii* and *C. cochlearium* produce lipase. *C. leptum* give variable lipase reactions but are usually lipase negative.

Urease test [UK SMI TP 36 - Urease Test](#)

The urease test is used to determine the ability of an organism to split urea, through the production of the enzyme urease.

C. sordellii are urease positive which distinguishes it from *C. bifermentans*, which it resembles and are urease negative.

8.4.2 Commercial identification Systems

Laboratories should follow manufacturer's instructions and rapid tests and kits and should be validated and be shown to be fit for purpose prior to use.

8.4.3 Matrix-Assisted Laser Desorption/Ionisation - Time of Flight (MALDI-TOF)

Matrix-assisted laser desorption ionization–time-of-flight mass spectrometry (MALDI-TOF MS), which can be used to analyse the protein composition of a bacterial cell, has emerged as a new technology for species identification. This has been shown to be a rapid and powerful tool because of its reproducibility, speed and sensitivity of analysis. The advantage of MALDI-TOF as compared with other identification methods is that the results of the analysis are available within a few hours rather than several days. The speed and the simplicity of sample preparation and result acquisition associated with minimal consumable costs make this method well suited for routine and high-throughput use³⁷.

This has been used for the identification of *Clostridium* species especially to discern different ribotypes among isolates of *C. difficile*. However, an extensive database is essential to identify species and closely related strains reliably and available databases needs to be optimised^{38,39}.

Other limitations to the use of this technique is the presence of spores of *Clostridium* species and so younger cultures are now used to minimize spectral interference³⁹.

8.4.4 Nucleic Acid Amplification Tests (NAATs)

PCR is usually considered to be a good method for bacterial detection as it is simple, rapid, sensitive and specific. The basis for PCR diagnostic applications in microbiology is the detection of infectious agents and the discrimination of non-pathogenic from pathogenic strains by virtue of specific genes. However, it does have limitations. Although the 16S rRNA gene is generally targeted for the design of species-specific PCR primers for identification, designing primers is difficult when the sequences of the homologous genes have high similarity.

This has been used successfully in the identification of *Clostridium* species eg *C. perfringens*, *C. botulinum*, *C. baratii* and *C. butyricum*, *C. novyi*, *C. difficile*⁴⁰⁻⁴⁴.

8.5 Further Identification

Rapid Methods

A variety of current rapid typing methods have been developed for isolates from clinical samples; these include molecular techniques such as Pulsed- Field Gel Electrophoresis (PFGE), Fluorescent Amplified Fragment Length Polymorphism (AFLP), 16S rDNA gene sequencing, PCR- restriction fragment length Polymorphism

(PCR-RFLP), Microarray analysis, Multiple-Locus Variable-Number Tandem-Repeat Analysis (MVLA), and even whole-genome sequencing (WGS). All of these approaches enable subtyping of strains, but do so with different accuracy, discriminatory power, and reproducibility.

However, some of these methods remain accessible to reference laboratories only and are difficult to implement for routine bacterial identification in a clinical laboratory.

Fluorescent Amplified Fragment Length Polymorphism (AFLP)

Fluorescent Amplified Fragment Length Polymorphism is a high-resolution whole genome methodology used as a tool for rapid and cost-effective analysis of genetic diversity within bacterial genomes. It is useful for a broad range of applications such as identification and subtyping of microorganisms from clinical samples, for identification of outbreak genotypes, for studies of micro and macro-variation, and for population genetics.

FAFLP has numerous advantages over other DNA fingerprinting techniques because it assesses the whole genome for both conserved and rapidly evolving sequences in a relatively unbiased way. The number of fragments obtained for comparative purposes between isolates is significantly greater than pulsed-field gel electrophoresis (PFGE), thus making it more discriminatory than PFGE and the FAFLP results are highly reproducible due to stringent PCR cycling parameters.

This relatively fast method can be applied to different clostridia and used for the generation of identification libraries. Libraries of AFLP profiles of well-defined *Clostridium* strains provide a valuable additional tool in the identification of *Clostridium* species.

This technique has been used to genotype *C. botulinum*, *C. difficile*, *C. novyi* and *C. perfringens*⁴⁵⁻⁴⁸. It has also been used to differentiate between *C. bifermentans* and *C. sordellii* strains (which closely resemble phenotypically) and between strains of *C. perfringens*⁴⁹.

Pulsed Field Gel Electrophoresis (PFGE)

PFGE detects genetic variation between strains using rare-cutting restriction endonucleases, followed by separation of the resulting large genomic fragments on an agarose gel. PFGE is known to be highly discriminatory and a frequently used technique for outbreak investigations. However, the stability of PFGE may be insufficient for reliable application in long-term epidemiological studies. Due to its time-consuming nature (30hr or longer to perform) and its requirement for special equipment and the interpretation of its results often being subjective, PFGE is not used widely outside reference laboratories^{50,51}.

