

# CHESTER AND WIRRAL MICROBIOLOGY SERVICE USER GUIDE

|                           |                   |
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## REVISION HISTORY

| REVISION | CHANGE DETAILS   | SECTION(S) | PAGE(S) | MADE BY | APPROVED BY | ACTIVE DATE |
|----------|--|------------|---------|---------|-------------|-------------|
| 4.1      | Changed Aptima swabs to Roche throughout.  |            |         | BDB     | BDB         | 24.07.23    |
| 4.1      | Covid details updated  | 10         | 28      | BDB     | BDB         | 24.07.23    |
| 4.1      | New IUCD information added   | 10         | 35      | BDB     | BDB         | 24.07.23    |
| 4.1      | <i>C.difficile</i> interpretation updated  | 13         | 48      | BDB     | BDB         | 24.07.23    |
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| 4.2      | Specimen transport added   | Throughout |         | BDB     | BDB         | 27.03.24    |
| 4.2      | Added micro duty senior email address  | 2          | 6       | BDB     | BDB         | 27.03.24    |
| 4.2      | Removal of in-house CT/NG testing/exelicare requesting   | Throughout |         | BDB     | BDB         | 28.03.24    |
| 4.2      | Added collection and storage conditions  | Throughout |         | BDB     | BDB         | 28.03.24    |
| 4.2      | Verify patient ID  | 7          | 14      | BDB     | BDB         | 28.03.24    |
| 4.2      | Number CSF samples   | 7.11       | 20      | BDB     | BDB         | 28.03.24    |
| 4.2      | Users can request UoM reports  | 11         | 47      | BDB     | BDB         | 28.03.24    |
| 4.2      | Changes to examination procedures which could affect interpretation will be notified to users. | 11         | 47      | BDB     | BDB         | 28.03.24    |
| 4.2      | Delayed results  | 3          | 7       | BDB     | BDB         | 28.03.24    |
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| 4.4      | Added $\beta$ -D-Glucan  | 6.14       | 49      | BJE     | BDB         | 30.12.24    |
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|          | Multiple specimens   | 5          | 13      | BJE     | BDB         | 12.02.25    |
| 4.6      | Changed PHE to UKHSA   | 10         | 42-43   | BJE     | BDB         | 22.04.25    |
| 5        | Full Review Added guidance on the labelling of high-risk samples                               | 6.1        | 10      | BJE     | BDB         | 14.07.25    |
|          | Added Gonorrhoea testing to eye swab. Removed VTM swab as now use Roche Uni swabs              | 10         | 29      | BJE     | BDB         | 14.07.25    |
| 5.1      | Added section on Consent   | 3.6        | 8       | BJE     | BDB         | 27.07.25    |
|          | Updated multiple samples   | 6.1        | 11      | BJE     | BDB         |             |

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## 1.0 GENERAL INFORMATION

The Medical Microbiology Service is part of collaboration between Wirral University Teaching Hospital NHS Foundation Trust and Countess of Chester NHS Foundation Trust. The main Microbiology Laboratory is located on the Croft Business Park, Bromborough.

The **Microbiology Laboratory** address is:

Chester and Wirral Microbiology Service  
11 Bassendale Road,  
Bromborough,  
Wirral.  
CH62 3QL

Tel 01244 362500

**24-hour cover is provided for ALL aspects of infectious diseases.**

### On-call service for urgent clinical advice

Consultant in Medical Microbiology

Between 17-00 – 09-00 Monday – Friday

Between 17-00 on Friday and 09-00 on Monday (weekends), please contact the Consultant.

Microbiologist on-call via the switchboard of each Trust. Please note that the on-call Consultant Microbiologist should only be contacted in response to requests from the following categories of personnel:

**Consultants** (in exceptional circumstances contact may be made by a registrar **if** the patient has been discussed with the consultant in the team, but the consultant is unable to make the call himself/herself).

### Pharmacists

### Laboratory staff

### General Practitioners

### Countess of Chester Hospital

There is a 24-hour onsite service providing Molecular testing for SARS-CoV-2 and Flu in Pathology at Countess of Chester Hospital

Blood Sciences Laboratory  
Department of Pathology  
CoCH  
Tel: 01244 365595

## 2.0 KEY CONTACTS AND THEIR TELEPHONE NUMBERS/ EXTENSIONS

### Laboratory Results/Enquiries

**Monday – Sunday 09-00hrs, - 19-00hrs 01244 362500**

#### Consultant Staff (Wirral)

|                   |                        |
|-------------------|------------------------|
| Dr Kavya Mohandas | 0151 482 7694          |
| Dr David Harvey   | 0151 604 7466          |
| Dr A Abdelrahman  | 0151 678 5111 Ext 7436 |

#### Clinical Scientist (Wirral)

|                      |                        |
|----------------------|------------------------|
| Mrs Sharon Bamber    | 0151 678 5111 Ext 2847 |
| Dr Elizabeth Thursby | 0151 552 1875          |

#### Medical Secretaries for clinical enquiries (Wirral)

|                                     |                       |
|-------------------------------------|-----------------------|
| Monday – Friday: 09-00hrs– 17-00hrs | 01244 362500 Option 3 |
|-------------------------------------|-----------------------|

Referrals for Microbiology clinical advice for inpatients should be sent via Cerner, using the ORDER "Refer Microbiology Inpatient."

This service will operate during routine working hours between 9:00 - 4.30pm for inpatients only.

For any inpatient referrals on weekdays between 4.30 pm - 5:00 pm, please call Ext 7875

For outpatients during working hours, please call Ext 7875.

#### Consultant Staff (Countess of Chester)

|                   |              |
|-------------------|--------------|
| Dr Ildiko Kustos  | 01244 366785 |
| Dr Jeremy Gardner | 01244 366788 |

#### Staff Grade Doctor (CoCH)

|                |              |
|----------------|--------------|
| Dr Muna Yousif | 01244 363518 |
|----------------|--------------|

#### Clinical Scientist (CoCH)

|                |              |
|----------------|--------------|
| Mrs Sarah Wood | 01244 364062 |
|----------------|--------------|

#### Trainee Clinical Scientist (CoCH)

|                |              |
|----------------|--------------|
| Eleanor Senior | 01244 366773 |
|----------------|--------------|

#### Medical Secretaries for clinical enquiries (Countess of Chester)

|                                      |              |
|--------------------------------------|--------------|
| Monday – Friday: 09-00hrs – 17-00hrs | 01244 366773 |
|--------------------------------------|--------------|

#### Laboratory Manager

|             |              |
|-------------|--------------|
| Ms N Duggan | 01244 362499 |
|-------------|--------------|

#### Technical Manager / Deputy Laboratory Manager

|               |              |
|---------------|--------------|
| Claire Greene |              |
| Dave Bond     | 01244 363352 |

#### Quality Management

|                  |              |
|------------------|--------------|
| Mrs Joanne Evans | 01244 363352 |
|------------------|--------------|

#### Senior Biomedical Scientist Team

wuth.microdutysenior@nhs.net

## 3.0 MEDICAL MICROBIOLOGY – PRINCIPAL SERVICES

### 3.1 Clinical Service

The principal diagnostic laboratory is based at The Croft Business Park, Bromborough. Limited molecular testing is also performed at APH and CoCH (refer to **Section 1.0**) Access to consultative and principal diagnostic service outlined below is available on a 24-hour basis.

### 3.2 Diagnostic Service

Chester and Wirral Microbiology Service provides a comprehensive microbiological service in medical bacteriology, mycology, virology, parasitology and serological investigations. Advice on the selection of appropriate diagnostic specimens, their collection and transport is available.

The laboratory is accredited by the United Kingdom Accreditation Service (UKAS) to ISO 15189 for the test repertoire stated on the Schedule of Accreditation, which can be accessed at: [9595 Medical Multiple \(ukas.com\)](https://www.ukas.com/9595/Medical/Multiple)



**Any tests reported that are not on the UKAS Schedule of Accreditation will have the following report code added.**

**“Test internally verified awaiting UKAS Accreditation”**

**Results of clinical significance** are phoned through to the surgery or relevant medical staff, irrespective of whether the original request is marked as urgent or routine.

**Delay in results** – if there is going to be a significant delay in the availability of results, the users will be informed.

#### Antimicrobial Therapy and Clinical Consultation

When a conclusive microbiological diagnosis has been reached, optimum therapeutic regimens are reported when necessary. They will be reported as:

- **S – Susceptible, standard dosing regimen:** A microorganism is categorised as *Susceptible, standard dosing regimen\**, when there is a high likelihood of therapeutic success using a standard dosing regimen of the agent
- **I –Susceptible, increased exposure\*:** increasing the dose may improve the chance of treatment success
- **R – Resistant:** A microorganism is categorised as *Resistant* when there is a high likelihood of therapeutic failure even when there is increased exposure\*

\* Exposure is a function of how the mode of administration, dose, dosing interval, infusion time, as well as distribution, metabolism and excretion of the antimicrobial agent will influence the infecting organism at the site of infection.

Ref: Redefining S, I and R 2019 – [www.eucast.org](http://www.eucast.org)

The serum concentration of relatively toxic antimicrobials and those used in critical infections are monitored.

The following empirical (blind / provisional) prescribing regimens can be found in the Wirral Prescribers' Guide and The Chester Joint Formulary

- (a) For patients with severe sepsis and
- (b) When the microbiological diagnosis is inconclusive

### **3.3 Teaching and Training**

The Department of Medical Microbiology supports scientific and professional training for its staff, as well as the teaching of science students attending local universities and colleges. It is also actively involved in providing work experience placements for BTEC students from local colleges. Placements are also given to students undertaking the Biomedical Science degree course at Liverpool John Moore's University and University of Chester.

### **3.4 Document Control**

All documents used in Microbiology are managed electronically via Q pulse and backed up on Countess of Chester Servers to protect their integrity.

There are policies, procedures and templates specific to Microbiology as well as shared directorate documents.

The department is obliged to follow Trust policy and procedures. To avoid duplication some of these policy and procedure documents are used in place of departmental ones.

The Laboratory is hosted by Wirral University Teaching Hospital NHS Foundation Trust policies and procedures are located on the intranet

### **3.5 Patient Confidentiality/Personal information**

Wirral Hospitals adopts the NHS Information Governance framework to ensure patient, staff and other confidential information is handled securely and safely. The Wirral Hospitals Information Governance policy (ref 095) relates to all information used by the Trust and its employees and to other NHS policies and legislation. Through its mandatory staff induction programme, it ensures staff are made aware and follow procedures documented in this policy and subsequently annual mandatory assessments are required to allow the trust to monitor its compliance.

### **3.6 Consent**

Informed consent for the venepuncture procedure and the testing of samples taken is required. For patients attending our Phlebotomy Clinics consent is assumed by them presenting themselves to Phlebotomy with a request form and then presenting their arm for venepuncture.

For patients that are in a hospital bed consent for the procedure is verbally checked by the phlebotomist before they are bled. They have an opportunity to refuse the procedure. For those who are unable to consent, i.e. are unconscious the decision to bleed and complete the relevant testing is taken by the clinical team in the best interests of the patient.

The consent for testing, is assumed to have been gained by the clinician or healthcare professional requesting the test and is inferred to the laboratory through the receipt of the request form with the relevant sample(s). Similarly, for samples that are received from GPs or other laboratories referring work to WUTH Pathology the consent for testing is gained by the clinician or healthcare professional requesting the test and is inferred to the laboratory through the receipt of the request form with the relevant sample(s).

