**Microbiology**

**Service User Manual**

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# introduction and Scope

The purpose of this document is to provide users with clear information on the Microbiology service provided by East Kent Hospital University Foundation Trust (EKHUFT). It includes information regarding the pre-examination activities of specimen collection and handling prior to analysis in the laboratory, to ensure that results are accurate and clinically useful.

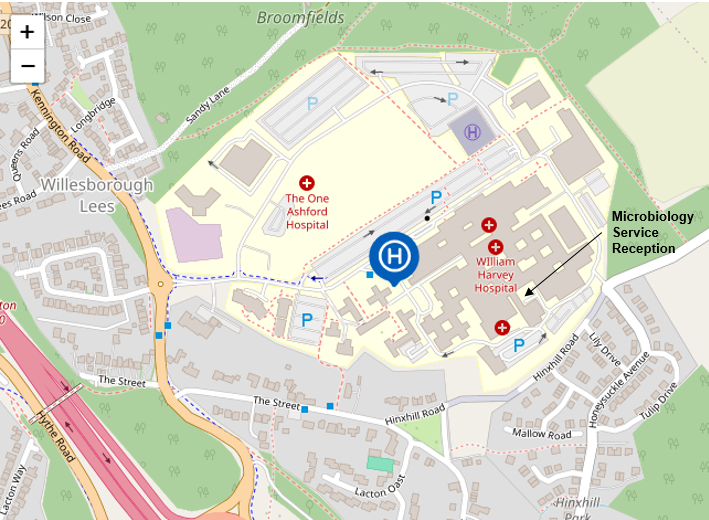
Ownership and accountability of the user manual lies with the Microbiology Service Head of Service.

The scope of this user guide includes routine, urgent samples, sample containers, transportation and turnaround times from specimen receipt to results becoming available. The intended audience for this user manual is healthcare professionals; however, instructions for self-taken specimen collection have been included for distribution to patients.

# general information on the use of the microbiology laboratory

The Microbiology Service is based at the William Harvey Hospital (WHH) in Ashford and provides diagnostic Microbiology services (Bacteriology, Parasitology and specialist Virology/Mycology investigations) and Infection Control services to EKHUFT and GP/Community services. The Microbiology Service also hold Hepatitis B (500iu and 200iu), and Rabies (2.5iu/ml) immunoglobulin on behalf of UK Health Security Agency (UKHSA).

## Locating and contacting the Microbiology Laboratory



Microbiology Service

The William Harvey Hospital

Kennington Road

Willesborough

Ashford

Kent TN24 0LZ

Tel: 01233 616760 (Mon – Sat for routine enquiries)

Sunday – urgent enquiries only, via switchboard (01233 633331)

## Laboratory opening hours

* Weekdays: 08:30 to 19:00
* Weekends and bank holidays: 08:30 to 17:00

## Medical advice

Consultation about the investigation and management of infection is welcomed. Early liaison over infection control matters, especially outbreaks is encouraged. Consultant Microbiologists are available to advise on examinations, the use of services (including the required type of sample), clinical indications and limitations of the examination procedure and the frequency of examination requests and interpretation of results (ISO 15189:2022 5.3.3a).

The laboratory staff includes five Consultant Microbiologists who provide a service across the three main hospital sites at Ashford, Canterbury and Margate.

The laboratory maintains close links with the University of Kent.

Medical advice can be obtained by telephoning the department (see section 2.1) or via the WHH switchboard out of hours: **ask for the On-Call Medical Microbiologist.**

## Senior members of the Microbiology Service

* Consultant Microbiologist / Virologist, Head of Service - Microbiology:

Dr S Moses MBBS, MRCP, FRCPath (Virology), CCT in Microbiology & Virology

* Consultant Microbiologist / Virologist and Antenatal & Newborn Screening Clinical Lead:

Dr M Strutt MBBS, MSc, FRCPath

* Consultant Microbiologist & Antimicrobial Stewardship Lead:

Dr S Glass MEng MBBch MRCP FRCPath

* Consultant Microbiologist:

Professor Fritz Muehlschlegel MD, FRCPath, FRSB

* Consultant Medical Microbiologist:

(Vacant)

* Specialist Doctor in Microbiology:

(Vacant)

* Consultant Clinical Scientist in Virology & Infection:

Dr E Meader PhD, FRCPath

* Head Biomedical Scientist, Service Lead:

Mrs R Arkley CSci, FIBMS

* Chief Biomedical Scientist (Bacteriology):

Mr Michael Dawson MSc, FIBMS

* Chief Biomedical Scientist (Virology) and Antenatal & Newborn Screening Laboratory Lead:

Mrs Claire Warren PGDip, MIBMS

* Chief Biomedical Scientist, Containment Level 3 Manager & Pathology H&S Lead:

Miss Samantha Sheppard MSc, FIBMS, MISTR

* Quality Lead:

Mrs Sarah Hogben MSc, BSc

# Comments, compliments & complaints

The Microbiology Service Laboratory is committed to offering high quality specialist microbiology services that meet and respond to the needs of all service users.

If something has gone wrong or you are not happy with any aspect of our services then please do let us know or alternatively if there is something we have done well we would be grateful for your feedback. There are two main ways that you can make a compliment, complaint or raise a concern:

1. Contact the Laboratory directly:

Contact the laboratory directly either by telephone, email or in writing as below.

Telephone the Laboratory - 01233 616760

Ask to speak to the Head Biomedical Scientist or the Chief Biomedical Scientist.

E-mail the Head Biomedical Scientist or Chief Biomedical Scientist on:

Head Biomedical Scientist: [rachael.arkley@nhs.net](mailto:rachael.arkley@nhs.net)

Bacteriology Chief Biomedical Scientist: [michaeldawson1@nhs.net](mailto:michaeldawson1@nhs.net)

Virology Chief Biomedical Scientist: [claire.warren@nhs.net](mailto:claire.warren@nhs.net)

Microbiology Chief Biomedical Scientist: [samantha.sheppard2@nhs.net](mailto:samantha.sheppard2@nhs.net)

Write to the Laboratory at:

Head Biomedical Scientist

Microbiology Service Laboratory

William Harvey Hospital

Kennington Road

Ashford

Kent

TN24 0LZ

Direct contact with the Laboratory is often the best way to make a complaint as it means that we can quickly understand the problem and take immediate action to investigate and resolve the situation.

1. Contact the Patient Advice and Liaison Service (PALS)

If you prefer you can also make a complaint via the Patient Advice and Liaison Service (PALS) on the contact details below:

Telephone: 01227 783145, 9am to 4pm Monday to Friday

E-mail:   [ekh-tr.PALS@nhs.net](mailto:ekh-tr.PALS@nhs.net)

Write: Patient Advice and Liaison Service (PALS), First Floor, Trust Offices, Kent and Canterbury Hospital, Ethelbert Road, Canterbury, Kent, CT1 3NG.

Please see the following link for the complaints process:



[Patient Advice and Liaison Service (PALS): Do you have a worry or concern? We are here to help. (Easy Read and Text only))](https://leaflets.ekhuft.nhs.uk/patient-advice-and-liaison-service-pals-do-you-have-a-worry-or-concern-we-are-here-to-help-easy-read-and-text-only/)

The Laboratory follows the Pathology Policy for the Management of Comments and Complaints Raised by Patients, Relatives or Users of the Service in line with the Trust Policy and national guidance. In all cases our aim is to ensure that complaints and concerns are resolved quickly and thoroughly with appropriate investigation and resolution.

# Quality assurance

The laboratory is accredited by the United Kingdom Accreditation Service (UKAS) to ISO 15189. Details are available on the UKAS website [www.ukas.com](https://www.ukas.com/find-an-organisation/), (type 9399 in the search function).

Where tests are not included on the laboratory’s scope of UKAS accreditation the following caveat is added to the report:

East Kent Microbiology is a UKAS accredited laboratory, this test however does not appear on our accredited scope of practice.

External quality assessment (EQA) schemes assess the Microbiology Service’s performance against other laboratories. The Microbiology Service participates in the UK National External Quality Assessment Scheme (UKNEQAS) for many of the microbiological investigations performed by the Microbiology Service where a scheme is provided.

**UKNEQAS schemes the Microbiology Service currently participate in:**

Anti-HBs detection Antifungal susceptibility Immunity screen

Antimicrobial susceptibility Carbapenemase-Producing organisms

*Clostridioides difficile* CMV DNA quantification

Cryptococcal antigen detection

Diagnostic serology hepatitis EBV DNA quantification

Faecal parasites (molecular) Fungal Biomarkers (Galactomannan)

General bacteriology Genital pathogens

Hepatitis B serology Hepatitis C serology

Hepatitis E detection HIV serology

Immunity screen Interferon Gamma Release Assay

Lyme serology (CSCQ) Measles & Mumps IgG serology

Molecular detection of SARS-CoV-2 Molecular detection of viruses in CSF

MRSA screening Mycology

Parvovirus B19 and Rubella serology Rubella IgG serology

Syphilis serology Toxoplasma serology

Urinary antigens Viral gastroenteritis (lateral flow – BCP)

**QCMD schemes the Microbiology Service participate in:**

Bacterial Gastroenteritis BK Virus DNA

Central Nervous System I (Viral Meningitis & Encephalitis) Hepatitis B Virus DNA

Hepatitis C Virus RNA Human Immunodeficiency Virus RNA MALDI-TOF Mycoplasma genitalium

Pneumocystis jirovecii pneumonia (PCP) DNA Trichomonas vaginalis

Viral Gastroenteritis Joint Infection EQA Pilot Study

**Labquality schemes the Microbiology Service participates in:**

Blood culture (incl. sepsis multiplex methods)

Cytomegalovirus antibodies

Helicobacter pylori, antigen detection in faeces

Hepatitis A antibodies

Human T-cell lymphotropic virus antibodies

Meningitis encephalitis multiplex nucleic acid detection

Mycoplasma pneumoniae, antibodies

Mycobacterium tuberculosis, drug resistance, nucleic acid detection

Neisseria gonorrhoeae (GC), culture & susceptibility testing

Respiratory infections multiplex nucleic acid detection

Sexually transmitted diseases multiplex nucleic acid detection

Surveillance for multidrug resistant bacteria, gram neg. rods

Urinary bacterial screening with automated analysers

Urine culture (quantitative) screening, ID & susceptibility

**Instand e.V. schemes the Microbiology Service participates in:**

Mycoserology 02 – Cryptococcus neoformans/ -gattii antigen

Virus Immunology – Hepatitis D Virus (Ab)

Internal quality assurance (1% of our work is re-submitted to determine reproducibility and repeatability), internal quality controls and a comprehensive audit schedule are used to monitor the quality of results.

# data Confidentiality and patient consent

The Microbiology Service complies with the requirements of the Data Protection Act, the Caldicott Principles on safeguarding patient confidentiality and information, and the Royal College of Pathologists’ guidance. All patient identifiable information is regarded as confidential and is only released for official purposes i.e. to professionals with responsibility for patient care or public health. All confidential data is stored securely and only held as long as necessary for operational purposes.

Consent, for the purposes of confidentiality, means that the service user understands and does not object to:

• the information being disclosed or shared;

• the reason for the disclosure;

• the people or organisations the information will be shared with; and

• how the information will be used.

For consent to be valid, it must be voluntary and informed, and the person giving consent must have the capacity to make the decision.

When a patient presents to a GP surgery, theatre or clinic (e.g. phlebotomy, outpatient, antenatal, etc.) and participates in a sample collecting procedure, it is assumed that the patient has already given consent upon receipt of the request in the microbiology laboratory. Refer to section 10.2 Request Form Labelling for additional information on patients’ right to decline certain tests on antenatal specimens.

# Routine samples

The laboratory is fully staffed during its routine opening hours (see section 2.2). Outside of these routine working hours, only essential work can be undertaken.

The laboratory aims to attend to all microbiological samples for routine culture on the day of receipt and results will be reported according to published turnaround times in section 26.

Some serology investigations are batched but may be processed urgently by arrangement with the laboratory.

# urgent samples

The laboratory should be contacted by telephone when an urgent sample is being sent.

Samples **MUST** be packaged in BS regulation transport containers provided at **K&CH Main Reception and** **QEQMH Emergency Department (ED)**. See section 11.

Please ensure that samples are transferred to the Microbiology service at WHH as soon after collection as possible. Outside of EKHUFT the requestor **MUST** arrange a taxi service to transport the samples to Microbiology WHH.

Within EKHUFT sites the requestor **MUST** arrange transportation to the on-site Pathology department. The Pathology department at the Kent and Canterbury (K&CH) and Queen Elizabeth The Queen Mother (QEQMH) hospitals will arrange transport to the Microbiology Service at WHH.

During normal working hours, please telephone the laboratory on the external number (section 2) or using the internal number 723 8370. Please mark the request as **‘Urgent’.**

Outside of normal working hours please telephone the on-call Biomedical Scientist via switchboard. Please mark the request as **‘Urgent’.**

# Specimen rejection

Specimens may be rejected by the laboratory if:

1. There is insufficient volume of sample.
2. The sample is received in an inappropriate container e.g. jam jar
3. The request is inappropriate for the specimen type.
4. The details on the sample do not match the request.
5. Unlabelled specimens (a minimum of three points of patient identification (full name, full date of birth and identification number) is required).
6. Requests containing insufficient information.
7. Leaking specimen containers
8. The sample is too old to process and give reliable results
9. The sample has been stored in an inappropriate environment i.e. excessive temperature.

However if blood cultures, tissues or joint, pleural, ascitic, CSF fluids are received as described above the ward would be contacted prior to the decision to discard the specimen.

# ORDERING CONSUMABLES (E.G. SPECIMEN CONTAINERS)

Please note that supplies of Blood Culture kits, Pernasal swabs, TB urine containers, and kits for flu surveys must be requested from the Microbiology Service – please see section 2.1 for contact details.

To order all other microbiology consumables for your department/surgery please complete the supplies form (please see embedded supplies form Appendix 16) and send the request to the Supplies and Procurement Team: [ekh-tr.PathologyOrder@nhs.net](mailto:ekh-tr.PathologyOrder@nhs.net) 01233 633331 Ext. 723-8403

# labelling specimens and request forms

Please refer to EKHUFT Policy Centre for Pathology Sample and Request Form Acceptance Policy.

**Details in bold are ESSENTIAL, these may be hand written or labels can be used.**

## 10.1 Specimen labelling

**Labelling must be on the body of the sample pot to be accepted, any labelling on the lid may be discounted. All samples MUST be labelled in the presence of the patient.**

* **Last name/Surname AND First Name** (or coded identifier as in GU patients);
* **Date of Birth and / or NHS/Hospital number**
* **Specimen type(s) and anatomical site(s) (essential for Microbiology).**

**Samples labelled with initials rather than full name will not be accepted unless BOTH DOB and NHS/Hospital number are present.**

**Any sample that has been relabelled (crossing out visible on the sample tube) will not be accepted**

For Microbiology **the site and specimen type is ESSENTIAL** on the samples.

If using more than one sample pot, each pot must be individually numbered and labelled as above.

## Request Form Labelling

**Details in bold are ESSENTIAL**

* **Last name/Surname (or coded identifier as in GU patients);**
* **First name;**
* **Date of Birth;**
* **NHS/Hospital number**
* **Date of sample;**
* **Tests required;**
* **Address/location for report;**
* **Anatomical site/type**
* **Clinical details (refer to section 15.1 Specimen requisition) e.g. relevant symptoms and duration, history of drug administration such as antimicrobial, pre- / immunosuppression therapy, history of alcohol / substance abuse, history of travel and vaccinations, underlying medical conditions. Failure to do so may affect clinical interpretation of results.**
* Consultant/GP name;
* Time and Date of Sample (essential for test specific samples)
* Contact/bleep number of requestor;
* Urgency of request.
* Initials of person who took sample (blood tubes)
* Source/location of patient
* **NB: all antenatal specimens must have patient consent indicated on the request form for full antenatal screen and can decline to have their specimen tested for Hepatitis B, Syphilis and/or HIV –** for all other specimens it is assumed that patient consent has been given to the clinician following an explanation about the tests proposed. This must include consent to disclose clinical information and patient history to relevant healthcare professionals where referral is required.

All samples MUST have a minimum of three points of patient identification, (full name, full date of birth and identification number)**.**

**NB: PLEASE ENSURE THAT THE SPECIMEN IS PLACED IN THE LARGE PLASTIC POCKET OF THE SAMPLE REQUEST FORM.**

# collection and Transport of specimens to the laboratory from GP surgeries / private hospitals / non-hospital based clinics

Please see appendices for instructions on specimen collection.

As soon as the specimen has been collected, the container (labelled correctly) must be placed in the integral transparent plastic transport bag immediately after taking the sample.

The transport bag must be sealed by means of an integral sealing strip as indicated on the request form. Bags must not be sealed with pins, staples or metal clips etc.



Samples are collected regularly from the GP surgeries / private hospitals / non-hospital based clinics by the EKHUFT transport vehicles. The specimens must be placed in robust, secure and safe Category B transport boxes designed specifically for transporting biological specimens.



