Detection and enumeration of bacteria in swabs and other environmental samples

National Infection Service
Food Water and Environmental Microbiology
Standard Method
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Citation for this document:
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Introduction

Scope

This method is applicable to the procedures used for examination of environmental samples including swabs from carcasses in meat processing plants, swabs of food preparation surfaces and other environmental samples such as cloths collected from the food manufacturing environment and bottle rinses.

This support method must be used in conjunction with accredited methods for the detection of bacteria in foods and includes the use of three different types of swab.

Background

The significance of food contact surfaces in food safety is highlighted in the Food Safety Act 1990\(^1\) which states under section 29 (c) that authorised officers can “take a sample from any food source, or a sample of any contact material, which is found by him on or in any such premises”. This implies that environmental samples are to be considered as part of legislation. EU Legislation also states that “Sampling of the production and processing environment can be a useful tool to identify and prevent the presence of pathogenic micro-organisms in foodstuffs\(^2\), and cleanliness standards have also been laid down for meat premises in European Law\(^3\). A recommendation to undertake environmental monitoring of the food preparation environment during the investigation of poor microbiological results is also given in the HPA Guidelines for Assessing the Microbiological Safety of Ready to Eat Foods Placed on the Market\(^4\).

The environment can be contaminated with a variety of micro-organisms derived from various sources. Estimation of the overall numbers of bacteria (Aerobic Colony Count, \textit{E.coli} and Enterobacteriaceae) present can provide useful information when assessing general hygiene. Environmental monitoring can also be used as part of routine inspections of food premises or in an investigation of a suspected food poisoning outbreak where surfaces are thought to be likely vehicles of cross-contamination or if no food is available for examination.

Determination of the number of aerobic viable micro-organisms and the number of Enterobacteriaceae on a specified area of a surface can provide an indication of the cleanliness and allows monitoring of cleaning procedures over time.

Detection of \textit{Listeria} species in a food processing premises can be used as an indicator of poor cleaning. The presence of foodborne pathogens such as \textit{Listeria}...
monocytogenes, Salmonella species, Campylobacter species and STEC O157 in a ready to eat food preparation area or retail setting is of particular concern.

The main value of determining the colony counts on a surface is to assess the cleanliness of that surface. If a swab is taken to demonstrate that procedures used for cleaning are effective, swabbing should be performed after the surface has been cleaned. If it is suspected that cleaning is not being done or the methods used for cleaning are not effective then swabbing can be done as part of a routine hygiene inspection without prior cleaning. A high count can be an early warning of inadequate cleaning practices in food premises and may highlight the need for further investigation but careful consideration must be given if the surface is in use at the time of sampling. Cleaning of a surface prior to swabbing during outbreak investigations should be avoided to maximise the possibility of detecting a pathogen.

The method of choice for examination of surfaces is swabbing of a known area (10-100cm²) using a sterile swab that has been moistened in 10mL of neutralising diluent. This semi-quantitative approach enables enumeration of the micro-organisms per cm² and can facilitate interpretation of the results. When qualitative or detection tests only are to be performed the area swabbed need not be known but this may limit the laboratory’s ability to interpret the significance of the results obtained. It is assumed that the sample has been taken by a competent sampling officer who has been trained to carry out this procedure.

Microbiological examination of cleaning cloths can also be useful both in monitoring general hygiene and in the investigation of outbreaks, since these cloths are frequently used to wipe food preparation surfaces, and therefore have the potential to spread microbial contamination over the surfaces. Since many cloths are stored wet and may contain food debris, they have the potential to support the growth of high numbers of bacteria if not sanitised appropriately. Both enumeration of hygiene indicators and detection of pathogens can be performed on these samples by rinsing the entire cloth in diluent or a growth medium.

1. Principle

Three types of commercially available swabs are issued to customers by PHE laboratories. Two contain a measured volume of commercially prepared neutralising buffer or neutralising diluent while the third has Amies charcoal media. Neutralising buffer/diluent is used to neutralise the effect of disinfectants that have been used to clean surfaces, charcoal containing media has a similar effect. Surfaces are generally swabbed by the customer (with the exception of complex investigations where
laboratory staff may be asked to assist with sampling). The laboratory provides the
customer with instructions and offers training for use of the swabs but the
effectiveness of the swabbing procedure used is often outside the control of the
laboratory.