For work referred to us, consent to disclose clinical information and family history to relevant healthcare professionals within WUTH Pathology is inferred by the receipt of the request form with the relevant sample(s). Similarly, work that we refer to other laboratories for specialist testing the consent to disclose is assumed and then inferred to those laboratories by the receipt of a request form with the relevant sample(s).

The consent for other collection procedures, e.g. Invasive, tissue biopsy is gained by the clinical team completing the procedure and the record kept as part of the patient's notes. This is covered by the Trust Consent Policy.

***The decision whether to consent or not is entirely the patient's decision to take.***

However, In the Trusts Consent Policy, it states:

In an emergency, the doctor is excused from conferring with the patient in circumstances of necessity, as for example where the patient requires treatment urgently but is unconscious or otherwise unable to make a decision. The laboratory will therefore accept and process samples in these circumstances.

Clinicians should also be aware that the laboratory may perform reflex tests where clinically indicated or to aid in interpretation.

## **4.0 'URGENT' SPECIMENS FOR MICROBIOLOGICAL INVESTIGATION**

Biomedical Scientists (BMSs) are available in the laboratory 24 hours a day, 7 days a week to process any samples that are considered urgent. Between 09-00 and 19-00 the BMS must be contacted in the laboratory on 01244 362500 with the details of the request.

### **4.1 Out of Hours Service**

19-00hrs to 09-00hrs 7 days a week

Requests for urgent specimens to be processed after 19-00hrs should be directed to the out of hours Biomedical Scientist through switchboard at either the Wirral site or Countess of Chester Hospital.



The following are out-of-hours requests that may be made via Biomedical Scientists:-

- Paediatric MSSUs for children **under** 3 years of age (Microscopy / Culture)
- Material from Sterile Sites – e.g. Synovial fluid, Peritoneal fluid (e.g. Ascites) CSF
- Pus from deep seated abscesses (Other pus swabs etc. contact the Consultant Medical Microbiologist)
- Pus
- Specimens from theatre (excluding bone/tissue)

## 4.2 Taxis

The Department operates a 24/7 service for urgent specimens – see the details above – in the unlikely event that a specimen may not be processed urgently because it has missed the transport run then a taxi may be required to transport the sample to the Laboratory at Bromborough between 17-00hrs and 09-00hrs. Samples must be packaged in a robust container and **no** patient details should be visible on the outer packaging (refer to **Sections 8.4 and 9.1 Packaging and Transport**). Refer to the relevant transport policy: WUTH Transport Goods SOP EFCO2 or

## 5.0 LABELLING REQUIREMENTS FOR REQUESTS (CERNER, ICE AND HANDWRITTEN FORMS)

Requests communicated to the laboratory are as follows:

- Cerner – order labels are attached to the sample (no request form required)
- ICE forms generated by the GP practices.
- Handwritten request forms from wards, clinics, or GPs
- All verbal requests to the laboratory **must** be accompanied by one of the above requests forms for the sample to be processed.

**All Locations within each Trust and GP practice should make requests via the above Order Entry Systems – otherwise results may not be viewable electronically.** The sample, request label, or request form should clearly state the following information for unequivocal identification of the patient and specimen:

- Patient name (in full – no abbreviations)
- Ward, Clinic, or GP name and number/address
- NHS number
- Date of Birth
- Sex
- Type of specimen
- Date and time specimen taken.

**NB** It is **ESSENTIAL** that the laboratory knows the date on which a specimen is taken: processing delayed specimens can yield unhelpful or frankly misleading results and they may be discarded (e.g., urine samples dated 2 days prior to day of receipt). When patients are given a request form and asked to provide a specimen **they should be asked to ensure that the date on which the specimen was collected is given on the container and the form.**

- Tests required (specify 'TB' if required)
- Only request 'Miscellaneous Microbiology' if the appropriate investigation is not listed on the screen and state the specific investigation required in the clinical details field.
- Antibiotic treatment (recent, current or intended)
- All relevant clinical details
- History of recent foreign travel, if applicable
- Risk status, if applicable \* **see section 6.2**
- Date of onset and duration of illness, particularly for serology
- Specify anatomical site from which "wound" specimens were taken.
- Key epidemiological information, e.g., for faeces
- Request 'OCP' (ova, cysts and parasites) if appropriate

### Cerner Requesting

CERNER is a paperless system that will not generate a request form.

All CERNER requests should have a status of collected in CERNER before sending to the laboratory. The specimens should be sent with the printed label on the specimen. All of the above details are necessary to include when making a CERNER or Excelicare request.

If uncertain about the exact test and terminology, please give a detailed clinical history as this can help the Laboratory staff to decide the most appropriate investigation.

## 6. LABELLING REQUIREMENTS FOR SPECIMENS

- The specimen must be labelled with the patient details as on the request form.
- Patient name (in full – no abbreviations)
- Ward, Clinic, or GP name and number/address
- NHS number
- Date of Birth
- Sex
- Type of specimen
- Date and time specimen taken (24-hr clock).

### 6.1 Multiple samples taken at different times or from different sites on a single patient must be labelled with:

- with the recommended patient identifiers,
- the specimen types e.g. sputum, urine,
- the time (24-hour clock) when each sample is taken
- the site of specimen. e.g. swabs taken from different areas of the body
- Each different specimen or test request must have its own unique order number.

#### **EXCEPTIONS:**

*Blood Cultures where 2 sample bottles are required. Label both samples with the patient identifiers and order only one unique order number.*

*MRSA samples where Swabs from various sites are combined into one sample collection. Label the one container with the patient identifiers and order only one unique order number.*

- Please note that unlabelled and mislabelled specimens cannot be processed and will be rejected.

**If the laboratory cannot unequivocally identify the sample and match it to a form, then it will be rejected.**

The laboratory will inform senders by means of an electronic or printed report when a specimen has been rejected for the above reasons.

In certain circumstances it may be possible to add tests to samples that the laboratory has already received. It is assumed by the laboratory that the request for additional tests and the sending of samples has patient consent.

### 6.2 Laboratory guidance on appropriate labelling of high-risk samples

#### **Danger of Infection - High Risk Samples**

High risk labels alert laboratory workers to samples which may need additional safety and handling precautions.

**High Risk Labelling NOT required for BBV positive samples**

It can now be confirmed that samples from BBV positive patients no longer need to be specially labelled or packaged.

The laboratories follow 'universal precautions' and treat all samples as potentially hazardous.

**High Risk Labelling IS REQUIRED**

All biological agents causing infection are classified into hazard groups 1-4 by the Health and Safety Executive (HSE). Those in groups 3 and 4 are deemed to be a serious hazard and infective samples taken from such should be labelled high risk. But the list of biological agents in these groups is extensive and includes many rare infections unlikely to be encountered in general practice in the UK. Below is a relevant summary of when labelling would be appropriate.

It is very important that the request contains **clear clinical details** so staff can help identify which samples may require additional precautions when processing.

**Clinical details that should alert the clinician to the possibility of a 'high risk' pathogen**

Suspected infection in these groups:

|                                   |  |
|-----------------------------------|--|
| <b>Bloody diarrhoea/dysentery</b> | E Coli O157 is common and very infectious to laboratory staff.   |
| <b>Injection site infections</b>  | in IV drug users - risk of anthrax   |
| <b>Haemoptysis</b>                | If TB is suspected   |
| <b>Laboratory workers</b>         | especially those handling 'hazard group 3 or 4' organisms (the individual worker should be able to inform you of this) |
| <b>Animal exposure</b>            | (farm animals/exotic animals). Animals and animal products or by-products from overseas are a particular concern.      |

**Apart from E Coli O157 diarrhoea and bovine TB, it is unusual to get high risk infections from UK farm animals**

**Foreign travel mainly outside Western Europe\*** – particularly if basic or rural conditions, animal exposure, volunteer/healthcare work (visiting family)

Consider viral haemorrhagic fever in endemic areas.

*(When should I suspect Viral Haemorrhagic fever (VHF)?*

*If the patient has a fever (>38 C) or history of a fever in the last 24hrs and has either returned from a viral haemorrhagic fever endemic area in the last 21 days or had contact with an individual, animal (or specimens from such) that is known or strongly suspected to have VHF).*

VHF endemic areas include East and West Africa, Central Asia and the Middle East.

A detailed list can be found on pages 35-39 of the following document: [Viral haemorrhagic fever: ACDP algorithm and guidance on management of patients - GOV.UK](#)

**\*In cases of suspected viral haemorrhagic fever, the Regional Infectious Diseases Unit should be contacted before ANY samples are taken.**

**What conditions caused by 'high risk' organisms can present in the UK?**

Anthrax  
Brucellosis  
Diarrhoea associated with E Coli O157  
Travel associated exotic fungal infections  
Meliodosis  
Tuberculosis  
Histoplasma  
Typhoid (Enteric Fever)  
Viral Haemorrhagic Fevers  
Rabies

**This list is not exhaustive.**

**If a high-risk pathogen is suspected: -**

- 1. Please document clinical details clearly on the request.**
- 2. Ensure containers are properly sealed and bagged.**
- 3. Label transport bags as “high-risk” with a yellow high-risk sticker \*\***

**\*\* The exception to this is patients with suspected viral haemorrhagic fever. These cases must be discussed with medical staff at the Regional Infectious Diseases Unit. They will provide a risk assessment and advise on sampling if necessary.**

The full Health and Safety Executive (HSE) guidance can be found at [Provision of key clinical information on laboratory specimen request forms - HSE](#)

A complete list of organisms and their Hazard Group can be found at [The Approved List of biological agents: Advisory Committee on Dangerous Pathogens](#)

Country specific information on recent outbreaks (that is updated regularly) is available on Travax: <http://www.travax.nhs.uk>

Any queries please send us an email on [wuth.microdutysenior@nhs.net](mailto:wuth.microdutysenior@nhs.net) or call us on 01244363351 and ask to speak to the Duty Senior or Specialist biomedical Scientist out of hours.

### **6.3 Storage of samples**

The table below indicates how long samples are kept in the laboratory before disposal. Requests for extra tests must be received within the sample storage period and must be accompanied by a request form. Please telephone the laboratory before requesting extra tests to ensure the sample is available and still viable.

| <b>Sample</b> | <b>Time Kept</b>  |
|---------------|---|
| Faeces        | 1 week after primary culture. Aliquot of C diff toxin positive samples – min 3 months |

|   |  |
|---|--|
| Respiratory samples   | 1 week after primary culture   |
| Flu samples   | 1 week after primary screening   |
| Swabs, fluids and aspirates   | 1 weeks after primary culture  |
| Urines  | 1 weeks after primary culture  |
| CSF samples   | 1 month after collection (? CJD samples) stores securely in -70°C freezer until Reference Laboratory report is received) |
| Blood cultures  | Positive bottles – until all follow up work is finalised.<br>Negative bottles – discarded after 5 days incubation.       |
| Tissue / bone / cartilage   | 1 month after primary culture  |
| Stained slides  | 1 week after primary culture   |
| Postmortem tissue   | 1 month after culture  |
| Mycology samples for culture  | Generally, all samples are processed in KOH  |
| MRSA/CPE/VRE screening swabs  | 1 week after PCR/primary culture   |
| Left over serum from first pregnancy booking  | -20°C 2 years  |
| Left over serum or plasma (other than transplant-related)                                 | Minimum 2 years -20°C  |
| Left over serum or plasma from transplantation, post transplantation donor/recipient sera | Minimum 11 years at -20°C  |
| Serum from accidental exposures to blood and bodily fluids                                | Minimum 2 years -20°C  |
| All clots   | Minimum 1 week at 4°C  |
| Unprocessed samples (e.g., spares)  | Minimum 1 week at 4°C  |
| Human milk and serum from milk donors   | Milk -10 years at -80°C<br>Sera – 10 years at -20 °C   |

## 7.0 STANDARD PROCEDURES FOR THE SAFE COLLECTION OF SPECIMENS

These procedures concern all clinical staff who are qualified to collect diagnostic specimens from patients.