# transporting specimens from ekhuft wards and hospital based clinics

Please see appendices for instructions on specimen collection.

**N.B**. To ensure samples are transported promptly and arrive at WHH microbiology the same day as collection, please note the following cut off times for sample arrival at the relevant pick-up point are as follows:

K&CH Pathology Reception 15:30

QEQMH Pathology Reception 14:00

BHD Phlebotomy 15:30

RVHF Phlebotomy 14:30

## Transport of high risk sample e.g Viral Haemorrhagic Fever (VHF)

Please refer to Viral Haemorrhagic Fever (VHF) – Ebola/Lassa/Marburg/CCHF

Operating Standard available from EKHUFT Policy Centre.

All specimens taken from suspected VHF infection patients must be transported to the laboratory in a sealed specimen bag placed in a closed appropriate transport box. Specimens MUST NOT be carried by hand to the onsite Pathology department or be put through the pneumatic tube system.

The clinician taking the specimens **MUST** inform the onsite Pathology department and Microbiology at the William Harvey Hospital. The clinician must also ensure the samples are transported to the onsite Pathology department **immediately** after the sample has been taken.

Items required for sending specimens to onsite Pathology department:



## Transport of all other specimens

The labelled sample container of all specimens must be placed in the integral transparent plastic transport bag immediately after taking the sample.

The transport bag must be sealed by means of an integral sealing strip as indicated on the request form. Bags must not be sealed with pins, staples or metal clips etc.



Blood cultures must be sent immediately to the Pathology department using the pneumatic air specimen transport system (POD).

All other specimens with the exception of:

CSF samples

Urine samples greater than 50ml

Faeces samples

may be transported using the POD or alternatively collected by Porters, who must place the packed specimen inside a red biological specimen transport box to transfer the samples to the Pathology department:



Pathology staff must transfer the specimens into the blue transport bags prior to placing in the appropriate area for collection by the EKHUFT transport service that transfers specimens to the Microbiology department at WHH.



# requesting additional investigations

Requesting additional tests on microbiology samples cannot usually be performed unless they are serum samples. The following serum samples are retained for a minimum of 2 years and appropriate additional investigations may be requested for these serum samples at any time during this period providing there is sufficient volume of sample:

1. Renal transplant screens
2. Antenatal screens
3. Needlestick
4. Any sample where the submitting clinician has specified the serum to be saved.

Please note: it is often not possible for additional investigations to be added to serum samples which have been sent for blood sciences investigations. If serology is required, an independent sample must be sent.

# ON-Call investigations

There is an ON-CALL service for processing samples and providing Medical advice for William Harvey Hospital (WHH), Kent & Canterbury Hospital (K&C) and Queen Elizabeth the Queen Mother Hospital (QEQM). Samples from K&C and QEQM are transported to Microbiology at WHH for processing (see section 14.1).

The On-Call Biomedical Scientist (BMS) and Medical Microbiologists can be contacted via the EKHUFT switchboard.

Please **DO NOT** contact the On-Call BMS when blood cultures are performed as there are procedures in place for hospital porters to take them directly to the Pathology reception area.

Blood cultures are continuously monitored by an automated detection system 24hrs a day. All positive results are telephoned to the appropriate medical staff. **Please DO NOT call the laboratory to check whether a blood culture is positive.**

Please complete request form (electronic or written) clearly giving the **Ward, Requesting Doctor and the bleep/contact number.** Staff on other sites may not be conversant with abbreviations used locally.

## Sending Urgent Samples to Microbiology Out of Hours

These procedures are for **urgent** samples only and may include:

1. CSF samples
2. Vitreous/aqueous aspirates
3. Joint fluids (septic arthritis)
4. Corneal scrapes

* Other samples at the discretion of the requesting Consultant Please provide full clinical details with all requests. This is vital for the safety of laboratory staff, ensures the right tests are performed and enables clinical interpretive advice to be provided.
* Please state if the patient is immunocompromised.

**Arrangements for Monday to Friday 17:00 to Midnight and 08:30 to Midnight at weekends or public holidays**

1. Telephone the on-call Microbiology Biomedical Scientist (BMS) **via switchboard** once the sample has been collected.
2. Transport the sample to Pathology reception (on the local site) **immediately.** This can be by porter or clinical staff member.
3. The microbiology laboratory is located at William Harvey Hospital (WHH). Any sample arriving in pathology on an alternative site will be transported to WHH.
4. Samples will be tested upon arrival at the microbiology laboratory, unless informed otherwise by the on-call BMS. Results will usually be available with 2 hours.
5. **Results will be available on Sunrise electronic patient records (preliminary results) or DART (authorised results)** and results will only be telephoned if clinically significant and could influence the management of the patient.

**Arrangements after midnight**

1. Transport the sample to Pathology reception (on the local site) **immediately.** This can be by porter or clinical staff member.
2. The microbiology laboratory is located at William Harvey Hospital. Any sample arriving in pathology on an alternative site will be transported to WHH.
3. Samples will be tested upon opening of the microbiology laboratory at 08.30, and results will be usually available with 2 hours.
4. **Results will be available on Sunrise electronic patient records (preliminary results) or DART (authorised results).** Results will only be telephoned if clinically significant and could influence the management of the patient.

## Microbiology On-Call Procedures – For Pathology Staff

These are the procedures for **URGENT** Microbiology samples that require testing between Monday to Friday 17:00 – midnight and public holidays.

1. The microbiology on-call sample will be taken to the pathology reception by the requesting doctor, nurse or porter.
2. The on-call Microbiology Biomedical Scientist will contact the Blood Science Continuous Processing Pattern (CPP) staff member to ALERT them that the sample is coming down to reception also **providing the name of the patient and sample type.**
3. K&CH contact number – Biochemistry Ext **722-3174** alternative **722-5065** or via switchboard **bleep 7022**
4. QEQMH contact number – Biochemistry Ext **725-4428** or **bleep** **6131.** Or before 1700 call Specimen Reception onExt **725-4250**
5. The Microbiology Biomedical Scientist will either order a taxi or contact SERV depending on the time of day.
6. The CPP blood science member of staff will locate the sample in pathology reception and then package sample up into the designated transport box (see below).



1. **Between 06:00 and 19:00 hours** the Microbiology Biomedical Scientist will contact switchboard to arrange a taxi for URGENT samples to be **collected from pathology reception** for delivery to pathology reception at WHH. (NEW arrangement:-taxis previously collected from switchboard or A&E)
2. **AFTER 19:00 hours and before 06:00 hours** the Microbiology Biomedical Scientist will contact SERV **(01227 200 608)** for URGENT samples to be **collected from pathology reception** for delivery to pathology reception at WHH.
3. CPP pathology staff to contact the on-call Microbiology Biomedical Scientist **(via EKHUFT switchboard)** to ALERT them that the **TAXI** or **SERV** driver has collected sample from pathology reception.

## Contacting the Consultant Microbiologist

The Consultant Microbiologist on-call duties are detailed as follows:

17:00- 09:00 Monday to Friday

24 hours on Saturday, Sunday and Bank Holidays.

### Before 21:00

Between the hours of 17:00 – 21:00 Monday to Friday and 09:00 – 21:00 on Saturday, Sunday and Bank Holidays, the on-call Consultant Microbiologist will accept calls from junior doctors and other staff seeking advice.

### Between 21:00 and 09:00

Between the hours of 21:00 – 09:00 from Monday to Sunday including Bank Holidays, please see the following criteria for communicating with the on-call Consultant Microbiologist:

1. During the above hours there is an advice and guidance service only available for all issues relating to Microbiology.
2. The Consultant Microbiologist on-call will only accept calls from on-call Consultants, registrars and staff grade doctors.
3. The Manager on-call will be able to speak to the Consultant Microbiologist if necessary for advice and guidance.
4. Switchboard are instructed to check the designation of the member of staff wishing to be connected to the Consultant Microbiologist’s mobile or home telephone number.
5. Switchboard will only connect Consultants, Registrars and staff grade doctors and/or on-call Managers during the hours specified above.

# Submitting samples to the laboratory

## Specimen requisition

Electronic requisition (DART for GPs and Sunrise for hospital users) should be used whenever possible to allow access to patient results in a timely manner.

Where DART is used for specimen request please either finalise the request by selecting ‘collection at surgery’ or ‘collect at home’. When ‘collect at home is selected, please print the form to give to the patient and ask the patient to record the date and time of collection on the form and return the specimen and form to the surgery. The surgery should then enter the date and time of collection onto DART and finalise the request by selecting ‘collection at surgery’. This will generate an adhesive barcoded label which must be adhered to the specimen container. Please do not send non-barcoded samples with the DART form to the laboratory.

Where electronic request cannot be made, handwritten forms should be completed ensuring they are legible and signed by the person requesting the investigation.

Adequate clinical details **must** be provided, including the following information when relevant:

1. Patient demographics, including NHS number
2. Reason for request
3. Date of onset of illness
4. Date and time the specimen was collected
5. System(s) involved
6. Site of infection (especially for wounds)
7. Date of surgery if applicable
8. Previous and planned antimicrobial therapy
9. History of travel and vaccinations (details of countries visited are essential)
10. If possible, a diagnosis or provisional diagnosis should be given. Any defects in host defences e.g. immunosuppression should be mentioned.
11. History of alcohol / substance abuse
12. Underlying medical conditions

Also refer to section 10 Labelling Specimens and Request Forms

## Preparation to collecting specimens

**NB:** Hands must be decontaminated with alcohol hand rub and the appropriate protective equipment (i.e. disposable plastic apron and gloves) worn prior to the SPECIMEN being collected.

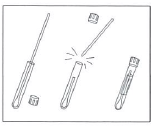
See also EKHUFT Infection Control Policy Document, which provides additional guidance/information on the collection of specimens.

All samples must be clearly labelled and dated (see section 10).

Specimens must be collected in the appropriate screw capped CE leak proof container. The containers (see sections 17 and 18) must be shatter proof, not overfilled, the lid secured tightly, and the specimen placed in a specimen bag.

The expiry date of the sample container must be checked before using for collecting the samples – please discard any sample containers that are past their expiry date.

Swabs must be placed fully into the specimen container and then the swab handle gently pulled / pushed against the sample container opening until the swab breaks at the swab breakpoint (this ensures that the swab is the correct height to fit in the sample container) – **DO NOT** force the swab into the specimen container as this causes the container to leak.



All specimens must be singularly – DO NOT place multiple samples in one bag.

For the safety of the laboratory staff, badly leaking samples will be discarded. It is assumed by the laboratory staff that a patient being investigated for HIV infection has been counselled by the doctor whose name appears on the request.

# optimising the validity of results

There are many factors that may affect the results obtained from laboratory investigations. Good laboratory practice and the laboratory quality management system minimises those factors that could occur within the laboratory. However the following factors that may contribute to erroneous laboratory results are out of the laboratory’s control:

| Factors that may affect results: | Reducing the risk: |
| --- | --- |
| Delays in transporting samples to the Microbiology Service @ WHH. Samples should be submitted for investigation as soon as possible following collection to prevent deterioration of cells and changes in the relative quantities of micro-organisms present.  The date of sample collections must be included with all requests. | * Ensure samples are transferred to pathology as soon as possible after collection to prevent deterioration of cells and changes in the relative quantities of micro-organisms. * Notify microbiology of any urgent samples prior to sending them. * Use the POD system to transport blood cultures to Pathology |
| Insufficient volume of sample sent to the laboratory | * Fill blood culture bottles appropriately – see Appendix 9. * Where possible adhere to fill lines on sample containers (please do not overfill) |
| Contaminated samples | * Adhere to sampling guidelines in EKHUFT Infection Control Policy Document and associated Operating Standards available on EKHUFT Policy Centre. * Maintain aseptic sampling technique * Use the appropriate screw capped CE leak proof container for collecting samples - see section 17 and 18. * Ensure sample containers are correctly sealed to prevent contamination * All sample transport containers must be securely closed to prevent sample leakage or contamination en route to the laboratory. |
| Inappropriate transport medium and/or sample container | * Use an appropriate plain screw capped CE leak proof specimen container to collect samples – please see section 17 and 18. |
| Inappropriate sample request | * Ensure that the information on the request is correct, include the patient’s clinical history – please see section 10. |
| Inappropriate storage conditions. Exposure to significantly raised or reduced room temperatures may affect the results obtained during laboratory investigations. | * If the appropriate sample transport media and containers are used, please refer to section 19 for further details. |
| Sample quality. The result of the laboratory investigation is dependent on the quality of the sample submitted to the laboratory. | * Ensure there is sufficient material or cells in the sample or on swabs. * Ensure the sample type is appropriate i.e. MSU or EMU * Ensure the sample site is clean * Maintain good aseptic techniques during sample collection. |
| Presence of inhibitory substance in the samples. The presence of antimicrobials or other inhibitory substances in the sample may affect the results of laboratory investigations and the subsequent recovery of micro-organisms. | * Any antimicrobial therapy the patient is on should be recorded in the clinical details section of the request. |

Recovery of micro-organisms or detection of cells, antigen or antibody is dependent upon the amount of micro-organisms, antigen or antibody present in the sample. Low levels in the sample may not be sufficient to detect by the microscopy, culture or assays employed by the laboratory.

**Please note that a single negative result does not exclude infection or immunity status and multiple repeat investigations may be required to obtain a definitive diagnosis.**

