# Microbiology Services User Handbook





Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 1 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# A – Z of tests A B C D E F G H I J K L M N O P Q R S T U V W X Y Z

0 0	ontents			
0	Contents	2		
1	INTRODUCTION	4		
2	LABORATORY LOCATION	5		
3	PATHOLOGY QUALITY POLICY	6		
4	OPENING HOURS, CLINICAL ADVICE AND RESULTS	7		
4.1	Laboratory Opening Hours	7		
4.2	Clinical advice	7		
4.3	Urgent samples	7		
4.4	Testing out of hours	7		
4.5	Additional tests	8		
4.6	Results	8		
4.7	Telephoned results	9		
4.8	Turnaround times	9		
4.9	Tests currently in scope of UKAS accreditation	9		
5	CONTACT DETAILS	10		
6	SAMPLE COLLECTION10			
6.1	Preparation of patient			
6.2	Optimum time of and conditions for collection			
6.3	Health and safety issues pertaining to sample collection			
7	SAMPLE CONTAINERS	12		
7.1	Supply of specimen containers			
7.2	Selection of appropriate container	13		
7.3	Labelling of sample containers	13		
8	REQUEST FORMS	14		
8.1	Electronic requesting (ICE)			
8.2	Handwritten request forms	14		
8.3	Anonymous/uniquely identified samples	15		
8.4	Microbiology department request forms	16		
9	TRANSPORTATION OF SAMPLES	17		
9.1	Transportation of routine samples to the laboratory	17		
9.2	Transportation of urgent samples			
10	HIGH RISK SAMPLES	18		
uthorised	by: C Frearson	DCN: MIC-P-006-12.7		
bate of iss	ıe: 07/02/2023	Page 2 of 192		

11 confirr	Samples from patients categorised as 'high possibility of vhf' and samples from pation med vhf	ents with 19				
12	SAMPLE ACCEPTANCE CRITERIA					
13	REPERTOIRE OF TESTS (A – Z)					
13.1	L Reference Intervals	21				
13.2	2 Referred Tests	21				
13.3	3 Repertoire index	22				
14	REFERENCE LABORATORIES	186				
14 15	REFERENCE LABORATORIES PATIENT CONSENT DISCLOSURE	186 191				
14 15 15.1	REFERENCE LABORATORIES         PATIENT CONSENT DISCLOSURE         L       Laboratory Policy on protection of personal information	<b>186</b> <b>191</b> 191				
14 15 15.1 15.2	<b>REFERENCE LABORATORIES PATIENT CONSENT DISCLOSURE</b> L       Laboratory Policy on protection of personal information         2       Patient consent	<b>186</b> <b>191</b> 191 				
14 15 15.1 15.2 15.3	REFERENCE LABORATORIES         PATIENT CONSENT DISCLOSURE         L       Laboratory Policy on protection of personal information         2       Patient consent         3       Medico-legal samples	<b>186</b> <b>191</b> 191 191 192				
14 15 15.1 15.2 15.3 15.4	REFERENCE LABORATORIES         PATIENT CONSENT DISCLOSURE         Laboratory Policy on protection of personal information         Patient consent         Patient consent         Medico-legal samples         The Human Tissue Act					

# **1** INTRODUCTION

The Clinical Microbiology Service is provided by the laboratories at the Great Western Hospital NHS Foundation Trust, Swindon, providing a formulary of tests reflecting the usual demands of a District General hospital service. Specialist and Reference test services are used where necessary.

Microbiology services are provided on a 24 hour basis, with a routine service available between 09:00 and 17:00 Monday to Friday, 08:00 and 13:00 on Saturday and 08:45 and 12:30 on Sunday and bank holidays. The laboratory provides an on-call bacteriology service outside of these hours. Virology services are provided Monday to Friday 09:00 to 17:00.

Consultant advice is available on-site on an open access basis during normal working hours and on an on-call basis at all other times.

We provide an analytical and interpretative service on a wide-range of clinical samples, processing over 270,000 requests each year. The efficiency of the service we provide is reliant on the cooperation of our users with the necessary policies relating to safety, sample transport and sample identification.

In its pursuit of excellence and as part of its continuous quality improvement programme the Microbiology department participates in all relevant internal and external quality assurance schemes. All laboratory work is carried out on up to date equipment in a modern laboratory which meets with all statutory requirements of a quality management system.

The repertoire of tests provided by Microbiology support the Trust in its diagnostic and screening programmes.

The laboratory is accredited by the Institute of Biomedical Science (IBMS) for Biomedical Scientist training and Biomedical Scientist Specialist training. We also support the University of Bristol in the provision of clinical undergraduate training and the development of junior doctors at Great Western Hospital.

The Pathology services are fully computerised with all laboratories using Clinisys Winpath laboratory information system. Pathology results are available electronically via the Trust network at ward level or via the GP electronics links. Hard copies (if required) are returned daily Monday-Friday.

We hope that this handbook contains all the information you require to use our service. However, please feel free to contact us to discuss any problems or issues you may have. Any comments or suggestions about the User Handbook should be addressed to the Laboratory Manager, by email to Mitchell.Reed@NHS.net.

# 2 LABORATORY LOCATION

The Microbiology Department is part of the Clinical Support and Specialist Services Division, within the Great Western Hospitals NHS Foundation Trust. The department is sited on the fourth floor of the main hospital building (see diagram).



Authorised by: C Frearson Date of issue: 07/02/2023

SN3 6BB

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# **3 PATHOLOGY QUALITY POLICY**

Great Western Hospitals NH	S Foundation Trus	 	PAT-P-012-8.0
<ul> <li>Department of Pathology</li> </ul>		 · · · · · · · · · · · Date of iss	ue: 14/06/2022
	• • • • • • • • • • • • • • • • • • • •		Page 1 of 1

### Quality Policy

The Pathology Department provides Microbiology, Cellular Pathology, Blood Sciences (incorporating Haematology, Biochemistry, Blood Transfusion and Point of Care Testing) and the Mortuary and Bereavement services to the Great Western Hospitals NHS Foundation Trust, Bath & North East Somerset, Swindon and Wiltshire Clinical Commissioning Group (BSW CCG) and other users where such arrangements have been made.

The management of the Pathology Department is committed to delivering a service that is compliant with the requirements for Medical Laboratories set by the International Standard Organisation (UKAS ISO 15189:2012), Health and Safety Executive (HSE), UK Health Security Agency (UKHSA), Medicines and Healthcare Products Regulatory Agency (MHRA) and the Human Tissue Authority (HTA).

The Pathology management team is fully committed to the on-going development and improvement of laboratory services through the continual assessment of the Pathology Quality Management System and the establishment by means of regular meetings, internal and external audits and annual review of quality objectives during the Pathology Annual Management Review.

The management of the Pathology Department is committed to good professional practice and the provision of examinations that are fit for intended use to ensure the delivery of a high quality service that meets the requirements of its users. This commitment is reflected in the core values of the Quality Management System:

- The development of a friendly working environment that supports training and encourages the retention and recruitment of committed, highly professional staff.
- A commitment to maintaining a laboratory environment, in compliance with relevant legislation, to ensure the health, safety and welfare of staff and visitors.
- The provision of information on the collection, transportation and handling of all specimens to ensure the validity of results of laboratory examinations.
- The review of test repertoire, in conjunction with users, to ensure it is fit for intended use.
- The procurement and maintenance of appropriate equipment, reagents and consumables to enable the provision of quality examinations of specimens.
- The reporting of high quality examination results in a timely, confidential, accurate and clinically useful manner.
- The provision of advice, in the context of clinical information, to support patient management.
- The engagement with users (e.g. by use of surveys, meetings, feedback, newsletters) to ensure that the Pathology service continues to meet their needs and requirements.
- Agreeing and monitoring quality indicators designed to improve our services to all our customers.
- To ensure all personnel are familiar with this quality policy and comply with the contents of the quality
  manual and all procedures relevant to their work to ensure user satisfaction.

Melanie Wilson

Dr Melanie Wilson Pathology Clinical Lead 14/06/2022 Norma Manzon

Noman Manzoor Pathology & Transfusion Services General Manager 14/06/2022

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 6 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# **4 OPENING HOURS, CLINICAL ADVICE AND RESULTS**

### 4.1 Laboratory Opening Hours

The laboratory is open:

Monday to Friday: 0900 – 1700

Saturday:0800 - 1300Sunday:0845 - 1230Bank Holidays:0845 - 1230

### 4.2 Clinical advice

Consultation about investigation and management of infections is welcomed. For advice on diagnosis and the interpretation of Microbiology results, use of antimicrobials or infection control (including the use of containment facilities) consultant advice is available on-site on an open access basis during normal working hours and on an on-call basis at all other times.

For advice during normal working hours: Telephone 01793 604798.

For advice out of hours: Telephone 01793 604020 (switchboard) and ask for the duty Consultant Medical Microbiologist.

- Internal users, please refer to the antibiotic guidelines, in the first instance, for the commoner microbiology enquiries. These are available on the intranet at the Antibiotic Home Page.
- New or junior doctors should discuss queries with their own clinical team, before calling the Medical Microbiologist.
- For Medical Microbiology advice for more complicated cases the Medical Microbiology team should be contacted on 01793 604798.
- For Infection Control advice alone, the Infection Control Nurses can be contacted on 01793 604554, or via switchboard.

### 4.3 Urgent samples

If a result is required urgently and the sample will arrive during normal working hours the laboratory MUST be notified by telephone so that we can prioritize the request.

Please ensure that the requesting doctor contact details are provided on the request form to enable the result to be telephoned to the requesting clinician.

### 4.4 Testing out of hours

The on-call service is available outside of normal Laboratory opening hours.

# The Microbiology out of hours service is an urgent service. Urgent samples out-of-hours should not be sent before agreement with the laboratory on-call staff.

Authorised by: C Frearson	
Date of issue: 07/02/2023	

To discuss an urgent sample with the duty Biomedical Scientist: Telephone 01793 604020 (switchboard) and ask for the duty Biomedical Scientist.

The use of the service should be restricted to those samples where it is essential to have a result before the next routine session. In general, samples normally accepted for the on-call service would include:

- Cerebrospinal fluid (CSF)
- Peritoneal dialysis (PD) fluid
- Fluids from sterile sites (joint fluids, pleural fluids, ascitic fluids etc.)
- Pus
- Tissue samples
- Corneal Scrapes
- Urine

### 4.5 Additional tests

All tests should be requested at the time of submitting the sample to the laboratory. However, amendments to requests, or the need for additional tests, can still be discussed with the laboratory after processing has started.

In general, additional tests must be requested within 48 hours of sample receipt within the laboratory. In some cases, additional tests may not be possible and a fresh sample will be required. Further advice can be obtained from the laboratory.

### 4.6 Results

Pathology results are available electronically immediately after authorisation via Medway PAS at ward level or via the GP electronic links. Hard copies of reports are produced and returned daily Monday – Friday.

All laboratory results are returned to the requesting clinician who has ultimate responsibility for ensuring that all results are actioned and communicated to the patient as appropriate.

In cases of difficulty or further clarification, the laboratory enquiry telephone number is 01793 604798.

Please note that we need to establish the caller's identity before giving results over the telephone. We are unable to give results directly to patients or their relatives.

General culture results are available 24 hours after sample receipt (at the earliest), and sensitivities after a further 24 hours. For samples such as blood cultures and CSF, the Medical Microbiologist will usually inform the clinicians of initial significant results as soon as they are known.

In general, results are not available until they have been authorised. In exceptional circumstances, preliminary results may be available direct from the relevant laboratory. However, please bear in mind that this may delay the testing of other samples.

### 4.7 Telephoned results

Results of urgent requests and results which may aid the immediate patient management will be telephoned. This includes all positive blood cultures, positive CSFs and specimens processed on-call.

All other results will only be telephoned (or faxed by arrangement) on request.

### 4.8 Turnaround times

The laboratory continually monitors its turnaround times to ensure that it complies with its responsibilities within the patient pathway. The laboratory measures its turnaround times as the time from receipt until the point at which the result is authorised (at this point the result is available through direct enquiry and is available for transmission via GP links).

The expected turnaround times for each test are indicated on the individual test sheets. For detailed turnaround times for each test and actual performance, please contact the laboratory.

Interrogation of the electronic systems allows for full audit of the reception, testing and reporting process, including time of report viewing and report printing.

### 4.9 Tests currently in scope of UKAS accreditation

The laboratory is a UKAS certified as a medical laboratory and we seek UKAS accreditation for all our tests. However, some tests are currently provided outside the scope of our current UKAS certificate. They are:

Test outside current UKAS accreditation	Additional
Helicobacter Pylori antibody (by manual EIA)	UKAS accreditation being sought
Pneumococcal Antigen (TruPneumo)	UKAS accreditation being sought
Legionella Antigen (TruLegionella)	UKAS accreditation being sought
Faecal Calprotectin (Diasorin Liaison)	UKAS accreditation being sought
Urinalysis by UF5000i	UKAS accreditation being sought
Blood Culture by BD BACTEC FX	UKAS accreditation being sought
Mumps (Diasorin Liaison)	UKAS accreditation being sought
Measles (Diasorin Liaison)	UKAS accreditation being sought
Varicella Zoster (Diasorin Liaison)	UKAS accreditation being sought
SARS COV2 (Hologic Panther)	UKAS accreditation being sought
Xpress SARS COV2 (Cepheid GeneXpert)	UKAS accreditation being sought

All unaccredited test results are provided with an indicator comment reflecting unaccredited status: *Please be aware this procedure is not within the laboratory scope of UKAS accreditation*'

We send some testing to external reference centers and, where possible, we use UKAS accredited laboratories. For details on our external laboratories and their accreditation status please see section 14: REFERENCE LABORATORIES

# **5 CONTACT DETAILS**

Position	External Number	Internal Number
Head of Pathology	01793605488	5488
Consultant Microbiologist, Joint Clinical Lead for Pathology	01793 604801	4801
Consultant Microbiologist, Infection Control Doctor	01793 604802	4802
Consultant Microbiologist	01793 605494	5494
Laboratory Manager	01793 604804	4804
Deputy Laboratory Manager	01793 604800	4800
Bacteriology Enquiries	01793 604798	4798
Virology Enquiries	01793 604799	4798
Laboratory	01793 604798	4798
Fax	01793 604803	4803
Hospital switchboard	01793 604020	0

# 6 SAMPLE COLLECTION

### 6.1 Preparation of patient

Adequate privacy during reception and sampling should be available and appropriate to the type of information being requested and primary sample being collected.

Information for patients regarding tests performed, including instruction for preparation of the patient and instructions for patient-collected samples, can be accessed at the 'Lab Tests Online UK' website.

For details of the Laboratory Policy on protection of personal information, patient consent, medico-legal samples and the Human Tissue Act refer to <u>Patient Consent Disclosure</u>.

### 6.2 Optimum time of and conditions for collection

Samples for bacterial culture, wherever possible, should be collected prior to commencement of antibiotic treatment.

Actual pus or tissue samples are always preferable to a swab.

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 10 of 192
	THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED	

To avoid inadvertent contamination of a specimen during collection, an aseptic technique must be used: use universal precautions at all times, wash hands and wear appropriate personal protective clothing.

Decontamination of the sampling site or equipment may be necessary e.g. skin antisepsis before taking blood cultures or Cerebro-spinal fluid (CSF), or catheter port antisepsis before collecting a specimen of urine via a catheter (CSU).

Specimens must be collected into sterile containers with close fitting lids (refer to <u>Selection of</u> <u>appropriate container</u>). The specimen must be clearly labelled. Once collected, place the specimen into a plastic specimen bag and seal the bag. Wash your hands and dispose of clinical waste into a yellow clinical waste collection bag. Sharps must be disposed of safely.

### 6.3 Health and safety issues pertaining to sample collection

Every clinical specimen sent for microbiology examination should be treated as potentially infectious. Standard precautions must be observed at all times. Use aseptic technique.

With patients known to be infected, or if there is a strong suspicion that they may be infected with a high-risk organism (e.g. tuberculosis), then procedures likely to produce aerosols must be conducted whilst wearing face masks, goggles or full facial visors as appropriate. Such investigations include cough inducing procedures and lancing of an abscess.

Used sharps must be disposed of according to Trust policy (see Safe Handling and Disposal of Sharps Policy & Guidelines). This is the responsibility of the individual(s) who generates them.

It is the responsibility of the person collecting the specimen to ensure that it is properly labelled and safe for transportation (see <u>Transportation of Samples</u>).

Refer to appropriate Trust policies for further information:

- Hand Hygiene and Skin Care Policy (including scrubbing gowning and gloving)
- Standard Infection Control Precautions Policy
- Safe Handling and Disposal of Sharps Policy & Guidelines
- TRANSPORTATION OF SAMPLES

# **7 SAMPLE CONTAINERS**

# 7.1 Supply of specimen containers

#### The following Microbiology consumables can be obtained from the following locations:

Consumable	Description	Issue from
	Green form (non-blood Microbiology requests, excluding Blood Cultures) For locations that do not have access to ICE only	Materials Management Team
	Ref form (blood Microbiology requests) For locations that do not have access to ICE only	Materials Management Team
	Yellow form (MRSA admission screen requests) For locations that do not have access to ICE only	Materials Management Team
	Universal containers (with boric acid) - for urine bacteriology specimens	Materials Management Team
OID Contract Nutra O	Bacteriology swabs in Amies transport swab	Materials Management Team
	Pernasal swab for whooping cough	Microbiology Department
	Virus swabs in virus transport medium	Materials Management Team Microbiology Department
	Faeces container	Materials Management Team
	Universal containers (sterile and empty)	Materials Management Team
	Sputum container	Materials Management Team
	Collection kits for Chlamydia trachomatis	Materials Management Team
	Vacutainer tubes for blood samples	Materials Management Team
	Blood culture bottles Pink = peadiatric (single bottle) Grey (aerobic) and purple (anaerobic) = adult set	Pathology Reception
	Pin worm collection kits	Microbiology Department
	TB Quantiferon gold	Microbiology Department

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 12 of 192

### 7.2 Selection of appropriate container

Please see <u>Repertoire Index</u> for the selection of appropriate container for test.

Sample containers must be CE marked. Specimen containers must be leak proof and be sufficiently robust to withstand stresses during transit. Only containers approved by the Microbiology Department may be used to ensure sample integrity during transit to the Laboratory. Samples that are sent in non-approved containers may not be processed by the Laboratory. It is the responsibility of the person sending the sample to the Laboratory to ensure that the container used for transportation is appropriate.

The container must be adequately closed to avoid leakage. Samples that have leaked in transit may not be processed by the Laboratory.

### 7.3 Labeling of sample containers

Clinical governance requires the sample container to be labelled with sufficient information to provide an unequivocal link with the request form and the patient from whom they are collected.

Pre-printed addressograph labels are acceptable on sample containers for Microbiology investigations.

Minimum Data Set for Identification:

- Patient's surname
- Patient's forename (initial is acceptable)
- Date of birth and /or district number / NHS number

Microbiology sample containers should additionally include type of sample and site of collection.

For antibiotic assay levels, for example Teicoplanin, the following information must be completed on the request form:

- Mg of last dose given
- Date and time of last dose
- Date and time that sample was taken

Failure to comply with correct guidance may result in the sample being rejected by the Microbiology department (refer to <u>Sample Acceptance Criteria</u>).

Multiple samples taken at different times on a patient MUST be labelled on the sample container with the time (24 hr clock) when the sample is taken. The request form should be labelled accordingly.

### 8 **REQUEST FORMS**

All samples must be accompanied by a properly completed request form. Failure to comply with correct guidance may result in the sample being rejected by the Microbiology department (refer to <u>Sample Acceptance Criteria</u>). Acceptance of a testing request by the laboratory acts as an agreement with the requestor. This means that a contract is established between the laboratory and the requester when the laboratory accepts a request. This will apply whether the request is written or electronic.

### 8.1 Electronic requesting (ICE)

Please use electronic requesting (ICE) order-comms where available. It is important to ensure that the correct sample accompanies the correct request form before placing in the inside the sample bag.

Please ensure that you order the correct test and select the correct sample type as failure to do so may lead to incorrect testing. The ICE requesting system will show those tests most commonly requested for the Microbiology Service; should the test you require not be visible please contact the laboratory to check that the test is available.

The information required is the same as that required on a handwritten request form and should include clinical details and symptoms, as well as information on antibiotic use, foreign travel, outbreaks, date of onset, etc.

Where ICE requesting is not available handwritten request forms must be used.

### 8.2 Handwritten request forms

Minimum Data Set for Identification:

- District number and/or NHS number
- Patient surname and forename (in full, not initials)
- Date of birth (DOB)
- Patient address if district number/NHS number not supplied

In addition to the minimum data set for patient identification please ensure all other relevant fields are completed:

- Ward/ Practice, Consultant/GP
- Patient address
- Patient gender
- Date and time of collection
- Specimen type
- Investigation(s) required

- Name of requesting clinician and bleep number
- Relevant clinical details \*
- Current drug therapy
- Copy reports, if required
- Patient category (PP/AQP/NHS)

\* To ensure samples can be safely and appropriately tested in the laboratory, information including details of foreign travel, symptoms and known or suspected contact with other patients known to have communicable disease is important. For example, samples likely to contain high risk pathogens as

described by the Advisory Committee for Dangerous Pathogens (refer to ACDP guidance) are handled at a higher containment level to safeguard both laboratory staff and other downstream workers (refer to <u>High Risk Samples</u>). The information is also of benefit to the patient ensuring that appropriate testing is performed.

Unnecessary confidential patient information, for example HIV, Hepatitis B or C status, should not be disclosed on the request form.

It is essential to use a ballpoint pen when completing request forms. Use of felt tip and fountain pens can lead to delay in processing samples, or requests being missed altogether, as carbon copies are often incomplete. When addressograph labels are used, please ensure that a label is fixed to EACH part of the request form.

### 8.3 Anonymous/uniquely identified samples

In certain circumstances patient identification details are intentionally hidden or substituted with particular ID numbers (for example, Sexual Health, donor samples, samples from unconscious or incoherent patients). In such instances, a properly coded identifier must be used in place of the patient last name and first name.

#### **Unidentified Patients**

Samples from unconscious or incoherent patients should be labelled with "UNKNOWN MALE OR FEMALE" and the emergency unit number.

All request forms must be signed.

#### **GUM Patients**

Where Patient name is not appropriate, then GUM number, patient gender and DOB is acceptable.