Firstly, check who the patient is before taking the sample, both verbally and using the patient's wristband.

**N.B.** *Staff must always follow aseptic techniques when handling blood, body fluids, excretions, or secretions, even when these have not been specified as infectious.*

### Objectives

All staff must be aware of the potential physical and infectious hazards, associated with these procedures, and should therefore collect specimens:

- 1) Being mindful of personal safety, without injury or exposure of themselves and of collective safety, without exposing colleagues who are involved with the handling, transport and laboratory investigations of specimens, to physical or infectious hazards.
- 2) Staff collecting specimens must take care to prevent contaminating themselves, their environment, the external surfaces of the specimen containers, or the accompanying test request forms. If gross contamination of the hands with blood, faeces or other biological fluids is anticipated, then gloves should be worn. Hands should always be washed after taking specimens. If splashing into the eyes or on to mucous membranes is anticipated goggles should be worn.
- 3) In addition, specimens should be collected aseptically, without allowing contamination by extraneous and, therefore, irrelevant micro-organisms. Contaminated specimens can adversely affect the validity of many laboratory results. For example, the microbiological investigation of contaminated blood or other materials from sites, which are normally sterile, can commit patients to unwarranted courses of expensive and potentially toxic treatment.

### Before you start

- 1) Ensure that the lighting conditions are adequate.
- 2) Select the correct specimen container (s), appropriate for the type of specimen, and keep the container close to the site from which the specimen is to be obtained.
- 3) Complete, legibly and fully, all section of the label on the specimen container and, check the details on the computer-generated request form are correct or, where used, the test request forms.
- 4) If you suspect, or are aware of, an infection with a Hazard Group 3 pathogen (example of relatively common Hazard Group 3 pathogens are Hepatitis B virus, Human



Immunodeficiency virus and *Mycobacterium tuberculosis*), or suspect Monkey Pox it must be mentioned in the clinical details sent with the specimen.

- 5) If you suspect, or are aware of, an infection with a Hazard Group 4 pathogen (Viral haemorrhagic fevers, e.g., Ebola and Lassa) do not attempt to collect any specimen. Inform the Infection Control Doctor for the Trust through switchboard.

**When you have finished all waste generated from obtaining a specimen should be disposed of according to Local Waste Disposal Protocols.**

### **7.1 Procedure for venepuncture to obtain a specimen of blood / blood cultures.**

Please refer to the Trust's intranet site for guidance on this procedure.

### **7.2 Procedure for the collection of pus or exudates**

Where there are clinical signs of infection i.e., inflammation, oedema, pyrexia, pain or purulent exudate, it is preferable to obtain a specimen of pus rather than to take a swab.

Pus or exudate can be drawn up in a syringe and transferred to a universal container.

Taking a Transwab (blue top) or Charcoal swab (black top), remove the swab and gently but firmly rotate it on the surface directly where infection is suspected. Do not take swabs from slough or necrotic tissue. Place the swab into the transport medium.

Ensure that the specimen containers are labelled accurately and place, with the completed request form, in the appropriate pockets of the clear mini-grip transport bag for transportation to the Department of Medical Microbiology (via Pathology).

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 is preferable to storage at ambient temperature.

### **7.3 Procedure for the collection of screening swabs (MRSA, CPE, VRE)**

These swabs should only be taken on the advice of the Community Infection Control Team or to comply with individual hospital protocols. They are taken to ascertain whether a patient is colonised with potentially pathogenic bacteria e.g., MRSA, VRE, CPE. If clinical infection is suspected, please send another swab from ulcers, wounds etc. separately for MC&S.

#### **MRSA culture**

For routine MRSA screens nose and groin swabs are required. Axillae swabs are only tested from pre-pacemaker insertion patients on CCU and CCS.



**Collection:**

**Nasal** – rotate the swab gently but firmly around the anterior nares of each nostril. One swab can be used for both nostrils.

**Groin** – rotate the swab gently but firmly over each area. One swab can be used for both groins.

**MRSA PCR (CoCH only)**

Only nose and groin swabs from patient on ITU at CoCH are tested for MRSA by PCR. Swabs from contact screens or any other patient are only tested by PCR if requested by the Infection Control Nurses at CoCH. **Pink topped liquid swabs are required for MRSA PCR** – contact Infection Control if guidance is required.

**CPE / VRE culture**

Using a Transwabs (blue or black top), take a rectal swab – refer to Trust guidelines for guidance on collection.

**CPE PCR (WUTH only)**

Refer to Infection Control policies for guidance on which patients require molecular testing for CPE. Using a **dual** swab (red top), take a rectal swab – refer to Trust guidelines for guidance on collection.

**VRE PCR (CoCH only)**

Refer to Infection Control policies for guidance on which patients require molecular testing for VRE. Using a **dual** swab (red top), take a rectal swab – refer to Trust guidelines for guidance on collection.

Ensure that swabs are labelled accurately. Place, with the completed request form, in the appropriate pockets of the clear minigrip transport bag for transportation to the Department of Medical Microbiology (via Pathology).

Specimens should be transported and processed as soon as possible.

**7.4 Procedures for the collection of Urogenital Samples**

**N.B.** If an expanded screen for sexually transmitted diseases is required, the patient should be referred to the local Sexual Health Service.

**7.4.1 Collection of urogenital samples for Microscopy, Culture & Sensitivity (MC&S)****High vaginal swabs and Cervical/Endocervical swabs**

Place the patient in dorsal position, supported by a pillow and ask her to bring her heels together, bend her legs and then draw her heels towards her bottom.

Moisten the speculum with warm water and insert into the vagina to separate the vaginal walls. Wipe away any excess cervical mucus with a tissue.

**HVS:** Using a blue or black topped swab, sample as high as possible into the vault and swab the vaginal wall.

**Cervical/endocervical swab:** Using a blue or black topped swab, gently insert a swab into the endocervical canal and rotate to obtain any exudate. Try to avoid contact with the vaginal mucosa when removing the swab.

Remove speculum and wipe vaginal / vulval area with a tissue.

Ensure that the swab is labelled accurately and place with the completed request form, in the appropriate pockets of the clear minigrip transport bag for transportation to the Department of Medical Microbiology (via Pathology).

**NB. Charcoal based transport swabs prolong the survival of gonococci compared to non-charcoal-based swabs.**

#### **Low vaginal swab**

Place the patient in dorsal position, supported by a pillow and ask her to bring her heels together, bend her legs and then draw her heels towards her bottom.

Insert the swab into the lower part of the vagina and rotate gently but firmly.

Ensure that the swab is labelled accurately and place, with the completed request form, in the appropriate pockets of the clear minigrip transport bag for transportation to the Department of Medical Microbiology (via Pathology).

#### **Urethral swabs**

Avoid contamination with micro-organisms from the vulva or the foreskin. Small swabs are available for the purpose. The patient should not have passed urine for at least 1 hour. For males, if discharge is not apparent attempt to "milk" it out of the penis. Pass the swab gently through the urethral meatus and roll around. Place the swab in the plastic transport sheath containing the black charcoal containing Amies medium. **NB specimens for Chlamydia investigations should be collected after the swab for MC&S.**

All specimens should be transported and processed as soon as possible. If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### **7.4.2 Collection of urogenital samples for molecular investigations**

Urogenital samples for molecular investigations should be collected as per collection for samples for MC&S, but the samples should be collected using the following specimen containers:

#### ***Chlamydia trichomonas* / *Neisseria gonorrhoea* (CT/NG) and *Trichomonas vaginalis* (TV) PCR**

##### **For all requestors**

If Chlamydia/gonorrhoea infection is suspected, swabs or urine can be submitted for analysis. The sample must be collected/placed into the appropriate Roche Cobas **Uni** Swab/urine container. (available from NHS Supplies or [wuth.ctngconsumables@nhs.net](mailto:wuth.ctngconsumables@nhs.net)).

- Roche Cobas **Uni** Swab (clinical and self-taken samples)
- Roche Cobas **Uni** Swab (urethral, oropharyngeal & rectal samples)
- Roche Cobas DUAL swabs (for endocervical samples)
- Roche Cobas Urine Collection tube (for first catch urine from males or females)

NB Investigation for *Trichomonas vaginalis* (TV) can be performed on self-taken vaginal swabs placed in a Roche Cobas Uni Swab **BUT** TV processing will only be performed if specifically requested.

Following specimen collection, store the Cobas PCR Media tube containing the specimen swab at 2-30°C. The specimen is viable for up to 3 months.

### **7.5 Procedure for the collection of sputum**

The material required is fresh sputum from the lower respiratory tract, expectorated by deep coughing. When the cough is dry, physiotherapy, postural drainage or inhalation of an aerosol may be helpful. Saliva and postnasal secretions are not suitable. Early morning specimens for examination of *Mycobacterium species* should be collected on at least 3 consecutive days.

Routine sputum microscopy is not worthwhile, but will be done urgently were Staphylococcal pneumonia is suspected or where specifically requested.

Ensure that the specimen container is labelled accurately and place, with the completed request form, in the appropriate specimen transport bag for transportation to the Department of Medical Microbiology (via Pathology).

Collect specimens before starting antimicrobial therapy where possible.

BAL and sputum should be processed promptly to give the best opportunity to culture pathogenic organisms and reduce the risk of overgrowth with contaminants.

If processing has to be delayed up to 24 hours, refrigeration is preferable to storage at ambient temperature.

#### **Bronchial Lavage**

Please inform the laboratory for urgent processing.

*NB. Legionella/PCP investigations – please contact the Laboratory if required.*

### **7.6 Collection of a mid-stream specimen of urine (MSSU) for culture and sensitivity**

**Ensure that the patient is physically clean.**

If the patient has had the perineum washed in the last 12 hours (i.e., has had a shower or bath), further cleansing of the perineal area before urine collection is not necessary.

If the patient:

- Is incontinent and / or:
- Has had their bowels opened since washing the area:

**The collection of urine must be postponed** until the perineal area has been washed.

Catch the middle portion of the urine in a clean wide-mouth receptacle. Such a receptacle need not be sterile: any container, previously washed thoroughly with detergent and hot water and stored dry, is suitable.

A sample of the middle portion of the urine must be poured into a 20ml **red** capped universal container (boric acid) with all sections on the label completed. Very small samples from paediatric patients only may be collected into a 20ml white capped sterile universal (white capped samples **MUST** be received in the lab **within 4 hours** of collection).

If processing is delayed for up to 48hr, refrigeration is essential. Alternatively, the specimen may be collected in a CE marked leak proof container with boric acid preservative. This increases the maximum permissible time for transport to the laboratory to up to 96hr.

### **7.7 Collection of a specimen of urine from a catheter (CSU)**

When small volumes of fresh urine are required for laboratory investigations, the distal end of the catheter, or preferably the sampling port if present, must be disinfected with 70% isopropyl alcohol and urine aspirated with a sterile syringe.

The urine must be transferred to a 20ml red capped universal container (boric acid) with all sections on the label completed.

If large volumes of urine for laboratory tests are required, these should be obtained aseptically from the drainage bag.