## Specimen Transportation and Sample Integrity

| **Sample Type** | **Container** | **Time collection – processing limit** | **Temperature Limits** | **Stabilising** |
| --- | --- | --- | --- | --- |
| **Blood culture** | Biomerieux BactAlert bottles | 4 hours | Room temp NOT incubated | Bottles inspected on arrival. Loaded out of hours. |
| **CSF** | Universal | 2 hour (SMI) | Room temp until microscopy and culture | On-call service. Out of hours transport. |
| **Parasites** | Faecal container | Specimens should be transported and processed as soon as possible. | Refrigeration at 2-8 °C | Parasep kit used Monday-Thursday. Faeces Friday-Monday prepared in Parasep on Monday. |
| **Faeces** | Faecal container | Enteric PCR – 48 hours | Refrigeration at 2-8 °C | Once in SPS + heat treated – can be stored between 2-8°C for up to 48 hours or at -20°C for 1 week. |
|  |  | Helico - 48 hours | Refrigeration at 2-8 °C | Helicobacter – stored at -20°C or below. |
| **Fluid** | Universal | Specimens should be transported and processed as soon as possible.  If acute infection is suspected and the result may affect medical management, receive and process the sample within 4 hours. The result for microscopy should be made available within 2hr of the Gram stain (SMI B26). | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature (SMI B26). | If result affects medical management it is dealt with using the out of hours procedure. |
| **Tissue** | Universal | Specimens should be transported and processed as soon as possible.  The volume of the specimen influences the transport time that is acceptable. Larger pieces of tissue maintain the viability of anaerobes for longer (SMI B17). | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature (SMI B17). | If result affects medical management it is dealt with using the out of hours procedure. |
| **Pus** | Universal | Specimens should be transported and processed as soon as possible.  The volume of specimen influences the transport time that is acceptable. Large volumes of purulent material maintain the viability of anaerobes for longer.  The recovery of anaerobes in particular is compromised if the transport time is delayed. | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature. | If result affects medical management it is dealt with using the out of hours procedure. |
| **Urine (for culture)** | Boric container | 3 days (SMI) | Room temperature | The use of a boric acid preservative increases the maximum permissible time for transport to the laboratory to up to 96hr.  Boric acid preservative at a concentration of 1–2% holds the bacterial population steady for 48–96 hours, and other cellular components remain intact. |
| **Wound / respiratory swab** | Medical wire | Specimens should be transported and processed as soon as possible (SMI). | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature. | Use of M40-A compliant Transwab® a self-contained gel transport swab for aerobes and anaerobes, for reliable collection and transport of microbiological specimens.  Amies medium with inorganic buffer ensures maintenance of microorganisms without overgrowth. The open weave rayon bud is non-toxic and allows best specimen uptake and release, particularly for microorganisms. |
| **Genital swab (for culture)** | Medical wire | Specimens should be transported and processed as soon as possible (SMI). | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature. | Transwab® self-contained gel transport swab for aerobes and anaerobes, for reliable collection and transport of microbiological specimens.  M40-A Compliant Amies medium with inorganic buffer ensures maintenance of microorganisms without overgrowth. The open weave rayon bud is non-toxic and allows best specimen uptake and release, particularly for microorganisms. |
| **MRSA screen liquid swab**  **VRE / CRO / ESBL G40-A transwab** | Medical wire | Specimens should be transported and processed as soon as possible (SMI). | If processing is delayed, refrigeration at 2-8°C is preferable to storage at ambient temperature. | Liquid culture swabs from MWE called ∑-Transwab®  M40-A Compliant**,** specimen is collected using ∑-Swab®, an open celled, polyurethane foam-tipped swab which allows complete flow through of reagents and micro-organisms. The specimen is then placed into the tube of liquid medium for transportation to the laboratory.  The micro-organisms in the specimen are dispersed through the medium, producing a uniform suspension ready for use, either conventionally or on an automatic specimen processing platform, or direct with many rapid molecular tests currently available. |
| **Mycology skin / hair / nail** | Envelope | 1 week | Storage in dry conditions at ambient temperature. | It is well documented that arthrospores remain in the environment indefinitely. |
| **Sputum / Bronchial alveolar lavage** | Universal | BAL and sputum should be processed promptly to give the best opportunity to culture pathogenic organisms and reduce the risk of overgrowth with contaminants. | Refrigeration at 2-8˚C | Semi-quantitative results to help determine contaminant.  Comment added to samples that have had delayed processing to explain that the overgrowth may occur post collection. |
| **Corneal scrape** | Media and slide inoculated at collection | Specimens should be transported and incubated in the appropriate conditions as soon as possible. | CO2 – 37˚C  Anaerobic – 37˚c  O2 – 37˚C | Samples are dealt with using the out of hours service. |
| **Serum** | Greiner red top serum tubes |  |  | Samples are frozen at –20˚C to store for longer periods |
|  |  | Parvo IgG & IgM for 3 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 4 times without affecting the result. |
|  |  | Anti-HBs, HBeAG, HBsAg,  Syphilis, Toxo IgG & IgM - 7 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 4 times without affecting the result. |
|  |  | Borrelia IgG, Anti-HBc,  Anti-HBe, Anti-HCV,  EBV-VCA IgM, HAV IgG & IgM, HBcIgM, HBsAg neut,  HIV Ag/Ab, HTLV I/II,  Rubella IgM, VZV IgG - 7 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 5 times without affecting the result. |
|  |  | CMV IgG & IgM, EBNA-IgG,  EBV-VCA IgG - 7 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 6 times without affecting the result. |
|  |  | Mycoplasma IgM - 8 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 5 times without affecting the result. |
|  |  | Rubella IgG - 8 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 6 times without affecting the result. |
|  |  | Measles IgG, Mumps IgG - 9 days | Refrigeration at 2-8˚C | Fresh serum and plasma can be frozen and thawed up to 6 times without affecting the result. |
|  |  | Cryptococcus - Non-haemolyzed serum. Plasma, whole blood and CSF specimens are acceptable. | Refrigeration at 2-8˚C |  |
| **EDTA** | Greiner | Freshly drawn specimens (whole blood) may be held at 2 to 30°C for up to 6 hours prior to centrifugation. | After centrifugation, serum or plasma may be removed from cells. Serum or plasma specimens may be stored:   * 15 to 30°C for up to 72 hours * 2 to 8°C for up to 14 days * -20°C or colder for longer term * Cadaveric blood specimens can be stored for up to 7 days at 2-8˚C or up to 3 days at 15-30˚C following collection | Fresh serum and plasma should not be subjected to repeated freeze-thaw cycles of more than three.  If frozen, thaw specimens at 15 to 30°C or at 2 to 8°C. Once thawed, if specimens are not being processed immediately, they can be stored at 2 to 8°C for up to 6 hours. |
| **Urine (for CMV PCR)** | Universal |  | Room temp (18-25 ˚C for 4 hours maximum or  2-8 ˚C for 3 days maximum. If not, sample must be frozen at -20 ˚C for 30 days max. or at -70 ˚C for longer periods |  |
| **NPA** | Universal | Within 48 hours | Refrigeration at 2-8˚C |  |
| **Viral transport media (all viral PCR)** | Medical wire |  |  | Viral transport medium: Virocult® and ∑-Virocult® offer combined collection and transport systems for viruses, incorporating a specially formulated liquid transport medium, which is M40-A Compliant.  Virocult® demonstrates survival of many types of virus at ambient temperatures, including Herpes Simplex Virus, Varicella-Zoster Virus, Influenza Type A (includes Novel H1N1v, H5N1, and H3N2), Influenza Type B, Respiratory syncytial virus, mumps virus, adenovirus, rhinovirus, and enteroviruses. |
| **Chlamydia / N. gonorrhoea / MGEN / TV**  Urogenital swab  Extragenital (throat and rectal) swab specimens)  Urine | Alinity-M Multi-Collect specimen collection kit | Specimens should be assayed with the Alinity-M STI assay and must be processed within 14 days of collection | 2°C to 30°C up to 14 days maximum. (Abbott multi-Collect Specimen Collection Kit, Ref: 9K12-01 / AL1016.6, October 2020) | If longer storage is required, store at  -20°C +/-5oC for up to 60 days prior to testing  (Abbott multi-Collect Specimen Collection Kit, Ref: 09N19-010 / 53-608180/R1, 2020) |
|  | Urine samples that are still in the primary collection container | Transfer the urine sample into the Alinity-M multi-collect specimen tube | Store at 2°C to 30°C |  |

# Collection of samples – routine bacteriology

## Blood culture specimens:

### PUO from tropical areas

Priority should be given to diagnosing and treating Malaria investigations, these are performed by Haematology. Please contact Haematology via the hospital switchboard – 01233 633331.

If the onset of fever >38˚C was less than 21 days after leaving a tropical area, the possibility of Viral Haemorrhagic Fevers (VHF) must be considered. Such cases must be discussed with the on-call Medical Microbiologist before admission to any general ward area and collection of blood cultures.

### Fever in HIV infected patients.

In addition to routine blood cultures cases of unexplained fever in HIV positive and severely immunocompromised patients may require additional investigations. These may include…

CMV

Toxoplasma

Syphilis

Hepatitis B

Cryptococcal antigen if CNS symptoms are present (CSF can also be screened).

Fungal investigations

Routine mid-stream urine (MSU)

MSU in viral transport medium to exclude CMV

Early morning urine (EMU) x 3 for TB if appropriate (only applicable in renal or if miliary TB is suspected)

If respiratory symptoms present:

Sputum for routine bacteriology and TB

Bronchial washings or ‘induced’ sputum for Pneumocystis carinii (sputum is not useful).

Faeces if diarrhoea or 10% baseline weight loss.

**Please refer to relevant sections of this user guide for full specimen requirements and collection details.**

### Specimen collection- Blood cultures

Collect specimens as soon as possible after the onset of clinical symptoms. Although blood can be sampled at any time, drawing blood at, or as soon as possible after a fever spike is optimal, except in endocarditis where timing is less important. Empirical treatment within the first hour has been shown to maximise the chances of survival.

Please refer to Eolas Medical 

Blood culture is a culture of blood collected from a single venepuncture site inoculated to one or multiple bottles. A blood culture set is defined as one aerobic and one anaerobic bottle. For infants and neonates, a single aerobic bottle may be requested. Whenever possible it should be performed before antimicrobial therapy is commenced, however do **NOT** delay administration.

Full instructions are provided with each blood culture set, **please note** that the blood culture collection kits **DO NOT** automatically prevent overfilling, please monitor until the correct volume has been collected.

In pyrexial patients with central or tunnelled IV lines, consider taking ‘central’ blood cultures via the IV line in addition to the normal peripheral vein blood cultures. This may help to implicate an infected line as the source of sepsis. If peripheral cultures cannot be collected, take a second set of cultures from the central line using a fresh syringe, having discarded 5ml aspirated from the line.

**NB:** **Care must be taken to avoid contamination.**  It is important to fill the blood cultures with the **correct volume** of blood:

**Adults**

10mls in each bottle (1 x Aerobic (green) & 1 x Anaerobic Bact/Alert (purple)



**Paediatric**

Please refer to EKHUFT Policy Centre for Blood Culture Collection Operating Standard for volume required (4mls maximum)



**NB**: Volumes greater than or less than these recommended volumes DO NOT maintain the optimal blood to medium ratio and may affect the recovery of organisms. Monitor until correct volume is achieved – there is NO automatic shut off to prevent overfilling.

After inoculation, the blood cultures (the entire blood culture pack must be returned) **MUST be transported to the laboratory immediately. Out of hours blood cultures should be placed in the main Pathology Reception on each acute hospital site.** Blood culture bottles **must not** be placed in a refrigerator.

See Appendix 9 – Collecting a Blood Culture

## Cerebral Spinal Fluid

### Meningitis and encephalitis

Suspected meningitis/encephalitis should be treated **IMMEDIATELY** with reference to Eolas Medical for empirical therapy without waiting for confirmation. In 80% of cases antigen detection or Polymerase Chain Reaction (PCR) will yield a positive result even after antibiotic administration. It is **not** acceptable to make a diagnosis of meningococcal septicaemia in A&E and then delay commencement of therapy pending admission and clerking on the ward.

Please inform the Medical Microbiologist **and** the UKHSA local Heath Protection Team (HPT) when a patient is admitted with suspected meningitis. When meningitis is suspected, telephone the laboratory (or on call Biomedical Scientist out of normal working hours) as soon as the sample is collected to minimise delay.

Lumbar puncture should not be attempted if raised intracranial pressure is likely. Absence of papilloedema does not exclude recent onset of raised intracranial pressure. If in doubt, arrange for a CT scan to exclude raised pressure before attempting the lumbar puncture. In suspected cases of meningitis/encephalitis the samples should be collected as soon as possible:

For routine CSF examination send the following samples to Microbiology:-

CSF: 1 to 3 ml in a sterile 30ml universal container (white cap) (x2 if possible in case further tests are needed). CSF MUST be transported to the laboratory immediately after collection – time between microscopy and collection should be a maximum of 2 hours (UKHSA, 2017)



Blood: **Adults** 10mls in each bottle (1 x Aerobic (green) & 1 x Anaerobic Bact/Alert (purple))



**Paediatric** Bact/Alert (yellow)

Please refer to EKHUFT Policy Centre for Blood Culture Collection Operating Standard for volume required (4mls maximum)



**Volumes greater than or less than these recommended volumes DO NOT maintain the optimal blood to medium ratio and may affect the recovery of organisms. Monitor until correct volume is achieved – there is NO automatic shut off to prevent overfilling.**

An EDTA (purple top vacutainer) blood SAMPLE for meningococcal PCR should always be sent to Microbiology when either meningococcal meningitis or septicaemia is suspected.



If viral meningitis or encephalitis is suspected the CSF samples may be examined in house or referred to a reference laboratory (please see section 20).These tests will be at the discretion of the Consultant Microbiologist.

Tests performed on CSF:

* Bacterial Culture
* White blood cell count (WBC) and differential if the white count is raised
* Red blood cell count.
* Gram stain if WBC count is raised
* Polymerase Chain Reaction (PCR) screen for common causes if WBC is raised or at the discretion of the Consultant Microbiologist
* Bacterial antigen screen for *Streptococcus pneumoniae* when indicated.

At the discretion of the Consultant Microbiologist samples may be sent to a reference laboratory for:

* PCR for meningococci

A CSF should always be submitted for PCR when *Herpes* encephalitisis suspected.

## Faeces

Faeces specimens are primarily submitted to aid the investigation of diarrhoeal disease.

This is defined as unusual frequency of bowel action (usually at least three times in a 24hr period), passing loose, watery, unformed faeces. The consistency of the stools is more important than the frequency: frequently passed formed stools are not considered to be diarrhoea. It may be associated with symptoms such as abdominal cramps, nausea, malaise, vomiting, fever and consequent dehydration.

### Gastroenteritis

1. Routine Investigations.

All routine specimens are investigated for nucleic acids detection of:

*Salmonella spp*

*Shigella spp/*Enteroinvasive *E.coli*

*Campylobacter spp (jejuni and coli)*

*Shiga toxins 1 and 2* found in Shiga producing *E.coli (*STX*)*

*Giardia* and *Cryptosporidium*

*Vibrio cholerae*

*Yersinia*

Further investigations may be indicated based upon clinical details provided and Consultant Microbiologist discretion. It is essential that full, clear and comprehensive clinical details are provided with all stool samples to ensure appropriate testing is performed (refer to section 15.1 Specimen requisition for further information on clinical details).

### Additional testing when clinically indicated

1. Rotavirus, Adenovirus and Sapovirus PCR is routinely performed on all samples from children <6 years of age
2. Travel associated diarrhoea.

Please ensure full travel details are provided

Ova, Cysts and Parasite investigation

1. Food poisoning: – Please discuss with Consultant Microbiologist prior to submitting samples.

### Clostridium difficile investigation.

Toxin B gene detection by PCR are available to aid the diagnosis of *Clostridium difficile* disease.

Testing is only performed on diarrhoeal stool specimens (liquid or semi-formed faeces, Bristol stool type 5-7), which should be collected as described in section 17.3.6.

Morbid anatomy samples should be received in appropriate plain screw capped CE leak proof specimen containers which are **free from preservative.**

#### Repeat Clostridium difficile testing

Repeat toxin testing need only be performed on previously Toxin negative samples.

Specimens from toxin positive patients need repeating only once every 28 days, unless the patient shows clinical symptoms of relapse.

Repeat testing for *C.difficile* toxin is NOT normally required within 28 days of a previous toxin positive result. However, toxin testing may on occasion be requested by medical staff if there is a clinical relapse after treatment in the period 14 – 28 days after a previous positive toxin result. Such samples still need be to be examined.

**NB**: Stool samples for *C.difficile* ‘clearance’ are **not** required. 20 – 30% of patients with *C.difficile* may relapse following treatment. The Infection Prevention and Control team **must** be contacted for advice before sending a repeat faecal specimen.

### Gastritis

*Helicobacter pylori*

A stool sample for antigen detection is the required sample for laboratory confirmation of current infection or treatment efficacy.

Treatment regimes are available in the British National Formulary (BNF) or in Eolas Medical 

### Norovirus infection

Norovirus PCR is available locally on liquid faecal samples for the investigation of outbreaks of diarrhoea and vomiting, please contact Infection Control prior to submitting requests if an outbreak is suspected.

### Faecal Specimen collection

Faecal specimens should be submitted to the laboratory in an appropriate plain screw capped CE leak proof specimen container (do not use flimsy plastic sputum cups) as soon as possible after collection.



Submit a sample that fills at least a third of the container if possible, but **please do not overfill the container**. Samples must be transported to the laboratory as soon as possible after collection as pH changes that occur in faecal specimens may affect the survival of important pathogens such as *Shigella*. The Patient should be given instructions on collecting a faecal specimen by the requesting clinician or directed to the EKHUFT website: [The Collection of a stool (faeces) sample (ekhuft.nhs.uk)](https://leaflets.ekhuft.nhs.uk/the-collection-of-a-stool-faeces-sample/) (see also Appendix 2).

Faecal specimens of Bristol stool consistency 1-4 will not be tested and specimens discarded unless indicated by a Consultant Microbiologist.

**CLINICAL DETAILS (refer to section 15.1 Specimen requisition) MUST BE PROVIDED, INCLUDING:**

Nature and duration of the illness

Onset date

Travel history (countries visited plus dates) Antibiotic history, if relevant

In the case of outbreak or suspected food poisoning, notify the Medical Microbiologists and the Consultant in Communicable Disease Control (CCDC) immediately.

**NB**: sputum, pus and urine samples may also be submitted to the laboratory for parasitic investigations. Please collect the specimen into an appropriate clean CE marked container.



## 

## Parasites

### Faecal microscopy

Faecal samples for microscopy must be collected as described in section 17.3.6 and transferred to the laboratory as soon as possible.

Please submit three faecal samples collected over a maximum of a 10-day period.



### Ectoparasites

Lice, ticks, bedbugs, larvae and pupa must be collected into an appropriate plain screw capped CE specimen container.

 Sterile container with metal cap

### Endoparasites

Whole tapeworm, roundworm and proglottids must be collected into an appropriate plain screw capped CE specimen container and enough saline added to the container to cover the specimen.

 Sterile container with metal cap

### Urine for Schistosomiasis

Total urine for Schistosomiasis must be collected between 10am and 2pm after exercise into an appropriate plain screw capped CE leak proof specimen container. Alternatively submit terminal collections over a 24-hour period. Please do not use boric acid containers.



### Jejunal aspirates

Jejunal aspirates must be collected into an appropriate plain screw capped CE leak proof specimen container.

 Sterile container with metal cap

### Puritis ani

Sellotape slides for threadworms (*Enterobius vermicularis*)

The specimen must be collected before the patient has defecated or washed in the morning:

Method of collection:

* Label a glass slide with a minimum of 2 of the following:

Patient’s name

Patients date of birth

Patient’s hospital number

* Open the gluteal folds to visualise the anus - using sellotape press the sticky surface of the tape on several places on the skin around the anus.
* Attach the sticky surface of the sellotape to the glass slide
* Send the slide to the laboratory in a slide container

## Tissue and fluid samples

Many aerobic and anaerobic organisms can be associated with infections of the subcutaneous tissue, joint fluids, prosthetics, and internal organs. These organisms may enter the body through various means e.g. breaks in the skin or mucous membranes through trauma, as a complication of medical treatments (surgery, implanted devices), and through the blood and lymphatic system from another site of infection. Optimal samples for culture are tissue or biopsies and aspiration by needle and syringe of the involved areas.