### 8.4 Microbiology department request forms

The following request forms are used by the Microbiology department (please do not mix with samples for other departments):

LINE LINE	PATHOLOGY REQUESTS TISSUE/SWABS/FLUIDS ETC.	LABORATORY NUM
	BLOCK LETTERS PLEASE USE BALLPOINT PEN BOXES IN BOLD PRINT MANDATORY	PLEASE SEND SEPARATE REQUEST AND SAMPLE FOR EACH DEPT.
	UNIT NUMBER	TIME & DATE TAKEN BY DATE RECEIVED
	SURNAME	SPECIMEN TYPE:- MICROBIOLOGY:- ANTIBIOTIC THERAPY-
erro HEI OOI	FORENAMES	DATE OF ONSET OF ILLNESS
IER IER •ount	SEX D.O.B. N.H.S. PRIVATE OTHER	
A K A K HIS F	HOSPITAL/CODE REPORT TO:- WARD/DEPT COPY TO	OTHER:-
Y ON A LE CAL	CONSULTANT/G.R/CODE SURNAME (PATIENTS) UNIT NUMBER	HAEMATOLOGY:- BONE MARROW MGG CYTOGENETICS
	PATIENT'S ADDRESS	IRON IMMUNOPHENOTYPING CSF CYTO
E YOU LAB	CLINICAL DETAILS INCLUDING RELEVANT DRUGS AND OPERATIONS	CHEMICAL PATHOLOGY URINE/FAECES/MISC. FLUIDS SPECIFY TESTS:-
REAT CO		HISTOPATHOLOGY/CYTOLOGY:- PREV. HIST. No.
G L R		PREVIOUS HISTOLOGY/CYTOLOGY Y/N PREV. CYT. No.
		PATHOLOGIST DATE PROCESSED BLOCKS
MAN AND	REQUESTING DOCTOR'S NAME (Please Print)	DEPARTMENT OF PATHOLOGY, THE GREAT WESTERN HOSPITAL,

### GREEN FORM (NON-BLOOD MICROBIOLOGY REQUESTS, EXCLUDING BLOOD CULTURES)

### **RED FORM (BLOOD MICROBIOLOGY REQUESTS)**

1208 B	TEAR	PATHOLOGY REQUESTS BLOOD SPECIMENS ONLY	PLEASE PRINT WITH BALL POINT PEN OR AFEIX PRINTED LARELS	GENERAL LAB NUMBER
222		BOXES IN BOLD PRINT MANDATORY	ON ALL FOUR COPIES	
T NO		NHS NUMBER	SEPARATE SPECIMEN IS REQUIRED FOR EACH DISCIPLIN	E
ATEN	D ISI		YELLOW TOP TUBE RED TOP TUBE	
e LI	A TR		CHEMISTRY SEROLOGY	
ORRE	CH L 30C B ATIOI	SURNAME	Glucose     Glucose     Liver Function     Glucose     Glucos	
IEN C	EAC KPH RIE	FORENAMES	Calcium Group Hepatitis B Thyroid HIV	
ECIN	A B A	SEX D.O.B.	Lipid Studies     Other	
E SP	C L	HOSPITAL/CODE REPORT TO:- WARD/DEPT COPY TO	GREEN TOP TUBE	BLACK TOP TUBE
ED TH	ML REA NEN	CONSULTANT/G.P./CODE SURNAME (PATIENTS) UNIT NO	(see reverse) Troponin	
ABELL	FIA SUF SUF SUF SUF	PATIENT'S ADDRESS	HAEMATOLOGY	MAUVE TOP TUBE HAEMATOLOGY
NON	ESS EN SPE	CLINICAL DETAILS INCLUDING RELEVANT DRUGS AND OPERATIONS	AIP     BLUE TOP TUBE     Glandular Fever     CLOTTING STUDIES	Full Blood Count     Other
HAVE	TO TO		Other INR (Warfarin) APTT	
		PATIENT WAITING FOR RESULT	GREY TOP TUBE (Unfractionated Heparie GP GLUGOSE Other	
19180	D427	HIGH INFECTION RISK NO / YES URGENT ROUTINE ROUTINE REQUESTING DOCTOR'S NAME (Please Print)	Glucose	
Ref:	GWH	CONTACTABLE ON BLEEP EXT.		

### YELLOW FORM (MRSA ADMISSION SCREEN REQUESTS)

	1	Pathology Requests		MRSA Admis	sion Screening Form
LTV?	A started	Specimens submitted on this for	orm will ONLY be tested for MF	RSA Date Taken:	Time Taken:
DRRECT		Unit Number		Taken By:	Bleep / Ext:
MEN CO	ORM	Surname		Specimen Types (r	max 4 per form)
SPECIA CH L ROC FR SPECIA		Forename(s)		Туре:	Lab No.:
V EA AKF	RRIE	Sex	DOB		
Y OI	I SCF	Ward	Consultant	Туре:	Lab No.:
RML RE /	MEN SSION	Screen Type (please tick)			
D THE S FI NSU		Elective admission screen		Туре:	Lab No.:
RES TO E	<i>SH</i> RSA /	Emergency admission screen			
	Ī			Туре:	Lab No.:
HAVE	6	For Lab Use Only			
	K		MSCR	Department of Micr Marlborough Road	obiology, The Great Western Hospital, d. Swindon, SN3 6BB (01793) 604798

### 9 TRANSPORTATION OF SAMPLES

Please refer to the Trust Specimen Transportation Policy for the correct procedures for submitting samples to the laboratory.

### 9.1 Transportation of routine samples to the laboratory

All sample containers for transport to the Laboratory must be sealed in a plastic bag attached to the request form.

Samples for microbiological investigation should be examined as soon as possible after collection to avoid compromising results. Samples may be transported via normal portering rounds during the normal working day.

Where this is not practicable due to delays in transportation samples should be kept refrigerated. Samples may be kept in a refrigerator at a temperature of 4-8°C for a maximum of 24 hours prior to transportation. There is a refrigerator in Pathology Reception for non-urgent samples that arrive outside the normal opening hours.

Samples taken for blood culture examination MUST NOT be refrigerated. These must be transported to the Laboratory as soon as possible for incubation at 37°C.

Certain samples may be sent direct to the laboratories using the pneumatic chute system:

Pathology address: 104 Microbiology address: 102

For transportation of samples to the laboratory from external sites or by post, and use of the pneumatic chute system, please refer to the Trust Specimen Transportation Policy.

In cases of difficulty or further clarification, the laboratory enquiry telephone number is 01793 604798.

### 9.2 Transportation of urgent samples

Urgent samples must be sent to the laboratory immediately and arrangements made with the portering service.

To discuss an urgent sample with the duty Biomedical Scientist: telephone 01793 604020 (switchboard) and ask for the duty Biomedical Scientist.

### **10 HIGH RISK SAMPLES**

All samples should be regarded as potentially infectious.

Certain samples from patients who are known or suspected to have the following diseases/conditions constitute a potential higher risk of infection to persons handling the samples:

- Typhoid/paratyphoid fever (faecal samples only)
- Dysentery (faecal samples only)
- Tuberculosis (samples from sites where tuberculosis infection is likely)
- Anthrax
- Brucellosis
- Transmissible Spongiform Encephalopathy (including CJD)
- Viral haemorrhagic fever
- Avian Flu
- MERS/SARS respiratory syndrome, including SARS-CoV2

To minimise the risks ensure that such samples are packaged as follows:

- Attach a "Danger of Infection" label to the sample container and request form for all qualifying samples (available from Phlebotomy Department, GWH)
- Specify the nature of the risk on the request form
- Use unambiguous and commonly recognised terminology
- Place the sample in a sealable plastic bag and close the seal

This is a necessary procedure to safeguard both laboratory staff and other downstream workers. The labels must be used in accordance with the Trust Specimen Transportation Policy

Samples thought to constitute a risk to laboratory staff because of inadequate packaging or warning may be rejected.

The Consultant Microbiologist must be contacted **BEFORE** collecting samples from a patient suspected of having a viral haemorrhagic fever, human avian influenza, MERS/SARS or CJD. These organisms require special transport arrangements and specialist laboratories designed for containment during manipulation of samples and cultures.

These lists are not exhaustive and rarely other biological agents that can cause severe human disease, and present a serious hazard to employees, may be present in samples. If there is any suspicion of a high risk atypical organism advice on sample collection and transport should be sought from the Consultant Microbiologist.

# 11 Samples from patients categorised as 'high possibility of vhf' and samples from patients with confirmed vhf

Instructions for sample transportation of suspected VHF samples are defined in the Trust Specimen Transportation Policy and are formulated in line with current ACDP guidance.

The laboratory MUST be notified prior to receipt of all samples.

In cases of difficulty or further clarification, the laboratory enquiry telephone number is 01793 604798.

### **12 SAMPLE ACCEPTANCE CRITERIA**

Sample acceptance criteria ensure adequate identification for Microbiology samples and request forms in order for them to be accepted by the laboratory for analysis.

The laboratory will make every effort to ensure requests are processed in a safe and timely manner but it is essential that request forms and samples are labelled appropriately and legibly in compliance with this policy. It is also important to clearly identify the investigations required with relevant supporting information. Inadequate or inaccurate labelling results in delays before Microbiology results are available and hence affect patient care. If you have any doubts regarding this policy please ring the relevant department for further information.

The requesting clinician is responsible for the correct completion of the request form and the correct labelling of the sample. It is recommended that samples collected by the patient (eg urine samples) are labelled first by the requesting clinician to minimise the risk of labelling errors.

It is the requester's responsibility to ensure that all details are correct, clearly written and that the sample details match those on the form and patient wrist band (if applicable).

Any labelling discrepancy will be included on the Microbiology report.

Samples will **not** be accepted for analysis if:

- There is no unique identification of the patient i.e. they do not meet the minimum data set for identification
- There is an incorrect sample type or tube
- Incorrect transportation conditions
- Sample is received in a hazardous condition e.g. leaking or sharps attached
- Sample or request form is unlabelled or incorrectly labelled with less than the minimum data sets for patient identification
- Mismatch of details between the form and sample(s)
- The information provided is illegible

Inadequately or inaccurately labelled samples or forms will not be accepted unless they are considered to be unrepeatable or reproducible. A classification of unrepeatable or unreproducible will be made by the Consultant Microbiologist and/or Microbiology Management staff on an individual basis. The risk to the patient of rejection of the sample will be weighed against the risk of acceptance of a wrongly labelled sample. Microbiology will accept no responsibility for samples analysed which initially failed to meet the acceptance criteria and will issue a disclaimer on such reports.

Where the sample is repeatable/ reproducible, no analysis will be performed and an appropriate comment will be included on the Microbiology report. The event may be reported as an incident on the Trust incident report system.

# **13** REPERTOIRE OF TESTS (A - Z)

This section covers the tests that the Microbiology department offers according to the service repertoire agreed with our users.

Find a test or clinical condition using the <u>A – Z list</u>. With each test we provide the following information where appropriate:

- Name of test
- Examinations offered
  - Which sample containers are required What specimen type is required What sample volume is required Which request form should be used
- Sample instructions
  - Collection of the specimen
  - Specimen transportation requirements
  - Specimen storage requirements
  - Special requirements for performing this examination
- Laboratory information
   What test will be performed
   Measurement units of examination performed
   Biological reference intervals of examination performed
   Turnaround time of examination performed
   When the test is available

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 20 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

- Clinical information
  - Factors known to significantly affect the results

For more information on any of these tests see the 'Lab Tests Online' UK website. Almost all examinations are based on NICE accredited UK Standards for Microbiology Investigations (SMI) hosted by Public Health England.

### **13.1** Reference Intervals

Reference intervals for any test are specific to that test and laboratory methodology. Reference intervals will be displayed with the patient results taking these factors into account.

These will be available, whether the result is sent via paper, through ward/web enquiries or via the electronic links to General Practice.

### 13.2 Referred Tests

The laboratory provides a range of specialist testing which is undertaken at reference centres. These tests are indicated within this section. Please contact the laboratory on Telephone 01793 604798 for details of the tests offered, name and location of the testing laboratory and information regarding any special sample requirements.

The parameters analysed in referred tests and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.

### 13.3 Repertoire index

# A B C D E F G H I J K L M N O P Q R S T U V W X Y Z

# Α

Abscesses and deep seated wound infections Adenovirus PCR Amniotic fluid Amoebic serology Antenatal serology Antibiotic levels Antistreptolysin (ASO) titres Aspergillus PCR Aspergillus serology Astrovirus Atypical pneumonia Avian influenza Avian precipitans

### В

Bacteraemia Bacteriuria Bartonella serology (no longer performed in the UK) B-glucan test **Biopsies BK virus PCR Blepharitis Blood cultures** Blood culture collection Bone donor bacteriology screen Bordetella pertussis culture Borrelia burgdorferi (Lyme) antibody Borrelia burgdorferi (Lyme) confirmation Brucella serology Burns Bursa fluid

# С

Campylobacter serology Candidosis Carbapenemase-producing Enterobacteriaceae (CPE) screen Cellulitis Chicken pox (diagnostic) Chicken pox IgG (immunity) Chicken pox PCR

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 22 of 192



Chikunguna, Murray, Ross River, O.Tsusu, Sandfly Chlamydia trachomatis antibody Chlamydia trachomatis PCR Chlamydia trachomatis PCR – collection of urine sample Chlamydia trachomatis PCR - collection of vaginal sample Chlamydia LGV PCR Clostridium difficile toxin Clostridium difficile toxin ribotyping Conjunctivitis Contact lens Continuous ambulatory peritoneal dialysis (CAPD) fluid **Corneal scrape** Cough swab COVID-19 PCR CPE screen Cryptococcal antigen CSF (Cerebro-spinal fluid) microscopy and culture CSF (Cerebro-spinal fluid) oligoclonal bands CSF (Cerebro-spinal fluid) virology PCR Culture Culture: Wounds (deep-seated) Culture: Wounds (skin, superficial, non-surgical) Cystic fibrosis Cytomegalovirus (CMV) serology Cytomegalovirus (CMV) PCR

# D

Dermatological specimens – hair, skin, nails Dengue and West Nile virus Diphtheria serology Dysuria

# E

Ear swab culture Ebola Enteric virus PCR Enterovirus PCR Epstein Barr virus (EBV) serology Epstein Barr virus (EBV) PCR Eye and canalicular pus culture

# F

Faeces culture Faeces: Calprotectin Faeces: Clostridium difficile Faeces: Enteric virus panel

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 23 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Faeces: Norovirus Faeces: Parasitology Faeces: Rotavirus Fluids from normally sterile sites Folliculitis

# G

Genital swab culture (female) Genital specimens (excluding female genital swabs) Glucan (Mycology)

### Η

Haematuria Haemophilus influenzae PCR Haemophilus influenzae type b IgG Helicobacter pylori IgG Hepatitis A virus (HAV) IgG Hepatitis A virus (HAV) IgM Hepatitis B virus (HBV) confirmation Hepatitis B virus (HBV) core IgG antibody Hepatitis B virus (HBV) core IgM antibody Hepatitis B virus (HBV) surface antibody Hepatitis B virus (HBV) surface antigen Hepatitis B virus (HBV) viral load (PCR) Hepatitis C virus (HCV) antibody Hepatitis C virus (HCV) confirmation Hepatitis C virus (HCV) genotype Hepatitis C virus (HCV) qualitative PCR Hepatitis C virus (HCV) viral load Hepatitis D (delta) virus antibody Hepatitis E (delta) virus antibody Herpes simplex virus (HSV) antibody Herpes simplex virus (HSV) type 1 and 2 PCR HIV-1 and 2 antigen/antibodies and p24 antigen **HIV** confirmation HIV resistance, integrase, tropism HIV vertical transmission (neonates) HIV viral load (PCR) Human Herpes 6 (HHV) PCR Human T lymphotrophic virus (HTLV) 1 and 2 serology Hydatid serology

# 

Impetigo Infective endocarditis Influenza A

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 24 of 192

### Influenza B Intravascular cannulae

### J

JC virus PCR Joint fluid

# Κ

# L

Legionella urinary antigen Leptospira serology Lyme disease

# Μ

Measles (diagnostic) Measles IgG (immunity) Meningitis Meningococcal antibody Meningococcal PCR Metapneumonvirus Mouth swab MRSA Mumps (diagnostic) Mumps IgG (immunity) Mycobacteria Mycobacteria PCR Mycology Mycology PCR Mycology serology

# Ν

Neisseria gonorrhoeae PCR Neonatal sepsis Norovirus PCR Nose swab

# 0

Otitis externa Otitis media Ova, cysts and parasites

### Ρ

Panfungal PCR Pan-valentine leukocidin (PVL) toxin detection

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 25 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Parainfluenza virus Parasitology (Bilharzia) Parasitology (Pinworm) Parasitology (serology) Parasitology (Stool) Parasitology (Worm identification) Paronychia Parotitis **Parvovirus PCR** Parvovirus serology Pericardial fluid Peritoneal dialysis fluid (PDF) Peritoneal fluid Pharyngitis Pleural fluid Pneumococcal PCR Pneumococcal serology Pneumococcal urinary antigen Pneumocystis (IF) Polyoma viruses (BK) Polyoma viruses (JC) Prosthetic valve endocarditis Pseudomonas serology Pus Pyuria

# Q

Q fever serology Quantiferon gold TB Quantiferon TB Gold - Instructions for Specimen Collection

# R

Respiratory samples for culture Respiratory syncytial virus (RSV) Respiratory virus PCR Rhinovirus Rotavirus Rubella (diagnostic) Rubella IgG (immunity)

# S

Sapovirus Sepsis Skin, superficial, non-surgical wounds Sputum Sterile fluid

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 26 of 192 Department of Microbiology

Streptococcal serology (ASO) Streptococcus pneumonia serology Syphilis antibody Syphilis confirmation

# T

TB examination TB (Quantiferon Gold) Tetanus antibody Throat swab Tips/intravascular cannulae Tissues and biopsies Toxoplasma (diagnostic) Toxoplasma IgG (immunity) Treponema pallidum antibody Treponema pallidum Confirmation Treponema pallidum PCR

# U

Ulcers Urinary tract infection Urines (microscopy and culture)

# V

Varizella zoster virus (VZV) IgG (immunity) Varicella zoster virus (VZV) PCR Viral haemorrhagic fever (VHF) Virus isolation

# W

West Nile virus Whooping cough Wounds (skin, superficial, non-surgical) Wounds (deep-seated)

# Х

# Υ

Z

Zika virus

# Abscesses and deep seated wound infections

Abscesses are accumulations of pus in the tissues and any organism isolated from them may be of significance. They occur in many parts of the body as superficial infections or as deep-seated infections associated with any internal organ.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Collection of pus or exudate	Minimum volume 1ml of pus			
Concernent Harrison (20)	Amies transport swab	Swabs should be well soaked in pus			
Sample instructions					
Collection	Optimally collected b Collection of pus or e tiny amounts, then s microflora.	Optimally collected before antimicrobial therapy started. Collection of pus or exudate is always preferable to swabs, except when in tiny amounts, then sample the deepest part of the wound avoiding superficial microflora.			
Specimen transport Specimens should be sent to the laboratory without delay dur working hours.			ut delay during normal		
Storage requirements	Outside of normal w Delays of over 48 ho	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.			
<b>Special requirements</b> Important to indicate site and nature of lesion.					
Laboratory information					
Tests	Microscopy for detec quantitative) (pus). General isolation and anaerobic micro-orga	ction of gram positive and neg d characterisation of aerobic, anisms (qualitative).	gative bacteria (semi- microaerophilic and		
Measurement units	Growth detected or not detected.				
Biological reference units					
Turnaround time	4 days, plus 2 days fo	4 days, plus 2 days for enrichment culture (pus).			
Availability	Routine hours and o	Routine hours and on-call (pus).			
Clinical information					
Factors known to significantlyThe recovery of anaerobes is compromised if transport time exceeds 3 houaffect the resultsDelays in transportation may affect the recovery of pathogens.			sport time exceeds 3 hours. of pathogens.		

# **Adenovirus PCR**

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	Minimum volume 500µl		
	Eye swab (virus transport medium)			
	Stool sample	<20ml		
Sample instructions				
Collection	Send a viral (green top) swab of vesicle fluid or affected mucous membranes. Faeces specimen may be passed into a clean, dry, disposable bedpan or similar container and transferred to an appropriate collection container.			
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives ma inappropriate timir of organism below detection of an ass New and emerging this assay.	y occur for a variety of reason ng of sample collection, inapp the detectable limit of the as ay sampling variation will res variants may also occur whic	ns, for example propriate sample, presence ssay. Towards the limit of ult in lower reproducibility. ch may not be detected by	
Back to index				

Authorised by: C Frearson Date of issue: 07/02/2023

# **Amoebic serology**

Used to determine past or current infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details and date of onset are essential for processing.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significan affect the results	tly Haemolysis.		

# **Antenatal serology**

Infectious Disease in Pregnancy (IDP) screening.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	Use an antenatal screening department approved request form
Sample instructions			
Collection	No special requir	rements.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Requests for blood borne virus testing must be clearly indicated as accepted by the patient and signed by the requesting clinician. Remaining serum sample stored for 2 years.		
Laboratory information			
Tests	Detection of Hepatitis B surface antigen (qualitative) Detection of HIV-1 and 2 antibodies and HIV antigen (qualitative) Detection of Treponema pallidum antibody (qualitative)		
Measurement units			
Biological reference units			
Turnaround time	7 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significat affect the results	ntly Haemolysis.		

# **Antibiotic levels**

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requiren	nents.		
Specimen transport	Specimens should t working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal v	Outside of normal working hours samples should be refrigerated.		
	Requests must be discussed with the Consultant Microbiologist.			
Special requirements	Gentamicin and Vancomycin assays:			
	These are performed by the Biochemistry department at GWH.			
Laboratory information				
Tests	Other Antibiotic Lev Contact the laborat required. The para these parameters w requestor.	Other Antibiotic Level tests are processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	3 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	Haemolysis.			

# Antistreptolysin (ASO) titres

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	itly Haemolysis.			