### **7.8 Procedure for the collection of a specimen of faeces**

When collecting a specimen of **Faeces**, it should be obtained in a convenient container and transferred into a sterile container with a wooden disposable spatula.

#### **Minimum volume of sample:**

- A liquid specimen of 1-2ml is sufficient.
- 1 gram (large pea-size) of solid specimen

Rectal swabs are not a reasonable substitute for faeces: except for CPE/VRE screening.

Faeces for parasites – the recommendation is 3 specimens taken on different days for optimum recovery.

Ensure that the specimen container is labelled accurately and place, with the completed request form, in the appropriate specimen transport bag for transportation to the Department of Medical Microbiology (via Pathology).

For the detection of ova of *Enterobius vermicularis* (threadworm): use with a plain swab, moisten with sterile saline, wipe firmly around the anal margin and place the swab into a sterile universal container.

Important pathogens such as *Shigella* species may not survive the pH changes that occur in faecal specimens if not promptly delivered to the laboratory, even if refrigerated.

If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### **7.9 Procedure for the collection of a pernasal swab**

Gently insert the fine, flexible pernasal swabs (sky blue top) swab horizontally to the back of the nose. If obstruction is encountered, withdraw and re-insert through the other nostril.

Ensure that the swab is labelled accurately and place, with the completed request form, in the appropriate specimen transport bag for transport to the Department of Medical Microbiology (via Pathology).

Collect specimens before antimicrobial therapy where possible.

Specimens should be transported and processed as soon as possible. If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### **7.10 Aspirates and fluids from normally sterile sites**

Collect the specimen with a sterile syringe. Transfer a maximum 20ml into a sterile universal container. Ensure the cap is tightly screwed on.

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. If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### **7.11 Cerebrospinal fluid**

? Meningitis - An adequate amount is essential – send at least 2-3ml. Label the samples as you collect them as the lab need to know which sample is the most valuable for cell count and culture. This is particularly important if *Mycobacterium tuberculosis* infection is suspected where small numbers of organisms may be present. For exclusion of mycobacterial CNS infection at least 6mls CSF is required: this will be processed by Microbiology at Manchester Royal Infirmary. If there are smaller volumes, then an automated comment will be produced indicating low volume. The results of microscopy and any positive cultures are always telephoned.

? Subarachnoid haemorrhage (SAH) if there is a clinical suspicion of SAH and the specimen is bloodstained send the 1<sup>st</sup> and 3<sup>rd</sup> samples so that differential red blood cell counts may be performed. The results of microscopy and any positive culture are always telephoned. Always inform the laboratory if SAH is a possibility.

Time between collection to microscopy and culture should occur within a maximum of 2 hours. Cells disintegrate and a delay may produce a cell count that does not reflect the clinical situation of the patient. Specimens should be transported and processed as soon as possible. Do not refrigerate specimen until after microscopy and bacterial culture have been performed. The specimen should then be refrigerated pending further investigation.

### **7.12 Wound Swabs**

Decontaminate the skin to remove as much of the superficial flora. Using a blue or black topped transwab, remove the swab and gently but firmly rotate it on the surface directly where infection is suspected. Do not take swabs from slough or necrotic tissue. Place the swab into the transport medium. If pus is present send as much as possible in a sterile universal container.

Specimens should be transported and processed as soon as possible. If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### **7.13 Collection of specimens for mycology investigations**

#### **Skin**

Patient's skin and nails can be swabbed with 70% alcohol prior to collection of the specimen, this is especially important if creams, lotions or powders have been applied. The edges of skin lesions yield the greatest quantities of viable fungus. Lesions should be scraped with a blunt scalpel blade.

#### **Nails**

Good nail samples are difficult to obtain. It should be specified whether the sample is from the fingernails or toenails. Material should be taken from any discoloured, dystrophic or brittle parts of the nail. The affected nail should be cut as far back as possible through the entire thickness and should include any crumbly material. Nail drills, scalpels and nail elevators may be helpful but must be sterilized between patients. When there is superficial involvement (as in white superficial onychomycosis) nail scrapings may be taken with a curette. If associated skin lesions are present, samples from these are likely to be infected with the same organism and are more likely to give a positive culture.

#### **Hair**

Samples from the scalp should include skin scales and plucked hairs or hair stumps. Cut hairs are not suitable for direct examination as the infected area is usually close to the scalp surface. Plastic hairbrushes, scalp massage pads or plastic toothbrushes may be used to sample scalps for culture where there is little obvious scaling, but such samples do not replace a scraping for direct examination.

Collect specimens before antifungal therapy where possible.  
Specimens should be transported and processed as soon as possible.

Specimens should be kept at room temperature and transported and processed as soon as possible although, provided the samples are kept dry, the fungus will remain viable for several months. Samples should be allowed to dry out and kept at room temperature.

### **7.14 Blood for Virology/Serology investigations**

For antibody and antigen assays collect blood in a blood collection tube (red or ochre cap) – 4mls (Wirral), - 8mls (Chester).

For viral DNA/RNA Polymerase Chain Reaction (PCR) tests please send two 4ml EDTA tubes (purple cap) – (Wirral and Chester).

$\beta$ -D-Glucan any tube with clotting accelerator (red or ochre cap). **The test is not validated for sputum specimens.**



### Paediatric blood tubes

- Chester EDTA = Red  
Clotted blood = Brown
- Wirral EDTA = Purple  
Clotted blood = Red

For the various virology/serology investigations available refer to **Section 10 Investigations and Turnaround Times**

Specimens should be transported and processed as soon as possible.

If processing is delayed, refrigeration is preferable to storage at ambient temperature.

### 7.15 Swabs for viral investigations

Moisten the plastic shafted swab with sterile saline never with Viral Transport Media (VTM) before swabbing. Using a sterile saline moistened plastic shafted swab, swab the area concerned or vesicles, if vesicles present burst vesicle with sterile needle and swab fluid released. Snap off the swab tips into VTM.

Transport as soon as possible at ambient temperature.

If transport is delayed, samples may be stored at room temperature for up to 24 hours.

### 7.16 Fluids and Pus for viral investigations

Collect as much fluid/pus as possible in a universal container.

### 7.17 Corneal Scrapes

#### Collection

A standard operating procedure is available in the Eye Clinic; the following is a summary of this document.

#### During Core Laboratory Hours – Mon – Sun 08-45 – 19-00.

1. Please contact the Consultant Microbiologist, to discuss the case.

Two corneal scrape kits (containing a bijoux of broth and a glass slide) are kept in the Ophthalmology Clinic and a further kit is kept in the A+E department. The lab sends these kits to Ophthalmology upon request. To request further kits please telephone the Microbiology Department between 9am-4pm Monday to Friday (**01244 362500**)

Specimens should be transported and processed as soon as possible.

If processing is delayed, refrigeration is preferable to storage at ambient temperature.

If specimens for investigation for amoebae cannot be processed within 8hr, it is preferable to store them at ambient temperature.

Do not freeze specimens.

### **7.18 Nose and throat swabs/Nasal Aspirations**

A single swab can be used. Swab the throat first then nose and place into one pot of viral transport medium: OR

If 2 swabs in the pack, place BOTH the throat & nose swab into the same VTM.

Taking throat swabs: Ask patient to open mouth wide. Using swab, vigorously swab only the posterior pharyngeal wall.

Taking nose swabs: Tilt head back slightly and gently insert swab along the medial part of the septum. Rotate swab several times and remove.

**The shafts of the swabs should be broken off at the break point so that they fit into the VTM container. Firmly secure cap.**

Aspirates should be placed in a dry sterile specimen container. Ensure the cap is tightly screwed on.

Order the correct PCR tests and label the specimen accordingly.

Specimens should be transported and processed as soon as possible.

## **8.0 TRANSPORT OF CLINICAL SPECIMENS FROM WIRRAL**

### **8.1 Specimens collected and sent from Arrowe Park Hospital**

#### **Monday to Friday**

**From 09-00hrs. until 14-00hrs daily** there is an hourly collection of Pathology specimens from wards by the portering staff.

**After 14-00hrs bleep porters on 2145 to pick up specimens** (except Blood cultures which should be sent by pneumatic tube or taken to Specimen Reception at Laboratory Medicine, Arrowe Park). Blood cultures must be sent to pathology immediately after collection to ensure they are stored in the correct conditions.

The last routine transport from Arrowe Park Hospital to the diagnostic Laboratory at Bromborough is at **16-45hrs**.

#### **Weekend and Bank Holidays**

**At approximately 09-00hrs** a single collection from the ward is made by the portering staff.

If specimens miss the collection, then they should be sent to the Arrowe Park Specimen Reception.



There is a scheduled pick-up of specimens, by van, from Arrowe Park Pathology Laboratory until 15-00hrs.

### **Urgent Specimens – Normal Hours (Mon-Sun)**

If specimens require urgent processing during normal working hours, then please contact the Microbiology Department and inform us of the patient, the ward and any tests required on Extension **4511**. Arrange delivery by telephoning the Porters to request urgent collection of samples to be taken directly to the Pathology Laboratory Specimen Reception. Refer to **Section 8.4 Packaging and Transport** for guidance on packaging samples for transport by taxi.

### **Urgent Specimens – Out of Hours (Mon-Sun)**

Weekdays 18-30 – 09-00hrs, Saturday, Sunday and Bank Holidays an out of hours service is available. The Biomedical Scientist can be reached by the Hospital switchboard. Clinical advice is always available from the Consultant Microbiologist (available through the Hospital switchboard).

**Do not send specimens to Microbiology Department during ‘Out of Hours’ unless instructed to do so by the Biomedical Scientist.**

## ***8.2 Specimens collected and sent from Clatterbridge General Hospital***

### **Monday to Sunday**

Specimens need to be taken to the first floor Blood Science Lab at CCC. Drivers will collect CGH samples every hour from 09-00hrs. This Service will run between CGH, APH and the Microbiology lab at Bromborough on a continuous loop. The last collection from CCC is at 16-30. Any non-urgent samples that will not be ready for transport by 16-30 should be refrigerated and made ready for transport the next day.

**For URGENT specimens only after 18-30hrs** please contact the Out of Hours BMS, via Arrowe Park Switchboard.

## ***8.3 Specimens collected and sent from GP Practices***

Samples from General Practice for Laboratory Medicine will be collected by a hospital courier or SRCL/ERS Courier Service Monday to Friday only.

## ***8.4 Packaging and transport***

Before the specimens are collected by porters, couriers, volunteers, nursing and support staff ensure that specimens and request forms are placed correctly into the min-grip plastic bags. Specimens should be placed in the pocket of the plastic bag and grip seal sealed. The request

form should be slid into the sleeve of the plastic bag. The specimen should then be placed in the large Blue Microbiology specimen bags (with an absorbent material to comply with United Nations standard Packing Instruction P620) for collection. All microbiology specimens that are not collected promptly should be refrigerated, unless otherwise stated in this guide.

Specimens that are to be transported by taxi from the hospital to the main Microbiology Laboratory must be packaged in a robust container and **no** patient details should be visible on the outer packaging – refer to the European Agreement Concerning the International Carriage of Dangerous Good by road (ADR2021).

**N.B** *The plastic transport bags, if properly sealed, are designed to contain accidental specimen leakage from the container. Spontaneous specimen discharge, due to defective materials, is rare. Most incidents of specimen leakage are due to the fact that neither the container nor the integral bag strips have been closed properly. If both container and transport bag are closed correctly, the practice of 'double bagging', even when an infection with a Hazard Group 3 pathogen is suspected, does not confer any additional safety advantage and is, therefore, unnecessary*

*The containers supplied, comply with standards BS4851 and BS5213 for leakage and spontaneous discharge. Leaked containers frequently result in irreplaceable loss of specimens and, equally as important, staff to unwarranted hazards of infection.*

Members of the public who come across large blue specimen bags containing specimens should telephone the lab.