### Tissue samples

Please collect tissue and biopsy specimens into an appropriate plain screw capped CE leak proof specimen container.

Gastric biopsies for Helicobacter pylori are sent to a reference laboratory for examination (please see section 20).

For theatre tissue samples and fluids e.g. joint revisions please use the specimen request form embedded in Appendix 15.

 Sterile container with metal cap

### Fluid samples

The detection of organisms in fluids that are normally sterile indicates significant infection, which can be life-threatening.

**NB:** for microscopy the absence of bacteria seen does not exclude infection.

### Ascitic fluid

Tests performed typically include WBC count, Gram stain and culture

Submit the specimen in an appropriate plain screw capped CE leak proof specimen container.



TB cultures should be requested when risk factors are present.

### Pleural and pericardial fluids

Tests performed include Gram stain and culture.

Submit in a CE Marked leak proof container; indicate whether a TB culture is required.



### Joint fluids

Tests performed include Gram stain and culture

Submit in an appropriate plain screw capped CE leak proof specimen container.



If an examination for crystals is required, submit a separate sample to the Histopathology/Cytology department where polarising microscopy is available.

### CAPD Fluids

CAPD fluids may be submitted in an appropriate plain screw capped CE leak proof specimen container. Blood culture bottles may also be used, but a separate fluid sample should be sent if a WBC estimation and differential cell count is required.

 and / or 

## Urine specimens

### Tests: Microscopy, culture and sensitivity

Urine microscopy is automated, a result of <50 leucocytes/mm3 is considered to be negative and these specimens are not routinely cultured. Samples with > 50 leucocytes/mm3 / 50-100 leucocytes/mm3 and >100 leucocytes/mm3 will be automatically cultured. Specimens from pregnant women, renal unit patients, ITU patients and children of 12 years of age and under will undergo routine bacterial culture irrespective of the microscopy results. For Legionella / Pneumococcal urine antigen testing please refer to section 17.16.

### Urine sample collection

1. **Mid-stream Clean Catch Urine**

This is suitable for most routine bacteriological examinations.

The patient should be given a red capped urine collection tube containing borate preservative, labelled with their name, date of birth and the sample collection date.



The urine collection tube must be filled to within the two lines indicated (minimum of 3 ml, maximum of 7ml) please do not obliterate the lines by adhering labels over them.

Instructions on collecting a mid-stream urine should be given to the patient by the requesting clinician or be directed to the EKHUFT website: [The Collection of a random urine sample (ekhuft.nhs.uk)](https://leaflets.ekhuft.nhs.uk/the-collection-of-a-random-urine-sample/) (see also Appendix 1)

1. **Bag Urine**

The baby’s external genitalia must be thoroughly washed, dried and a self-adhesive plastic collecting bag applied. This is removed when sufficient urine has been collected (please see Appendix 1). Interpretation of results can be complicated because of the high incidence of contamination. A Supra Pubic Aspirate (SPA) or ‘clean catch’ sample may be useful in such cases. Please submit a minimum of 7ml.

1. **Suprapubic Bladder Aspirate (SPA)**

A needle and syringe is used to sample bladder urine directly using a suprapubic percutaneous approach (please see Appendix 1). This produces an ideal sample for bacteriology, because the problem of urethral contamination is avoided. It is particularly useful for sampling baby’s urine. Transfer the urine specimen into an appropriate CE marked specimen container.



Please submit a minimum of 7ml.

1. **Catheter Urine (CSU)**

A catheter SAMPLE of urine should only be sent for laboratory culture if the patient has **signs of clinical sepsis** (i.e. fever or chill, associated localised loin or suprapubic tenderness) and **not** because the appearance or smell of the urine suggest that bacteriuria (bacteria in the urine).

Using a strict aseptic technique, a syringe is used to aspirate 10ml of urine from one of the self-sealing ports in the drainage tube. Urine from the collecting bag is unsuitable for culture. The closed circuit drainage system should never be disconnected to collect a urine sample (please see Appendix 1).

For MC&S, please use a red capped urine collection tube containing borate preservative



For all other catheter urine tests (with the exception of MRSA screening) please use an appropriate plain CE marked specimen container.



Even closed circuit urine collection systems become colonised with bacteria about 5 days after catheter insertion. Presence of bacteria in catheter urine is therefore a normal finding after about 5 days and is not an indication for antimicrobial therapy, unless there are clinical signs of systemic sepsis. CSUs are often of doubtful clinical value and should not be collected routinely.

1. **Urine samples for Chlamydia**

Please ensure that a first catch is collected using the Alinity m multi-Collect Specimen Collection Kit – please see Appendix 6 for instructions



Alternatively, a minimum of 10mls first catch must be collected in a plain universal (**not** a red top containing borate). Please ensure that the sample is received by the laboratory within 24hours.



## Wound swabs and Pus

The skin is colonised by normally non-harmful flora. Infections of the skin and subcutaneous tissues are caused by a wide range of organisms, however the majority are caused by Staphylococcus aureus and β haemolytic streptococci groups A, C and G. Particular organisms are often typically associated with specific clinical conditions. Microbiological cultures may be undertaken to establish the causative organism enabling antibiotic sensitivity testing which is essential to ensure optimal treatment regimens.

Abscesses are accumulations of pus in tissue and any organism isolated from them may be of significance. They occur in many parts of the body as superficial infections or as deep-seated infections associated with any internal organ. Many abscesses are caused by Staphylococcus aureus alone, but others are mixed infections associated with a wide range of organisms. The site of the wound and **clinical details (refer to section 15.1 Specimen requisition) must be stated** on the request form so that the appropriate media can be used. Please take **only one swab** from a wound on any one occasion.

Tests performed: Gram stain (where appropriate) and culture

Pus should be sent to the laboratory in an appropriate plain CE marked specimen container rather than on a swab whenever possible.



Wound swabs – please use Charcoal swabs (black lid)



Chronic leg ulcers with cellulitis – collect swabs after removing necrotic debris, please see Appendix 11.

Chronic leg ulcers / pressure sores with no cellulitis and no fever – it is not useful to send swabs from these.

**NB:** Swabs are usually unsuitable for TB Culture; please consult the laboratory if TB is suspected.

## 

## Throat swabs

Throat related specimens are one of the most commonly performed procedures in patients with upper respiratory tract infections. Whilst the majority of throat infection are viral in nature throat swabs can be useful in identifying or eliminating bacterial causes.

For routine bacteriology culture use charcoal swabs (black lid), please see Appendix 11.



In cases where diphtheria is suspected consultation with the Consultant Microbiologist is recommended and relevant clinical details must be provided to ensure appropriate processing (refer to section 15.1 Specimen requisition).

## Nose swabs

Of limited diagnostic value nose swabs may be useful for detecting the carriage of certain pathogens such as:-

Group A Beta Haemolytic Streptococci

Corynebacterium diphtheria

Staphylococcus aureus

Dry swabs are not suitable. In the absence of discharge, moisten a swab in a bacterial transport medium before taking the swab. Please use a thin wire Charcoal swab (orange lid), please see Appendix 11.



## Pernasal swab (Pertussis culture)

Bordetella culture only

For the isolation of *Bordetella pertussis* in cases of whooping cough.

A special wire pernasal swab and transport medium is required, (Thin wire Charcoal swab (blue lid) Available from Microbiology WHH).



Please see Appendix 11.

If a pernasal swab is not available, a throat swab using a charcoal swab (black lid) can be used.



Please see Appendix 11.

Nose swabs are not suitable for Bordetella culture and will not be processed by the laboratory.

## Eye swabs

Infections of the eye can be caused by a variety of microorganisms which may be introduced to the eye via hands, fomites (e.g. contact lenses), traumatic injury or following surgery.

Tests performed include Gram stain and culture

Purulent discharge is suitable for bacterial culture.

Please use a Charcoal swab (black lid), please see Appendix 11.



## Corneal scrapes

Eye kits for corneal scraping are available from the Microbiology reception upon request.

Pus from cases of endophthalmitis can be sent in a syringe.

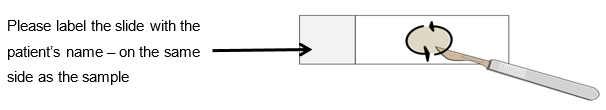
Collect sample and inoculate plates by spreading the sample on the **surface** of the agar plate on the top portion of the plate only (see diagram below).

(Please do not cut into agar surface – this makes it difficult if not impossible to interpret any growth that may occur)

Inoculate the top portion of plate only

Please inoculate ALL plates if possible

Inoculate slide in circular motion:



Return plates & slide in box with a request form to Pathology Reception.

**Contact On-Call Biomedical Scientist for Microbiology via switchboard if sample collected after 7 pm.**

Specimens requiring Acanthamoeba culture are referred by the Microbiology laboratory to the London School of Hygiene & Tropical Medicine. Please contact the laboratory if clinically indicated.

## Ear swabs

Swabs may be taken and submitted to the laboratory to aid the diagnosis of both otitis externa and media.

For ear swabs, please use thin wire Charcoal transport swabs (orange lid), please see Appendix 11.



## Genital swabs

Appropriate specimens are often difficult to obtain, particularly from women, and incorrect or sub-optimal specimens are often received. It is important to avoid contamination with faecal flora during collection of specimens.

Vaginal/Endocervical discharge

A high vaginal swab (HVS) in bacterial transport medium (Charcoal swab, black lid) is used for routine detection of the following pathogens:

• *Candida* sp

• Βeta haemolytic Streptococci (including Group B streptococci in pregnant females)



The following can be tested if specifically requested:

• Bacterial vaginosis

* *Trichomonas vaginalis*

Gonorrhoea is diagnosed by culturing endocervical, urethral and rectal swabs using the normal charcoal bacteriology swabs. (High vaginal swabs are not suitable because Neisseria gonorrhoea does not grow in the squamous epithelium of the vagina).

For Herpes simplex, swab any visible lesion and break off the swab into viral transport medium (VTM). An endocervical swab in VTM may be useful in patients who have a past history suggestive of Herpes simplex but do not have identifiable lesions.



Pelvic Inflammatory Disease – Pouch of Douglas fluid is the best sample for the diagnosis of deep PID. Collect into an appropriate plain screw capped CE marked leak-proof specimen container.



## Rectal swabs

Rectal swabs may be useful for the detection of carriage of antibiotic resistant organisms such as Carbapenemase Resistant Enterobacteriaceae, Vancomycin Resistant Enterococci and ESBL producing organisms.

Rectal swabs must be taken using black lidded charcoal swabs (there is no need to moisten the swab with normal saline).



The swab must be gently inserted through the anal sphincter, rotated through one full turn and then withdrawn.

## Pneumonia

Sputum culture

Routine culture to detect common respiratory pathogens includes:

* *Streptococcus pneumoniae*
* *Haemophilus influenza*
* *Staphylococcus aureus*
* *Moraxella catarrhalis*

Fungal culture and microscopy may also be performed where clinically indicated.

Specimens of sputum must be freshly collected into sterile containers.

 Sterile container with metal cap

Where TB investigations are required please refer to section 17.16.1 of this user manual for details.

When Legionella infection is suspected, please indicate the diagnosis clearly on the request **and request an urgent urine examination for Legionella antigen.**

* Urine samples for Legionella / Pneumococcus antigen detection are only processed when agreed by a Consultant Microbiologist. Urine must be collected in an appropriate plain CE marked screw-capped specimen container – a minimum of 3ml. Please ensure that the sample is received by the laboratory within 24 hours.



* In cases of suspected pneumonia, serology (using red top serum tube) may be useful for the detection of non-culturable and viral pathogens:
* *Mycoplasma sp*
* *Chlamydia sp* (including Psittacosis)
* *Coxiella burnetti* (Q Fever)
* *Avian precipitins*



Pneumonia (submitted with blood culture)

Infective exacerbation of COPD

Chest infections unresponsive to antibiotic therapy (give details)

Screening of Immunosuppressed and ITU patients

Sterile container with metal cap 60ml container (silver lid)

Samples for CYTOLOGY should be submitted separately with the appropriate request to the Cellular Pathology Department

Patients should be instructed that true sputum is required and not saliva (i.e. collected after a deep cough; a post physiotherapy sample is ideal.

When sputum is unobtainable, a bronchial washing obtained via bronchoscope is an alternative.

Gastric washings may occasionally be useful in young children when sputum is unobtainable.

### Mycobacterium (TB)

When TB is suspected, at least three samples should be submitted on consecutive days, in addition to a routine sample if required. Please refer to ‘appropriate samples for TB below. TB culture result may take 6 weeks. Microscopy for Acid Fast Bacilli in the urine is unlikely to be helpful and will not be performed routinely.

**Appropriate samples for TB**

1. Sputum/broncho-alveolar lavage

Sputum specimens should be less than 1 day old to minimise contamination. Approximately 5ml per sample early in the morning on three consecutive days should be collected. When the cough is dry, physiotherapy, postural drainage or inhalation of nebulised saline before expectoration may be helpful.

Sterile container with metal cap 60ml container (silver lid)

1. Broncho-alveolar washings / bronchial washings (BAL)

– please collect a minimum sample size of 5ml if possible.



1. Early morning urine examination

Three consecutive COMPLETE early morning urine samples are collected on separate days into separate appropriate plain 250 ml CE marked specimen containers, available from Microbiology at WHH.



These must be clearly labelled for TB culture. The three containers must be transported to the laboratory together (container for day 1 and 2 collections must be stored in a refrigerator prior to sending to the laboratory with day 3 sample). If there are no appropriate containers for whole early morning urine (EMU) sample, a midstream EMU sample is an acceptable, but not ideal alternative.

EMUs are usually requested when ‘sterile pyuria’ has already been demonstrated on routine urine culture.

1. Pus examination e.g. from cold abscesses

Investigation for Mycobacteria may be appropriate from selected sites where routine culture is negative. Pus or pus swabs should be collected aseptically and the largest practical sample submitted in an appropriate plain screw capped CE leak proof specimen container

1. Faeces examination

Mycobacteria investigation may be requested on selected samples only. The value of such a sample is questionable and they are not commonly submitted though they may be valuable where disseminated disease is considered in patients who are immunocompromised. It is true however that Mycobacteria (of several species) have been incriminated in bowel disease. Culture is not the best technique available and should only be undertaken if the clinician fully understands the problems.



1. Tissue examination

Skin, tissue e.g. curetting, lung, lymph node or post mortem specimens should be collected aseptically in an appropriate plain screw capped CE leak proof specimen container without preservatives and add sterile distilled water to prevent desiccation.

Sterile container with metal cap 

A representative portion should be selected if possible: the majority of organisms will be found in the periUKHSAry of a caseous lesion. As large a sample as possible should be sent to the laboratory.

1. Gastric lavage / washings examination

Gastric lavage / washings are usually taken instead of sputum where a patient is unable to expectorate e.g. for children where there are problems obtaining sputum. Ideally 3 early morning samples prior to food intake (before breakfast) are collected over 3 consecutive days. Preferably a minimum volume of 5ml should be provided. Aspirates should be promptly delivered to the laboratory to avoid acidic deterioration of organisms.



1. Sterile fluid examination e.g. CSF, pleural fluid, joint fluid, ascitic fluid

Gross fluid production in these sites is abnormal. Where no microbiological or other underlying cause is proven, investigation for Mycobacteria may be appropriate. Collect aseptically as much CSF sample as possible into an appropriate plain screw capped CE leak proof specimen container.



A minimum of 1ml of other fluids is required. If a small volume is available after the initial lumbar puncture and the findings of cell counts and protein suggest TB meningitis, a second procedure should be considered to obtain a larger volume to improve the chance of achieving a positive culture.

1. Blood / bone marrow + blood samples

Investigation for disseminated mycobacterial infection will be performed on selected patients only; usually immunocompromised, often HIV positive patients.

1. QuantiFERON/ T Spot test:

Please refer to serology section for specimen details.

## MRSA Screen

Pre-admission screens, admission screens and long-stay (ward) areas – nose and groin using clear broths with white caps (refer to EKHUFT Policy Centre for Management and Control of Methicillin Resistant Staphylococcus aureus (MRSA) for guidance on the screening process).



For wounds / skin lesions, and sites of invasive devices, charcoal swabs (black lid) must be used.



Catheter urine – add 1ml of catheter urine to MRSA screening broth (white cap).



GP Screens

We recommend using a routine charcoal swab (black lid) to collect a sample from the nose and groin.



Routine screens should only be carried out according to protocols in the Infection Control Manual.

Staphylococcus aureus screen (renal unit)

We recommend collecting screening swabs from the nose and groin, combining them into a staphylococcus screening broth (white cap)



See EKHUFT Policy Centre for Management and Control of Methicillin Resistant Staphylococcus aureus (MRSA) policy document.

## Mycology samples

Submit:

* Hair plucked close to root (for scalp infection) and/or plastic sampling brushes available from large Pharmacies.
* Nail clippings (plain CE marked screw-capped specimen container).