# Aspergillus PCR

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	Minimum volume 5ml		
	Sputum/BAL	Minimum volume 1ml		
	CSF	Minimum volume 0.5ml		
Sample instructions				
Collection	Sputum specimens Refer to <u>Respirator</u> Cerebrospinal fluid Refer to <u>CSF micro</u>	s/ bronchoalveolar lavage/br ry samples for culture. d (CSF) scopy and culture.	onchial washings	
Specimen transport	Specimens should working hours.	be sent to the laboratory with	nout delay during normal	
Storage requirements	Outside of normal	working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details are	essential for processing.		
Laboratory information				
Tests	This test is process laboratory on Tele parameters analys parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significa affect the results	False negatives ma inappropriate timin ntly of organism below detection of an ass New and emerging this assay.	y occur for a variety of reason ng of sample collection, inapp the detectable limit of the as ay sampling variation will res y variants may also occur whic	ns, for example ropriate sample, presence say. Towards the limit of ult in lower reproducibility. h may not be detected by	
Back to index				

# Aspergillus serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	juirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is pro laboratory on The paramete parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significa affect the results	ntly Haemolysis.	ly Haemolysis.		

# **Avian precipitans**

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on The paramete parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significa affect the results	ntly Haemolysis.			
## **Bartonella serology**

Used to determine past or current infection. *No longer available as a test in this country*. Contact Microbiology consultant if required.

## **BetaGlucan test**

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	One whole red t along with othe test.	op is required for this test in tests, one separate red to	.e. if BetaGlucan is requested p will be needed just for this	
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details a	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is proce laboratory on Te The parameters parameters will requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units	pg/mL	pg/mL		
Biological reference units				
Turnaround time	48-96 hours			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significa affect the results	ntly Lipemic samples	nples. S		

### **BK virus PCR**

Diagnosis of acute disease.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	EDTA blood	Minimum volume 5ml	
	CSF	Minimum volume 0.5ml	
	Urine	Minimum volume 5ml	
Sample instructions			
Collection	<b>Cerebrospinal f</b> Refer to <u>CSF min</u> <b>Urine</b> Refer to <u>Urine (</u>	luid (CSF) croscopy and culture. microscopy and culture).	
Specimen transport	Specimens shou working hours.	Ild be sent to the laboratory witl	hout delay during normal
Storage requirements	Outside of norn	nal working hours samples shoul	d be refrigerated.
Special requirements	Clinical details a	are essential for processing.	
Laboratory information			
Tests	This test is proc laboratory on T parameters ana parameters will requestor.	essed at an external reference c elephone 01793 604798 if furthe lysed in this test and any referen be displayed on the report whe	entre. Contact the er details are required. The nce ranges for these n it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significa affect the results	False negatives inappropriate ti ntly of organism bel detection of an New and emerg this assay.	may occur for a variety of reaso ming of sample collection, inapp ow the detectable limit of the as assay sampling variation will res ging variants may also occur whic	ns, for example propriate sample, presence ssay. Towards the limit of sult in lower reproducibility. ch may not be detected by
Back to index			

Authorised by: C Frearson Date of issue: 07/02/2023

### **Blood cultures**

Bacteria are not normally found in the blood - any growth is usually significant *however* contamination from normal skin flora can easily take place. A strict aseptic technique is essential.

Blood cultures are not a 'routine' investigation. Take only when active clinical infection is suspected and where possible before antibiotics have been given. Take during or as soon as possible after a spike of temperature. Do not remove or cover up barcode labels as these are required in the laboratory.

The following list serves as a guide for when blood cultures should be considered:

- Fever ≥ 38°C (suspected bacterial or fungal cause)
- Pyrexia of unknown origin (PUO)
- Rigors
- Febrile convulsion (paediatrics)
- Sepsis, septicaemia or septic shock
- Febrile neutropenia
- Pneumonia
- Meningitis
- Meningococcaemia/petechial, purpuric or non-blanching rash
- Enteric fever (typhoid)
- Infective endocarditis or other endovascular infection
- Pyelonephritis
- Pancreatitis
- Septic arthritis
- Intravascular catheter/cannula infection
- Enteric fever (e.g. typhoid)

Examinations offered			
Collection container	Specimen	Sample volume	Request form
Children – yellow top bottle. Adults – grey top and purple top bottle.	Venous blood, arterial blood, blood via IV line. Ascetic fluid, pleural fluid.	Children – inoculate up to 4ml. Adults – inoculate up to 10ml in each bottle.	

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 40 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

Sample instructions	
Collection	A blood culture set is defined as one aerobic and one anaerobic bottle. For infants and neonates a single aerobic bottle may be requested. For patients with suspected endocarditis collect 2 sets from separate venepunctures at different times. Refer to <u>Blood Culture Method Options</u> .
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours. Do not use pneumatic chute system.
Storage requirements	Inoculated bottles should be incubated as soon as possible. Outside of normal working hours samples must be stored in the incubator in Pathology Reception. DO NOT refrigerate blood cultures.
Special requirements	Collect specimens before antimicrobial therapy where possible. Samples should be taken as soon as possible after a spike of fever.
Laboratory information	
Tests	Detection of gram positive and negative bacteria (semi-quantitative). General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).
Measurement units	Growth detected or not detected.
Biological reference units	
Turnaround time	1 – 5 days, depending on positivity. Significant positive results are communicated to clinicians as and when they arise.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	Any recent antimicrobial therapy can have a significant effect on blood culture results by decreasing the sensitivity of the test. This may be of particular importance in those patients receiving prophylactic antibiotics and who are at high risk of bloodstream infections. If patients have received previous antimicrobial treatment, bacteraemia should be considered even if blood culture results are negative. There is a direct relationship between blood volume and yield, with approximately a 3% increase in yield per ml of blood cultured. False negatives may occur if inadequate blood culture volumes are submitted.

Authorised by: C Frearson

Date of issue: 07/02/2023

# Great Western Hospitals MHS

**NHS Foundation Trust** 

## **Blood culture collection**





When collecting blood using a wingset, it is recommended that you use a discard tube to prime the wingset tubing first. Then collect blood into blood culture bottles. Push and hold the BD Vacutainer\* holder over the top of the bottle. Fill aerobic bottle first, then anaerobic bottle, holding them upright. Collect blood to indicated fill level.

If required, BD Vacutainer\* blood collection tubes may be drawn at this time by inserting them into the BD Vacutainer® holder.

For maximum safety, the device is designed to be activated while still in the patient's vein. Place a gauze pad on the venepuncture site. Allow it to cover the front barrel. After collection, grasp the body of the device with thumb and middle finger, and activate the push button using your index finger. Do not impede device retraction.

Apply pressure to the venepuncture site in accordance with your institution's protocol.

Make sure that the needle is fully retracted and is in the

shielded position.

DCN: MIC-P-006-12.7 Page 42 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Great Western Hospitals NHS

**NHS Foundation Trust** 



Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 43 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Great Western Hospitals NHS

**NHS Foundation Trust** 

### 5. Additional Cultures



It is generally recommended to collect 2-3 blood culture sets for children (> 36 kg body weight) and adult patients. Additional cultures may be collected in a similar way. I deally, a different venepuncture site should be used for each culture set collected. The clinical status of the patient should be the primary guide to the timing of blood cultures.'

Repeat steps 1-4 for additional cultures.

## For BD Customer Service, please call 01865 781666, Option 1

**Back to index** 

Authorised by: C Frearson

Date of issue: 07/02/2023



Page 44 of 192

DCN: MIC-P-006-12.7 THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Bone donor bacteriology screen

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
OID canner turi (O) ()	Femoral head swab (Amies transport swab	)		
	Bone chips			
Sample instructions				
Collection	Swabs and bone during surgery.	are taken from the patient o	r from the donor femoral head,	
Specimen transport	Specimens shoul working hours.	d be sent to the laboratory v	vithout delay during normal	
Storage requirements	Outside of norm Delays of over 48	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	No special requir	No special requirements.		
Laboratory information				
Tests	General isolatior anaerobic micro-	and characterisation of aero organisms (qualitative).	bbic, microaerophilic and	
Measurement units	Growth detected	l or not detected.		
Biological reference units				
Turnaround time	Femoral head 7 o Bone chips 14 da	days. Iys.		
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Delays in transpo	ortation may affect the recov	ery of pathogens.	

## Bordetella pertussis culture

Whooping cough is a highly contagious disease that is caused by the fastidious Gram-negative coccobacillus *Bordetella pertussis*. In some cases this syndrome may also be caused by *Mycoplasma pneumoniae*, and by viruses such as adenoviruses and enteroviruses. It is advisable to take two pernasal swabs: one for the culture of Bordetella species and the other for viral culture; however nasal swabs for PCR are preferred.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Pernasal swab			
Sample instructions				
Collection	A pernasal sw the nose until Optimally coll	ab is inserted through a nostril it reaches the nasopharynx. ected before antimicrobial ther	and advanced along the floor of rapy started.	
Specimen transport	Specimens sho working hours	ould be sent to the laboratory v s.	without delay during normal	
Storage requirements	Outside of nor Delays of over	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	No special req	juirements.		
Laboratory information				
Tests	General isolat	ion and characterisation of Bor	detella species.	
Measurement units	Growth detec	ted or not detected.		
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours			
Clinical information				
Factors known to significar affect the results	ntly Delays in trans	sportation may affect the recov	very of pathogens.	

# Borrelia burgdorferi (Lyme) antibody

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirem	nents.		
Specimen transport	Specimens should b working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details, date of onset and bite/travel history are essential for processing.			
Laboratory information				
Tests	Detection of Lymes IgM antibody (qualitative). Detection of Lymes IgG antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	tly Haemolysis.			

# Borrelia burgdorferi (Lyme) confirmation

Lyme confirmation would only be performed on a Lyme positive sample.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	No special req	No special requirements.		
Laboratory information				
Tests	This test is pro laboratory on parameters ar parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

# Brucella serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	juirements.		
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and any history of travel or occupational exposure are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

# **Campylobacter serology**

Used to determine past or current infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special rec	quirements.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours	5.	
Clinical information			
Factors known to significar affect the results	ntly Haemolysis.		

## Carbapenemase-producing Enterobacteriaceae (CPE) screen

In response to the increasing numbers of CPE producing clinical isolates of Enterobacteriaceae the Infection Control Team and Microbiology department have produced a protocol for CPE screening and detection. The isolation of a clinical CPE isolate prompts the Infection Control Team to screen all possible patient contacts to reduce the transmission of resistance enzymes within the Trust.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
OID Carrier War O D	Rectal swab (Amies transport swab)			
	Stool sample	<20ml		
Sample instructions				
Collection	Optimally collecte Faeces specimen similar container a Rectal swabs mus	d before antimicrobial ther may be passed into a clean, and transferred to an appro t have evidence of stool on	apy started. dry, disposable bedpan or priate collection container. swab for optimal sensitivity.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.			
Special requirements	No special require	No special requirements.		
Laboratory information				
Tests	General isolation Enterobacteriacea	and characterisation of carb ae (qualitative).	apenemase producing	
Measurement units	Growth detected	or not detected.		
Biological reference units				
Turnaround time	Negative screen 24 hours. Positive result 4 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly			

## Chikungunya, Murray, Ross River, O.Tsusu, Sandfly

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details, date of onset and travel history are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours			
Clinical information				
Factors known to significat affect the results	Haemolysis.			

# Chlamydia trachomatis antibody

Used to determine past infection during investigations for infertility in women.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special requ	irements.	
Specimen transport	Specimens shou working hours.	uld be sent to the laborator	y without delay during normal
Storage requirements	Outside of norn	nal working hours samples	should be refrigerated.
Special requirements	Clinical details i	.e. Respiratory / Infertility a	re essential for processing.
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significan affect the results	Haemolysis.		

# Chlamydia trachomatis PCR

Examinations offered			
Collection container	Specimen	Sample volume	Request form
Arrithment and Arristed and Arrithment	Eye, cervical, urethral, throat, rectal swab (Chlamydia transport medium)		
Annual Annua	Urine (first void) (Chlamydia transport medium)	Minimum volume 2ml	
	Urine (first void)	Minimum volume 2ml	
Sample instructions			
Collection	guidelines on the Refer to <u>Chlamyd</u> collection of urine Urine specimens s submitted in whit transport medium <b>Endocervical or s</b> An endocervical or s trachomatis as it vaginal swab. Wh negative result ar endocervix with t NB. Only one swa swab must not be <b>Men</b> The patient shoul approximately 10 universal containe <b>Eye swabs</b> Do not use fluore Apply a local anae from a female PC remaining swab, f collect epithelial o medium. Place sy cap.	collection packs. <u>ia PCR – collection of vaginal set sample</u> . submitted from non-Sexual H te topped universal containers in the laboratory. <b>elf-taken vaginal swab</b> wab is the specimen of choice has a higher sensitivity than a hite cells and blood can produce the accompanying swab prior b is required for a self-taken we te used and should be discarded d not have urinated for at lea -20 mls of first voided urine in er. scein as this can interfere witt esthetic. Remove excess exucor R sample kit; discard the clean firmly swab the inner surface cells. Do NOT pre-moisten the wab in sample tube, snap off a	eample and <u>Chlamydia PCR –</u> ealth Clinic locations can be s for transfer into Chlamydia urine sample or a self-taken ce either an invalid or false buld be removed from the to taking the sample. vaginal swab; the cleaning d. st one hour. Collect nto a sterile white capped h the test. late using one of the swabs ning swab. Using the of upper and lower eyelids to e swab in the transport at the score line and replace

Department of Microbiology

Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.
Storage requirements	Outside of normal working hours samples should be refrigerated.
Special requirements	Urine – patient should not have urinated for 2 hours prior to sample collection.
Laboratory information	
Tests	Detection of Chlamydia trachomatis nucleic acid (qualitative).
Measurement units	Presence detected or not detected.
Biological reference units	
Turnaround time	4 days.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. White cells and blood can produce either an invalid or false negative result. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.

## Chlamydia trachomatis PCR - collection of urine sample

# Aptima<sup>®</sup> urine collection kit Collection procedure guide

#### Collection for male and female urine specimens

Patient should not have urinated for at least 1 hour prior to specimen collection.



Direct patient to provide first-catch urine (approximately 20 to 30 mL of initial urine stream) into urine collection cup free of any preservatives. Collection of larger volumes of urine may result in specimen dilution that may reduce test sensitivity. Female patients should not cleanse labial area prior to providing specimen.

#### Urine specimen collection guide for:

- Chlamydia trachomatis (CT)
- Neisseria gonorrhoeae (GC)
- Trichomonas vaginalis (TV) for female only

luid is etween black fill lines

into urine specimen transport tube using the disposable pipette provided. The correct volume of urine has been added when the fluid level is between the black fill lines on urine specimen transport tube label. Ter Alt

Remove cap from urine specimen transport tube and transfer 2 mL of urine

Re-cap urine specimen transport tube tightly. This is now known as the "processed urine specimen."

## Chlamydia trachomatis PCR – collection of vaginal sample



#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 57 of 192

#### THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

## Chlamydia LGV PCR

Chlamydia LGV PCR would only be performed on a Chlamydia positive rectal sample.

In order to diagnose LGV, different samples from those listed may be indicated; please discuss with Consultant Medical Microbiologist.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
Second areas and a second areas and a second areas and a second areas and a second areas	Rectal swab (Chlamydia transport medium)		
Sample instructions			
Collection	Specimens shoul guidelines on the	d be collected and handled collection packs.	following the recommended
Specimen transport	Specimens shoul working hours.	d be sent to the laboratory	without delay during normal
Storage requirements	Outside of norma	al working hours samples sl	hould be refrigerated.
Special requirements	No special requir	ements.	
Laboratory information			
Tests	This test is proce laboratory on Tel parameters analy parameters will b requestor.	ssed at an external referen lephone 01793 604798 if fu ysed in this test and any ref pe displayed on the report v	ce centre. Contact the irther details are required. The ference ranges for these when it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significat affect the results	False negatives n inappropriate tin of organism belo detection of an a New and emergin this assay.	nay occur for a variety of re ning of sample collection, ir w the detectable limit of th ssay sampling variation wil ng variants may also occur	easons, for example happropriate sample, presence he assay. Towards the limit of I result in lower reproducibility. which may not be detected by

### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

## **Clostridium difficile toxin**

*C. difficile* is a Gram positive, spore forming, strictly anaerobic rod, so named because of the difficulty in original culture and characterisation. Toxigenic strains produce large protein toxins A and B, both being major virulence factors. Most disease associated with *C. difficile* is intestinal though *C. difficile* may be isolated from blood or tissues.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Stool sample	<20ml	
Sample instructions			
Collection	Specimen may container and	be passed into a clean, dry, d transferred to an appropriate	isposable bedpan or similar collection container.
Specimen transport	Specimens sho working hours	ould be sent to the laboratory .	without delay during normal
Storage requirements	Outside of nor Delays of over	mal working hours samples sh 48 hours are undesirable.	ould be refrigerated.
Special requirements	Formed stools are unsuitable for investigation for C.difficile. Clostridium difficile toxin test performed on in-patient samples, patients over 65yrs or if history of antibiotic-associated diarrhoea. Children less than 2 years old are unsuitable for investigation for C.difficile. Investigation not performed if a positive result within previous 28 days.		
Laboratory information			
Tests	Glutamate dehydrogenase (GDH) detection (qualitative), Clostridium difficile toxin A and B detection (qualitative) and PCR ribotyping of Clostridium difficile (qualitative).		
Measurement units	Toxin detected	l or not detected.	
Biological reference units			
Turnaround time	1 day.		
Availability	Routine hours		
Clinical information			
Factors known to significat affect the results	ntly		

# Clostridium difficile toxin ribotyping

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Stool sample	<20ml	
Sample instructions			
Collection	Specimen may container and	be passed into a clean, dry, dis transferred to an appropriate c	posable bedpan or similar collection container.
Specimen transport	Specimens sho working hours.	uld be sent to the laboratory w	vithout delay during normal
Storage requirements	Outside of nor Delays of over	mal working hours samples sho 48 hours are undesirable.	ould be refrigerated.
Special requirements	Investigation p Consultant dur	erformed at request of Infection in the structure in the section in the section of the section o	on Control Microbiology
Laboratory information			
Tests	This test is pro on Telephone ( analysed in this displayed on th	cessed at an external reference 01793 604798 if further details s test and any reference ranges ne report when it is returned to	e centre. Contact the laboratory are required. The parameters 5 for these parameters will be 9 the requestor.
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
<b>Clinical information</b>			
Factors known to significan affect the results	ntly		

## **Contact lens**

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Contact lens case or sterile container with saline		
Sample instructions			
Collection	No special requirem	ents.	
Specimen transport	Specimens should b working hours.	e sent to the laboratory witho	out delay during normal
Storage requirements	Outside of normal w Delays of over 48 hc	vorking hours samples should ours are undesirable.	be refrigerated.
Special requirements	No special requirem	ents.	
Laboratory information			
Tests	Gram stain and cult	ure.	
Measurement units	Growth detected or	not detected.	
Biological reference units			
Turnaround time	5 days.		
Availability	Routine hours and o	n-call.	
Clinical information			
Factors known to significant	tly Delays in transporta	tion may affect the recovery	of pathogens.

## **Corneal scrape**

Keratitis is an inflammation of the cornea which is a serious condition requiring prompt and meticulous investigation, and may progress to perforation and blindness if treatment is unsuccessful. Predisposing factors include prior ocular disease, wearing contact lenses and use of topical corticosteroids. The condition may be caused by a wide range of bacteria, fungi and parasites.

Collection container S	pecimen	Sample volume	Request form
Chocolate agar A SAB agar s FAA agar C Acanthamoeba plate c Microscope slide n	Aqueous and vitreous Jumour, corneal crapings. Direct inoculation onto ulture plates and nicroscope slide	Sufficient quantity to make a visible deposit on to a microscope slide and to inoculate agar plates	
Sample instructions			
Collection	Optimally collected by Performed by trained Performed after Use sterile need Carefully spread marker) for Grar Carefully smear If insufficient specim cultures should be th	before antimicrobial therapy s d staff according to Trust poli instillation of local anaesther le or loop to scrape base of u material onto glass slide (circ n staining and/or material onto agar plate en to make an impression sm he priority.	started. cy: cic eye drops lcer cle area with permanent clear and inoculate plates,
Specimen transport	Specimens should be working hours and o	e sent to the laboratory withon n-call.	ut delay during normal
Storage requirements	Outside of normal we Delays of over 48 ho	orking hours samples should urs are undesirable.	be refrigerated.
Special requirements	Contact the laborato required for Acantham	ry (Telephone 01793 604798 oeba culture,24 hours in advanc	) if Acanthamoeba plate e of specimen collection.
Laboratory information			
Tests	Gram stain and cultu	re.	
Measurement units	Growth detected or	not detected.	
Biological reference units			
Turnaround time	5 days.		
Availability	Routine hours and or	n-call.	
Clinical information			
Factors known to significantly affect the results	Where media and sm transported immedia Delays in transportat	nears are inoculated at the pa ately to the laboratory for pro ion may affect the recovery o	itient's side they must be ocessing. of pathogens.
Back to index			

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

## **COVID-19 PCR**

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Viral swab in transport media	Nose and throat swab	
Sample instructions			
Collection	Nose and Throat sw bagged. Do <b>not</b> rem	ab collected wearing correct ove viral transport media fro	PPE. Swabs should be double m sample container.
Specimen transport	Specimens should b where appropriate t samples should be t	e taken directly to Microbiolo to prevent delay of results. O aken to Pathology Reception	ogy during working hours utside working hours
Storage requirements	Outside of normal w	vorking hours samples should	be refrigerated.
Special requirements	Clinical details are e Do <b>not</b> ring the labo For rapid testing ple	ssential for processing. pratory for results. pase speak to the Site Manag	ers.
Laboratory information			
Tests	SARS-CoV2 PCR Test	t	
Measurement units	N/A		
Biological reference units	N/A		
Turnaround time	Rapid: 2 hours* Routine: 6-8 hours* *From receipt in lab	oratory	
Availability	Weekday: Routine h Weekend: Routine h	nours nours with scope for site app	roved rapid testing at 16:00
<b>Clinical information</b>			
Factors known to significar affect the results	Results may be affe viral material pres processing times. Detection of low-lev Results produced of bacterial pathogens	ected by improperly collected ent in the specimen and/ rel viral RNA may not be of cl cannot rule out infections/c	d samples, low or insufficient 'or delays in transport and inical significance. disease from other viral and

# Cryptococcal antigen

Used to determine past or current infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
	CSF	Minimum volume 0.5ml	
Sample instructions			
Collection	<b>Cerebrospinal fluic</b> Refer to <u>CSF micros</u>	I (CSF) scopy and culture.	
Specimen transport	Specimens should I working hours.	be sent to the laboratory with	out delay during normal
Storage requirements	Outside of normal	working hours samples should	d be refrigerated.
Special requirements	Clinical details are	essential for processing.	
Laboratory information			
Tests	This test is process laboratory on Telep parameters analyse parameters will be requestor.	ed at an external reference ce ohone 01793 604798 if furthe ed in this test and any referen displayed on the report wher	entre. Contact the r details are required. The ce ranges for these n it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significa affect the results	ntly Haemolysis.		