## 9.0 TRANSPORT OF CLINICAL SPECIMENS FROM CHESTER

### Routine Specimens

Specimens are delivered to the Microbiology Dept (via Pathology) throughout the working day, Monday – Friday, from the Pathology Laboratory reception via hospital transport vans. However, the last collection of the day leaves there at 17-00hrs. Therefore, specimens must have reached the pathology laboratory reception by the porters, well in advance of this time. Samples will be collected from Pathology throughout the day on Saturday and Sunday up until 15-00hrs. Routine samples may be transported to Pathology via the pneumatic air tube system providing the samples are correctly packaged with secure lids, except for CSF samples.

### Urgent Specimens – Normal Hours

If specimens require urgent processing during normal working hours, then please contact the Microbiology Department and inform us of the patient, the ward and any tests required on Extension 2500. Arrange delivery by telephoning the Porters to request urgent collection of samples to be taken directly to the Pathology Laboratory Specimen Reception. Refer to **Section 9.1 Packaging and Transport** for guidance on packaging samples for transport by taxi.

### Urgent Specimens – Out of Hours

Weekdays 18-30 – 09-00hrs, Saturday, Sunday and Bank Holidays an out of hours service is available. The out of hours Biomedical Scientist can be reached by the Hospital switchboard.

Clinical advice is always available from the Consultant Microbiologist (available through the Hospital switchboard).

**Do not send specimens to Microbiology Department during 'Out of Hours' unless instructed to do so by the Biomedical Scientist.**

### 9.1 Packaging and transport

See Section 8.4

## 10.0 INVESTIGATIONS AND TURNAROUND TIMES

For specimen collection see **Section 7**

| Specimen Investigation  | Specimens and Comments  | Referred to Ref Lab* | Turnaround times |
|---|---|----------------------|------------------|
| Adenovirus PCR  | Broncho-alveolar lavage<br>CSF<br>Eye swab<br>Urine<br>Urethral swab<br>EDTA blood preferred. | 2                    | 2-5 days         |
| Amoebic Antibodies  | Clotted blood   | 6                    | 5-7 days         |
| Antenatal Screen:<br>Hepatitis B surface antigen/HIV/Syphilis/ (Rubella)  | Clotted blood<br><br>Antenatal rubella no longer advised                                      |                      | 1-5 days         |
| Staphylococcal Antibody<br><br>(Reference lab test-availability limited so needs discussion with Consultant Microbiologist) | Clotted blood   | 1                    | 5-7 days         |
| Anti Streptolysin-O antibody and anti streptodornase  | Clotted blood   | 4                    | 5-7 days         |
| Arbovirus<br><br>(Includes Flaviviruses such as West Nile, yellow fever, dengue   | Clotted blood<br>EDTA Blood<br>Urine<br><br>REQUEST CAN ONLY BE MADE IN                       | 3                    | 5-7 days         |

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| and Alphavirus such as chikungunya, Ross River, EEE, WEE   | CONSULTATION WITH A MICROBIOLOGY CONSULTANT   |    |  |
| Aspergillus Antibodies   | Clotted blood (Immunology Request at Chester)   |    | 5-7 days   |
| Aspergillus antigen (Galactomannan)  | Clotted blood   | 10 | 2-5 days   |
| Aspergillus PCR  | Bronchoalveolar Lavage<br>Sputum  | 2  | 2-5 days   |
|  | EDTA Blood  | 4  | 5-7 days   |
| Aspirates and fluids from normally sterile sites (joint, ascites, peritoneal and pleural fluids) |   |    | 2-5 days<br><br>Urgent Cell Count/Gram Stain 1 Hour from receipt |
| Avian Precipitins  | Clotted blood (Immunology requests at Chester)  |    | 10 days  |
| Atypical Pneumonia   | For Legionella, please send urine sample for <b>Urinary Legionella Antigen</b><br><br>Mycoplasma pneumoniae serology available: clotted blood<br><br>Molecular techniques i.e., PCR looking for Viruses, mycoplasma, pertussis etc. require.<br>respiratory tract samples e.g., Sputum, BAL. NPA: In clear universal container, or trap.<br><br>Nose and throat swab / NP swab in VTM | 2  | 2-4 days   |
| Bartonella (cat scratch)   | Specific test for Bartonella no longer available – please contact Consultant Microbiologist for further advice.   |    |  |
| BK/JC PCR<br><br>Haemorrhagic cystitis<br><br>Progressive multifocal leukoencephalopathy         | EDTA Blood<br>CSF   | 2  | 2-5 days   |

|   |   |   |  |
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| Renal transplant  |   |   |  |
| <p>Blood Cultures for diagnosis of sepsis, bacteraemia and infective endocarditis</p> <p>Blood cultures must be sent to pathology immediately after collection to ensure they are stored in the correct conditions.</p> | <p>Blood culture set = Aerobic (blue or green top) AND anaerobic (purple top) blood culture bottles (adults / adolescent) 4-10ml blood per bottle <b>Inoculate O<sub>2</sub> bottle first</b></p> <p>Paediatric bottle (yellow top) 1-4ml blood</p> <p>Culture is no longer extended beyond 5 days, if endocarditis is suspected contact the Consultant Microbiologist as it may be appropriate to refer an aliquot of the blood culture samples for 16s (pan-bacterial PCR)</p> <p>Normally sterile body fluids may also be inoculated into blood culture bottles.</p> |   | Up to 5 days (incubated for 5 days before being discarded as negative) |
| <p>Bordetella pertussis Serology</p> <p>Pertussis (whooping cough)</p>  | Clotted blood   | 1 | 5-7 days   |
| <p>Bordetella PCR</p> <p>Pertussis (whooping cough)</p>   | Throat swab in viral transport media or pernasal swab   | 2 | 5-7 days   |
| Brucella serology   | Clotted blood   | 8 | 5-7 days   |
| Bronchoalveolar Lavage (Washings)   | Send washings in a sterile universal container  |   | 2-5 days   |
| Campylobacter Serology  | Clotted blood   | 1 | 5-7 days   |
| Candida PCR   | EDTA Blood BAL  | 2 | 5-7 days   |
| Candida Precipitins   | Clotted blood   | 4 | 5-7 days   |
| Catheter specimen of urine (CSU)  | Transfer urine to a sterile universal container (>3ml)  |   | 48 hours   |
| Cervical swab   | For the culture of gonorrhoea use a Black topped Microbiology (Charcoal) swab and transport to the laboratory   |   | 2-4 days   |

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|  | <p>Immediately. (Urethral, rectal and throat swabs may also be collected and sent for gonorrhoea culture).</p> <p>For virology investigation send a swab in virus transport medium</p>  |   |  |
| Chlamydia (MIF)<br>Serology  | Clotted blood   |   | 3-5 days   |
| Chlamydia &<br>Gonorrhoea eye<br>swabs   | For investigation of <i>C.trachomatis</i> Infection/Gonorrhoea in the eye, send a swab from the conjunctiva in a Roche Cobas UNI swabs collection tube.   |   | 3-5 days   |
| <p>Clostridium<br/>difficile toxin*</p> <p><i>C.difficile</i> PCR*</p> <p><b>*See <a href="#">Appendix 1</a> for interpretation of results</b></p> | <p>Detection of Clostridium difficile cytotoxin in faeces of patients with antibiotic associated diarrhoea, antibiotic-associated colitis, or Pseudomembranous colitis.</p> <p>Transfer 1 gram (large pea-size) portion of faeces, or 1-2 ml volume of liquid specimen, into a sterile universal container.</p> <p>Only diarrhoeal stools will be tested.</p> |   | 24 hours   |
| CMV IgM  | Clotted blood   |   | 1-4 days   |
| CMV IgG<br>CMV IgG avidity   | Clotted blood   |   | 1-4 days   |
| CMV PCR  | <p>EDTA Blood preferred.</p> <p>Sputum,<br/>Placenta<br/>Urine<br/>Amniotic fluid<br/>Tissue</p>  | 2 | 2-5 days   |
| Corneal scrapes  | Corneal scrape kits (containing a bijoux of broth and a glass slide). Please indicate on the slide which side has been inoculated using a pencil to write on the frosted area. Kits can be obtained from Microbiology during 9am – 4pm Monday to Friday.  |   | <p>2-7 days</p> <p>Urgent Gram stain 1 hour from receipt</p> |
| Covid PCR  | <p>Nose/Throat swabs in viral transport media (both swabs in same tube)</p> <p>Nasopharyngeal aspirate</p>  | 9 | Cepheid<br>1-3 hours   |

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|  | NB testing performed 24/7 at CoCH.  |   |  |
| Covid antibodies<br>IgM and IgG<br>combined  | Clotted blood   |   | 5 days   |
| Coxsackie B virus<br>serology – no longer<br>available, but serum<br>may be tested for<br>enteroviruses by PCR   | Clotted blood<br>EDTA Blood   | 2 | 3-5 days   |
| Culture for bacterial<br>infections  | Pus is the ideal specimen or a biopsy of<br>the infected tissue. Send in a sterile<br>universal container. If only a small sample<br>of tissue is available, add a few drops of<br>sterile normal saline to prevent drying. If<br>swabs are taken, use black topped<br>Microbiology (Charcoal) swab or blue<br>topped Microbiology swab (Transtube) –<br>refer to Wound and Ulcer Swabs |   | 2-5 days   |
| Creutzfeldt-Jakob<br>Disease (CJD)<br><br>ONLY IN<br>CONSULTATION<br>WITH FIRSTLY THE<br>NATION CJD<br>REFERENCE<br>UNIT (0131 537<br>2128) and<br>SECONDLY<br>MEDICAL<br>MICROBIOLOGY<br>CONSULTANT | >1ml CSF, only accepted if <150 RBC on<br>microscopy. CSF sent to lab for cell count.<br>Lab will freeze at -80°C and send via<br>courier to:<br><br>The National Creutzfeldt-Jakob Disease<br>Research & Surveillance Unit<br>Western General Hospital<br>Crewe Road<br>Edinburgh EH4 2XU<br><br>See <a href="http://www.cjd.ed.ac.uk">http://www.cjd.ed.ac.uk</a> for Contacts        |   | 1-2 weeks  |
| Cryptococcus<br>antigen testing  | Clotted blood<br>>1ml CSF in a sterile universal container  | 2 | 2-3 days   |
| Cerebro Spinal Fluid<br>(CSF)  | <ul style="list-style-type: none"> <li>Bacterial Meningitis &gt;1ml CSF in a<br/>sterile universal container</li> <li>Viral Meningitis/Encephalitis &gt;1ml<br/>CSF in a sterile universal container</li> <li>Sub Arachnoid Haemorrhage<br/>please send the first and third</li> </ul>  | 2 | 48 hours<br>Urgent<br>Microscopy<br>1 hour from<br>receipt<br><br>3 days<br><br>48 hours<br>Urgent<br>Microscopy |