When enumeration tests are required a template enclosing a known surface area
(eg 10cm x 10cm) is used. Swabs in diluent are refrigerated at 2-8°C and are
submitted to the laboratory as soon as possible to ensure that they are examined on
the day of collection or at least within 24 hours of collection. The swabs are vortex
mixed or homogenised to aid release of organisms into the diluent. The swab diluent
and serial dilutions if necessary are subjected to the same methods as used for
examination of food homogenates.

The target count for enumeration tests is in the range of 0 to 300 colony forming units
(CFU) per plate. Samples likely to have counts that exceed the countable limit per
plate (method dependent) must be diluted.

Detection of pathogens from a known surface area can be performed by addition of
enrichment media to the swab sample. In outbreak situation a large random area can
also be sampled to maximise the likelihood of detection.

Enumeration and detection test can be performed on cloths and other environmental
samples such as bottle rinses by adding measured volumes of diluent or enrichment
media.

2. Definitions

For definitions of target organisms refer to the corresponding PHE standard methods
for foods for the bacteria of interest.

3. Safety considerations

3.1 General safety considerations

Normal laboratory precautions apply. All laboratory activities associated with this Standard Method must be risk assessed
locally to identify hazards. Appropriate controls must be in place to reduce the risk
to staff or other groups. Staff must be trained to perform the activities described and must be provided with any personal protective equipment (PPE) specified in this method. Review of this method must also include a review of local risk assessments to ensure that controls are appropriate and effective.

**Information Note:** Throughout this method hazards are identified using red text. Where a means of controlling a hazard has been identified this is shown in green text.

### 3.2 Specific safety considerations

Refer to individual food methods for specific safety considerations

### 3.3 Laboratory containment

Swabs, cloths or other environmental samples that are considered to be “high risk” for the isolation of STEC O157, must be handled in a CL3 Laboratory.

### 4. Equipment

- top pan balance capable of weighing to 0.1g
- stomacher
- vortex mixer
- stomacher bags (sterile)
- other equipment as detailed in PHE, MS, FW&E Standard Methods

### 5. Media, reagents and issue of sampling equipment

#### 5.1 Commercially prepared Dacron swabs with neutralising buffer

These are stored at ambient temperature and must be used before the displayed expiry date. Instructions for use and a template are issued to the customer (see Appendix 1).
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- 10mL neutralising buffer with dacron tipped swab (eg Technical Service Consultants Ltd, product code TS5-42)
- sterile templates (eg 10cm x 10cm plastic templates, Technical Service Consultants Ltd, product code TS15-T)

5.2 Commercially prepared sponge swabs with neutralising buffer

These are stored at ambient or under refrigeration (2 – 25°C) and must be used before the displayed expiry date. They should not be used if they appear to have dried out during storage (the chance of swabs drying out during their shelf-life may be reduced by storing between 2 and 8°C). Instructions for use with a template are issued to the customer (see Appendix 2).

- SpongeSicle™ swabs or sponge swabs containing 10mL neutralising buffer (eg 3M, product code SSL-10NB)
- sterile templates (eg 10cm x 10cm plastic templates, Technical Service Consultants Ltd, product code TS15-T)

5.3 Commercially prepared viscose tipped swabs with Amies charcoal medium

These swabs are used for the detection of a single pathogen only. They are stored at ambient or under refrigeration (2 – 25°C) and must be used before the displayed expiry date.

- amies charcoal swabs (eg Technical Service Consultants Ltd, product TS-5 18)

5.4 Sampling bags for collection of other environmental samples

Other environmental samples including cloths or utensils can be collected in sterile sampling bags. Bottles to be rinsed should be capped (if possible) and placed in a food grade bag. Food grade bags are not provided by the laboratory.