## CSF (Cerebro-spinal fluid) microscopy and culture

Meningitis is defined as inflammation of the meninges. This process may be acute or chronic and infective or non-infective. Many infective agents have been shown to cause meningitis, including viruses, bacteria, fungi and parasites.

Collection container	Specimen	Sample volume	Request form	
	CSF	Minimum volume 1ml		
Sample instructions				
	Optimally co Do not delay	llected before antimicrobial thera antibiotic administration if clinica	apy started. ally indicated.	
	Sample taker with Trust pr	n using a strict aseptic technique ocedure.	by trained medical staff in line	
Collection	Dispense CSF containers an estimation o	- (minimum 0.5ml in each bottle) nd label in order of removal 1 to 3 f glucose levels.	into 3 sterile single use 3, plus a fluoride bottle for the	
	Where meni antibiotics al Bacteria EDTA blo	<ul> <li>Where meningococcal meningitis/septicaemia is suspected (particularly if antibiotics already give in community) also send:</li> <li>Bacterial throat swab and request meningococcal culture</li> <li>EDTA blood for meningococcal DNA PCR</li> </ul>		
Specimen transport	Specimens sl hours. Outsi reception frie through swit Do not use p required.	hould be sent to the laboratory w de of normal hours samples shou dge and the on-call Microbiology chboard (Telephone 01793 60402 neumatic chute system if investig	ithout delay during normal Id be placed in the pathology Biomedical Scientist contacted 20). ation for Xanthochromia	
Storage requirements	See above.			
Special requirements	Always cont Ideally collec 1 to 3 accord	act the laboratory when sending t the CSF sample in 3 consecutive lingly.	specimens. e universal containers, labelled	
Laboratory information				
Tests	Presence of v Differential o Detection of Detection of General isola anaerobic m	white blood cells and red blood co of white blood cells (qualitative). Cryptococcus neoformans capsul gram positive and negative bacte ation and characterisation of aero icro-organisms (qualitative).	ells (quantitative). es (qualitative). eria (semi-quantitative). bic, microaerophilic and	
Measurement units	Cell count x :	10 <sup>6</sup> /l		

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

Biological reference units	Leucocytes: Neonates 1 – 12 months Adults Erythrocytes:	0 – 30 cells x 10 <sup>6</sup> /l 0 – 20 cells x 10 <sup>6</sup> /l 0 – 5 cells x 10 <sup>6</sup> /l No red cells should be present in normal CSF*
Turnaround time	Microscopy 2 hours. Culture 2 days. Significant positive results are communicated to clinicians as and when they arise.	
Availability	Routine hours and on-call.	
Clinical information		
Factors known to significantly affect the results	Cells disintegrate not reflective of t Delays in transpo	A delay in transportation may produce a cell count that is he clinical situation of the patient. Article recovery of pathogens.

**Back to index** 

Authorised by: C Frearson Date of issue: 07/02/2023

## CSF (Cerebro-spinal fluid) Oligoclonal bands

The presence of Oligoclonal bands in cerebrospinal fluid combined with their absence in blood serum often indicates that immunoglobulins are produced in central nervous system.

Oligoclonal bands are an important indicator in the diagnosis of multiple sclerosis.

The presence of one band (a monoclonal band) may be considered serious, such as lymphoproliferative disease, or may simply be normal — it must be interpreted in the context of each specific patient. More bands may reflect the presence of a disease. The bands tend to disappear from the cerebrospinal fluid as a person recovers from the neurological disease

Examinations offered	<u>Canadiman</u>	Comula visitiones	Dominant form
Collection container	Specimen	Sample volume	Request form
	CSF	Minimum volume 1ml	
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	Refer to <u>CSF microscopy and culture</u> .		
Specimen transport	Refer to <u>CSF microscopy and culture</u> .		
Storage requirements	Refer to <u>CSF microscopy and culture</u> .		
Special requirements	CSF and a paired venous blood sample are required for testing. Refer to <u>CSF microscopy and culture</u> .		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units	mg/dL		
Biological reference units	N/A		
Turnaround time	6 weeks.		
Availability	Routine hours.		
Clinical information			
Factors known to significan affect the results	ntly		
sack to index			
orised by: C Frearson of issue: 07/02/2023			DCN: MIC-P-00 Page 67
			ED

# CSF (Cerebro-spinal fluid) Viral PCR

Discuss all molecular/PCR requests with Microbiology Consultant or Senior Laboratory Biomedical Scientist.

The standard viral PCR panel includes Enterovirus, Herpes simplex virus and Varicella-Zoster.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	CSF	Minimum volume 1ml		
Sample instructions				
Collection	Refer to <u>CSF micr</u>	oscopy and culture.		
Specimen transport	Refer to <u>CSF micr</u>	oscopy and culture.		
Storage requirements	Refer to <u>CSF micr</u>	oscopy and culture.		
Special requirements	Refer to <u>CSF micr</u>	oscopy and culture.		
Laboratory information				
Tests	This test is proces laboratory on Tel parameters analy parameters will b requestor. Detec nucleic acid, Herp (HSV-2) nucleic ac	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor. Detection of Enterovirus nucleic acid, Varizella-Zoster virus nucleic acid, Herpes Simplex Virus (HSV) type 1 (HSV-1) and HSV type 2 (HSV-2) nucleic acid (qualitative).		
Measurement units	N/A			
Biological reference units	N/A			
Turnaround time	4 days Significant positiv arise.	e results are communicated to	o clinicians as and when they	
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives m inappropriate tim ntly of organism below detection of an a New and emergin this assay.	ay occur for a variety of reaso ing of sample collection, inap w the detectable limit of the a ssay sampling variation will res g variants may also occur whi	ons, for example propriate sample, presence ssay. Towards the limit of sult in lower reproducibility. ch may not be detected by	

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 68 of 192
	THIS DOCUMENT IS LINCONTROLLED WHEN PRINTED	

## Culture

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
California, Russian (California)	Amies transport swab			
	Collection of pus or exudate			
	Collection of pus or exudate			
Sample instructions				
Collection	Optimally collec	Optimally collected before antimicrobial therapy started.		
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm Delays of over 4	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Please state ana	Please state anatomical site and nature of lesion on request form		
Laboratory information				
Tests	Detection of white blood cells, gram positive and negative bacteria (semi- quantitative) (fluids/pus). General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).		and negative bacteria (semi- obic, microaerophilic and	
Measurement units	Growth detected	d or not detected.		
Biological reference units				
Turnaround time	4 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Delays in transp	ortation may affect the reco	very of pathogens.	

## **Cystic fibrosis**

Cystic fibrosis (CF) is caused by a defect in the CF transmembrane conductance regulator gene that affects the transport of ions and water across the epithelium. This leads to progressive pulmonary disease associated with pulmonary infections, which are the major cause of morbidity and mortality in CF patients. The major pathogens are *S. aureus*, *H. influenza* (usually non-encapsulated in CF patients), *S. pneumoniae*, *Burkholderia* and pseudomonads, particularly mucoid *P. aeruginosa* strains. Strains of *P. aeruginosa* with differing antibiotic susceptibilities may be isolated from a single sample.

Collection container	Specimen	Sample volume	Request form	
Of Di Courses Hurse (D) (D)	Cough swab (Amies transport swab)			
	Sputum	Minimum volume 5ml		
Sample instructions				
	Optimally collecte	d before antimicrobial therap	y started.	
Collection	<b>Sputum specimer</b> Refer to <u>Respirato</u>	is bry samples for culture.		
	<b>Cough swabs</b> Younger patients from the upper ai	do not usually expectorate ar rway as an alternative to sputi	nd cough swabs may be taken um samples.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of norma Delays of over 48	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	No special requirements.			
Laboratory information				
Tests	General isolation anaerobic micro-c	and characterisation of aerobi organisms (qualitative).	c, microaerophilic and	
Measurement units	Growth detected	Growth detected or not detected.		
Biological reference units				
Turnaround time	4 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significant affect the results	Iy Delays in transportation may affect the recovery of pathogens. The recovery rate of Haemophilus is reduced the longer the time taken to transport the specimen.			

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

A positive cough swab is a strong predictor of a positive sputum sample; however, a negative cough swab cannot rule out lower airway infection and persistent symptoms should be further investigated, for example by BAL.

# Cytomegalovirus (CMV) serology

Diagnosis of acute/recent or reactivated disease (IgM) or if evidence of past infection/exposure required (IgG).

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special rec	quirements.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.		
Laboratory information			
Tests	Detection of CMV IgM and IgG antibody (qualitative). This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	7 days.		
Availability	Routine hours	5.	
Clinical information			
Factors known to significat affect the results	ntly Haemolysis.		
# Cytomegalovirus (CMV) PCR

Diagnosis of acute disease.

For diagnosis of congenital CMV send neonatal urine sample within first three weeks of life.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	EDTA blood	Minimum volume 500µl	
	Urine	Minimum volume 5ml	
Sample instructions			
Collection	<b>Urine</b> Refer to <u>Urine</u>	(microscopy and culture).	
Specimen transport	Specimens sho working hours	ould be sent to the laboratory with	nout delay during normal
Storage requirements	Outside of nor	mal working hours samples shoul	d be refrigerated.
Special requirements	Clinical details CMV DNA PCR the Consultant	are essential for processing. is a specialist test – outside of the Microbiologist.	ese specialties discuss with
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days		
Availability	Routine hours.		
Clinical information			
Factors known to significan affect the results	False negatives inappropriate of organism be detection of ar New and emer this assay.	s may occur for a variety of reaso timing of sample collection, inapp clow the detectable limit of the as n assay sampling variation will res ging variants may also occur whic	ns, for example propriate sample, presence ssay. Towards the limit of ult in lower reproducibility. ch may not be detected by

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 73 of 192
	THIS DOCUMENT IS LINCONTROLLED WHEN PRINTED	

### **Dengue and West Nile virus**

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hour	ould be sent to the laborato s.	ry without delay during normal	
Storage requirements	Outside of no	rmal working hours samples	s should be refrigerated.	
Special requirements	Clinical detail	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours	S.		
Clinical information				
Factors known to signification affect the results	ntly Haemolysis.			

# Diphtheria serology

Used to determine past or current infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special rec	quirements.	
Specimen transport	Specimens sh working hours	ould be sent to the laborato s.	ry without delay during normal
Storage requirements	Outside of no	rmal working hours samples	should be refrigerated.
Special requirements	Clinical details and any history of travel or occupational exposure are essential for processing.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours	5.	
Clinical information			
Factors known to significat affect the results	Haemolysis.		

### Ear swab culture

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Ear swab (Amies transport swab)			
Sample instructions				
Collection	Optimally collec	ted before antimicrobial ther	apy started.	
Specimen transport	Specimens shou working hours.	ld be sent to the laboratory v	vithout delay during normal	
Storage requirements	Outside of norm Delays of over 4	al working hours samples she 8 hours are undesirable.	ould be refrigerated.	
Special requirements	For investigatior are preferred, al	For investigation of fungal infection, scrapings of material from the ear canal are preferred, although swabs can also be used.		
Laboratory information				
Tests	General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).		obic, microaerophilic and	
Measurement units	Growth detected	Growth detected or not detected.		
Biological reference units				
Turnaround time	4 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significan affect the results	Delays in transp	ortation may affect the recov	very of pathogens.	

### **Enteric virus PCR**

Diagnosis of acute disease.

Enteric virus screen including:

- Adenovirus
- Astrovirus
- Rotavirus
- Sapovirus
- Norovirus

Rotavirus, sapovirus, astrovirus and adenovirus are major causes of acute gastroenteritis. The majority of infections occur in infants and young children. Infections in the elderly are also reported for these agents, and chronic infections can result in immunocompromised patients. Norovirus is the cause of epidemic gastroenteritis.

Collection container	Specimen	Sample volume	Request form
	Stool sample	Liquid specimen: 1 – 2ml Formed specimen: large pea size sample	
Sample instructions			
Collection	Specimen ma container and	y be passed into a clean, dry, dispo d transferred to an appropriate colle	sable bedpan or similar ection container.
Specimen transport	Specimens sh working hour	nould be sent to the laboratory with s.	out delay during normal
Storage requirements	Outside of no	ormal working hours samples should	be refrigerated.
Special requirements	Clinical detail	s are essential for processing.	
Laboratory information			
Tests	This test is pr laboratory on parameters a parameters w requestor.	ocessed at an external reference ce n Telephone 01793 604798 if furthe nalysed in this test and any referen vill be displayed on the report wher	entre. Contact the r details are required. The ce ranges for these n it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days		
Availability	Routine hour	S.	
arised by C Freezen			

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Clinical information	
Factors known to significantly affect the results	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.

### **Enterovirus PCR**

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	Minimum volume 500µl		
Sample instructions				
Collection	No special requir	ements.		
Specimen transport	Specimens should working hours.	d be sent to the laboratory with	nout delay during normal	
Storage requirements	Outside of norma	I working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details an	e essential for processing.		
Laboratory information				
Tests	This test is proce laboratory on Tel parameters analy parameters will b requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives m inappropriate tim of organism belo detection of an a New and emergin this assay.	hay occur for a variety of reason ning of sample collection, inapp w the detectable limit of the as ssay sampling variation will res ng variants may also occur whic	ns, for example propriate sample, presence say. Towards the limit of ult in lower reproducibility. th may not be detected by	

### Epstein Barr virus (EBV) serology

Assay useful in distinguishing individuals who have acquired the infection recently from those who have not (EBV IgM, EBV IgG, EBV confirmation (EBNA)). Detection of EBV IgM is consistent with acute disease, but may also be detectable in chronic or reactivated disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	working hours samples s	hould be refrigerated.	
Special requirements	State whether te (IgM) or if evidenc Clinical details are	State whether test for diagnosis of acute/recent or reactivated disease (IgM) or if evidence of past exposure required (IgG). Clinical details are essential to allow for interpretation.		
Laboratory information				
Tests				
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	Haemolysis.			

# Epstein Barr virus (EBV) PCR

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	Minimum volume 500µl		
Sample instructions				
Collection	No special requirer	nents.		
Specimen transport	Specimens should working hours.	be sent to the laboratory with	nout delay during normal	
Storage requirements	Outside of normal	working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details are EBV DNA PCR is a s the Consultant Mic	Clinical details are essential for processing. EBV DNA PCR is a specialist test – outside of these specialties discuss with the Consultant Microbiologist.		
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives ma inappropriate timin of organism below detection of an ass New and emerging this assay.	y occur for a variety of reason ng of sample collection, inapp the detectable limit of the as ay sampling variation will res y variants may also occur whic	ns, for example ropriate sample, presence say. Towards the limit of ult in lower reproducibility. h may not be detected by	

# Eye and canalicular pus culture

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Collection of pus or exudate	Minimum volume 1ml of pus		
QID Counter Married () ()	Eye swab (Amies transport swab)			
Sample instructions				
Collection	Optimally collected b Collection of pus or e tiny amounts, then sa microflora. Hold the swab paralle lower eyelid.	before antimicrobial therapy s exudate is always preferable t ample the deepest part of the el to the cornea and gently ru	started. so swabs, except when in e wound avoiding superficial ub the conjunctiva in the	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.			
Special requirements	Separate samples she detection of <u>viruses</u> of	Separate samples should be collected into appropriate transport media for detection of <u>viruses</u> or <u>C.trachomatis</u> .		
Laboratory information				
Tests	Detection of white bl quantitative). General isolation and anaerobic micro-orga	ood cells, gram positive and characterisation of aerobic, anisms (qualitative).	negative bacteria (semi- microaerophilic and	
Measurement units	Growth detected or	not detected.		
Biological reference units				
Turnaround time	4 days, plus 2 days fo	or enrichment culture (pus).		
Availability	Routine hours and or	n-call (pus).		
Clinical information				
Factors known to significat affect the results	Delays in transportat	ion may affect the recovery o	of pathogens.	

### **Faeces culture**

Collection container	Specimen	Sample volume	Request form
	Stool sample	Liquid specimen: 1 – 2ml Formed specimen: large pea size sample	
Sample instructions			
Collection	Optimally collec Specimen may b container and tr	ted before antimicrobial therapy so be passed into a clean, dry, disposa ransferred to an appropriate collec	tarted. able bedpan or similar tion container.
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.	
Storage requirements	Outside of norm Delays of over 4	nal working hours samples should 8 hours are undesirable.	be refrigerated.
Special requirements	Please provide i	nformation regarding recent forei	gn travel and antibiotic us
Laboratory information			
Tests	Macroscopic ass Presence and id Detection of Cyo (qualitative). General isolatio anaerobic micro <u>Clostridium diffi</u> 65yrs or if histor <u>Rotavirus</u> test p <u>Norovirus</u> test p the investigation <u>Parasitology</u> tes clinical syndrom Repeat samples Microbiologists Investigations n culture, within t	sessment of consistency/appearan entification Cryptosporidium and o clospora sp, Isospora sp and Crypto n and characterisation of aerobic, p-organisms (qualitative). cile toxin test performed on in-par ry of antibiotic-associated diarrhoo erformed on samples from childre performed only on instruction by th n of outbreaks. t performed on samples depender ne. for microbiological clearance not will advise if necessary. ot performed on in-patient stools the same in-patient episode.	ce. Giardia lamblia (qualitativ osporidium sp oocysts microaerophilic and tient samples, patients ov ea. n <5 years. ne Infection Control Team nt on travel history and usually required – within 30 days of a previo
Measurement units	Growth detecte	d or not detected.	
Biological reference units			
Turnaround time	4 days. Significant posit arise.	ive results are communicated to c	linicians as and when they

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

#### **Clinical information**

**Factors known to significantly** Delays in transportation may affect the recovery of pathogens.

affect the results

# **Faecal Calprotectin**

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Stool sample	Nose and throat swab		
Sample instructions				
Collection	Specimen may be passed into a clean, dry, disposable bedpan or similar container and transferred to an appropriate collection container.			
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm Delays of over 4	al working hours samples should 8 hours are undesirable.	d be refrigerated.	
Special requirements	Faecal Calproted Childrens Unit.	Faecal Calprotectin is only available for GP patients, Gastroenterology and Childrens Unit.		
Laboratory information				
Tests	Faecal Calproted	ctin		
Measurement units	µg/g			
Biological reference units				
Turnaround time	7 days			
Availability	Routine hours.	Routine hours.		
Clinical information				
	Liquid stools a	re processed by the Immunol	ogy Department in Bristol.	
Factors known to significar affect the results	Patients who htly (NSAIDs) may l	Patients who are taking non-steroidal anti-inflammatory drugs y (NSAIDs) may have elevations in their faecal calprotectin levels.		
	Assay results s and laboratory decisions.	should be interpreted in conj v data to assist clinicians in m	junction with other clinical aking patient management	

### Fluids from normally sterile sites

The detection of organisms in fluids that are normally sterile indicates significant infection, which can be life-threatening. Specimens may be taken primarily for culture or this may be incidental to the prime reason for obtaining the specimen.

Blood cultures may be positive with the same infecting organism, and occasionally may be positive when culture of the fluid fails to reveal the organism.

Fluids will be sterile in the absence of infection, as will "sympathetic effusions", and those of immunological or traumatic origin and those due to metabolic disease or heart failure.

Signs of infection may be difficult to detect clinically in patients whose joints are already inflamed due to rheumatological conditions. This is important because these patients are at increased risk of joint sepsis. Do not remove or cover barcodes on bottles as these are required by the laboratory.