Sterile container with metal cap 

Clean the site using a 70% alcohol wipe, removing any varnish that may be present. Using a pair of sterile nail clippers, clip small samples from the edge of the nail, collect into a sterile pot labelled with the patient details. If the lesion is not at the edge of the nail it can be scraped with a sterile scalpel.

* Skin scrapings – Dermapak containers (available from Microbiology WHH).

Clean the lesion of loose material, debris and any creams that may be present using a 70% alcohol wipe. Scrape the edge of the lesion with a sterile scalpel to remove loose skin scales and collect into a Dermapak container. If vesicles are present, remove the roofs with sterile scissors or scalpel and collect in the Dermapak container.

**NB:** Wound swabs are unsuitable for the diagnosis of dermatophyte fungi

**PLEASE DO NOT STICK SAMPLES TO SLIDES WITH SELLOTAPE. IF NO DERMAPAK AVAILABLE PLEASE USE A SUITABLE CE MARKED SPECIMEN CONTAINER.**

# collection of sample - virology / serology

If unclear which sample type is required for a test, please consult a Medical Microbiologist or the Senior BMS in Virology before collecting samples.

A virology diagnosis may be achieved by:

* 1. Serological methods including
     1. Detection of specific IgG and IgM
     2. Detection and quantification of antibody for immunity levels (titres)
     3. Detection of viral antigen in blood e.g. Hepatitis B surface antigen
     4. Detection of viral antigens in faeces e.g. Norovirus
  2. Nucleic acid detection by qualitative methods (Respiratory and herpes group PCR) or quantitative methods (HIV, HCV, HBV and CMV viral load testing)

## Blood samples for serology/virology.

The majority of serology/virology blood tests can be performed on serum (red top) specimens, SST (yellow top) specimens or EDTA (purple tops). For all PCR tests e.g. HCV Viral load an EDTA sample **MUST** be submitted.

It is often not possible to add tests on to samples from blood sciences due to short retention times and volume of sample used. Wherever possible it is preferable to send an independent sample for serological testing.

Refer to EKHUFT Policy Centre for Venepuncture Guidelines on taking blood specimens appropriately and safely.

## Antenatal screening specimens

Microbiology Service offer an Antenatal Screening Service on serum (yellow top SST tubes)

SST (yellow top) specimen tubes:



Please ensure patient consent is recorded on the request form

## QuantiFERON/ T spot samples for diagnosis of latent TB

QuantiFERON and T spot testing requires a strict coherence to the protocol:

1. Tests can be sent Monday to Friday inclusive, with the exception of Bank Holidays and the period between Christmas and New Year, when the reference laboratory facility is shut. Any variances from this will be communicated by Trust-wide email.
2. It is recommended that the case is discussed with a respiratory physician prior to requesting. The significance of both positive and negative results needs to be carefully assessed both in the light of clinical findings and the results of other investigations. Positive tests should be interpreted with caution in low risk populations. A negative result does not exclude active infection.
3. QuantiFERON and T Spot testing requires lithium heparin blood (green top tubes):



Adults and Children ≥10 years old (QuantiFERON): 1x 6ml or 2x 4ml tube (if insufficient is sent the sample will not be processed)

Children 2-9 years old: 2x 4ml tube

Children up to 2 years old (T SPOT): 2x 2ml paediatric tube

1. Specimens **MUST** be received by the WHH Microbiology laboratory by **3pm** on the same day the specimen is taken.

This is because:

* 1. The test is a timed test and specimens must be tested within **12 hours** of being taken.
  2. The specimens need to be processed for booking onto the laboratory system and paediatric specimens for T SPOT need to be packaged for courier collection.

1. Deadlines for collecting samples to ensure they arrive in Microbiology by 15:00 are:

WHH: 14:30

K&CH: 14:00 (to catch 14:15 transport van, which arrives at the WHH at 15:00)

QEQMH: 13:30 (to catch 13:45 transport van, which arrives at the WHH at 14:55)

1. If there is an unavoidable delay in sending the lithium heparin tubes for receipt the same day, they may be stored at 2-8 °C for 16-48 hours, provided this is within 3 hours of collection.
2. Results will be available from 16:30, 2 days after the specimen is sent, e.g. if the specimen is sent on a Monday the results are available from 16:30 on Wednesday. Results for specimens received on a Thursday or Friday are available from 16:30 on the following Monday.

## HIV Viral Load Monitoring (specialist test)

Submit two fresh EDTA (purple top) blood samples to the laboratory with details of all current anti-viral therapies and current CD4 count.



The country of origin of HIV infections is useful to help exclude spurious low results caused by variant HIV strains.

## HCV, HBV and CMV viral load monitoring

Submit a fresh EDTA blood to the laboratory (cannot be added onto other EDTA tests).



These tests are run daily with the exception of planned and unplanned downtime, however if an urgent result is required, please contact the laboratory to arrange urgent testing.

## Hepatitis

Serological tests

**First line tests:**   
Hepatitis A IgM

Hepatitis B surface antigen

Hepatitis C virus

(Consider Hepatitis E if patient has high LFTs).

**Second line Serology:**

CMV

Toxoplasmosis

EB virus

Leptospirosis

Parvovirus

Specialist Hepatitis investigations by special arrangement with the laboratory.

Note: These tests are expensive and we reserve the right to decline requests if thought inappropriate by the Consultant Microbiologist. Hepatitis C testing should be in line with ‘NICE’ treatment guidelines.

## BAL/lower respiratory samples for Pneumocystis.

If respiratory virus PCR is required from a BAL/sputum please make this clear on the request form, otherwise it will be processed for routine bacteriology.

Samples must be in sterile universals with no additives.



In cases where Pneumocystis infection is suspected a BAL sample is preferred. Sputum can be tested but the results are less well defined. The BAL can be tested after routine culture/TB testing as long as the need is clear on the request form. Pneumocystis can be detected via PCR (via reference laboratory). Please discuss with a consultant microbiologist before requesting.

## NPA (Nasopharyngeal aspirates) for RSV and PCR

NPA is the sample of choice for respiratory PCR on neonates and young children. Around 5 mls (if possible) should be collected into a conical bottom tube.



Routine samples will be initially tested for Influenza A, Influenza B, RSV and COVID-19 RNA/PCR. If this is negative a full respiratory PCR will be performed on immunocompromised patients or SCBU specimens.

## Throat / nose swabs in viral transport media (VTM) or Sigma MM transwabs for PCR

Submit a nose and/or throat swab in VTM (clear fluid, green cap)



or using Sigma MM transwab (white cap, green swab),



available from microbiology on request, to look for respiratory viruses (influenza / COVID / RSV / Extended Respiratory Panel) via PCR (as with NPA samples). See Appendix 13 for instructions on taking influenza and COVID samples.

Samples are retained for 1 week after testing in case further viral studies are needed.

In cases of suspected influenza outbreaks, please discuss with infection control or a consultant microbiologist prior to screening.

## Genital, lesion and mucosal swabs for Herpes Simplex in VTM.

Swabs for HSV 1 + 2 testing should be taken in VTM (clear fluid, green cap).



If Enterovirus is required (i.e. vesicular rash,?hand foot and mouth) the sample will be sent to a reference laboratory for PCR. Please discuss with a consultant microbiologist.

## Other swabs in VTM (eye swabs, skin swabs, vesicle swabs/fluid etc)

The relevant tests will be performed, dependant on the site of the swab and clinical details provided. Eye swabs will be tested for Adenovirus, HSV, Enterovirus and VZV, skin swabs will be tested for HSV, VZV and Enterovirus (if required).



## CSF

All CSF samples tested by microbiology will then be triaged by the consultant microbiologists to decide if viral PCR is required. The transport and retention is described in section 17.2 of this document. In cases of raised WBC counts (>50 WBC seen) a Biofire film-array PCR test will be performed automatically to test for common causes of bacterial, viral and fungal meningitis. Further PCR tests and CSF’s with low WBC counts will be tested at the Consultant Microbiologists’ discretion.

Cryptococcal antigen testing can also be performed from CSF (or serum).

## Urine for PCR (CMV)

Submit an MSU in a plain white-top universal to the lab, promptly after collection



Red top urines **cannot** be used as the boric acid interferes with the PCR. A minimum of 1 ml is required.

## Faeces samples for virology

Tests for Enterovirus, Adenovirus, Rotavirus, Norovirus, Astrovirus and Poliovirus can be performed from faeces. Adenovirus, Rotavirus and Norovirus are performed in-house via the enteric section. If further viral studies are required, the sample can be sent to the reference laboratory for investigation.



## Salivary samples for Mumps testing

When mumps is suspected a salivary sample can be sent for Mumps PCR. This test is NOT carried out by the laboratory and arrangement must be made with the Kent UKHSA.

Telephone 0344 225 3861 option 1 to arrange delivery of a collection kit and instructions.

## Samples for Chlamydia, Gonorrhoea and HSV PCR testing.

Only use Abbott Alinity m swabs for any Chlamydia and/or Gonorrhoea testing. Any other swab types will **NOT** be tested.

Routine genital swabs for Chlamydia and/or Gonorrhoea and/or HSV1+2 PCR testing should be taken according to the instruction insert (see Appendix 3, 4, 5, 7 & 8).

For chlamydial eye swabs first use a fine wire swab (e.g. ENT thin wire Charcoal swab with orange lid) to wipe the purulent discharge from the lower conjunctivae, this can then be sent for bacterial examination if required (NOT viral). Secondly, take an Abbott Alinity m swab collection kit and firmly wipe the now clean epithelium of the lower lid. Replace the swab into the container and send to the laboratory.



For urine samples please submit the sample in the Abbott Alinity m urine collection tube. Please see Appendix 6.

If this is not possible a clean white leak proof universal container may be used, but please do not submit urine in white top universals on Fridays.

 or 

## 18.17 Summary table of sample collection for in-house tests:

| **Sample type** | **Pathogens investigated in-house** | **Referred tests** | **Sample container** |
| --- | --- | --- | --- |
| Blood | HIV  Syphilis  CMV  Rubella  Toxoplasma  Hepatitis A  Hepatitis B  Hepatitis C  Hepatitis E  HTLV  EBV  Parvovirus  Mycoplasma  VZV IgG  Measles IgG  Mumps IgG  Lymes IgG | ASO titres  Amoebic IFAT  Anti Hyaluronidase  Arbovirus antibodies  BD glucan  Brucella serology  Cat scratch serology (Bartonella)  Candida ppts  Coccidiomycosis Cystercercosis IFAT  Entamoeba histolytica  Electron microscopy  Ethambutol levels  Farmers lung ppt  Filaria  Hepatitis D Abs  HIB antibody  Histoplasma  HSV 1 serology  HSV 2 serology  Hydatid CFT  Hydatid Elisa  Itraconazole level  Isoniazid levels  Leishmaniasis  Leptospira serology  Lyme IgM serology  Malaria IFAT  Meningococcal C post vax level  Mumps IgM  Measles IgM  Netilmicin level  Pertussis Ab level  Posaconazole levels  Pneumococcal abs  Poliovirus ab  Q fever  Rabies ab level  Rickettsia Ab  Rifampicin levels  SARS studies  Schistosomal Elisa  Strongyloides Elisa  Teicoplanin level  Tetanus ab level  Toxocara  Trichinella IFAT  Yersinia enterocolitica abs  Yersinia Pseudo TB ab | Serum red top Vacutainer,  SST yellow top Vacutainer |
| Blood / serum/ BAL |  | Galactomannan ag | Serum red top Vacutainer,  SST yellow top Vacutainer |
| Blood | HIV1 viral load  CMV viral load  HCV viral load  HBV viral load  VZV PCR | Adenovirus PCR  BK virus PCR  CMV PCR (in conjunction with other tests)  EBV PCR  HCV and HBV Genotype  HHV6 PCR  Therapeutic drug monitoring (HIV)  HHV7 PCR  HIV2 viral load  HIV pro-viral DNA  HLA testing  HSV PCR  Parvovirus PCR  Polyomavirus PCR  Anti-retroviral resistance | EDTA purple top Vacutainer |
| NPA, Nose/throat swabs, BAL, sputum | RSV A + B  Influenza A H  Influenza B  Mycoplasma pneumoniae  Rhinovirus A/B/C  Human Metapneumovirus  Parainfluenza 1,2,3,4  Coronavirus OC43, 229E, NL63  Bocavirus 1/2/3/4  Enterovirus  Adenovirus  CMV | MERS Coronavirus  Pneumocystis PCR  Influenza typing | Sterile conical bottom universal,  Sterile Universal  VTM (green top swab)  Sterile universal |
| Genital swabs | HSV | Syphilis  Haemophilus ducreyii | VTM (green top swab) |
| Skin swab | HSV  VZV | Enterovirus | VTM (green top swab) |
| Eye swabs | VZV | Adenovirus, HSV, Enterovirus | VTM (green top swab) |
| CSF | HSV 1+2  VZV,  CMV,  Enterovirus  Parechovirus  E.Coli K1,  Haemophilus influenza  Streptococcus pneumonia  Streptococcus aggalactiae,  Listeria monocytogenes,  Neisseria meningitides  Cryptococcus neoformans/gattii | EBV  JC/BK virus  16S bacterial PCR  18s fungal PCR  Whipples  TB PCR fastrack | Sterile universal |
| Pericardial fluid |  | Enterovirus, CMV, EBV, Adeno | Sterile universal |
| Faeces |  | Enterovirus, CMV | Blue topped faeces collection pot |

# storage of collected specimens prior to transferring to the laboratory

Exposure to significantly raised or reduced room temperatures may affect the results obtained during laboratory investigations. If the appropriate sample transport media or containers are used, samples for microbiological investigation should generally be stored in dry conditions at room temperature and not refrigerated. To ensure that results are clinically useful please store samples as shown in the table below:

| **Sample type** | **Storage conditions** |
| --- | --- |
| Swabs | Store between 2-30˚C in dry conditions away from heat and out of direct sunlight. Transfer to the laboratory as soon as possible after collection. |
| Faeces  Blood  Aspirate, washing, pus, tissue, Fluid | Store at ambient temperature in dry conditions away from heat and out of direct sunlight. Transfer to the laboratory as soon as possible after collection. |
| Urine:  Boric acid container (red cap) – DO NOT under or overfill – adhere to the fill lines on the side of the sample container (UKHSA, 2018). | Urine samples should be transferred to the laboratory as soon as possible after collection.  Sample collected in a boric acid container should maintain the sample quality for up to 96 hours prior to processing at ambient temperature in dry conditions. |
| Non-boric acid container (white cap) | All urine sample collected in non-boric acid containers should be refrigerated to preserve sample quality. |

# referring samples to other laboratories

It may be necessary to refer samples to the reference laboratory for full identification of the samples which are not within the remit of this laboratory. The table below indicates which isolates need to be referred.