5.5 Media and reagents

Equivalent commercial dehydrated media may be used; follow the manufacturer’s instructions.

*Peptone Saline Diluent (PSD)*
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Peptone 1.0 g
Sodium chloride 8.5 g
Water 1L

pH 7.0 ± 0.2 at 25°C

Information note: This media is often referred to as Maximum Recovery Diluent (MRD) in the laboratory

Buffered peptone water (ISO formulation)

Enzymatic digest of casein 10.0 g
Sodium chloride 5.0 g
Disodium hydrogen phosphate dodecahydrate 9.0 g
or anhydrous disodium hydrogen phosphate 3.5 g
Potassium dihydrogen phosphate 1.5 g
Water 1 L

pH 7.0 ± 0.2 at 25°C

Refer to individual food test methods for culture media and reagents.

6. Sample processing

The diluent used will depend on the tests to be performed. If enumeration tests only are being performed on a swab Peptone Saline Diluent (PSD) or Buffered Peptone Water (BPW) can be used as the diluent. If the swab will be subjected to detection of Salmonella species, in addition to enumeration tests BPW can be used. Detection of pathogens other than Salmonella from swabs requires the addition of selective enrichment media. Due to this only one pathogen can be sought per swab. Cloths for detection of a single pathogen are prepared by adding 500mL of the appropriate enrichment medium. If a cloth or other environmental sample is being investigated for more than one pathogen by enrichment or if enumeration tests are being performed in addition to pathogen testing PSD should be added, the sample homogenised and the resulting liquid aliquoted to perform these tests. The resulting suspension can be mixed with the enrichment media for the pathogens of interest. If this is performed a comment must be added to the report to reflect that the entire sample was not tested (see section 9).

6.1 Sample preparation and dilutions
6.1.1 Preparation of dacron swabs

The sample comprises a swab in a tube of 10 mL neutralising buffer. For the purpose of this method this is considered to be a $10^0$ dilution or neat sample.

*Preparation for enumeration tests*
Transfer the neutralising buffer and swab tip to a sterile bag with wire closures. Perform a 1 in 10 dilution by adding 90mL of PSD or BPW. The sample should be homogenised for between 30 seconds and 3 minutes in a stomacher. The homogenisation time required will depend on the manufacturer instructions and the sample. Transfer 20mL to a universal container. This is equivalent to a $10^{-1}$ dilution and gives a lower limit of detection of 100 CFU per swab if 1mL is plated. Further dilutions can be performed if required.

*Preparation for detection tests*
Transfer the contents of the universal (including the swab tip) to a sterile bag with wire closures or sterile sample container. Add the appropriate enrichment medium for the test required to gravimetrically prepare a 1:10 dilution. Incubate and sub-culture as described in individual methods.

6.1.2 Preparation of sponge swabs

The sample comprises of a sponge moistened in 10mL of neutralising buffer inside a plastic bag with wire closures. For the purpose of this method this is considered to be a $10^0$ dilution or neat sample. The handle should have been aseptically removed at the time of use. If all, or a portion of the handle is still in place this should be aseptically removed by pushing it up through the bag and snapping it off. It must be noted on the report that the handle was not fully removed.

*Preparation for enumeration tests*
Add 90 mL of PSD or BPW to the bag containing the sponge swab to prepare a $10^{-1}$ dilution. Place the plastic bag with sponge swab inside a stomacher and homogenise for between 30 seconds and 3 minutes. The homogenisation time required will depend on the manufacturer instructions. Transfer 20mL of diluent to a universal. This gives rise to a lower limit of detection of 100 CFU per swab if 1mL is plated.

*Preparation for detection tests*
Add 90mL of the required enrichment medium to the swab bag and homogenise for between 30 seconds and 3 minutes in a stomacher. The homogenisation time required will depend on the manufacturer instructions. Incubate and sub-culture as described in individual methods.
6.1.3 Preparation of charcoal swabs

The sample comprises a swab in a tube of Amies Charcoal medium.