Collection container	Specimen	Sample volume	Request form		
	Collection of amniotic fluid, bursa pericardial fluid, synovial (joint) fluid, peritoneal fluid (ascites), pleural fluid.	Minimum volume 1ml			
		Inoculate up to 10ml in each bottle			
Sample instructions					
	Optimally collected	before antimicrobial therap	by started.		
	Samples include:				
	Ascitic fluid: ?spontaneous bacterial peritonitis				
	CAPD fluid: ?PD peritonitis				
	Pleural fluid: ?empyema				
Collection	Synovial or bursa fluid: ?septic arthritis or bursitis				
Concernon	Vitreous fluid: ?endophthalmitis				
	Samples taken using strict aseptic technique – by trained medical staff in				
	line with Trust procedure.				
	Ideally a minimum volume of 1ml should be collected.				
	Where adequate sa	mple, inoculate also into bl	ood culture bottle set.		
norised by: C Frearson			DCN: MIC-P-006-12		
e of issue: 07/02/2023			Page 86 of 19		
	THIS DOCUMENT IS LINCO		FD		

Specimen transport	Note: Fluids from existing indwelling drains are not considered to be 'sterile'. As with urinary catheters, drains commonly become colonised and any culture of fluid taken through them may simply reflect colonisation rather than infection. Drain fluid samples should be sent only where there is a high degree of suspicion of infection. Specimens should be sent to the laboratory without delay during normal working hours.
Storage requirements	Outside of normal working hours samples should be refrigerated.
Special requirements	Clinical details are essential for processing. Total cell counts performed on Ascitic fluid SBP patients only.
Laboratory information	
Tests	Presence of white blood cells (quantitative) (ascitic fluid only). Detection of crystals (qualitative).(synovial fluid only). Detection of white blood cells, gram positive and negative bacteria (semi- quantitative). General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).
Measurement units	Cell count x 10 <sup>6</sup> /l
Biological reference units	Total white cell <500 cells x 10 <sup>6</sup> /l count
Turnaround time	Microscopy 2 hours. Culture 5 days.
Availability	Routine hours and on-call.
Clinical information	
Factors known to significantly affect the results	<ul> <li>Small volumes – fluids such as synovial fluids may be received inadequate volumes which may impede the recovery of organisms.</li> <li>Large volumes – specimens such as peritoneal fluid and ascetic fluid may contain very low numbers or organisms which are usually received in adequate quantities and require concentration to increase likelihood of successful culture.</li> <li>Cells disintegrate. A delay in transportation may produce a cell count that is not reflective of the clinical situation of the patient.</li> <li>Delays in transportation may affect the recovery of pathogens.</li> </ul>

# **Genital swab culture (female)**

Sample instructions	HVS, vaginal discharge, vulval swab, labial swab, cervical swab, endocervical swab, urethral swab (Amies transport swab) Optimally collected Genital tract swabs Cervical and high va is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	vulval swab, labial swab, cervical swab, endocervical swab, urethral swab (Amies transport swab) Optimally collected Genital tract swabs Cervical and high va, is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	cervical swab, endocervical swab, urethral swab (Amies transport swab) Optimally collected Genital tract swabs Cervical and high va is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	endocervical swab, urethral swab (Amies transport swab) Optimally collected Genital tract swabs Cervical and high va is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspec	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	urethral swab (Amies transport swab) Optimally collected Genital tract swabs Cervical and high va- is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	transport swab) Optimally collected Genital tract swabs Cervical and high va- is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
Sample instructions	Optimally collected Genital tract swabs Cervical and high va- is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspect	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
	Optimally collected Genital tract swabs Cervical and high va- is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	pefore antimicrobial therapy ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspec	y started. with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
	Genital tract swabs Cervical and high va- is important to avoid posterior fornix, incl pelvic infection, incl swabbed. High vaginal swabs After the introductio the surface of the va	ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspec	with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
	Cervical and high va is important to avoid posterior fornix, incl pelvic infection, incl swabbed. <b>High vaginal swabs</b> After the introductio the surface of the va	ginal swabs should be taken I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspec	with the aid of a speculum. It e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
	is important to avoid posterior fornix, incl pelvic infection, incl swabbed. <b>High vaginal swabs</b> After the introductio the surface of the va	I vulval contamination of the uding any obvious candidal uding gonorrhoea, is suspec	e swab. For Trichomonas, the plaques should be swabbed. If ted, the cervical os should be			
	posterior fornix, incl pelvic infection, incl swabbed. <b>High vaginal swabs</b> After the introductio the surface of the va	uding any obvious candidal uding gonorrhoea, is suspec	plaques should be swabbed. If ted, the cervical os should be			
	pelvic infection, incl swabbed. <b>High vaginal swabs</b> After the introductio the surface of the va	uding gonorrhoea, is suspec	ted, the cervical os should be			
	High vaginal swabs After the introduction the surface of the va	n of the speculum, the swal				
	After the introduction the surface of the value of the va	n of the speculum the swal				
	the surface of the va		Fight vaginal swaps After the introduction of the speculum, the swah should be rolled firmly over			
	the surface of the va	the surface of the vaginal vault				
Collection	Cervical swabs					
	After introduction of	the speculum to the vaging	the swah should be rotated			
	inside the endocervi	v	i, the swab should be rotated			
	Urethral swabs	···				
	Contamination with	micro-organisms from the v	ulva should be avoided. Thin			
	swabs are available for collection of specimens. The patient should not have					
	passed urine for at least one hour.					
	Please send endoce	vical swab if gonococcal cul	ture is required.			
	Separate samples should be collected into appropriate transport media for					
	detection of viruses	or <u>C. trachomatis</u> .				
Specimen transport	Specimens should be	e sent to the laboratory with	nout delay during normal			
specimen transport	working hours.					
Storage requirements	Outside of normal w	orking hours samples shoul	d be refrigerated.			
Storage requirements	Delays of over 48 ho	urs are undesirable.				
Special requirements	Clinical details are estimated	ssential for processing.				
special requirements	Female genital swabs for gonococcal investigation should not be refrigerated.					
Laboratory information						
	Presence of white b	ood cells, red blood cells, ep	oithelial cells, candida,			
Tests	Trichomonas vaginialis, clue cells (quantitative).					
	General Isolation an	a characterisation of aerobio	c, microaerophilic and			
Measurement units	Growth detected or	not detected.				

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

Biological reference units	
Turnaround time	4 days.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	Delays in transportation may affect the recovery of pathogens. Female genital swabs for gonococcal investigation should not be refrigerated as this significantly reduces the recovery rate. Delays in transportation may reduce the recovery of Neisseria gonorrhoea.

# Genital specimens (excluding female genital swabs)

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
OBCANNELIUN OB	Penile swab, urethral swab, screening swabs for Neisseria gonorrhoea (Amies transport swab)			
	Intra-uterine contraceptive device (IUCD)	Entire device should be sent		
	Collection of pus or exudate	Minimum volume 1ml		
Sample instructions				

Optimally collected before antimicrobial therapy started.

#### **Urethral swabs**

	Contamination with micro-organisms from the vulva or the foreskin should be avoided. Thin swabs are available for collection of specimens. The patient should not have passed urine for at least one hour. For males, if a discharge is not apparent, attempts should be made to "milk" exudate from the penis. The swab is gently passed through the urethral meatus and rotated.		
Collection	Intrauterine contraceptive devices (IUCDs) The entire device should be sent.		
Conection	Rectal swabs Rectal swabs are taken via a proctoscope.		
	<b>Throat swabs</b> Throat swabs should be taken from the tonsillar area and/or posterior pharynx avoiding the tongue and uvula.		
	<b>Fluids and pus</b> These are taken from the fallopian tubes, tubo-ovarian and Bartholin's abscesses, etc, taken during surgery.		
	Separate samples should be collected into appropriate transport media for detection of <u>viruses</u> or <u>C. trachomatis</u> .		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Clinical details are essential for processing. Genital swabs for gonococcal investigation should not be refrigerated.		
Laboratory information			
Tests	Detection of white blood cells, gram positive and negative bacteria (semi- quantitative) (fluids and pus only). General isolation and characterisation of aerobic, microaerophilic and		
Authorised by: C Frearson Date of issue: 07/02/2023	DCN: MIC-P-006-12.7 Page 90 of 192		

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

	anaerobic micro-organisms (qualitative).
Measurement units	Growth detected or not detected.
Biological reference units	
Turnaround time	4 days.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	Delays in transportation may affect the recovery of pathogens. Genital swabs for gonococcal investigation should not be refrigerated as this significantly reduces the recovery rate. Delays in transportation may reduce the recovery of Neisseria gonorrhoea.

# Haemophilus influenzae PCR

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	CSF	Minimum volume 0.5ml		
Sample instructions				
Collection	<b>Cerebrospinal fl</b> Refer to <u>CSF mic</u>	uid (CSF) roscopy and culture.		
Specimen transport	Specimens shoul working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm	al working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details an	Clinical details are essential for processing.		
Laboratory information				
This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are requirTestsparameters analysed in this test and any reference ranges for thes parameters will be displayed on the report when it is returned to t requestor.		entre. Contact the er details are required. The nce ranges for these n it is returned to the		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	vailability Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives r inappropriate tir of organism belo detection of an a New and emergi this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility New and emerging variants may also occur which may not be detected by this assay.		

# Haemophilus influenzae type b IgG

#### Determination of immunity.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special re	quirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

## Helicobacter pylori IgG

Infection with *H. pylori* is associated with peptic ulceration. There is evidence that it may play an important role in non-ulcer dyspepsia.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	Detection of Helicobacter pylori IgG antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

## Hepatitis A virus (HAV) IgG

Used to screen for Hepatitis past infection or immunity. Positive result indicates exposure at some time. Test is performed on the assumption that this is a screening test for immunity. If patient acutely icteric or acute infection suspected then request Hepatitis A IgM.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	Detection of Hepatitis A IgG antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

### Hepatitis A virus (HAV) IgM

For diagnosis of acute Hepatitis A infection (jaundice in adults). Hepatitis A in adults does NOT present as abnormal liver functions. It invariably presents as an acute icteric disease (jaundice). It does not cause chronic disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirer	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	Detection of Hepat	Detection of Hepatitis A IgM antibody (qualitative).		
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

### Hepatitis B virus (HBV) confirmation

Routinely performed on sample if newly detected HBV surface Ag, for confirmatory purposes and to help assess timing and infectivity of disease. Also used to monitor response to treatment.

The test consists of HBV surface antigen, HBV surface antigen confirmation, HBV core total antibody, HBV core IgM, HBV e antigen and HBV e antibody.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Venous blood	2 – 6 mls			
Sample instructions					
Collection	No special rec	No special requirements.			
Specimen transport	Specimens sh working hours	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.				
Special requirements	Clinical details are essential for processing.				
Laboratory information					
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.				
Measurement units					
Biological reference units	Biological reference units				
Turnaround time	14 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significat affect the results	tly Haemolysis.				

### Hepatitis B virus (HBV) core IgG antibody

HBV core antibody serves as a marker of past infection.

Where HBV core antibody is detected, further testing for presence of HBV surface antibody will automatically be performed if sufficient serum.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirements.			
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	Detection of Hepatitis B core IgG antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	tly Haemolysis.			

### Hepatitis B virus (HBV) core IgM antibody

HBV core antibody serves as a marker of past infection. Investigation performed during routine infectious disease screening for patients undergoing infertility treatment.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Venous blood	2 – 6 mls			
Sample instructions					
Collection	No special require	No special requirements.			
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.				
Storage requirements	Outside of normal working hours samples should be refrigerated.				
Special requirements	Clinical details and date of onset are essential for processing.				
Laboratory information					
Tests	Detection of Hepatitis B core IgM antibody (qualitative).				
Measurement units					
Biological reference units	Biological reference units				
Turnaround time	7 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significat affect the results	ntly Haemolysis.				

### Hepatitis B virus (HBV) surface antibody

Test to determine if protective immunity has been achieved following immunisation.

Low levels HBV surface antibody may be found in patients who have past infection.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Venous blood	2 – 6 mls			
Sample instructions					
Collection	No special rec	quirements.			
Specimen transport	Specimens she working hours	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.			
Special requirements	Should be tested 6-8 weeks after final dose of Hepatitis B vaccination. Accurate interpretation of this result is reliant upon detailed vaccination history and clinical details.				
Laboratory information					
Tests	Detection of Hepatitis B surface antibody (qualitative).				
Measurement units	IU/L	IU/L			
Biological reference units	Current national recommendations (as per DOH <u>Green Book</u> ) are that a level of $\geq$ 10 IU/L indicates adequate immunity, although a post vaccination level of $\geq$ 100 IU/L is desirable.				
Turnaround time	7 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significat affect the results	tly Haemolysis.				

### Hepatitis B virus (HBV) surface antigen

For diagnosis of acute or recent hepatitis or carrier state.

If first diagnosis of HBV infection a repeat venous blood sample from patient is required to confirm the result.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Venous blood	2 – 6 mls			
Sample instructions					
Collection	No special requirements.				
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.				
Storage requirements	Outside of normal working hours samples should be refrigerated.				
Special requirements	Clinical details and date of onset are essential for processing.				
Laboratory information					
Tests	Detection of Hepatitis B surface antigen (qualitative).				
Measurement units					
Biological reference units	Biological reference units				
Turnaround time	7 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significat affect the results	ntly Haemolysis.				

## Hepatitis B virus (HBV) viral load (PCR)

Indications for testing:

- Detection of viraemia in patients with chronic hepatitis B infection.
- Investigation of possible transmission of hepatitis B e.g. following exposure to blood or body fluids of an infected patient.
- Monitoring effectiveness of anti-viral therapy in patients with chronic hepatitis B infection.
- Measurement of hepatitis B viral load in e antigen negative hepatitis B infected health care workers who perform exposure prone procedures (Health Service Circular 2000/020).

	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	No special re	quirements.		
Specimen transport	Specimens sh working hour	hould be sent to the laboratory rs.	without delay during normal	
Storage requirements	Outside of no	ormal working hours samples s	hould be refrigerated.	
Special requirements	Clinical detail	s are essential for processing.		
Laboratory information				
Tests	This test is pr laboratory or parameters a parameters v requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hour	Routine hours.		
Clinical information				
Factors known to significa affect the results	False negativ inappropriate antly of organism b detection of a New and eme	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

Authorised by: C Frearson Date of issue: 07/02/2023

#### THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Hepatitis C virus (HCV) antibody

Marker of infection at some time.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirements.			
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	Detection of Hepatitis C antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	tly Haemolysis.			

### Hepatitis C virus (HCV) confirmation

HCV confirmation would only be performed on a HCV positive sample.

The test consists of HCV antibody, HCV antibody confirmation, and may include HCV RNA (qualitative PCR).

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of nor	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on parameters an parameters wi requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significar affect the results	ntly Haemolysis.	Haemolysis.		

## Hepatitis C virus (HCV) genotype

Assay used to determine the HCV genotype of patients known to be HCV positive and who are undergoing treatment.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ements.		
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norma	I working hours samples sh	nould be refrigerated.	
Special requirements	Clinical details are	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is proces laboratory on Tele parameters analy parameters will b requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	False negatives m inappropriate tim of organism below detection of an as New and emergin this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

## Hepatitis C virus (HCV) qualitative PCR

Performed on first positive HCV antibody diagnoses or on all other patients HCV antibody positive to confirm active disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requir	ements.		
Specimen transport	Specimens shoul working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm	al working hours samples sl	nould be refrigerated.	
Special requirements	Clinical details ar	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is proce laboratory on Te parameters anal parameters will l requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	False negatives r inappropriate tir of organism belo detection of an a New and emergi this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

## Hepatitis C virus (HCV) viral load

Quantitative assay used for monitoring patients known to be HCV positive and who are undergoing treatment.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	rmal working hours samples sh	nould be refrigerated.	
Special requirements	Clinical detail	s are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on parameters a parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	False negative inappropriate of organism b detection of a New and eme this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

# Hepatitis D (delta) virus antibody

Only appropriate for patients known to be HBV surface Ag positive.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is pro laboratory on parameters a parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.	y Haemolysis.		
# Hepatitis E virus antibody

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	<b>tly</b> Haemolysis.			

# Herpes simplex virus (HSV) antibody

Used to determine past infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special rec	juirements.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	rmal working hours samples s	should be refrigerated.
Special requirements	Clinical details are essential for processing.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significar affect the results	ntly Haemolysis.		

# Herpes simplex virus (HSV) type 1 and 2 PCR

Examinations offered			
Collection container	Specimen	Sample volume	Request form
Annual Article State Annual State State State State	Lesion swab (virus transport medium)/effected mucous membranes		
	EDTA	2 – 6 mls	
Sample instructions			
Collection	Swab: Send an o membranes. Blood: EDTA, no	orange Aptima swab of vest	icle fluid or affected mucous
Specimen transport	Specimens shou working hours.	Ild be sent to the laborator	y without delay during normal
Storage requirements	Outside of norn	nal working hours samples	should be refrigerated.
Special requirements	Clinical details a For HSV in CSF r	re essential for processing efer to <u>CSF (Cerebro-spinal</u>	<u>I fluid) virology PCR</u> .
Laboratory information			
Tests	Detection of HS HSV PCR from b the laboratory o The parameters parameters will requestor.	V type 1 (HSV-1) and HSV t lood is processed at an ext on Telephone 01793 60479 analysed in this test and a be displayed on the report	ype 2 (HSV-2) nucleic acid. ernal reference centre. Contact 8 if further details are required. ny reference ranges for these : when it is returned to the
Measurement units	Qualitative		
Biological reference units			
Turnaround time	Swab: 7 days Blood: 14 days		
Availability	Routine hours.		
Clinical information			
Factors known to significar affect the results	False negatives inappropriate ti of organism bel detection of an New and emerg this assay.	may occur for a variety of r ming of sample collection, ow the detectable limit of t assay sampling variation w ing variants may also occu	reasons, for example inappropriate sample, presence the assay. Towards the limit of ill result in lower reproducibility. r which may not be detected by

## **Back to index**

Authorised by: C FrearsonDCN: MIC-P-006-12.7Date of issue: 07/02/2023Page 111 of 192THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# HIV-1 and 2 antigen/antibodies and p24 antigen

For diagnosis of HIV infection.

If first diagnosis of HIV infection a repeat venous blood sample from patient is required to confirm the result.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing. All requests for HIV investigations must include the Doctor's signature on the request form.			
Laboratory information				
Tests	Detection of HIV-1 and 2 antigen/antibodies plus p24 antigen (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.	ly Haemolysis.		

# **HIV confirmation**

HIV confirmation would only be performed on a HIV positive sample.

The test consists of HIV antigen/antibody confirmation, HIV antigen and HIV antibody, and may include a HIV line immunoassay.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	ould be sent to the laborator 5.	y without delay during normal	
Storage requirements	Outside of nor	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on parameters ar parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significar affect the results	ntly Haemolysis.			

## HIV resistance, integrase, tropism

HIV resistance markers would only be performed on a HIV positive sample. This test is exclusively only available to the Great Western Hospital Sexual Health department.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	EDTA blood	2 – 6 mls			
Sample instructions					
Collection	No special requir	ements.			
Specimen transport	Specimens shoul working hours.	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of norma	al working hours samples sl	nould be refrigerated.		
Special requirements	Clinical details ar	e essential for processing.			
Laboratory information					
Tests	This test is proce laboratory on Te parameters analy parameters will b requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units					
Biological reference units					
Turnaround time	14 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significat affect the results	False negatives n inappropriate tin of organism belo detection of an a New and emergin this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.			

# HIV vertical transmission (neonates)

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	EDTA blood	2 – 6 mls	
Sample instructions			
Collection	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	ormal working hours samples s	hould be refrigerated.
Special requirements	<ul> <li>Requires:</li> <li>a single maternal EDTA at birth</li> <li>neonatal EDTA samples at birth, 3, 6 and 9 months of age.</li> </ul>		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hour	S.	
Clinical information			
Factors known to significar affect the results	ntly		

# **HIV viral load**

HIV viral load would only be performed on a HIV positive sample. This test is exclusively only available to the Great Western Hospital Sexual Health department.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	No special requir	ements.		
Specimen transport	Specimens shoul working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norma	al working hours samples s	should be refrigerated.	
Special requirements	Clinical details ar	Clinical details are essential for processing.		
Laboratory information				
Tests	Detection of HIV	Detection of HIV viral copies (Quantitative )		
Measurement units	Copies / ml	Copies / ml		
Biological reference units				
Turnaround time	48 hours	48 hours		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	False negatives n inappropriate tin ntly of organism belo detection of an a New and emergin this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

# Human herpes virus 6 (HHV) PCR

For diagnosis of HHV infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	No special requirer	nents.		
Specimen transport	Specimens should I working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details are	essential for processing.		
Laboratory information				
Tests	This test is process laboratory on Telep parameters analyse parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	False negatives ma inappropriate timir of organism below detection of an ass New and emerging this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

# Human T lymphotrophic virus (HTLV) 1 and 2 serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	rmal working hours samples	should be refrigerated.	
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significar affect the results	ntly Haemolysis.			

# Hydatid serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	rmal working hours samples	should be refrigerated.	
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significar affect the results	ntly Haemolysis.			

# Influenza A/B rapid PCR

Diagnosis of acute disease. (Clincally suspected influenza cases - Hospital in-patients only)

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Virus swab in transport media	Throat swab	
Sample instructions			
Collection	No special requirer	nents.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.		
Laboratory information			
Tests	Influenza A/B rapic	PCR test	
Measurement units			
Biological reference units			
Turnaround time	2 hours		
Availability	Routine hours.		
Clinical information			
Factors known to significat affect the results	ntly		

# **JC virus PCR**

Diagnosis of acute disease.

Collection container	Specimen	Sample volume	Request form
	EDTA blood	2 – 6 mls	
	Urine	Minimum volume 5ml	
	CSF	Minimum volume 0.5ml	
Sample instructions			
Collection	Cerebrospinal fluid Refer to <u>CSF microso</u> Urine Refer to <u>Urine (micr</u>	(CSF) copy and culture.	
Specimen transport	specimens should b working hours.	e sent to the laboratory with	lout delay during normal
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are e	essential for processing.	
Laboratory information			
Tests	This test is processe laboratory on Telep parameters analyse parameters will be o requestor.	ed at an external reference ce hone 01793 604798 if furthe d in this test and any referen displayed on the report wher	entre. Contact the r details are required. The ice ranges for these n it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significant affect the results	False negatives may inappropriate timin of organism below t detection of an assa New and emerging this assay.	voccur for a variety of reasor g of sample collection, inapp the detectable limit of the as ay sampling variation will resu variants may also occur whic	ns, for example ropriate sample, presence say. Towards the limit of ult in lower reproducibility. h may not be detected by
orised by: C Frearson			DCN: MIC-P-00
of issue: 07/02/2023			Page 121

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Back to index Legionella urinary antigen

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Urine	Minimum volume 5ml		
	Urine	Minimum volume 1ml		
Sample instructions				
Collection	A minimum of 5m If less than 5ml o white topped uni Refer to <u>Urines (</u> N	nl is required. f urine is anticipated, or collecti versal container. Microscopy and Culture).	ng from a child, collect in to a	
Specimen transport	Specimens should working hours.	d be sent to the laboratory with	out delay during normal	
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.			
Special requirements	Clinical details are essential for processing. The British Thoracic Society do not recommend testing unless moderate to high severity pneumonia in hospitalised patients. Will be tested only if clinical details indicate severe pneumonia on request form or where epidemiologically indicated (e.g. atypical features or associated with known <i>Legionella</i> outbreak)			
Laboratory information				
Tests	Detection of Legionell	onella pneumophila antigen (qu a pneumophila serotype 01 only	alitative). /.	
Measurement units	Antigen detected	or not detected.		
Biological reference units				
Turnaround time	1 day.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly			

## Back to index

Authorised by: C Frearson Date of issue: 07/02/2023

# Leptospira serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requ	uirements.		
Specimen transport	Specimens sho working hours.	uld be sent to the laborato	ry without delay during normal	
Storage requirements	Outside of norr	mal working hours samples	should be refrigerated.	
Special requirements	State date of or for processing.	State date of onset, nature of symptoms and exposure history are essential for processing.		
Laboratory information				
Tests	This test is proo laboratory on T parameters and parameters wil requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	ntly Antibody detec Haemolysis.	Antibody detection earliest at 7 days post onset of symptomatic disease. Haemolysis.		

# Measles (diagnostic)

To determine recent/acute disease. For patients who present later into the rash phase of illness.

Diagnosis of measles can usually be made clinically. Characteristic 3-5 days prodromal illness of fever, coryzal symptoms, cough and conjunctivitis. Maculo-papular rash then develops starting behind the ears and spreading down to trunk and arms. Viral shedding from upper respiratory tract is highest from 4 days before to 4 days post onset of rash.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirer	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significan affect the results	Haemolysis.			

# Measles IgG (immunity)

To determine serological evidence of past infection/vaccination where history is uncertain.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requiren	nents.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	No special requirements.			
Laboratory information				
Tests	Detection of Measl	es IgG antibody (semi-quanti	tative).	
Measurement units	AU/mL			
Biological reference units	<13.5 – Susceptible 13.5-16.5 – Equivocal, treat as susceptible >16.5 – Immune			
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	tly Haemolysis.			