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|  | specimen >1ml CSF in a sterile universal container<br><br><ul style="list-style-type: none"> <li>• Mycobacterial Meningitis 6ml CSF in a sterile universal container</li> <li>• PCR Screen: &gt;1ml CSF in a sterile universal container</li> </ul> Preliminary cell counts and Gram Stain<br>Clotted blood results will be telephoned to the sending location as soon as possible after receipt of the specimen and released as preliminary results for viewing on <b>CERNER</b> |   | 1 hour from receipt<br><br>7-14 days<br><br>3 days |
| Delta Antibody (Hepatitis D)   | Clotted blood   | 2 | 5-7 days   |
| Delta PCR (Hepatitis D RNA)  | EDTA Blood  | 1 | 7-10 days  |
| Dengue Virus   | Clotted blood (Antibody)<br>EDTA Blood (PCR)<br>Urine (PCR)   | 3 | 5-7 days   |
| Diphtheria Antibodies  | Clotted blood   | 2 | 1-2 weeks  |
| Ear swab   | Send a swab in black topped Microbiology (Charcoal) swab or in blue topped Microbiology swab (Transtube)  |   | 2-4 days   |
| Early morning urine for tuberculosis   | First catch urine in the morning collect in a sterile universal container, must send 3 consecutive samples  |   | 6-8 weeks  |
| Enterobius vermicularis (Threadworm)   | With a plain swab, moistened with sterile saline, wipe firmly around the anal margin and place the swab into a universal container  |   | 24 hours   |
| Epstein-Barr virus serology – detection of EBV VCA IgG, VCA IgM, EBNA antibodies | Clotted blood   |   | 1-4 days   |
| Epstein Barr Virus PCR   | EDTA Blood<br>CSF<br>Tissue<br>Throat swab in VTM   | 2 | 2-5 days   |



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| Eye swab   | <p>Send a swab in black topped Microbiology (Charcoal) swab or blue topped Microbiology swab (Transtube)</p> <p>For detection of <i>Chlamydia trachomatis</i> and/or <i>Neisseria gonorrhoea</i> by PCR refer to Sexual Health Screening</p> <p>Send a swab in virus transport medium for virology if required.</p> |    | 2-4 days  |
| Enterovirus PCR  | <p>Clotted blood</p> <p>EDTA Blood</p> <p>CSF</p> <p>Vesicle swab in VTM</p> <p>Throat swab in VTM</p> <p>Faeces</p>  | 2  | 2-5 days  |
| Farmers Lung   | <p>Clotted blood</p> <p>(Immunology in Chester)</p>   |    | 5-7 days  |
| <p>Faeces PCR<br/>(VTEC E.coli, Shigella, Salmonella, Campylobacter, Cryptosporidia, Giardia)</p> <p>Faeces culture</p> <p>Ova, cysts and parasites.</p> | <p>Transfer 1 gram (large pea-size) portion of faeces, or 1-2 ml volume of liquid specimen, into a sterile universal container</p> <p>E.g. to confirm positive PCR results or clearance of food handlers</p> <p>Please indicate any foreign travel / country of travel</p>  |    | 2-3 days  |
| Filariasis Antibodies  | Clotted blood   | 6  | 5-7 days  |
| Galactomannan<br>(See Aspergillus antigen above)   |   | 10 |           |
| Glucan ( $\beta$ -Dglucan)   | <p>Invasive fungal infection</p> <p>Clotted blood</p>   | 10 | 2-5 days  |
| Haemophilus Influenzae B (HIB) Antibodies  | Clotted blood   | 2  | 2-3 weeks |
| HIB PCR  | <p>EDTA Blood</p> <p>CSF</p>  | 2  | 5-7 days  |

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| Helicobacter Pylori<br>Stool Antigen | <p>Transfer 1 gram (large pea-size) portion of faeces, or 1-2 ml volume of liquid specimen, into a sterile universal container</p> <p>The test must be carried out within 72 hours of taking the specimen.</p> <p>PLEASE ENSURE THE PATIENT RECORDS THE DATE AND TIME OF COLLECTION ON THE FORM/ SAMPLE</p> |   | 1-3 days  |
| Hepatitis A IgM                      | <p>Serology: detection of IgM antibody in jaundiced patients</p> <p>Clotted blood</p> <p>Positive serology may indicate recent infection. Confirmation will follow – HAV PCR on serum</p>   |   | 1-4 days  |
| Hepatitis A IgG                      | <p>Immune status assessment: IgG serum antibody, when considering active or passive immunisation</p> <p>Clotted blood</p>   |   | 1-4 days  |
| Hepatitis B infection                | <p>Serology: detection of surface antigen and core IgM antibody in jaundiced patient</p> <p>Clotted blood</p>   |   | <p>1-4 days</p> <p>Urgent 1-3 hours from receipt (if received between 9:00 – 17:30)</p> |
| Hepatitis B PCR                      | <p>Quantitative PCR: To indicate viral load</p> <p>Sequencing for genotype and antiviral Resistance</p> <p>EDTA Blood</p> <p>If EDTA not available use serum</p>  | 2 | 1-4 days  |
| Hepatitis B e Antibody/Antigen       | Clotted blood   |   | 1-4 days  |
| Hepatitis B immunity                 | <p>Immune status: Assessment of surface antibody, or to verify sero-conversion, following vaccination</p> <p>Clotted blood</p>  |   | 1-4 days  |

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| Hepatitis B Surface Antigen        | Clotted blood   |   | 1-4 days<br><br>Urgent<br>1-3 hours<br>(if received between 9:00 – 17:30) |
| Hepatitis C screen                 | Serology, detection of viral antibody<br><br>Clotted blood  |   | 1-4 days<br><br>Urgent<br>1-3 hours<br>(if received between 9:00 – 17:30) |
| Hepatitis C PCR/ Genotype          | Quantitative PCR: To indicate viral load<br><br>Qualitative PCR for genotype<br><br>EDTA Blood<br>If EDTA not available use serum   | 2 | 1-4 days  |
| Hepatitis E IgM Antibody           | Clotted blood   | 2 | 3-5 days  |
| High Vaginal swab (HVS)            | Send a swab in black topped Microbiology (Charcoal) swab or blue topped Microbiology swab (Transtube)<br><br>For suspected PID or detection of <i>Chlamydia trachomatis</i> , <i>Neisseria gonorrhoea</i> or <i>Trichomonas vaginalis</i> by PCR refer to Sexual Health Screening |   | 2-4 days  |
| HIV 1 & 2 Antibody and P24 Antigen | Clotted blood   |   | 1-4 days<br><br>Urgent<br>1-3 hours<br>(if received between 9:00 – 17:30) |
| HIV Viral Load                     | EDTA blood<br><br>(For neonates pro-viral DNA may be requested-whole blood on EDTA, must not be separated)  | 2 | 2-5 days  |
| HIV Resistance                     | Characterising the genotype of the HIV  | 2 | 7-10 days   |

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| Testing   | virus and enhancing this by matching to the database of phenotypes for the HIV gives a virtual phenotype of the HIV virus. This leads to an understanding of the resistance mechanisms that might be present.<br><br>EDTA blood |   |          |
| HSV 1 & 2 Antibody  | Clotted blood<br><br>Both HSV total antibody and type specific antibody available.  | 2 | 2-5 days |
| HSV 1 & 2 PCR   | >1ml CSF<br>Neonatal blood<br>Vesicles<br>Respiratory samples<br>EDTA blood tube  | 2 | 1-4 days |
| HTLV I/II Antibody  | Clotted blood   |   | 2-5 days |
| HTLV-1 PCR  | EDTA Blood  | 1 | 5-7 days |
| Human Herpes Virus 6 PCR  | EDTA Blood<br>CSF   | 2 | 5-7 days |
| Human Herpes Virus 7 PCR  | EDTA Blood<br>CSF   | 2 | 5-7 days |
| Human Herpes Virus 8 PCR  | EDTA Blood<br>Tissue<br>Fluid (effusion)  | 1 | 5-7 days |
| Hydatid, Malaria, Schistosoma and Amoeba antibody tests                   | Clotted blood   | 6 | 5-7 days |
| Infection Control screen (MRSA screen)                                    | Nose and groin swabs<br><br>Send a swab in black topped Microbiology (Charcoal) swab or blue topped Microbiology swab (Transtube)   |   | 1-4 days |
| Infection Control screen (Resistant Gram Negative Organisms (CPE) screen) | Rectal swab<br><br>Send a swab in black topped Microbiology (Charcoal) swab or blue topped Microbiology swab (Transtube)  |   | 1-4 days |

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| Infection Control screen (VRE screen)   | Rectal swab<br><br>Send a swab in black topped Microbiology (Charcoal) swab or blue topped Microbiology swab (Transtube)   |   | 2-4 days   |
| Infection Control Screen rapid molecular testing (Cepheid)                    | CPE PCR:<br>Red top dual swab (rectal swab) (APH only)<br><br>MRSA PCR:<br>Pink top liquid swab (nose & groin swabs) (Chester-ITU only)<br><br>VRE PCR:<br>Red top dual swab (rectal swab) (Chester-ITU only)  |   | 24 hours<br><br>2-3 hours (if received between 9:00 – 17:30) |
| Influenza Virus A/B PCR<br><br>+<br><br>Respiratory Syncytial Virus (RSV) PCR | Nose/Throat swabs in viral transport media (both swabs in same tube)<br><br>Nasopharyngeal aspirate<br><br>NB testing performed 24/7 at CoCH   |   | 24 hours<br><br>1-3 hours (if received between 9:00 – 17:30) |
| Intrauterine contraceptive device (IUCD)                                      | Send the device in a sterile universal Container.<br><b>NB.</b> IUCDs are not routinely processed and will only be processed if clinical details state there is a suspicion of infection.<br><br>Culture for <i>Actinomyces sp</i>   |   | 2-4 days<br><br>10-12 days                                   |
| Joint Fluids  | >1ml in a sterile universal container + lithium heparin tube for 'hot' joints ('hot joint' packs available from specimen reception at WUTH only)<br><br>If the specimen is urgent preliminary cell count and gram stain will be telephoned to the sending location as soon as possible after receipt of the specimen and preliminary report released electronically. |   | 2-5 days<br><br>Urgent Microscopy<br>1-2 hours from receipt  |
| Legionella Antigen (detects <i>Legionella pneumophila</i> Serogroup 1 only)   | >3ml Urine in a sterile universal  |   | Same day as receipt  |
| Leptospira Antibody   | Clotted blood  | 3 | 5-7 days   |

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| Leptospira PCR   | EDTA Blood<br>Urine  |       | 5-7 days  |
| Lyme Disease<br>(Borrelia burgdorferi)<br>Antibody   | Clotted blood  |       | 1-4 days  |
| Malaria Antibody   | Clotted blood  | 6     | 5-7 days  |
| Measles IgM Antibody   | Clotted blood  | 2     | 2-4 days  |
| Measles IgG Antibody   | Clotted blood  |       | 5-7 days<br><br>Urgent<br>1-3 hours<br>from receipt |
| Measles PCR<br><br>Please inform lab<br>about sample<br>dispatch   | >1ml CSF<br>EDTA blood   | 2     | 1-4 days  |
| Meningococcal<br>Antibodies (vaccine<br>Functional antibodies:<br>Serological tests<br>for specific antibodies<br>after vaccination) | Clotted blood  | 2     | 2-3 weeks   |
| Meningococcal &<br>Pneumococcal PCR<br><br>If urgent, please<br>inform lab about<br>sample dispatch                                  | CSF<br><br>EDTA Blood  | 2     | 1-4 days<br><br>Urgent 1 day                        |
| Mouth swab   | Send a swab in Blue topped Microbiology<br>swab (Transtube) or Black topped<br>Microbiology (Charcoal) swab<br><br>For virology send the swab in a virus<br>transport medium |       | 2-4 days<br><br>1-4 days                            |
| Mumps IgG Antibody   | Testing mumps immunity/past infection<br><br>Clotted blood   |       | 5-7 days  |
| Monkeypox PCR  | Diagnosis of monkeypox virus<br>Viral swab of vesicles and placed in viral<br>transport media  | 1 & 8 | 7 days  |