Preparation for detection tests
Transfer the swab tip only to a universal container approximately 20mL of the appropriate enrichment medium for the test required. Incubate and sub-culture as described in individual methods.

6.1.4 Preparation of cloths and other environmental samples

The cloth samples that will be submitted are likely to vary considerably in their size and degree of moisture content. It may also be the case that the cloth contains disinfectant and it is important that an attempt to dilute this disinfectant is made. It is therefore recommended that 500mL of diluent is added to the cloth regardless of size and moisture content. The diluent added will depend on the tests to be performed. It is recommended that if enumeration tests only are being performed PSD or BPW is used. If Salmonella and hygiene indicators are sought dilute with 500mL of BPW remove 20mL for enumeration tests and incubate the rest of the sample as described in method FNES16 (F13). Where enumeration is being performed in addition to the detection of pathogens PSD or BPW should be used and 25mL, aliquoted to 225mL of the appropriate enrichment media for each test of interest. If a single pathogen is being sought in a cloth consider adding 500mL of the appropriate enrichment media to the cloth and incubating the whole sample. The sample should be homogenised for between 30 seconds and 3 minutes in a stomacher. The homogenisation time required will depend on the manufacturer instructions and the sample. Where this is not possible the sample should be manipulated manually for at least 2 minutes. This resulting suspension is considered to be the neat or 10^0 dilution.

Other environmental samples including scrubbing brushes, nozzles and utensils are placed in a sampling bag. When enumeration tests are being performed 100mL of the appropriate diluent is added. These samples must be manipulated manually for at least 2 minutes, again the resulting suspension is considered to be the neat or 10^0 sample. When carrying out a pathogen detection test, sufficient enrichment media must be added to fully immerse the sample.

6.1.5 Preparation of bottle rinses

The bottle submitted for rinsing is removed from the sample bag aseptically. If the bottle is intended for single use, add 20mL of PSD. The internal surfaces of the bottle
are thoroughly rinsed by rotating the bottle gently 12 times in one direction. The bottle is allowed to stand for 15-30 minutes. Rinse the internal surfaces of the bottle again by rotating the bottle gently 12 times in one direction. Using a sterile pipette transfer the bottle rinse fluid to a sterile universal container. This is the neat or $10^0$ sample. When carrying out pathogen detection tests, add 100 mL of enrichment media to the bottle, rinse the bottle as described above and transfer the broth to a closure bag or sterile container.

6.2 Inoculation and incubation

6.2.1 Enumeration methods

For all samples, dilutions should be selected that will give colony counts within the appropriate range for the test being performed.

Proceed using PHE MS FW&E Standard methods for Foods.

Using a separate sterile pipette for each dilution, aseptically transfer 0.5mL or 1mL volumes (pour plates methods only) of the initial dilution ($10^{-1}$) of the sample to the appropriate medium. If necessary repeat the procedure with further decimal dilutions prepared in PSD.

6.2.2 Detection methods

For all samples gravimetrically prepare a $10^{-1}$ dilution in the appropriate enrichment medium. Enumeration tests can also be performed when samples have been diluted in BPW. If multiple pathogen tests are to be performed sample homogenates can be aliquoted to the appropriated enrichment media to achieve a $10^{-1}$ dilution.

Proceed using PHE MS FW&E Standard methods for Foods.

7. Quality control

Quality control of media and internal quality assurance checks should be performed according to in-house procedures. See PHE MS FW&E Standard methods for Foods for each method.
8. Calculations of results

Calculations occur automatically in the StarLims system as described in National Method FNES6 (Q12) Sample processing and result entry in Starlims. Calculations are performed as described below.

For enumeration tests counts should be calculated, where possible, using dilutions giving 15 to 150 or 300 colonies on the plate. The upper countable limit used is method dependent; please refer to individual food methods.