Serological tests referred to reference laboratories:-

| **SEROLOGICAL TEST** | **REFERENCE LAB** |
| --- | --- |
| Actinomyces  Pertussis | Bacterial Reference Department (BRD) UKHSA Colindale |
| HLA B\*5701  Therapeutic Drug Monitoring | Cambridge Clinical Laboratories 184 Cambridge Science Park |
| Isoniazid  Ethambutol | Cardiff Toxicology Laboratory Wales |
| Amoebic IFAT  Entamoeba histolytica  Filaria  Giardia IFAT  Malarial Antibody  Toxocara | Department of Parasitology HSL The Halo London |
| HIV2 viral load | Department of Virology University College London Hospitals NHS Foundation Trust (UCLH) |
| Hydatid | Hydatid Reference Centre School of Tropical Medicine Liverpool |
| Haemophilus influenza B  Pneumococcal Abs  HIB Post Vax  Tetanus | Immunology Laboratory Churchill Hospital Oxford |
| Syphilis serology | Infections Sciences Pathology Sciences Building Southmead Hospital UKHSA Bristol |
| Pneumococcal (Invasive disease) | Meningococcal Reference Unit (MRU) UKHSA Manchester |
| Adenovirus PCR  BK Virus  CMV PCR (only used if can’t do in-house)  EBV PCR  Enterovirus PCR  Galactomannan screening (BAL)  Hepatitis B PCR (if not tested in-house)  Hepatitis C genotype  Human Herpes Virus 6 PCR  Human Herpes Virus 7 PCR  Human Herpes Virus 8 PCR  JC Virus  Measles PCR  Mumps PCR  Parvovirus B19  Syphilis PCR  Viral PCR (CSF) (only used if can’t do in-house)  Whipples | Micropathology |
| Candida precipitins  Coccidiomycosis  Galactomannan (serum)  Histoplasma | Mycology Reference Laboratory (MRL) UKHSA South West Bristol |
| T-SPOT | Oxford Diagnostic Laboratories Oxford |
| Rabies | Rabies Section Animal & Plant Health Agency (APHA) Weybridge Surrey |
| Arboviruses (includes West Nile, Dengue, Hantavirus, Flavivirus, Ross River Virus)  Leptospirosis  Lyme Confirmation CSF/Serum  Rickettsia  Viral Haemorrhagic Fever (Ebola, Marburg, CCHF) | Rare & Imported Pathogens Laboratory (RIPL) UKHSA Porton Down |
| Itraconazole  Rifampicin  Posaconazole  Teicoplanin | Regional Antimicrobial Reference Laboratory Bristol |
| Diphtheria | Respiratory & Vaccine Preventable Bacteria Reference Unit (RVPBRU) UKHSA Colindale |
| Cystercercosis IFAT  Leishmaniasis  Parasitic Infections  Schistosoma  Strongyloides | The Department of Clinical Parasitology The Hospital for Tropical Diseases London |
| Toxoplasma | Toxoplasma Reference Unit Public Health Wales Microbiology ABM Singleton Hospital Swansea |
| Psittacosis | UKHSA South West Bristol |
| EBV PCR  Adenovirus PCR (infant) | Virology Department Great Ormond Street Hospital London |
| Brucella | Virology Department Royal Liverpool and Broadgreen Hospital |
| Electron Microscopy  Hepatitis C resistance testing  HIV pro-viral DNA  Measles IgG/M  Mumps IgG/M  Polio  Rubella | Virus Reference Department (VRD) UKHSA Colindale |

Bacteriological tests referred to reference laboratories:-

| **ORGANISM / TEST** | **REFERENCE LABORATORY** |
| --- | --- |
| Acanthamoeba culture | UKHSA Malaria Reference Laboratory  London School of Hygiene & Tropical Medicine |
| Antibiotic susceptibility testing | UKHSA Antimicrobial Resistance and Healthcare Associated  Infections Unit (AMRHAI) Colindale |
| *Bacillus spp* (? Anthrax)  *Listeria monocytogenes*  *Helicobacter pylori* | UKHSA Gastrointestinal Bacteria Reference Unit (GBRU) Colindale |
| Fungal identification and susceptibility testing e.g. *Aspergillus fusaria* | Mycology Reference Laboratory (MRL) UKHSA Bristol |
| *Haemophilus influenza*  *S.pyogenes* Group A  *Streptococcus pneumoniae* | UKHSA Respiratory & Vaccine Preventable Bacteria ReferenceUnit (RVPBRU) Whitechapel, London |
| *Neisseria meningitidis* | Meningococcal reference unit (MRU) UKHSA Manchester |

Other reference laboratories that may be used:

Public Health Wales Microbiology

UKHSA Bacteria Reference Unit Department (BRD) Colindale

UKHSA Meningococcal Reference Unit (MRU) Manchester

UKHSA Mycology Reference Laboratory (MRL) Bristol

UKHSA National Mycobacterium Reference Service (NMRS-South) National Infection Service, Colindale

UKHSA Rare and Imported Pathogens Laboratory (RIPL) Porton Down

UKHSA The Central Sequencing Laboratory (CSL) Colindale

UKHSA Virus Reference Department (VRD) Colindale

# results

Enquiry terminals are available on all wards and in ED giving access to the Sunrise and DART OCM electronic results service and, in some cases, directly to the laboratory computer system. Hard copies of reports are only issued if appropriate. Most bacteriology culture reports are issued within 2-5 days depending on the investigation. Serology and Virology turnaround times depend on the frequency that assays are performed and the urgency of the request. Most assays are performed at least once a week.

GP surgeries subscribing to the local PMIP/DART OCM initiative should receive electronic copies of all reports on samples originating from their surgery. Hard copies of reports are only issued if appropriate.

# telephoned results

Results of urgent requests and positive results that may influence patient management are telephoned to the requesting medical team; this includes all positive blood cultures.

# reference laboratory results

Reports that contain results obtained from a third party reference laboratory will have the name of the referral laboratory clearly stated on the electronic and hard copies.

Please see section 20 for the names of the third party reference laboratories that may be used.

# reports

All results are electronically reported and the user can access them through the DART and PAS systems. Interim reports are issued for TB microscopy, Actinomyces investigation and where isolates are sent to reference laboratories for further identification. Final reports are issued when all the examinations are complete. Occasionally it may be necessary to amend a result and a further report clearly marked as amended will be issued; in these circumstances the user will be contacted by phone.

Reports contain the following information:

Patient details Name, sex, DOB and hospital number, address, consultant/GP

Laboratory number e.g. MR000111

Prefixed with M (for Microbiology) and another letter used to denote sample type e.g. R – respiratory sample, U – urine sample

Source Name of hospital or GP

Ward/surgery Name of ward or surgery

Sample type e.g. Mid-stream Urine, Blood Culture

Result section includes:

The type of test e.g. bacterial culture, microscopy & culture

Organism(s) isolated

Sensitivity results S = Sensitive

R = Resistant

I = Intermediate (can be successfully treated if high doses of antibiotics are used)

Certain samples may have suppressed antibiotic sensitivity tests e.g. catheter urines, urine samples with leucocytes <50/mm3 and leg ulcers. These can be available on request after consultation with a Medical Microbiologist.

Other information:

Authorised: The name of the person who authorised the report

Dates: Collected, received and reported

# interpretation of results

Where appropriate laboratory reports will have interpretive comments added. The comments may be automatic, rule-based comments added by the laboratory computer system or individual comments added by the Biomedical Scientists or Consultant Microbiologists.

Biological reference intervals, reference ranges, clinical decision values, where applicable and the uncertainty of measurement associated with the microbiological examinations performed are available from the laboratory on request.

# clinical tests available and turnaround times

Some specimens may be sent to reference laboratories for further confirmation / typing /grouping– see tables below:

| Bacteriology Samples | Tests | Turn-around time (working days) | Comments |
| --- | --- | --- | --- |
| Urine | Routine Microscopy & Culture  Negative urine | 1 day | Urgent Microscopy 1hr |
|  | Routine Microscopy & Culture  Positive urine | 3 days | Urgent Microscopy 1hr |
|  | Legionella and (pneumococcal antigen) | 1 day | Urgent antigen 1hr (pneumococcal antigen request ONLY after discussion with consultant microbiologist) |
|  | TB | 6 weeks |  |
| MRSA broth screening (nose + groin swabs) | Culture | 2-3 days | Negative screen |
|  |  | 4 days | Positive screen |
| *S.aureus* Renal screening | Culture | 2 days | Negative test |
|  |  | 2-3 days | Positive test |
| Ear swab | Culture | 2-3 days |  |
| Pus samples | Routine culture | 1-6 days |  |
|  | Actinomyces/fungi | 10 days |  |
| Nose swab | Culture | 2-3 days | generally used to check for staphylococcal carriage (including MRSA) and sometimes streptococcal and *C. diphtheria* |
| General wound swabs | Routine culture | 2-6 days | Urgent microscopy 2hrs |
|  | Anaerobic culture | 6 days |  |
| Throat swabs | Routine culture | 1-3 days | for Beta haemolytic Streptococci |
|  | Meningococcal | 1-3 days | On advice of microbiologist |
| Genito urinary swabs | Vaginal swab | 1-2 days | Useful for Gardnerella vaginalis, thrush and bacterial vulvovaginitis. Also Gp. B Strep. in pregnancy |
|  | Endocervical swab (female)  Urethral swabs (Male) | 2-3 days | Useful for Chlamydia and Gonococcus. A special swab and transport medium are needed for Chlamydia |
|  | Urethral swabs (Female) | 2-3 days | May also be examined for Chlamydia but are not recommended because collection of the SAMPLE is uncomfortable |
|  | Rectal swab | 2-3 days | May be examined for chlamydia and/or gonococcus. A special transport medium is needed for Chlamydia |
| Pernasal swab | for Bordetella pertussis | 5 days | A thin wire swab must be passed along the floor of the nasal passage until the resistance of the soft palate is met. |
| Sputum | Routine Microscopy & Culture | 2-3 days |  |
|  | AFB microscopy | 2-3 days | Referred test |
|  | AFB culture | 6 weeks | Referred test  Fast Track PCR available for TB and Rif. Resistance on serum samples |
|  | Bronchial washings (and induced sputum) |  | All samples undergo routine bacteriology. Virology, microscopy for PCP and TB examination are optional |
| Faeces/stools | C.difficile Toxin studies | 1 day | Repeat specimens from patients with previous positive toxin results should not be tested within 28 days of initial diagnosis, unless clinical details indicate a relapse of infection. |
|  | Culture | 3-5 days | Salmonella and E coli O157 |
|  | Helicobacter pylori (PCR) | 3-7 days | Not suitable for monitoring treatment. Indicates exposure to infection at some time  **Faecal sample only** |
|  | Rota/Adeno/Sapo virus | 1-3 days | Children <6 years old |
|  | Norovirus (SRSV) | 1-3 days | Only requested in outbreak situations |
|  | Microscopy (for ova & parasites) | 1-3 days | Available on request. Full travel history must be provided |
|  | Nucleic acid detection (PCR) | 3 days | Cryptosporidium / Giardia, Salmonella, Campylobacter, E.coli 0157, Vibrio and Yersinia. |
| Tissue and sterile fluids | Culture | 14 days |  |
| Blood culture | Aerobic & Anaerobic | 7 days |  |
| CSF | Microscopy & culture | 3 days | Microscopy 2hrs |
|  | DNA amplification screen (PCR) | 14 days | Also available on EDTA blood SAMPLE |
| Skin scrapings | Mycology microscopy and culture | 8 weeks |  |
| Any specimen type (preferably from sterile site) | 16s rRNA bacterial gene detection/sequencing | 5 days | Referred test |
| Any specimen type (preferably from sterile site) | Pan fungal (18s) DNA detection/sequencing | 5 days | Referred test |
| Faeces, urine and other specimen types | Parasite identification | 14 days | Referred test |

| Non-viral Serology | Tests | Turn-around time (working days) | Comments |
| --- | --- | --- | --- |
| Amoebic inf. | Amoebic IFAT | 10-15 days | Referred test |
| Antenatal screening | Syphilis | 8 days |  |
| ASOT | Antistreptolysin O | 7-10 days | Referred test |
| Brucellosis | Br.abortus & melitensis microagg | 14 days | Exposure history must be provided. Brucellosis is no longer endemic in the UK  Referred test |
| Candida | Candida antigen | 10 days | Discuss with microbiologist  Referred test |
| Cat scratch disease  (Bartonella henselae) | Antibodies | 3 months | Referred test – only available on agreement with a Consultant Microbiologist as no referral laboratory within UK. |
| Cryptococcal antigen |  | 2 days |  |
| Filariasis | Filaria Elisa | 7 days | Exposure history must be provided  Referred test |
| Hydatid | Hydatid ELISA | 10 days | clinical information essential  Referred test |
| Leptospirosis | Leptospira IgM/ELISA | 7 days | 24-48 hr result if urgent.  Referred test |
| Lyme disease | IgG  Western blot | 10 days | Exposure history must be provided  Referred test |
| Malaria | Malaria IFAT | 7 days | Blood film examination is preferred investigation. Antibody detection is rarely helpful. EDTA to Blood Sciences |
| Meningococcal meningitis | Meningococcal PCR | 3-7 days | Referred test |
|  | Meningococcal typing | 7-10 days | Referred test |
|  | Meningococcal antibodies | 28 days | Referred test |
| Mycobacteria | Interferon gamma release assay (QFT) | 3 days |  |
| Mycoplasma sp | Mycoplasma pneumoniae IgM | 7 days | Test ran twice each week |
| Mycoplasma genitalium | Resistance testing | 7 days | Test ran once weekly |
| Mycoplasma pneumoniae | PCR | 3 days | Referred test |
| Mycoplasma genus | PCR | 6 days | Referred test |
| Pertussis serology |  | 14 days | Referred test |
| Q fever | Q fever serology | 12 days | Referred test |
| Rickettsial or Arbovirus infection | Arbovirus and Rickettsial antibody | 7 days | travel history required  Referred test |
| Schistosomiasis | Schistosomal ELISA | 7-14 days | travel history required |
| Strongyloidiasis | Strongyloides ELISA | 7-14 days | travel history required + clinical details  Referred test |
| Toxocara |  | 7-14 days | Referred test |
| Toxoplasmosis | IgG, IgM,  Confirmation Dye test  PCR | 14 days | In the case of pregnancy contact, please contact Microbiology to arrange same day testing |
| Syphilis | Antibodies | 3 days if negative | In-house test for IgG/IgM combined. |
|  | RPR, TPPA and IgM | 7 days | In-house test for RPR and TPHA for confirmation and infection stage. IgM testing is a referred test |
|  | PCR | 7 days | Referred test |
| Vaccine screen | Haemophilus, pneumococcus, tetanus antibodies | 9 days | Referred test |
| Antimicrobial therapeutic drug monitoring |  | 5 days | Referred test |
| Fungal serology including Galactomannan and B-D-Glucan |  | 10 days | Referred test |

| Virology | Tests | Turn-around time (working days) | Comments |
| --- | --- | --- | --- |
| Antenatal Screening | Hepatitis B  HIV | 8 days |  |
| BK Virus | Molecular detection | 3-5 days | Referred test |
| Cytomegalovirus | IgG  IgM | 1-5 days | Screens for immunity or recent infection available. |
|  | Avidity load  Viral load | 3 days | CMV viral load (EDTA) can be estimated by PCR and is indicated for routine follow up of BMT patients and in selected immunosuppressed patients by arrangement with microbiologist |
| EB virus | EBV nuclear IgG, EBV capsid IgM | 1-5 days | Can determine a current, recent or historic infection |
|  | PCR | 3-5 days | Referred test |
| Enterovirus RNA | Coxsackievirus, Enterovirus, Echovirus | 3-5 days | Referred test |
| Herpes simplex  1 & 2 | Serology not routinely available. | 4 days | PCR is preferred diagnostic method. Can be sent in a VTM tube. |
| Herpes simplex  1 & 2 | Serology | 4 days | Referred test |
| Hepatitis A | IgM and total antibody (IgG/IgM) for immunity | 3 days | Indicates recent infection or immunisation |
| Hepatitis B | HBsAg | 3 days if negative | 4hrs if urgent +ve = Hepatitis B carrier |
|  | HBeAg | 3 days | high level carrier |
|  | anti-HBs | 3 days | Immunity to Hepatitis B (vaccine or past infection) |
|  | anti-HBe | 3 days | low level carrier (if present with HBsAg) |
|  | anti-HBc | 3 days | past or current Hepatitis B (not vaccine response) |
|  | anti-HBc IgM | 3 days | Suggests recent infection |
|  | Hep B viral load | 7 days | In-house test |
|  | HBV Genotype | 14 days | Referred test |
| Hepatitis C | Anti-HCV.  Hepatitis C viral load, Hepatitis C Ag/Ab | 3 days if negative | Initial screen for Hepatitis C antibody, if positive, viral load is performed from the same sample |
|  | Resistance testing | 18 days | Referred test |
|  | HCV Genotype | 5 days | Referred test |
| Hepatitis D | Serology | 17 days | Referred test |
| Hepatitis E | IgG  IgM | 3 days | Test run daily where possible |
|  | RNA | 3 days |  |
| HIV | 1+ 2 Ag/Ab Combo  HIV typing | 3 days if negative | If positive a repeat sample for confirmation of patient ID 2 x EDTA blood samples for viral load will be required |
|  | HIV viral load / serum confirmatory | 3 days | If positive a repeat sample for confirmation of patient ID 2 x EDTA blood samples for viral load will be required |
| HTLV | 1 & 2 antibodies | 3 days |  |
| Human Herpes Virus (HHV) 6/7 | In vesicles / skin lesions (swabs in VTM) | 7 days | Referred test |
| Human Herpes Virus (HHV) 6/7/8 | DNA in blood | 3-5 days | Referred test |
| Influenza | Typing | 6 days (season dependent) | Referred test |
| Measles | IgG | 3 days | IgG performed in-house |
|  | IgM | 10 days | IgM is a referred test, please provide contact symptoms and treatment information if requesting IgM testing |
| Mumps | IgG | 3 days | IgG performed in-house |
|  | IgM | 10-15 days | IgM is a referred test, please provide contact symptoms and treatment information if requesting IgM testing |
| Rubella | IgG and IgM | 3 days |  |
|  |  |  |  |
| Parvovirus | IgG | 3 days | Test run weekly, more frequent if urgent samples |
|  | IgM | 3 days |  |
|  | PCR | 3-5 days | PCR is carried out on IgM positive samples if clinically required – Referred test |
| Varicella Zoster | IgG | 3 days (same day possible if urgent) | In cases of pregnancy contact, please contact Microbiology to arrange same day testing (needed for administering VZIG). |

| Virus investigations | Tests | Turn-around time (working days) | Comments |
| --- | --- | --- | --- |
| Throat swab in VTM for PCR | Adenovirus | 1-3 days | Virus tested as routine |
|  | Bocavirus | 1-3 days | Virus tested as routine |
|  | Corona viruses | 1-3 days | Virus tested as routine |
|  | Enterovirus | 1-3 days | Virus tested as routine |
|  | Human Metapneumo virus | 1-3 days | Virus tested as routine |
|  | Influenzavirus A & B | 1-3 days | Virus tested as routine |
|  | Parainfluenza 1-4 | 1-3 days | Virus tested as routine |
|  | Rhinovirus | 1-3 days | Virus tested as routine |
|  | RSV A & B | 1-3 days | Virus tested as routine |
| Nasopharyngeal aspirate (NPA) | Adenovirus | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Bocavirus | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Corona viruses | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Enterovirus | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Human Metapneumo virus | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Influenzavirus A & B | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Parainfluenza 1-4 | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Rhinovirus | 1-3 days | Virus tested as routine on RSV negative samples or by special request |
|  | Respiratory syncytial virus (RSV) in babies | 24 hours | PCR |
| CSF | Enterovirus | 3 days | In-house test |
|  | Parechovirus | 3 days | In-house test |
|  | VZ virus | 3 days | In-house test |
|  | Herpes simplex 1 and 2 PCR | 3 days | In-house test |
|  | Adenovirus PCR | 3 days | In-house test |
|  | CMV PCR | 3 days | Referred test |
|  | EBV PCR | 3 days | Referred test |
| Vesicles / skin lesions  (swabs in VTM) | Herpes simplex | 2-4 days | Swabs in VTM taken from ulcerated genital and non-genital lesions give good results. Dry lesions need to be scraped to expose the base of ulcer. |
|  | Enterovirus | 3 days | Referred test |
|  | Varicella zoster | 5 days | Virus is in cells lining vesicle rather than vesicle fluid. |
| CMV PCR (Urine & Saliva) | CMV | 3 days | In-house test  Urine - In plain white top universal **not** boric acid  Saliva – Dry Swab or VTM |