# Meningococcal antibody

Neisseria meningitidis functional antibody to serogroups A, C, W, Y and B.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	4 weeks.			
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	Haemolysis.			

# Meningococcal PCR

Meningococcal DNA detection by PCR. Discuss all molecular/PCR requests with Microbiology Consultant or Senior Laboratory Biomedical Scientist.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
	CSF	Minimum volume 0.5ml		
Sample instructions				
Collection	<b>Cerebrospinal fluid</b> Refer to <u>CSF micros</u>	Cerebrospinal fluid (CSF) Refer to <u>CSF microscopy and culture</u> .		
Specimen transport	Specimens should I working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	Outside of normal working hours samples should be refrigerated.		
Special requirements	Where a CSF sample is available, this should be sent in addition to an EDTA blood sample. Clinical details are essential for processing.			
Laboratory information				
Tests	This test is process laboratory on Telep parameters analyse parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	The likelihood of a ntly starting antibiotics after commenceme results. CSF may re	The likelihood of a positive result decreases as the interval of sampling after starting antibiotics lengthens. Samples for PCR taken more than 48 hours after commencement of antibiotic therapy are unlikely to give useful results. CSF may remain "positive" for longer periods.		

#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

# Mouth swab

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Mouth swab (Amies transport swab)				
Sample instructions					
Collection	Optimally collected To assure that the comparable it is ad Eat or drink wi Brush their tee Use any mouth Sample pus if prese A tongue depresso contamination from	before antimicrobial ther preconditions of the samp vised that patients should thin 2 hours th within 2 hours n rinse of disinfectant with ent otherwise sample any l r or spatula may be helpfu n other parts of the mouth	apy started. ling for oral infections are not: in 2 hours prior to sampling esions or inflamed areas. I to aid vision and avoid h.		
Specimen transport	Specimens should I working hours.	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal Delays of over 48 h	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.			
Special requirements	No special requirer	No special requirements.			
Laboratory information					
Tests	General isolation a anaerobic micro-or	nd characterisation of aero ganisms (qualitative).	bbic, microaerophilic and		
Measurement units	Growth detected o	r not detected.			
Biological reference units					
Turnaround time	4 days.				
Availability	Routine hours.				
Clinical information					
Factors known to significan affect the results	tly Delays in transport	ation may affect the recov	ery of pathogens.		

## **MRSA**

Most MRSA infections are healthcare-associated, but an increasing number of infections are communityacquired, with patients having no established risk factors for acquisition of MRSA.

Collection container S	pecimen	Sample volume	Request form
Ν	lose swab, groin swab,		Admission screen:
p	erineum swab,		
espectadore Marter (G)(2)   m	nanipulated wound site		
S	wabs (Amies transport		1 - * - ***** Displantation
S	wab)		— Discharge screen:
	Irine	Minimum volume 1ml	
Sample instructions			
	Optimally collected b	pefore antimicrobial therapy	v started.
	MRSA screen swabs	should be obtained from no	se, groin/perineum and other
	wounds, skin lesions	or invasive devices. Specim	ens from other sites will be
	rejected.		
Collection	Only one request for	m needs to be sent per pati	ent.
	Refer to GWH Trust	MRSA Policy.	
	Urine		
	Refer to Urine (micro	oscopy and culture).	
	Specimens should be	e sent to the laboratory with	out delay during normal
Specimen transport	working hours.	,	, 0
Storago roquiromonto	Outside of normal w	orking hours samples should	d be refrigerated.
Storage requirements	Delays of over 48 ho	urs are undesirable.	
Special requirements	No special requirem	ents.	
Laboratory information			
Tests	General isolation and	d characterisation of MRSA	(qualitative).
Measurement units	Growth detected or	not detected.	
Biological reference units			
Turnaround time	Negative results 24 h	nours.	
	Positive results 3 day	/S.	
Availability	Routine hours.		
Clinical information			
Factors known to significantly affect the results	Delays in transporta	tion may affect the recover	y of pathogens.
ack to index			

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Mumps (diagnostic)

Used to determine disease progression in individuals infected with mumps.

Examinations offered					
Collection container	Specimen	Sample volume	Request form		
	Venous blood	2 – 6 mls			
Sample instructions					
Collection	No special rec	No special requirements.			
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing.				
Laboratory information					
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.				
Measurement units					
Biological reference units					
Turnaround time	14 days.				
Availability	Routine hours	Routine hours.			
Clinical information					
Factors known to significat affect the results	ntly Haemolysis.				

# Mumps IgG (immunity)

Used to determine immune status to mumps.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requiren	nents.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	No special requirements.			
Laboratory information				
Tests	Detection of Mump	os IgG antibody (semi-quantit	ative).	
Measurement units	AU/mL			
Biological reference units	<9.0 – Susceptible 9.0-11.0 – Equivocal, treat as susceptible >11.0 – Immune			
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	tly Haemolysis.			

# Mycobacteria

Collection container	Specimen	Sample volume	Request form	
	Sputum, gastric washing, sterile site body fluids (CSF, pleural fluids etc), skin or tissue biopsies, bone marrow, bronchoalveolar washings, bone and bone marrow, lymph node and tissue samples	1mL of Sputum 5mL of BAL 6mL of CSF	$M = \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} \sum_{j=1}^{n} \sum_{i=1}^{n} $	
	Urine	Early morning urine on three consecutive days, 250ml container		
	Heparin blood	2 – 6 mls		
Sample instructions				
	contamination. Purulent specimens are best. Three samples of ≥5 be collected approximately 8-24 hours apart with at least one from morning. Samples taken early morning (ie shortly after patient waking) have greatest yield. When the cough is dry, physiotherapy, postural drai inhalation of nebulised saline ('sputum induction') before expector be helpful.			
Collection	Bronchoalveolar lava These may be sent if specimens are AFB so bronchoscope with to Mycobacterium spec preferably 5mL.	age/bronchial washings spontaneous or induced spu mear negative. Note: Contai ap water, which may contair ies, should be avoided. Mini	utum is unavailable or if such mination of the n environmental mum sample size is	
	<b>Urine specimens</b> Whole urine specimens should be collected in the early morning on three consecutive days in a 250ml CE marked leak proof container (that does not contain boric acid), and placed in a sealed plastic bag. Urine specimens received in 20ml universal containers will be rejected.			
	Sterile site body fluid Collect aseptically as	<b>ds</b> much (eg >6mL in adults) CS	F sample as possible If only	

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

a small volume is available after initial lumbar puncture, and the findings of cell counts and protein suggest TB meningitis, a second procedure should be considered to obtain a larger volume to improve chances of achieving positive cultures.

It should be noted that pleural or pericardial fluids are not very sensitive samples for the detection of *M. tuberculosis,* and that a concurrent pleural or pericardial biopsy taken with the fluid is more useful. A negative result on these fluids does not rule out the diagnosis.

#### Lymph node and tissue samples

Send in sterile container. A small amount of sterile water or saline may be added to prevent the sample from dehydrating.

#### **Cerebrospinal fluid (CSF)**

For CSF refer to CSF microscopy and culture.

#### **Blood culture**

In patients where disseminated mycobacterial disease is suspected (e.g. *Mycobacterium avium intracellulare* complex in HIV infected patients) send a peripheral blood sample in a Lithium heparin tube (green top vacuette).

#### The following are specialist tests:

	Molecular tests (PCR)
	Gamma Interferon Tests
	Specimens should be sent to the laboratory without delay during normal
Specimen transport	working hours.
	Do not use pneumatic chute system if investigation for Mycobacteria required.
Storage requirements	Outside of normal working hours samples should be refrigerated.
	For the initial diagnosis of mycobacterial infection all specimens should be
	fresh and taken, whenever possible, before anti-tubercular treatment is
Special requirements	started. 'Other' antimicrobials may also have significant anti-mycobacterial
	activity, notably the fluoroquinolones such as ciprofloxacin, levofloxacin or
	moxifloxacin, and the macrolides such as clarithromycin or azithromycin.
Laboratory information	
	No microscopy performed on urine samples for Mycobacteria investigations.
	If sample volume is insufficient for both microscopy and culture, culture is
	usually preferred to microscopy due to greater sensitivity.
Tests	
	This test is processed at an external reference centre. Contact the laboratory
	on Telephone 01793 604798 if further details are required. The parameters
	analysed in this test and any reference ranges for these parameters will be
	displayed on the report when it is returned to the requestor.
Measurement units	
Biological reference units	
Turnaround time	6 weeks.
norised by: C Frearson	DCN: MIC-P-006-12
e of issue: 07/02/2023	Page 133 of 1
	THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

	Significant positive results are communicated to clinicians as and when they arise.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	EDTA, even in trace amounts, inhibits the growth of some <i>Mycobacterium</i> species. Some antimicrobials have significant anti-mycobacterial activity, notably the fluoroquinolones such as ciprofloxacin, levofloxacin or moxifloxacin, and the macrolides such as clarithromycin or azithromycin.

# Mycobacteria PCR

May be appropriate under certain circumstances. Usually performed on smear positive samples where drug resistance is strongly suspected. Requests must be discussed with Consultant Microbiologist.

Collection container	Specimen	Sample volume	Request form
	Sputum, gastric washing, sterile site body fluids (CSF, pleural fluids etc), skin or tissue biopsies, bone marrow, bronchoalveolar washings, bone and bone marrow, lymph node and tissue samples	1mL of Sputum 5mL of BAL 6mL of CSF	
	Urine	Early morning urine on three consecutive days, 250ml container	
	Heparin blood	2 – 6 mls	
Sample instructions			
Collection Specimen transport	Cerebrospinal fluid ( Refer to <u>CSF microsco</u> Specimens should be working hours. Do not use pneumati	<b>CSF)</b> <u>opy and culture</u> . sent to the laboratory with c chute system if investigation	out delay during normal
Storage requirements	Outside of normal we	orking hours samples should	be refrigerated.
Special requirements	No special requirements.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	2 weeks. Significant positive results are communicated to clinicians as and when they arise.		

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Clinical information	
Factors known to significantly affect the results	EDTA, even in trace amounts, inhibits the growth of some <i>Mycobacterium</i> species. Some antimicrobials have significant anti-mycobacterial activity, notably the fluoroquinolones such as ciprofloxacin, levofloxacin or moxifloxacin, and the macrolides such as clarithromycin or azithromycin.

# Mycology

Infection by dermatophytes is cutaneous and generally restricted to the non-living cornified layers in patients who are immunocompetent. This is because the dermatophyte group of fungi are generally unable to penetrate tissues which are not fully keratinised (ie deeper tissues and organs). However, reactions to such infections can range from mild to severe, depending upon the host's immune response, the virulence of the infecting species, the site of infection and environmental factors.

Collection container	Specimen	Sample volume	Request form
	Skin, hair, nails		
DECEMBENT 2000     Second	Skin, hair, nails		
Sample instructions			
Collection	Sterile Universispecifically for Skin Material from outer edges of a scalpel blade <b>Hair</b> Scalp scraping may be plucket as infection is be transporter <b>Nails</b> Clippings shou cut back as fait the lower part supplement th present. Whol container.	sal or commercially available par the collection and transport o skin lesions is collected by gen f the lesion, usually with the ed e. The edge is most likely to cor s are obtained as above but sh ed from the scalp with forceps, usually below the surface near d to the laboratory as for skin s all be taken from the discolour r as possible from the free edge ts. Scrapings can also be taken f ne clippings. Nail clippings often le nails can be sent to the Labo	ackets e.g. Dermapak, designed f skin, nail and hair samples. tly scraping off material from the lge of a glass microscope slide or ntain viable fungus. ould include hair stubs. Hairs but cut hairs are unsatisfactory the scalp. The material should crapings. ed or brittle parts of the nail and e as some fungi are restricted to from under the nail to n fail to grow fungi even if ratory in a sterile Universal
	Invasive funga BAL, tissue bio Serological tes Consultant.	<b>al disease</b> opsy, blood cultures, CSF, urine sts may be appropriate – please	for culture as clinically indicated. e discuss with the Microbiology
Specimen transport	Specimens sho	ould be transported and proces	ssed as soon as possible.
Storage requirements	Samples shou Provided the s months.	ld be allowed to dry out and ke samples are kept dry, the fungu	pt at room temperature. Is will remain viable for several

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

Special requirements	No special requirements.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	Microscopy 1 week. Culture 4 weeks.		
Availability	Routine hours.		
Clinical information			
Factors known to significantly affect the results			

# Mycology serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	ould be sent to the laborator 5.	y without delay during normal	
Storage requirements	Outside of nor	rmal working hours samples	should be refrigerated.	
Special requirements	Clinical details essential for p	Clinical details and any history of travel or occupational exposure are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on parameters ar parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	ly Haemolysis.			

# Neisseria gonorrhoeae PCR

This test is exclusively only available to the Great Western Hospital Sexual Health department.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
Sector Artest	Eye, cervical, urethral, throat, rectal swab			
Partie	Urine (first void)	Minimum volume 2ml		
Sample instructions				
Collection	Specimens should guidelines on the c Refer to <u>Chlamydia</u> <u>collection of urine</u>	Specimens should be collected and handled following the recommended guidelines on the collection packs. Refer to <u>Chlamydia PCR – collection of vaginal sample</u> and <u>Chlamydia PCR – collection of vaginal sample</u> .		
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	working hours samples shou	ld be refrigerated.	
Special requirements	Urine – patient sho collection.	Urine – patient should not have urinated for 2 hours prior to sample collection.		
Laboratory information				
Tests	Detection of Neisseria gonorrhoeae nucleic acid (qualitative).			
Measurement units	Presence detected or not detected.			
Biological reference units				
Turnaround time	4 days.	4 days.		
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	<ul> <li>False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.</li> </ul>			

# **Norovirus PCR**

Norovirus test performed only on instruction by the Infection Control Team in the investigation of outbreaks.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Stool sample	Liquid specimen: 1 – 2ml Formed specimen: large pea size sample		
Sample instructions				
Collection	Specimen may be p container and tran	bassed into a clean, dry, dispo sferred to an appropriate coll	sable bedpan or similar ection container.	
Specimen transport	Specimens should working hours.	be sent to the laboratory with	out delay during normal	
Storage requirements	Outside of normal	working hours samples should	be refrigerated.	
Special requirements	Clinical details are Repeat samples for Microbiologists wil	Clinical details are essential for processing. Repeat samples for microbiological clearance not usually required – Microbiologists will advise if necessary.		
Laboratory information				
Tests	Detection of Norovirus nucleic acid (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	1 day.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.			

## Nose swab

Nasal colonisation with *Staphylococcus aureus* increases the risk of staphylococcal infections at other sites of the body such as postoperative wounds and dialysis access sites.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Nose swab (Amies transport swab)			
Sample instructions				
Collection	Optimally collecte Plain sterile cotto the swab over the	ed before antimicrobial the n wool swab. Sample the a e mucosal surface.	rapy started. Interior nares by gently rotating	
Specimen transport	Specimens should working hours.	be sent to the laboratory	without delay during normal	
Storage requirements	Outside of norma Delays of over 48	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Nasal swabs shou <u>pertussis</u> .	Nasal swabs should NOT be taken to investigate the presence of <u>Bordetella</u> pertussis.		
Laboratory information				
Tests	General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).			
Measurement units	Growth detected	Growth detected or not detected.		
Biological reference units				
Turnaround time	4 days.	4 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	ntly Delays in transpo	Delays in transportation may affect the recovery of pathogens.		

# Panfungal PCR

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	Minimum volume 500µl		
Sample instructions				
Collection	No special require	ments.		
Specimen transport	Specimens should working hours.	be sent to the laboratory with	nout delay during normal	
Storage requirements	Outside of normal	working hours samples shoul	d be refrigerated.	
Special requirements	Clinical details are	essential for processing.		
Laboratory information				
Tests	This test is process laboratory on Tele parameters analys parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days	14 days		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	False negatives ma inappropriate timi of organism below detection of an as New and emerging this assay.	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.		

# Pan-valentine leukocidin (PVL) toxin detection

Testing for the PVL toxin gene on isolates will be directed by the Consultant Microbiologist, based on clinical presentation and/or antibiotic sensitivity patterns. Generally, PVL toxin testing will be carried out on the following:

- S.aureus cultured from individuals with recurrent boils/abscesses
- S.aureus cultured from individuals with necrotising skin and soft tissue infections
- S.aureus pneumonia
- Ciprofloxacin sensitive MRSA
- Any other S.aureus isolate as indicated by the Consultant Microbiologist

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	S.aureus isolated by laboratory, as directed by Consultant Microbiologist			
Sample instructions				
Collection	No special require	ements.		
Specimen transport	Specimens should working hours.	be sent to the laboratory	without delay during normal	
Storage requirements	Outside of normal	l working hours samples s	hould be refrigerated.	
Special requirements	No special require	ements.		
Laboratory information				
Tests	Detection of PVL toxin nucleic acid (qualitative): This test is processed at an external reference centre.			
Measurement units				
Biological reference units				
Turnaround time	4 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	False negatives may occur for a variety of reasons, for example inappropriate timing of sample collection, inappropriate sample, presence of organism below the detectable limit of the assay. Towards the limit of detection of an assay sampling variation will result in lower reproducibility. New and emerging variants may also occur which may not be detected by this assay.			

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 144 of 192
	THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED	
# Parasitology (Bilharzia)

Diagnosis of acute infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Urine sample	Sample collected between 1000 and 1400. Alternatively a 24hr collection of terminal samples of urine may be obtained.		
Sample instructions				
	First 3 months pos water exposure in terminal urine three stool sar 3 months or more terminal urine three stool sar clotted blood	t exposure, if suspecting schist endemic area, send: – not mid-stream nples, 2 days apart <b>post exposure</b> : – not mid-stream nples, 2 days apart for <u>Schistosoma serology</u>	osomiasis and has fresh	
Collection	Send also a FBC for	Send also a FBC for detection of eosinophilia.		
	Collect a urine spec concentration of ep Ask patient to urin voided and collect 20ml of urine) in a Alternatively a 24h It is also recommen specimen is collect of stairs).	cimen between 1000 and 1400 ggs is found. ate as normal. Halt the process the remaining end-stream urin sterile container. Send 3 such r collection of terminal samples nded that a little light exercise s ed (e.g. 20 rapid knee bends, o	, as this is when the highest before bladder completely e sample (the last 10 to samples. s of urine may be obtained. should be taken before the r running up & down a flight	
Specimen transport	Specimens should working hours.	be sent to the laboratory witho	ut delay during normal	
Storage requirements	Outside of normal Delays of over 48 h	working hours samples should nours are undesirable.	be refrigerated.	
Special requirements	Please provide info	ormation regarding recent forei	gn travel.	
Laboratory information				
Tests	Presence of Schisto	osoma haematobium (qualitativ	/e).	
Measurement units				

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 145 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Biological reference units		
Turnaround time	2 days.	
Availability	Routine hours.	
Clinical information		
Factors known to significantly affect the results		

# Parasitology (Pinworm)

Diagnosis of acute infection.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
Please contact the laboratory on 01793 604798 for collection kits	Sellotape from perianal region		
Sample instructions			
Collection	Please contact the la "Sellotape" slides ar should be carried ou firmly against the pe	aboratory on 01793 604 e used in the diagnosis o It first thing in the morn prianal skin.	798 for collection kits. of threadworm and the procedure ing. Press the sticky middle 1-2"
Specimen transport	Specimens should be working hours.	e sent to the laboratory	without delay during normal
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	No special requirem	ents.	
Laboratory information			
Tests	Presence of Enterob	ius vermicularis ova (qu	alitative).
Measurement units			
Biological reference units			
Turnaround time	2 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significar affect the results	ntly		

# Parasitology (serology)

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	Clotted blood	sample – at least 12 weeks po	ost exposure.	
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Please include and travel his Send stool sa	Please include relevant clinical details, including reason for investigations and travel history. Send stool sample.		
Laboratory information				
Tests	This test is pro- laboratory on parameters a parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Persist for sev Haemolysis.	ay take up to 3 months to dev veral months after successful t	elop. Once detectable may reatment	

## Parasitology (Stool)

There is no need to request OCP for Cryptosporidium and Giardia lamblia; all stool samples for culture will be automatically tested for these.

Information required for other parasitic infections:

- Foreign travel history
- Blood eosinophil count
- Duration of diarrhoea
- Presence/absence of abdominal symptoms
- Evidence of malabsorption

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Stool sample	3 stool samples over a period of 10 days. Liquid specimen: 1 – 2ml Formed specimen: large pea size sample.	
Sample instructions			
Collection	Specimen may be pa container and transf	ssed into a clean, dry, disposa erred to an appropriate colled	able bedpan or similar ction container.
Specimen transport	Specimens should be working hours.	e sent to the laboratory witho	ut delay during normal
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	For examination of amoebic trophozoites the specimen must reach the laboratory within 1 hour of its production. It is advisable to arrange this examination with the Departments in advance.		
Laboratory information			
Tests	Presence and identif	ication of ova and parasites (	qualitative).
Measurement units			
Biological reference units			
Turnaround time	4 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significan affect the results	tly		

#### **Back to index**

Authorised by: C FrearsonDCN: MIC-P-006-12.7Date of issue: 07/02/2023Page 149 of 192THIS DOCUMENT IS UNCONTROLLED WHEN DRINTED

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Parasitology (Worm identification)

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Worm	Please send actual worm seen	
Sample instructions			
Collection	Please send a	ctual worm seen.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	No special requirements.		
Laboratory information			
Tests	Parasite ident	ification (qualitative).	
Measurement units			
Biological reference units			
Turnaround time	2 days.		
Availability	Routine hours.		
Clinical information			
Factors known to significar affect the results	ntly		

### **Parvovirus PCR**

Diagnosis of acute disease. DNA detection may be indicated if significant immuno-suppression (e.g. HIV disease or organ transplant).

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	No special re	quirements.		
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	ormal working hours samples sl	hould be refrigerated.	
Special requirements	No special re	quirements.		
Laboratory information				
Tests	This test is pr laboratory or parameters a parameters v requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hour	S.		
Clinical information				
Factors known to significat affect the results	False negativ inappropriate of organism b detection of New and emo this assay.	es may occur for a variety of re e timing of sample collection, ir below the detectable limit of th an assay sampling variation wil erging variants may also occur	easons, for example nappropriate sample, presence ne assay. Towards the limit of I result in lower reproducibility. which may not be detected by	

### Parvovirus serology

Please state whether test required for acute disease (IgM/DNA) or if evidence of past exposure (immunity) required (IgG).