Calculate count per swab or cloth as follows:

\[
\text{Count} = \frac{C}{v (n_1 + 0.1n_2)d} \times n_3
\]

where:

- \(C\) is the sum of colonies on all plates counted;
- \(v\) is the volume applied to each plate;
- \(n_1\) is the number of plates counted at the first dilution;
- \(n_2\) is the number of plates counted at the second dilution;
- \(n_3\) is the original volume of neat suspension (i.e. 10 for swab, 500 or 100 for other samples);
- \(d\) is the dilution from which the first count was obtained (e.g. 10\(^{-2}\) is 0.01;

If the swab is from a measured area the count can be divided by the area swabbed in cm\(^2\).

*eg* Sponge swab, area swabbed 100cm\(^2\), tested for Enterobacteriaceae.

45 colonies confirmed at 10\(^{-2}\) dilution.

\[
\text{Count} = \frac{45}{1(1 + 0)0.01} \times 10
\]

=4.5x10\(^4\)

This count is the count per swab so to calculate the result per cm\(^2\) it is necessary to divide this by the area swabbed which in this case is 100.
Result per cm\(^2\) is 4.5x10\(^2\) CFU per cm\(^2\).

\textit{eg} Dacron swab, area swabbed 100 cm\(^2\) tested for \textit{E.coli}.

Example: 15 colonies on plate 1 and 18 colonies on plate 2 at 10\(^0\) dilution.

\begin{equation*}
\text{Count} = \frac{33}{0.5 (2 + 0)0.1} x 10
\end{equation*}

=3.3x10\(^2\)

This count is the count per swab so to calculate the result per cm\(^2\) it is necessary to divide this by the area swabbed which in this case is 100.

Result per cm\(^2\) is 33 CFU per cm\(^2\)

\section*{9. Reporting of results}

All results are reported using the StarLims system as described in method FNES17 (Q13) Technical Validation and release of result in StarLims\(^{10}\). The test report must specify the method used, all details necessary for complete identification of the sample and details of any incidents that may have influenced the result.

Take as the result the count per swab, per cloth, per item or per cm\(^2\) expressed as a whole number to two significant figures (if less than 100).

If count is 100 or more, report counts with one figure before and one figure after the decimal point in the form of:

\begin{equation*}
a x 10^b \text{ CFU per swab, cloth, item, or cm}^2
\end{equation*}

where a is never less than 1.0 or greater than 9.9 and b represents the appropriate power of ten. Round counts up if the last figure is 5 or more, and down if the last figure is 4 or less.

\textit{eg}: 1920 CFU per swab = 1.9 \times 10^3 \text{ CFU per swab}
235,000 CFU per swab = 2.4 \times 10^5 \text{ CFU per swab}

\subsection*{9.1 Enumeration}
9.1.1 Area swabbed not specified

Report the count as CFU per swab.

9.1.2 Area specified

Report as CFU per cm². Include a comment on the report (eg Results expressed per cm² are calculated using the area tested as stated by the customer).

9.1.3 Cloth or other environmental samples

Report the count as CFU per cloth or per sample.

9.1.4 Bottle rinses

Report the count as CFU per bottle.

9.2 Detection

If a single pathogen test has been carried out only ie no other tests performed on the sample

Report as:-

Not Detected per swab, per cloth or per item.

or

DETECTED per swab, per cloth or per item.

If more than one pathogen test has been carried out (cloths or other samples only) or enumeration tests have been performed in addition to Salmonella (swabs) report as in swab or in item with the comment “Entire sample not tested”
10. Reference facilities and referral of cultures

Any pathogen detected in a swab or in other environmental samples, particularly during the investigation of an outbreak must be referred to a reference laboratory for further typing. Please refer to advice provided in individual food methods.

11. Acknowledgements and contacts

This Standard Method has been developed, reviewed and revised by National Infection Service, Food, Water and Environmental Microbiology Methods Working Group.

The contributions of many individuals in Food, Water and Environmental laboratories, reference laboratories and specialist organisations who have provided information and comment during the development of this document are acknowledged.

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References


Appendix 1: Swabbing for pathogens and indicator organisms using dacron tipped swabs

This procedure can be used after cleaning to assess cleaning standards it can also be used if no cleaning protocol is in place. During an outbreak investigation swab surfaces prior to cleaning when possible.