For Chlamydia trachomatis, Neisseria gonorrhoea, Mycoplasma genitalium & Trichomonas vaginalis investigation, **please use Abbott Alinity M multi-collect specimen collection kits only**

For MGEN & TV – please use urine and genital swabs only

| CT, GC, MGEN & TV PCR | Tests | Turn-around time (working days) | Comments |
| --- | --- | --- | --- |
| Urine | PCR | 1-5 days | Decant into urine collection tube in the Abbott Alinity M multi-collection specimen collection kit as soon as the sample is taken. If no Alinity M urine collection tubes are available, please ensure that the sample reaches Microbiology the same day the sample was taken. |
| Urethral swab |  |  | Abbott Alinity M multi-collect specimen collection kit |
| Cervical / vaginal swab |  |  | Abbott Alinity M multi-collect specimen collection kit |
| Throat |  |  | Abbott Alinity M multi-collect specimen collection kit |
| Rectal |  |  | Abbott Alinity M multi-collect specimen collection kit |
| Other sites including eye |  |  | Samples from these other sites are tested, but please note that they have not been validated on the Alinity M instrument used to process samples. |

If any test is urgent, please contact Microbiology.

If a test is required that is not listed above, please contact Microbiology and we will endeavour to identify a laboratory that is able to perform the examination.

# REferences

Public Health England (2017) Investigations of Specimens other than Blood for Parasites UK Standards for Microbiology Investigations B31 Issue 5.1

[B 31 - Investigation of specimens other than blood for parasites (rcpath.org)](https://www.rcpath.org/static/35769984-d03b-4727-abaaf8c547ae519c/UK-SMI-B-31i51-June-2017-Investigation-of-specimens-other-than-blood-for-parasites.pdf)

Public Health England (2017) Investigations of Cerebrospinal Fluid, UK Standards for Microbiology Investigations B27 Issue 6.1

[B 27 - Investigation of Cerebrospinal Fluid (rcpath.org)](https://www.rcpath.org/static/34f0577c-2ff0-4ed6-90a3848e7511d634/UK-SMI-B-27i61-May-2017-Investigation-of-cerebrospinal-fluid.pdf)

Public Health England (2016) Investigation of pus and exudates, UK Standards for Microbiology Investigations B14 Issue 6.2

[B 14 - Investigation of pus and exudates (rcpath.org)](https://www.rcpath.org/static/e2148c4a-af12-4864-8e2918759c69d35b/UK-SMI-B-14i62-Investigation-of-pus-and-exudates-November-2016.pdf)

Public Health England (2019) Investigation of urine, UK Standards for Microbiology Investigations B41 Issue 8.7

[B 41 - Investigation of urine (rcpath.org)](https://www.rcpath.org/static/de4a6639-b118-46ea-9d3b8a0be4014944/UK-SMI-B-41i87-January-2019-Investigation-of-urine.pdf)

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[U 1 - National user manual template (rcpath.org)](https://www.rcpath.org/static/56879556-0b0a-4cd9-9ed7cae2859735ce/uk-smi-u-1i1-national-user-manual-template-october-2016-pdf.pdf)

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[Pernasal swab for whooping cough ( Bordetella Pertussis ) (gloshospitals.nhs.uk)](https://www.gloshospitals.nhs.uk/our-services/services-we-offer/pathology/tests-and-investigations/pernasal-swab-whooping-cough-pertussis/)

Howard, M (2010) *Medical Microbiology*, Oxford University Press pages 9-10

Public Health England (2020), Investigations of Gastroenteritis, UK Standards for Microbiology Investigations S7, Issue 2

[S 7 - Gastroenteritis (rcpath.org)](https://www.rcpath.org/static/fbfe4d59-a044-4285-8cd3782922f3971a/fe32b496-b264-4296-9805f4f52dbef55f/uk-smi-s-7i2-gastroenteritis-october-2020-pdf.pdf)

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[DPL Handbook (lshtm.ac.uk)](https://www.lshtm.ac.uk/files/dpl-handbook.pdf)

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[Consent and confidentiality | (hcpc-uk.org)](https://www.hcpc-uk.org/standards/meeting-our-standards/confidentiality/guidance-on-confidentiality/consent-and-confidentiality/)

Public Health England (2018) Investigation of tissues and biopsies from deep-seated sites and organs, UK Standards for Microbiology Investigations B17 Issue 6.3

[B 17 - Investigation of tissues and biopsies from deep seated sites and organs](https://www.rcpath.org/static/47722f8c-8c43-4ad2-be5785b454021795/UK-SMI-B-17i63-Investigation-of-tissues-and-biopsies-from-deep-seated-sites-and-organs-January-2018.pdf)

Public Health England (2018) Investigation of Fluids from Normally Sterile Sites, UK Standards for Microbiology Investigations B26 Issue 6.2

[B 26 - Investigation of fluids from normally sterile sites](https://www.rcpath.org/static/88930ee2-cb77-4a12-91b1d6aadacd93c9/UK-SMI-B-26i62-Investigation-of-fluids-from-normally-sterile-sites-October-2018.pdf)

# appendices

## Appendix 1 – Collection of urine samples

1. Mid-stream urine

Refer to [The Collection of a random urine sample (ekhuft.nhs.uk)](https://leaflets.ekhuft.nhs.uk/the-collection-of-a-random-urine-sample/)

1. Urine collection pads:

Remove the infant’s nappy and clean the perineum or prepuce with soap and water. Do not apply any creams.

Place the urine collection pad across vulva or penis in a lengthwise fashion.

Remove the adhesive backing from the pad and secure to the nappy.

Change urine collection pad every 30-45 minutes and also when the child has passed a stool, to reduce the risk of contamination with skin or faecal flora.

When the infant has passed urine, remove the nappy with the urine collection pad in it.

Lay the pad down wet side up on an appropriate clean surface.

Take sterile 5 ml syringe and place the tip on the pad and extract the urine by pulling up the plunger and empty the syringe into an appropriate clean CE marked container.

Repeat this process until the required amount of urine has been obtained.

1. Urine collection bags:

Select the correct size sterile urine bag to avoid leakage or contamination with faeces.

Remove the infant’s nappy and clean the perineum or prepuce with soap and water, dry the area thoroughly and do not apply any creams.

Remove the protective backing from the bag, and follow the following procedure:

For females, place the bag over the vulva, starting from the perineum and work upwards, pressing the adhesive to perineum and symphysis.

For male, insert penis and scrotum into the opening of the bag and press adhesive to perineum and symphysis.

Cut a hole in the nappy and pull the urine bag through the opening.

When the infant has passed urine, perform the hand hygiene procedures before putting on disposable gloves and removing the bag.

Maintain a good aseptic technique whilst holding the bag over a suitable clean urine specimen container and cut off the tip of the bottom corner of the bag using clean scissors and empty the urine into an appropriate clean CE marked container.

Wash the infant’s genitalia after the procedure to prevent soreness of the skin.

1. Suprapubic aspirate

Collection of urine by a supra-pubic aspirate should be considered when a sterile sample is required. Ultrasound guidance should be used to indicate the presence of urine in the bladder before a suprapubic aspirate is attempted (NICE, 2007).

1. Catheter specimen

Catheter samples must be collected from the self-sealing valve of the urinary drainage tubing. Do not disconnect the closed drainage system as infection may be introduced (DH 2001). Similarly do not take the sample from the urinary drainage bag as the specimen may be contaminated.

Using an aseptic non-touch technique, clean the catheter sampling site with 2% chlorhexidine/70% alcohol wipe (e.g. Clinell®) and allow to dry.

Use a sterile syringe and needle to access the self-sealing valve, insert the needle into the valve at an angle of 45 degrees; this will minimise penetration of the wall of the tubing and subsequent needle stick injury.

Gently pull the syringe plunger out to transfer the urine into the syringe.

Remove the needle and syringe, wipe the area with the alcohol swab and allow to dry (the valve will self-seal) and transfer the urine into an appropriate clean CE marked container.

Discard the needle and syringe into a sharps container.

## Appendix 2 – Collection of a faecal sample

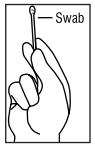
Refer to [The Collection of a stool (faeces) sample (ekhuft.nhs.uk)](https://leaflets.ekhuft.nhs.uk/the-collection-of-a-stool-faeces-sample/)

## Appendix 3 - Abbott Alinity M Endocervical Swab Specimen Collection Guide

CAUTION: Do NOT expose swab to Transport Buffer prior to collection.

1. Discard disposable transfer pipette; it is not required for endocervical swab specimen collection.

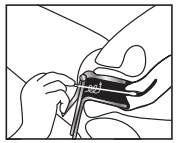
2. Remove the sterile swab from the wrapper, taking care not to touch swab tip or lay it down on any surface.

****

3. Insert the white tip of the specimen collection swab into the endocervix canal.

4. Gently rotate the swab for 15 to 30 seconds to ensure adequate sampling.

5. Withdraw the swab carefully.

****

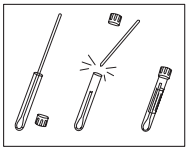
6. Handle the cap and tube carefully to avoid contamination, including the outside of the transport tube and cap. If necessary, change gloves.

7. Unscrew the transport tube cap and immediately place the specimen collection swab into the transport tube so that the white tip is down.

8. Carefully break the swab at the scored line on the shaft; use care to avoid splashing of contents.

9. Recap the transport tube carefully. Ensure the cap seals tightly or leakage may occur.

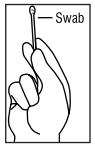
10. Label the transport tube with sample identification information, including date of collection using an adhesive label.

****

## Appendix 4 – Abbott Alinity M Clinician-Collected Vaginal Swab Specimen Collection Guide

CAUTION: Do NOT expose swab to Transport Buffer prior to collection.

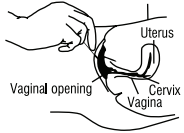
1. Discard disposable transfer pipette; it is not required for vaginal swab specimen collection. 2. Remove the sterile swab from the wrapper, taking care not to touch swab tip or lay it down on any surface.

****

3. Insert the white tip of the specimen collection swab about two inches (5 cm) into the opening of the vagina without touching the skin or labia external to the vagina.

4. Gently rotate the swab for 15 to 30 seconds against the sides of the vagina.

5. Withdraw the swab carefully. Do not touch the tip of the swab to area outside the vagina.



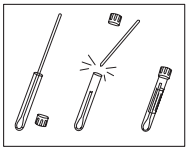
6. Handle the cap and tube carefully to avoid contamination, including the outside of the transport tube and cap. If necessary, change gloves.

7. Unscrew the transport tube cap and immediately place the specimen collection swab into the transport tube so that the white tip is down.

8. Carefully break the swab at the scored line on the shaft; use care to avoid splashing of contents.

9. Recap the transport tube carefully. Ensure the cap seals tightly or leakage may occur.

10. Label the transport tube with sample identification information, including date of collection using an adhesive label.

****

## Appendix 5 - Abbott Alinity M Self-Collected Vaginal Swab Specimen Collection Guide

Your kit contains the following:

• One Transport Tube containing a liquid

• One Sterile Specimen Collection Swab

• One disposable Transfer Pipette

CAUTION: DO NOT touch the white tip of the swab or lay the swab down. If the white tip is touched or the swab is laid down or dropped, your results may not be accurate. You need to request a new Alinity m multi-Collect Specimen Collection Kit.

DO NOT pre-wet the collection swab with the liquid in the Transport Tube before collecting a sample.

DO NOT ingest or expose skin/eyes to the liquid in the Transport Tube.

IF exposed or concerned: Get medical advice/attention.

**Pre-Collection Steps**

1. Wash your hands with soap and water thoroughly before starting and after completing all steps.

2. In the privacy of the examination room or restroom, you will need to undress from the waist down. You will need to position yourself to maintain balance during the collection procedure.

3. Open the kit package. Discard the transfer pipette (Shown in Diagram 1). Do not open the transport tube. Set the tube aside on a clean, dry surface before beginning collection.

**Collection Steps**

4. Remove the swab from the wrapper with your clean hands. Hold the swab with the white tip up (Shown in Diagram 2). Do not touch the tip of the swab to anything.

5. Holding the swab with one hand, gently spread the vaginal labia with your other hand. Insert the white tip of the swab about two inches (5 cm) into the opening of your vagina (Shown in Diagram 3). Rotate the swab for 15 to 30 seconds. Make sure the swab touches the sides of your vagina. Remove the swab from your vagina being careful not to touch your skin. Do not set the swab down.

6. While still holding the swab, unscrew and remove the cap from the transport tube without setting the cap down. Place the swab into the tube with the white tip down (Shown in Diagram 4 and 5). If the transport tube spills or liquid splashes out, you will need to request a new Specimen Collection Kit.

7. Break off the top of the swab along the score line. (The score line is made to break easily). Try not to spill or splash any of the liquid out of the transport tube. Screw the cap back onto the transport tube tightly (Shown in Diagram 6). The cap must be tight or leakage may occur.

8. Return the transport tube containing the swab to the nurse or doctor.



## Appendix 6 – Abbott Alinity M Urine Specimen Collection Guide (males and females)

1. The patient should not have urinated for at least one hour prior to sample collection.

2. Discard specimen collection swab; it is not required for urine specimen collection.



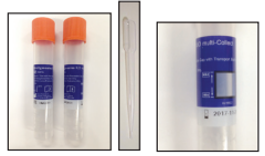
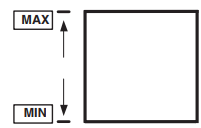
3. Using a urine specimen collection cup, the patient should collect the first 20 to 30 mL of voided urine (the first part of the stream).



4. Unscrew the transport tube cap, taking care not to spill the transport buffer within.

5. Handle the cap and tube carefully to avoid contamination, including the outside of the transport tube and cap. If necessary, change gloves.

6. Use the plastic transfer pipette to transfer urine from the collection cup into the transport tube until the liquid level in the tube falls within the clear fill window of the transport tube label or else a new specimen should be collected. Do not overfill.

7. Recap the transport tube carefully. Ensure the cap seals tightly or leakage may occur.

8. Label the transport tube with sample identification information, including date of collection using an adhesive label. Take care not to obscure the fill window on the transport tube.

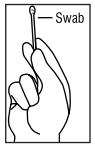
9. Decontaminate and dispose of all specimens, reagents, and other potentially contaminated materials in accordance with local, state, and federal regulations.

## Appendix 7 – Abbott Alinity M Oropharyngeal Swab Specimen Collection Guide

CAUTION: Do NOT expose swab to Transport Buffer prior to collection.

1. Discard disposable transfer pipette; it is not required for oropharyngeal swab collection.

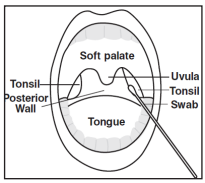
2. Remove the sterile swab from the wrapper, taking care not to touch swab tip or lay it down on any surface.

****

3. The white tip of the swab should be rubbed against each tonsillar pillar and the posterior wall and rotated at least one time. Do NOT collect specimens from the tongue.

4. If pharyngeal exudate is present, this should also be sampled.

5. Withdraw the swab carefully.



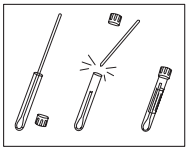
6. Handle the cap and tube carefully to avoid contamination, including the outside of the transport tube and cap. If necessary, change gloves.

7. Unscrew the transport tube cap and immediately place the specimen collection swab into the transport tube so that the white tip is down.