IgM is usually positive at time of presentation with acute symptoms. May remain detectable for up to 3 months.

DNA detection may be indicated if significant immuno-suppression (e.g. HIV disease or organ transplant).

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirer	nents.		
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details and Indicate if patient i exposure.	Clinical details and date of onset are essential for processing. Indicate if patient is pregnant and gestation, and date of contact or exposure.		
Laboratory information				
Tests	This test is process laboratory on Teley parameters analyse parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significan affect the results	tly Haemolysis.			

#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

DCN: MIC-P-006-12.7 Page 152 of 192

# Peritoneal dialysis fluid (PDF)

Collection container	Specimen	Sample volume	Request form	
	Peritoneal dialysis fluid	Minimum volume 1ml		
		Inoculate up to 10ml in each bottle		
Sample instructions				
Collection	Blood culture both	tles ture Method Options		
Specimen transport	Specimens should working hours.	be sent to the laboratory wit	hout delay during normal	
Storage requirements	Outside of normal	working hours samples shou	ld be refrigerated.	
Special requirements	No special require	No special requirements.		
Laboratory information				
Tests	Presence of white Detection of gram General isolation a anaerobic micro-o	blood cells (quantitative). positive and negative bacter and characterisation of aerob rganisms (qualitative).	ia (semi-quantitative). ic, microaerophilic and	
Measurement units	Cell count x 10 <sup>6</sup> /l Growth detected c	or not detected.		
Biological reference units	Total white cell count	<500 cells x 10 <sup>6</sup> /l		
Turnaround time	Microscopy 2 hour Culture 5 days.	rs.		
Availability	Routine hours and	l on-call.		
Clinical information				
Factors known to significant affect the results	Large volumes of f are usually receive increase likelihood Cells disintegrate. not reflective of th Delays in transpor	iluid may contain very low nu ed in adequate quantities and d of successful culture. A delay in transportation ma ne clinical situation of the pat tation may affect the recover	mbers or organisms which require concentration to y produce a cell count that is ient. y of pathogens	
ack to index	- ,	.,	, <u>, , , , , , , , , , , , , , , , , , </u>	
-				

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

### **Pneumococcal PCR**

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	Minimum volume 5ml		
	CSF	Minimum volume 0.5ml		
Sample instructions				
Collection	<b>Cerebrospinal fluic</b> Refer to <u>CSF micros</u>	l <b>(CSF)</b> scopy and culture.		
Specimen transport	Specimens should I working hours.	pe sent to the laboratory with	out delay during normal	
Storage requirements	Outside of normal	working hours samples should	d be refrigerated.	
Special requirements	Clinical details are	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is process laboratory on Telep parameters analyse parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	False negatives ma inappropriate timir ntly of organism below detection of an ass New and emerging this assay.	y occur for a variety of reasor og of sample collection, inapp the detectable limit of the as ay sampling variation will reso variants may also occur whic	ns, for example ropriate sample, presence say. Towards the limit of ult in lower reproducibility. h may not be detected by	

# Pneumococcal serology

Used to determine immunity.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requi	rements.		
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details a	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is proce laboratory on Te The parameters parameters will requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	Haemolysis.			

# Pneumococcal urinary antigen

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Urine	Minimum volume 5ml		
	Urine	Minimum volume 1ml		
Sample instructions				
Collection	A minimum of 5ml is If less than 5ml of ur white topped univer Refer to <u>Urines (Mic</u>	s required. ine is anticipated, or collectin sal container. roscopy and Culture).	g from a child, collect in to a	
Specimen transport	Specimens should be working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal w Delays of over 48 ho	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Clinical details are es The British Thoracic high severity pneum Will be tested only if form.	Clinical details are essential for processing. The British Thoracic Society do not recommend testing unless moderate to high severity pneumonia in hospitalised patients. Will be tested only if clinical details indicate severe pneumonia on request form.		
Laboratory information				
Tests	Detection of Pneum	ococcal antigen (qualitative).		
Measurement units	Antigen detected or	not detected.		
Biological reference units				
Turnaround time	1 day.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	ntly Pneumococcal vacci	nation within previous week r	nay give positive result.	

# Pneumocystis jirovecii (IF)

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Sputum/BAL	Minimum volume 1ml		
Sample instructions				
Collection	<b>Sputum speci</b> Refer to <u>Respi</u>	mens/ bronchoalveolar lavage/b ratory samples for culture.	ronchial washings	
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of nor	rmal working hours samples shou	ld be refrigerated.	
Special requirements	Clinical details	are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on parameters ar parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significa affect the results	False negative inappropriate of organism be detection of a New and eme this assay.	es may occur for a variety of reasc timing of sample collection, inap elow the detectable limit of the a n assay sampling variation will re rging variants may also occur whi	ons, for example propriate sample, presence ssay. Towards the limit of sult in lower reproducibility. ch may not be detected by	

Note: This test is not accredited by UKAS 15189

### **Pseudomonas serology**

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requ	uirements.		
Specimen transport	Specimens sho working hours.	uld be sent to the laborato	ry without delay during normal	
Storage requirements	Outside of norr	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

# **Q** fever serology

Used to determine past or current infection.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special req	uirements.		
Specimen transport	Specimens sho working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of nor	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details	Clinical details are essential for processing.		
Laboratory information				
Tests	This test is pro laboratory on The paramete parameters wi requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.	Haemolysis.		

# Quantiferon gold TB

These tests are used primarily for the diagnosis of latent infection in the context of contact tracing. They do not differentiate between latent and active disease.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	Please contac Refer to <u>Quan</u>	t the laboratory on 01793 60 <u>ntiferon TB Gold - Instruction</u>	04798 for collection kits. <u>s for Specimen Collection</u> .
Specimen transport	All quantiferon samples should be returned directly to the microbiology laboratory within 2 hours of sample being taken.		
Storage requirements	See above.		
Special requirements	Clinical details are essential for processing. Samples will be receipted Monday – Thursday up to 1700 hrs. Please return samples in the box supplied with a completed request form.		
Laboratory information			
This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for thes parameters will be displayed on the report when it is returned to the requestor.			ence centre. Contact the further details are required. Iny reference ranges for these t when it is returned to the
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Due to pre-analytical requirements, quantiferon test kits are only availabl Monday – Thursday between 0845 and 1500.		
<b>Clinical information</b>			
Factors known to significat affect the results	ntly		

#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

### **Quantiferon TB Gold - Instructions for Specimen Collection**

#### ALL QUANTIFERON SAMPLES SHOULD BE RETURNED DIRECTLY TO THE MICROBIOLOGY LABORATORY WITHIN 2 HOURS OF SAMPLE BEING TAKEN.

#### SAMPLES WILL BE RECEIPTED MONDAY – THURSDAY UP TO **1700 HRS**. DUE TO PRE-ANALYTICAL REQUIREMENTS, QUANTIFERON TEST KITS WILL ONLY BE AVAILABLE MONDAY – THURSDAY BETWEEN 0845 AND 1500

#### PLEASE RETURN SAMPLES IN THE BOX SUPPLIED WITH A COMPLETED REQUEST FORM.

#### Specimen Collection

The Quantiferon-TB Gold uses the following collection tubes:-

Grey Cap	Nil Control
Green Cap	TB Antigen 1
Yellow Cap	TB Antigen 2
Purple Cap	Mitogen Control

The black mark on the side of these tubes indicates a 1ml fill volume.

For each patient, collect 1 ml of blood by venepuncture directly into **EACH** of the 3 tubes contained within the Quantiferon kit. Ensure <u>each</u> tube is labelled with patient details.

The Quantiferon kit tubes will draw blood relatively slowly, and so it is important that the needle is kept on the tube for 2-3 seconds once the tube appears to have completed filling. This is to ensure that the correct volume is drawn.

If a butterfly needle is being used to collect blood, a purge tube should be used to ensure that the tubing is filled with blood prior to the quantiferon TB tubes being used.

Mix the tubes by turning the tube end over end 8 - 10 times, or shaking the tube for 5 seconds ensuring that the entire inner surface of the tube has been coated with blood. Thorough mixing is required to ensure complete mixing of the blood with the tube's contents.

# **Respiratory samples for culture**

Collection container	Specimen	Sample volume	Request form	
	Bronchial aspirate, transthoracic aspirate, bronchoalveolar lavage, transtracheal aspirate, bronchial brushings, protected catheter specimens, bronchial washings, endotracheal tube specimens, sputum –	Minimum volume 1ml		
	expectorated			
Sample instructions				
Collection	therapy started. <b>Sputum specimens</b> Sputum specimens sl contamination. Puru (ie shortly after patie dry, physiotherapy, p ('sputum induction') <b>Bronchoalveolar lava</b> These may be sent if Minimum sample size A BAL is required for infection. For <u>Legionella</u> or <u>Pne</u> sample in a plain univ	nould be relatively fresh (les lent specimens are best. Sa ont waking) have the greates oostural drainage or inhalati before expectoration may b age/bronchial washings spontaneous or induced sp e is preferably 5mL. microbiological diagnosis o <u>sumococcal antigen</u> is to be versal container.	ss than 1 day old) to minimise amples taken early morning st yield. When the cough is on of nebulised saline be helpful. utum is unavailable. f invasive fungal respiratory excluded, please send a urine	
	alveolar lavage (BAL) co-infected with HIV.	alveolar lavage (BAL) is required. Induced sputum is acceptable in patients co-infected with HIV.		
Specimen transport	Specimens should be working hours.	sent to the laboratory with	out delay during normal	
Storage requirements	Outside of normal we Delays of over 48 hou	orking hours samples should urs are undesirable.	d be refrigerated.	
	Calivary engeimone a	re not processed on the bas	is of macroscopic description	

Aut Date of issue: 07/02/2023

2.7 Page 162 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Laboratory information	
Tests	General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative). Refer to <u>Cystic fibrosis</u> for cough swab specimens. Extended culture for <i>Burkholderia cepacia</i> performed where requests indicate Cystic Fibrosis.
Measurement units	Growth detected or not detected.
Biological reference units	
Turnaround time	4 days.
Availability	Routine hours.
Clinical information	
Factors known to significantly affect the results	All samples are suitable for overnight refrigeration only, they must not be stored over a weekend. Delays in transportation may affect the recovery of pathogens. Sputum may be refrigerated for up to 2-3 h without an appreciable loss of pathogens. Any delay beyond this time may allow overgrowth of Gram- negative bacilli, and Haemophilus species and <i>S. pneumoniae</i> may be rendered non-viable.

# **Respiratory syncytial virus (RSV)**

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	NPA	Minimum volume 1ml		
Sample instructions				
Collection	No special	requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing.			
Laboratory information				
Tests	RSV PCR te	st		
Measurement units				
Biological reference units				
Turnaround time	2 hours			
Availability	Routine hours.			
Clinical information	Clinical information			
Factors known to significa affect the results	ntly			

## **Respiratory virus PCR**

Respiratory screen for at risk patient groups including:

- Influenza A inc H1N1 (avian types: contact lab)
- Influenza B
- Parainfluenza viruses 1,2,3
- Respiratory syncytial virus
- Metapneumovirus
- Adenovirus
- Rhinovirus

. . .

Collection container	Specimen	Sample volume	Request form	
	Nose and/or throat swab (virus transport medium)			
	Sputum/BAL	Minimum volume 1ml		
	NPA	Minimum volume 1ml		
Sample instructions				
Collection	Sputum specimens Refer to <u>Respirator</u>	/ bronchoalveolar lavage/b y samples for culture.	bronchial washings	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details are essential for processing. Please contact the laboratory (Telephone 01793 604798) if urgent processing for PCP is required.			
Laboratory information				
Tests	This test is processe laboratory on Telep parameters analyse parameters will be requestor.	ed at an external reference whone 01793 604798 if furth ed in this test and any refere displayed on the report who	centre. Contact the ner details are required. The ence ranges for these en it is returned to the	
Measurement units				
Biological reference units				
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
orised by: C Frearson			ηςνι Μις-δ-υ	
of issue: 07/02/2023			Page 16	

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

	False negatives may occur for a variety of reasons, for example
	inappropriate timing of sample collection, inappropriate sample, presence
Factors known to significantly	of organism below the detectable limit of the assay. Towards the limit of
affect the results	detection of an assay sampling variation will result in lower reproducibility.
	New and emerging variants may also occur which may not be detected by
	this assay.

## Rotavirus

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Stool sample	Liquid specimen: 1 – 2ml Formed specimen: large pea size sample		
Sample instructions				
Collection	Specimen may be part container and transfe	Specimen may be passed into a clean, dry, disposable bedpan or similar container and transferred to an appropriate collection container.		
Specimen transport	Specimens should be working hours.	sent to the laboratory witho	ut delay during normal	
Storage requirements	Outside of normal wo Delays of over 48 hou	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Rotavirus test perfor	Rotavirus test performed on samples from children <5 years.		
Laboratory information				
Tests	Rotavirus antigen de	Rotavirus antigen detection (qualitative).		
Measurement units	Growth detected or r	Growth detected or not detected.		
Biological reference units				
Turnaround time	2 days.	2 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significan affect the results	Collect specimens be Specimens should be A positive rotavirus la likely to reflect vaccir	fore antimicrobial therapy whe transported and processed a aboratory result within 15 day nation status and NOT active	nere possible. Is soon as possible. ys of Rotarix vaccination is infection.	

# Rubella (diagnostic)

Used to determine disease progression in individuals infected with rubella.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Please indicate if patient is pregnant and gestation with contact history.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hour	S.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

# Rubella IgG (immunity)

Test is for evidence of past exposure or vaccination/immunity (IgG).

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requirements.			
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Please indicate if patient is pregnant and gestation with contact history.			
Laboratory information				
Tests	Detection of Rubella IgG antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	Haemolysis.			

## Syphilis antibody

For diagnosis of acute or recent Syphilis.

If first diagnosis of Syphilis infection a repeat venous blood sample from patient is required to confirm the result.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and CSF sample if neur Microbiologist.	Clinical details and date of onset are essential for processing. CSF sample if neurosyphilis suspected – discuss with the Consultant Microbiologist.		
Laboratory information				
Tests	Detection of Treponema pallidum antibody (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

## Syphilis confirmation

Syphilis confirmation would only be performed on a Syphilis positive sample. The test consists of RPR titre, T.pallidum particle agglutination test, Syphilis total antibody and Syphilis IgM.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hour	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Syphilis confirmation would only be performed on a Syphilis positive sample. Clinical details are essential for processing.			
Laboratory information				
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.			
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours	5.		
Clinical information				
Factors known to significat affect the results	ntly Haemolysis.			

### **Tetanus antibody**

Tetanus IgG antibody determination.

Examinations offered			
Collection container	Specimen	Sample volume	Request form
	Venous blood	2 – 6 mls	
Sample instructions			
Collection	No special req	uirements.	
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing.		
Laboratory information			
Tests	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units			
Biological reference units			
Turnaround time	14 days.		
Availability	Routine hours		
Clinical information			
Factors known to significat affect the results	ntly Haemolysis.		

### **Throat swab**

Bacterial throat swabs will be routinely cultured for primary pathogens i.e. Groups A, C and G  $\beta$ -haemolytic streptococci. Where other potential pathogens such as *Staph. aureus* are predominant or pure growth, they will be reported.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
Q3D counter territo (1)	Throat swab (Amies transport swab)			
Sample instructions				
Collection	Optimally collecte Throat swab take be taken avoiding	ed before antimicrobial thera n from the tonsillar area and g the tongue and uvula.	apy started. I/or posterior pharynx, should	
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norma Delays of over 48	l working hours samples sho hours are undesirable.	ould be refrigerated.	
Special requirements	Throat swabs sho pertussis. Isolation of Neiss Ideally, inoculatic on to culture med without delay. Tr Culture for <i>Coryi</i> clinical or epidem Anaerobic infecti case please doc anaerobically.	<ul> <li>Throat swabs should NOT be taken to investigate the presence of <u>Bordetella pertussis</u>.</li> <li>Isolation of Neisseria sp only on request.</li> <li>Ideally, inoculation of specimens for <i>N. gonorrhoeae</i> should be made directly on to culture media at the time of collection and these should be incubated without delay. Transport time should be as short as possible.</li> <li>Culture for <i>Corynebacterium diphtheriae</i> is only performed where relevant clinical or epidemiological details are provided.</li> <li>Anaerobic infection can present with very severe symptoms – if this is the case please document on request form and specimen will be cultured and specimen will be specimen will be specimen will be cultured and specimen will be specim</li></ul>		
Laboratory information				
Tests	General isolation anaerobic micro-	and characterisation of aerc organisms (qualitative).	bic, microaerophilic and	
Measurement units	Growth detected	or not detected.		
Biological reference units				
Turnaround time	4 days.	4 days.		
Availability	Routine hours.			
Clinical information				
Factors known to significan affect the results	tly Delays in transpo	ortation may affect the recov	very of pathogens.	

### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023 DCN: MIC-P-006-12.7 Page 173 of 192

# Tips/intravascular cannulae

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Line tips (eg CVP or Hickman lines)	End of cannulae tip (2 – 5 cm in length)		
QID counter ture O()	Swab of cannula insertion sites (Amies transport swab)			
Sample instructions				
Collection	Optimally collected b Tips are preferable to Disinfect the skin aro technique, and cut of proof container using	Optimally collected before antimicrobial therapy started. Tips are preferable to swabs. Disinfect the skin around the cannula entry site, remove cannula using aseptic technique, and cut off 2 – 5 cm of the tip into an appropriate CE marked leak proof container using sterile scissors.		
Specimen transport	Specimens should be working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal wo Delays of over 48 hou	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	Cannulae should only Where line related in and peripheral taken Do NOT send line tips NOT suspected. Urinary catheter tips microbiology investig	Cannulae should only be sent if there is evidence of infection. Where line related infection/sepsis suspected, send blood cultures (central and peripheral taken simultaneously), prior to line removal. Do NOT send line tips if they are being removed routinely and infection is NOT suspected. Urinary catheter tips and drain tips are not appropriate samples for microbiology investigation and will not be processed.		
Laboratory information				
Tests	General isolation and anaerobic micro-orga	characterisation of aerobic, inisms (qualitative).	microaerophilic and	
Measurement units	Growth detected or r	not detected.		
Biological reference units				
Turnaround time	4 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	Delays in transportat	tion may affect the recovery of	of pathogens.	

### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

# **Tissues and biopsies**

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Tissue and biopsies			
Sample instructions				
Collection	Optimally collected	ed before antimicrobial ther	apy started.	
Specimen transport	Specimens should working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norma Delays of over 48	Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Special requirements	If specimen is sm	If specimen is small place it in sterile water to prevent desiccation.		
Laboratory information				
Tests	Microscopy for detection of Gram positive and negative bacteria (semi- quantitative). General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).			
Measurement units	Growth detected	Growth detected or not detected.		
Biological reference units				
Turnaround time	4 days, plus 2 day	4 days, plus 2 days for enrichment culture.		
Availability	Routine hours an	Routine hours and on-call.		
Clinical information				
Factors known to significan affect the results	tly Specimens receiv Delays in transpo	ed in formal-saline are not s rtation may affect the recov	uitable for culture. ery of pathogens.	

## Toxoplasma diagnostic

Toxoplasma confirmation would only be performed on a Toxoplasma IgG positive sample. The test consists of Toxoplasma dye test and Toxoplasma IgM.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special rec	quirements.		
Specimen transport	Specimens sh working hours	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of no	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details are essential for processing. Please indicate if patient is pregnant and gestation with contact history.			
Laboratory information				
Tests	This test is pro laboratory on parameters a parameters w requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours	Routine hours.		
Clinical information				
Factors known to significar affect the results	Haemolysis.			

## Toxoplasma IgG (immunity)

In addition, if congenital infection suspected – amniotic fluid, fetal whole blood, neonatal cord blood can be tested – discuss with Consultant Microbiologist.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special require	No special requirements.		
Specimen transport	Specimens should be sent to the laboratory without delay during normal working hours.			
Storage requirements	Outside of normal working hours samples should be refrigerated.			
Special requirements	Clinical details and date of onset are essential for processing. Please indicate if patient is pregnant and gestation, with date of contact and exposure history.			
Laboratory information				
Tests	Detection of Toxoplasma gondii IgG (qualitative).			
Measurement units				
Biological reference units				
Turnaround time	7 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significat affect the results	htly Haemolysis.			

### Urines (microscopy and culture)

Send sample for microscopy and culture if clinically suspected UTI and any of the following:

- Pregnancy
- Signs of systemic or upper urinary tract infection (e.g.: fever, loin pain, renal angle tenderness)
- Immunocompromised or diabetic patients
- Male patients
- Children
- Female patients  $\geq$  65 years old
- Anatomically abnormal urinary/renal tract
- Failure to respond to empirical therapy
- History of recurrent UTIs (≥ 3 episodes/year)
- Patients with indwelling catheters ONLY if symptoms or signs of infection.