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| Mumps IgM Antibody                 | Diagnosis of recent mumps<br><br>Clotted blood  |   | 5-7 days                                       |
| Mycobacteria                       | See TB ( <i>Mycobacterium tuberculosis</i> / other Mycobacteria)  | 2 |  |
| Mycology culture                   | For skin, nail and hair clippings use black card, Dermapak or sterile universal.<br><br>For investigation of Candida infections in superficial wound/ENT sites, send a swab in Blue topped Microbiology swab (Transtube) or Black topped Microbiology (Charcoal) swab, request routine culture <b>including</b> Candida |   | 2-4 weeks<br><br>2-4 days                      |
| Mycology PCR                       | For information on: <ul style="list-style-type: none"> <li>Aspergillus antigen (Galactomannan)</li> <li>Aspergillus PCR</li> <li>Glucan (<math>\beta</math>-Dglucan)</li> </ul> refer to individual tests in this table.<br><br>For panfungal PCR/ 18s – contact the Consultant Microbiologist                          |   |  |
| Mycoplasma pneumoniae PCR          | Respiratory sample (Sputum / BAL / NPA)   | 2 | 5-7 days                                       |
| Mycoplasma pneumoniae IgM Antibody | Clotted blood   |   | 3-5 days                                       |
| Nasal swab                         | Send a swab in Blue topped Microbiology swab (Transtube) or Black topped Microbiology (Charcoal) swab<br><br>For virology, send a plastic shafted dacron swab in virus transport medium   |   | 2-4 days<br><br>3-5 days                       |
| Norovirus PCR                      | Only performed on unformed stools (Bristol stool chart 6 & 7) from in-patients (only performed on WUTH in-patients after Consultation with the Infection Control team)  |   | 24 hours<br><br>Urgent<br>4 hours from receipt |

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|  | <p>Or after discussion with Consultant Microbiologist in Immunocompromised patients</p> <p>Transfer 1 gram (large pea-size) portion of faeces, or 1-2 ml volume of liquid specimen, into a sterile universal container</p> |   |   |
| Parvovirus B19 IgG                     | Clotted blood  |   | 1-4 days  |
| Parvovirus B19 IgM                     | Clotted blood  |   | 1-4 days  |
| Parvovirus B19 PCR                     | EDTA blood<br>Clotted blood<br>Amniotic fluid  | 2 | 1-4 days  |
| Pertussis culture (whooping cough)     | Use a thin wired pernasal swab (pale blue top) and transport immediately to the laboratory for pertussis culture/PCR   |   | 5 days  |
| Pleural Fluid                          | Culture and Sensitivity >1ml in a sterile universal container  |   | 2-5 days  |
|  | Tuberculosis culture >1ml in a sterile universal container   |   | 6-8 weeks   |
| Pneumococcal Antibodies (Quantitative) | Clotted blood  | 2 | 2-3 weeks   |
| Pneumococcal Antigen                   | >3ml Urine in a sterile universal  |   | Same day as receipt                                 |
| Pneumocystis jirovecii PCR             | <p>Broncho-alveolar lavage</p> <p>Sputum (ideally induced)</p> <p>If no respiratory samples: EDTA Blood</p> <p>Urgent Request must be discussed with a Consultant Medical Microbiologist</p>                               | 2 | <p>3-5 days</p> <p>Urgent 24 hours from receipt</p> |
| Polio Antibodies                       | Clotted blood  | 1 | 5-7 days  |
| Proviral DNA HIVPCR                    | EDTA blood   | 1 | 5-7 days  |
| Pus                                    | <p>Transfer into a sterile universal container</p> <p>If pus cannot be obtained then send a swab in Blue topped Microbiology</p>   |   | 2-5days   |



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|   | swab (Transtube) or Black topped Microbiology (Charcoal) swab – refer to Wound and Ulcer Swabs  |   | 5-7 days for extended culture                        |
| Rabies  | For any request for Rabies diagnosis, prevention or immunity testing contact Consultant Microbiology  |   |  |
| Respiratory PCR (full panel)<br><br>Urgent screens performed in outbreak situations and by arrangement with Consultant Medical Microbiologist | <p>PCR for:<br/>Influenza A and B, Parainfluenza 1-4, Human Metapneumovirus, Adenovirus, Respiratory Syncytial Virus, Rhinovirus/Enterovirus, MERS, Coronavirus 229E, HKU1, NL63, OC43, SARS-CoV-2, Bordetella pertussis/parapertussis, Chlamydia pneumoniae, Mycoplasma pneumoniae</p> <p>NB. Biofire not verified for immunosuppressed patients.</p> <p>Nasopharyngeal aspirates and Broncho alveolar lavage (send in a sterile universal container)</p> <p>Nose/Throat swabs in viral transport media (both swabs in same tube (can be processed from same sample as flu screen)</p> <p>Swab of nasal secretions or throat swab, send plastic shafted Dacron or rayon swab in viral transport media (swab and media supplies as a pack)</p> <p>Additional virus PCRs can be arranged by request through the Consultant Medical Microbiologists e.g. bocavirus.</p> <p>Supplies of swabs and Viral Transport media can be obtained from Pathology Specimen Reception (In outbreak situations supplies may be placed in other locations)</p> | 2 | 1 day<br><br>Urgent Biofire<br>24 hours from receipt |
| Respiratory Syncytial Virus (RSV) PCR   | Refer to information for Influenza Virus PCR  |   |  |
| Rotavirus antigen   | Detection of rotavirus in faeces. Transfer 1 gram (large pea-size) portion of faeces, or  |   | 24 hours   |

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|   | 1-2 ml volume of liquid specimen, into a sterile universal container  |   |                                 |
| Rickettsia Antibodies   | Clotted blood   | 3 | 5-7 days                        |
| Rubella IgG   | Immunity check (routine antenatal screening no longer routine) or past infection<br><br>Clotted blood   |   | 2-5 days                        |
| Rubella IgM   | IgM antibody: to diagnose recent acute infection in symptomatic or asymptomatic individuals<br><br>Clotted blood  |   | 2-5 days                        |
| Schistosoma serology  | Clotted blood   | 6 | 5-7 days                        |
| Schistosoma detection (from urine)                                      | It is preferable to obtain total urine collected over the time period between 10.00h and 14.00h. Alternatively, a 24h collection of terminal samples of urine may be obtained.  |   | 24 hours                        |
| Screening swabs and surface swabs                                       | Send swab in Blue topped Microbiology swab (Transtube), or Black topped Microbiology (Charcoal) swab  |   | 2-4 days                        |
| Seminal fluid culture   | Sterile universal container   |   | 2-4 days                        |
| Sexual Health Screening<br><br>Chlamydia /<br>Gonorrhoea PCR<br>(CT/NG) | <p><b>Samples from all requesters (excluding Wirral Sexual Health)</b></p> <p>For investigation of <i>C.trachomatis</i>/Gonorrhoea Infection in the eye, send a Roche Uni swab from the conjunctiva in a Roche collection tube.</p> <p>For detection of CT/NG from the throat, vaginal, urethral &amp; anorectal sites use a Roche Cobas <b>Uni</b> swab.</p> <p>For detection in urine first catch urine is required, optimally this should be collected in a Roche urine collection tube.</p> | 9 | <p>3-5 days</p> <p>3-5 days</p> |

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| <i>Trichomonas vaginalis</i> (TV)<br><br><i>Mycoplasma genitalia</i> | Self-taken HVS in a Roche Cobas <b>Uni</b> Swab.<br><br>Not routinely available – contact Consultant Microbiologist for advice  | 9 | 3-5 days   |
| Sputum culture   | Sputum from deep expectoration and not saliva is required. Send specimen in a 30ml sputum container or universal  |   | 2-4 days<br><br>(5-7 days if fungal culture also required) |
| Strongyloides serology   | Clotted blood   | 6 | 5-7 days   |
| Syphilis serology  | A combination of non-Treponema and specific Treponema antibody screening tests for Treponema pallidum<br><br>Clotted blood  |   | 1-4 days   |
| Syphilis PCR   | Virology swab in VTM:<br>Ulcer<br>Mouth Swab  | 2 | 3-5 days   |
| Tetanus Antibodies (Quantitative)                                    | Clotted blood   | 1 | 4 weeks  |
| Throat swab  | Send a swab in Blue topped Microbiology swab (Transtube), or Black topped Microbiology (Charcoal) swab<br><br>For virology send the swab in virus transport medium (refer to respiratory PCR) |   | 2-4 days<br><br>1-3 days                                   |
| Tissue / bone / cartilage and biopsies                               | Sterile universal container<br><br>If biopsy is small add 0.5ml of sterile saline to prevent it from drying out. Ensure   |   | 2-5 days<br><br>5-7 days for extended culture              |

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|   | <p>there is NO formalin or other preservative in samples for culture: samples in formalin can still be used for bacterial, fungal and viral PCRs.</p> <p>(Depending on the nature of the tissue / bone / cartilage enrichment and extended culture maybe required which will take up to 7 days)</p>  |   |  |
| TORCH Screen  | Clotted blood  |   | 3-5 days   |
| Toxocara Antibodies   | Clotted blood  | 6 | 5-7 days   |
| Toxoplasma Antibodies   | Clotted blood  | 7 | 2-4 days   |
| Toxoplasma PCR  | EDTA Blood<br>Amniotic fluid<br>Tissue   | 2 | 1-4 days   |
| TB ( <i>Mycobacterium tuberculosis</i> ) / other Mycobacteria | <p>Recommended specimens are sputum, urine, pus or tissue. For sputum and urine send 3 early morning specimens taken on consecutive days</p> <p>Blood culture bottles available to send out to wards for direct inoculation of blood or bone marrow for TB culture, please contact the Laboratory for special bottle (inoculate ONE bottle a day for 3 consecutive days)</p> | 2 | <p>6-8 weeks</p> <p>Microscopy<br/>2 days</p> <p>6-8 weeks</p> |
| Urethral swab   | <p>For the investigation of gonorrhoea use a swab (orange topped Microbiology swab) transport to the laboratory immediately.</p> <p>For detection of <i>Chlamydia trachomatis</i> and/or <i>Neisseria gonorrhoeae</i> by PCR refer to Sexual Health Screening</p> <p>Do NOT order urethral swabs for MRSA culture (order MRSA screen and add urethra as the site)</p>        |   | 2-4 days   |
| Urine   | Collect mid-stream of urine in a red capped (boric acid) sterile universal container (>3ml)  |   |  |