PFGE is considered a very useful tool for molecular epidemiological analysis of proteolytic *C. botulinum* types A and B as it enabled discrimination between them but this has not been very successful with non-proteolytic *C. botulinum*⁵². It has been used for typing *C. difficile* although a considerable proportion of strains are non-typable by this technique due to degradation of the DNA during the procedure; making uninterpretable gel smears or likely spore formation⁵³.

PFGE has also been used to establish *C. perfringens* as the etiological agent in food-borne outbreaks and to reveal its wide genetic diversity from different sources⁵⁴.

16S rDNA gene sequencing analysis

A genotypic identification method, 16S rDNA gene sequencing is used for phylogenetic studies and has subsequently been found to be capable of re-classifying bacteria into completely new species, or even genera. It has also been used to describe new species that have never been successfully cultured.

This has been used to differentiate between *Clostridium* species eg *C. novyi* type A and *C. botulinum* type C that are closely related⁹.

Microarrays

DNA microarray technology can provide detailed, clinically relevant information on the isolate by detecting the presence or absence of a large number of virulence-associated genes simultaneously in a single assay; however, their clinical value has been limited by a complicated methodology that is unsuitable for routine use in diagnostic microbiology laboratories.

This technique has been used and it demonstrates the high-throughput detection and identification of pathogenic *Clostridium* species and it has advantages over the conventional traditional methods. This has also been particularly useful in efficiently and specifically identifying all *Clostridium* species present in a mixed bacterial population. The high-throughput feature of this technique is very useful in the detection and analysis of outbreak strains and for epidemiologic studies of *Clostridium* infections⁷.

Multiple-Locus Variable Number Tandem Repeat Analysis (MVLA)

Multiple-Locus Variable number tandem repeat Analysis (MLVA) is a method used to perform molecular typing of particular microorganisms. It utilizes the naturally occurring variation in the number of tandem repeated DNA sequences found in many different loci in the genome of a variety of organisms. The molecular typing profiles are used to study transmission routes, to assess sources of infection and also to assess the impact of human intervention such as vaccination and use of antibiotics on the composition of bacterial populations.

This has been used successfully in the typing of *C. perfringens* and newly emerging variants of *C. difficile*^{55,56}.

Whole Genome Sequencing (WGS)

This is also known as full genome sequencing, complete genome sequencing, or entire genome sequencing. It is a laboratory process that determines the complete DNA sequence of an organism's genome at a single time. There are several high-throughput techniques that are available and used to sequence an entire genome such as pyrosequencing, nanopore technology, illumina sequencing, ion torrent sequencing, etc. This sequencing method holds great promise for rapid, accurate, and comprehensive identification of bacterial transmission pathways in hospital and community settings, with concomitant reductions in infections, morbidity and costs.

This has been used successfully to explore the phylogeny, horizontal gene transfer, recombination, and micro and macroevolution of the major hospital-acquired pathogen, *C. difficile* as well as proteolytic *C. botulinum* and *C. perfringens*⁵⁷⁻⁵⁹.

8.6 Storage and Referral

If further identification is required, refer to the appropriate reference laboratory user manual for details on referral.

Frozen storage (-20°C) of toxin positive faecal samples is recommended for retrospective culture should the need for further investigation arise^{60,61}.

9 Reporting

9.1 Presumptive Identification

If appropriate growth characteristics, colonial appearances and Gram stain of the culture are demonstrated and the isolate is metronidazole susceptible.

9.2 Confirmation of Identification

Following identification processes as outlined in this document and/or Reference Laboratory report.

9.3 Medical Microbiologist

Inform the medical microbiologist of all positive cultures from normally sterile sites.

According to local protocols, the medical microbiologist should also be informed of a presumptive and confirmed *Clostridium* species. When the request card bears relevant information eg:

- Cases of trauma, penetrating injury, compound fracture or retained foreign body, or known injecting drug abuse (especially heroin)
- Septic abortion
- Suspicion of clostridial myonecrosis, (necrotising) myofasciitis, surgical wound infection (especially in cases with occlusive peripheral vascular disease and/or diabetes mellitus)
- Other serious medical conditions eg alcohol or substance abuse, immunodeficiency, cancer, or persons receiving treatment for cancer (including neutropenia and/or mucositis)
- Food poisoning (especially involving descending paralysis with cranial nerve involvement) and/or consumption of unusual or imported foods (suspicion of botulism)
- Investigation of outbreaks
- Pseudomembranous colitis or antibiotic related diarrhoea
- Suspicion of tetanus

Follow local protocols for reporting to clinician.

9.4 CCDC

Refer to local Memorandum of Understanding.

9.5 UK Health Security Agency⁶²

Refer to current guidelines on CIDSC and COSURV reporting.

9.6 Infection Prevention and Control Team

Inform the infection prevention and control team of presumptive and confirmed isolates of *C. botulinum* and *C. difficile* according to local protocols.

10 Referral to reference laboratories

For information on the tests offered, turnaround times, transport procedure and the other requirements of the reference laboratory [see user manuals and request forms](#)

Contact appropriate reference laboratory for information on the tests available, turnaround times, transport procedure and any other requirements for sample submission:

[England](#)

[Wales](#)

[Scotland](#)

[Northern Ireland](#)

Note: In case of sending away to laboratories for processing, ensure that specimen is placed in appropriate package and transported accordingly.