Collection container	Specimen	Sample volume	Request form
	Urine, MSU, Bladder urine, SPA	Minimum volume 5ml	
	Urine, MSU, Bladder urine, SPA	Minimum volume 1ml	
Sample instructions			
	Optimally collected Fill the container t 5ml is required. If less than 5ml of white topped univ	d before antimicrobial therap o the marked line (adults app urine is anticipated, or collect ersal container.	y started. rox 20-30 ml). A minimum of ing from a child, collect in to a
	MSU and clean catch urines are the most commonly collected space are recommended for routine use.		only collected specimens and
	Mid-stream specimen (MSU):		
Collection	Wash the genital a retract the foreskin water or sterile sal	rea in women with soap and n and wash skin surrounding t line	water or sterile saline. In men, he meatus with soap and
	Ask patient to pass	s a small amount of urine into	a bottle, bedpan or toilet.
	Using a clean conta Transfer the specir marked line, minin	3 a clean container collect a mid-stream specimen of urine afer the specimen into a sterile red-topped boric acid contain and line, minimum of 2ml) and send to the laboratory.	
	Catheter Specime	n of Urine (CSU	
	Do not use dipstic	ks for screening for infection,	this invariably gives a positive
result due to catheter colonisation. Request culture only when there are symptoms of infection.		of infection – document this	
norised by: C Frearson	•	, , ,	DCN: MIC-P-006-:
e of issue: 07/02/2023			Page 178 of

	clearly on the request form.
	Collect the specimen from the catheter self-sealing rubber sampling port using an asentic technique. The sample must not be obtained from the bag
	Disinfect the port using an alcohol or Chlorhexidine 2% swab, allow to the
	port to dry then use a sterile needle and syringe withdraw urine.
	Transfer the specimen into a sterile red-topped boric acid container (fill to
	marked line, minimum of 2ml) and send to the laboratory.
	Suprapubic aspirate (SPA)
	SPA is seen as the "gold standard" but is usually reserved for clarification of
	equivocal results from voided urine in infants and small children. Before SPA
	is attempted it is preferable to use ultrasound guidance to determine the
	presence of unite in the bladder.
	For <u>Mycobacteria</u> ; early morning urine on three consecutive days in 3 x 250ml container.
	For Schistosomiasis; Sample collected between 1000 and 1400. Alternatively
	a 24hr collection of terminal samples of urine may be obtained.
	Please note that urinary catheter tips will not be processed as they do not
	provide helpful microbiological information.
Specimen transport Specimens should be sent to the laboratory without delay during norr working hours.	
Storage requirements	Outside of normal working hours samples should be refrigerated.
	Delays of over 48 hours are undesirable.
Special requirements No special requirements.	
Laboratory information	
	Presence of white blood cells, red blood cells, epithelial cells and casts (semi-
Tests	quantitative).
	General isolation and characterisation of aerobic, microaerophilic and
Measurement units	Cell count x 10 <sup>6</sup> /l
Biological reference units	
	2 days
Turnaround time	3 days.
Availability	Routine hours and on-call (by arrangement).
Clinical information	
Frankright States	Bacteria multiply rapidly in urine – delays in transportation may affect the
Factors known to significantly	recovery of pathogens.
	misleading results.
	0

### **Back to index**

Authorised by: C FrearsonDCN: MIC-P-006-12.7Date of issue: 07/02/2023Page 179 of 192THIS DOCUMENTIS UNCONTROLLED MULEIN DRIVIEND

### Varicella zoster virus (VZV) IgG (immunity)

To determine past chickenpox infection (or vaccination); indicates immunity.

Chickenpox/zoster contact in susceptible persons (e.g. pregnant, immunocompromised, neonates): If an urgent VZV IgG is required after exposure, the laboratory must be notified, and information provided on nature of contact and date of exposure.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
Sample instructions				
Collection	No special requ	uirements.		
Specimen transport	Specimens sho working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of nor	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details Please indicate	Clinical details and date of onset are essential for processing. Please indicate if patient is pregnant and gestation with contact history.		
Laboratory information				
Tests	Detection of V2	ZV IgG (quantitative).		
Measurement units	IU/mL			
Biological reference units	<100 IU/mL - N 100-150 IU/mL >150 IU/mL - E	<100 IU/mL - No evidence of immunity 100-150 IU/mL – Evidence of immunity in the immunocompetent >150 IU/mL – Evidence of immunity in the immunocompromised		
Turnaround time	7 days.	7 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significar affect the results	Haemolysis. <b>htly</b> The performan have not been be interpreted	ce characteristics of the tes established. Results in imm with caution	t in newborns or in vaccinees unosuppressed subjects should	
# Varicella zoster virus (VZV) PCR

Diagnosis of acute disease.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
	Lesion swab (virus transport medium)			
Sample instructions				
Collection	Send a viral (gre membranes.	Send a viral (green top) swab of vesicle fluid or affected mucous membranes.		
Specimen transport	Specimens shou working hours.	Specimens should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of norm	Outside of normal working hours samples should be refrigerated.		
Special requirements	Clinical details a For VZV in CSF r	Clinical details are essential for processing. For VZV in CSF refer to <u>CSF (Cerebro-spinal fluid) virology PCR</u> .		
Laboratory information				
Tests	This test is proc laboratory on Te parameters ana parameters will requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units				
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significat affect the results	False negatives inappropriate ti of organism belo detection of an New and emerg this assay.	may occur for a variety of re ming of sample collection, in ow the detectable limit of th assay sampling variation will ing variants may also occur v	asons, for example happropriate sample, presence e assay. Towards the limit of result in lower reproducibility. which may not be detected by	

#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

# Viral haemorrhagic fever (VHF)

#### Used to determine past or current infection.

Collection container	Specimen	Sample volume	Request form	
	EDTA blood	2 – 6 mls		
Sample instructions				
Collection	If VHF other tha first discussing VHF Policy). Refer to curren	an Dengue fever suspected D with the Consultant Microbio t ACDP guidance.	O NOT TAKE SAMPLES without ologist (refer to the GWH Trust	
Specimen transport	Instructions fo defined in the C Specimens sho working hours. Do not use pne	r sample transportation o GWH Trust Specimen Transpo Id be sent to the laboratory umatic chute system if invest	f suspected VHF samples are ortation Procedure. without delay during normal igation for VHF required.	
Storage requirements	Outside of norr	nal working hours samples sl	nould be refrigerated.	
Special requirements	Samples from a the Microbiolog been performe Scientist has be Consultant Mic	Samples from a patient suspected of having VHF WILL NOT be processed by the Microbiology Department until a diagnosis VHF risk assessment has been performed by the Consultant Microbiologist, and the Biomedical Scientist has been authorised to proceed with processing the sample by the Consultant Microbiologist.		
Laboratory information				
Tests	This test is proc laboratory on T parameters and parameters wil requestor.	essed at an external referen elephone 01793 604798 if fu alysed in this test and any ref be displayed on the report v	ce centre. Contact the irther details are required. The erence ranges for these when it is returned to the	
Measurement units	·			
Biological reference units				
Turnaround time	14 days.			
Availability	Routine hours.			
Clinical information				
Factors known to significant affect the results	False negatives inappropriate t of organism bel detection of an New and emerg this assay.	may occur for a variety of re iming of sample collection, ir ow the detectable limit of th assay sampling variation wil ging variants may also occur	easons, for example nappropriate sample, presence ne assay. Towards the limit of I result in lower reproducibility. which may not be detected by	
ack to index				
orised by: C Frearson of issue: 07/02/2023			DCN: MIC-P-00 Page 182	

# Wounds (skin, superficial, non-surgical)

Swabs of acute wounds will be routinely cultured for primary pathogens i.e. *Staph aureus*,  $\beta$ -haemolytic streptococci. Where other potential pathogens are isolated in predominant or pure culture they will be reported. Growth of bacteria alone does not indicate the presence of infection, unless other factors such as inflammation, pus, erythema or fever are exhibited.

Chronic wounds are invariably colonised with bacteria. When processed, primary pathogens, potential pathogens in predominant or pure culture are reported as above as well as organisms likely to be simply colonising the wound (e.g. skin flora and faecal flora). This is because chronic wound management is influenced by degree of wound colonisation. Where heavy colonisation is identified this is invariably an indication for enhanced local wound care and not an immediate indication for systemic antibiotics.

Collection of pus or exudate	Minimum volume 1ml of	· · · · · · · · · · · · · · · · · · ·
	pus	
Amies transport swab	Swabs should be well soaked in pus	
Optimally collected Sample a represent unlikely to yield the If specimens are tak and the ulcer should aspiration of the ed irrigation-aspiration	before antimicrobial therapy ative part of the lesion. Swab causative pathogen. en from ulcers, the debris on d be cleaned with saline. A bi- ge of the wound should then method may be preferred.	started. bing dry crusted areas is the ulcer should be removed opsy or, preferably, a needle be taken. A less invasive
Specimens should be sent to the laboratory without delay during normal working hours.		
Outside of normal working hours samples should be refrigerated. Delays of over 48 hours are undesirable.		
Important to indicate site and nature of lesion.		
Microscopy for detection of gram positive and negative bacteria (semi- quantitative) (pus). General isolation and characterisation of aerobic, microaerophilic and anaerobic micro-organisms (qualitative).		
Growth detected or	not detected.	
4 days, plus 2 days f	or enrichment culture (pus).	
Routine hours and c	on-call (pus).	
		DCN: MIC-P-00 Page 183
	Amies transport swab Optimally collected Sample a representa unlikely to yield the If specimens are tak and the ulcer should aspiration of the ed, irrigation-aspiration Specimens should b working hours. Outside of normal w Delays of over 48 ho Important to indicat Microscopy for dete quantitative) (pus). General isolation an anaerobic micro-org Growth detected or 4 days, plus 2 days f Routine hours and c THIS DOCUMENT IS UNC	Amies transport swab       Swabs should be well soaked in pus         Optimally collected before antimicrobial therapy Sample a representative part of the lesion. Swab unlikely to yield the causative pathogen.         If specimens are taken from ulcers, the debris on and the ulcer should be cleaned with saline. A bid aspiration of the edge of the wound should then irrigation-aspiration method may be preferred.         Specimens should be sent to the laboratory without working hours.         Outside of normal working hours samples should Delays of over 48 hours are undesirable.         Important to indicate site and nature of lesion.         Microscopy for detection of gram positive and ne quantitative) (pus).         General isolation and characterisation of aerobic, anaerobic micro-organisms (qualitative).         Growth detected or not detected.         4 days, plus 2 days for enrichment culture (pus).         Routine hours and on-call (pus).         THIS DOCUMENT IS UNCONTROLLED WHEN PRINTE

#### **Clinical information**

Factors known to significantly	The recovery of anaerobes is compromised if transport time exceeds 3 hours.
affect the results	Delays in transportation may affect the recovery of pathogens.

#### **Back to index**

## Zika Virus

Zika virus testing is only available via PHE's Rare and Imported Pathogens Laboratory (RIPL). Please read PHE's Zika virus sample testing advice (link below) before collecting and sending a specimen to the laboratory.

Examinations offered				
Collection container	Specimen	Sample volume	Request form	
	Venous blood	2 – 6 mls		
	Urine (within 21 days of symptom onset)	1-5 mls		
Sample instructions				
Collection	Please read PHE's Z collecting and send	lika virus sample testing adv ling a specimen to the labora	ice (link below) before atory.	
Specimen transport	Specimens which d laboratory without	Specimens which do meet testing requirements should be sent to the laboratory without delay during normal working hours.		
Storage requirements	Outside of normal	Outside of normal working hours samples should be refrigerated.		
Special requirements	Comprehensive clir processing.	Comprehensive clinical details, including travel history, are essential for processing.		
Laboratory information				
Tests	This test is process laboratory on Telep parameters analyse parameters will be requestor.	This test is processed at an external reference centre. Contact the laboratory on Telephone 01793 604798 if further details are required. The parameters analysed in this test and any reference ranges for these parameters will be displayed on the report when it is returned to the requestor.		
Measurement units	N/A			
Biological reference units	N/A			
Turnaround time	14 days.	14 days.		
Availability	Routine hours.	Routine hours.		
Clinical information				
Factors known to significan affect the results	itly Haemolysis. Urine samples mus	t be taken within 21 days of	the onset of symptoms.	

Please refer to PHE's <u>Zika virus: sample testing advice</u> for further information.

#### **Back to index**

Authorised by: C Frearson Date of issue: 07/02/2023

**NHS Foundation Trust** 

#### **14 REFERENCE LABORATORIES**

As part of the testing process, it may be necessary to refer some, or all, of the sample to an external reference laboratory which has the necessary expertise. In some cases there will be only one specialist laboratory in the whole country which performs a particular test, meaning using referral laboratories is essential.

There is a detailed policy in place to govern how we choose these referral laboratories. They are selected for their expertise and their quality standards, and are regularly checked for their accreditation status.

The name of the reference laboratory used will be indicated on the Microbiology report. The reference laboratories currently used are:

Laboratory	Address	CPA/UKAS accreditation	Examinations offered
Anaerobe reference unit (ARU	Public Health Wales Microbiology	UKAS 9510	Anaerobe identification of
	Cardiff		Bacteroides, Clostridia,
	University Hospital of Wales		Fusobacteria, Actinomyces spp
	Heath Park		
	Cardiff		
	CF14 4XW		
Animal and Plant Health Agency	Virology Department	UKAS 1769	Diagnostic service for Rabies
	Woodham Lane	Accredited to ISO/IEC 17025:2005	
	New Haw		
	Addleston		
	Surrey		
	KT15 3NB		
Antimicrobial reference unit	North Bristol NHS Trust	UKAS 8099	Antimicrobial assay service
	Southmead Hospital		
	Southmead Road		
	Bristol		
	BS10 5NB		
Antimicrobial resistance and	Public Health England	UKAS 8197	National reference laboratory for
healthcare associated infections	61 Colindale Avenue		investigating antibiotic resistance in
reference unit (AMRHAI)	London		healthcare associated bacterial

Authorised by: C Frearson Date of issue: 07/02/2023 DCN: MIC-P-006-12.7

Page 186 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

# Great Western Hospitals NHS Foundation Trust

	NW9 5EQ		pathogens
<u>Clostridium difficile ribotyping</u> network (CDRN)	North Bristol NHS Trust Southmead Hospital Southmead Road Bristol BS10 5NB	UKAS 8099	Clostridium difficile culture and ribotyping
<u>Cryptosporidium reference unit</u> (CRU)	Public Health Wales Microbiology ABM, Singleton Hospital Sgeti Road Swansea SA2 8QA	UKAS 9510	Cryptosporidium typing and confirmation services
Epsom and St Helier University Hospitals NHS Trust	Microbiology Department West Park Hospital Morton Lane Epsom KT19 8PB	UKAS 8598	Enterovirus serology and PCR
Gastrointestinal bacteria reference unit (GBRU)	Public Health England 61 Colindale Avenue London NW9 5EQ	UKAS 8197	National reference laboratory for a range of gastrointestinal pathogens and providing specialist testing of clinical, food, water and environmental samples
<u>Genomic research unit</u>	Public Health England 61 Colindale Avenue London NW9 5EQ	UKAS 8727	Genome sequencing, transcription and proteogenome analysis, pathogen discovery and metagenomics
<u>Great Ormond Street Hospital for</u> Children NHS Foundation Trust	Bacteriology Laboratory Level 4 Camelia Botnar Laboratories Great Ormond Street London WC1N 3JH	UKAS 8675	Diagnostic service for Whipples disease
Insect Research and Development, Cambridge	6 Quy Court Colliers Lane Stow - cum- Quy	No accreditation status Laboratory work recognised in civil litigation and criminal prosecutions,	Identification of insect and animal foreign bodies

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 187 of 192
	THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED	

Department of Microbiology

Great Western Hospitals NHS

**NHS Foundation Trust** 

	Cambridge	or defence	
Liverpool Clinical Laboratories	Liverpool Clinical Laboratories Royal Liverpool and Broadgreen Univerisity Hospitals NHS Trust Prescot Street Liverpool L7 8XP	UKAS 9755	Brucella Serology
Lab 21	Park House Winship Road Milton Cambridge Cambridgeshire CB24 6BQ	UKAS 9325	Therapeutic drug monitoring for HIV patients
Meningococcal reference unit (MRU)	Clinical Sciences Building 2 Manchester Royal Infirmary Oxford Road Manchester M13 9WL	UKAS 10175	Meningococcal PCR and Serology Pneumococcal PCR
Mycology reference laboratory	Public Health England South West Laboratory Myrtle Road Bristol BS2 8EL	UKAS 8043	Laboratory services for the diagnosis and management of fungal infections
National CJD research and surveillance unit (NCJDRSU)	Western General Hospital Crewe Road Edinburgh EH4 2XU	Laboratory work recognised by WHO, inspected by HSE and perform well in European EQA schemes	Diagnostic service for CJD
National parasitology reference laboratory (NPRL)	Department of Clinical Parasitology Hospital for Tropical Diseases 3rd floor Mortimer Market Centre Mortimer Market London WC1E 6JB	UKAS 9702	Laboratory reference services for parasites and amoeba Various parasitology serology

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 188 of 192
	THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED	

Great Western Hospitals NHS

**NHS Foundation Trust** 

North Bristol NHS Trust	Immunology and Immunogenetics	UKAS 8067	Various viral PCR and serology
	Pathology Sciences Building		Faecal Calprotectin (Liquid stools
	Southmead Hospital		only)
	Westbury-on-Trym		CD4 counts
	Bristol		
	BS10 5NB		
Oxford University Hospitals NHS	Immunology Department	UKAS 8782	HIB serology
Trust	Churchill Hospital		Pneumococcal serology
	Old Road		
	Headington		
	Oxford		
	OX3 7⊔		
Royal Preston Hospital	Microbiology	UKAS 8545	Campylobacter serology
	Royal Preston Hospital		
	Sharoe Green Lane		
	Fulwood		
	Preston PR2 9HT		
Rare and imported pathogens	Public Health England	UKAS 9304	Diagnosis and management of
laboratory (RIPL)	Manor Farm Road		unusual or hazardous infectious
	Porton Down		diseases present in the UK or
	Salisbury		imported into the country.
	Wiltshire		Clinical diagnostic and reference
	SP4 0JG		leptospirosis service.
Respiratory and vaccine preventable	Public Health England	UKAS 8727	National and international reference
bacteria reference unit (RVPBRU)	61 Colindale		laboratory services for a number of
	London		bacteria causing respiratory,
	NW9 5EQ		systemic and vaccine preventable
			bacterial infections
Royal Devon and Exeter NHS	Microbiology Department	UKAS 9018	Diagnostic service for TB
Foundation Trust	Barrack Road		Quantiferon
	Exeter		
	EX2 5DW		
Sexually transmitted bacteria	Public Health England	UKAS 8727	National and international
reference unit (STBRU)	61 Colindale avenue		laboratory reference services for a
Authorised by: C Frearson		DCN: M	IIC-P-006-12.7

Date of issue: 07/02/2023

Page 189 of 192

THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

Department of Microbiology

# Great Western Hospitals NHS

**NHS Foundation Trust** 

	London		number of sexually transmitted
	NW9 5EQ		infections (STIs)
The Midlands public health	Heart of England NHS Foundation	UKAS 8213	HIV resistance service
laboratory services	Trust		
	Bordesley Green East		
	Birmingham		
	B9 5SS		
Toxoplasma reference laboratory	Department of Microbiology	UKAS 9510	Diagnostic service for toxoplasma
<u>(TRL)</u>	Singleton Hospital		infection
	Sgeti		
	Swansea		
	SA2 8QA		
University Hospital	Microbiology Department	UKAS 8403	Laboratory services for the diagnosis
Southampton NHS Foundation Trust	Tremona Road		and management of fungal
	Southampton		infections and mycobacterial
	Hampshire		infections
	SO16 6YD		
Virus reference department (VRD)	Public Health England	UKAS 8825	Clinical advice and laboratory
	61 Colindale Avenue		investigations for a wide range of
	London		human virus infections
	NW9 5EQ		

Authorised by: C FrearsonDCN: MIC-P-006-12.7Date of issue: 07/02/2023Page 190 of 192THIS DOCUMENT IS UNCONTROLLED WHEN PRINTED

### **15 PATIENT CONSENT DISCLOSURE**

#### 15.1 Laboratory Policy on protection of personal information

The Microbiology Department regards the lawful and correct treatment of patients' personal information as vital to successful operations and to maintaining the confidence of users of the service. Request form information may additionally be used for billing purposes, financial audit, resource management and utilization reviews.

Our policy is that we will treat personal information lawfully and correctly in adherence to the principles of data protection described in the <u>Data Protection Act 1998</u>.

As part of the Great Western Hospital NHS Foundation Trust we also work to its governance and data protection policies which incorporate the Data Protection Act, the <u>Department of Health Confidentiality</u> <u>NHS Code of Practice</u>, and <u>Department of Health Security Management NHS Code of Practise</u>, as listed below:

- Information Governance Strategy and Policy
- Information Protection and Security Policy
- Information Asset Register Procedure
- Data Protection Policy
- Data Transfer Policy
- Data Quality Policy
- Code of Conduct for Employees in Respect of Confidentiality Policy
- Freedom of Information Requests Procedure
- Consent to Treatment Policy

All the above Trust policy documentation is available upon request to the Laboratory Manager on 01793 604804.

#### **15.2** Patient consent

Consent to a specimen being taken and analysed is implied by the patient presenting to the point of specimen collection. The responsibility for obtaining informed consent for the tests(s) resides with the individual ordering the test. Informed consent should cover all the tests being done, implications of their results and disclosure of clinical and personal details to personnel (in the requesting organisation and any other healthcare organisations involved in providing the test). Special procedures, including more invasive procedures, or those with an increased risk of complications to the procedure, will need a more detailed explanation and, in some cases, written consent.

Patients in a hospital bed should normally be given the opportunity to refuse.

The laboratory procedures the laboratory assumes that patient consent has been obtained for the investigations requested, as the patient has presented themselves and willingly submitted to the usual

Authorised by: C Frearson		DCN: MIC-P-006-12.7
Date of issue: 07/02/2023		Page 191 of 192
	THIS DOCUMENT IS LINCONTROLLED WHEN PRINTED	

collecting procedure. The exception to this being antenatal requests, which always require requests for blood borne virus testing must be clearly indicated as accepted by the patient and signed by the requesting clinician.

In emergency situations consent may not be possible. Under these circumstances the laboratory will carry out the necessary investigations provided they are in the patient's best interest.

#### **15.3 Medico-legal samples**

Any specimens submitted for medico – legal purposes should have documentation accompanying these specimens to provide an unbroken chain of evidence.

#### **15.4** The Human Tissue Act

Great Western Hospitals NHS Foundation Trust are licensed by the Human Tissue Act (HTA) to undertake examinations of post mortem samples submitted by clinical consultants and pathologists. Under the license, the samples may be retained until the examination has been completed and in line with the sample retention policies.

It is the obligation of the requesting clinician or pathologist to ensure that examination of samples they submit have been requested by the coroner or appropriate consent has been obtained from the deceased person or their relatives.

Only the specific examinations requested by the sending clinician or pathologist may be performed. It must be assumed that the coroner has not asked for any other examinations to be performed and consent has not been obtained for any other work and so this would be outside the scope of the licence. If additional work on samples from the deceased is thought necessary by the medical microbiologist or virologist they must obtain written confirmation of consent from the sending departments.

All relevant material is stored securely and under conditions which maintain the integrity of the sample if possible and confidentiality is maintained in compliance with Caldicott principles, as are all samples received. Following processing, relevant material is only retained for the period of time specified by the retention policy.

#### **16 FEEDBACK ON OUR MICROBIOLOGY SERVICE AND COMPLAINTS PROCEDURE**

Any complaints should be directed to the Laboratory Manager or Clinical Lead. Also any suggestions from users on how this user guide could be improved would be welcome for inclusion in future editions. Please forward suggestions to the Laboratory Manager. Please also let us know about new services you would wish to see developed.