8. Carefully break the swab at the scored line on the shaft; use care to avoid splashing of contents.

9. Recap the transport tube carefully. Ensure the cap seals tightly or leakage may occur.

10. Label the transport tube with sample identification information, including date of collection using an adhesive label.

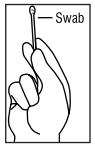
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## Appendix 8 – Abbott Alinity M Rectal Swab Specimen Collection Guide

CAUTION: Do NOT expose swab to Transport Buffer prior to collection.

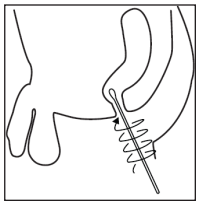
1. Discard disposable transfer pipette; it is not required for rectal swab collection.

2. Remove the sterile swab from the wrapper, taking care not to touch swab tip or lay it down on any surface.

****

3. The white tip of the swab should be inserted 1-2.5 cm into the anal canal and rotated at least one time.

4. Withdraw the swab carefully



5. Handle the cap and tube carefully to avoid contamination, including the outside of the transport tube and cap. If necessary, change gloves.

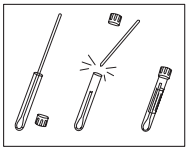
6. Unscrew the transport tube cap and immediately place the specimen collection swab into the transport tube so that the white tip is down.

7. Carefully break the swab at the scored line on the shaft; use care to avoid splashing of contents.

8. Recap the transport tube carefully. Ensure the cap seals tightly or leakage may occur.

9. Specimens discoloured by stool or blood are acceptable.

10. Label the transport tube with sample identification information, including date of collection using an adhesive label.

****

## Appendix 9 – Collecting a Blood Culture

**See EKHUFT Policy Centre for Blood Culture Collection Operating Standard.**

Please ensure correct volume collected:

**Adults** 10mls in each bottle (1 x Aerobic (green) & 1 x Anaerobic Bact/Alert (purple))



**Paediatric** - Please refer to EKHUFT Policy Centre for Blood Culture Collection Operating Standard for volume required (4mls maximum)



Volumes greater than or less than these recommended volumes DO NOT maintain the optimal blood to medium ratio and may affect the recovery of organisms. Monitor until correct volume is achieved – there is NO automatic shut off to prevent overfilling

**5 Important Points for the Prevention of Contaminated Blood Cultures (MIC-WI-196 rev. 6)**

1. Effective hand hygiene using soap and water and/or alcohol hand rub prior to:

* Undertaking the procedure
* After preparing the skin (before taking the blood culture)
* After removing gloves on completion of the procedure

1. 2% chlorhexidine in 70% alcohol (e.g. Chloraprep Frepp) **must** be used for skin disinfection and allowed to dry before undertaking the procedure
2. Tops of the blood culture bottles **must** be disinfected using 2% chlorhexidine in 70% alcohol (e.g. Clinell wipe)
3. A fresh stab must be made in patients with suspected bacteraemia. Avoid femoral vein puncture. Blood cultures **MUST NOT** be taken from old peripheral lines, but may be taken at the insertion of a line if the patient is difficult to bleed.
4. If blood is being collected for other tests, always **collect** **the blood culture first**.

Adults = 10ml blood for each bottle (1x green top, 1x purple top)

**DO NOT EXCEED 10 mL**

Paediatrics = 4ml (maximum) blood in a single yellow top bottle

NB: Blood culture collection MUST ONLY be undertaken by a Doctor, Registered Nurse or Phlebotomist who has successfully completed the EKHUFT blood culture collection E-learning package (mandatory training)

## Appendix 10 Phlebotomy Mycobacteria Blood Culture Collection Instructions

Patients will arrive in the Phlebotomy clinic with BD BACTEC™ Myco/F Lytic culture bottles supplied by the General Practice (Microbiology will provide the bottles on request to the GP’s). Patients will also have a microbiology form requesting “Mycobacterium chimaera culture”.

Three mycobacteria blood culture bottles must be inoculated on separate days and submitted to the Microbiology Department.

Pre-collection

* DO NOT USE any vial showing evidence of contamination, leakage or damage.
* Prior to use, each **glass** vial must be checked for damage. On rare occasions, the glass bottle neck may be cracked and the neck may break during removal of the flip-off cap or in handling.
* Each vial should be examined for evidence of contamination such as cloudiness, bulging or depressed septum, or leakage. If a contaminated vial is used for direct draw, gas or contaminated culture media could be refluxed into the patient’s vein.

Collection

* The specimen must be collected using sterile technique to reduce the chance of contamination.
* The range of blood volume which can be cultured is 1 ml to 5 ml, with optimum recovery obtained at **3 ml to 5 ml.**
* Prior to inoculation, the medium fill volume should be noted on the label with a pen or marker to indicate the starting point of specimen collection. The vacuum in the bottle will usually exceed 5 mL, so the user should monitor the volume collected by means of the 5 mL graduation marks on the vial label.
* The bottle should be transported as quickly as possible to the laboratory and, if available, using the pneumatic air specimen transport system (POD).

## Appendix 11 – Collecting Swab Samples (Howard, 2010)

A good aseptic technique must be used during the swab sample collection.

Ensure the specimen is labelled correctly as stated in section 10

1. Wound swabs

Remove the swab from the packaging and gently rotate the swab tip in the area where as much material as possible may be collected, collect fresh pus if present. To prevent contaminants affecting the validity of the result please remove any old pus that might be present before taking the sample. Replace the swab into the swab transport tube.

1. Nasal swabs

Remove the swab from the packaging and moistened the swab in sterile saline solution. Ask the patient to tilt their heads backwards as this will make it easier to collect the sample. Insert the swab into the nose, taking care not to touch the outer sides of the nose and gently rotate the swab against the inner surfaces of the nose and any lesions that may be present. Withdraw the swab and replace in the swab transport tube.

1. Throat swabs

Place the patient facing the light; depress the patient’s tongue with a spatula to prevent contamination from the tongue in the event of the patient gagging. Remove the swab from the packaging and insert into the patient’s mouth taking care not to touch the swab against the sides of the mouth. Gently rub the swab over any areas at the back of the throat that are inflamed including any lesions that may be present. Withdraw the swab and replace it in the swab transport tube.

1. Eye swabs

Ask the patient to try not to flinch during the procedure. Ask them to look up. Remove the swab from the packaging and moisten with sterile saline solution. Wearing sterile disposable gloves expose the pink conjunctiva by pulling gently on the eyelid. Collect the sample by gently rubbing the swab across the lower eyelid starting at the inner corner and working outwards. Replace the swab in the swab transport tube.

1. Ear swabs

Remove the swab from the packaging and place the swab in the outer ear (taking care not to push the swab in too far and damaging the ear drum). Gently rotate the swab to collect any pus or discharge. Replace the swab in the swab transport tube.

1. Pernasal swab

Seat the patient, looking upwards with the neck fully extended. Insert the pernasal swab through a nostril and advance along the floor of the nose until it reaches the nasopharynx. It has been suggested that the swab be held against the posterior nasopharynx for up to 30 seconds or until the patient coughs. In practice, it is more likely that a patient will only be able to tolerate this for a few seconds.



Remove the swab and plunge into transport tube.

## Appendix 12 - Acanthamoeba sample collection instructions

(LSHTM Handbook March 2022) [Diagnostic Laboratory Parasitology Laboratory User Handbook (lshtm.ac.uk)](https://www.lshtm.ac.uk/files/dpl-handbook.pdf)

Suitable sample types & sample preparation:

All specimens should be submitted for testing together with a completed Acanthamoeba referral form available to download and print from: [www.parasite-referencelab.co.uk](http://www.parasite-referencelab.co.uk))

Please ensure all containers are tightly screwed and use Parafilm (NOT Sellotape) to prevent leakage during transit.

• **Clinical samples** (corneal scrapes, biopsies, fluids, swabs etc.) should be sent in a small volume (1 – 2 millilitre ideal) of sterile saline or sterile distilled water in a small (less than 5 millilitre) sterile vial or tube.

Material from a **needle or blade scrape** should be rinsed into the saline or water. Please remove blades or needles after rinsing. Do NOT leave the blade in the tube as it rusts: this inhibits our PCR and may have a detrimental effect on culture isolation.

**Swabs or washings** appear to be less efficient in detecting the organism. If swabs must be sent then please add a small volume of sterile saline or sterile distilled water to the swab to prevent drying. Please do not send dry swabs.

Punch **biopsies** or portions of excised cornea may also be submitted: put sample into a small volume of sterile saline or distilled water in a small sterile vial.

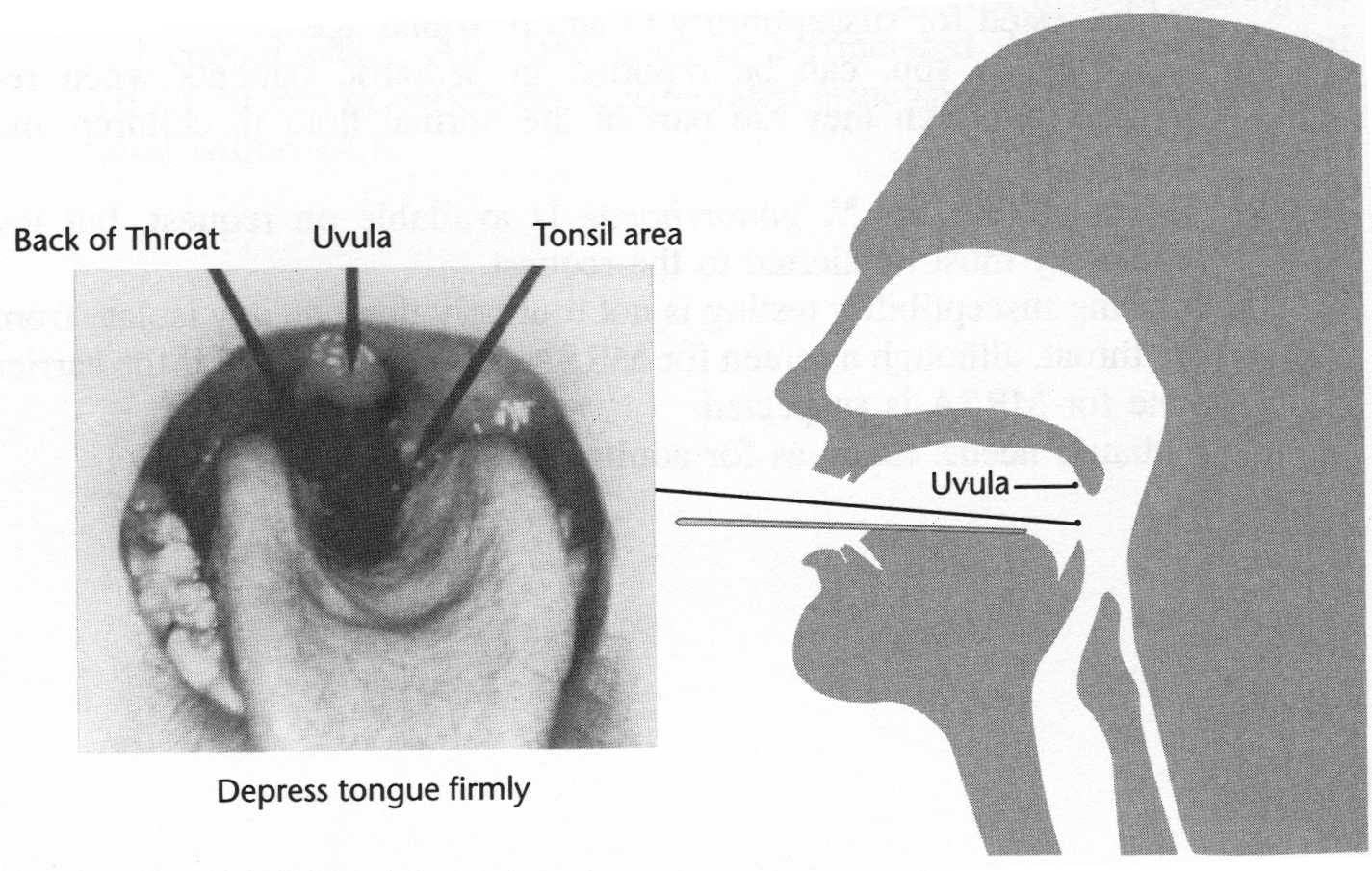
• **Non-clinical samples** - contact lenses: should be sent in their lens cases (i.e. in used contact lens fluid.) Please note: we do not test commercial contact lens solutions (other than that already in patients’ contact lens cases). Culture is performed on lenses and fluids; PCR is performed on fluids only. N.B. Isolation of Acanthamoeba from contact lens-related specimens, whilst suggestive, does not necessarily implicate the amoeba as causing the patient’s symptoms. Amoebic genera (other than Acanthamoeba), flagellates, ciliates and other organisms may be found in contaminated washing fluids and on lenses, particularly with poor lens hygiene.

• **Culture-positive samples**: please send original culture plate if possible, or blocks of agar from the plate in a sterile vial.

## Appendix 13 – Collection of influenza and COVID swabs

1. **Influenza:**

Cases of suspected influenza (acute influenza-like illness or acute bronchitis within five days of onset of illness) are asked for a combined nose & throat swab specimen. A good specimen for the detection of influenza must contain a substantial number of respiratory epithelial cells, which are mainly obtained from the nasal swab. A throat swab alone will contain mainly squamous epithelial cells in which influenza does not replicate.

1. A single swab with cotton wool bud is inserted in one nostril and rubbed against and above the nasal turbinate.
2. A second swab is used to abrade the tonsils and pharynx 

* Both swabs are broken off into a **single** vial of viral transport medium (VTM)
* Replace the lid of VTM vial and screw up firmly
* Label the vial with patient’s name and date of birth

Please use the test request form supplied with each vial. Please fill in a separate form for each patient and please include:

1. Patient name
2. Date of birth
3. Sex
4. Whether the patient has influenza-like illness or acute bronchitis
5. Date of symptom onset
6. Influenza vaccine status, whether patient falls into Department of Health ‘Green Book’ Risk Group and details about any anti-influenza drugs (currently or in previous 14 days)
7. Date of swab

If you are not able to courier samples back immediately then they should be kept in a fridge at +4 °C and sent the next day.

Unused collection bag with swabs, request form and virus transport medium bottle should be stored at room temperature.

Any further information or help can be obtained by phoning your **Microbiology WHH** on 01233 616760

1. **COVID**

Routine sample should be collected preferably using Sigma MM transwabs (white cap, green swab) or alternatively using VTM.

1. **Prepare to test:**

Clean and dry a surface before unpacking the contents from the test kit

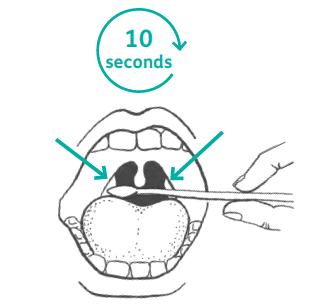
Ask the patient to blow their nose with a tissue before taking the test, throwing the tissue in a bin.

Before taking the test, wash your hands with soap and water

1. **Test Procedure**

Open the swab package and remove the swab holding it at the stick end – do not let the swab touch other surfaces.

Ask the patient to open their mouth and gently rub the swab over their tonsils for 10 seconds

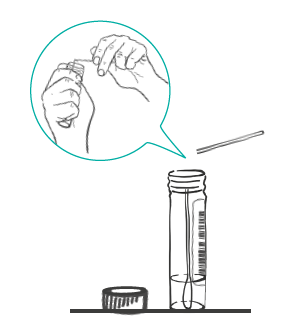


Then place the same swab inside the patient’s nostril and gently push upwards, then gently turn the swab for 10-15 seconds



**If unable to swab the patient’s throat, please swab both nostrils.**

Remove the lid of the sample tube containing the transport fluid (lay the lid on the work surface – open end facing up). Place the swab fully into the fluid before gently pulling/pushing the swab handle against the sample container opening until the swab breaks at the swab breakpoint (this ensures that the swab is the correct height to fit in the sample container) – (DO NOT force/bend the swab into the specimen container as this causes the container to leak) so that the swab fits into the sample tube without bending.



Replace the lid of the sample tube and ensure there are no leaks.

1. **Packaging samples**

All specimens must be singularly – DO NOT place multiple samples in one bag.

## Appendix 14 – Collection of Sputum

* The patient should be encouraged to drink plenty of fluid the evening before the sputum collection
* Do not allow the patient to clean their teeth or use mouthwash before collecting the specimen as this may kill any bacteria present.
* Collect the specimen where possible before the patient eats or drinks (particularly important if TB is suspected.
* Instruct the patient that true sputum is required and not saliva e.g. collected after a deep cough or following physiotherapy.
* Ask the patient to take some deep breaths (or use a nebuliser) to loosen the secretions before coughing hard to bring the sputum into the mouth.
* The patient should then spit the sputum into the sample collection container
* Transfer the labelled specimen with the sample request form as soon as possible after collection.

## Appendix 15 - Orthopaedic Theatre Specimens Request Form





## Appendix 16 – supplies order form