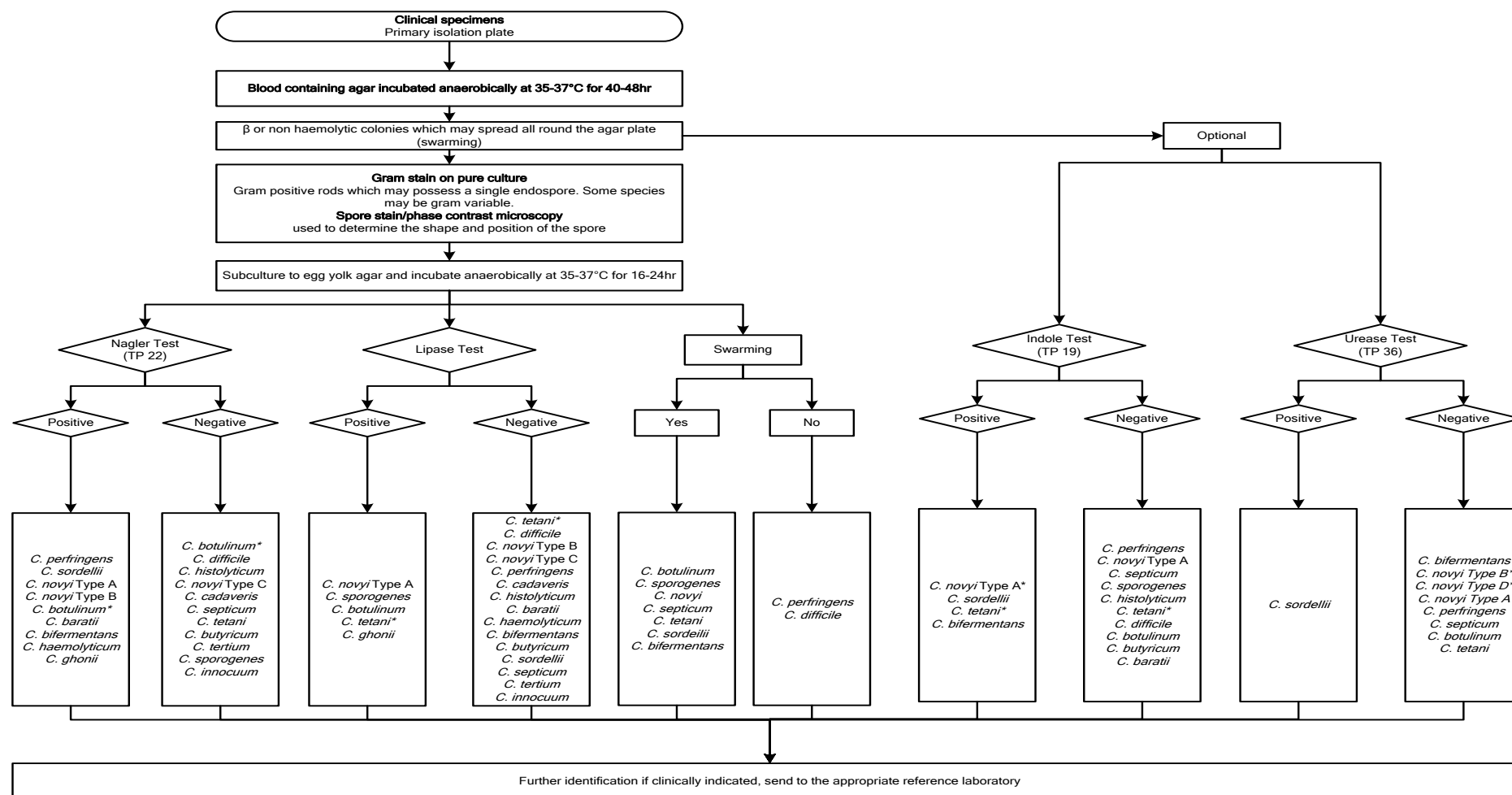
11 Public Health responsibilities of diagnostic laboratories

Diagnostic laboratories have public health responsibility as part of their duties. Amongst these are additional local testing, or referral, to further characterise the organism, as required, primarily for public health purposes e.g. routine cryptosporidium detection; serotyping or microbial subtyping; and a duty to refer appropriate specimens and isolates of public health importance to a reference laboratory.

Diagnostic laboratory outputs inform public health intervention, and surveillance data is required to develop policy and guidance, forming an essential component of healthcare. It is recognised that additional testing and referral of samples may entail some costs that has to be borne by the laboratory but in certain jurisdictions these costs are covered centrally.

Diagnostic laboratories should be mindful of the impact of laboratory investigations on public health and consider requests from the reference laboratories for specimen referral or enhanced information.

Identification of *Clostridium* species



* These give variable test results

The flowchart is for guidance only.

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An explanation of the reference assessment used is available in the [scientific information section on the UK SMI website](#).

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