**Consumables** (supplied by the laboratory on request)  
**Supplier/product code**

1 x sterile swab with 10ml Neutralising Buffer  
TSC Ltd, TS5-42  

1 x sterile 100cm² template  
TSC Ltd, TS 15-T

1. Put on a clean overcoat and wash hands thoroughly prior to sampling. Sterile gloves are not necessary unless the sample is likely to lead to legal proceedings.
2. Select the area to be investigated. If the surface is not flat and indicator organisms are being sought estimate the area to be swabbed.
3. If the area is flat remove a template from its bag by the handle and hold it between the thumb and index finger of one hand. Ensure that the template does not come into contact with any other object and place in the selected position.
4. Open the swab pack (do not remove the sterile swab from the pack at this stage).
5. Remove the lid from the tube containing neutralising buffer.
6. Remove the swab from the packaging and insert it into the neutralising buffer to moisten it. Squeeze excess liquid from the swab by pressing the cotton tip against the side of the tube.
7. Replace the lid on the neutralising buffer tube.
8. Using the moistened swab, swab the entire template area for 30 seconds as shown in Figure 1 using firm even pressure and rotation of the swab.
9. Remove the cap of the tube containing neutralising buffer and insert the swab. The top of the swab then becomes the lid of the tube. Push swab firmly in place until it is fully engaged—it should “click”.
10. The sample should be labelled appropriately and placed in a suitable cool box in an upright position and transported to the laboratory as soon as possible to ensure it is tested on the same day or at least within 24 hours of collection.

If indicator organisms are being sought the area swabbed in cm² or per 100cm² needs to be known to enable meaningful examination. If pathogens are being sought the area swabbed is less critical in the examination.

It is essential to clearly indicate which investigations are required.

**Figure 1**

A  

B  

C  

D

If indicator organisms are being sought the area swabbed in cm² or per 100cm² needs to be known to enable meaningful examination. If pathogens are being sought the area swabbed is less critical in the examination.

It is essential to clearly indicate which investigations are required.
Appendix 2: Sponge swabs
instructions for use

1. Check that the swab is within its shelf-life, the swab appears wet and that the bag is intact.

   Please note: before swabbing ensure that the sponge is moist but not over wet by squeezing excess diluents into the sample bag.

2. Tear open the bag along the perforation. Feed the Sponge handle through the opening, by pushing up from outside the bag. Grasp the handle above the thumbstop and remove the Sponge from the bag. To avoid contamination do not touch the space between the thumbstop and sponge, or the sponge itself.

3. Using both sides of the sponge swab the surface back and forth for a period of approximately 30 seconds. If a template is being used swab in all directions within the template area.

4. Return the sponge part of the Sponge to the sterile sample bag. Do not insert the handle past the thumbstop. Break off the handle by holding the sponge in place from outside the bag and bending the handle back and forth at the point where it joins onto the sponge (do not try to break at thumbstop). The sponge will drop to the bottom of the bag.

5. Discard the handle. Roll down the top edge of the bag and secure with wire tabs. Sample sponge is ready for laboratory examination and must be transported at 2-8°C.
Appendix 3: Detection and enumeration of bacteria in swabs and other environmental samples

**Enumeration methods**

For swabs test neat or prepare a 1:2 or 10⁻¹ dilution in PSD or BPW.
For cloths add 500mL of PSD

Mix by vortexing or in a stomacher for 30secs to 3mins

Prepare further dilutions in PSD if required

Inoculate appropriate medium for organisms of interest and proceed as for PHE Standard method for food

Count and confirm colonies as for the PHE Standard method for food

Calculate Count per swab, per cm², per cloth or per item

**Detection methods**

For swabs prepare an approximate 10⁻¹ dilution of the sample in the required enrichment medium.
For cloths add 500mL of PSD, homogenise and aliquot to the required enrichment media

Mix by vortexing or in a stomacher for 30secs to 3mins

Inoculate appropriate medium for organisms of interest and proceed as for PHE Standard method for food

Report as pathogen DETECTED or Not Detected

If required refer isolate to the appropriate reference laboratory