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|  | <p>Very small samples from paediatric patients only may be collected into a 20ml white capped sterile universal (white capped samples <b>MUST</b> be received in the lab <b>within 4 hours</b> of collection).</p> <p>Negative urine</p> <p>Positive urine (1 organism isolated)<br/>Positive urine (2 organisms isolated)</p> <p>For detection of <i>Chlamydia trachomatis</i> and/or <i>Neisseria gonorrhoeae</i> by PCR refer to Sexual Health Screening</p> |   | <p>24 hours</p> <p>2-4 days<br/>4-6 days</p>                                       |
| Urinary Legionella antigen & Urinary Pneumococcal Antigen                                  | Collect mid-stream of urine in a sterile universal container (>3ml)   |   | Same day as receipt  |
| Varicella zoster IgG Antibodies  | <p>Varicella zoster IgG. Positive result indicates infection with VZV at some time or vaccination</p> <p>Antibody 'negative' high risk groups with exposure &lt;10 days can be offered human zoster immunoglobulin for pregnant women or immunocompromised patients if level less than 150iu/l.</p> <p>Clotted blood</p>  |   | <p>2-4 days</p> <p>Urgent<br/>1-3 hours from receipt (provisional result only)</p> |
| Varicella zoster IgM Antibodies  | Clotted blood   |   | 3-5 days   |
| Varicella zoster PCR   | EDTA blood<br>CSF<br>Lesions<br>Vesicle fluid   | 2 | 2-4 days   |
| VDRL   | Refer to Syphilis serology  |   |  |
| Vesicles, ulcers and genital lesions   | Send a swab in virus transport medium for PCR   | 2 | 2-4 days   |
| Wound and ulcer swabs  | Send a swab in Blue topped Microbiology swab (Transtube), or Black topped Microbiology (Charcoal) swab  |   | 2-4 days   |
| Yersinia antibody tests (reference lab test- availability limited so needs discussion with | Clotted blood   | 1 | 5-7 days   |

|                               |  |  |  |
|-------------------------------|--|--|--|
| Consultant<br>Microbiologist) |  |  |  |
|-------------------------------|--|--|--|

**Specimens should be transported to the Laboratory as soon as possible after they are taken, even overnight so they can be placed in the correct storage conditions**

### \*Reference Lab Referrals

Tests are referred to a number of outside Laboratories. These are listed below:

1. UK Health Security Agency (UKHSA) Colindale, 61 Colindale Avenue, London NW9 5HT
2. UK Health Security Agency (UKHSA) Manchester Laboratory, Manchester Royal Infirmary, Oxford Road, Manchester. M13 9WL
3. UK Health Security Agency (UKHSA) Porton Down, Salisbury, Wiltshire. SP4 0JG
4. UKHSA Mycology Reference Laboratory, National Infection Services, UKHSA Southwest Laboratory, Science Quarter, Southmead Hospital, Bristol, BS10 5NB
5. UK Anaerobe Reference Laboratory, Public Health Wales Microbiology Cardiff, University Hospital of Wales, Heath Park Cardiff. CF14 4XW
6. Liverpool School of Tropical Medicine, Pembroke Place, Liverpool L3 5QA
7. Toxoplasma Reference Laboratory, Department of Microbiology, Singleton Hospital, Sgeti, Swansea SA2 8QA
8. Brucella Special Diagnostic Unit, Virology Department, Liverpool Clinical Laboratories, Mount Vernon Street Liverpool L7 8YE
9. Virology Department, Liverpool Clinical Laboratories, Mount Vernon Street Liverpool L7 8YE
10. Mycology Reference Centre Manchester, 2nd Floor Laboratory, Education and Research Centre, Wythenshawe Hospital, Southmoor Road, Manchester. M23 9LT

## 11.0 KEY FACTORS WHICH AFFECT THE PERFORMANCE AND OR RESULT OF A MICROBIOLOGY TEST

- The technical competency, bias and experience of the staff performing the test.
- The patient sample, how it is taken, stored and transported to the laboratory.
- The homogeneity of the patient sample i.e., is there an even distribution of micro-organisms within the sample?
- Dilutions, how they are performed, what volume is used and how accurate is the equipment used to perform the dilution.
- Media and reagents, if they are not stored correctly, used in the correct way, expired and are not sensitive and specific enough they will have a detrimental effect on the result.
- Inoculation of media, volume of inoculum, media selection and spreading of inoculum will affect the result.
- Incubation conditions such as duration, temperature and humidity.
- Reading and interpretation of results.
- The uncertainty of measurement (UoM) is a quantitative indication of the quality of the result and how reliable and reproducible it is. Reports for UoM are generated every 12 months. All UoM reports are reviewed by the Consultant Microbiologists for clinical impact. If the reports indicate the results may have a clinical impact the users will be informed accordingly. Users can request to view the UoM information.
- If a change is made to an examination procedure which could affect the interpretation of a result, this will be communicated to users.



## CONTAINERS APPROPRIATE FOR TRANSPORT OF SPECIMENS FOR MICROBIOLOGICAL INVESTIGATIONS

| SPECIMEN  | CONTAINERS  |
|---|---|
| Clotted blood for serology:   | Wirral: 4ml OCHRE capped, clear plastic blood tube<br>Chester: 8ml clotted blood (Red or Gold top) clear plastic tube   |
| Blood for PCR (Viral DNA/RNA)   | Wirral: 4ml LAVENDER capped clear plastic EDTA blood tube.<br>Chester: 8ml EDTA (Purple top)  |
| Blood Cultures  | Wirral: Green (O <sub>2</sub> ) and purple (ANO <sub>2</sub> ) bottles<br>Chester: Blue (O <sub>2</sub> ) and purple (ANO <sub>2</sub> ) bottles<br>Paediatric (both sites): Single yellow bottle |
| Urine for MC&S and urinary antigens   | Red capped boric acid container<br>20ml universal bottle (white cap) – <b>Paediatric urine only where a small volume is collected</b>   |
| Body fluids eg joint fluid, CSF, pus  | 20ml universal bottle (white cap)   |
| For faeces, pus, tissue / bone / cartilage and other semisolid specimens                      | 50ml wide-mouth container (Wirral)<br>Blue 30ml container with spoon (Chester)  |
| For the collection and transport of specimens, when <i>Chlamydia trachomatis</i> is suspected | <b>All requestors including Wirral Sexual Health</b><br>Roche Cobas Uni Swab/urine collection tube  |
| For the aerobic collection of small amounts of fluids, or exudates                            | Standard cotton wool-tipped rigid stem swab (blue top) & transport medium<br>or<br>Standard cotton wool-tipped rigid stem swab (black top) & charcoal transport medium                            |
| For sampling ear, nose or throat, or urethral discharge                                       | Special cotton wool-tipped fine rigid wire swab (orange top) & transport medium   |
| For sampling post-nasal space   | Special cotton wood-tipped fine flexible wire swab (sky blue top)   |
| For the anaerobic collection of pus, or exudates  | Sterile universal container   |
| For the collection and transport of swabs for virus tissue culture/PCR                        | Use viral collection kit (2 female swabs and viral transport media)   |
| For the collection and transport of urine for cytomegalovirus (CMV) tissue culture            | Early morning urine in 20ml universal bottle (white cap)  |

## • REFERENCE RANGES

| SPECIMEN                     | REFERENCE RANGE   | COMMENT   |
|------------------------------|---|---|
| URINE                        | White Blood cells <10x10 <sup>6</sup> /L<br>Red Blood cells <17x10 <sup>6</sup> /L  | Normal reference range<br>WBCs <10 RBCs <17 (x10 <sup>6</sup> /L)<br>Pyuria is defined as WBC >10 x10 <sup>6</sup> /L<br>WBC >100 x10 <sup>6</sup> /L is considered more suggestive of infection<br><br>The urinalyser does not reliably detect or measure microscopic haematuria. Please correlate clinically. |
| CSF                          | <b>Neonates/Newborns:</b>   |   |
|                              | <b>White Blood cells</b><br>Neonates<br>(<28 days old including pre-term)<br>0 - 30 (x10 <sup>6</sup> /L)<br><br><b>Red Blood cells</b><br>0 - 675 cells x 10 <sup>6</sup> /L                                   | Normal reference range<br>Preterm <28 days old - WBC 0-30 (x10 <sup>6</sup> /L)<br><br>(A WBC:RBC ratio of 1:500 to 1:1000 is suggestive of a traumatic tap)  |
|                              | <b>Children and adults:</b>   |   |
|                              | <b>White Blood cells</b><br>Infants 1-12 mths<br>0 - 15 (x10 <sup>6</sup> /L)<br><br>Children/Adults (1+ yrs)<br>0 - 5 (x10 <sup>6</sup> /L)<br><br><b>Red Blood Cells</b><br>0 - 10 cells x 10 <sup>6</sup> /L | Normal reference range<br>Infants 1-12 months<br>WBCs 0-15 (x10 <sup>6</sup> /L)<br><br>Normal reference range<br>Children/Adults (1+ yrs)<br>WBCs 0-5 (x10 <sup>6</sup> /L)<br><br>(A WBC:RBC ratio of 1:500 to 1:1000 is suggestive of a traumatic tap.)  |
| Anti Streptolysin-O antibody | <b>Adult:</b> 0 – 200 IU/MI<br><b>Children &lt;4yrs:</b> 0-100 IU/MI  |   |
| β-D-Glucan                   | <b>Measurement range 6-500-pg/ml</b><br><br><6 pg/ml Negative (N)<br><br>≥7 pg/ml<br>Positive (P)   | ≥6.00 ≤6.99 pg/ml - Indeterminate (I)   |

|  |  |
|--|--|
| <b>Hepatitis B immunity (anti-HBs)</b> | <p>An antibody level below 10m IU/ml is classified as non-immune (a non-response to vaccine).</p> <p>Responders with anti-HBs levels of 10 to 100m IU/ml should receive one additional dose of vaccine at that time.</p> <p>Responders with anti-HBs levels greater than or equal to 100m IU/ml do not require any further primary</p> |
|--|--|

## • APPENDIX 1

### Interpretation of *Clostridium difficile* testing results

CWMS microbiology laboratory combines two different tests to optimize sensitivity and specificity of *C. difficile* testing. The polymerase chain reaction (PCR) test (detects the genes encoding the *C. difficile* toxins) is followed by a toxin EIA test.

If the PCR is positive, it implies that the patient has *C. difficile* in their gastrointestinal tract, with the capability to produce toxins. However, this method cannot determine if the *C. difficile* strain actually produces toxins. The EIA toxin test is suitable for the detection of the *C. difficile* toxins in the stool specimens. Please note, according to National Guidance the *C. difficile* toxin EIA test is not suitable as a stand-alone test for the diagnosis of *C. difficile* infection because of low sensitivity (that results in a high rate of false negatives).

Reporting and interpretation of *C. difficile* results:

#### 1. Toxin gene DETECTED/toxin production DETECTED

Both the PCR and toxin EIA tests are positive. In symptomatic patients this result confirms the diagnosis of *C. difficile* infection. Appropriate antibiotic treatment and infection control precautions are required.

This will be reported as:

*C.difficile* toxin gene detected by PCR

*C.difficile* toxin production detected by EIA

Positive for *C.difficile* toxin, likely to be significant in the appropriate clinical context.

#### 2. Toxin gene DETECTED/toxin production NOT detected

The PCR test is positive, that means the patient carries toxigenic *C.difficile* strains in the gastrointestinal tract. However, the EIA toxin test remained negative.

The patient might be an asymptomatic carrier (in which case the toxin test is true negative), or the patient has toxigenic *C.difficile* infection with a false negative EIA test.

Clinical review and correlation are required. If the patient is symptomatic, appropriate treatment and infection control precautions are required.

This will be reported as:

*C.difficile* toxin gene detected by PCR

*C.difficile* toxin production NOT detected by EIA

Interpretation: *C.difficile* bacteria detected. Potential for

*C.difficile* toxin to be produced but not detectable at

Present. Clinical correlation required to determine whether

*C.difficile* infection causing symptoms or *C.difficile*

colonisation with another cause of the symptoms.

#### 3. *Clostridium difficile* NOT detected

The PCR test is negative. Alternative diagnosis should be considered. If high suspicion of *C. difficile* infection, sending a repeat specimen can be considered.

This will be reported as: Toxigenic *Clostridioides difficile* NOT detected